# Package 'mxnet'

May 5, 2020

```
Type Package
Title MXNet: A Flexible and Efficient Machine Learning Library for Heterogeneous Distributed Sys-
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Repository apache/incubator-mxnet
Description MXNet is a deep learning framework designed for both efficiency
      and flexibility. It allows you to mix the flavours of deep learning programs
      together to maximize the efficiency and your productivity.
License Apache License (== 2.0)
URL https://github.com/apache/incubator-mxnet/tree/master/R-package
BugReports https://github.com/apache/incubator-mxnet/issues
Imports methods,
      Rcpp (>= 0.12.1),
      DiagrammeR (>= 0.9.0),
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```

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# $\mathsf{R}$ topics documented:

rguments	
ıs.array.MXNDArray	
ıs.matrix.MXNDArray	
hildren	
tx	. 17
lim.MXNDArray	
graph.viz	. 17
m2rec	. 18
nternals	. 19
s.mx.context	. 20
s.mx.dataiter	. 20
s.mx.ndarray	. 20
s.mx.symbol	. 21
s.serialized	
ength.MXNDArray	. 22
nx.apply	. 22
nx.callback.early.stop	. 23
nx.callback.log.speedometer	. 23
nx.callback.log.train.metric	. 24
nx.callback.save.checkpoint	
nx.cpu	. 25
nx.ctx.default	
nx.exec.backward	
nx.exec.forward	. 26
nx.exec.update.arg.arrays	. 26
nx.exec.update.aux.arrays	
nx.exec.update.grad.arrays	
nx.gpu	
nx.infer.rnn	. 28
nx.infer.rnn.one	
nx.infer.rnn.one.unroll	
nx.init.create	
nx.init.internal.default	
nx.init.normal	
nx.init.uniform	
nx.init.Xavier	
nx.io.arrayiter	
nx.io.bucket.iter	
nx.io.CSVIter	. 32
nx.io.extract	
nx.io.ImageDetRecordIter	
nx io ImageRecordInt8Iter	38

mx.io.ImageRecordIter	41
	14
	<del>1</del> 7
	50
	53
mx.io.MNISTIter	55
mx.kv.create	56
mx.lr_scheduler.FactorScheduler	56
mx.lr_scheduler.MultiFactorScheduler	57
mx.metric.accuracy	58
mx.metric.custom	58
mx.metric.logistic_acc	58
mx.metric.logloss	59
mx.metric.mae	59
mx.metric.mse	59
mx.metric.Perplexity	50
mx.metric.rmse	50
mx.metric.rmsle	50
mx.metric.top_k_accuracy	51
	51
mx.model.buckets	52
mx.model.FeedForward.create	53
mx.model.init.params	55
	55
mx.model.save	56
mx.nd.abs	56
mx.nd.Activation	57
mx.nd.adam.update	57
mx.nd.add.n	58
mx.nd.all.finite	59
mx.nd.amp.cast	59
mx.nd.amp.multicast	70
mx.nd.arccos	70
mx.nd.arccosh	71
mx.nd.arcsin	71
mx.nd.arcsinh	72
mx.nd.arctan	72
	73
mx.nd.argmax	73
mx.nd.argmax.channel	74
mx.nd.argmin	75
mx.nd.argsort	75
mx.nd.array	76
mx.nd.batch.dot	77
mx.nd.batch.take	78
mx.nd.BatchNorm	78
mx.nd.BatchNorm.v1	30
my nd Bilinear Camplar	21

mx.nd.BlockGrad	82
mx.nd.broadcast.add	83
mx.nd.broadcast.axes	84
mx.nd.broadcast.axis	84
mx.nd.broadcast.div	85
mx.nd.broadcast.equal	86
mx.nd.broadcast.greater	86
mx.nd.broadcast.greater.equal	87
mx.nd.broadcast.hypot	87
mx.nd.broadcast.lesser	88
mx.nd.broadcast.lesser.equal	89
mx.nd.broadcast.like	89
mx.nd.broadcast.logical.and	90
mx.nd.broadcast.logical.or	91
mx.nd.broadcast.logical.xor	91
mx.nd.broadcast.maximum	92
mx.nd.broadcast.minimum	92
mx.nd.broadcast.minus	93
mx.nd.broadcast.mod	94
mx.nd.broadcast.mul	94
mx.nd.broadcast.not.equal	95
mx.nd.broadcast.plus	95
mx.nd.broadcast.power	
mx.nd.broadcast.sub	
mx.nd.broadcast.to	
mx.nd.Cast	
mx.nd.cast	
mx.nd.cast.storage	
mx.nd.cbrt	
mx.nd.ceil	
mx.nd.choose.element.0index	
mx.nd.clip	
mx.nd.col2im	
mx.nd.Concat	
mx.nd.concat	
mx.nd.Convolution	
mx.nd.Convolution.v1	
mx.nd.copyto	
mx.nd.Correlation	
mx.nd.cos	
mx.nd.cosh	
-	
mx.nd.crop	
mx.nd.ctc.loss	
mx.nd.CTCLoss	
mx.nd.cumsum	
mx.nd.Custom	
mx.nd.Deconvolution	117

mx.nd.ones	. 187
mx.nd.ones.like	
mx.nd.Pad	
mx.nd.pad	
mx.nd.pick	
mx.nd.Pooling	
mx.nd.Pooling.v1	
mx.nd.preloaded.multi.mp.sgd.mom.update	
mx.nd.preloaded.multi.mp.sgd.update	
mx.nd.preloaded.multi.sgd.mom.update	
$mx.nd.preloaded.multi.sgd.update \\ \ldots \\ $	
mx.nd.prod	
mx.nd.radians	
mx.nd.random.exponential	. 198
mx.nd.random.gamma	. 199
mx.nd.random.generalized.negative.binomial	. 199
mx.nd.random.negative.binomial	
mx.nd.random.normal	
mx.nd.random.pdf.dirichlet	
mx.nd.random.pdf.exponential	
mx.nd.random.pdf.gamma	
mx.nd.random.pdf.generalized.negative.binomial	
mx.nd.random.pdf.negative.binomial	
mx.nd.random.pdf.normal	
mx.nd.random.pdf.poisson	
mx.nd.random.pdf.uniform	
mx.nd.random.poisson	
mx.nd.random.randint	
mx.nd.random.uniform	
mx.nd.ravel.multi.index	
mx.nd.rcbrt	. 211
mx.nd.reciprocal	. 211
mx.nd.relu	
mx.nd.repeat	. 212
mx.nd.reset.arrays	
mx.nd.Reshape	
mx.nd.reshape	
mx.nd.reshape.like	
mx.nd.reverse	
mx.nd.rint	
mx.nd.rmsprop.update	
mx.nd.rmspropalex.update	
mx.nd.RNN	
mx.nd.ROIPooling	
mx.nd.round	
mx.nd.rsqrt	
mx.nd.sample.exponential	. 225
mx.nd.sample.gamma	. 226

8

mx.nd.sample.generalized.negative.binomial	227
mx.nd.sample.multinomial	
mx.nd.sample.negative.binomial	
mx.nd.sample.normal	
mx.nd.sample.poisson	
mx.nd.sample.uniform	
mx.nd.save	
mx.nd.scatter.nd	
mx.nd.SequenceLast	
mx.nd.SequenceMask	
mx.nd.SequenceReverse	
mx.nd.sgd.mom.update	
mx.nd.sgd.update	
mx.nd.shape.array	
mx.nd.shuffle	
mx.nd.sigmoid	
mx.nd.sign	
mx.nd.signsgd.update	
mx.nd.signum.update	
mx.nd.sin	
mx.nd.sinh	
mx.nd.size.array	
mx.nd.slice.axis	
mx.nd.slice.like	
mx.nd.SliceChannel	
mx.nd.smooth.l1	
mx.nd.Softmax	
mx.nd.softmax	
mx.nd.softmax.cross.entropy	
mx.nd.SoftmaxActivation	
mx.nd.SoftmaxOutput	
mx.nd.softmin	
mx.nd.softsign	
mx.nd.sort	
mx.nd.space.to.depth	
mx.nd.SpatialTransformer	
mx.nd.split	
mx.nd.sqrt	
mx.nd.square	
mx.nd.squeeze	
mx.nd.stack	
mx.nd.stop.gradient	
mx.nd.sum	
mx.nd.sum.axis	
mx.nd.SVMOutput	
mx.nd.swapaxes	
mx.nd.SwapAxis	
mx.nd.take	. 265
	. 400

mx.nd.tan	. 266
mx.nd.tanh	. 267
mx.nd.tile	. 268
mx.nd.topk	. 268
mx.nd.transpose	
mx.nd.trunc	
mx.nd.uniform	. 271
mx.nd.unravel.index	. 271
mx.nd.UpSampling	. 272
mx.nd.where	. 273
mx.nd.zeros	. 274
mx.nd.zeros.like	. 275
mx.opt.adadelta	. 275
mx.opt.adagrad	. 276
mx.opt.adam	
mx.opt.create	. 277
mx.opt.get.updater	
mx.opt.nag	
mx.opt.rmsprop	
mx.opt.sgd	
mx.profiler.config	
mx.profiler.state	
mx.rnorm	
mx.runif	
mx.serialize	
mx.set.seed	
mx.simple.bind	
mx.symbol.abs	
mx.symbol.Activation	
mx.symbol.adam_update	
mx.symbol.add_n	
mx.symbol.all_finite	
mx.symbol.amp_cast	
mx.symbol.amp_multicast	
mx.symbol.arccos	
mx.symbol.arccosh	
mx.symbol.arcsin	
mx.symbol.arcsinh	
mx.symbol.arctan	
mx.symbol.arctanh	
mx.symbol.argmax	
mx.symbol.argmax_channel	
mx.symbol.argmin	
mx.symbol.argsort	
mx.symbol.BatchNorm	
mx.symbol.BatchNorm v1	
mx.symbol.batch_dot	
mx.symbol.batch_take	

mx.symbol.BilinearSampler	. 301
mx.symbol.BlockGrad	. 302
mx.symbol.broadcast_add	. 303
mx.symbol.broadcast_axes	. 304
mx.symbol.broadcast_axis	. 305
mx.symbol.broadcast_div	
mx.symbol.broadcast_equal	
mx.symbol.broadcast_greater	
mx.symbol.broadcast_greater_equal	
mx.symbol.broadcast_hypot	
mx.symbol.broadcast_lesser	
mx.symbol.broadcast_lesser_equal	
mx.symbol.broadcast_like	
mx.symbol.broadcast_logical_and	
mx.symbol.broadcast_logical_or	
mx.symbol.broadcast_logical_xor	
mx.symbol.broadcast_maximum	
mx.symbol.broadcast_minimum	
mx.symbol.broadcast_minus	
mx.symbol.broadcast_mod	
mx.symbol.broadcast_mul	
mx.symbol.broadcast_not_equal	
mx.symbol.broadcast_plus	
mx.symbol.broadcast_power	
mx.symbol.broadcast_sub	. 320
mx.symbol.broadcast_to	. 321
mx.symbol.Cast	. 322
mx.symbol.cast	. 322
mx.symbol.cast_storage	. 323
mx.symbol.cbrt	. 324
mx.symbol.ceil	. 324
mx.symbol.choose_element_0index	
mx.symbol.clip	
mx.symbol.col2im	
mx.symbol.Concat	
mx.symbol.concat	
mx.symbol.Convolution	
mx.symbol.Convolution v1	
mx.symbol.Correlation	
mx.symbol.cos	
mx.symbol.cosh	
mx.symbol.Crop	
•	
mx.symbol.crop	
mx.symbol.CTCLoss	
mx.symbol.ctc_loss	
mx.symbol.cumsum	
mx.symbol.Custom	
mx symbol Deconvolution	341

mx.symbol.degrees	
mx.symbol.depth_to_space	
mx.symbol.diag	. 344
mx.symbol.digamma	. 345
mx.symbol.dot	346
mx.symbol.Dropout	347
mx.symbol.ElementWiseSum	. 348
mx.symbol.elemwise_add	. 349
mx.symbol.elemwise_div	349
mx.symbol.elemwise_mul	. 350
mx.symbol.elemwise_sub	350
mx.symbol.Embedding	. 351
mx.symbol.erf	. 352
mx.symbol.erfinv	. 353
mx.symbol.exp	. 354
mx.symbol.expand_dims	. 354
mx.symbol.expm1	. 355
mx.symbol.fill_element_0index	
mx.symbol.fix	
mx.symbol.Flatten	
mx.symbol.flatten	. 358
mx.symbol.flip	358
mx.symbol.floor	
mx.symbol.ftml_update	
mx.symbol.ftrl_update	
mx.symbol.FullyConnected	
mx.symbol.gamma	
mx.symbol.gammaln	
mx.symbol.gather_nd	
mx.symbol.GridGenerator	
mx.symbol.Group	
mx.symbol.GroupNorm	
mx.symbol.hard_sigmoid	
mx.symbol.identity	
mx.symbol.IdentityAttachKLSparseReg	
mx.symbol.im2col	
mx.symbol.infer.shape	
mx.symbol.InstanceNorm	
mx.symbol.khatri_rao	
mx.symbol.L2Normalization	
mx.symbol.lamb_update_phase1	
mx.symbol.lamb_update_phase2	
mx.symbol.LayerNorm	
mx.symbol.LeakyReLU	
mx.symbol.linalg_det	
mx.symbol.linalg_extractdiag	
mx.symbol.linalg_extracttrian	
mx.symbol.linalg_gelqf	

12

mx.symbol.linalg_gemm	. 381
mx.symbol.linalg_gemm2	. 382
mx.symbol.linalg_inverse	
mx.symbol.linalg_makediag	
mx.symbol.linalg_maketrian	
mx.symbol.linalg_potrf	
mx.symbol.linalg_potri	
mx.symbol.linalg_slogdet	
mx.symbol.linalg_sumlogdiag	
mx.symbol.linalg_syrk	
mx.symbol.linalg_trmm	
mx.symbol.linalg_trsm	
mx.symbol.LinearRegressionOutput	
mx.symbol.load	
mx.symbol.load.json	
mx.symbol.log	
·	
mx.symbol.log10	
mx.symbol.log1p	
mx.symbol.log2	
mx.symbol.logical_not	
mx.symbol.LogisticRegressionOutput	
mx.symbol.log_softmax	
mx.symbol.LRN	
mx.symbol.MAERegressionOutput	
mx.symbol.MakeLoss	
mx.symbol.make_loss	
mx.symbol.max	
mx.symbol.max_axis	
mx.symbol.mean	
mx.symbol.moments	. 405
mx.symbol.mp_lamb_update_phase1	
mx.symbol.mp_lamb_update_phase2	. 407
mx.symbol.mp_nag_mom_update	. 408
mx.symbol.mp_sgd_mom_update	. 409
mx.symbol.mp_sgd_update	
mx.symbol.multi_all_finite	. 410
mx.symbol.multi lars	
mx.symbol.multi_mp_sgd_mom_update	. 412
mx.symbol.multi_mp_sgd_update	
mx.symbol.multi_sgd_mom_update	
mx.symbol.multi_sgd_update	
mx.symbol.multi_sum_sq	
mx.symbol.nag_mom_update	
mx.symbol.nanprod	
mx.symbol.nansum	
mx.symbol.negative	
mx.symbol.norm	
mx.symbol.normal	
majamoumonma	. <del>+</del> ∠U

mx.symbol.ones_like	
$mx.symbol.one\_hot \ . \ . \ . \ . \ . \ . \ . \ . \ . \ $	. 422
mx.symbol.Pad	. 423
$mx.symbol.pad \dots \dots$	. 424
mx.symbol.pick	. 425
mx.symbol.Pooling	. 427
mx.symbol.Pooling_v1	. 428
mx.symbol.preloaded_multi_mp_sgd_mom_update	. 430
mx.symbol.preloaded_multi_mp_sgd_update	
mx.symbol.preloaded_multi_sgd_mom_update	. 431
mx.symbol.preloaded_multi_sgd_update	
mx.symbol.prod	
mx.symbol.radians	
mx.symbol.random_exponential	
mx.symbol.random_gamma	
mx.symbol.random_generalized_negative_binomial	
mx.symbol.random_negative_binomial	
mx.symbol.random_normal	
mx.symbol.random_pdf_dirichlet	
mx.symbol.random_pdf_exponential	
mx.symbol.random_pdf_gamma	
mx.symbol.random_pdf_generalized_negative_binomial	
mx.symbol.random_pdf_negative_binomial	
mx.symbol.random_pdf_normal	
mx.symbol.random_pdf_poisson	
mx.symbol.random_pdf_uniform	
mx.symbol.random_poisson	
mx.symbol.random_randint	
mx.symbol.random_uniform	
mx.symbol.ravel_multi_index	
mx.symbol.rcbrt	
mx.symbol.reciprocal	
mx.symbol.relu	
mx.symbol.repeat	
mx.symbol.reset_arrays	
mx.symbol.Reshape	
mx.symbol.reshape	
mx.symbol.reshape_like	
mx.symbol.reverse	
mx.symbol.rint	
$mx.symbol.rmspropalex\_update  .  .  .  .  .  .  .  .  .  $	
$mx.symbol.rmsprop\_update \qquad . \qquad $	
mx.symbol.RNN	
mx.symbol.ROIPooling	
$mx.symbol.round \ \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	
$mx.symbol.rsqrt \dots \dots$	
mx.symbol.sample_exponential	
mx.symbol.sample gamma	

mx.symbol.sample_generalized_negative_binomial	
mx.symbol.sample_multinomial	469
mx.symbol.sample_negative_binomial	470
mx.symbol.sample_normal	471
mx.symbol.sample_poisson	472
mx.symbol.sample_uniform	
mx.symbol.save	
mx.symbol.scatter_nd	
mx.symbol.SequenceLast	475
mx.symbol.SequenceMask	
mx.symbol.SequenceReverse	
mx.symbol.sgd_mom_update	
mx.symbol.sgd_update	
mx.symbol.shape_array	
mx.symbol.shuffle	
mx.symbol.sigmoid	
mx.symbol.sign	
mx.symbol.signsgd_update	
mx.symbol.signum_update	
mx.symbol.sin	
mx.symbol.sinh	
mx.symbol.size_array	
mx.symbol.slice	
mx.symbol.SliceChannel	
mx.symbol.slice_axis	
mx.symbol.slice_like	
mx.symbol.smooth_11	
mx.symbol.Softmax	
mx.symbol.softmax	
mx.symbol.SoftmaxActivation	
mx.symbol.SoftmaxOutput	
mx.symbol.softmax_cross_entropy	
mx.symbol.softmin	
mx.symbol.softsign	
mx.symbol.sort	
mx.symbol.space_to_depth	
mx.symbol.SpatialTransformer	
mx.symbol.split	
mx.symbol.sqrt	
mx.symbol.square	
mx.symbol.squeeze	
mx.symbol.stack	
mx.symbol.stop gradient	
mx.symbol.sum	
mx.symbol.sum_axis	
mx.symbol.SVMOutput	
mx.symbol.swapaxes	
mx.symbol.SwapAxis	

arguments	15

	mx.symbol.take			
	<i>U</i> 1			
Index				528

arguments

Get the arguments of symbol.

# Description

Get the arguments of symbol.

# Usage

arguments(x)

# Arguments

Х

The input symbol

16 children

as.array.MXNDArray

as.array operator overload of mx.ndarray

# Description

as.array operator overload of mx.ndarray

# Usage

```
## S3 method for class 'MXNDArray'
as.array(nd)
```

#### **Arguments**

nd

The mx.ndarray

as.matrix.MXNDArray

as.matrix operator overload of mx.ndarray

# Description

as.matrix operator overload of mx.ndarray

# Usage

```
## S3 method for class 'MXNDArray'
as.matrix(nd)
```

#### **Arguments**

nd

The mx.ndarray

children

Gets a new grouped symbol whose output contains inputs to output nodes of the original symbol.

#### Description

Gets a new grouped symbol whose output contains inputs to output nodes of the original symbol.

#### Usage

```
children(x)
```

#### **Arguments**

Х

The input symbol

ctx 17

ctx

Get the context of mx.ndarray

# Description

Get the context of mx.ndarray

# Usage

ctx(nd)

# Arguments

nd

The mx.ndarray

dim.MXNDArray

Dimension operator overload of mx.ndarray

# Description

Dimension operator overload of mx.ndarray

# Usage

```
## S3 method for class 'MXNDArray'
dim(nd)
```

# Arguments

nd

The mx.ndarray

graph.viz

Convert symbol to Graphviz or visNetwork visualisation.

# Description

Convert symbol to Graphviz or visNetwork visualisation.

im2rec

#### Usage

```
graph.viz(
  symbol,
  shape = NULL,
  direction = "TD",
  type = "graph",
  graph.width.px = NULL,
  graph.height.px = NULL)
```

#### **Arguments**

```
symbol a string representing the symbol of a model.

shape a numeric representing the input dimensions to the symbol.

direction a string representing the direction of the graph, either TD or LR.

type a string representing the rendering engine of the graph, either graph or vis.

graph.width.px a numeric representing the size (width) of the graph. In pixels

graph.height.px

a numeric representing the size (height) of the graph. In pixels
```

#### Value

a graph object ready to be displayed with the print function.

im2rec

Convert images into image recordio format

#### Description

Convert images into image recordio format

#### Usage

```
im2rec(
  image_lst,
  root,
  output_rec,
  label_width = 1L,
  pack_label = 0L,
  new_size = -1L,
  nsplit = 1L,
  partid = 0L,
  center_crop = 0L,
  quality = 95L,
  color_mode = 1L,
```

internals 19

```
unchanged = 0L,
inter_method = 1L,
encoding = ".jpg"
)
```

# Arguments

image_lst	The image 1st file
root	The root folder for image files
output_rec	The output rec file
label_width	The label width in the list file. Default is 1.
pack_label	Whether to also pack multi dimenional label in the record file. Default is 0.
new_size	The shorter edge of image will be resized to the newsize. Original images will be packed by default.
nsplit	It is used for part generation, logically split the image.lst to NSPLIT parts by position. Default is 1.
partid	It is used for part generation, pack the images from the specific part in image.lst. Default is 0.
center_crop	Whether to crop the center image to make it square. Default is 0.
quality	JPEG quality for encoding (1-100, default: 95) or PNG compression for encoding (1-9, default: 3).
color_mode	Force color (1), gray image (0) or keep source unchanged (-1). Default is 1.
unchanged	Keep the original image encoding, size and color. If set to 1, it will ignore the others parameters.
inter_method	NN(0), BILINEAR(1), CUBIC(2), AREA(3), LANCZOS4(4), AUTO(9), RAND(10). Default is 1.
encoding	The encoding type for images. It can be '.jpg' or '.png'. Default is '.jpg'.

internals	Get a symbol that contains all the internals

# Description

Get a symbol that contains all the internals

# Usage

```
internals(x)
```

# Arguments

x The input symbol

20 is.mx.ndarray

is.mx.context

Check if the type is mxnet context.

# Description

Check if the type is mxnet context.

# Usage

```
is.mx.context(x)
```

#### Value

Logical indicator

is.mx.dataiter

Judge if an object is mx.dataiter

# Description

Judge if an object is mx.dataiter

# Usage

```
is.mx.dataiter(x)
```

#### Value

Logical indicator

is.mx.ndarray

Check if src.array is mx.ndarray

# Description

Check if src.array is mx.ndarray

# Usage

```
is.mx.ndarray(src.array)
```

# Value

Logical indicator

is.mx.symbol 21

# **Examples**

```
mat = mx.nd.array(1:10)
is.mx.ndarray(mat)
mat2 = 1:10
is.mx.ndarray(mat2)
```

is.mx.symbol

Judge if an object is mx.symbol

# Description

Judge if an object is mx.symbol

# Usage

```
is.mx.symbol(x)
```

#### Value

Logical indicator

is.serialized

Check if the model has been serialized into RData-compatiable format.

# Description

Check if the model has been serialized into RData-compatiable format.

# Usage

```
is.serialized(model)
```

#### Value

Logical indicator

22 mx.apply

length.MXNDArray

Length operator overload of mx.ndarray

# Description

Length operator overload of mx.ndarray

# Usage

```
## S3 method for class 'MXNDArray'
length(nd)
```

# Arguments

nd

The mx.ndarray

mx.apply

Apply symbol to the inputs.

# Description

Apply symbol to the inputs.

# Usage

```
mx.apply(x, ...)
```

# Arguments

Χ

The symbol to be applied

kwargs

The keyword arguments to the symbol

mx.callback.early.stop 23

```
mx.callback.early.stop
```

Early stop with different conditions

#### **Description**

Early stopping applying different conditions: hard thresholds or epochs number from the best score. Tested with "epoch.end.callback" function.

# Usage

```
mx.callback.early.stop(
   train.metric = NULL,
   eval.metric = NULL,
   bad.steps = NULL,
   maximize = FALSE,
   verbose = FALSE
)
```

#### **Arguments**

train.metric	Numeric. Hard threshold for the metric of the training data set (optional)		
eval.metric	Numeric. Hard threshold for the metric of the evaluating data set (if set, optional)		
bad.steps	Integer. How much epochs should gone from the best score? Use this option with evaluation data set		
maximize	Logical. Do your model use maximizing or minimizing optimization?		
verbose	Logical		

```
mx.callback.log.speedometer
```

Calculate the training speed

# Description

Calculate the training speed

#### Usage

```
mx.callback.log.speedometer(batch.size, frequency = 50)
```

# Arguments

```
frequency The frequency of the training speed update
```

batch\_size The batch size

```
\verb|mx.callback.log.train.metric| \\
```

Log training metric each period

# Description

Log training metric each period

### Usage

```
mx.callback.log.train.metric(period, logger = NULL)
```

# Arguments

period The number of batch to log the training evaluation metric

logger The logger class

mx.callback.save.checkpoint

Save checkpoint to files each period iteration.

# Description

Save checkpoint to files each period iteration.

# Usage

```
mx.callback.save.checkpoint(prefix, period = 1)
```

# Arguments

prefix The prefix of the model checkpoint.

mx.cpu 25

mx.cpu

Create a mxnet CPU context.

#### **Description**

Create a mxnet CPU context.

#### **Arguments**

dev.id

optional, default=0 The device ID, this is meaningless for CPU, included for interface compatibility.

#### Value

The CPU context.

mx.ctx.default

Set/Get default context for array creation.

# Description

Set/Get default context for array creation.

#### Usage

```
mx.ctx.default(new = NULL)
```

#### **Arguments**

new

optional takes mx.cpu() or mx.gpu(id), new default ctx.

#### Value

The default context.

mx.exec.backward

Peform an backward on the executors This function will MUTATE the state of exec

# Description

Peform an backward on the executors This function will MUTATE the state of exec

#### Usage

```
mx.exec.backward(exec, ...)
```

mx.exec.forward

Peform an forward on the executors This function will MUTATE the state of exec

# Description

Peform an forward on the executors This function will MUTATE the state of exec

#### Usage

```
mx.exec.forward(exec, is.train = TRUE)
```

```
mx.exec.update.arg.arrays
```

Update the executors with new arrays This function will MUTATE the state of exec

# Description

Update the executors with new arrays This function will MUTATE the state of exec

#### Usage

```
mx.exec.update.arg.arrays(
  exec,
  arg.arrays,
  match.name = FALSE,
  skip.null = FALSE
)
```

```
mx.exec.update.aux.arrays
```

Update the executors with new arrays This function will MUTATE the state of exec

#### **Description**

Update the executors with new arrays This function will MUTATE the state of exec

#### Usage

```
mx.exec.update.aux.arrays(
  exec,
  arg.arrays,
  match.name = FALSE,
  skip.null = FALSE
)
```

mx.exec.update.grad.arrays

Update the executors with new arrays This function will MUTATE the state of exec

### **Description**

Update the executors with new arrays This function will MUTATE the state of exec

# Usage

```
mx.exec.update.grad.arrays(
  exec,
  arg.arrays,
  match.name = FALSE,
  skip.null = FALSE
)
```

mx.gpu

Create a mxnet GPU context.

# Description

Create a mxnet GPU context.

# Arguments

dev.id

optional, default=0 The GPU device ID, starts from 0.

#### Value

The GPU context.

28 mx.infer.rnn.one

mx.infer.rnn

Inference of RNN model

# Description

Inference of RNN model

# Usage

```
mx.infer.rnn(infer.data, model, ctx = mx.cpu())
```

# Arguments

infer.data DataIter

model Model used for inference

ctx

mx.infer.rnn.one

Inference for one-to-one fusedRNN (CUDA) models

# Description

Inference for one-to-one fusedRNN (CUDA) models

# Usage

```
mx.infer.rnn.one(
  infer.data,
  symbol,
  arg.params,
  aux.params,
  input.params = NULL,
  ctx = mx.cpu()
)
```

# Arguments

infer.data Data iterator created by mx.io.bucket.iter

symbol Symbol used for inference

ctx

mx.infer.rnn.one.unroll 29

```
mx.infer.rnn.one.unroll
```

Inference for one-to-one unroll models

# Description

Inference for one-to-one unroll models

# Usage

```
mx.infer.rnn.one.unroll(
   infer.data,
   symbol,
   num_hidden,
   arg.params,
   aux.params,
   init_states = NULL,
   ctx = mx.cpu()
)
```

#### **Arguments**

mx.init.create

Create initialization of argument like arg.array

#### **Description**

Create initialization of argument like arg.array

#### Usage

```
mx.init.create(initializer, shape.array, ctx = NULL, skip.unknown = TRUE)
```

#### **Arguments**

initializer The initializer.

shape.array A named list that represents the shape of the weights

ctx mx.context The context of the weights skip.unknown Whether skip the unknown weight types

30 mx.init.uniform

```
mx.init.internal.default
```

Internal default value initialization scheme.

#### **Description**

Internal default value initialization scheme.

# Usage

```
mx.init.internal.default(name, shape, ctx, allow.unknown = FALSE)
```

#### **Arguments**

name the name of the variable.

shape the shape of the array to be generated.

mx.init.normal

Create a initializer that initialize the weight with normal(0, sd)

#### **Description**

Create a initializer that initialize the weight with normal(0, sd)

#### Usage

```
mx.init.normal(sd)
```

#### **Arguments**

sd

The standard deviation of normal distribution

 $\verb|mx.init.uniform| \\$ 

Create a initializer that initialize the weight with uniform [-scale, scale]

#### **Description**

Create a initializer that initialize the weight with uniform [-scale, scale]

#### Usage

```
mx.init.uniform(scale)
```

#### Arguments

scale

The scale of uniform distribution

mx.init.Xavier 31

# Description

Create a initializer which initialize weight with Xavier or similar initialization scheme.

#### Usage

```
mx.init.Xavier(rnd_type = "uniform", factor_type = "avg", magnitude = 3)
```

#### Arguments

rnd\_type A string of character indicating the type of distribution from which the weights

are initialized.

factor\_type A string of character.

magnitude A numeric number indicating the scale of random number range.

mx.io.arrayiter Create MXDataIter compatible iterator from R's array

#### **Description**

Create MXDataIter compatible iterator from R's array

#### Usage

```
mx.io.arrayiter(data, label, batch.size = 128, shuffle = FALSE)
```

#### **Arguments**

data The data array.

label The label array.

batch.size The batch size used to pack the array.

shuffle Whether shuffle the data

32 mx.io.CSVIter

mx.io.bucket.iter

Create Bucket Iter

# Description

Create Bucket Iter

#### Usage

```
mx.io.bucket.iter(
  buckets,
  batch.size,
  data.mask.element = 0,
  shuffle = FALSE,
  seed = 123
)
```

# Arguments

buckets The data array.

batch.size The batch size used to pack the array.

data.mask.element

The element to mask

shuffle Whether shuffle the data

seed The random seed

mx.io.CSVIter

Returns the CSV file iterator.

#### **Description**

In this function, the 'data\_shape' parameter is used to set the shape of each line of the input data. If a row in an input file is '1,2,3,4,5,6" and 'data\_shape' is (3,2), that row will be reshaped, yielding the array [[1,2],[3,4],[5,6]] of shape (3,2).

#### Usage

```
mx.io.CSVIter(...)
```

mx.io.CSVIter 33

#### **Arguments**

data.csv string, required The input CSV file or a directory path. data.shape Shape(tuple), required The shape of one example. string, optional, default='NULL' The input CSV file or a directory path. If label.csv NULL, all labels will be returned as 0. label.shape Shape(tuple), optional, default=[1] The shape of one label. batch.size int (non-negative), required Batch size. round.batch boolean, optional, default=1 Whether to use round robin to handle overflow batch or not. prefetch.buffer long (non-negative), optional, default=4 Maximum number of batches to prefetch. 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for. ctx None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional, dtype default='None' Output data type. "None" means no change.

#### **Details**

By default, the 'CSVIter' has 'round\_batch' parameter set to "True". So, if 'batch\_size' is 3 and there are 4 total rows in CSV file, 2 more examples are consumed at the first round. If 'reset' function is called after first round, the call is ignored and remaining examples are returned in the second round.

If one wants all the instances in the second round after calling 'reset', make sure to set 'round\_batch' to False.

If "data\_csv = 'data/'" is set, then all the files in this directory will be read.

"reset()" is expected to be called only after a complete pass of data.

By default, the CSVIter parses all entries in the data file as float32 data type, if 'dtype' argument is set to be 'int32' or 'int64' then CSVIter will parse all entries in the file as int32 or int64 data type accordingly.

#### Examples::

3.]]

```
// Contents of CSV file "data/data.csv". 1,2,3 2,3,4 3,4,5 4,5,6

// Creates a 'CSVIter' with 'batch_size'=2 and default 'round_batch'=True. CSVIter = mx.io.CSVIter(data_csv = 'data/data.csv', data_shape = (3,), batch_size = 2)

// Two batches read from the above iterator are as follows: [[ 1. 2. 3.] [ 2. 3. 4.]] [[ 3. 4. 5.] [ 4. 5. 6.]]

// Creates a 'CSVIter' with default 'round_batch' set to True. CSVIter = mx.io.CSVIter(data_csv = 'data/data.csv', data_shape = (3,), batch_size = 3)

// Two batches read from the above iterator in the first pass are as follows: [[ 1. 2. 3.] [ 2. 3. 4.] [ 3. 4. 5.]]

[[ 4. 5. 6.] [ 1. 2. 3.] [ 2. 3. 4.]]

// Now, 'reset' method is called. CSVIter.reset()

// Batch read from the above iterator in the second pass is as follows: [[ 3. 4. 5.] [ 4. 5. 6.] [ 1. 2.
```

// Creates a 'CSVIter' with 'round\_batch'=False. CSVIter = mx.io.CSVIter(data\_csv = 'data/data.csv', data\_shape = (3,), batch\_size = 3, round\_batch=False)

// Contents of two batches read from the above iterator in both passes, after calling // 'reset' method before second pass, is as follows: [[1. 2. 3.] [2. 3. 4.] [3. 4. 5.]]

```
[[4. 5. 6.] [2. 3. 4.] [3. 4. 5.]]
```

// Creates a 'CSVIter' with 'dtype'='int32' CSVIter = mx.io.CSVIter(data\_csv = 'data/data.csv', data\_shape = (3,), batch\_size = 3, round\_batch=False, dtype='int32')

// Contents of two batches read from the above iterator in both passes, after calling // 'reset' method before second pass, is as follows: [[1 2 3] [2 3 4] [3 4 5]]

```
[[4 5 6] [2 3 4] [3 4 5]]
```

Defined in src/io/iter\_csv.cc:L308

#### Value

iter The result mx.dataiter

mx.io.extract

Extract a certain field from DataIter.

#### Description

Extract a certain field from DataIter.

#### Usage

```
mx.io.extract(iter, field)
```

mx.io.ImageDetRecordIter

Create iterator for image detection dataset packed in recordio.

#### **Description**

Create iterator for image detection dataset packed in recordio.

#### Usage

```
mx.io.ImageDetRecordIter(...)
```

#### **Arguments**

path.imglist string, optional, default="Dataset Param: Path to image list.

path.imgrec string, optional, default='./data/imgrec.rec' Dataset Param: Path to image record

file.

aug.seq string, optional, default='det\_aug\_default' Augmentation Param: the augmenter

names to represent sequence of augmenters to be applied, seperated by comma. Additional keyword parameters will be seen by these augmenters. Make sure

you don't use normal augmenters for detection tasks.

label.width int, optional, default='-1' Dataset Param: How many labels for an image, -1 for

variable label size.

preprocess.threads

int, optional, default='4' Backend Param: Number of thread to do preprocess-

ing.

verbose boolean, optional, default=1 Auxiliary Param: Whether to output parser infor-

mation.

num.parts int, optional, default='1' partition the data into multiple parts

part.index int, optional, default='0' the index of the part will read

shuffle.chunk.size

long (non-negative), optional, default=0 the size(MB) of the shuffle chunk, used

with shuffle=True, it can enable global shuffling

shuffle.chunk.seed

int, optional, default='0' the seed for chunk shuffling

label.pad.width

int, optional, default='0' pad output label width if set larger than 0, -1 for auto

estimate

label.pad.value

float, optional, default=-1 label padding value if enabled

shuffle boolean, optional, default=0 Augmentation Param: Whether to shuffle data.

seed int, optional, default='0' Augmentation Param: Random Seed.

batch.size int (non-negative), required Batch size.

round.batch boolean, optional, default=1 Whether to use round robin to handle overflow

batch or not.

prefetch.buffer

long (non-negative), optional, default=4 Maximum number of batches to prefetch.

ctx 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for.

dtype None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

resize int, optional, default='-1' Augmentation Param: scale shorter edge to size before

applying other augmentations, -1 to disable.

rand.crop.prob float, optional, default=0 Augmentation Param: Probability of random cropping,

 $\leq 0$  to disable

min.crop.scales

tuple of <float>, optional, default=[0] Augmentation Param: Min crop scales.

max.crop.scales

tuple of <float>, optional, default=[1] Augmentation Param: Max crop scales.

min.crop.aspect.ratios

tuple of <float>, optional, default=[1] Augmentation Param: Min crop aspect ratios.

max.crop.aspect.ratios

tuple of <float>, optional, default=[1] Augmentation Param: Max crop aspect ratios.

min.crop.overlaps

tuple of <float>, optional, default=[0] Augmentation Param: Minimum crop IOU between crop box and ground-truths.

max.crop.overlaps

tuple of <float>, optional, default=[1] Augmentation Param: Maximum crop IOU between crop\_box and ground-truth.

min.crop.sample.coverages

tuple of <float>, optional, default=[0] Augmentation Param: Minimum ratio of intersect/crop\_area between crop box and ground-truths.

max.crop.sample.coverages

tuple of <float>, optional, default=[1] Augmentation Param: Maximum ratio of intersect/crop\_area between crop box and ground-truths.

min.crop.object.coverages

tuple of <float>, optional, default=[0] Augmentation Param: Minimum ratio of intersect/gt\_area between crop box and ground-truths.

max.crop.object.coverages

tuple of <float>, optional, default=[1] Augmentation Param: Maximum ratio of intersect/gt area between crop box and ground-truths.

num.crop.sampler

int, optional, default='1' Augmentation Param: Number of crop samplers.

crop.emit.mode 'center', 'overlap',optional, default='center' Augmentation Param: Emition mode for invalid ground-truths after crop. center: emit if centroid of object is out of crop region; overlap: emit if overlap is less than emit\_overlap\_thresh.

emit.overlap.thresh

float, optional, default=0.300000012 Augmentation Param: Emit overlap thresh for emit mode overlap only.

max.crop.trials

Shape(tuple), optional, default=[25] Augmentation Param: Skip cropping if fail crop trail count exceeds this number.

rand.pad.prob float, optional, default=0 Augmentation Param: Probability for random padding.

max.pad.scale float, optional, default=1 Augmentation Param: Maximum padding scale.

max.random.hue int, optional, default='0' Augmentation Param: Maximum random value of H channel in HSL color space.

random.hue.prob

float, optional, default=0 Augmentation Param: Probability to apply random hue.

max.random.saturation

int, optional, default='0' Augmentation Param: Maximum random value of S channel in HSL color space.

random.saturation.prob

float, optional, default=0 Augmentation Param: Probability to apply random saturation.

max.random.illumination

int, optional, default='0' Augmentation Param: Maximum random value of L channel in HSL color space.

random.illumination.prob

float, optional, default=0 Augmentation Param: Probability to apply random illumination.

max.random.contrast

float, optional, default=0 Augmentation Param: Maximum random value of delta contrast.

random.contrast.prob

float, optional, default=0 Augmentation Param: Probability to apply random contrast.

rand.mirror.prob

float, optional, default=0 Augmentation Param: Probability to apply horizontal flip aka. mirror.

fill.value int, optional, default='127' Augmentation Param: Filled color value while padding. inter.method int, optional, default='1' Augmentation Param: 0-NN 1-bilinear 2-cubic 3-area

4-lanczos4 9-auto 10-rand.

data.shape Shape(tuple), required Dataset Param: Shape of each instance generated by the

DataIter.

resize.mode 'fit', 'force', 'shrink',optional, default='force' Augmentation Param: How image data fit in data\_shape. force: force reshape to data\_shape regardless of aspect ratio; shrink: ensure each side fit in data\_shape, preserve aspect ratio; fit:

fit image to data\_shape, preserve ratio, will upscale if applicable.

mean.img string, optional, default="Augmentation Param: Mean Image to be subtracted."

mean.r float, optional, default=0 Augmentation Param: Mean value on R channel.

mean.g float, optional, default=0 Augmentation Param: Mean value on G channel.

mean.b float, optional, default=0 Augmentation Param: Mean value on B channel.
mean.a float, optional, default=0 Augmentation Param: Mean value on Alpha channel.

std.r float, optional, default=0 Augmentation Param: Standard deviation on R chan-

nei.

std.g float, optional, default=0 Augmentation Param: Standard deviation on G channel.

std.b float, optional, default=0 Augmentation Param: Standard deviation on B chan-

std.a float, optional, default=0 Augmentation Param: Standard deviation on Alpha channel.

scale float, optional, default=1 Augmentation Param: Scale in color space.

# Value

iter The result mx.dataiter

mx.io.ImageRecordInt8Iter

Iterating on image RecordIO files

# Description

 $..\ note:: "ImageRecordInt8Iter" is deprecated.\ Use\ ImageRecordIter(dtype='int8')\ instead.$ 

# Usage

```
mx.io.ImageRecordInt8Iter(...)
```

# Arguments

seed.aug

path.imglist	string, optional, default=" Path to the image list (.lst) file. Generally created with tools/im2rec.py. Format (Tab separated): <index of="" record=""> <one labels="" more="" or=""> <relative folder="" from="" path="" root="">.</relative></one></index>
path.imgrec	string, optional, default=" Path to the image RecordIO (.rec) file or a directory path. Created with tools/im2rec.py.
path.imgidx	string, optional, default=" Path to the image RecordIO index (.idx) file. Created with tools/im2rec.py.
aug.seq	string, optional, default='aug_default' The augmenter names to represent sequence of augmenters to be applied, seperated by comma. Additional keyword parameters will be seen by these augmenters.
label.width	int, optional, default='1' The number of labels per image.
preprocess.thr	eads
	int, optional, default='4' The number of threads to do preprocessing.
verbose	boolean, optional, default=1 If or not output verbose information.
num.parts	int, optional, default='1' Virtually partition the data into these many parts.
part.index	int, optional, default='0' The *i*-th virtual partition to be read.
device.id	int, optional, default='0' The device id used to create context for internal NDArray. Setting device_id to -1 will create Context::CPU(0). Setting device_id to valid positive device id will create Context::CPUPinned(device_id). Default is 0.
shuffle.chunk.size	
	long (non-negative), optional, default=0 The data shuffle buffer size in MB. Only valid if shuffle is true.
shuffle.chunk.seed	

int, optional, default='0' The random seed for shuffling

int or None, optional, default='None' Random seed for augmentations.

shuffle boolean, optional, default=0 Whether to shuffle data randomly or not.

seed int, optional, default='0' The random seed.

batch.size int (non-negative), required Batch size.

round.batch boolean, optional, default=1 Whether to use round robin to handle overflow

batch or not.

prefetch.buffer

long (non-negative), optional, default=4 Maximum number of batches to prefetch.

ctx 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for.

dtype None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

resize int, optional, default='-1' Down scale the shorter edge to a new size before

applying other augmentations.

rand.crop boolean, optional, default=0 If or not randomly crop the image

random.resized.crop

boolean, optional, default=0 If or not perform random resized cropping on the image, as a standard preprocessing for resnet training on ImageNet data.

max.rotate.angle

int, optional, default='0' Rotate by a random degree in "[-v, v]"

max.aspect.ratio

float, optional, default=0 Change the aspect (namely width/height) to a random value. If min\_aspect\_ratio is None then the aspect ratio ins sampled from [1 - max\_aspect\_ratio, 1 + max\_aspect\_ratio], else it is in "[min\_aspect\_ratio, max\_aspect\_ratio]"

min.aspect.ratio

float or None, optional, default=None Change the aspect (namely width/height) to a random value in "[min\_aspect\_ratio, max\_aspect\_ratio]"

max.shear.ratio

float, optional, default=0 Apply a shear transformation (namely "(x,y)->(x+my,y)") with "m" randomly chose from "[-max\_shear\_ratio, max\_shear\_ratio]"

max.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

min.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

max.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]". Ignored if "random\_resized\_crop" is True.

min.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]"Ignored if "random\_resized\_crop" is True.

max.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

min.random.area		
	float, optional, default=1 Change the area (namely width * height) to a random value in "[min_random_area, max_random_area]". Ignored if "random_resized_crop" is False.	
max.img.size	float, optional, default=1e+10 Set the maximal width and height after all resize and rotate argumentation are applied	
min.img.size	float, optional, default=0 Set the minimal width and height after all resize and rotate argumentation are applied	
brightness	float, optional, default=0 Add a random value in "[-brightness, brightness]" to the brightness of image.	
contrast	float, optional, default=0 Add a random value in "[-contrast, contrast]" to the contrast of image.	
saturation	float, optional, default=0 Add a random value in "[-saturation, saturation]" to the saturation of image.	
pca.noise	float, optional, default=0 Add PCA based noise to the image.	
random.h	int, optional, default='0' Add a random value in "[-random_h, random_h]" to the H channel in HSL color space.	
random.s	int, optional, default='0' Add a random value in "[-random_s, random_s]" to the S channel in HSL color space.	
random.l	int, optional, default='0' Add a random value in "[-random_l, random_l]" to the L channel in HSL color space.	
rotate	int, optional, default='-1' Rotate by an angle. If set, it overwrites the "max_rotate_angle" option.	
fill.value	int, optional, default='255' Set the padding pixels value to "fill_value".	
data.shape	Shape(tuple), required The shape of a output image.	
inter.method	int, optional, default='1' The interpolation method: 0-NN 1-bilinear 2-cubic 3-area 4-lanczos4 9-auto 10-rand.	
pad	int, optional, default='0' Change size from "[width, height]" into "[pad + width + pad, pad + height + pad]" by padding pixes	

# **Details**

This iterator is identical to "ImageRecordIter" except for using "int8" as the data type instead of "float".

Defined in src/io/iter\_image\_recordio\_2.cc:L948

# Value

iter The result mx.dataiter

 $\verb|mx.io.ImageRecordIter| Iterates on image RecordIO files|$ 

# Usage

```
mx.io.ImageRecordIter(...)
```

# Arguments

_	
path.imglist	string, optional, default=" Path to the image list (.lst) file. Generally created with tools/im2rec.py. Format (Tab separated): <index of="" record=""> <one labels="" more="" or=""> <relative folder="" from="" path="" root="">.</relative></one></index>
path.imgrec	string, optional, default=" Path to the image RecordIO (.rec) file or a directory path. Created with tools/im2rec.py.
path.imgidx	string, optional, default=" Path to the image RecordIO index (.idx) file. Created with tools/im2rec.py.
aug.seq	string, optional, default='aug_default' The augmenter names to represent sequence of augmenters to be applied, seperated by comma. Additional keyword parameters will be seen by these augmenters.
label.width	int, optional, default='1' The number of labels per image.
preprocess.thr	eads
	int, optional, default='4' The number of threads to do preprocessing.
verbose	boolean, optional, default=1 If or not output verbose information.
num.parts	int, optional, default='1' Virtually partition the data into these many parts.
part.index	int, optional, default='0' The *i*-th virtual partition to be read.
device.id	int, optional, default='0' The device id used to create context for internal NDArray. Setting device_id to -1 will create Context::CPU(0). Setting device_id to valid positive device id will create Context::CPUPinned(device_id). Default is 0.
shuffle.chunk.	**
	long (non-negative), optional, default=0 The data shuffle buffer size in MB. Only valid if shuffle is true.
shuffle.chunk.	seed
	int, optional, default='0' The random seed for shuffling
seed.aug	int or None, optional, default='None' Random seed for augmentations.
shuffle	boolean, optional, default=0 Whether to shuffle data randomly or not.
seed	int, optional, default='0' The random seed.
batch.size	int (non-negative), required Batch size.
round.batch	boolean, optional, default=1 Whether to use round robin to handle overflow batch or not.
prefetch.buffe	
	long (non-negative), optional, default=4 Maximum number of batches to prefetch.

ctx 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for.

dtype None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

resize int, optional, default='-1' Down scale the shorter edge to a new size before

applying other augmentations.

rand.crop boolean, optional, default=0 If or not randomly crop the image

random.resized.crop

boolean, optional, default=0 If or not perform random resized cropping on the image, as a standard preprocessing for resnet training on ImageNet data.

max.rotate.angle

int, optional, default='0' Rotate by a random degree in "[-v, v]"

max.aspect.ratio

float, optional, default=0 Change the aspect (namely width/height) to a random value. If min\_aspect\_ratio is None then the aspect ratio ins sampled from [1 - max\_aspect\_ratio, 1 + max\_aspect\_ratio], else it is in "[min\_aspect\_ratio, max\_aspect\_ratio]"

min.aspect.ratio

float or None, optional, default=None Change the aspect (namely width/height) to a random value in "[min\_aspect\_ratio, max\_aspect\_ratio]"

max.shear.ratio

float, optional, default=0 Apply a shear transformation (namely "(x,y)->(x+my,y)") with "m" randomly chose from "[-max\_shear\_ratio, max\_shear\_ratio]"

max.crop.size int,

int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

min.crop.size

int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

max.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]". Ignored if "random\_resized\_crop" is True.

min.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]"Ignored if "random\_resized\_crop" is True.

max.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

min.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

max.img.size float, optional, default=1e+10 Set the maximal width and height after all resize and rotate argumentation are applied

min.img.size float, optional, default=0 Set the minimal width and height after all resize and rotate argumentation are applied

brightness	float, optional, default=0 Add a random value in "[-brightness, brightness]" to the brightness of image.
contrast	float, optional, default=0 Add a random value in "[-contrast, contrast]" to the contrast of image.
saturation	float, optional, default=0 Add a random value in "[-saturation, saturation]" to the saturation of image.
pca.noise	float, optional, default=0 Add PCA based noise to the image.
random.h	int, optional, default='0' Add a random value in "[-random_h, random_h]" to the H channel in HSL color space.
random.s	int, optional, default='0' Add a random value in "[-random_s, random_s]" to the S channel in HSL color space.
random.l	int, optional, default='0' Add a random value in "[-random_l, random_l]" to the L channel in HSL color space.
rotate	int, optional, default='-1' Rotate by an angle. If set, it overwrites the "max_rotate_angle" option.
fill.value	int, optional, default='255' Set the padding pixels value to "fill_value".
data.shape	Shape(tuple), required The shape of a output image.
inter.method	int, optional, default='1' The interpolation method: 0-NN 1-bilinear 2-cubic 3-area 4-lanczos4 9-auto 10-rand.
pad	int, optional, default='0' Change size from "[width, height]" into "[pad + width + pad, pad + height + pad]" by padding pixes
mirror	boolean, optional, default=0 Whether to mirror the image or not. If true, images are flipped along the horizontal axis.
rand.mirror	boolean, optional, default=0 Whether to randomly mirror images or not. If true, 50
	\itemmean.imgstring, optional, default="Filename of the mean image.
	\itemmean.rfloat, optional, default=0 The mean value to be subtracted on the R channel
	\itemmean.gfloat, optional, default=0 The mean value to be subtracted on the G channel
	\itemmean.bfloat, optional, default=0 The mean value to be subtracted on the B channel
	\itemmean.afloat, optional, default=0 The mean value to be subtracted on the alpha channel
	\itemstd.rfloat, optional, default=1 Augmentation Param: Standard deviation on R channel.
	\itemstd.gfloat, optional, default=1 Augmentation Param: Standard deviation on G channel.
	\itemstd.bfloat, optional, default=1 Augmentation Param: Standard deviation on B channel.
	\itemstd.afloat, optional, default=1 Augmentation Param: Standard deviation on Alpha channel.
	\itemscalefloat, optional, default=1 Multiply the image with a scale value.

\itemmax.random.contrastfloat, optional, default=0 Change the contrast with a value randomly chosen from "[-max\_random\_contrast, max\_random\_contrast]" \itemmax.random.illuminationfloat, optional, default=0 Change the illumination with a value randomly chosen from "[-max\_random\_illumination, max\_random\_illumination]" iter The result mx.dataiter

Reads batches of images from .rec RecordIO files. One can use "im2rec.py" tool (in tools/) to pack raw image files into RecordIO files. This iterator is less flexible to customization but is fast and has lot of language bindings. To iterate over raw images directly use "ImageIter" instead (in Python).

#### Example::

data\_iter = mx.io.ImageRecordIter( path\_imgrec="./sample.rec", # The target record file. data\_shape=(3, 227, 227), # Output data shape; 227x227 region will be cropped from the original image. batch\_size=4, # Number of items per batch. resize=256 # Resize the shorter edge to 256 before cropping. # You can specify more augmentation options. Use help(mx.io.ImageRecordIter) to see all the options. ) # You can now use the data\_iter to access batches of images. batch = data\_iter.next() # first batch. images = batch.data[0] # This will contain 4 (=batch\_size) images each of 3x227x227. # process the images ... data\_iter.reset() # To restart the iterator from the beginning.

Defined in src/io/iter\_image\_recordio\_2.cc:L911

mx.io.ImageRecordIter\_v1

Iterating on image RecordIO files

#### **Usage**

```
mx.io.ImageRecordIter_v1(...)
```

#### Arguments

path.imglist	string, optional, default=" Path to the image list (.lst) file. Generally created with tools/im2rec.py. Format (Tab separated): <index of="" record=""> <one labels="" more="" or=""> <relative folder="" from="" path="" root="">.</relative></one></index>	
path.imgrec	string, optional, default=" Path to the image RecordIO (.rec) file or a directory path. Created with tools/im2rec.py.	
path.imgidx	string, optional, default=" Path to the image RecordIO index $(.idx)$ file. Created with tools/im2rec.py.	
aug.seq	string, optional, default='aug_default' The augmenter names to represent sequence of augmenters to be applied, seperated by comma. Additional keyword parameters will be seen by these augmenters.	
label.width	int, optional, default='1' The number of labels per image.	
preprocess.threads		

int, optional, default='4' The number of threads to do preprocessing.

verbose boolean, optional, default=1 If or not output verbose information.

num.parts int, optional, default='1' Virtually partition the data into these many parts.

part.index int, optional, default='0' The \*i\*-th virtual partition to be read.

device.id int, optional, default='0' The device id used to create context for internal NDAr-

ray. Setting device\_id to -1 will create Context::CPU(0). Setting device\_id to valid positive device id will create Context::CPUPinned(device\_id). Default is

0.

shuffle.chunk.size

long (non-negative), optional, default=0 The data shuffle buffer size in MB. Only

valid if shuffle is true.

shuffle.chunk.seed

int, optional, default='0' The random seed for shuffling

seed.aug int or None, optional, default='None' Random seed for augmentations. shuffle boolean, optional, default=0 Whether to shuffle data randomly or not.

seed int, optional, default='0' The random seed.
batch.size int (non-negative), required Batch size.

round.batch boolean, optional, default=1 Whether to use round robin to handle overflow

batch or not.

prefetch.buffer

long (non-negative), optional, default=4 Maximum number of batches to prefetch.

ctx 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for.

dtype None, 'bfloat16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

resize int, optional, default='-1' Down scale the shorter edge to a new size before

applying other augmentations.

rand.crop boolean, optional, default=0 If or not randomly crop the image

random.resized.crop

boolean, optional, default=0 If or not perform random resized cropping on the image, as a standard preprocessing for resnet training on ImageNet data.

max.rotate.angle

int, optional, default='0' Rotate by a random degree in "[-v, v]"

max.aspect.ratio

float, optional, default=0 Change the aspect (namely width/height) to a random value. If min\_aspect\_ratio is None then the aspect ratio ins sampled from [1 - max\_aspect\_ratio, 1 + max\_aspect\_ratio], else it is in "[min\_aspect\_ratio, max\_aspect\_ratio]"

min.aspect.ratio

float or None, optional, default=None Change the aspect (namely width/height) to a random value in "[min\_aspect\_ratio, max\_aspect\_ratio]"

max.shear.ratio

float, optional, default=0 Apply a shear transformation (namely "(x,y)->(x+my,y)") with "m" randomly chose from "[-max\_shear\_ratio, max\_shear\_ratio]"

max.crop.size int, optional, default='-1' Crop both width and height into a random size in

"[min\_crop\_size, max\_crop\_size]. "Ignored if "random\_resized\_crop" is True.

min.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

max.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]". Ignored if "random\_resized\_crop" is True.

min.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]" Ignored if "random\_resized\_crop" is True.

max.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

min.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

max.img.size float, optional, default=1e+10 Set the maximal width and height after all resize and rotate argumentation are applied

min.img.size float, optional, default=0 Set the minimal width and height after all resize and rotate argumentation are applied

brightness float, optional, default=0 Add a random value in "[-brightness, brightness]" to the brightness of image.

contrast float, optional, default=0 Add a random value in "[-contrast, contrast]" to the contrast of image.

saturation float, optional, default=0 Add a random value in "[-saturation, saturation]" to the saturation of image.

pca.noise float, optional, default=0 Add PCA based noise to the image.

random.h int, optional, default='0' Add a random value in "[-random\_h, random\_h]" to the H channel in HSL color space.

the 11 chamier in 113L color space.

random.s int, optional, default='0' Add a random value in "[-random\_s, random\_s]" to the S channel in HSL color space.

random. 1 int, optional, default='0' Add a random value in "[-random\_l, random\_l]" to the L channel in HSL color space.

rotate int, optional, default='-1' Rotate by an angle. If set, it overwrites the "max\_rotate\_angle" option.

fill.value int, optional, default='255' Set the padding pixels value to "fill\_value".

data. shape Shape(tuple), required The shape of a output image.

inter.method int, optional, default='1' The interpolation method: 0-NN 1-bilinear 2-cubic

3-area 4-lanczos4 9-auto 10-rand.

pad int, optional, default='0' Change size from "[width, height]" into "[pad + width

+ pad, pad + height + pad]" by padding pixes

mirror boolean, optional, default=0 Whether to mirror the image or not. If true, images

are flipped along the horizontal axis.

rand.mirror boolean, optional, default=0 Whether to randomly mirror images or not. If true,

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\itemmean.imgstring, optional, default="Filename of the mean image.

\itemmean.rfloat, optional, default=0 The mean value to be subtracted on the R channel

\itemmean.gfloat, optional, default=0 The mean value to be subtracted on the G channel

\itemmean.bfloat, optional, default=0 The mean value to be subtracted on the B channel

\itemmean.afloat, optional, default=0 The mean value to be subtracted on the alpha channel

\itemstd.rfloat, optional, default=1 Augmentation Param: Standard deviation on R channel.

\itemstd.gfloat, optional, default=1 Augmentation Param: Standard deviation on G channel.

\itemstd.bfloat, optional, default=1 Augmentation Param: Standard deviation on B channel.

\itemstd.afloat, optional, default=1 Augmentation Param: Standard deviation on Alpha channel.

\itemscalefloat, optional, default=1 Multiply the image with a scale value.

\itemmax.random.contrastfloat, optional, default=0 Change the contrast with a value randomly chosen from "[-max\_random\_contrast, max\_random\_contrast]"

\itemmax.random.illuminationfloat, optional, default=0 Change the illumination with a value randomly chosen from "[-max\_random\_illumination, max\_random\_illumination]" iter The result mx.dataiter

.. note::

"ImageRecordIter\_v1" is deprecated. Use "ImageRecordIter" instead.

Read images batches from RecordIO files with a rich of data augmentation options.

One can use "tools/im2rec.py" to pack individual image files into RecordIO files.

Defined in src/io/iter\_image\_recordio.cc:L352

mx.io.ImageRecordUInt8Iter

Iterating on image RecordIO files

## **Description**

.. note:: ImageRecordUInt8Iter is deprecated. Use ImageRecordIter(dtype='uint8') instead.

#### **Usage**

```
mx.io.ImageRecordUInt8Iter(...)
```

#### **Arguments**

path.imglist string, optional, default="Path to the image list (.lst) file. Generally created with tools/im2rec.py. Format (Tab separated): <index of record> <one or more labels> <relative path from root folder>. string, optional, default=" Path to the image RecordIO (.rec) file or a directory path.imgrec path. Created with tools/im2rec.py. string, optional, default="Path to the image RecordIO index (.idx) file. Created path.imgidx with tools/im2rec.py. string, optional, default='aug\_default' The augmenter names to represent seaug.seq quence of augmenters to be applied, seperated by comma. Additional keyword parameters will be seen by these augmenters. label.width int, optional, default='1' The number of labels per image. preprocess.threads int, optional, default='4' The number of threads to do preprocessing. boolean, optional, default=1 If or not output verbose information. verbose int, optional, default='1' Virtually partition the data into these many parts. num.parts part.index int, optional, default='0' The \*i\*-th virtual partition to be read. device.id int, optional, default='0' The device id used to create context for internal NDArray. Setting device\_id to -1 will create Context::CPU(0). Setting device\_id to

0.

shuffle.chunk.size

long (non-negative), optional, default=0 The data shuffle buffer size in MB. Only

valid positive device id will create Context::CPUPinned(device\_id). Default is

valid if shuffle is true.

shuffle.chunk.seed

int, optional, default='0' The random seed for shuffling

int or None, optional, default='None' Random seed for augmentations. seed.aug shuffle boolean, optional, default=0 Whether to shuffle data randomly or not.

int, optional, default='0' The random seed. seed batch.size int (non-negative), required Batch size.

boolean, optional, default=1 Whether to use round robin to handle overflow round.batch

batch or not.

prefetch.buffer

long (non-negative), optional, default=4 Maximum number of batches to prefetch.

'cpu', 'gpu', optional, default='gpu' Context data loader optimized for. ctx

dtype None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

resize int, optional, default='-1' Down scale the shorter edge to a new size before

applying other augmentations.

rand.crop boolean, optional, default=0 If or not randomly crop the image

random.resized.crop

boolean, optional, default=0 If or not perform random resized cropping on the image, as a standard preprocessing for resnet training on ImageNet data.

max.rotate.angle

int, optional, default='0' Rotate by a random degree in "[-v, v]"

max.aspect.ratio

float, optional, default=0 Change the aspect (namely width/height) to a random value. If min\_aspect\_ratio is None then the aspect ratio ins sampled from [1 - max\_aspect\_ratio, 1 + max\_aspect\_ratio], else it is in "[min\_aspect\_ratio, max\_aspect\_ratio]"

min.aspect.ratio

float or None, optional, default=None Change the aspect (namely width/height) to a random value in "[min\_aspect\_ratio, max\_aspect\_ratio]"

max.shear.ratio

float, optional, default=0 Apply a shear transformation (namely "(x,y)->(x+my,y)") with "m" randomly chose from "[-max\_shear\_ratio, max\_shear\_ratio]"

max.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

min.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

max.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]". Ignored if "random\_resized\_crop" is True.

min.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]"Ignored if "random\_resized\_crop" is True.

max.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

min.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

max.img.size float, optional, default=1e+10 Set the maximal width and height after all resize and rotate argumentation are applied

min.img.size float, optional, default=0 Set the minimal width and height after all resize and rotate argumentation are applied

brightness float, optional, default=0 Add a random value in "[-brightness, brightness]" to the brightness of image.

contrast float, optional, default=0 Add a random value in "[-contrast, contrast]" to the contrast of image.

saturation	float, optional, default=0 Add a random value in "[-saturation, saturation]" to the saturation of image.
pca.noise	float, optional, default=0 Add PCA based noise to the image.
random.h	int, optional, default='0' Add a random value in "[-random_h, random_h]" to the H channel in HSL color space.
random.s	int, optional, default='0' Add a random value in "[-random_s, random_s]" to the S channel in HSL color space.
random.l	int, optional, default='0' Add a random value in "[-random_l, random_l]" to the L channel in HSL color space.
rotate	int, optional, default='-1' Rotate by an angle. If set, it overwrites the "max_rotate_angle" option.
fill.value	int, optional, default='255' Set the padding pixels value to "fill_value".
data.shape	Shape(tuple), required The shape of a output image.
inter.method	int, optional, default='1' The interpolation method: 0-NN 1-bilinear 2-cubic 3-area 4-lanczos4 9-auto 10-rand.
pad	int, optional, default='0' Change size from "[width, height]" into "[pad + width + pad, pad + height + pad]" by padding pixes

# **Details**

This iterator is identical to "ImageRecordIter" except for using "uint8" as the data type instead of "float".

Defined in src/io/iter\_image\_recordio\_2.cc:L930

## Value

iter The result mx.dataiter

```
mx.io.ImageRecordUInt8Iter_v1
```

Iterating on image RecordIO files

# Description

.. note::

# Usage

```
mx.io.ImageRecordUInt8Iter_v1(...)
```

#### **Arguments**

path.imglist string, optional, default="Path to the image list (.lst) file. Generally created

with tools/im2rec.py. Format (Tab separated): <index of record> <one or more

labels> < relative path from root folder>.

path.imgrec string, optional, default="Path to the image RecordIO (.rec) file or a directory

path. Created with tools/im2rec.py.

path.imgidx string, optional, default="Path to the image RecordIO index (.idx) file. Created

with tools/im2rec.py.

aug.seq string, optional, default='aug\_default' The augmenter names to represent se-

quence of augmenters to be applied, seperated by comma. Additional keyword

parameters will be seen by these augmenters.

label.width int, optional, default='1' The number of labels per image.

preprocess.threads

int, optional, default='4' The number of threads to do preprocessing.

verbose boolean, optional, default=1 If or not output verbose information.

num.parts int, optional, default='1' Virtually partition the data into these many parts.

part.index int, optional, default='0' The \*i\*-th virtual partition to be read.

device.id int, optional, default='0' The device id used to create context for internal NDAr-

ray. Setting device\_id to -1 will create Context::CPU(0). Setting device\_id to valid positive device id will create Context::CPUPinned(device\_id). Default is

0.

shuffle.chunk.size

long (non-negative), optional, default=0 The data shuffle buffer size in MB. Only

valid if shuffle is true.

shuffle.chunk.seed

int, optional, default='0' The random seed for shuffling

seed.aug int or None, optional, default='None' Random seed for augmentations. shuffle boolean, optional, default=0 Whether to shuffle data randomly or not.

seed int, optional, default='0' The random seed.

batch.size int (non-negative), required Batch size.

round.batch boolean, optional, default=1 Whether to use round robin to handle overflow

batch or not.

prefetch.buffer

long (non-negative), optional, default=4 Maximum number of batches to prefetch.

ctx 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for.

dtype None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

resize int, optional, default='-1' Down scale the shorter edge to a new size before

applying other augmentations.

rand.crop boolean, optional, default=0 If or not randomly crop the image

random.resized.crop

boolean, optional, default=0 If or not perform random resized cropping on the

image, as a standard preprocessing for resnet training on ImageNet data.

max.rotate.angle

int, optional, default='0' Rotate by a random degree in "[-v, v]"

max.aspect.ratio

float, optional, default=0 Change the aspect (namely width/height) to a random value. If min\_aspect\_ratio is None then the aspect ratio ins sampled from [1 - max\_aspect\_ratio, 1 + max\_aspect\_ratio], else it is in "[min\_aspect\_ratio, max\_aspect\_ratio]"

min.aspect.ratio

float or None, optional, default=None Change the aspect (namely width/height) to a random value in "[min\_aspect\_ratio, max\_aspect\_ratio]"

max.shear.ratio

float, optional, default=0 Apply a shear transformation (namely "(x,y)->(x+my,y)") with "m" randomly chose from "[-max\_shear\_ratio, max\_shear\_ratio]"

max.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

min.crop.size int, optional, default='-1' Crop both width and height into a random size in "[min\_crop\_size, max\_crop\_size]."Ignored if "random\_resized\_crop" is True.

max.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]". Ignored if "random\_resized\_crop" is True.

min.random.scale

float, optional, default=1 Resize into "[width\*s, height\*s]" with "s" randomly chosen from "[min\_random\_scale, max\_random\_scale]"Ignored if "random\_resized\_crop" is True.

max.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

min.random.area

float, optional, default=1 Change the area (namely width \* height) to a random value in "[min\_random\_area, max\_random\_area]". Ignored if "random\_resized\_crop" is False.

max.img.size float, optional, default=1e+10 Set the maximal width and height after all resize and rotate argumentation are applied

min.img.size float, optional, default=0 Set the minimal width and height after all resize and rotate argumentation are applied

brightness float, optional, default=0 Add a random value in "[-brightness, brightness]" to the brightness of image.

contrast float, optional, default=0 Add a random value in "[-contrast, contrast]" to the contrast of image.

saturation float, optional, default=0 Add a random value in "[-saturation, saturation]" to the saturation of image.

pca.noise float, optional, default=0 Add PCA based noise to the image.

random.h int, optional, default='0' Add a random value in "[-random\_h, random\_h]" to the H channel in HSL color space.

mx.io.LibSVMIter 53

random.s	int, optional, default='0' Add a random value in "[-random_s, random_s]" to the S channel in HSL color space.
random.l	int, optional, default='0' Add a random value in "[-random_l, random_l]" to the L channel in HSL color space.
rotate	int, optional, default='-1' Rotate by an angle. If set, it overwrites the "max_rotate_angle" option.
fill.value	int, optional, default='255' Set the padding pixels value to "fill_value".
data.shape	Shape(tuple), required The shape of a output image.
inter.method	int, optional, default='1' The interpolation method: 0-NN 1-bilinear 2-cubic 3-area 4-lanczos4 9-auto 10-rand.
pad	int, optional, default='0' Change size from "[width, height]" into "[pad + width + pad, pad + height + pad]" by padding pixes

#### **Details**

"ImageRecordUInt8Iter\_v1" is deprecated. Use "ImageRecordUInt8Iter" instead.

This iterator is identical to "ImageRecordIter" except for using "uint8" as the data type instead of "float".

Defined in src/io/iter\_image\_recordio.cc:L377

#### Value

iter The result mx.dataiter

mx.io.LibSVMIter	Returns the LibSVM iterator which returns data with 'csr' storage
	type. This iterator is experimental and should be used with care.

# Description

The input data is stored in a format similar to LibSVM file format, except that the \*\*indices are expected to be zero-based instead of one-based, and the column indices for each row are expected to be sorted in ascending order\*\*. Details of the LibSVM format are available 'here. <a href="https://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/">https://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/</a>.

## Usage

```
mx.io.LibSVMIter(...)
```

54 mx.io.LibSVMIter

### **Arguments**

data.libsvm string, required The input zero-base indexed LibSVM data file or a directory path. data.shape Shape(tuple), required The shape of one example. string, optional, default='NULL' The input LibSVM label file or a directory label.libsvm path. If NULL, all labels will be read from "data\_libsvm". label.shape Shape(tuple), optional, default=[1] The shape of one label. int, optional, default='1' partition the data into multiple parts num.parts int, optional, default='0' the index of the part will read part.index batch.size int (non-negative), required Batch size. round.batch boolean, optional, default=1 Whether to use round robin to handle overflow batch or not. prefetch.buffer long (non-negative), optional, default=4 Maximum number of batches to prefetch. 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for. ctx None, 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional, dtype

#### **Details**

The 'data\_shape' parameter is used to set the shape of each line of the data. The dimension of both 'data\_shape' and 'label\_shape' are expected to be 1.

default='None' Output data type. "None" means no change.

The 'data\_libsvm' parameter is used to set the path input LibSVM file. When it is set to a directory, all the files in the directory will be read.

When 'label\_libsvm' is set to "NULL", both data and label are read from the file specified by 'data\_libsvm'. In this case, the data is stored in 'csr' storage type, while the label is a 1D dense array.

The 'LibSVMIter' only support 'round\_batch' parameter set to "True". Therefore, if 'batch\_size' is 3 and there are 4 total rows in libsym file, 2 more examples are consumed at the first round.

When 'num\_parts' and 'part\_index' are provided, the data is split into 'num\_parts' partitions, and the iterator only reads the 'part\_index'-th partition. However, the partitions are not guaranteed to be even.

"reset()" is expected to be called only after a complete pass of data.

#### Example::

# Contents of libsym file "data.t". 1.0 0:0.5 2:1.2 -2.0 -3.0 0:0.6 1:2.4 2:1.2 4 2:-1.2

# Creates a 'LibSVMIter' with 'batch\_size'=3. »> data\_iter = mx.io.LibSVMIter(data\_libsvm = 'data.t', data\_shape = (3,), batch\_size = 3) # The data of the first batch is stored in csr storage type »> batch = data\_iter.next() »> csr = batch.data[0] <CSRNDArray 3x3 @cpu(0)> »> csr.asnumpy() [[ 0.5 0. 1.2 ] [ 0. 0. 0. ] [ 0.6 2.4 1.2]] # The label of first batch »> label = batch.label[0] »> label [ 1. -2. -3.] <NDArray 3 @cpu(0)>

>> second\_batch = data\_iter.next() # The data of the second batch >> second\_batch.data[0].asnumpy() [[ 0. 0. -1.2 ][ 0.5 0. 1.2 ][ 0. 0. 0. ]] # The label of the second batch >> second\_batch.label[0].asnumpy() [ 4. 1. -2.]

mx.io.MNISTIter 55

»> data\_iter.reset() # To restart the iterator for the second pass of the data

When 'label\_libsvm' is set to the path to another LibSVM file, data is read from 'data\_libsvm' and label from 'label\_libsvm'. In this case, both data and label are stored in the csr format. If the label column in the 'data\_libsvm' file is ignored.

#### Example::

- # Contents of libsym file "label.t" 1.0 -2.0 0:0.125 -3.0 2:1.2 4 1:1.0 2:-1.2
- # Creates a 'LibSVMIter' with specified label file »> data\_iter = mx.io.LibSVMIter(data\_libsvm = 'data.t', data\_shape = (3,), label\_libsvm = 'label.t', label\_shape = (3,), batch\_size = 3)
- # Both data and label are in csr storage type »> batch = data\_iter.next() »> csr\_data = batch.data[0] <CSRNDArray 3x3 @cpu(0)> »> csr\_data.asnumpy() [[ 0.5 0. 1.2 ] [ 0. 0. 0. ] [ 0.6 2.4 1.2 ]] 
  >> csr\_label = batch.label[0] <CSRNDArray 3x3 @cpu(0)> »> csr\_label.asnumpy() [[ 0. 0. 0. ] [ 0.125 0. 0. ] [ 0. 0. 1.2 ]]

Defined in src/io/iter\_libsvm.cc:L298

#### Value

iter The result mx.dataiter

mx.io.MNISTIter	Iterating on the MNIST dataset.	
-----------------	---------------------------------	--

#### **Description**

One can download the dataset from http://yann.lecun.com/exdb/mnist/

# Usage

```
mx.io.MNISTIter(...)
```

# Arguments

image	string, optional, default='./train-images-idx3-ubyte' Dataset Param: Mnist image path.
label	string, optional, default='./train-labels-idx1-ubyte' Dataset Param: Mnist label path.
batch.size	int, optional, default='128' Batch Param: Batch Size.
shuffle	boolean, optional, default=1 Augmentation Param: Whether to shuffle data.
flat	boolean, optional, default=0 Augmentation Param: Whether to flat the data into 1D.
seed	int, optional, default='0' Augmentation Param: Random Seed.
silent	boolean, optional, default=0 Auxiliary Param: Whether to print out data info.
num.parts	int, optional, default='1' partition the data into multiple parts
part.index	int, optional, default='0' the index of the part will read

```
prefetch.buffer
```

long (non-negative), optional, default=4 Maximum number of batches to prefetch.

ctx 'cpu', 'gpu', optional, default='gpu' Context data loader optimized for.

dtype None, 'bfloat16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='None' Output data type. "None" means no change.

#### **Details**

Defined in src/io/iter\_mnist.cc:L265

#### Value

iter The result mx.dataiter

mx.kv.create

Create a mxnet KVStore.

# Description

Create a mxnet KVStore.

## Arguments

type

string(default="local") The type of kvstore.

### Value

The kystore.

```
mx.lr_scheduler.FactorScheduler
```

 $Learning\ rate\ scheduler.\ Reduction\ based\ on\ a\ factor\ value.$ 

# Description

Learning rate scheduler. Reduction based on a factor value.

# Usage

```
mx.lr_scheduler.FactorScheduler(
   step,
   factor_val,
   stop_factor_lr = 1e-08,
   verbose = TRUE
)
```

### **Arguments**

```
step (integer) Schedule learning rate after n updates
factor (double) The factor for reducing the learning rate
```

#### Value

scheduler function

```
mx.lr_scheduler.MultiFactorScheduler

Multifactor learning rate scheduler. Reduction based on a factor value
at different steps.
```

# Description

Multifactor learning rate scheduler. Reduction based on a factor value at different steps.

## Usage

```
mx.lr_scheduler.MultiFactorScheduler(
   step,
   factor_val,
   stop_factor_lr = 1e-08,
   verbose = TRUE
)
```

## **Arguments**

```
step (array of integer) Schedule learning rate after n updates
factor (double) The factor for reducing the learning rate
```

#### Value

scheduler function

mx.metric.accuracy

Accuracy metric for classification

# Description

Accuracy metric for classification

# Usage

```
mx.metric.accuracy
```

#### **Format**

An object of class mx.metric of length 3.

mx.metric.custom

Helper function to create a customized metric

# Description

Helper function to create a customized metric

# Usage

```
mx.metric.custom(name, feval)
```

```
mx.metric.logistic_acc
```

Accuracy metric for logistic regression

# Description

Accuracy metric for logistic regression

# Usage

```
{\tt mx.metric.logistic\_acc}
```

## **Format**

An object of class mx.metric of length 3.

mx.metric.logloss 59

mx.metric.logloss

LogLoss metric for logistic regression

# Description

LogLoss metric for logistic regression

## Usage

```
mx.metric.logloss
```

## **Format**

An object of class mx.metric of length 3.

mx.metric.mae

MAE (Mean Absolute Error) metric for regression

# Description

MAE (Mean Absolute Error) metric for regression

## Usage

mx.metric.mae

### **Format**

An object of class mx.metric of length 3.

mx.metric.mse

MSE (Mean Squared Error) metric for regression

# Description

MSE (Mean Squared Error) metric for regression

## Usage

mx.metric.mse

#### **Format**

An object of class mx.metric of length 3.

60 mx.metric.rmsle

mx.metric.Perplexity Perplexity metric for language model

# Description

Perplexity metric for language model

## Usage

```
mx.metric.Perplexity
```

## **Format**

An object of class mx.metric of length 3.

mx.metric.rmse

RMSE (Root Mean Squared Error) metric for regression

# Description

RMSE (Root Mean Squared Error) metric for regression

#### Usage

mx.metric.rmse

### **Format**

An object of class mx.metric of length 3.

mx.metric.rmsle

RMSLE (Root Mean Squared Logarithmic Error) metric for regression

# Description

RMSLE (Root Mean Squared Logarithmic Error) metric for regression

## Usage

mx.metric.rmsle

#### **Format**

An object of class mx.metric of length 3.

```
mx.metric.top_k_accuracy
```

Top-k accuracy metric for classification

## **Description**

Top-k accuracy metric for classification

## Usage

```
mx.metric.top_k_accuracy
```

#### **Format**

An object of class mx.metric of length 3.

mx.mlp

Convenience interface for multiple layer perceptron

## **Description**

Convenience interface for multiple layer perceptron

### Usage

```
mx.mlp(
   data,
   label,
   hidden_node = 1,
   out_node,
   dropout = NULL,
   activation = "tanh",
   out_activation = "softmax",
   ctx = mx.ctx.default(),
   ...
)
```

## **Arguments**

data the input matrix. Only mx.io.DataIter and R array/matrix types supported.

label the training label. Only R array type supported.

hidden\_node a vector containing number of hidden nodes on each hidden layer as well as the

output layer.

out\_node the number of nodes on the output layer.

62 mx.model.buckets

dropout a number in [0,1) containing the dropout ratio from the last hidden layer to the output layer.

activation either a single string or a vector containing the names of the activation functions.

out\_activation a single string containing the name of the output activation function.

ctx whether train on cpu (default) or gpu.

other parameters passing to mx.model.FeedForward.create/

eval.metric the evaluation metric/

## **Examples**

mx.model.buckets

Train RNN with bucket support

### Description

Train RNN with bucket support

## Usage

```
mx.model.buckets(
    symbol,
    train.data,
    eval.data = NULL,
    metric = NULL,
    arg.params = NULL,
    aux.params = NULL,
    fixed.params = NULL,
    num.round = 1,
    begin.round = 1,
    initializer = mx.init.uniform(0.01),
    optimizer = "sgd",
    ctx = NULL,
    batch.end.callback = NULL,
```

```
epoch.end.callback = NULL,
kvstore = "local",
verbose = TRUE,
metric_cpu = TRUE
)
```

## **Arguments**

```
symbol Symbol or list of Symbols representing the model train.data Training data created by mx.io.bucket.iter eval.data Evaluation data created by mx.io.bucket.iter num.round int, number of epoch verbose
```

mx.model.FeedForward.create

Create a MXNet Feedforward neural net model with the specified training.

## **Description**

Create a MXNet Feedforward neural net model with the specified training.

## Usage

```
mx.model.FeedForward.create(
  symbol,
 Χ,
  y = NULL,
  ctx = NULL,
 begin.round = 1,
  num.round = 10,
  optimizer = "sgd",
  initializer = mx.init.uniform(0.01),
  eval.data = NULL,
  eval.metric = NULL,
  epoch.end.callback = NULL,
  batch.end.callback = NULL,
  array.batch.size = 128,
  array.layout = "auto",
  kvstore = "local",
  verbose = TRUE,
  arg.params = NULL,
  aux.params = NULL,
  input.names = NULL,
  output.names = NULL,
```

```
fixed.param = NULL,
allow.extra.params = FALSE,
metric_cpu = TRUE,
...
)
```

#### **Arguments**

symbol The symbolic configuration of the neural network.

X mx.io.DataIter or R array/matrix The training data.

y R array, optional label of the data This is only used when X is R array.

ctx mx.context or list of mx.context, optional The devices used to perform training.
begin.round integer (default=1) The initial iteration over the training data to train the model.
num.round integer (default=10) The number of iterations over training data to train the

model.

optimizer string, default="sgd" The optimization method.

initializer, initializer object. default=mx.init.uniform(0.01) The initialization scheme for

parameters.

eval.data mx.io.DataIter or list(data=R.array, label=R.array), optional The validation set

used for validation evaluation during the progress

eval.metric function, optional The evaluation function on the results.

epoch.end.callback

function, optional The callback when iteration ends.

batch.end.callback

function, optional The callback when one mini-batch iteration ends.

array.batch.size

integer (default=128) The batch size used for R array training.

array.layout can be "auto", "colmajor", "rowmajor", (detault=auto) The layout of array. "row-

major" is only supported for two dimensional array. For matrix, "rowmajor" means  $\dim(X) = c(\text{nexample}, \text{nfeatures})$ , "colmajor" means  $\dim(X) = c(\text{nfeatures}, \text{nexample})$  "auto" will auto detect the layout by match the feature size, and will report error when X is a square matrix to ask user to explicitly specify layout.

kvstore string (default="local") The parameter synchronization scheme in multiple de-

vices.

verbose logical (default=TRUE) Specifies whether to print information on the iterations

during training.

arg.params list, optional Model parameter, list of name to NDArray of net's weights.

aux.params list, optional Model parameter, list of name to NDArray of net's auxiliary states.

input.names optional The names of the input symbols. output.names optional The names of the output symbols.

fixed.param The parameters to be fixed during training. For these parameters, not gradients

will be calculated and thus no space will be allocated for the gradient.

allow.extra.params

Whether allow extra parameters that are not needed by symbol. If this is TRUE, no error will be thrown when arg\_params or aux\_params contain extra parameters that is not needed by the executor.

mx.model.init.params 65

## Value

model A trained mxnet model.

mx.model.init.params Parameter initialization

# Description

Parameter initialization

## Usage

```
mx.model.init.params(symbol, input.shape, output.shape, initializer, ctx)
```

### **Arguments**

symbol The symbolic configuration of the neural network.

input.shape The shape of the input for the neural network.

 $\hbox{output.shape} \qquad \hbox{The shape of the output for the neural network. It can be NULL}.$ 

 $initializer\,,\qquad initializer\,object.\ The\ initialization\ scheme\ for\ parameters.$ 

ctx mx.context. The devices used to perform initialization.

mx.model.load Load model checkpoint from file.

## **Description**

Load model checkpoint from file.

#### Usage

```
mx.model.load(prefix, iteration)
```

## **Arguments**

prefix string prefix of the model name

iteration integer Iteration number of model we would like to load.

66 mx.nd.abs

mx.model.save

Save model checkpoint into file.

## Description

Save model checkpoint into file.

## Usage

```
mx.model.save(model, prefix, iteration)
```

# Arguments

model The feedforward model to be saved.

prefix string prefix of the model name

iteration integer Iteration number of model we would like to load.

mx.nd.abs

Returns element-wise absolute value of the input.

# Description

Example::

# Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

```
abs([-2, 0, 3]) = [2, 0, 3]
```

The storage type of "abs" output depends upon the input storage type:

-  $abs(default) = default - abs(row\_sparse) = row\_sparse - abs(csr) = csr$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L721

## Value

mx.nd.Activation 67

mx.nd.Activation	Applies an activation function element-wise to the input.	
mx.nd.Activation	Applies an activation function element-wise to the input.	

## **Description**

The following activation functions are supported:

# Arguments

data NDArray-or-Symbol The input array.

act.type 'relu', 'sigmoid', 'softrelu', 'softsign', 'tanh', required Activation function to be

applied.

#### **Details**

- 'relu': Rectified Linear Unit, :math: 'y =  $\max(x, 0)$ ' - 'sigmoid': :math: 'y =  $\frac{1}{+ \exp(-x)}$ ' - 'tanh': Hyperbolic tangent, :math: 'y =  $\frac{1}{+ \exp(x)}$  -  $\exp(-x)\exp(x)$  +  $\exp(-x)$ ' - 'softrelu': Soft ReLU, or SoftPlus, :math: 'y =  $\frac{1}{+ \exp(x)}$ ' - 'softsign': :math: 'y =  $\frac{1}{+ \exp(x)}$ '

Defined in src/operator/nn/activation.cc:L165

## Value

out The result mx.ndarray

mx.nd.adam.update	Update function for Adam optimizer. Adam is seen as a generalization of AdaGrad.
-------------------	--

# Description

Adam update consists of the following steps, where g represents gradient and m, v are 1st and 2nd order moment estimates (mean and variance).

## Arguments

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
mean	NDArray-or-Symbol Moving mean
var	NDArray-or-Symbol Moving variance
lr	float, required Learning rate
beta1	float, optional, default=0.899999976 The decay rate for the 1st moment estimates.

68 mx.nd.add.n

beta2 float, optional, default=0.999000013 The decay rate for the 2nd moment esti-

mates.

epsilon float, optional, default=9.9999994e-09 A small constant for numerical stability.

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row\_sparse and all of w, m and v have the same stype

#### **Details**

.. math::

 $g_t = \adjustified J(W_t-1) \ m_t = \beta_1 \ m_t-1 + (1 - beta_1) \ g_t \ v_t = \beta_2 \ v_t-1 + (1 - beta_2) \ g_t^2 \ W_t = W_t-1 - \adjustified \ frac \ m_t \ v_t + \ensuremath{\mbox{\\mbox{\mbox{\mbox{\s\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\m\m\m\m\s\m\\\m\s\m\m\s\m\m\s\m\s\m\m\s\m\m\s\m\s\m\s\m\s\s\m\s\m\s\s\m\s\s\$ 

It updates the weights using::

 $m = beta1*m + (1-beta1)*grad v = beta2*v + (1-beta2)*(grad**2) w += - learning_rate * m / (sqrt(v) + epsilon)$ 

However, if grad's storage type is "row\_sparse", "lazy\_update" is True and the storage type of weight is the same as those of m and v, only the row slices whose indices appear in grad.indices are updated (for w, m and v)::

for row in grad.indices: m[row] = beta1\*m[row] + (1-beta1)\*grad[row] v[row] = beta2\*v[row] + (1-beta2)\*(grad[row]\*\*2) w[row] += - learning\_rate \* m[row] / (sqrt(v[row]) + epsilon)

Defined in src/operator/optimizer\_op.cc:L679

#### Value

out The result mx.ndarray

mx.nd.add.n

Adds all input arguments element-wise.

### **Description**

```
.. math:: add\_n(a_1, a_2, ..., a_n) = a_1 + a_2 + ... + a_n
```

### Arguments

args

NDArray-or-Symbol[] Positional input arguments

mx.nd.all.finite 69

#### **Details**

"add\_n" is potentially more efficient than calling "add" by 'n' times.

The storage type of "add\_n" output depends on storage types of inputs

- add\_n(row\_sparse, row\_sparse, ..) = row\_sparse - add\_n(default, csr, default) = default - add\_n(any input combinations longer than 4 (>4) with at least one default type) = default - otherwise, "add\_n" falls all inputs back to default storage and generates default storage

Defined in src/operator/tensor/elemwise\_sum.cc:L155

#### Value

out The result mx.ndarray

mx.nd.all.finite

Check if all the float numbers in the array are finite (used for AMP)

### **Description**

Defined in src/operator/contrib/all\_finite.cc:L101

## **Arguments**

data NDArray Array

init.output boolean, optional, default=1 Initialize output to 1.

#### Value

out The result mx.ndarray

 ${\sf mx.nd.amp.cast}$ 

Cast function between low precision float/FP32 used by AMP.

## Description

It casts only between low precision float/FP32 and does not do anything for other types.

#### **Arguments**

data NDArray-or-Symbol The input.

dtype 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', required

Output data type.

### **Details**

Defined in src/operator/tensor/amp\_cast.cc:L121

70 mx.nd.arccos

#### Value

out The result mx.ndarray

mx.nd.amp.multicast Cast function used by AMP, that casts its inputs to the common widest

type.

## **Description**

It casts only between low precision float/FP32 and does not do anything for other types.

## Arguments

data NDArray-or-Symbol[] Weights

num.outputs int, required Number of input/output pairs to be casted to the widest type.

cast.narrow boolean, optional, default=0 Whether to cast to the narrowest type

#### **Details**

Defined in src/operator/tensor/amp\_cast.cc:L165

#### Value

out The result mx.ndarray

mx.nd.arccos

Returns element-wise inverse cosine of the input array.

### **Description**

The input should be in range '[-1, 1]'. The output is in the closed interval :math: '[0, \pi]'

# **Arguments**

data

NDArray-or-Symbol The input array.

## **Details**

```
.. math:: arccos([-1, -.707, 0, .707, 1]) = [\pi, 3\pi/4, \pi/4, \pi/4, 0]
```

The storage type of "arccos" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L233

### Value

mx.nd.arccosh 71

mx.nd.arccosh	Returns the element-wise inverse hyperbolic cosine of the input array, \computed element-wise.
	Computed etement-wise.

## **Description**

The storage type of "arccosh" output is always dense

## Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L535

#### Value

out The result mx.ndarray

mx.nd.arcsin

Returns element-wise inverse sine of the input array.

## **Description**

The input should be in the range '[-1, 1]'. The output is in the closed interval of [:math:'-\pi/2', :math:'\pi/2'].

## Arguments

data

NDArray-or-Symbol The input array.

### **Details**

```
.. math:: \arcsin([-1, -.707, 0, .707, 1]) = [-\pi/2, -\pi/4, 0, \pi/4, \pi/2]
```

The storage type of "arcsin" output depends upon the input storage type:

-  $\arcsin(\text{default}) = \text{default} - \arcsin(\text{row\_sparse}) = \text{row\_sparse} - \arcsin(\text{csr}) = \text{csr}$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L187

#### Value

72 mx.nd.arctan

 $\begin{tabular}{ll} mx.nd.arcsinh & Returns the element-wise inverse hyperbolic sine of the input array, $$ computed element-wise. \\ \end{tabular}$ 

## **Description**

The storage type of "arcsinh" output depends upon the input storage type:

### **Arguments**

data

NDArray-or-Symbol The input array.

#### **Details**

```
- arcsinh(default) = default - arcsinh(row_sparse) = row_sparse - arcsinh(csr) = csr
Defined in src/operator/tensor/elemwise_unary_op_trig.cc:L494
```

#### Value

out The result mx.ndarray

mx.nd.arctan

Returns element-wise inverse tangent of the input array.

## **Description**

The output is in the closed interval :math: '[-\pi/2, \pi/2]'

#### **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

```
.. math:: \arctan([-1, 0, 1]) = [-\pi/4, 0, \pi/4]
```

The storage type of "arctan" output depends upon the input storage type:

- arctan(default) = default - arctan(row\_sparse) = row\_sparse - arctan(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L282

## Value

mx.nd.arctanh 73

mx.nd.arctanh	Returns the element-wise inverse hyperbolic tangent of the input array, \computed element-wise.

# Description

The storage type of "arctanh" output depends upon the input storage type:

## Arguments

data

NDArray-or-Symbol The input array.

## **Details**

```
- arctanh(default) = default - arctanh(row\_sparse) = row\_sparse - arctanh(csr) = csr
Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L579
```

### Value

out The result mx.ndarray

mx.nd.argmax Returns indices of the maximum values along an axis.	mx.nd.argmax	Returns indices of the maximum values along an axis.	
---	--------------	--	--

# Description

In the case of multiple occurrences of maximum values, the indices corresponding to the first occurrence are returned.

data	NDArray-or-Symbol The input
axis	int or None, optional, default='None' The axis along which to perform the reduction. Negative values means indexing from right to left. "Requires axis to be set as int, because global reduction is not supported yet."
keepdims	boolean, optional, default=0 If this is set to 'True', the reduced axis is left in the result as dimension with size one.

## **Details**

```
Examples::
```

```
 x = [[\ 0.,\ 1.,\ 2.],\ [\ 3.,\ 4.,\ 5.]]  // argmax along axis 0 argmax(x, axis=0) = [\ 1.,\ 1.,\ 1.] 
 // argmax along axis 1 argmax(x, axis=1) = [\ 2.,\ 2.] 
 // argmax along axis 1 keeping same dims as an input array argmax(x, axis=1, keepdims=True) = [[\ 2.],\ [\ 2.]]
```

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L52

### Value

out The result mx.ndarray

mx.nd.argmax.channel Returns argmax indices of each channel from the input array.

## **Description**

The result will be an NDArray of shape (num\_channel,).

## Arguments

data

NDArray-or-Symbol The input array

### **Details**

In case of multiple occurrences of the maximum values, the indices corresponding to the first occurrence are returned.

Examples::

```
x = [[0., 1., 2.], [3., 4., 5.]]

argmax\_channel(x) = [2., 2.]
```

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L97

#### Value

mx.nd.argmin 75

mx.nd.argmin	Returns indices of the minimum values along an axis.	

## Description

In the case of multiple occurrences of minimum values, the indices corresponding to the first occurrence are returned.

## Arguments

data	NDArray-or-Symbol The input
axis	int or None, optional, default='None' The axis along which to perform the reduction. Negative values means indexing from right to left. "Requires axis to be set as int, because global reduction is not supported yet."
keepdims	boolean, optional, default=0 If this is set to 'True', the reduced axis is left in the result as dimension with size one.

## **Details**

```
Examples::
```

```
x = [[ 0., 1., 2.], [ 3., 4., 5.]]

// argmin along axis 0 argmin(x, axis=0) = [ 0., 0., 0.]

// argmin along axis 1 argmin(x, axis=1) = [ 0., 0.]

// argmin along axis 1 keeping same dims as an input array argmin(x, axis=1, keepdims=True) = [[ 0.], [ 0.]]
```

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L77

## Value

out The result mx.ndarray

mx.nd.argsort	Returns the indices that would sort an input array along the given axis.
---------------	--

# Description

This function performs sorting along the given axis and returns an array of indices having same shape as an input array that index data in sorted order.

76 mx.nd.array

## Arguments

data NDArray-or-Symbol The input array

axis int or None, optional, default='-1' Axis along which to sort the input tensor. If

not given, the flattened array is used. Default is -1.

is ascend boolean, optional, default=1 Whether to sort in ascending or descending order.

dtype 'float16', 'float32', 'float64', 'int32', 'int64', 'uint8', optional, default='float32'

DType of the output indices. It is only valid when ret\_typ is "indices" or "both".

An error will be raised if the selected data type cannot precisely represent the indices.

### **Details**

## Examples::

```
x = [[ 0.3, 0.2, 0.4], [ 0.1, 0.3, 0.2]]
```

// sort along axis -1 argsort(x) = [[ 1., 0., 2.], [ 0., 2., 1.]]

// sort along axis 0 argsort(x, axis=0) = [[1., 0., 1.][0., 1., 0.]]

// flatten and then sort argsort(x, axis=None) = [3., 1., 5., 0., 4., 2.]

Defined in src/operator/tensor/ordering\_op.cc:L185

### Value

out The result mx.ndarray

mx.nd.array

Create a new mx.ndarray that copies the content from src on ctx.

## Description

Create a new mx.ndarray that copies the content from src on ctx.

## Usage

```
mx.nd.array(src.array, ctx = NULL)
```

## Arguments

src.array Source array data of class array, vector or matrix.

ctx optional The context device of the array. mx.ctx.default() will be used in default.

#### Value

An mx.ndarray

An Rcpp\_MXNDArray object

mx.nd.batch.dot 77

## **Examples**

```
mat = mx.nd.array(x)
mat = 1 - mat + (2 * mat)/(mat + 0.5)
as.array(mat)
```

mx.nd.batch.dot

Batchwise dot product.

## **Description**

"batch\_dot" is used to compute dot product of "x" and "y" when "x" and "y" are data in batch, namely N-D ( $N \ge 3$ ) arrays in shape of '( $B0, ..., B_i, ..., ...$ ).

## **Arguments**

1hs NDArray-or-Symbol The first input

rhs NDArray-or-Symbol The second input

transpose.a boolean, optional, default=0 If true then transpose the first input before dot.

transpose.b boolean, optional, default=0 If true then transpose the second input before dot.

forward.stype None, 'csr', 'default', 'row\_sparse',optional, default='None' The desired stor-

age type of the forward output given by user, if the combination of input storage types and this hint does not matchany implemented ones, the dot operator will perform fallback operationand still produce an output of the desired storage type.

## **Details**

For example, given "x" with shape ' $(B_0, ..., B_i, N, M)$ ' and "y" with shape ' $(B_0, ..., B_i, M, K)$ ', the result array will have shape ' $(B_0, ..., B_i, N, K)$ ', which is computed by::

```
batch\_dot(x,y)[b\_0, ..., b\_i, :, :] = dot(x[b\_0, ..., b\_i, :, :], y[b\_0, ..., b\_i, :, :])
```

Defined in src/operator/tensor/dot.cc:L127

## Value

78 mx.nd.BatchNorm

mx.nd.batch.take

Takes elements from a data batch.

## **Description**

.. note:: 'batch\_take' is deprecated. Use 'pick' instead.

## **Arguments**

a NDArray-or-Symbol The input array indices NDArray-or-Symbol The index array

#### **Details**

Given an input array of shape "(d0, d1)" and indices of shape "(i0,)", the result will be an output array of shape "(i0,)" with::

output[i] = input[i, indices[i]]

Examples::

```
x = [[1., 2.], [3., 4.], [5., 6.]]
```

// takes elements with specified indices batch\_take(x, [0,1,0]) = [1.4.5.]

Defined in src/operator/tensor/indexing\_op.cc:L841

### Value

out The result mx.ndarray

mx.nd.BatchNorm

Batch normalization.

## **Description**

Normalizes a data batch by mean and variance, and applies a scale "gamma" as well as offset "beta".

### **Arguments**

data NDArray-or-Symbol Input data to batch normalization

gamma NDArray-or-Symbol gamma array beta NDArray-or-Symbol beta array

moving.mean NDArray-or-Symbol running mean of input moving.var NDArray-or-Symbol running variance of input

eps double, optional, default=0.0010000000474974513 Epsilon to prevent div 0.

Must be no less than CUDNN\_BN\_MIN\_EPSILON defined in cudnn.h when

using cudnn (usually 1e-5)

mx.nd.BatchNorm 79

momentum float, optional, default=0.899999976 Momentum for moving average

fix.gamma boolean, optional, default=1 Fix gamma while training

use.global.stats

boolean, optional, default=0 Whether use global moving statistics instead of local batch-norm. This will force change batch-norm into a scale shift operator.

output.mean.var

boolean, optional, default=0 Output the mean and inverse std

axis int, optional, default='1' Specify which shape axis the channel is specified

cudnn.off boolean, optional, default=0 Do not select CUDNN operator, if available

min.calib.range

float or None, optional, default=None The minimum scalar value in the form of float32 obtained through calibration. If present, it will be used to by quantized batch norm op to calculate primitive scale.Note: this calib\_range is to calib bn output.

max.calib.range

float or None, optional, default=None The maximum scalar value in the form of float32 obtained through calibration. If present, it will be used to by quantized batch norm op to calculate primitive scale.Note: this calib\_range is to calib bn output.

#### **Details**

Assume the input has more than one dimension and we normalize along axis 1. We first compute the mean and variance along this axis:

.. math::

```
data\_mean[i] = mean(data[:,i,:,...]) \setminus data\_var[i] = var(data[:,i,:,...])
```

Then compute the normalized output, which has the same shape as input, as following:

.. math::

```
out[:,i,:,...] = \frac{\text{ata}_mean[i]} \operatorname{ata}_var[i] + \operatorname{beta[i]}
```

Both \*mean\* and \*var\* returns a scalar by treating the input as a vector.

Assume the input has size \*k\* on axis 1, then both "gamma" and "beta" have shape \*(k,)\*. If "output\_mean\_var" is set to be true, then outputs both "data\_mean" and the inverse of "data\_var", which are needed for the backward pass. Note that gradient of these two outputs are blocked.

Besides the inputs and the outputs, this operator accepts two auxiliary states, "moving\_mean" and "moving\_var", which are \*k\*-length vectors. They are global statistics for the whole dataset, which are updated by::

moving\_mean = moving\_mean \* momentum + data\_mean \* (1 - momentum) moving\_var = moving\_var \* momentum + data\_var \* (1 - momentum)

If "use\_global\_stats" is set to be true, then "moving\_mean" and "moving\_var" are used instead of "data\_mean" and "data\_var" to compute the output. It is often used during inference.

The parameter "axis" specifies which axis of the input shape denotes the 'channel' (separately normalized groups). The default is 1. Specifying -1 sets the channel axis to be the last item in the input shape.

80 mx.nd.BatchNorm.v1

Both "gamma" and "beta" are learnable parameters. But if "fix\_gamma" is true, then set "gamma" to 1 and its gradient to 0.

.. Note:: When "fix\_gamma" is set to True, no sparse support is provided. If "fix\_gamma is" set to False, the sparse tensors will fallback.

Defined in src/operator/nn/batch\_norm.cc:L547

#### Value

out The result mx.ndarray

mx.nd.BatchNorm.v1

Batch normalization.

## **Description**

This operator is DEPRECATED. Perform BatchNorm on the input.

#### **Arguments**

data NDArray-or-Symbol Input data to batch normalization

gamma NDArray-or-Symbol gamma array beta NDArray-or-Symbol beta array

eps float, optional, default=0.00100000005 Epsilon to prevent div 0

momentum float, optional, default=0.899999976 Momentum for moving average

fix.gamma boolean, optional, default=1 Fix gamma while training

use.global.stats

boolean, optional, default=0 Whether use global moving statistics instead of

local batch-norm. This will force change batch-norm into a scale shift operator.

output.mean.var

boolean, optional, default=0 Output All,normal mean and var

#### **Details**

Normalizes a data batch by mean and variance, and applies a scale "gamma" as well as offset "beta".

Assume the input has more than one dimension and we normalize along axis 1. We first compute the mean and variance along this axis:

.. math::

```
data\underline{nean[i]} = mean(data[:,i,:,...]) \cdot data\underline{var[i]} = var(data[:,i,:,...])
```

Then compute the normalized output, which has the same shape as input, as following:

.. math::

```
out[:,i,:,...] = \frac{\text{fracdata}[:,i,:,...] - \text{data}_mean[i]}{\text{var}[i] + \text{var}[i] + \text{var}[i] + \text{beta}[i]}
```

Both \*mean\* and \*var\* returns a scalar by treating the input as a vector.

Assume the input has size \*k\* on axis 1, then both "gamma" and "beta" have shape \*(k,)\*. If "output\_mean\_var" is set to be true, then outputs both "data\_mean" and "data\_var" as well, which are needed for the backward pass.

Besides the inputs and the outputs, this operator accepts two auxiliary states, "moving\_mean" and "moving\_var", which are \*k\*-length vectors. They are global statistics for the whole dataset, which are updated by::

moving\_mean = moving\_mean \* momentum + data\_mean \* (1 - momentum) moving\_var = moving\_var \* momentum + data\_var \* (1 - momentum)

If "use\_global\_stats" is set to be true, then "moving\_mean" and "moving\_var" are used instead of "data\_mean" and "data\_var" to compute the output. It is often used during inference.

Both "gamma" and "beta" are learnable parameters. But if "fix\_gamma" is true, then set "gamma" to 1 and its gradient to 0.

There's no sparse support for this operator, and it will exhibit problematic behavior if used with sparse tensors.

Defined in src/operator/batch\_norm\_v1.cc:L95

#### Value

out The result mx.ndarray

mx.nd.BilinearSampler Applies bilinear sampling to input feature map.

## Description

Bilinear Sampling is the key of [NIPS2015] \"Spatial Transformer Networks\". The usage of the operator is very similar to remap function in OpenCV, except that the operator has the backward pass.

#### **Arguments**

data NDArray-or-Symbol Input data to the BilinearsamplerOp.

grid NDArray-or-Symbol Input grid to the BilinearsamplerOp.grid has two channels:

x\_src, y\_src

cudnn.off boolean or None, optional, default=None whether to turn cudnn off

## Details

Given :math: 'data' and :math: 'grid', then the output is computed by

.. math::  $x\_src = grid[batch, 0, y\_dst, x\_dst] \setminus y\_src = grid[batch, 1, y\_dst, x\_dst] \setminus output[batch, channel, y\_dst, x\_dst] = G(data[batch, channel, y\_src, x\_src)$ 

:math: ' $x_dst$ ', :math: ' $y_dst$ ' enumerate all spatial locations in :math: 'output', and :math: 'G()' denotes the bilinear interpolation kernel. The out-boundary points will be padded with zeros. The shape of the output will be (data.shape[0], data.shape[1], grid.shape[2], grid.shape[3]).

82 mx.nd.BlockGrad

The operator assumes that :math: 'data' has 'NCHW' layout and :math: 'grid' has been normalized to [-1, 1].

BilinearSampler often cooperates with GridGenerator which generates sampling grids for BilinearSampler. GridGenerator supports two kinds of transformation: "affine" and "warp". If users want to design a CustomOp to manipulate :math: 'grid', please firstly refer to the code of GridGenerator.

```
Example 1::
```

```
## Zoom out data two times data = array([[[1, 4, 3, 6], [1, 8, 8, 9], [0, 4, 1, 5], [1, 0, 1, 3]]]])

affine_matrix = array([[2, 0, 0], [0, 2, 0]])

affine_matrix = reshape(affine_matrix, shape=(1, 6))

grid = GridGenerator(data=affine_matrix, transform_type='affine', target_shape=(4, 4))

out = BilinearSampler(data, grid)

out [[[[ 0, 0, 0, 0], [ 0, 3.5, 6.5, 0], [ 0, 1.25, 2.5, 0], [ 0, 0, 0, 0]]]

Example 2::

## shift data horizontally by -1 pixel

data = array([[[[1, 4, 3, 6], [1, 8, 8, 9], [0, 4, 1, 5], [1, 0, 1, 3]]]])

warp_maxtrix = array([[[[1, 1, 1, 1], [1, 1, 1, 1], [1, 1, 1, 1], [1, 1, 1, 1]], [[0, 0, 0, 0], [0, 0, 0, 0], [0, 0, 0, 0]]])

grid = GridGenerator(data=warp_matrix, transform_type='warp') out = BilinearSampler(data, grid) out [[[[ 4, 3, 6, 0], [ 8, 8, 9, 0], [ 4, 1, 5, 0], [ 0, 1, 3, 0]]]

Defined in src/operator/bilinear_sampler.cc:L256
```

### Value

out The result mx.ndarray

mx.nd.BlockGrad

Stops gradient computation.

## **Description**

Stops the accumulated gradient of the inputs from flowing through this operator in the backward direction. In other words, this operator prevents the contribution of its inputs to be taken into account for computing gradients.

### **Arguments**

data

NDArray-or-Symbol The input array.

mx.nd.broadcast.add 83

### **Details**

```
Example::
```

```
v1 = [1, 2] v2 = [0, 1] a = Variable('a') b = Variable('b') b_stop_grad = stop_gradient(3 * b) loss = MakeLoss(b_stop_grad + a) executor = loss.simple_bind(ctx=cpu(), a=(1,2), b=(1,2)) executor.forward(is_train=True, a=v1, b=v2) executor.outputs [ 1. 5.] executor.backward() executor.grad_arrays [ 0. 0.] [ 1. 1.]

Defined in src/operator/tensor/elemwise_unary_op_basic.cc:L326
```

#### Value

out The result mx.ndarray

mx.nd.broadcast.add

Returns element-wise sum of the input arrays with broadcasting.

## **Description**

'broadcast\_plus' is an alias to the function 'broadcast\_add'.

### **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
Example::
```

```
x = [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] y = [[\ 0.],\ [\ 1.]] broadcast_add(x, y) = [[\ 1.,\ 1.,\ 1.],\ [\ 2.,\ 2.,\ 2.]] broadcast_plus(x, y) = [[\ 1.,\ 1.,\ 1.],\ [\ 2.,\ 2.,\ 2.]]
Supported sparse operations: broadcast_add(csr, dense(1D)) = dense broadcast_add(dense(1D), csr) = dense Defined in src/operator/tensor/elemwise_binary_broadcast_op_basic.cc:L58
```

### Value

84 mx.nd.broadcast.axis

mx.nd.broadcast.axes Broadcasts the input array over particular axes.

## **Description**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

### **Arguments**

data	NDArray-or-Symbol The input
axis	Shape(tuple), optional, default=[] The axes to perform the broadcasting.
size	Shape(tuple), optional, default=[] Target sizes of the broadcasting axes.

### **Details**

'broadcast\_axes' is an alias to the function 'broadcast\_axis'.

### Example::

```
// given x of shape (1,2,1) x = [[[ 1.], [ 2.]]]
```

// broadcast x on on axis 2 broadcast\_axis(x, axis=2, size=3) = [[[ 1., 1., 1.], [ 2., 2., 2.]]] // broadcast x on on axes 0 and 2 broadcast\_axis(x, axis=(0,2), size=(2,3)) = [[[ 1., 1., 1.], [ 2., 2., 2.]], [[ 1., 1., 1.], [ 2., 2., 2.]]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L93

## Value

out The result mx.ndarray

mx.nd.broadcast.axis Broadcasts the input array over particular axes.

## Description

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

data	NDArray-or-Symbol The input
axis	Shape(tuple), optional, default=[] The axes to perform the broadcasting.
size	Shape(tuple), optional, default=[] Target sizes of the broadcasting axes.

mx.nd.broadcast.div 85

### **Details**

'broadcast\_axes' is an alias to the function 'broadcast\_axis'.

Example::

```
// given x of shape (1,2,1) x = [[[ 1.], [ 2.]]]
```

// broadcast x on on axis 2 broadcast\_axis(x, axis=2, size=3) = [[[ 1., 1., 1.], [ 2., 2., 2.]]] // broadcast x on on axes 0 and 2 broadcast\_axis(x, axis=(0,2), size=(2,3)) = [[[ 1., 1., 1.], [ 2., 2., 2.]], [[ 1., 1., 1.], [ 2., 2., 2.]]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L93

### Value

out The result mx.ndarray

mx.nd.broadcast.div

Returns element-wise division of the input arrays with broadcasting.

## **Description**

Example::

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

$$x = [[6., 6., 6.], [6., 6., 6.]]$$

y = [[2.], [3.]]

broadcast\_div(x, y) = [[ 3., 3., 3.], [ 2., 2., 2.]]

Supported sparse operations:

broadcast\_div(csr, dense(1D)) = csr

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L187

#### Value

mx.nd.broadcast.equal Returns the result of element-wise \*\*equal to \*\* (==) comparison operation with broadcasting.

## Description

Example::

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
x = [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]]

y = [[\ 0.],\ [\ 1.]]

broadcast_equal(x, y) = [[\ 0.,\ 0.,\ 0.],\ [\ 1.,\ 1.,\ 1.]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_logic.cc:L46
```

## Value

out The result mx.ndarray

```
mx.nd.broadcast.greater
```

Returns the result of element-wise \*\*greater than\*\* (>) comparison operation with broadcasting.

## Description

Example::

### **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
broadcast_greater(x, y) = [[ 1., 1., 1.], [ 0., 0., 0.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L82

### Value

out The result mx.ndarray

```
mx.nd.broadcast.greater.equal
```

Returns the result of element-wise \*\*greater than or equal to\*\* (>=) comparison operation with broadcasting.

## Description

Example::

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_greater\_equal(x,y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L100 \end{aligned}
```

## Value

out The result mx.ndarray

mx.nd.broadcast.hypot Returns the hypotenuse of a right angled triangle, given its "legs" with broadcasting.

## Description

It is equivalent to doing :math:  $\frac{1^2 + x_2^2}{\cdot}$ .

### **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

88 mx.nd.broadcast.lesser

## **Details**

```
Example::
```

```
x = [[ 3., 3., 3.]]

y = [[ 4.], [ 4.]]

broadcast_hypot(x, y) = [[ 5., 5., 5.], [ 5., 5., 5.]]

z = [[ 0.], [ 4.]]

broadcast_hypot(x, z) = [[ 3., 3., 3.], [ 5., 5., 5.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L158

## Value

out The result mx.ndarray

mx.nd.broadcast.lesser

Returns the result of element-wise \*\*lesser than\*\* (<) comparison operation with broadcasting.

## **Description**

Example::

# Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]

y = [[ 0.], [ 1.]]

broadcast_lesser(x, y) = [[ 0., 0., 0.], [ 0., 0., 0.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L118

#### Value

mx.nd.broadcast.lesser.equal

Returns the result of element-wise \*\*lesser than or equal to\*\* (<=) comparison operation with broadcasting.

## **Description**

Example::

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
x = [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]]

y = [[\ 0.],\ [\ 1.]]

broadcast_lesser_equal(x, y) = [[\ 0.,\ 0.,\ 0.],\ [\ 1.,\ 1.,\ 1.]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_logic.cc:L136
```

### Value

out The result mx.ndarray

mx.nd.broadcast.like Broadcasts lhs to have the same shape as rhs.

## Description

Broadcasting is a mechanism that allows NDArrays to perform arithmetic operations with arrays of different shapes efficiently without creating multiple copies of arrays. Also see, 'Broadcasting <a href="https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html">https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html</a> '\_ for more explanation.

lhs	NDArray-or-Symbol First input.
rhs	NDArray-or-Symbol Second input.
lhs.axes	Shape or None, optional, default=None Axes to perform broadcast on in the first input array
rhs.axes	Shape or None, optional, default=None Axes to copy from the second input array

## **Details**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

For example::

```
broadcast\_like([[1,2,3]], [[5,6,7],[7,8,9]]) = [[1., 2., 3.], [1., 2., 3.]]) \\ broadcast\_like([9], [1,2,3,4,5], lhs\_axes=(0,), rhs\_axes=(-1,)) = [9,9,9,9,9] \\ lhs\_axes=(0,0,0,0) = [1,0,0,0] \\ lhs\_
```

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L180

### Value

out The result mx.ndarray

```
{\tt mx.nd.broadcast.logical.and}
```

Returns the result of element-wise \*\*logical and\*\* with broadcasting.

## Description

Example::

# **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
broadcast_logical_and(x, y) = [[ 0., 0., 0.], [ 1., 1., 1.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L154

#### Value

```
mx.nd.broadcast.logical.or
```

Returns the result of element-wise \*\*logical or\*\* with broadcasting.

### **Description**

Example::

## Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

```
x = [[1, 1, 0], [1, 1, 0]]

y = [[1], [0]]

broadcast_logical_or(x, y) = [[1, 1, 1, 1], [1, 1, 0]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_logic.cc:L172
```

### Value

out The result mx.ndarray

```
mx.nd.broadcast.logical.xor
```

Returns the result of element-wise \*\*logical xor\*\* with broadcasting.

## **Description**

Example::

## Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

```
x = [[1., 1., 0.], [1., 1., 0.]]

y = [[1.], [0.]]

broadcast_logical_xor(x, y) = [[0., 0., 1.], [1., 1., 0.]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_logic.cc:L190
```

92 mx.nd.broadcast.minimum

### Value

out The result mx.ndarray

```
mx.nd.broadcast.maximum
```

Returns element-wise maximum of the input arrays with broadcasting.

## **Description**

This function compares two input arrays and returns a new array having the element-wise maxima.

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

```
Example::
```

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
broadcast_maximum(x, y) = [[ 1., 1., 1.], [ 1., 1., 1.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L81

#### Value

out The result mx.ndarray

```
mx.nd.broadcast.minimum
```

Returns element-wise minimum of the input arrays with broadcasting.

## **Description**

This function compares two input arrays and returns a new array having the element-wise minima.

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function

mx.nd.broadcast.minus 93

### **Details**

```
Example::
```

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
```

broadcast\_maximum(x, y) = [[0., 0., 0.], [1., 1., 1.]]

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L117

### Value

out The result mx.ndarray

mx.nd.broadcast.minus Returns element-wise difference of the input arrays with broadcasting.

# Description

'broadcast\_minus' is an alias to the function 'broadcast\_sub'.

# Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

Example::

```
x = [[1., 1., 1.], [1., 1., 1.]]

y = [[0.], [1.]]

broadcast_sub(x, y) = [[1., 1., 1.], [0., 0., 0.]]

broadcast_minus(x, y) = [[1., 1., 1.], [0., 0., 0.]]
```

Supported sparse operations:

 $broadcast\_sub/minus(csr, dense(1D)) = dense \ broadcast\_sub/minus(dense(1D), csr) = dense \ dense(1D)$ 

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L106

### Value

94 mx.nd.broadcast.mul

mx.nd.broadcast.mod

Returns element-wise modulo of the input arrays with broadcasting.

## **Description**

Example::

## Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

## **Details**

```
x = [[8., 8., 8.], [8., 8., 8.]]

y = [[2.], [3.]]

broadcast_mod(x, y) = [[0., 0., 0.], [2., 2., 2.]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_basic.cc:L222
```

## Value

out The result mx.ndarray

mx.nd.broadcast.mul

Returns element-wise product of the input arrays with broadcasting.

## **Description**

Example::

## Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

#### **Details**

## Value

out The result mx.ndarray

```
mx.nd.broadcast.not.equal
```

Returns the result of element-wise \*\*not equal to \*\* (!=) comparison operation with broadcasting.

## Description

Example::

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_not\_equal(x,\ y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 0.,\ 0.,\ 0.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L64 \end{aligned}
```

## Value

out The result mx.ndarray

mx.nd.broadcast.plus Returns element-wise sum of the input arrays with broadcasting.

## **Description**

'broadcast\_plus' is an alias to the function 'broadcast\_add'.

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function

## **Details**

```
Example::
```

```
x = [[1., 1., 1.], [1., 1., 1.]]
y = [[0.], [1.]]
```

broadcast\_add(x, y) = [[ 1., 1., 1.], [ 2., 2., 2.]]

broadcast\_plus(x, y) = [[ 1., 1., 1.], [ 2., 2., 2.]]

Supported sparse operations:

broadcast\_add(csr, dense(1D)) = dense broadcast\_add(dense(1D), csr) = dense

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L58

## Value

out The result mx.ndarray

mx.nd.broadcast.power Returns result of first array elements raised to powers from second array, element-wise with broadcasting.

## **Description**

Example::

## **Arguments**

NDArray-or-Symbol First input to the function 1hs NDArray-or-Symbol Second input to the function rhs

#### **Details**

```
x = [[1., 1., 1.], [1., 1., 1.]]
y = [[0.], [1.]]
broadcast_power(x, y) = [[ 2., 2., 2.], [ 4., 4., 4.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L45

### Value

mx.nd.broadcast.sub 97

mx.nd.broadcast.sub

Returns element-wise difference of the input arrays with broadcasting.

## Description

'broadcast\_minus' is an alias to the function 'broadcast\_sub'.

## **Arguments**

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the function

### **Details**

```
Example::
```

```
x = [[1., 1., 1.], [1., 1., 1.]]

y = [[0.], [1.]]

broadcast_sub(x, y) = [[1., 1., 1.], [0., 0., 0.]]

broadcast_minus(x, y) = [[1., 1., 1.], [0., 0., 0.]]
```

Supported sparse operations:

broadcast\_sub/minus(csr, dense(1D)) = dense broadcast\_sub/minus(dense(1D), csr) = dense Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L106

## Value

out The result mx.ndarray

mx.nd.broadcast.to

Broadcasts the input array to a new shape.

#### **Description**

Broadcasting is a mechanism that allows NDArrays to perform arithmetic operations with arrays of different shapes efficiently without creating multiple copies of arrays. Also see, 'Broadcasting <a href="https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html">https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html</a> '\_ for more explanation.

## **Arguments**

data	NDArray-or-Symbol The input

shape Shape(tuple), optional, default=[] The shape of the desired array. We can set the

dim to zero if it's same as the original. E.g 'A = broadcast\_to(B, shape=(10, 0,

0)) has the same meaning as 'A = broadcast\_axis(B, axis=0, size=10)'.

98 mx.nd.Cast

### **Details**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

For example::

```
broadcast_to([[1,2,3]], shape=(2,3)) = [[1., 2., 3.], [1., 2., 3.]])
```

The dimension which you do not want to change can also be kept as '0' which means copy the original value. So with 'shape=(2,0)', we will obtain the same result as in the above example.

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L117

#### Value

out The result mx.ndarray

mx.nd.Cast

Casts all elements of the input to a new type.

## **Description**

```
.. note:: "Cast" is deprecated. Use "cast" instead.
```

### **Arguments**

data NDArray-or-Symbol The input.

dtype 'bfloat16', 'bool', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8',

required Output data type.

## Details

## Example::

```
cast([0.9, 1.3], dtype='int32') = [0, 1] cast([1e20, 11.1], dtype='float16') = [inf, 11.09375] cast([300, 11.1, 10.9, -1, -3], dtype='uint8') = [44, 11, 10, 255, 253]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L665

### Value

mx.nd.cast 99

## Description

```
.. note:: "Cast" is deprecated. Use "cast" instead.
```

## **Arguments**

data NDArray-or-Symbol The input.

dtype 'bfloat16', 'bool', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8',

required Output data type.

## **Details**

## Example::

```
cast([0.9, 1.3], dtype='int32') = [0, 1] cast([1e20, 11.1], dtype='float16') = [inf, 11.09375] cast([300, 11.1, 10.9, -1, -3], dtype='uint8') = [44, 11, 10, 255, 253]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L665

## Value

out The result mx.ndarray

mx.nd.cast.storage Casts tensor storage type to the new type.

## Description

When an NDArray with default storage type is cast to csr or row\_sparse storage, the result is compact, which means:

## **Arguments**

data NDArray-or-Symbol The input.

stype 'csr', 'default', 'row\_sparse', required Output storage type.

100 mx.nd.cbrt

### **Details**

- for csr, zero values will not be retained - for row\_sparse, row slices of all zeros will not be retained The storage type of "cast\_storage" output depends on stype parameter:

```
    - cast_storage(csr, 'default') = default - cast_storage(row_sparse, 'default') = default - cast_storage(default, 'csr') = csr - cast_storage(default, 'row_sparse') = row_sparse - cast_storage(csr, 'csr') = csr - cast_storage(row_sparse, 'row_sparse') = row_sparse
```

## Example::

```
dense = [[0., 1., 0.], [2., 0., 3.], [0., 0., 0.], [0., 0., 0.]]
```

# cast to row\_sparse storage type rsp = cast\_storage(dense, 'row\_sparse') rsp.indices = [0, 1] rsp.values = [[ 0., 1., 0.], [ 2., 0., 3.]]

# cast to csr storage type csr = cast\_storage(dense, 'csr') csr.indices = [1, 0, 2] csr.values = [1, 2, 3.] csr.indptr = [0, 1, 3, 3, 3]

Defined in src/operator/tensor/cast\_storage.cc:L71

#### Value

out The result mx.ndarray

mx.nd.cbrt

Returns element-wise cube-root value of the input.

## **Description**

```
.. math:: cbrt(x) = \sqrt{3}x
```

## **Arguments**

data

NDArray-or-Symbol The input array.

#### **Details**

Example::

```
cbrt([1, 8, -125]) = [1, 2, -5]
```

The storage type of "cbrt" output depends upon the input storage type:

- cbrt(default) = default - cbrt(row\_sparse) = row\_sparse - cbrt(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L270

## Value

mx.nd.ceil 101

mx.nd.ceil

Returns element-wise ceiling of the input.

## Description

The ceil of the scalar x is the smallest integer i, such that  $i \ge x$ .

## **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

## Example::

```
ceil([-2.1, -1.9, 1.5, 1.9, 2.1]) = [-2., -1., 2., 2., 3.]
```

The storage type of "ceil" output depends upon the input storage type:

- ceil(default) = default - ceil(row\_sparse) = row\_sparse - ceil(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L818

#### Value

out The result mx.ndarray

mx.nd.choose.element.0index

Picks elements from an input array according to the input indices along the given axis.

### Description

Given an input array of shape "(d0, d1)" and indices of shape "(i0,)", the result will be an output array of shape "(i0,)" with::

### Arguments

data	NDArray-or-Symbol The input array
index	NDArray-or-Symbol The index array

axis int or None, optional, default='-1' int or None. The axis to picking the elements.

Negative values means indexing from right to left. If is 'None', the elements in

the index w.r.t the flattened input will be picked.

keepdims boolean, optional, default=0 If true, the axis where we pick the elements is left

in the result as dimension with size one.

mode 'clip', 'wrap', optional, default='clip' Specify how out-of-bound indices behave.

Default is "clip". "clip" means clip to the range. So, if all indices mentioned are too large, they are replaced by the index that addresses the last element along an

axis. "wrap" means to wrap around.

102 mx.nd.clip

#### **Details**

```
output[i] = input[i, indices[i]]
```

By default, if any index mentioned is too large, it is replaced by the index that addresses the last element along an axis (the 'clip' mode).

This function supports n-dimensional input and (n-1)-dimensional indices arrays.

#### Examples::

```
x = [[1., 2.], [3., 4.], [5., 6.]]
```

// picks elements with specified indices along axis 0 pick(x, y=[0,1], 0) = [1., 4.]

// picks elements with specified indices along axis 1 pick(x, y=[0,1,0], 1) = [1., 4., 5.]

// picks elements with specified indices along axis 1 using 'wrap' mode // to place indicies that would normally be out of bounds pick(x, y=[2,-1,-2], 1, mode='wrap') = [1,, 4,, 5.]

$$y = [[1.], [0.], [2.]]$$

// picks elements with specified indices along axis 1 and dims are maintained pick(x, y, 1, keep-dims=True) = [[2.], [3.], [6.]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L151

#### Value

out The result mx.ndarray

mx.nd.clip

Clips (limits) the values in an array. Given an interval, values outside the interval are clipped to the interval edges. Clipping "x" between 'a\_min' and 'a\_max' would be:: .. math::  $clip(x, a_min, a_max) = \max(\min(x, a_max), a_min)$ ) Example:: x = [0, 1, 2, 3, 4, 5, 6, 7, 8, 9] clip(x,1,8) = [1, 1, 2, 3, 4, 5, 6, 7, 8, 8.] The storage type of "clip" output depends on storage types of inputs and the a\_min,  $a_max \ge 0$  a\_max 0 = 0

#### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L677

data	NDArray-or-Symbol Input array
a.min	float, required Minimum value
a.max	float, required Maximum value

mx.nd.col2im

## Value

out The result mx.ndarray

mx.nd.col2im	Combining the output column matrix of im2col back to image array.

## Description

Like :class: '~mxnet.ndarray.im2col', this operator is also used in the vanilla convolution implementation. Despite the name, col2im is not the reverse operation of im2col. Since there may be overlaps between neighbouring sliding blocks, the column elements cannot be directly put back into image. Instead, they are accumulated (i.e., summed) in the input image just like the gradient computation, so col2im is the gradient of im2col and vice versa.

# Arguments

data	NDArray-or-Symbol Input array to combine sliding blocks.
output.size	Shape(tuple), required The spatial dimension of image array: $(w,)$ , $(h, w)$ or $(d, h, w)$ .
kernel	Shape(tuple), required Sliding kernel size: (w,), (h, w) or (d, h, w).
stride	Shape(tuple), optional, default=[] The stride between adjacent sliding blocks in spatial dimension: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] The spacing between adjacent kernel points: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
pad	Shape(tuple), optional, default=[] The zero-value padding size on both sides of spatial dimension: (w,), (h, w) or (d, h, w). Defaults to no padding.

## **Details**

Using the notation in im2col, given an input column array of shape :math: '(N, C \times \prod(\textkernel), W)', this operator accumulates the column elements into output array of shape :math: '(N, C, \textout-put\_size[0], \textoutput\_size[1], ...)'. Only 1-D, 2-D and 3-D of spatial dimension is supported in this operator.

Defined in src/operator/nn/im2col.cc:L182

#### Value

104 mx.nd.concat

arrays along a giyen a	ıxis.
$\pi$	rrays along a given a

## Description

.. note:: 'Concat' is deprecated. Use 'concat' instead.

## **Arguments**

data NDArray-or-Symbol[] List of arrays to concatenate

num.args int, required Number of inputs to be concated.

dim int, optional, default='1' the dimension to be concated.

#### **Details**

The dimensions of the input arrays should be the same except the axis along which they will be concatenated. The dimension of the output array along the concatenated axis will be equal to the sum of the corresponding dimensions of the input arrays.

The storage type of "concat" output depends on storage types of inputs

- concat(csr, csr, ..., csr, dim=0) = csr - otherwise, "concat" generates output with default storage

Example::

```
x = [[1,1],[2,2]] y = [[3,3],[4,4],[5,5]] z = [[6,6],[7,7],[8,8]]

concat(x,y,z,dim=0) = [[1,1,1],[2,2,1],[3,3,1],[4,4,1],[5,5,1],[6,6,1],[7,7,1],[8,8,1]
```

Note that you cannot concat x,y,z along dimension 1 since dimension 0 is not the same for all the input arrays.

```
concat(y,z,dim=1) = [[ 3., 3., 6., 6.], [ 4., 4., 7., 7.], [ 5., 5., 8., 8.]]
```

Defined in src/operator/nn/concat.cc:L385

## Value

out The result mx.ndarray

mx.nd.concat	Joins input arrays along a given axis.
--------------	--

## Description

.. note:: 'Concat' is deprecated. Use 'concat' instead.

mx.nd.Convolution 105

## **Arguments**

data NDArray-or-Symbol[] List of arrays to concatenate

num.args int, required Number of inputs to be concated.

dim int, optional, default='1' the dimension to be concated.

### **Details**

The dimensions of the input arrays should be the same except the axis along which they will be concatenated. The dimension of the output array along the concatenated axis will be equal to the sum of the corresponding dimensions of the input arrays.

The storage type of "concat" output depends on storage types of inputs

- concat(csr, csr, ..., csr, dim=0) = csr - otherwise, "concat" generates output with default storage Example::

```
x = [[1,1],[2,2]] y = [[3,3],[4,4],[5,5]] z = [[6,6],[7,7],[8,8]] concat(x,y,z,dim=0) = [[1,1,1],[2,2],[3,3,1],[4,4,1],[5,5,1],[6,6,1],[7,7,1],[8,8,1]
```

Note that you cannot concat x,y,z along dimension 1 since dimension 0 is not the same for all the input arrays.

```
concat(y,z,dim=1) = [[3., 3., 6., 6.], [4., 4., 7., 7.], [5., 5., 8., 8.]]
```

Defined in src/operator/nn/concat.cc:L385

### Value

out The result mx.ndarray

mx.nd.Convolution Compute \*N\*-D convolution on \*(N+2)\*-D input.

## **Description**

In the 2-D convolution, given input data with shape \*(batch\_size, channel, height, width)\*, the output is computed by

data	NDArray-or-Symbol Input data to the ConvolutionOp.
weight	NDArray-or-Symbol Weight matrix.
bias	NDArray-or-Symbol Bias parameter.
kernel	Shape(tuple), required Convolution kernel size: (w,), (h, w) or (d, h, w)
stride	Shape(tuple), optional, default=[] Convolution stride: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] Convolution dilate: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.

106 mx.nd.Convolution

pad Shape(tuple), optional, default=[] Zero pad for convolution: (w,), (h, w) or (d,

h, w). Defaults to no padding.

num.filter int (non-negative), required Convolution filter(channel) number num.group int (non-negative), optional, default=1 Number of group partitions.

workspace long (non-negative), optional, default=1024 Maximum temporary workspace al-

lowed (MB) in convolution. This parameter has two usages. When CUDNN is not used, it determines the effective batch size of the convolution kernel. When CUDNN is used, it controls the maximum temporary storage used for tuning the

best CUDNN kernel when 'limited\_workspace' strategy is used.

no.bias boolean, optional, default=0 Whether to disable bias parameter.

cudnn.tune None, 'fastest', 'limited\_workspace', 'off',optional, default='None' Whether to

pick convolution algo by running performance test.

cudnn.off boolean, optional, default=0 Turn off cudnn for this layer.

layout None, 'NCDHW', 'NCHW', 'NCW', 'NDHWC', 'NHWC', optional, default='None'

Set layout for input, output and weight. Empty for default layout: NCW for 1d, NCHW for 2d and NCDHW for 3d.NHWC and NDHWC are only supported on

GPU.

#### **Details**

.. math::

 $out[n,i,:,:] = bias[i] + \sum_{j=0}^{n} data[n,j,:,:] \operatorname{star weight}[i,j,:,:]$ 

where :math: '\star' is the 2-D cross-correlation operator.

For general 2-D convolution, the shapes are

- \*\*data\*\*: \*(batch\_size, channel, height, width)\* - \*\*weight\*\*: \*(num\_filter, channel, kernel[0], kernel[1])\* - \*\*bias\*\*: \*(num\_filter,)\* - \*\*out\*\*: \*(batch\_size, num\_filter, out\_height, out\_width)\*.

Define::

f(x,k,p,s,d) = floor((x+2\*p-d\*(k-1)-1)/s)+1

then we have::

out\_height=f(height, kernel[0], pad[0], stride[0], dilate[0]) out\_width=f(width, kernel[1], pad[1], stride[1], dilate[1])

If "no\_bias" is set to be true, then the "bias" term is ignored.

The default data "layout" is \*NCHW\*, namely \*(batch\_size, channel, height, width)\*. We can choose other layouts such as \*NWC\*.

If "num\_group" is larger than 1, denoted by \*g\*, then split the input "data" evenly into \*g\* parts along the channel axis, and also evenly split "weight" along the first dimension. Next compute the convolution on the \*i\*-th part of the data with the \*i\*-th weight part. The output is obtained by concatenating all the \*g\* results.

1-D convolution does not have \*height\* dimension but only \*width\* in space.

```
- **data**: *(batch_size, channel, width)* - **weight**: *(num_filter, channel, kernel[0])* - **bias**: *(num_filter,)* - **out**: *(batch_size, num_filter, out_width)*.
```

mx.nd.Convolution.v1 107

3-D convolution adds an additional \*depth\* dimension besides \*height\* and \*width\*. The shapes are

- \*\*data\*\*: \*(batch\_size, channel, depth, height, width)\* - \*\*weight\*\*: \*(num\_filter, channel, kernel[0], kernel[1], kernel[2])\* - \*\*bias\*\*: \*(num\_filter,)\* - \*\*out\*\*: \*(batch\_size, num\_filter, out\_depth, out\_height, out\_width)\*.

Both "weight" and "bias" are learnable parameters.

There are other options to tune the performance.

- \*\*cudnn\_tune\*\*: enable this option leads to higher startup time but may give faster speed. Options are
- \*\*off\*\*: no tuning \*\*limited\_workspace\*\*:run test and pick the fastest algorithm that doesn't exceed workspace limit. \*\*fastest\*\*: pick the fastest algorithm and ignore workspace limit. \*\*None\*\* (default): the behavior is determined by environment variable "MXNET\_CUDNN\_AUTOTUNE\_DEFAULT". 0 for off, 1 for limited workspace (default), 2 for fastest.
- \*\*workspace\*\*: A large number leads to more (GPU) memory usage but may improve the performance.

Defined in src/operator/nn/convolution.cc:L469

#### Value

out The result mx.ndarray

mx.nd.Convolution.v1 This operator is DEPRECATED. Apply convolution to input then add a bias.

## **Description**

This operator is DEPRECATED. Apply convolution to input then add a bias.

data	NDArray-or-Symbol Input data to the ConvolutionV1Op.
weight	NDArray-or-Symbol Weight matrix.
bias	NDArray-or-Symbol Bias parameter.
kernel	Shape(tuple), required convolution kernel size: (h, w) or (d, h, w)
stride	Shape(tuple), optional, default=[] convolution stride: (h, w) or (d, h, w)
dilate	Shape(tuple), optional, default=[] convolution dilate: (h, w) or (d, h, w)
pad	Shape(tuple), optional, default=[] pad for convolution: (h, w) or (d, h, w)
num.filter	int (non-negative), required convolution filter(channel) number
num.group	int (non-negative), optional, default=1 Number of group partitions. Equivalent to slicing input into num_group partitions, apply convolution on each, then concatenate the results

108 mx.nd.copyto

workspace long (non-negative), optional, default=1024 Maximum temporary workspace al-

lowed for convolution (MB). This parameter determines the effective batch size of the convolution kernel, which may be smaller than the given batch size. Also, the workspace will be automatically enlarged to make sure that we can run the

kernel with batch\_size=1

no.bias boolean, optional, default=0 Whether to disable bias parameter.

cudnn.tune None, 'fastest', 'limited\_workspace', 'off',optional, default='None' Whether to

pick convolution algo by running performance test. Leads to higher startup time but may give faster speed. Options are: 'off': no tuning 'limited\_workspace': run test and pick the fastest algorithm that doesn't exceed workspace limit. 'fastest': pick the fastest algorithm and ignore workspace limit. If set to None

(default), behavior is determined by environment variable MXNET\_CUDNN\_AUTOTUNE\_DEFAULT:

0 for off, 1 for limited workspace (default), 2 for fastest.

cudnn.off boolean, optional, default=0 Turn off cudnn for this layer.

layout None, 'NCDHW', 'NCHW', 'NDHWC', 'NHWC', optional, default='None' Set

layout for input, output and weight. Empty for default layout: NCHW for 2d

and NCDHW for 3d.

#### Value

out The result mx.ndarray

mx.nd.copyto

Generate an mx.ndarray object on ctx, with data copied from src

## **Description**

Generate an mx.ndarray object on ctx, with data copied from src

#### Usage

```
mx.nd.copyto(src, ctx)
```

## **Arguments**

src The source mx.ndarray object.

ctx The target context.

mx.nd.Correlation 109

Correlation Applies correlation to inputs.
--

### **Description**

The correlation layer performs multiplicative patch comparisons between two feature maps.

## **Arguments**

	data1	NDArray-or-Symbol Input data1 to the correlation.
	data2	NDArray-or-Symbol Input data2 to the correlation.
	kernel.size	int (non-negative), optional, default=1 kernel size for Correlation must be an odd number
max.displacement		nt
		int (non-negative), optional, default=1 Max displacement of Correlation
	stride1	int (non-negative), optional, default=1 stride1 quantize data1 globally
	stride2	int (non-negative), optional, default=1 stride2 quantize data2 within the neighborhood centered around data1 $$
	pad.size	int (non-negative), optional, default=0 pad for Correlation
	is.multiply	boolean, optional, default=1 operation type is either multiplication or subduction

### **Details**

Given two multi-channel feature maps :math:'f\_1, f\_2', with :math:'w', :math:'h', and :math:'c' being their width, height, and number of channels, the correlation layer lets the network compare each patch from :math:'f\_1' with each patch from :math:'f\_2'.

For now we consider only a single comparison of two patches. The 'correlation' of two patches centered at :math: 'x\_1' in the first map and :math: 'x\_2' in the second map is then defined as:

.. math::

```
c(x_1, x_2) = \sum_{k=0}^{\infty} [-k,k] \le [-k,k] \le [-k,k] \le [-k,k]
```

for a square patch of size :math: 'K:=2k+1'.

Note that the equation above is identical to one step of a convolution in neural networks, but instead of convolving data with a filter, it convolves data with other data. For this reason, it has no training weights.

Computing :math:  $c(x_1, x_2)$  involves :math:  $k^2$  multiplications. Comparing all patch combinations involves :math:  $k^2$  such computations.

Given a maximum displacement :math: 'd', for each location :math: ' $x_1$ ' it computes correlations :math: ' $(x_1, x_2)$ ' only in a neighborhood of size :math: ' $(x_1, x_2)$ ' by limiting the range of :math: ' $(x_2)$ '. We use strides :math: ' $(x_1, x_2)$ ', to quantize :math: ' $(x_1, x_2)$ ' within the neighborhood centered around :math: ' $(x_1, x_2)$ '.

The final output is defined by the following expression:

```
.. math:: out[n, q, i, j] = c(x_i, j, x_q)
```

110 mx.nd.cosh

where :math:'i' and :math:'j' enumerate spatial locations in :math:'f\_1', and :math:'q' denotes the :math:'q^th' neighborhood of :math:'x\_i,j'.

Defined in src/operator/correlation.cc:L198

# Value

out The result mx.ndarray

mx.nd.cos

Computes the element-wise cosine of the input array.

# **Description**

The input should be in radians (:math: '2\pi' rad equals 360 degrees).

# **Arguments**

data

NDArray-or-Symbol The input array.

# **Details**

```
.. math:: cos([0, \pi/4, \pi/2]) = [1, 0.707, 0]
```

The storage type of "cos" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L90

# Value

out The result mx.ndarray

mx.nd.cosh

Returns the hyperbolic cosine of the input array, computed elementwise.

# **Description**

```
.. math:: cosh(x) = 0.5 \times (exp(x) + exp(-x))
```

# **Arguments**

data

NDArray-or-Symbol The input array.

## **Details**

The storage type of "cosh" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L409

mx.nd.Crop 111

# Value

out The result mx.ndarray

mx.nd.Crop	note:: 'Crop' is deprecated. Use 'slice' instead.	

# Description

Crop the 2nd and 3rd dim of input data, with the corresponding size of h\_w or with width and height of the second input symbol, i.e., with one input, we need h\_w to specify the crop height and width, otherwise the second input symbol's size will be used

# Arguments

data	Symbol or Symbol[] Tensor or List of Tensors, the second input will be used as crop_like shape reference
num.args	int, required Number of inputs for crop, if equals one, then we will use the h_wfor crop height and width, else if equals two, then we will use the heightand width of the second input symbol, we name crop_like here
offset	Shape(tuple), optional, default= $[0,0]$ crop offset coordinate: $(y, x)$
h.w	Shape(tuple), optional, default=[0,0] crop height and width: (h, w)
center.crop	boolean, optional, default=0 If set to true, then it will use be the center_crop,or it will crop using the shape of crop_like

# Details

Defined in src/operator/crop.cc:L50

# Value

112 mx.nd.crop

mx.nd.crop

Slices a region of the array. .. note:: "crop" is deprecated. Use "slice" instead. This function returns a sliced array between the indices given by 'begin' and 'end' with the corresponding 'step'. For an input array of "shape=(d\_0, d\_1, ..., d\_n-1)", slice operation with "begin=(b\_0, b\_1...b\_m-1)", "end=(e\_0, e\_1, ..., e\_m-1)", and " $step=(s_0, s_1, ..., s_m-1)$ ", where  $m \le n$ , results in an array with the shape "(|e\_0-b\_0|/|s\_0|, ..., |e\_m-1-b\_m-1|/|s\_m-1|, d\_m, ..., d\_n-1)". The resulting array's \*k\*-th dimension contains elements from the \*k\*-th dimension of the input array starting from index "b\_k" (inclusive) with step "s\_k" until reaching "e\_k" (exclusive). If the \*k\*-th elements are 'None' in the sequence of 'begin', 'end', and 'step', the following rule will be used to set default values. If 's\_k' is 'None', set  $s_k=1$ . If  $s_k>0$ , set  $b_k=0$ ,  $e_k=d_k$ ; else, set  $b_k=d_k-1$ , 'e\_k=-1'. The storage type of "slice" output depends on storage types of inputs - slice(csr) = csr - otherwise, "slice" generates output with default storage .. note:: When input data storage type is csr, it only supports step=(), or step=(None,), or step=(1,) to generate a csr output. For other step parameter values, it falls back to slicing a dense tensor. Example:: x = [[1., 2., 3., 4.], [5., 6., 7., 8.], [9., 10., 11.]12.]] slice(x, begin=(0,1), end=(2,4)) = [[2., 3., 4.], [6., 7., 8.]]slice(x, begin=(None, 0), end=(None, 3), step=(-1, 2)) = [[9., 11.],[5., 7.], [1., 3.]]

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L482

### **Arguments**

data	NDArray-or-Symbol Source input
begin	Shape(tuple), required starting indices for the slice operation, supports negative indices.
end	Shape(tuple), required ending indices for the slice operation, supports negative indices.
step	Shape(tuple), optional, default=[] step for the slice operation, supports negative values.

### Value

mx.nd.ctc.loss 113

mx.nd.ctc.loss	Connectionist Temporal Classification Loss.

### **Description**

.. note:: The existing alias "contrib\_CTCLoss" is deprecated.

# **Arguments**

data NDArray-or-Symbol Input ndarray

label NDArray-or-Symbol Ground-truth labels for the loss.

data.lengths NDArray-or-Symbol Lengths of data for each of the samples. Only required

when use\_data\_lengths is true.

label.lengths NDArray-or-Symbol Lengths of labels for each of the samples. Only required

when use\_label\_lengths is true.

use.data.lengths

boolean, optional, default=0 Whether the data lenghts are decided by 'data\_lengths'.

If false, the lengths are equal to the max sequence length.

use.label.lengths

boolean, optional, default=0 Whether the label lengths are decided by 'label\_lengths', or derived from 'padding\_mask'. If false, the lengths are derived from the first occurrence of the value of 'padding\_mask'. The value of 'padding\_mask' is "0" when first CTC label is reserved for blank, and "-1" when last label is reserved

for blank. See 'blank\_label'.

blank.label 'first', 'last', optional, default='first' Set the label that is reserved for blank la-

bel.If "first", 0-th label is reserved, and label values for tokens in the vocabulary are between "1" and "alphabet\_size-1", and the padding mask is "-1". If "last", last label value "alphabet\_size-1" is reserved for blank label instead, and label values for tokens in the vocabulary are between "0" and "alphabet\_size-2", and

the padding mask is "0".

#### **Details**

The shapes of the inputs and outputs:

```
- **data**: '(sequence_length, batch_size, alphabet_size)' - **label**: '(batch_size, label_sequence_length)' - **out**: '(batch_size)'
```

The 'data' tensor consists of sequences of activation vectors (without applying softmax), with ith channel in the last dimension corresponding to i-th label for i between 0 and alphabet\_size-1 (i.e always 0-indexed). Alphabet size should include one additional value reserved for blank label. When 'blank\_label' is ""first"", the "0"-th channel is be reserved for activation of blank label, or otherwise if it is "last", "(alphabet\_size-1)"-th channel should be reserved for blank label.

"label" is an index matrix of integers. When 'blank\_label' is ""first"", the value 0 is then reserved for blank label, and should not be passed in this matrix. Otherwise, when 'blank\_label' is ""last"", the value '(alphabet\_size-1)' is reserved for blank label.

114 mx.nd.CTCLoss

If a sequence of labels is shorter than \*label\_sequence\_length\*, use the special padding value at the end of the sequence to conform it to the correct length. The padding value is '0' when 'blank\_label' is ""first"", and '-1' otherwise.

For example, suppose the vocabulary is '[a, b, c]', and in one batch we have three sequences 'ba', 'cbb', and 'abac'. When 'blank\_label' is ""first"", we can index the labels as 'a': 1, 'b': 2, 'c': 3', and we reserve the 0-th channel for blank label in data tensor. The resulting 'label' tensor should be padded to be::

[[2, 1, 0, 0], [3, 2, 2, 0], [1, 2, 1, 3]]

When 'blank\_label' is ""last"", we can index the labels as 'a': 0, 'b': 1, 'c': 2', and we reserve the channel index 3 for blank label in data tensor. The resulting 'label' tensor should be padded to be::

[[1, 0, -1, -1], [2, 1, 1, -1], [0, 1, 0, 2]]

"out" is a list of CTC loss values, one per example in the batch.

See \*Connectionist Temporal Classification: Labelling Unsegmented Sequence Data with Recurrent Neural Networks\*, A. Graves \*et al\*. for more information on the definition and the algorithm.

Defined in src/operator/nn/ctc\_loss.cc:L100

#### Value

out The result mx.ndarray

mx.nd.CTCLoss Con

Connectionist Temporal Classification Loss.

### **Description**

.. note:: The existing alias "contrib\_CTCLoss" is deprecated.

# **Arguments**

data NDArray-or-Symbol Input ndarray

label NDArray-or-Symbol Ground-truth labels for the loss.

data.lengths NDArray-or-Symbol Lengths of data for each of the samples. Only required

when use\_data\_lengths is true.

label.lengths NDArray-or-Symbol Lengths of labels for each of the samples. Only required

when use\_label\_lengths is true.

use.data.lengths

boolean, optional, default=0 Whether the data lenghts are decided by 'data\_lengths'.

If false, the lengths are equal to the max sequence length.

use.label.lengths

boolean, optional, default=0 Whether the label lengths are decided by 'label\_lengths', or derived from 'padding\_mask'. If false, the lengths are derived from the first occurrence of the value of 'padding\_mask'. The value of 'padding\_mask' is "0" when first CTC label is reserved for blank, and "-1" when last label is reserved for blank. See 'blank\_label'.

mx.nd.CTCLoss 115

blank.label

'first', 'last', optional, default='first' Set the label that is reserved for blank label.If "first", 0-th label is reserved, and label values for tokens in the vocabulary are between "1" and "alphabet\_size-1", and the padding mask is "-1". If "last", last label value "alphabet\_size-1" is reserved for blank label instead, and label values for tokens in the vocabulary are between "0" and "alphabet\_size-2", and the padding mask is "0".

### **Details**

The shapes of the inputs and outputs:

```
- **data**: '(sequence_length, batch_size, alphabet_size)' - **label**: '(batch_size, label_sequence_length)' - **out**: '(batch_size)'
```

The 'data' tensor consists of sequences of activation vectors (without applying softmax), with ith channel in the last dimension corresponding to i-th label for i between 0 and alphabet\_size-1 (i.e always 0-indexed). Alphabet size should include one additional value reserved for blank label. When 'blank\_label' is ""first"", the "0"-th channel is be reserved for activation of blank label, or otherwise if it is "last", "(alphabet\_size-1)"-th channel should be reserved for blank label.

"label" is an index matrix of integers. When 'blank\_label' is ""first"", the value 0 is then reserved for blank label, and should not be passed in this matrix. Otherwise, when 'blank\_label' is ""last"", the value '(alphabet\_size-1)' is reserved for blank label.

If a sequence of labels is shorter than \*label\_sequence\_length\*, use the special padding value at the end of the sequence to conform it to the correct length. The padding value is '0' when 'blank\_label' is ""first"", and '-1' otherwise.

For example, suppose the vocabulary is '[a, b, c]', and in one batch we have three sequences 'ba', 'cbb', and 'abac'. When 'blank\_label' is ""first"", we can index the labels as 'a': 1, 'b': 2, 'c': 3', and we reserve the 0-th channel for blank label in data tensor. The resulting 'label' tensor should be padded to be::

```
[[2, 1, 0, 0], [3, 2, 2, 0], [1, 2, 1, 3]]
```

When 'blank\_label' is ""last"", we can index the labels as "a': 0, 'b': 1, 'c': 2', and we reserve the channel index 3 for blank label in data tensor. The resulting 'label' tensor should be padded to be::

```
[[1, 0, -1, -1], [2, 1, 1, -1], [0, 1, 0, 2]]
```

"out" is a list of CTC loss values, one per example in the batch.

See \*Connectionist Temporal Classification: Labelling Unsegmented Sequence Data with Recurrent Neural Networks\*, A. Graves \*et al\*. for more information on the definition and the algorithm.

Defined in src/operator/nn/ctc\_loss.cc:L100

### Value

116 mx.nd.Custom

mx.nd.cumsum	Return the cumulative sum of the elements along a given axis.

### **Description**

Defined in src/operator/numpy/np\_cumsum.cc:L70

# **Arguments**

a NDArray-or-Symbol Input ndarray

axis int or None, optional, default='None' Axis along which the cumulative sum is

computed. The default (None) is to compute the cumsum over the flattened

array.

dtype None, 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', optional, default='None'

Type of the returned array and of the accumulator in which the elements are summed. If dtype is not specified, it defaults to the dtype of a, unless a has an integer dtype with a precision less than that of the default platform integer. In

that case, the default platform integer is used.

#### Value

out The result mx.ndarray

mx.nd.Custom	Apply a custom operator implemented in a frontend language (like
	Python).

# Description

Custom operators should override required methods like 'forward' and 'backward'. The custom operator must be registered before it can be used. Please check the tutorial here: https://mxnet.incubator.apache.org/api/faq/new\_

### Arguments

data NDArray-or-Symbol[] Input data for the custom operator.

op. type string Name of the custom operator. This is the name that is passed to 'mx.operator.register'

to register the operator.

### **Details**

Defined in src/operator/custom/custom.cc:L547

### Value

mx.nd.Deconvolution 117

mx.nd.Deconvolution

Computes 1D or 2D transposed convolution (aka fractionally strided convolution) of the input tensor. This operation can be seen as the gradient of Convolution operation with respect to its input. Convolution usually reduces the size of the input. Transposed convolution works the other way, going from a smaller input to a larger output while preserving the connectivity pattern.

# Description

Computes 1D or 2D transposed convolution (aka fractionally strided convolution) of the input tensor. This operation can be seen as the gradient of Convolution operation with respect to its input. Convolution usually reduces the size of the input. Transposed convolution works the other way, going from a smaller input to a larger output while preserving the connectivity pattern.

# **Arguments**

8	
data	NDArray-or-Symbol Input tensor to the deconvolution operation.
weight	NDArray-or-Symbol Weights representing the kernel.
bias	NDArray-or-Symbol Bias added to the result after the deconvolution operation.
kernel	Shape(tuple), required Deconvolution kernel size: (w,), (h, w) or (d, h, w). This is same as the kernel size used for the corresponding convolution
stride	Shape(tuple), optional, default=[] The stride used for the corresponding convolution: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] Dilation factor for each dimension of the input: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
pad	Shape(tuple), optional, default=[] The amount of implicit zero padding added during convolution for each dimension of the input: (w,), (h, w) or (d, h, w). "(kernel-1)/2" is usually a good choice. If 'target_shape' is set, 'pad' will be ignored and a padding that will generate the target shape will be used. Defaults to no padding.
adj	Shape(tuple), optional, default=[] Adjustment for output shape: (w,), (h, w) or (d, h, w). If 'target_shape' is set, 'adj' will be ignored and computed accordingly.
target.shape	Shape(tuple), optional, default=[] Shape of the output tensor: $(w,)$ , $(h, w)$ or $(d, h, w)$ .
num.filter	int (non-negative), required Number of output filters.
num.group	int (non-negative), optional, default=1 Number of groups partition.
workspace	long (non-negative), optional, default=512 Maximum temporary workspace allowed (MB) in deconvolution. This parameter has two usages. When CUDNN is not used, it determines the effective batch size of the deconvolution kernel.

When CUDNN is used, it controls the maximum temporary storage used for tuning the best CUDNN kernel when 'limited\_workspace' strategy is used.

mx.nd.degrees

no.bias boolean, optional, default=1 Whether to disable bias parameter.

cudnn.tune None, 'fastest', 'limited\_workspace', 'off',optional, default='None' Whether to

pick convolution algorithm by running performance test.

cudnn.off boolean, optional, default=0 Turn off cudnn for this layer.

layout None, 'NCDHW', 'NCHW', 'NCW', 'NDHWC', 'NHWC', optional, default='None'

Set layout for input, output and weight. Empty for default layout, NCW for 1d, NCHW for 2d and NCDHW for 3d.NHWC and NDHWC are only supported on

GPU.

### Value

out The result mx.ndarray

mx.nd.degrees

Converts each element of the input array from radians to degrees.

# **Description**

# **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

The storage type of "degrees" output depends upon the input storage type:

- degrees(default) = default - degrees(row\_sparse) = row\_sparse - degrees(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L332

## Value

119 mx.nd.depth.to.space

mx.nd.depth.to.space

Rearranges(permutes) data from depth into blocks of spa-Similar to ONNX DepthToSpace operator: https://github.com/onnx/onnx/blob/master/docs/Operators.md#DepthToSpace.The output is a new tensor where the values from depth dimension are moved in spatial blocks to height and width dimension. The reverse of this operation is "space to depth". .. math:: \begingather\* x \prime =  $reshape(x, [N, block\_size, block\_size, C / (block\_size ^ 2), H *$  $block \leq v + block \leq v + bloc$ [0, 3, 4, 1, 5, 2]\\\\\y = reshape(x \prime \prime, [N, C / (block\\_size \^ 2),  $H * block \land size$ ,  $W * block \land size$ )  $\land endgather * where :math: 'x' is$ an input tensor with default layout as :math: '[N, C, H, W]': [batch, channels, height, width] and :math: 'y' is the output tensor of layout :math: '[N, C / ( $block \le ^ 2$ ),  $H * block \le W * block \le '$ Example:: x = [[[0, 1, 2], [3, 4, 5]], [[6, 7, 8], [9, 10, 11]], [[12, 13, 1]]14], [15, 16, 17]], [[18, 19, 20], [21, 22, 23]]]] depth\_to\_space(x, 2) = [[[0, 6, 1, 7, 2, 8], [12, 18, 13, 19, 14, 20], [3, 9, 4, 10, 5, 11], [15, 12]]21, 16, 22, 17, 23]]]]

# **Description**

Defined in src/operator/tensor/matrix\_op.cc:L972

# **Arguments**

data NDArray-or-Symbol Input ndarray

block.size int, required Blocks of [block\_size. block\_size] are moved

### Value

out The result mx.ndarray

mx.nd.diag

Extracts a diagonal or constructs a diagonal array.

# **Description**

"diag"'s behavior depends on the input array dimensions:

#### **Arguments**

data NDArray-or-Symbol Input ndarray

k int, optional, default='0' Diagonal in question. The default is 0. Use k>0 for diagonals above the main diagonal, and k<0 for diagonals below the main diag-

onal. If input has shape (S0 S1) k must be between -S0 and S1

120 mx.nd.digamma

axis1	int, optional, default='0' The first axis of the sub-arrays of interest. Ignored when the input is a 1-D array.
axis2	int, optional, default='1' The second axis of the sub-arrays of interest. Ignored when the input is a 1-D array.

### **Details**

- 1-D arrays: constructs a 2-D array with the input as its diagonal, all other elements are zero. - N-D arrays: extracts the diagonals of the sub-arrays with axes specified by "axis1" and "axis2". The output shape would be decided by removing the axes numbered "axis1" and "axis2" from the input shape and appending to the result a new axis with the size of the diagonals in question.

For example, when the input shape is '(2, 3, 4, 5)', "axis1" and "axis2" are 0 and 2 respectively and "k" is 0, the resulting shape would be '(3, 5, 2)'.

# Examples::

```
x = [[1, 2, 3], [4, 5, 6]]
diag(x) = [1, 5]
diag(x, k=1) = [2, 6]
diag(x, k=-1) = [4]
x = [1, 2, 3]
diag(x) = [[1, 0, 0], [0, 2, 0], [0, 0, 3]]
diag(x, k=1) = [[0, 1, 0], [0, 0, 2], [0, 0, 0]]
diag(x, k=-1) = [[0, 0, 0], [1, 0, 0], [0, 2, 0]]
x = [[[1, 2], [3, 4]],
[[5, 6], [7, 8]]]
diag(x) = [[1, 7], [2, 8]]
diag(x, k=1) = [[3], [4]]
diag(x, axis1=-2, axis2=-1) = [[1, 4], [5, 8]]
Defined in src/operator/tensor/diag_op.cc:L87
```

## Value

out The result mx.ndarray

mx.nd.digamma	Returns element-wise log derivative of the gamma function \ of the
J	input.
	npm.

# **Description**

The storage type of "digamma" output is always dense

mx.nd.dot

## Arguments

data NDArray-or-Symbol The input array.

#### Value

out The result mx.ndarray

mx.nd.dot Dot product of two arrays.

## **Description**

"dot"'s behavior depends on the input array dimensions:

### **Arguments**

1hs NDArray-or-Symbol The first input
rhs NDArray-or-Symbol The second input

transpose.a boolean, optional, default=0 If true then transpose the first input before dot. transpose.b boolean, optional, default=0 If true then transpose the second input before dot. forward.stype None, 'csr', 'default', 'row\_sparse', optional, default='None' The desired stor-

age type of the forward output given by user, if the combination of input storage types and this hint does not matchany implemented ones, the dot operator will perform fallback operationand still produce an output of the desired storage type.

#### **Details**

- 1-D arrays: inner product of vectors - 2-D arrays: matrix multiplication - N-D arrays: a sum product over the last axis of the first input and the first axis of the second input

For example, given 3-D "x" with shape '(n,m,k)' and "y" with shape '(k,r,s)', the result array will have shape '(n,m,r,s)'. It is computed by::

```
dot(x,y)[i,j,a,b] = sum(x[i,j,:]*y[:,a,b])
```

# Example::

```
x = reshape([0,1,2,3,4,5,6,7], shape=(2,2,2)) y = reshape([7,6,5,4,3,2,1,0], shape=(2,2,2)) dot(x,y)[0,0,1,1] = 0 sum(x[0,0,:]*y[:,1,1]) = 0
```

The storage type of "dot" output depends on storage types of inputs, transpose option and forward\_stype option for output storage type. Implemented sparse operations include:

- dot(default, default, transpose\_a=True/False, transpose\_b=True/False) = default - dot(csr, default, transpose\_a=True) = default - dot(csr, default, transpose\_a=True) = row\_sparse - dot(csr, default) = default - dot(csr, row\_sparse) = default - dot(default, csr) = csr (CPU only) - dot(default, csr, forward\_stype='default') = default - dot(default, csr, transpose\_b=True, forward\_stype='default') = default

If the combination of input storage types and forward\_stype does not match any of the above patterns, "dot" will fallback and generate output with default storage.

122 mx.nd.Dropout

.. Note::

If the storage type of the lhs is "csr", the storage type of gradient w.r.t rhs will be "row\_sparse". Only a subset of optimizers support sparse gradients, including SGD, AdaGrad and Adam. Note that by default lazy updates is turned on, which may perform differently from standard updates. For more details, please check the Optimization API at: https://mxnet.incubator.apache.org/api/python/optimization/optimization.html Defined in src/operator/tensor/dot.cc:L77

#### Value

out The result mx.ndarray

mx.nd.Dropout	Applies dropout operation to input array.

## **Description**

- During training, each element of the input is set to zero with probability p. The whole array is rescaled by :math:  $\frac{1}{1-p}$  to keep the expected sum of the input unchanged.

## **Arguments**

~	9	
	data	NDArray-or-Symbol Input array to which dropout will be applied.
	p	float, optional, default= $0.5$ Fraction of the input that gets dropped out during training time.
	mode	'always', 'training',optional, default='training' Whether to only turn on dropout during training or to also turn on for inference.
	axes	Shape(tuple), optional, default=[] Axes for variational dropout kernel.
	cudnn.off	boolean or None, optional, default=0 Whether to turn off cudnn in dropout operator. This option is ignored if axes is specified.

### **Details**

- During testing, this operator does not change the input if mode is 'training'. If mode is 'always', the same computation as during training will be applied.

## Example::

```
random.seed(998) input_array = array([[3., 0.5, -0.5, 2., 7.], [2., -0.4, 7., 3., 0.2]]) a = symbol.Variable('a') dropout = symbol.Dropout(a, p = 0.2) executor = dropout.simple_bind(a = input_array.shape)
## If training executor.forward(is_train = True, a = input_array) executor.outputs [[ 3.75 0.625 -0. 2.5 8.75 ] [ 2.5 -0.5 8.75 3.75 0. ]]
## If testing executor.forward(is_train = False, a = input_array) executor.outputs [[ 3. 0.5 -0.5 2. 7. ] [ 2. -0.4 7. 3. 0.2 ]]
```

Defined in src/operator/nn/dropout.cc:L96

mx.nd.ElementWiseSum 123

#### Value

out The result mx.ndarray

mx.nd.ElementWiseSum Adds all input arguments element-wise.

### Description

```
.. math:: add\_n(a_1, a_2, ..., a_n) = a_1 + a_2 + ... + a_n
```

# **Arguments**

args

NDArray-or-Symbol[] Positional input arguments

#### **Details**

"add\_n" is potentially more efficient than calling "add" by 'n' times.

The storage type of "add\_n" output depends on storage types of inputs

- add\_n(row\_sparse, row\_sparse, ..) = row\_sparse - add\_n(default, csr, default) = default - add\_n(any input combinations longer than 4 (>4) with at least one default type) = default - otherwise, "add\_n" falls all inputs back to default storage and generates default storage

Defined in src/operator/tensor/elemwise\_sum.cc:L155

# Value

out The result mx.ndarray

mx.nd.elemwise.add

Adds arguments element-wise.

## **Description**

The storage type of "elemwise\_add" output depends on storage types of inputs

## **Arguments**

1hs NDArray-or-Symbol first inputrhs NDArray-or-Symbol second input

#### **Details**

- elemwise\_add(row\_sparse, row\_sparse) = row\_sparse - elemwise\_add(csr, csr) = csr - elemwise\_add(default, csr) = default - elemwise\_add(csr, default) = default - elemwise\_add(default, rsp) = default - elemwise\_add(rsp, default) = default - otherwise, "elemwise\_add" generates output with default storage

124 mx.nd.elemwise.mul

# Value

out The result mx.ndarray

mx.nd.elemwise.div

Divides arguments element-wise.

# **Description**

The storage type of "elemwise\_div" output is always dense

# **Arguments**

1hs NDArray-or-Symbol first inputrhs NDArray-or-Symbol second input

### Value

out The result mx.ndarray

mx.nd.elemwise.mul

Multiplies arguments element-wise.

# **Description**

The storage type of "elemwise\_mul" output depends on storage types of inputs

# **Arguments**

1hs NDArray-or-Symbol first inputrhs NDArray-or-Symbol second input

### **Details**

- elemwise\_mul(default, default) = default - elemwise\_mul(row\_sparse, row\_sparse) = row\_sparse - elemwise\_mul(default, row\_sparse) = row\_sparse - elemwise\_mul(row\_sparse, default) = row\_sparse - elemwise\_mul(csr, csr) = csr - otherwise, "elemwise\_mul" generates output with default storage

## Value

mx.nd.elemwise.sub

# Description

The storage type of "elemwise\_sub" output depends on storage types of inputs

# **Arguments**

lhs	NDArray-or-Symbol first input
rhs	NDArray-or-Symbol second input

### **Details**

- elemwise\_sub(row\_sparse, row\_sparse) = row\_sparse - elemwise\_sub(csr, csr) = csr - elemwise\_sub(default, csr) = default - elemwise\_sub(csr, default) = default - elemwise\_sub(default, rsp) = default - elemwise\_sub(rsp, default) = default - otherwise, "elemwise\_sub" generates output with default storage

### Value

out The result mx.ndarray

mx.nd.Embedding Maps integer indices to vector representations (embeddings).	
--	--

# **Description**

This operator maps words to real-valued vectors in a high-dimensional space, called word embeddings. These embeddings can capture semantic and syntactic properties of the words. For example, it has been noted that in the learned embedding spaces, similar words tend to be close to each other and dissimilar words far apart.

### **Arguments**

data	NDArray-or-Symbol The input array to the embedding operator.
weight	NDArray-or-Symbol The embedding weight matrix.
input.dim	int, required Vocabulary size of the input indices.
output.dim	int, required Dimension of the embedding vectors.
dtype	'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional, default='float32' Data type of weight.
sparse.grad	boolean, optional, default=0 Compute row sparse gradient in the backward calculation. If set to True, the grad's storage type is row_sparse.

mx.nd.erf

#### **Details**

For an input array of shape (d1, ..., dK), the shape of an output array is (d1, ..., dK, output\_dim). All the input values should be integers in the range [0, input\_dim).

If the input\_dim is ip0 and output\_dim is op0, then shape of the embedding weight matrix must be (ip0, op0).

When "sparse\_grad" is False, if any index mentioned is too large, it is replaced by the index that addresses the last vector in an embedding matrix. When "sparse\_grad" is True, an error will be raised if invalid indices are found.

# Examples::

 $input\_dim = 4 output\_dim = 5$ 

// Each row in weight matrix y represents a word. So, y = (w0,w1,w2,w3) y = [[0., 1., 2., 3., 4.], [5., 6., 7., 8., 9.], [10., 11., 12., 13., 14.], [15., 16., 17., 18., 19.]]

// Input array x represents n-grams(2-gram). So,  $x = [(w_1, w_3), (w_0, w_2)] x = [[1, 3, 3], [0, 2, 2]]$ 

// Mapped input x to its vector representation y. Embedding(x, y, 4, 5) = [[[ 5., 6., 7., 8., 9.], [ 15., 16., 17., 18., 19.]],

The storage type of weight can be either row\_sparse or default.

.. Note:

If "sparse\_grad" is set to True, the storage type of gradient w.r.t weights will be "row\_sparse". Only a subset of optimizers support sparse gradients, including SGD, AdaGrad and Adam. Note that by default lazy updates is turned on, which may perform differently from standard updates. For more details, please check the Optimization API at: https://mxnet.incubator.apache.org/api/python/optimization/optimization.html

Defined in src/operator/tensor/indexing\_op.cc:L603

### Value

out The result mx.ndarray

mx.nd.erf

Returns element-wise gauss error function of the input.

# Description

Example::

# Arguments

data

NDArray-or-Symbol The input array.

### **Details**

```
erf([0, -1., 10.]) = [0., -0.8427, 1.]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L887

mx.nd.erfinv 127

# Value

out The result mx.ndarray

mx.nd.erfinv

Returns element-wise inverse gauss error function of the input.

# **Description**

Example::

# **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

```
erfinv([0, 0.5., -1.]) = [0., 0.4769, -inf]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L909

### Value

out The result mx.ndarray

mx.nd.exp

Returns element-wise exponential value of the input.

# Description

```
.. math:: exp(x) = e^x \cdot 2.718^x
```

# **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

Example::

```
\exp([0, 1, 2]) = [1., 2.71828175, 7.38905621]
```

The storage type of "exp" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L64

### Value

128 mx.nd.expm1

mx.nd.expand.dims	Inserts a new axis of size 1 into the array shape For example, given " $x$ " with shape " $(2,3,4)$ ", then "expand_dims $(x, axis=1)$ " will return
	a new array with shape " $(2,1,3,4)$ ".

# **Description**

Defined in src/operator/tensor/matrix\_op.cc:L395

# **Arguments**

data NDArray-or-Symbol Source input

axis int, required Position where new axis is to be inserted. Suppose that the in-

put 'NDArray''s dimension is 'ndim', the range of the inserted axis is '[-ndim,

ndim]'

#### Value

out The result mx.ndarray

mx.nd.expm1 Returns "exp(x) - 1" computed element-wise on the input.

# Description

This function provides greater precision than "exp(x) - 1" for small values of "x".

# Arguments

data NDArray-or-Symbol The input array.

# **Details**

The storage type of "expm1" output depends upon the input storage type:

- expm1(default) = default - expm1(row\_sparse) = row\_sparse - expm1(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L244

# Value

mx.nd.fill.element.0index 129

mx.nd.fill.element.0index

Fill one element of each line(row for python, column for R/Julia) in lhs according to index indicated by rhs and values indicated by mhs. This function assume rhs uses 0-based index.

# Description

Fill one element of each line(row for python, column for R/Julia) in lhs according to index indicated by rhs and values indicated by mhs. This function assume rhs uses 0-based index.

# Arguments

1hs NDArray Left operand to the function.mhs NDArray Middle operand to the function.rhs NDArray Right operand to the function.

#### Value

out The result mx.ndarray

mx.nd.fix

Returns element-wise rounded value to the nearest \ integer towards zero of the input.

# Description

Example::

# **Arguments**

data

NDArray-or-Symbol The input array.

## **Details**

```
fix([-2.1, -1.9, 1.9, 2.1]) = [-2., -1., 1., 2.]
```

The storage type of "fix" output depends upon the input storage type:

- fix(default) = default - fix(row\_sparse) = row\_sparse - fix(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L875

### Value

mx.nd.flatten

mx.nd.Flatten

Flattens the input array into a 2-D array by collapsing the higher dimensions. .. note:: 'Flatten' is deprecated. Use 'flatten' instead. For an input array with shape "(d1, d2, ..., dk)", 'flatten' operation reshapes the input array into an output array of shape "(d1, d2\*...\*dk)". Note that the behavior of this function is different from numpy.ndarray.flatten, which behaves similar to mxnet.ndarray.reshape((-1, 1)). Example:: x = [[1,2,3], [4,5,6], [7,8,9]], [1,2,3], [4,5,6], [7,8,9]], flatten(x = [1,2,3], [4,5,6], [7,8,9]], flatten(x = [1,2,3], [4,5,6], [7,8,9]]

## **Description**

Defined in src/operator/tensor/matrix\_op.cc:L250

### **Arguments**

data

NDArray-or-Symbol Input array.

### Value

out The result mx.ndarray

mx.nd.flatten

Flattens the input array into a 2-D array by collapsing the higher dimensions. .. note:: 'Flatten' is deprecated. Use 'flatten' instead. For an input array with shape "(d1, d2, ..., dk)", 'flatten' operation reshapes the input array into an output array of shape "(d1, d2\*...\*dk)". Note that the behavior of this function is different from numpy.ndarray.flatten, which behaves similar to mxnet.ndarray.reshape((-1,)). Example:: x = [[1,2,3], [4,5,6], [7,8,9]], [1,2,3], [4,5,6], [7,8,9]], flatten(x) = x0 = x1 = x3 = x4 = x5 = x5 = x6 = x7 = x8 = x9 =

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L250

# **Arguments**

data

NDArray-or-Symbol Input array.

#### Value

mx.nd.flip

reverse(x, axis=1) = [[4., 3., 2., 1., 0.], [9., 8., 7., 6., 5.]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L832

## **Arguments**

data NDArray-or-Symbol Input data array

axis Shape(tuple), required The axis which to reverse elements.

### Value

out The result mx.ndarray

mx.nd.floor

Returns element-wise floor of the input.

# Description

The floor of the scalar x is the largest integer i, such that  $i \le x$ .

# Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

Example::

floor([-2.1, -1.9, 1.5, 1.9, 2.1]) = [-3., -2., 1., 1., 2.]

The storage type of "floor" output depends upon the input storage type:

- floor(default) = default - floor(row\_sparse) = row\_sparse - floor(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L837

## Value

mx.nd.ftml.update

mx.nd.ftml.update	The	FTML	optimizer	described	in	*FTM	L -	Follow
	the	Moving	Leader i	n Deep	Learn	ing*,	availa	ble at
	http:/	//proceedii	ngs.mlr.press	/v70/zheng1	7a/zhei	ng17a.p	df.	

# Description

.. math::

# Arguments

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
d	NDArray-or-Symbol Internal state "d_t"
V	NDArray-or-Symbol Internal state "v_t"
z	NDArray-or-Symbol Internal state "z_t"
lr	float, required Learning rate.
beta1	float, optional, default=0.600000024 Generally close to 0.5.
beta2	float, optional, default=0.999000013 Generally close to 1.
epsilon	double, optional, default=9.9999999392252903e-09 Epsilon to prevent div 0.
t	int, required Number of update.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.grad	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).

# **Details**

```
 g_t = \quad J(W_{t-1}) \cdot v_t = \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2 \cdot d_t = \frac{1 - \beta_1 \cdot v_{t-1} \cdot v_
```

# Value

mx.nd.ftrl.update 133

mx.nd.ftrl.update	Update function for Ftrl optimizer. Referenced from *Ad
	Click Prediction: a View from the Trenches*, available at http://dl.acm.org/citation.cfm?id=2488200.

# **Description**

It updates the weights using::

# **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
z	NDArray-or-Symbol z
n	NDArray-or-Symbol Square of grad
lr	float, required Learning rate
lamda1	float, optional, default=0.00999999978 The L1 regularization coefficient.
beta	float, optional, default=1 Per-Coordinate Learning Rate beta.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).

#### **Details**

```
rescaled\_grad = clip(grad * rescale\_grad, clip\_gradient) \ z += rescaled\_grad - (sqrt(n + rescaled\_grad **2) - sqrt(n)) * weight / learning\_rate n += rescaled\_grad **2 w = (sign(z) * lamda1 - z) / ((beta + sqrt(n)) / learning\_rate + wd) * (abs(z) > lamda1)
```

If w, z and n are all of "row\_sparse" storage type, only the row slices whose indices appear in grad.indices are updated (for w, z and n)::

for row in grad.indices: rescaled\_grad[row] = clip(grad[row] \* rescale\_grad, clip\_gradient) z[row] += rescaled\_grad[row] - (sqrt(n[row] + rescaled\_grad[row]\*\*2) - sqrt(n[row])) \* weight[row] / learning\_rate n[row] += rescaled\_grad[row]\*\*2 w[row] = (sign(z[row]) \* lamda1 - z[row]) / ((beta + sqrt(n[row])) / learning\_rate + wd) \* (abs(z[row]) > lamda1)

Defined in src/operator/optimizer\_op.cc:L867

#### Value

mx.nd.FullyConnected Applies a linear transformation: :math:  $Y = XW^T + b'$ .

## **Description**

If "flatten" is set to be true, then the shapes are:

### **Arguments**

data NDArray-or-Symbol Input data.

weight NDArray-or-Symbol Weight matrix.

bias NDArray-or-Symbol Bias parameter.

num.hidden int, required Number of hidden nodes of the output.

no.bias boolean, optional, default=0 Whether to disable bias parameter.

flatten boolean, optional, default=1 Whether to collapse all but the first axis of the input

data tensor.

#### **Details**

```
- **data**: '(batch_size, x1, x2, ..., xn)' - **weight**: '(num_hidden, x1 * x2 * ... * xn)' - **bias**: '(num_hidden,)' - **out**: '(batch_size, num_hidden)'
```

If "flatten" is set to be false, then the shapes are:

```
- **data**: '(x1, x2, ..., xn, input_dim)' - **weight**: '(num_hidden, input_dim)' - **bias**: '(num_hidden,)' - **out**: '(x1, x2, ..., xn, num_hidden)'
```

The learnable parameters include both "weight" and "bias".

If "no\_bias" is set to be true, then the "bias" term is ignored.

.. Note::

The sparse support for FullyConnected is limited to forward evaluation with 'row\_sparse' weight and bias, where the length of 'weight.indices' and 'bias.indices' must be equal to 'num\_hidden'. This could be useful for model inference with 'row\_sparse' weights trained with importance sampling or noise contrastive estimation.

To compute linear transformation with 'csr' sparse data, sparse.dot is recommended instead of sparse.FullyConnected.

Defined in src/operator/nn/fully\_connected.cc:L287

# Value

mx.nd.gamma 135

mx.nd.gamma

Returns the gamma function (extension of the factorial function \ to the reals), computed element-wise on the input array.

# Description

The storage type of "gamma" output is always dense

# Arguments

data

NDArray-or-Symbol The input array.

# Value

out The result mx.ndarray

mx.nd.gammaln

Returns element-wise log of the absolute value of the gamma function \ of the input.

# Description

The storage type of "gammaln" output is always dense

# Arguments

data

NDArray-or-Symbol The input array.

# Value

mx.nd.GridGenerator

mx.nd.gather.nd	Gather elements or slices from 'data' and store to a tensor whose shape is defined by 'indices'.

### **Description**

Given 'data' with shape ' $(X_0, X_1, ..., X_{N-1})$ ' and indices with shape ' $(M, Y_0, ..., Y_{K-1})$ ', the output will have shape ' $(Y_0, ..., Y_{K-1}, X_M, ..., X_{N-1})$ ', where ' $M \le N$ '. If 'M = N', output shape will simply be ' $(Y_0, ..., Y_{K-1})$ '.

### **Arguments**

data NDArray-or-Symbol data indices NDArray-or-Symbol indices

# **Details**

The elements in output is defined as follows::

```
 \begin{aligned} & \text{output}[y\_0, ..., y\_K-1, x\_M, ..., x\_N-1] = \text{data}[\text{indices}[0, y\_0, ..., y\_K-1], ..., \text{indices}[M-1, y\_0, ..., y\_K-1], x\_M, ..., x\_N-1] \end{aligned}
```

Examples::

```
 \begin{array}{l} data = [[0,1],[2,3]] \ indices = [[1,1,0],[0,1,0]] \ gather\_nd(data,indices) = [2,3,0] \\ data = [[[1,2],[3,4]],[[5,6],[7,8]]] \ indices = [[0,1],[1,0]] \ gather\_nd(data,indices) = [[3,4],[5,6]] \\ \end{array}
```

### Value

out The result mx.ndarray

mx.nd.GridGenerator Generates 2D sampling grid for bilinear sampling.

#### **Description**

Generates 2D sampling grid for bilinear sampling.

### Arguments

data NDArray-or-Symbol Input data to the function.

transform.type 'affine', 'warp', required The type of transformation. For 'affine', input data

should be an affine matrix of size (batch, 6). For 'warp', input data should be an

optical flow of size (batch, 2, h, w).

target.shape Shape(tuple), optional, default=[0,0] Specifies the output shape (H, W). This is

required if transformation type is 'affine'. If transformation type is 'warp', this

parameter is ignored.

mx.nd.GroupNorm

### Value

out The result mx.ndarray

mx.nd.GroupNorm

Group normalization.

# **Description**

The input channels are separated into "num\_groups" groups, each containing "num\_channels / num\_groups" channels. The mean and standard-deviation are calculated separately over the each group.

# **Arguments**

data NDArray-or-Symbol Input data

gamma NDArray-or-Symbol gamma array

beta NDArray-or-Symbol beta array

num.groups int, optional, default='1' Total number of groups.

eps float, optional, default=9.99999975e-06 An 'epsilon' parameter to prevent divi-

sion by 0.

output.mean.var

boolean, optional, default=0 Output the mean and std calculated along the given

axis.

### **Details**

.. math::

 $\label{eq:data} $$ data = data.reshape((N, num\_groups, C // num\_groups, ...)) out = \frac{a - mean(data, axis)}{qata - mean(data, axis)} $$ axis) + epsilon * gamma + beta $$ data - mean(data, axis) + me$ 

Both "gamma" and "beta" are learnable parameters.

Defined in src/operator/nn/group\_norm.cc:L77

## Value

mx.nd.identity

mx.nd.hard.sigmoid

Computes hard sigmoid of x element-wise.

# Description

```
.. math:: y = max(0, min(1, alpha * x + beta))
```

# Arguments

data NDArray-or-Symbol The input array.

alpha float, optional, default=0.200000003 Slope of hard sigmoid

beta float, optional, default=0.5 Bias of hard sigmoid.

#### **Details**

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L161

### Value

out The result mx.ndarray

mx.nd.identity

Returns a copy of the input.

# Description

From:src/operator/tensor/elemwise\_unary\_op\_basic.cc:244

# Arguments

data

NDArray-or-Symbol The input array.

#### Value

mx.nd.IdentityAttachKLSparseReg

Apply a sparse regularization to the output a sigmoid activation function.

### **Description**

Apply a sparse regularization to the output a sigmoid activation function.

# Arguments

data NDArray-or-Symbol Input data.

sparseness.target

float, optional, default=0.100000001 The sparseness target

penalty float, optional, default=0.00100000005 The tradeoff parameter for the sparse-

ness penalty

momentum float, optional, default=0.899999976 The momentum for running average

#### Value

out The result mx.ndarray

mx.nd.im2col	Extract sliding blocks from input array.

# Description

This operator is used in vanilla convolution implementation to transform the sliding blocks on image to column matrix, then the convolution operation can be computed by matrix multiplication between column and convolution weight. Due to the close relation between im2col and convolution, the concept of \*\*kernel\*\*, \*\*stride\*\*, \*\*dilate\*\* and \*\*pad\*\* in this operator are inherited from convolution operation.

## Arguments

data	NDArray-or-Symbol Input array to extract sliding blocks.	
kernel	Shape(tuple), required Sliding kernel size: (w,), (h, w) or (d, h, w).	
stride	Shape(tuple), optional, default=[] The stride between adjacent sliding blocks in spatial dimension: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.	
dilate	Shape(tuple), optional, default=[] The spacing between adjacent kernel points: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.	
pad	Shape(tuple), optional, default=[] The zero-value padding size on both sides of spatial dimension: (w,), (h, w) or (d, h, w). Defaults to no padding.	

140 mx.nd.InstanceNorm

#### **Details**

Given the input data of shape :math: '(N, C, \*)', where :math: 'N' is the batch size, :math: 'C' is the channel size, and :math: '\*' is the arbitrary spatial dimension, the output column array is always with shape :math: '(N, C \times \prod(\textkernel), W)', where :math: 'C \times \prod(\textkernel)' is the block size, and :math: 'W' is the block number which is the spatial size of the convolution output with same input parameters. Only 1-D, 2-D and 3-D of spatial dimension is supported in this operator.

Defined in src/operator/nn/im2col.cc:L100

### Value

out The result mx.ndarray

mx.nd.InstanceNorm

Applies instance normalization to the n-dimensional input array.

### **Description**

This operator takes an n-dimensional input array where (n>2) and normalizes the input using the following formula:

### **Arguments**

data	NDArray-or-Symbol An n-dimensional input array $(n > 2)$ of the form [batch, channel, spatial_dim1, spatial_dim2,].
gamma	NDArray-or-Symbol A vector of length 'channel', which multiplies the normalized input.
beta	NDArray-or-Symbol A vector of length 'channel', which is added to the product of the normalized input and the weight.
eps	float, optional, default= $0.00100000005$ An 'epsilon' parameter to prevent division by $0$ .

### Details

.. math::

out = \fracx - mean[data] \sqrtVar[data] + \epsilon \* gamma + beta

This layer is similar to batch normalization layer ('BatchNorm') with two differences: first, the normalization is carried out per example (instance), not over a batch. Second, the same normalization is applied both at test and train time. This operation is also known as 'contrast normalization'.

If the input data is of shape [batch, channel, spacial\_dim1, spacial\_dim2, ...], 'gamma' and 'beta' parameters must be vectors of shape [channel].

This implementation is based on this paper [1]\_

.. [1] Instance Normalization: The Missing Ingredient for Fast Stylization, D. Ulyanov, A. Vedaldi, V. Lempitsky, 2016 (arXiv:1607.08022v2).

mx.nd.khatri.rao 141

### Examples::

```
// Input of shape (2,1,2) x = [[[ 1.1, 2.2]], [[ 3.3, 4.4]]]
// gamma parameter of length 1 gamma = [1.5]
```

// beta parameter of length 1 beta = [0.5]

// Instance normalization is calculated with the above formula InstanceNorm(x,gamma,beta) = [[[-0.997527, 1.99752665]], [[-0.99752653, 1.99752724]]]

Defined in src/operator/instance\_norm.cc:L95

### Value

out The result mx.ndarray

mx.nd.khatri.rao

Computes the Khatri-Rao product of the input matrices.

## **Description**

Given a collection of :math: 'n' input matrices,

### **Arguments**

args

NDArray-or-Symbol[] Positional input matrices

### **Details**

```
.. math:: A_1 \in \mathbbR^M_1 \times M, ..., A_n \in \mathbbR^M_n \times N,
```

the (column-wise) Khatri-Rao product is defined as the matrix,

.. math::  $X = A_1 \cdot A_n \in A_$ 

where the :math: 'k' th column is equal to the column-wise outer product :math: ' $A_1_k$  \otimes \cdots \otimes  $A_n_k$ ' where :math: ' $A_i_k$ ' is the kth column of the ith matrix.

### Example::

```
»> A = mx.nd.array([[1, -1], »> [2, -3]]) »> B = mx.nd.array([[1, 4], »> [2, 5], »> [3, 6]]) »> C =
mx.nd.khatri_rao(A, B) »> print(C.asnumpy()) [[ 1. -4.] [ 2. -5.] [ 3. -6.] [ 2. -12.] [ 4. -15.] [ 6.
-18.]]
```

Defined in src/operator/contrib/krprod.cc:L108

### Value

142 mx.nd.L2Normalization

mx.nd.L2Normalization Normalize the input array using the L2 norm.

### **Description**

For 1-D NDArray, it computes::

### **Arguments**

data NDArray-or-Symbol Input array to normalize.

eps float, optional, default=1.00000001e-10 A small constant for numerical stability.

mode 'channel', 'instance', 'spatial',optional, default='instance' Specify the dimen-

sion along which to compute L2 norm.

#### **Details**

```
out = data / sqrt(sum(data ** 2) + eps)
```

For N-D NDArray, if the input array has shape (N, N, ..., N),

with "mode" = "instance", it normalizes each instance in the multidimensional array by its L2 norm.::

for i in 0...N out[i,:,:,...,:] = data[i,:,:,...,:] / sqrt(sum(data[i,:,:,...,:] \*\* 2) + eps)

with "mode" = "channel", it normalizes each channel in the array by its L2 norm.::

for i in 0...N out[:,i,:,...,:] = data[:,i,:,...,:] / sqrt(sum(data[:,i,:,...,:] \*\* 2) + eps)

with "mode" = "spatial", it normalizes the cross channel norm for each position in the array by its L2 norm.::

for dim in 2...N for i in 0...N out[....,i,...] = take(out, indices=i, axis=dim) / sqrt(sum(take(out, indices=i, axis=dim) \*\* 2) + eps) -dim-

### Example::

```
x = [[[1,2], [3,4]], [[2,2], [5,6]]]
```

L2Normalization(x, mode='instance') =[[[ 0.18257418 0.36514837] [ 0.54772252 0.73029673]] [[ 0.24077171 0.24077171] [ 0.60192931 0.72231513]]]

L2Normalization(x, mode='channel') =[[[ 0.31622776 0.44721359] [ 0.94868326 0.89442718]] [[ 0.37139067 0.31622776] [ 0.92847669 0.94868326]]]

 $L2Normalization(x, mode='spatial') = [[[ 0.44721359 \ 0.89442718] \ [ 0.60000002 \ 0.80000001]] \ [[ 0.70710677 \ 0.70710677] \ [ 0.6401844 \ 0.76822126]]]$ 

Defined in src/operator/l2\_normalization.cc:L196

### Value

mx.nd.lamb.update.phase1

Phase I of lamb update it performs the following operations and returns g:.

# **Description**

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

### **Arguments**

weight NDArray-or-Symbol Weight NDArray-or-Symbol Gradient grad NDArray-or-Symbol Moving mean mean NDArray-or-Symbol Moving variance var float, optional, default=0.899999976 The decay rate for the 1st moment estibeta1 mates. beta2 float, optional, default=0.999000013 The decay rate for the 2nd moment estifloat, optional, default=9.9999997e-07 A small constant for numerical stability. epsilon int, required Index update count. bias.correction boolean, optional, default=1 Whether to use bias correction. wd float, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight. float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad. rescale.grad clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient] If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

### **Details**

```
.. math::\begingather*\grad = \text{grad} * \text{rescale_grad} \text{if} (\text{grad} < -\text{clip_gradient}) \text{then grad} = -\text{clip_gradient} \text{if} (\text{grad} > \text{clip_gradient}) \text{then grad} = \text{clip_gradient} \text{mean} + (1 - \text{beta1}) * \text{grad}; \text{variance} = \text{beta2} * \text{variance} + (1. - \text{beta2}) * \text{grad} ^ 2; \text{if} (\text{bias_correction}) \text{then mean_hat} = \text{mean / (1. - \text{beta1^t}); \text{var_hat} = \text{var/(1 - \text{beta2^t}); \text{g} = \text{mean_hat} / (\text{var_hat^(1/2)} + \text{epsilon}) + \text{wd} * \text{weight;} \text{\text{endgather*}} \text{Defined in src/operator/optimizer_op.cc:L944}
```

clip\_gradient), -clip\_gradient).

### Value

144 mx.nd.LayerNorm

mx.nd.lamb.update.phase2

Phase II of lamb update it performs the following operations and updates grad.

# Description

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

# **Arguments**

weight	NDArray-or-Symbol Weight	
g	NDArray-or-Symbol Output of lamb_update_phase 1	
r1	NDArray-or-Symbol r1	
r2	NDArray-or-Symbol r2	
lr	float, required Learning rate	
lower.bound	float, optional, default=-1 Lower limit of norm of weight. If lower_bound <= 0, Lower limit is not set	
upper.bound	float, optional, default=-1 Upper limit of norm of weight. If upper_bound <= 0, Upper limit is not set	

#### **Details**

```
.. math:: \begingather* if (lower_bound >= 0) then r1 = max(r1, lower_bound) if (upper_bound >= 0) then r1 = max(r1, upper_bound) if (r1 == 0 \text{ or } r2 == 0) then lr = lr \text{ else } lr = lr * (r1/r2) \text{ weight} = \text{weight} - lr * g \endgather* Defined in src/operator/optimizer_op.cc:L983
```

# Value

out The result mx.ndarray

mx.nd.LayerNorm	Layer normalization.	
-----------------	----------------------	--

# Description

Normalizes the channels of the input tensor by mean and variance, and applies a scale "gamma" as well as offset "beta".

mx.nd.LeakyReLU 145

#### **Arguments**

data NDArray-or-Symbol Input data to layer normalization

gamma NDArray-or-Symbol gamma array beta NDArray-or-Symbol beta array

axis int, optional, default='-1' The axis to perform layer normalization. Usually, this

should be be axis of the channel dimension. Negative values means indexing

from right to left.

eps float, optional, default=9.99999975e-06 An 'epsilon' parameter to prevent divi-

sion by 0.

output.mean.var

boolean, optional, default=0 Output the mean and std calculated along the given

axis.

#### **Details**

Assume the input has more than one dimension and we normalize along axis 1. We first compute the mean and variance along this axis and then compute the normalized output, which has the same shape as input, as following:

.. math::

out = \fracdata - mean(data, axis)\sqrtvar(data, axis) + \epsilon \* gamma + beta

Both "gamma" and "beta" are learnable parameters.

Unlike BatchNorm and InstanceNorm, the \*mean\* and \*var\* are computed along the channel dimension.

Assume the input has size  $k^*$  on axis 1, then both "gamma" and "beta" have shape k,k. If "output\_mean\_var" is set to be true, then outputs both "data\_mean" and "data\_std". Note that no gradient will be passed through these two outputs.

The parameter "axis" specifies which axis of the input shape denotes the 'channel' (separately normalized groups). The default is -1, which sets the channel axis to be the last item in the input shape.

Defined in src/operator/nn/layer\_norm.cc:L158

### Value

out The result mx.ndarray

mx.nd.LeakyReLU

Applies Leaky rectified linear unit activation element-wise to the input.

#### **Description**

Leaky ReLUs attempt to fix the "dying ReLU" problem by allowing a small 'slope' when the input is negative and has a slope of one when input is positive.

146 mx.nd.linalg.det

## Arguments

data	NDArray-or-Symbol Input data to activation function.
gamma	NDArray-or-Symbol Input data to activation function.
act.type	'elu', 'gelu', 'leaky', 'prelu', 'rrelu', 'selu', optional, default='leaky' Activation function to be applied.
slope	float, optional, default=0.25 Init slope for the activation. (For leaky and elu only)
lower.bound	float, optional, default=0.125 Lower bound of random slope. (For rrelu only)
upper.bound	float, optional, default= $0.333999991$ Upper bound of random slope. (For rrelu only)

### **Details**

The following modified ReLU Activation functions are supported:

- \*elu\*: Exponential Linear Unit. 'y = x > 0 ? x : slope \* (exp(x)-1)' - \*selu\*: Scaled Exponential Linear Unit. 'y = lambda \* (x > 0 ? x : alpha \* (exp(x) - 1))' where \*lambda = 1.0507009873554804934193349852946\* and \*alpha = 1.6732632423543772848170429916717\*. - \*leaky\*: Leaky ReLU. 'y = x > 0 ? x : slope \* x' - \*prelu\*: Parametric ReLU. This is same as \*leaky\* except that 'slope' is learnt during training. - \*rrelu\*: Randomized ReLU. same as \*leaky\* but the 'slope' is uniformly and randomly chosen from \*[lower\_bound, upper\_bound)\* for training, while fixed to be \*(lower\_bound+upper\_bound)/2\* for inference.

Defined in src/operator/leaky\_relu.cc:L161

#### Value

out The result mx.ndarray

mx.nd.linalg.det	Compute the determinant of a matrix. Input is a tensor *A* of dimen-
mx.nu.linaig.uet	Compute the determinant of a matrix. Input is a tensor A · of atmen-
	sion *n >= 2*.

## **Description**

If \*n=2\*, \*A\* is a square matrix. We compute:

### **Arguments**

A NDArray-or-Symbol Tensor of square matrix

#### **Details**

```
*out* = *det(A)*
```

If \*n>2\*, \*det\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only. .. note:: There is no gradient backwarded when A is non-invertible (which is equivalent to det(A) = 0) because zero is rarely hit upon in float point computation and the Jacobi's formula on determinant gradient is not computationally efficient when A is non-invertible.

Examples::

Single matrix determinant A = [[1., 4.], [2., 3.]] det(A) = [-5.]

Batch matrix determinant A = [[[1., 4.], [2., 3.]], [[2., 3.], [1., 4.]]] det(A) = [-5., 5.]

Defined in src/operator/tensor/la\_op.cc:L975

### Value

out The result mx.ndarray

mx.nd.linalg.extractdiag

Extracts the diagonal entries of a square matrix. Input is a tensor  $A^*$  of dimension  $n \ge 2$ .

### Description

If \*n=2\*, then \*A\* represents a single square matrix which diagonal elements get extracted as a 1-dimensional tensor.

### **Arguments**

A NDArray-or-Symbol Tensor of square matrices

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

#### **Details**

If n>2, then A represents a batch of square matrices on the trailing two dimensions. The extracted diagonals are returned as an n-1-dimensional tensor.

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix diagonal extraction A = [[1.0, 2.0], [3.0, 4.0]]

extractdiag(A) = [1.0, 4.0]

extractdiag(A, 1) = [2.0]

Batch matrix diagonal extraction A = [[[1.0, 2.0], [3.0, 4.0]], [[5.0, 6.0], [7.0, 8.0]]]

```
extractdiag(A) = [[1.0, 4.0], [5.0, 8.0]]
Defined in src/operator/tensor/la_op.cc:L495
```

#### Value

out The result mx.ndarray

mx.nd.linalg.extracttrian

Extracts a triangular sub-matrix from a square matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

### **Description**

If \*n=2\*, then \*A\* represents a single square matrix from which a triangular sub-matrix is extracted as a 1-dimensional tensor.

## **Arguments**

A NDArray-or-Symbol Tensor of square matrices

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

lower boolean, optional, default=1 Refer to the lower triangular matrix if lower=true,

refer to the upper otherwise. Only relevant when offset=0

#### Details

If \*n>2\*, then \*A\* represents a batch of square matrices on the trailing two dimensions. The extracted triangular sub-matrices are returned as an \*n-1\*-dimensional tensor.

The \*offset\* and \*lower\* parameters determine the triangle to be extracted:

- When \*offset = 0\* either the lower or upper triangle with respect to the main diagonal is extracted depending on the value of parameter \*lower\*. When \*offset = k > 0\* the upper triangle with respect to the k-th diagonal above the main diagonal is extracted. When \*offset = k < 0\* the lower triangle with respect to the k-th diagonal below the main diagonal is extracted.
- .. note:: The operator supports float32 and float64 data types only.

#### Examples::

Single triagonal extraction A = [[1.0, 2.0], [3.0, 4.0]]

extracttrian(A) = [1.0, 3.0, 4.0] extracttrian(A, lower=False) = [1.0, 2.0, 4.0] extracttrian(A, 1) = [2.0] extracttrian(A, -1) = [3.0]

Batch triagonal extraction A = [[[1.0, 2.0], [3.0, 4.0]], [[5.0, 6.0], [7.0, 8.0]]]

extracttrian(A) = [[1.0, 3.0, 4.0], [5.0, 7.0, 8.0]]

Defined in src/operator/tensor/la\_op.cc:L605

mx.nd.linalg.gelqf 149

#### Value

out The result mx.ndarray

mx.nd.linalg.gelqf LQ factorization for general matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

# **Description**

If \*n=2\*, we compute the LQ factorization (LAPACK \*gelqf\*, followed by \*orglq\*). \*A\* must have shape \*(x, y)\* with \*x <= y\*, and must have full rank \*=x\*. The LQ factorization consists of \*L\* with shape \*(x, x)\* and \*Q\* with shape \*(x, y)\*, so that:

#### **Arguments**

A NDArray-or-Symbol Tensor of input matrices to be factorized

#### **Details**

```
*A* = *L* \ *Q*
```

Here, \*L\* is lower triangular (upper triangle equal to zero) with nonzero diagonal, and \*Q\* is row-orthonormal, meaning that

is equal to the identity matrix of shape \*(x, x)\*.

If \*n>2\*, \*gelqf\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

## Examples::

Single LQ factorization A = [[1., 2., 3.], [4., 5., 6.]] Q, L = gelqf(A) Q = [[-0.26726124, -0.53452248, -0.80178373], [0.87287156, 0.21821789, -0.43643578]] L = [[-3.74165739, 0.], [-8.55235974, 1.96396101]]

Batch LQ factorization A = [[[1, 2., 3.], [4., 5., 6.]], [[7., 8., 9.], [10., 11., 12.]]] Q, L = gelqf(A) Q = [[[-0.26726124, -0.53452248, -0.80178373], [0.87287156, 0.21821789, -0.43643578]], [[-0.50257071, -0.57436653, -0.64616234], [0.7620735, 0.05862104, -0.64483142]]] L = [[[-3.74165739, 0.], [-8.55235974, 1.96396101]], [[-13.92838828, 0.], [-19.09768702, 0.52758934]]]

Defined in src/operator/tensor/la\_op.cc:L798

## Value

150 mx.nd.linalg.gemm

mx.nd.linalg.gemm	Performs general matrix multiplication and accumulation. Input are tensors $*A*$ , $*B*$ , $*C*$ , each of dimension $*n >= 2*$ and having the same shape on the leading $*n-2*$ dimensions.

### **Description**

If \*n=2\*, the BLAS3 function \*gemm\* is performed:

# Arguments

Α	NDArray-or-Symbol Tensor of input matrices
В	NDArray-or-Symbol Tensor of input matrices
С	NDArray-or-Symbol Tensor of input matrices
transpose.a	boolean, optional, default=0 Multiply with transposed of first input (A).
transpose.b	boolean, optional, default=0 Multiply with transposed of second input (B).
alpha	double, optional, default=1 Scalar factor multiplied with A*B.
beta	double, optional, default=1 Scalar factor multiplied with C.
axis	int, optional, default='-2' Axis corresponding to the matrix rows.

#### **Details**

```
*out* = *alpha* \ **op* \ (*A*) \ **op* \ (*B*) + *beta* \ **C*
```

Here, \*alpha\* and \*beta\* are scalar parameters, and \*op()\* is either the identity or matrix transposition (depending on \*transpose\_a\*, \*transpose\_b\*).

If \*n>2\*, \*gemm\* is performed separately for a batch of matrices. The column indices of the matrices are given by the last dimensions of the tensors, the row indices by the axis specified with the \*axis\* parameter. By default, the trailing two dimensions will be used for matrix encoding.

For a non-default axis parameter, the operation performed is equivalent to a series of swapaxes/gemm/swapaxes calls. For example let \*A\*, \*B\*, \*C\* be 5 dimensional tensors. Then gemm(\*A\*, \*B\*, \*C\*, axis=1) is equivalent to the following without the overhead of the additional swapaxis operations::

```
A1 = swapaxes(A, dim1=1, dim2=3) B1 = swapaxes(B, dim1=1, dim2=3) C = swapaxes(C, dim1=1, dim2=3) C = gemm(A1, B1, C) C = swapaxis(C, dim1=1, dim2=3)
```

When the input data is of type float32 and the environment variables MXNET\_CUDA\_ALLOW\_TENSOR\_CORE and MXNET\_CUDA\_TENSOR\_OP\_MATH\_ALLOW\_CONVERSION are set to 1, this operator will try to use pseudo-float16 precision (float32 math with float16 I/O) precision in order to use Tensor Cores on suitable NVIDIA GPUs. This can sometimes give significant speedups.

.. note:: The operator supports float32 and float64 data types only.

### Examples::

```
Single matrix multiply-add A = [[1.0, 1.0], [1.0, 1.0]] B = [[1.0, 1.0], [1.0, 1.0], [1.0, 1.0]] C = [[1.0, 1.0, 1.0], [1.0, 1.0], [1.0, 1.0]] gemm(A, B, C, transpose_b=True, alpha=2.0, beta=10.0) = [[14.0, 14.0, 14.0], [14.0, 14.0, 14.0]]
```

mx.nd.linalg.gemm2 151

```
Batch matrix multiply-add A = [[[1.0, 1.0]], [[0.1, 0.1]]] B = [[[1.0, 1.0]], [[0.1, 0.1]]] C = [[[10.0]], [[0.01]]] gemm(A, B, C, transpose_b=True, alpha=2.0, beta=10.0) = [[[104.0]], [[0.14]]] Defined in src/operator/tensor/la_op.cc:L89
```

### Value

out The result mx.ndarray

mx.nd.linalg.gemm2	Performs general matrix multiplication. Input are tensors $*A*$ , $*B*$ , each of dimension $*n >= 2*$ and having the same shape on the leading $*n-2*$ dimensions.
--------------------	---

### Description

If \*n=2\*, the BLAS3 function \*gemm\* is performed:

#### Arguments

A	NDArray-or-Symbol Tensor of input matrices
В	NDArray-or-Symbol Tensor of input matrices
transpose.a	boolean, optional, default=0 Multiply with transposed of first input (A).
transpose.b	boolean, optional, default=0 Multiply with transposed of second input (B).
alpha	double, optional, default=1 Scalar factor multiplied with A*B.
axis	int, optional, default='-2' Axis corresponding to the matrix row indices.

#### **Details**

```
*out* = *alpha* \ **op* \ (*A*) \ **op* \ (*B*)
```

Here \*alpha\* is a scalar parameter and \*op()\* is either the identity or the matrix transposition (depending on \*transpose\_a\*, \*transpose\_b\*).

If \*n>2\*, \*gemm\* is performed separately for a batch of matrices. The column indices of the matrices are given by the last dimensions of the tensors, the row indices by the axis specified with the \*axis\* parameter. By default, the trailing two dimensions will be used for matrix encoding.

For a non-default axis parameter, the operation performed is equivalent to a series of swapaxes/gemm/swapaxes calls. For example let \*A\*, \*B\* be 5 dimensional tensors. Then gemm(\*A\*, \*B\*, axis=1) is equivalent to the following without the overhead of the additional swapaxis operations::

```
A1 = \text{swapaxes}(A, \text{dim1}=1, \text{dim2}=3) B1 = \text{swapaxes}(B, \text{dim1}=1, \text{dim2}=3) C = \text{gemm2}(A1, B1) C = \text{swapaxis}(C, \text{dim1}=1, \text{dim2}=3)
```

When the input data is of type float32 and the environment variables MXNET\_CUDA\_ALLOW\_TENSOR\_CORE and MXNET\_CUDA\_TENSOR\_OP\_MATH\_ALLOW\_CONVERSION are set to 1, this operator will try to use pseudo-float16 precision (float32 math with float16 I/O) precision in order to use Tensor Cores on suitable NVIDIA GPUs. This can sometimes give significant speedups.

.. note:: The operator supports float32 and float64 data types only.

152 mx.nd.linalg.inverse

## Examples::

Single matrix multiply  $A = [[1.0, 1.0], [1.0, 1.0]] B = [[1.0, 1.0], [1.0, 1.0], [1.0, 1.0]] gemm2(A, B, transpose_b=True, alpha=2.0) = [[4.0, 4.0, 4.0], [4.0, 4.0, 4.0]]$ 

Batch matrix multiply  $A = [[[1.0, 1.0]], [[0.1, 0.1]]] B = [[[1.0, 1.0]], [[0.1, 0.1]]] gemm2(A, B, transpose_b=True, alpha=<math>2.0$ ) = [[[4.0]], [[0.04]]]

Defined in src/operator/tensor/la\_op.cc:L163

#### Value

out The result mx.ndarray

```
mx.nd.linalg.inverse Compute the inverse of a matrix. Input is a tensor *A* of dimension *n >= 2*.
```

## **Description**

If \*n=2\*, \*A\* is a square matrix. We compute:

# **Arguments**

Α

NDArray-or-Symbol Tensor of square matrix

## **Details**

```
*out* = *A* \ :sup:'-1'
```

If \*n>2\*, \*inverse\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix inverse A = [[1., 4.], [2., 3.]] inverse(A) = [[-0.6, 0.8], [0.4, -0.2]]

Batch matrix inverse A = [[[1., 4.], [2., 3.]], [[1., 3.], [2., 4.]]] inverse(A) = [[[-0.6, 0.8], [0.4, -0.2]], [[-2., 1.5], [1., -0.5]]]

Defined in src/operator/tensor/la\_op.cc:L920

### Value

mx.nd.linalg.makediag Constructs a square matrix with the input as diagonal. Input is a tensor \*A\* of dimension \*n >= 1\*.

### **Description**

If \*n=1\*, then \*A\* represents the diagonal entries of a single square matrix. This matrix will be returned as a 2-dimensional tensor. If \*n>1\*, then \*A\* represents a batch of diagonals of square matrices. The batch of diagonal matrices will be returned as an \*n+1\*-dimensional tensor.

## **Arguments**

A NDArray-or-Symbol Tensor of diagonal entries

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

#### **Details**

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single diagonal matrix construction A = [1.0, 2.0]

makediag(A) = [[1.0, 0.0], [0.0, 2.0]]

makediag(A, 1) = [[0.0, 1.0, 0.0], [0.0, 0.0, 2.0], [0.0, 0.0, 0.0]]

Batch diagonal matrix construction A = [[1.0, 2.0], [3.0, 4.0]]

makediag(A) = [[[1.0, 0.0], [0.0, 2.0]], [[3.0, 0.0], [0.0, 4.0]]]

Defined in src/operator/tensor/la\_op.cc:L547

#### Value

out The result mx.ndarray

mx.nd.linalg.maketrian

Constructs a square matrix with the input representing a specific triangular sub-matrix. This is basically the inverse of \*linalg.extracttrian\*. Input is a tensor \*A\* of dimension \*n >= 1\*.

### **Description**

If \*n=1\*, then \*A\* represents the entries of a triangular matrix which is lower triangular if \*off-set<0\* or \*offset=0\*, \*lower=true\*. The resulting matrix is derived by first constructing the square matrix with the entries outside the triangle set to zero and then adding \*offset\*-times an additional diagonal with zero entries to the square matrix.

mx.nd.linalg.potrf

### Arguments

A NDArray-or-Symbol Tensor of triangular matrices stored as vectors

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

lower boolean, optional, default=1 Refer to the lower triangular matrix if lower=true,

refer to the upper otherwise. Only relevant when offset=0

#### **Details**

If \*n>1\*, then \*A\* represents a batch of triangular sub-matrices. The batch of corresponding square matrices is returned as an \*n+1\*-dimensional tensor.

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix construction A = [1.0, 2.0, 3.0]

maketrian(A) = [[1.0, 0.0], [2.0, 3.0]]

maketrian(A, lower=false) = [[1.0, 2.0], [0.0, 3.0]]

maketrian(A, offset=1) = [[0.0, 1.0, 2.0], [0.0, 0.0, 3.0], [0.0, 0.0, 0.0]] maketrian(A, offset=-1) = [[0.0, 0.0, 0.0], [1.0, 0.0, 0.0], [2.0, 3.0, 0.0]]

Batch matrix construction A = [[1.0, 2.0, 3.0], [4.0, 5.0, 6.0]]

maketrian(A) = [[[1.0, 0.0], [2.0, 3.0]], [[4.0, 0.0], [5.0, 6.0]]]

maketrian(A, offset=1) = [[[0.0, 1.0, 2.0], [0.0, 0.0, 3.0], [0.0, 0.0, 0.0]], [[0.0, 4.0, 5.0], [0.0, 0.0, 6.0], [0.0, 0.0, 0.0]]]

Defined in src/operator/tensor/la\_op.cc:L673

#### Value

out The result mx.ndarray

 ${\tt mx.nd.linalg.potrf}$ 

Performs Cholesky factorization of a symmetric positive-definite matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

## **Description**

If \*n=2\*, the Cholesky factor \*B\* of the symmetric, positive definite matrix \*A\* is computed. \*B\* is triangular (entries of upper or lower triangle are all zero), has positive diagonal entries, and:

## Arguments

A NDArray-or-Symbol Tensor of input matrices to be decomposed

mx.nd.linalg.potri 155

#### **Details**

```
*A* = *B* \ *B* \ :sup: `T` if *lower* = *true* *A* = *B* \ :sup: `T` \ *B* if *lower* = *false* If *n>2*, *potrf* is performed separately on the trailing two dimensions for all inputs (batch mode).
```

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix factorization A = [[4.0, 1.0], [1.0, 4.25]] potrf(A) = [[2.0, 0], [0.5, 2.0]]

Batch matrix factorization A = [[[4.0, 1.0], [1.0, 4.25]], [[16.0, 4.0], [4.0, 17.0]]] potrf(A) = [[[2.0, 0], [0.5, 2.0]], [[4.0, 0], [1.0, 4.0]]]

Defined in src/operator/tensor/la\_op.cc:L214

#### Value

out The result mx.ndarray

mx.nd.linalg.potri

Performs matrix inversion from a Cholesky factorization. Input is a tensor \*A\* of dimension \*n >= 2\*.

## **Description**

If \*n=2\*, \*A\* is a triangular matrix (entries of upper or lower triangle are all zero) with positive diagonal. We compute:

### **Arguments**

A

NDArray-or-Symbol Tensor of lower triangular matrices

#### **Details**

```
*out* = *A*\ :sup: `-T` \ *A*\ :sup: `-1` if *lower* = *true* *out* = *A*\ :sup: `-1` \ *A*\ :sup: `-T` if *lower* = *false*
```

In other words, if \*A\* is the Cholesky factor of a symmetric positive definite matrix \*B\* (obtained by \*potrf\*), then

```
*out* = *B*\ :sup:'-1'
```

If \*n>2\*, \*potri\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

.. note:: Use this operator only if you are certain you need the inverse of  ${}^*B^*$ , and cannot use the Cholesky factor  ${}^*A^*$  (\*potrf\*), together with backsubstitution (\*trsm\*). The latter is numerically much safer, and also cheaper.

#### Examples::

```
Single matrix inverse A = [[2.0, 0], [0.5, 2.0]] potri(A) = [[0.26563, -0.0625], [-0.0625, 0.25]]
Batch matrix inverse A = [[[2.0, 0], [0.5, 2.0]], [[4.0, 0], [1.0, 4.0]]] potri(A) = [[[0.26563, -0.0625], [-0.0625, 0.25]], [[0.06641, -0.01562], [-0.01562, 0.0625]]]
```

Defined in src/operator/tensor/la\_op.cc:L275

mx.nd.linalg.slogdet

### Value

out The result mx.ndarray

mx.nd.linalg.slogdet Compute the sign and log of the determinant of a matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

## **Description**

If \*n=2\*, \*A\* is a square matrix. We compute:

## Arguments

Α

NDArray-or-Symbol Tensor of square matrix

### **Details**

```
*sign* = *sign(det(A))* *logabsdet* = *log(abs(det(A)))*
```

If \*n>2\*, \*slogdet\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only. .. note:: The gradient is not properly defined on sign, so the gradient of it is not backwarded. .. note:: No gradient is backwarded when A is non-invertible. Please see the docs of operator det for detail.

# Examples::

Single matrix signed log determinant A = [[2., 3.], [1., 4.]] sign, logabsdet = slogdet(A) sign = [1.] logabsdet = [1.609438]

Batch matrix signed log determinant A = [[[2., 3.], [1., 4.]], [[1., 2.], [2., 4.]], [[1., 2.], [4., 3.]]] sign, logabsdet = slogdet(A) sign = [1., 0., -1.] logabsdet = [1.609438, -inf, 1.609438]

Defined in src/operator/tensor/la\_op.cc:L1034

# Value

mx.nd.linalg.sumlogdiag

Computes the sum of the logarithms of the diagonal elements of a square matrix. Input is a tensor \*A\* of dimension  $*n \ge 2*$ .

## Description

If \*n=2\*, \*A\* must be square with positive diagonal entries. We sum the natural logarithms of the diagonal elements, the result has shape (1,).

## **Arguments**

A NDArray-or-Symbol Tensor of square matrices

#### **Details**

If \*n>2\*, \*sumlogdiag\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix reduction A = [[1.0, 1.0], [1.0, 7.0]] sumlogdiag(A) = [1.9459]

Batch matrix reduction A = [[[1.0, 1.0], [1.0, 7.0]], [[3.0, 0], [0, 17.0]]] sumlogdiag(A) = [1.9459, 3.9318]

Defined in src/operator/tensor/la\_op.cc:L445

### Value

out The result mx.ndarray

mx.nd.linalg.syrk Multiplication of matrix with its transpose. Input is a tensor \*A\* of dimension \*n >= 2\*.

### **Description**

If \*n=2\*, the operator performs the BLAS3 function \*syrk\*:

#### **Arguments**

A NDArray-or-Symbol Tensor of input matrices

transpose boolean, optional, default=0 Use transpose of input matrix.

alpha double, optional, default=1 Scalar factor to be applied to the result.

158 mx.nd.linalg.trmm

## **Details**

```
*out* = *alpha* \* *A* \* *A* \: sup: `T` if *transpose=False*, or
```

if \*transpose=True\*.

If \*n>2\*, \*syrk\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

## Examples::

Single matrix multiply A = [[1., 2., 3.], [4., 5., 6.]] syrk(A, alpha=1., transpose=False) = [[14., 32.], [32., 77.]] syrk(A, alpha=1., transpose=True) = [[17., 22., 27.], [22., 29., 36.], [27., 36., 45.]]

Batch matrix multiply A = [[[1., 1.]], [[0.1, 0.1]]] syrk(A, alpha=2., transpose=False) = [[[4.]], [[0.04]]]

Defined in src/operator/tensor/la\_op.cc:L730

### Value

out The result mx.ndarray

mx.nd.linalg.trmm	Performs multiplication with a lower triangular matrix. Input are ten-
	sors $*A*$ , $*B*$ , each of dimension $*n \ge 2*$ and having the same shape
	on the leading *n-2* dimensions.

# Description

If \*n=2\*, \*A\* must be triangular. The operator performs the BLAS3 function \*trmm\*:

# Arguments

A	NDArray-or-Symbol Tensor of lower triangular matrices
В	NDArray-or-Symbol Tensor of matrices
transpose	boolean, optional, default=0 Use transposed of the triangular matrix
rightside	boolean, optional, default=0 Multiply triangular matrix from the right to non-triangular one.
lower	boolean, optional, default=1 True if the triangular matrix is lower triangular, false if it is upper triangular.
alpha	double, optional, default=1 Scalar factor to be applied to the result.

mx.nd.linalg.trsm 159

### **Details**

```
*out* = *alpha* \* *op*\ (*A*) \* *B*
if *rightside=False*, or
*out* = *alpha* \* *B* \* *op*\ (*A*)
```

if \*rightside=True\*. Here, \*alpha\* is a scalar parameter, and \*op()\* is either the identity or the matrix transposition (depending on \*transpose\*).

If \*n>2\*, \*trmm\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

## Examples::

Single triangular matrix multiply A = [[1.0, 0], [1.0, 1.0]] B = [[1.0, 1.0, 1.0], [1.0, 1.0]] trmm(A, B, alpha=2.0) = [[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]]

Batch triangular matrix multiply A = [[[1.0, 0], [1.0, 1.0]], [[1.0, 0], [1.0, 1.0]]] B = [[[1.0, 1.0, 1.0], [1.0, 1.0]], [[0.5, 0.5, 0.5], [0.5, 0.5, 0.5]]] trmm(A, B, alpha=2.0) = [[[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]], [[1.0, 1.0, 1.0], [2.0, 2.0, 2.0]]]

Defined in src/operator/tensor/la\_op.cc:L333

### Value

out The result mx.ndarray

mx.nd.linalg.trsm	Solves matrix equation involving a lower triangular matrix. Input are tensors $*A*$ , $*B*$ , each of dimension $*n >= 2*$ and having the same
	shape on the leading *n-2* dimensions.

# **Description**

If \*n=2\*, \*A\* must be triangular. The operator performs the BLAS3 function \*trsm\*, solving for \*out\* in:

### **Arguments**

A	NDArray-or-Symbol Tensor of lower triangular matrices
В	NDArray-or-Symbol Tensor of matrices
transpose	boolean, optional, default=0 Use transposed of the triangular matrix
rightside	boolean, optional, default=0 Multiply triangular matrix from the right to non-triangular one.
lower	boolean, optional, default=1 True if the triangular matrix is lower triangular, false if it is upper triangular.
alpha	double, optional, default=1 Scalar factor to be applied to the result.

#### **Details**

```
*op*\(*A*)\* *out* = *alpha* \* *B*
if *rightside=False*, or
*out* \* *op*\(*A*) = *alpha* \* *B*
```

if \*rightside=True\*. Here, \*alpha\* is a scalar parameter, and \*op()\* is either the identity or the matrix transposition (depending on \*transpose\*).

If \*n>2\*, \*trsm\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

#### Examples::

Single matrix solve A = [[1.0, 0], [1.0, 1.0]] B = [[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]] trsm(A, B, alpha=0.5) = [[1.0, 1.0, 1.0], [1.0, 1.0, 1.0]]

Batch matrix solve A = [[[1.0, 0], [1.0, 1.0]], [[1.0, 0], [1.0, 1.0]]] B = [[[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]], [[4.0, 4.0, 4.0], [8.0, 8.0, 8.0]]] trsm(A, B, alpha=0.5) = [[[1.0, 1.0, 1.0], [1.0, 1.0, 1.0]], [[2.0, 2.0, 2.0], [2.0, 2.0, 2.0]]]

Defined in src/operator/tensor/la\_op.cc:L396

#### Value

out The result mx.ndarray

mx.nd.LinearRegressionOutput

Computes and optimizes for squared loss during backward propagation. Just outputs "data" during forward propagation.

# Description

If :math: '\haty\_i' is the predicted value of the i-th sample, and :math: 'y\_i' is the corresponding target value, then the squared loss estimated over :math: 'n' samples is defined as

#### **Arguments**

data NDArray-or-Symbol Input data to the function.

1abel NDArray-or-Symbol Input label to the function.

grad.scale float, optional, default=1 Scale the gradient by a float factor

#### **Details**

 $: math: `\text{textSquaredLoss}(\text{textbfY}, \text{hat}\text{textbfY}) = \frac{n-1 \ensuremath{\lowert_2^{-1} \ensure$ 

.. note:: Use the LinearRegressionOutput as the final output layer of a net.

The storage type of "label" can be "default" or "csr"

mx.nd.load 161

 $- Linear Regression Output (default, \ default) = default - Linear Regression Output (default, \ csr) = default$ 

By default, gradients of this loss function are scaled by factor '1/m', where m is the number of regression outputs of a training example. The parameter 'grad\_scale' can be used to change this scale to 'grad\_scale/m'.

Defined in src/operator/regression\_output.cc:L92

### Value

out The result mx.ndarray

mx.nd.load

Load an mx.nd.array object on disk

## **Description**

Load an mx.nd.array object on disk

### Usage

```
mx.nd.load(filename)
```

## **Arguments**

filename

the filename (including the path)

### **Examples**

```
mat = mx.nd.array(1:3)
mx.nd.save(mat, 'temp.mat')
mat2 = mx.nd.load('temp.mat')
as.array(mat)
as.array(mat2)
```

mx.nd.log

Returns element-wise Natural logarithmic value of the input.

## **Description**

The natural logarithm is logarithm in base  $e^*$ , so that "log(exp(x)) = x"

### **Arguments**

data

NDArray-or-Symbol The input array.

162 mx.nd.log.softmax

### **Details**

The storage type of "log" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L77

#### Value

out The result mx.ndarray

mx.nd.log.softmax Computes the log softmax of the input. This is equivalent to computing

softmax followed by log.

# **Description**

Examples::

# **Arguments**

data NDArray-or-Symbol The input array.

axis int, optional, default='-1' The axis along which to compute softmax.

temperature double or None, optional, default=None Temperature parameter in softmax

dtype None, 'float16', 'float32', 'float64',optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to the same as input's dtype if not

defined (dtype=None).

use.length boolean or None, optional, default=0 Whether to use the length input as a mask

over the data input.

### **Details**

```
\gg x = mx.nd.array([1, 2, .1]) \gg mx.nd.log_softmax(x).asnumpy() array([-1.41702998, -0.41702995, -2.31702995], dtype=float32)
```

```
»> x = mx.nd.array( [[1, 2, .1],[.1, 2, 1]] ) »> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-0.34115392, -0.69314718, -1.24115396], [-1.24115396, -0.69314718, -0.34115392]], dtype=float32)
```

## Value

mx.nd.log10 163

mx.nd.log10

Returns element-wise Base-10 logarithmic value of the input.

## **Description**

```
10**log10(x) = x
```

# Arguments

data

NDArray-or-Symbol The input array.

### **Details**

The storage type of "log10" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L94

#### Value

out The result mx.ndarray

mx.nd.log1p

Returns element-wise "log(1 + x)" value of the input.

# **Description**

This function is more accurate than "log(1 + x)" for small "x" so that :math: '1+x\approx 1'

# Arguments

data

NDArray-or-Symbol The input array.

## **Details**

The storage type of "log1p" output depends upon the input storage type:

 $-\log 1p(\text{default}) = \text{default} - \log 1p(\text{row\_sparse}) = \text{row\_sparse} - \log 1p(\text{csr}) = \text{csr}$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L199

### Value

mx.nd.logical.not

mx.nd.log2

Returns element-wise Base-2 logarithmic value of the input.

# Description

$$"2**log2(x) = x"$$

# Arguments

data

NDArray-or-Symbol The input array.

## **Details**

The storage type of "log2" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L106

### Value

out The result mx.ndarray

mx.nd.logical.not

Returns the result of logical NOT (!) function

# Description

Example:  $logical_not([-2., 0., 1.]) = [0., 1., 0.]$ 

# Arguments

data

NDArray-or-Symbol The input array.

## Value

mx.nd.LogisticRegressionOutput

Applies a logistic function to the input.

### **Description**

The logistic function, also known as the sigmoid function, is computed as :math: '\frac11+exp(-\textbfx)'.

## **Arguments**

data NDArray-or-Symbol Input data to the function.

1abel NDArray-or-Symbol Input label to the function.

grad.scale float, optional, default=1 Scale the gradient by a float factor

#### **Details**

Commonly, the sigmoid is used to squash the real-valued output of a linear model :math:'wTx+b' into the [0,1] range so that it can be interpreted as a probability. It is suitable for binary classification or probability prediction tasks.

.. note:: Use the LogisticRegressionOutput as the final output layer of a net.

The storage type of "label" can be "default" or "csr"

- LogisticRegressionOutput(default, default) = default - LogisticRegressionOutput(default, csr) = default

The loss function used is the Binary Cross Entropy Loss:

```
:math: -(y\log(p) + (1 - y)\log(1 - p))
```

Where 'y' is the ground truth probability of positive outcome for a given example, and 'p' the probability predicted by the model. By default, gradients of this loss function are scaled by factor '1/m', where m is the number of regression outputs of a training example. The parameter 'grad\_scale' can be used to change this scale to 'grad\_scale/m'.

Defined in src/operator/regression\_output.cc:L152

#### Value

mx.nd.LRN	Applies local response normalization to the input.

### **Description**

The local response normalization layer performs "lateral inhibition" by normalizing over local input regions.

## **Arguments**

data	NDArray-or-Symbol Input data to LRN
alpha	float, optional, default=9.99999975e-05 The variance scaling parameter :math: '\alpha' in the LRN expression.
beta	float, optional, default=0.75 The power parameter :math: '\beta' in the LRN expression.
knorm	float, optional, default=2 The parameter :math: 'k' in the LRN expression.
nsize	int (non-negative), required normalization window width in elements.

#### **Details**

If :math: 'a\_x,y^i' is the activity of a neuron computed by applying kernel :math: 'i' at position :math: '(x, y)' and then applying the ReLU nonlinearity, the response-normalized activity :math: 'b\_x,y^i' is given by the expression:

```
.. math:: b_x,y^i = \fraca_x,y^i\Bigg(k + \frac\alphan \sum_j = max(0, i-\fracn2)^min(N-1, i+\fracn2) (a_x,y^j)^2\Bigg)^\beta
```

where the sum runs over :math: 'n' "adjacent" kernel maps at the same spatial position, and :math: 'N' is the total number of kernels in the layer.

Defined in src/operator/nn/lrn.cc:L158

#### Value

out The result mx.ndarray

 ${\tt mx.nd.MAERegressionOutput}$ 

Computes mean absolute error of the input.

## **Description**

MAE is a risk metric corresponding to the expected value of the absolute error.

mx.nd.make.loss 167

#### **Arguments**

data NDArray-or-Symbol Input data to the function.

1abel NDArray-or-Symbol Input label to the function.

grad. scale float, optional, default=1 Scale the gradient by a float factor

#### **Details**

If :math: '\haty\_i' is the predicted value of the i-th sample, and :math: 'y\_i' is the corresponding target value, then the mean absolute error (MAE) estimated over :math: 'n' samples is defined as :math: '\textMAE(\textbfY, \hat\textbfY) = \frac1n \sum\_i=0^n-1 \lVert \textbfy\_i - \hat\textbfy\_i \rVert 1'

.. note:: Use the MAERegressionOutput as the final output layer of a net.

The storage type of "label" can be "default" or "csr"

- MAERegressionOutput(default, default) = default - MAERegressionOutput(default, csr) = default By default, gradients of this loss function are scaled by factor '1/m', where m is the number of regression outputs of a training example. The parameter 'grad\_scale' can be used to change this scale to 'grad\_scale/m'.

Defined in src/operator/regression\_output.cc:L120

#### Value

out The result mx.ndarray

mx.nd.make.loss

Make your own loss function in network construction.

### **Description**

This operator accepts a customized loss function symbol as a terminal loss and the symbol should be an operator with no backward dependency. The output of this function is the gradient of loss with respect to the input data.

#### Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

For example, if you are a making a cross entropy loss function. Assume "out" is the predicted output and "label" is the true label, then the cross entropy can be defined as::

```
cross\_entropy = label * log(out) + (1 - label) * log(1 - out) loss = make\_loss(cross\_entropy)
```

We will need to use "make\_loss" when we are creating our own loss function or we want to combine multiple loss functions. Also we may want to stop some variables' gradients from backpropagation. See more detail in "BlockGrad" or "stop\_gradient".

168 mx.nd.MakeLoss

The storage type of "make\_loss" output depends upon the input storage type:

- make\_loss(default) = default - make\_loss(row\_sparse) = row\_sparse

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L359

#### Value

out The result mx.ndarray

mx.nd.MakeLoss

Make your own loss function in network construction.

### **Description**

This operator accepts a customized loss function symbol as a terminal loss and the symbol should be an operator with no backward dependency. The output of this function is the gradient of loss with respect to the input data.

#### **Arguments**

data NDArray-or-Symbol Input array.

grad.scale float, optional, default=1 Gradient scale as a supplement to unary and binary

operators

valid.thresh float, optional, default=0 clip each element in the array to 0 when it is less than

"valid\_thresh". This is used when "normalization" is set to "'valid'".

normalization 'batch', 'null', 'valid',optional, default='null' If this is set to null, the output

gradient will not be normalized. If this is set to batch, the output gradient will be divided by the batch size. If this is set to valid, the output gradient will be

divided by the number of valid input elements.

### **Details**

For example, if you are a making a cross entropy loss function. Assume "out" is the predicted output and "label" is the true label, then the cross entropy can be defined as::

cross entropy = label \* log(out) + (1 - label) \* log(1 - out) loss = MakeLoss(cross entropy)

We will need to use "MakeLoss" when we are creating our own loss function or we want to combine multiple loss functions. Also we may want to stop some variables' gradients from backpropagation. See more detail in "BlockGrad" or "stop\_gradient".

In addition, we can give a scale to the loss by setting "grad\_scale", so that the gradient of the loss will be rescaled in the backpropagation.

.. note:: This operator should be used as a Symbol instead of NDArray.

Defined in src/operator/make\_loss.cc:L71

## Value

mx.nd.max 169

mx.nd.max	Computes the max of array elements over given axes.	

## **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L32

## **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

## Value

out The result mx.ndarray

mx.nd.max.axis	Computes the max of array elements over given axes.
----------------	---

## **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L32

### **Arguments**

data NDArray-or-Symbol The input

170 mx.nd.mean

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

#### Value

out The result mx.ndarray

mx.nd.mean Computes the mean of array elements over given axes.

### **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L84

# Arguments

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

#### Value

mx.nd.min 171

mx.nd.min	Computes the min of array elements over given axes.	

## **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L47

## Arguments

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

# Value

out The result mx.ndarray

my nd min avic	L'ammutag th	a min of array	alamante avar avian avac
mx.nd.min.axis	Computes in	e min oi arrav	elements over given axes.
		j	

### **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L47

### **Arguments**

data NDArray-or-Symbol The input

172 mx.nd.moments

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

#### Value

out The result mx.ndarray

mx.nd.moments Calculate the mean and variance of 'data'.

### **Description**

The mean and variance are calculated by aggregating the contents of data across axes. If x is 1-D and axes = [0] this is just the mean and variance of a vector.

### Arguments

data NDArray-or-Symbol Input ndarray

axes Shape or None, optional, default=None Array of ints. Axes along which to

compute mean and variance.

keepdims boolean, optional, default=0 produce moments with the same dimensionality as

the input.

### Details

#### Example:

x = [[1, 2, 3], [4, 5, 6]] mean, var = moments(data=x, axes=[0]) mean = [2.5, 3.5, 4.5] var = [2.25, 2.25, 2.25] mean, var = moments(data=x, axes=[1]) mean = [2.0, 5.0] var = [0.66666667] mean, var = moments(data=x, axis=[0, 1]) mean = [3.5] var = [2.9166667]

Defined in src/operator/nn/moments.cc:L54

# Value

```
mx.nd.mp.lamb.update.phase1
```

Mixed Precision version of Phase I of lamb update it performs the following operations and returns g:.

### **Description**

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

#### **Arguments**

weight NDArray-or-Symbol Weight
grad NDArray-or-Symbol Gradient
mean NDArray-or-Symbol Moving mean
var NDArray-or-Symbol Moving variance
weight32 NDArray-or-Symbol Weight32

beta1 float, optional, default=0.899999976 The decay rate for the 1st moment esti-

mates.

beta2 float, optional, default=0.999000013 The decay rate for the 2nd moment esti-

mates.

epsilon float, optional, default=9.99999997e-07 A small constant for numerical stability.

t int, required Index update count.

bias.correction

boolean, optional, default=1 Whether to use bias correction.

wd float, required Weight decay augments the objective function with a regulariza-

tion term that penalizes large weights. The penalty scales with the square of the

magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip\_gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

#### **Details**

```
.. math:: \begingather* grad32 = grad(float16) * rescale_grad if (grad < -clip_gradient) then grad = -clip_gradient if (grad > clip_gradient) then grad = clip_gradient mean = beta1 * mean + (1 - beta1) * grad; variance = beta2 * variance + (1. - beta2) * grad ^ 2; if (bias_correction) then mean_hat = mean / (1. - beta1^t); var_hat = var / (1 - beta2^t); g = mean_hat / (var_hat^(1/2) + epsilon) + wd * weight32; else g = mean / (var_data^(1/2) + epsilon) + wd * weight32; \endgather*
```

Defined in src/operator/optimizer\_op.cc:L1024

#### Value

```
mx.nd.mp.lamb.update.phase2
```

Mixed Precision version Phase II of lamb update it performs the following operations and updates grad.

# Description

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

# Arguments

weight	NDArray-or-Symbol Weight
g	NDArray-or-Symbol Output of mp_lamb_update_phase 1
r1	NDArray-or-Symbol r1
r2	NDArray-or-Symbol r2
weight32	NDArray-or-Symbol Weight32
lr	float, required Learning rate
lower.bound	float, optional, default=-1 Lower limit of norm of weight. If lower_bound <= 0, Lower limit is not set
upper.bound	float, optional, default=-1 Upper limit of norm of weight. If upper_bound <= 0, Upper limit is not set

# **Details**

```
.. math:: \begingather* if (lower_bound >= 0) then r1 = max(r1, lower_bound) if (upper_bound >= 0) then r1 = max(r1, upper_bound) if (r1 == 0 \text{ or } r2 == 0) then lr = lr \text{ else } lr = lr * (r1/r2) \text{ weight32} = \text{weight32} - lr * g \text{ weight(float16)} = \text{weight32} \setminus \frac{1}{r} Defined in src/operator/optimizer_op.cc:L1066
```

### Value

mx.nd.mp.nag.mom.update

*Update function for multi-precision Nesterov Accelerated Gradient(NAG) optimizer.* 

# Description

Defined in src/operator/optimizer\_op.cc:L736

## **Arguments**

weight NDArray-or-Symbol Weight NDArray-or-Symbol Gradient grad NDArray-or-Symbol Momentum mom weight32 NDArray-or-Symbol Weight32 lr float, required Learning rate momentum float, optional, default=0 The decay rate of momentum estimates at each epoch. float, optional, default=0 Weight decay augments the objective function with a wd regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight. float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad. rescale.grad clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient] If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip\_gradient), -clip\_gradient).

### Value

out The result mx.ndarray

mx.nd.mp.sgd.mom.update

Updater function for multi-precision sgd optimizer

### **Description**

Updater function for multi-precision sgd optimizer

176 mx.nd.mp.sgd.update

#### **Arguments**

weight NDArray-or-Symbol Weight
grad NDArray-or-Symbol Gradient
mom NDArray-or-Symbol Momentum
weight32 NDArray-or-Symbol Weight32
lr float, required Learning rate

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row\_sparse and both weight and momentum have the same stype

### Value

out The result mx.ndarray

# **Description**

Updater function for multi-precision sgd optimizer

### **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol gradient weight32 NDArray-or-Symbol Weight32 1r float, required Learning rate

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row\_sparse.

mx.nd.multi.all.finite

# Value

out The result mx.ndarray

```
mx.nd.multi.all.finite
```

Check if all the float numbers in all the arrays are finite (used for AMP)

## **Description**

Defined in src/operator/contrib/all\_finite.cc:L133

## **Arguments**

data NDArray-or-Symbol[] Arrays

num.arrays int, optional, default='1' Number of arrays.

init.output boolean, optional, default=1 Initialize output to 1.

#### Value

out The result mx.ndarray

their sums of square"

# Description

Defined in src/operator/contrib/multi\_lars.cc:L37

## **Arguments**

1rs NDArray-or-Symbol Learning rates to scale by LARS coefficient

weights.sum.sq NDArray-or-Symbol sum of square of weights arrays grads.sum.sq NDArray-or-Symbol sum of square of gradients arrays

wds NDArray-or-Symbol weight decays

eta float, required LARS eta eps float, required LARS eps

rescale.grad float, optional, default=1 Gradient rescaling factor

#### Value

mx.nd.multi.mp.sgd.mom.update

Momentum update function for multi-precision Stochastic Gradient Descent (SGD) optimizer.

# Description

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

## **Arguments**

data	NDArray-or-Symbol[] Weights
lrs	tuple of <float>, required Learning rates.</float>
wds	tuple of <float>, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.</float>
momentum	float, optional, default=0 The decay rate of momentum estimates at each epoch.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
num.weights	int, optional, default='1' Number of updated weights.

### **Details**

```
.. math:: v_1 = \alpha pha * \beta J(W_0) v_t = \gamma v_{-1} - \alpha pha * \beta J(W_{-1}) W_t = W_{-1} + v_t It updates the weights using:: v = momentum * v - learning\_rate * gradient weight += v Where the parameter "momentum" is the decay rate of momentum estimates at each epoch. Defined in src/operator/optimizer_op.cc:L463
```

### Value

mx.nd.multi.mp.sgd.update

Update function for multi-precision Stochastic Gradient Descent (SDG) optimizer.

# Description

It updates the weights using::

# Arguments

data	NDArray-or-Symbol[] Weights
lrs	tuple of <float>, required Learning rates.</float>
wds	tuple of <float>, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.</float>
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
num.weights	int, optional, default='1' Number of updated weights.

#### **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
Defined in src/operator/optimizer_op.cc:L408
```

#### Value

out The result mx.ndarray

```
mx.nd.multi.sgd.mom.update
```

Momentum update function for Stochastic Gradient Descent (SGD) optimizer.

# Description

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

#### **Arguments**

data NDArray-or-Symbol[] Weights, gradients and momentum

1rs tuple of <float>, required Learning rates.

wds tuple of <float>, required Weight decay augments the objective function with

a regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip\_gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

#### **Details**

.. math::

 $v_1 = \alpha * \Lambda J(W_0) \ v_t = \gamma v_1 - \alpha * \Lambda J(W_{t-1}) \ W_t = W_{t-1} + v_t$ 

It updates the weights using::

v = momentum \* v - learning\_rate \* gradient weight += v

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

Defined in src/operator/optimizer\_op.cc:L365

#### Value

out The result mx.ndarray

mx.nd.multi.sgd.update

Update function for Stochastic Gradient Descent (SDG) optimizer.

### **Description**

It updates the weights using::

## **Arguments**

data NDArray-or-Symbol[] Weights

1rs tuple of <float>, required Learning rates.

wds tuple of <float>, required Weight decay augments the objective function with

a regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

mx.nd.multi.sum.sq 181

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

#### **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
Defined in src/operator/optimizer_op.cc:L320
```

### Value

out The result mx.ndarray

mx.nd.multi.sum.sq

Compute the sums of squares of multiple arrays

# **Description**

Defined in src/operator/contrib/multi\_sum\_sq.cc:L36

## Arguments

data NDArray-or-Symbol[] Arrays

num.arrays int, required number of input arrays.

### Value

out The result mx.ndarray

 $\verb|mx.nd.nag.mom.update| \\$ 

Update function for Nesterov Accelerated Gradient( NAG) optimizer. It updates the weights using the following formula,

## **Description**

```
.. math:: v_t = \gamma v_{t-1} + \epsilon * \Lambda J(W_{t-1} - \gamma v_{t-1}) W_t = W_{t-1} - v_t
```

182 mx.nd.nanprod

#### **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient mom NDArray-or-Symbol Momentum 1r float, required Learning rate

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch. wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

#### **Details**

Where :math: '\eta' is the learning rate of the optimizer :math: '\gamma' is the decay rate of the momentum estimate :math: '\v\_t' is the update vector at time step 't' :math: '\W\_t' is the weight vector at time step 't'

Defined in src/operator/optimizer\_op.cc:L717

#### Value

out The result mx.ndarray

mx.nd.nanprod Computes the product of array elements over given axes treating Not a Numbers ("NaN") as one.

### **Description**

Computes the product of array elements over given axes treating Not a Numbers ("NaN") as one.

### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

mx.nd.nansum 183

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

## **Details**

Defined in src/operator/tensor/broadcast\_reduce\_prod\_value.cc:L47

## Value

out The result mx.ndarray

mx.nd.nansum Computes the sum of array elements over given axes treating Not a

Numbers ("NaN") as zero.

## **Description**

Computes the sum of array elements over given axes treating Not a Numbers ("NaN") as zero.

## **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

### **Details**

Defined in src/operator/tensor/broadcast\_reduce\_sum\_value.cc:L102

### Value

184 mx.nd.norm

mx.nd.negative Numerical negative of the argument, element-wise.
--

# Description

The storage type of "negative" output depends upon the input storage type:

# **Arguments**

data NDArray-or-Symbol The input array.

## **Details**

- negative(default) = default - negative(row\_sparse) = row\_sparse - negative(csr) = csr

### Value

out The result mx.ndarray

mx.nd.norm Computes the norm on an NDArray.
---

# Description

This operator computes the norm on an NDArray with the specified axis, depending on the value of the ord parameter. By default, it computes the L2 norm on the entire array. Currently only ord=2 supports sparse ndarrays.

## **Arguments**

data	NDArray-or-Symbol The input
ord	int, optional, default='2' Order of the norm. Currently ord=1 and ord=2 is supported.
axis	Shape or None, optional, default=None The axis or axes along which to perform the reduction. The default, 'axis=()', will compute over all elements into a scalar array with shape '(1,)'. If 'axis' is int, a reduction is performed on a particular axis. If 'axis' is a 2-tuple, it specifies the axes that hold 2-D matrices, and the matrix norms of these matrices are computed.
out.dtype	None, 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', optional, default='None' The data type of the output.
keepdims	boolean, optional, default=0 If this is set to 'True', the reduced axis is left in the result as dimension with size one.

mx.nd.normal 185

## **Details**

```
Examples::

x = [[[1, 2], [3, 4]], [[2, 2], [5, 6]]]

norm(x, ord=2, axis=1) = [[3.1622777 4.472136] [5.3851647 6.3245554]]

norm(x, ord=1, axis=1) = [[4., 6.], [7., 8.]]

rsp = x.cast_storage('row_sparse')

norm(rsp) = [5.47722578]

csr = x.cast_storage('csr')

norm(csr) = [5.47722578]

Defined in src/operator/tensor/broadcast_reduce_norm_value.cc:L89
```

### Value

out The result mx.ndarray

mx.nd.normal

Draw random samples from a normal (Gaussian) distribution.

### **Description**

.. note:: The existing alias "normal" is deprecated.

## **Arguments**

loc	float, optional, default=0 Mean of the distribution.
scale	float, optional, default=1 Standard deviation of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

### **Details**

Samples are distributed according to a normal distribution parametrized by \*loc\* (mean) and \*scale\* (standard deviation).

Example::

```
normal(loc=0, scale=1, shape=(2,2)) = [[ 1.89171135, -1.16881478], [-1.23474145, 1.55807114]] Defined in src/operator/random/sample_op.cc:L113
```

## Value

mx.nd.one.hot

mx.nd.one.hot	Returns a one-hot array.
---------------	--------------------------

## **Description**

The locations represented by 'indices' take value 'on\_value', while all other locations take value 'off\_value'.

## **Arguments**

indices	NDArray-or-Symbol array of locations where to set on_value
depth	int, required Depth of the one hot dimension.
on.value	double, optional, default=1 The value assigned to the locations represented by indices.
off.value	double, optional, default=0 The value assigned to the locations not represented by indices.
dtype	'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional, default='float32' DType of the output

# **Details**

## Value

out The result mx.ndarray

Defined in src/operator/tensor/indexing\_op.cc:L888

mx.nd.ones 187

mx.nd.ones

Generate an mx.ndarray object with ones

# Description

Generate an mx.ndarray object with ones

## Usage

```
mx.nd.ones(shape, ctx = NULL)
```

# Arguments

shape the dimension of the mx.ndarray

ctx optional The context device of the array. mx.ctx.default() will be used in default.

## **Examples**

```
mat = mx.nd.ones(10)
as.array(mat)
mat2 = mx.nd.ones(c(5,5))
as.array(mat)
mat3 = mx.nd.ones(c(3,3,3))
as.array(mat3)
```

mx.nd.ones.like

Return an array of ones with the same shape and type as the input array.

# Description

Examples::

## **Arguments**

data

NDArray-or-Symbol The input

# **Details**

```
x = [[ 0., 0., 0.], [ 0., 0., 0.]]
ones_like(x) = [[ 1., 1., 1.], [ 1., 1., 1.]]
```

### Value

mx.nd.Pad

	mx.nd.Pad	Pads an input array with a constant or edge values of the array.
--	-----------	--

### **Description**

.. note:: 'Pad' is deprecated. Use 'pad' instead.

### Arguments

data NDArray-or-Symbol An n-dimensional input array.

mode 'constant', 'edge', 'reflect', required Padding type to use. "constant" pads with

'constant\_value' "edge" pads using the edge values of the input array "reflect"

pads by reflecting values with respect to the edges.

pad.width Shape(tuple), required Widths of the padding regions applied to the edges of

each axis. It is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". It should be of length "2\*N" where "N" is the number of dimensions of the array. This is equivalent to pad\_width in

numpy.pad, but flattened.

constant.value double, optional, default=0 The value used for padding when 'mode' is "con-

stant".

### **Details**

.. note:: Current implementation only supports 4D and 5D input arrays with padding applied only on axes 1, 2 and 3. Expects axes 4 and 5 in 'pad\_width' to be zero.

This operation pads an input array with either a 'constant\_value' or edge values along each axis of the input array. The amount of padding is specified by 'pad\_width'.

'pad\_width' is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". The 'pad\_width' should be of length "2\*N" where "N" is the number of dimensions of the array.

For dimension "N" of the input array, "before\_N" and "after\_N" indicates how many values to add before and after the elements of the array along dimension "N". The widths of the higher two dimensions "before\_1", "after\_1", "before\_2", "after\_2" must be 0.

# Example::

```
x = [[[[ 1. 2. 3.] [ 4. 5. 6.]]

[[ 7. 8. 9.] [ 10. 11. 12.]]]

[[[ 11. 12. 13.] [ 14. 15. 16.]]

[[ 17. 18. 19.] [ 20. 21. 22.]]]]

pad(x,mode="edge", pad_width=(0,0,0,0,1,1,1,1)) =

[[[[ 1. 1. 2. 3. 3.] [ 1. 1. 2. 3. 3.] [ 4. 4. 5. 6. 6.] [ 4. 4. 5. 6. 6.]]

[[ 7. 7. 8. 9. 9.] [ 7. 7. 8. 9. 9.] [ 10. 10. 11. 12. 12.] [ 10. 10. 11. 12. 12.]]]

[[[ 11. 11. 12. 13. 13.] [ 11. 11. 12. 13. 13.] [ 14. 14. 15. 16. 16.] [ 14. 14. 15. 16. 16.]]
```

mx.nd.pad 189

```
[[ 17. 17. 18. 19. 19.] [ 17. 17. 18. 19. 19.] [ 20. 20. 21. 22. 22.] [ 20. 20. 21. 22. 22.]]]]

pad(x, mode="constant", constant_value=0, pad_width=(0,0,0,0,1,1,1,1)) =

[[[[ 0. 0. 0. 0. 0. 0.] [ 0. 1. 2. 3. 0.] [ 0. 4. 5. 6. 0.] [ 0. 0. 0. 0. 0.]]

[[ 0. 0. 0. 0. 0.] [ 0. 7. 8. 9. 0.] [ 0. 10. 11. 12. 0.] [ 0. 0. 0. 0. 0.]]]

[[[ 0. 0. 0. 0. 0.] [ 0. 11. 12. 13. 0.] [ 0. 14. 15. 16. 0.] [ 0. 0. 0. 0. 0.]]

[[ 0. 0. 0. 0. 0.] [ 0. 17. 18. 19. 0.] [ 0. 20. 21. 22. 0.] [ 0. 0. 0. 0. 0.]]]]

Defined in src/operator/pad.cc:L766
```

#### Value

out The result mx.ndarray

mx.nd.pad

Pads an input array with a constant or edge values of the array.

### Description

.. note:: 'Pad' is deprecated. Use 'pad' instead.

### **Arguments**

data NDArray-or-Symbol An n-dimensional input array.

mode 'constant', 'edge', 'reflect', required Padding type to use. "constant" pads with

'constant\_value' "edge" pads using the edge values of the input array "reflect"

pads by reflecting values with respect to the edges.

pad.width Shape(tuple), required Widths of the padding regions applied to the edges of

each axis. It is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". It should be of length "2\*N" where "N" is the number of dimensions of the array. This is equivalent to pad\_width in

numpy.pad, but flattened.

constant.value double, optional, default=0 The value used for padding when 'mode' is "con-

stant".

### **Details**

.. note:: Current implementation only supports 4D and 5D input arrays with padding applied only on axes 1, 2 and 3. Expects axes 4 and 5 in 'pad\_width' to be zero.

This operation pads an input array with either a 'constant\_value' or edge values along each axis of the input array. The amount of padding is specified by 'pad\_width'.

'pad\_width' is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". The 'pad\_width' should be of length "2\*N" where "N" is the number of dimensions of the array.

190 mx.nd.pick

For dimension "N" of the input array, "before\_N" and "after\_N" indicates how many values to add before and after the elements of the array along dimension "N". The widths of the higher two dimensions "before\_1", "after\_1", "before\_2", "after\_2" must be 0.

### Example::

```
x = [[[[ 1. 2. 3.] [ 4. 5. 6.]]
[[ 7. 8. 9.] [ 10. 11. 12.]]]
[[[ 11. 12. 13.] [ 14. 15. 16.]]
[[ 17. 18. 19.] [ 20. 21. 22.]]]]
pad(x,mode="edge", pad_width=(0,0,0,0,1,1,1,1)) =
[[[ 1. 1. 2. 3. 3.] [ 1. 1. 2. 3. 3.] [ 4. 4. 5. 6. 6.] [ 4. 4. 5. 6. 6.]]
[[ 7. 7. 8. 9. 9.] [ 7. 7. 8. 9. 9.] [ 10. 10. 11. 12. 12.] [ 10. 10. 11. 12. 12.]]]
[[[ 11. 11. 12. 13. 13.] [ 11. 11. 12. 13. 13.] [ 14. 14. 15. 16. 16.] [ 14. 14. 15. 16. 16.]]
[[ 17. 17. 18. 19. 19.] [ 17. 17. 18. 19. 19.] [ 20. 20. 21. 22. 22.] [ 20. 20. 21. 22. 22.]]]]
pad(x, mode="constant", constant_value=0, pad_width=(0,0,0,0,1,1,1,1)) =
[[[ 0. 0. 0. 0. 0.] [ 0. 1. 2. 3. 0.] [ 0. 4. 5. 6. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 11. 12. 13. 0.] [ 0. 14. 15. 16. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 11. 12. 13. 0.] [ 0. 14. 15. 16. 0.] [ 0. 0. 0. 0. 0.]]]
[[ 0. 0. 0. 0. 0.] [ 0. 17. 18. 19. 0.] [ 0. 20. 21. 22. 0.] [ 0. 0. 0. 0. 0.]]]]
Defined in src/operator/pad.cc:L766
```

## Value

out The result mx.ndarray

mx.nd.pick	Picks elements from an input array according to the input indices along the given axis.

## **Description**

Given an input array of shape "(d0, d1)" and indices of shape "(i0,)", the result will be an output array of shape "(i0,)" with::

## Arguments

data	NDArray-or-Symbol The input array
index	NDArray-or-Symbol The index array
axis	int or None, optional, default='-1' int or None. The axis to picking the elements.
	Negative values means indexing from right to left. If is 'None', the elements in
	the index w.r.t the flattened input will be picked.

mx.nd.Pooling

keepdims boolean, optional, default=0 If true, the axis where we pick the elements is left

in the result as dimension with size one.

mode 'clip', 'wrap',optional, default='clip' Specify how out-of-bound indices behave.

Default is "clip". "clip" means clip to the range. So, if all indices mentioned are too large, they are replaced by the index that addresses the last element along an

axis. "wrap" means to wrap around.

#### **Details**

output[i] = input[i, indices[i]]

By default, if any index mentioned is too large, it is replaced by the index that addresses the last element along an axis (the 'clip' mode).

This function supports n-dimensional input and (n-1)-dimensional indices arrays.

Examples::

```
x = [[1., 2.], [3., 4.], [5., 6.]]
```

// picks elements with specified indices along axis 0 pick(x, y=[0,1], 0) = [1., 4.]

// picks elements with specified indices along axis 1 pick(x, y=[0,1,0], 1) = [1., 4., 5.]

// picks elements with specified indices along axis 1 using 'wrap' mode // to place indicies that would normally be out of bounds pick(x, y=[2,-1,-2], 1, mode='wrap') = [1,,4,,5]

$$y = [[1.], [0.], [2.]]$$

// picks elements with specified indices along axis 1 and dims are maintained pick(x, y, 1, keep-dims=True) = [[2.], [3.], [6.]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L151

#### Value

out The result mx.ndarray

mx.nd.Pooling	Performs pooling on the input.	
---------------	--------------------------------	--

### **Description**

The shapes for 1-D pooling are

### **Arguments**

data	NDArray-or-Syn	ibol Input data to t	he pooling operator.
------	----------------	----------------------	----------------------

kernel Shape(tuple), optional, default=[] Pooling kernel size: (y, x) or (d, y, x)

pool.type 'avg', 'lp', 'max', 'sum',optional, default='max' Pooling type to be applied.

global.pool boolean, optional, default=0 Ignore kernel size, do global pooling based on cur-

rent input feature map.

mx.nd.Pooling

cudnn.off boolean, optional, default=0 Turn off cudnn pooling and use MXNet pooling

operator.

pooling.convention

'full', 'same', 'valid', optional, default='valid' Pooling convention to be applied.

stride Shape(tuple), optional, default=[] Stride: for pooling (y, x) or (d, y, x). Defaults

to 1 for each dimension.

pad Shape(tuple), optional, default=[] Pad for pooling: (y, x) or (d, y, x). Defaults to

no padding.

p. value int or None, optional, default='None' Value of p for Lp pooling, can be 1 or 2,

required for Lp Pooling.

count.include.pad

boolean or None, optional, default=None Only used for AvgPool, specify whether to count padding elements for averagecalculation. For example, with a 5\*5 kernel on a 3\*3 corner of a image, the sum of the 9 valid elements will be divided by 25 if this is set to true, or it will be divided by 9 if this is set to false. Defaults

to true.

layout None, 'NCDHW', 'NCHW', 'NCW', 'NDHWC', 'NHWC', 'NWC', optional,

default='None' Set layout for input and output. Empty for default layout: NCW

for 1d, NCHW for 2d and NCDHW for 3d.

#### **Details**

- \*\*data\*\* and \*\*out\*\*: \*(batch\_size, channel, width)\* (NCW layout) or \*(batch\_size, width, channel)\* (NWC layout).

The shapes for 2-D pooling are

- \*\*data\*\* and \*\*out\*\*: \*(batch\_size, channel, height, width)\* (NCHW layout) or \*(batch\_size, height, width, channel)\* (NHWC layout),

out\_height = f(height, kernel[0], pad[0], stride[0]) out\_width = f(width, kernel[1], pad[1], stride[1])

The definition of \*f\* depends on "pooling\_convention", which has two options:

- \*\*valid\*\* (default)::

f(x, k, p, s) = floor((x+2\*p-k)/s)+1

- \*\*full\*\*, which is compatible with Caffe::

f(x, k, p, s) = ceil((x+2\*p-k)/s)+1

When "global\_pool" is set to be true, then global pooling is performed. It will reset "kernel=(height, width)" and set the appropriate padding to 0.

Three pooling options are supported by "pool\_type":

- \*\*avg\*\*: average pooling - \*\*max\*\*: max pooling - \*\*sum\*\*: sum pooling - \*\*lp\*\*: Lp pooling

For 3-D pooling, an additional \*depth\* dimension is added before \*height\*. Namely the input data and output will have shape \*(batch\_size, channel, depth, height, width)\* (NCDHW layout) or \*(batch\_size, depth, height, width, channel)\* (NDHWC layout).

Notes on Lp pooling:

mx.nd.Pooling.v1

Lp pooling was first introduced by this paper: https://arxiv.org/pdf/1204.3968.pdf. L-1 pooling is simply sum pooling, while L-inf pooling is simply max pooling. We can see that Lp pooling stands between those two, in practice the most common value for p is 2.

For each window "X", the mathematical expression for Lp pooling is:

```
:math: f(X) = \sqrt{p}\sum_x^X x^p
```

Defined in src/operator/nn/pooling.cc:L414

#### Value

out The result mx.ndarray

mx.nd.Pooling.v1

This operator is DEPRECATED. Perform pooling on the input.

## **Description**

The shapes for 2-D pooling is

### **Arguments**

data	NDArray-or-Symbol Input data to the pooling operator.	
kernel	Shape(tuple), optional, default=[] pooling kernel size: $(y, x)$ or $(d, y, x)$	
pool.type	'avg', 'max', 'sum',optional, default='max' Pooling type to be applied.	
global.pool	boolean, optional, default=0 Ignore kernel size, do global pooling based on current input feature map.	
pooling.convention		
	'full', 'valid',optional, default='valid' Pooling convention to be applied.	
stride	Shape(tuple), optional, default=[] stride: for pooling $(y, x)$ or $(d, y, x)$	

## **Details**

pad

```
- **data**: *(batch_size, channel, height, width)* - **out**: *(batch_size, num_filter, out_height, out_width)*, with::
```

Shape(tuple), optional, default=[] pad for pooling: (y, x) or (d, y, x)

 $out\_height = f(height, kernel[0], pad[0], stride[0]) \ out\_width = f(width, kernel[1], pad[1], stride[1])$ 

The definition of \*f\* depends on "pooling\_convention", which has two options:

- \*\*valid\*\* (default)::

```
f(x, k, p, s) = floor((x+2*p-k)/s)+1
```

- \*\*full\*\*, which is compatible with Caffe::

```
f(x, k, p, s) = ceil((x+2*p-k)/s)+1
```

But "global\_pool" is set to be true, then do a global pooling, namely reset "kernel=(height, width)".

Three pooling options are supported by "pool\_type":

```
- **avg**: average pooling - **max**: max pooling - **sum**: sum pooling
```

1-D pooling is special case of 2-D pooling with \*weight=1\* and \*kernel[1]=1\*.

For 3-D pooling, an additional \*depth\* dimension is added before \*height\*. Namely the input data will have shape \*(batch\_size, channel, depth, height, width)\*.

Defined in src/operator/pooling\_v1.cc:L104

#### Value

out The result mx.ndarray

```
mx.nd.preloaded.multi.mp.sgd.mom.update
```

Momentum update function for multi-precision Stochastic Gradient Descent (SGD) optimizer.

### **Description**

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

# Arguments

decays

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

# Details

```
.. math::
```

```
v = \alpha y + 1 =
```

It updates the weights using::

```
v = momentum * v - learning_rate * gradient weight += v
```

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

Defined in src/operator/contrib/preloaded\_multi\_sgd.cc:L200

# Value

mx.nd.preloaded.multi.mp.sgd.update

Update function for multi-precision Stochastic Gradient Descent (SDG) optimizer.

## **Description**

It updates the weights using::

## **Arguments**

data NDArray-or-Symbol[] Weights, gradients, learning rates and weight decays

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

## **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
```

Defined in src/operator/contrib/preloaded\_multi\_sgd.cc:L140

### Value

out The result mx.ndarray

 ${\tt mx.nd.preloaded.multi.sgd.mom.update}$ 

Momentum update function for Stochastic Gradient Descent (SGD) optimizer.

## Description

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

### **Arguments**

data NDArray-or-Symbol[] Weights, gradients, momentum, learning rates and weight

decays

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

#### **Details**

.. math::

 $v_1 = \alpha V_t - 1 - \alpha V_t - \Omega V_t = \gamma V_t - 1 - \alpha V_t$ 

It updates the weights using::

v = momentum \* v - learning\_rate \* gradient weight += v

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

Defined in src/operator/contrib/preloaded\_multi\_sgd.cc:L91

### Value

out The result mx.ndarray

mx.nd.preloaded.multi.sgd.update

Update function for Stochastic Gradient Descent (SDG) optimizer.

### **Description**

It updates the weights using::

## **Arguments**

data NDArray-or-Symbol[] Weights, gradients, learning rates and weight decays

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

## Details

weight = weight - learning\_rate \* (gradient + wd \* weight)

Defined in src/operator/contrib/preloaded\_multi\_sgd.cc:L42

mx.nd.prod 197

### Value

out The result mx.ndarray

mx.nd.prod

Computes the product of array elements over given axes.

### **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L31

## **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

## Value

out The result mx.ndarray

mx.nd.radians

Converts each element of the input array from degrees to radians.

## Description

```
.. math:: radians([0, 90, 180, 270, 360]) = [0, \pi/2, \pi/2, 2\pi]
```

### **Arguments**

data

NDArray-or-Symbol The input array.

## **Details**

The storage type of "radians" output depends upon the input storage type:

- radians(default) = default - radians(row\_sparse) = row\_sparse - radians(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L351

## Value

out The result mx.ndarray

mx.nd.random.exponential

Draw random samples from an exponential distribution.

# Description

Samples are distributed according to an exponential distribution parametrized by \*lambda\* (rate).

# Arguments

lam	float, optional, default=1 Lambda parameter (rate) of the exponential distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

# **Details**

```
Example::
```

```
exponential(lam=4, shape=(2,2)) = [[ 0.0097189 , 0.08999364], [ 0.04146638, 0.31715935]] Defined in src/operator/random/sample_op.cc:L137
```

### Value

mx.nd.random.gamma 199

mx.nd.random.ga
-----------------

Draw random samples from a gamma distribution.

# Description

Samples are distributed according to a gamma distribution parametrized by \*alpha\* (shape) and \*beta\* (scale).

## Arguments

alpha	float, optional, default=1 Alpha parameter (shape) of the gamma distribution.
beta	float, optional, default=1 Beta parameter (scale) of the gamma distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

### **Details**

## Example::

```
gamma(alpha=9, beta=0.5, shape=(2,2)) = [[ 7.10486984, 3.37695289], [ 3.91697288, 3.65933681]] Defined in src/operator/random/sample_op.cc:L125
```

### Value

out The result mx.ndarray

```
mx.nd.random.generalized.negative.binomial
```

Draw random samples from a generalized negative binomial distribution.

# Description

Samples are distributed according to a generalized negative binomial distribution parametrized by \*mu\* (mean) and \*alpha\* (dispersion). \*alpha\* is defined as \*1/k\* where \*k\* is the failure limit of the number of unsuccessful experiments (generalized to real numbers). Samples will always be returned as a floating point data type.

## Arguments

mu	float, optional, default=1	Mean of the negative binomial distribution.	
----	----------------------------	---	--

alpha float, optional, default=1 Alpha (dispersion) parameter of the negative binomial

distribution.

shape Shape(tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

#### **Details**

### Example::

```
generalized_negative_binomial(mu=2.0, alpha=0.3, shape=(2,2)) = [[ 2., 1.], [ 6., 4.]]
```

Defined in src/operator/random/sample\_op.cc:L179

### Value

out The result mx.ndarray

```
mx.nd.random.negative.binomial
```

Draw random samples from a negative binomial distribution.

### **Description**

Samples are distributed according to a negative binomial distribution parametrized by \*k\* (limit of unsuccessful experiments) and \*p\* (failure probability in each experiment). Samples will always be returned as a floating point data type.

# **Arguments**

k	int, optional, default='1' Limit of unsuccessful experiments.
р	float, optional, default=1 Failure probability in each experiment.
shape	Shape(tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to float 32 if not defined (dtype=None).

# **Details**

#### Example::

```
negative_binomial(k=3, p=0.4, shape=(2,2)) = [[ 4., 7.], [ 2., 5.]]
```

Defined in src/operator/random/sample\_op.cc:L164

mx.nd.random.normal 201

## Value

out The result mx.ndarray

mx.nd.random.normal

Draw random samples from a normal (Gaussian) distribution.

# Description

.. note:: The existing alias "normal" is deprecated.

# Arguments

loc	float, optional, default=0 Mean of the distribution.
scale	float, optional, default=1 Standard deviation of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

# **Details**

Samples are distributed according to a normal distribution parametrized by \*loc\* (mean) and \*scale\* (standard deviation).

## Example::

```
normal(loc=0, scale=1, shape=(2,2)) = [[ 1.89171135, -1.16881478], [-1.23474145, 1.55807114]]
Defined in src/operator/random/sample_op.cc:L113
```

## Value

mx.nd.random.pdf.dirichlet

Computes the value of the PDF of \*sample\* of Dirichlet distributions with parameter \*alpha\*.

### **Description**

The shape of \*alpha\* must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*alpha\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the value of \*alpha\* at index \*i\*.

### **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
alpha	NDArray-or-Symbol Concentration parameters of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.

#### **Details**

### Examples::

```
random_pdf_dirichlet(sample=[[1,2],[2,3],[3,4]], alpha=[2.5, 2.5]) = [38.413498, 199.60245, 564.56085] sample = [[[1, 2, 3], [10, 20, 30], [100, 200, 300]], [[0.1, 0.2, 0.3], [0.01, 0.02, 0.03], [0.001, 0.002, 0.003]]] random_pdf_dirichlet(sample=sample, alpha=[0.1, 0.4, 0.9]) = [[2.3257459e-02, 5.8420084e-04, 1.4674458e-05], [9.2589635e-01, 3.6860607e+01, 1.4674468e+03]] Defined in src/operator/random/pdf_op.cc:L316
```

#### Value

out The result mx.ndarray

```
mx.nd.random.pdf.exponential
```

Computes the value of the PDF of \*sample\* of exponential distributions with parameters \*lam\* (rate).

## **Description**

The shape of \*lam\* must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*lam\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the value of \*lam\* at index \*i\*.

### **Arguments**

sample NDArray-or-Symbol Samples from the distributions.

lam NDArray-or-Symbol Lambda (rate) parameters of the distributions.

is.log boolean, optional, default=0 If set, compute the density of the log-probability

instead of the probability.

#### **Details**

### Examples::

```
random_pdf_exponential(sample=[[1, 2, 3]], lam=[1]) = [[0.36787945, 0.13533528, 0.04978707]] sample = [[1,2,3], [1,2,3], [1,2,3]]
```

random\_pdf\_exponential(sample=sample, lam=[1,0.5,0.25]) = [[0.36787945, 0.13533528, 0.04978707], [0.30326533, 0.18393973, 0.11156508], [0.1947002, 0.15163267, 0.11809164]]

Defined in src/operator/random/pdf\_op.cc:L305

## Value

out The result mx.ndarray

```
{\tt mx.nd.random.pdf.gamma}
```

Computes the value of the PDF of \*sample\* of gamma distributions with parameters \*alpha\* (shape) and \*beta\* (rate).

### **Description**

\*alpha\* and \*beta\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*alpha\* and \*beta\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*alpha\* and \*beta\* at index \*i\*.

### **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
alpha	NDArray-or-Symbol Alpha (shape) parameters of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
beta	NDArray-or-Symbol Beta (scale) parameters of the distributions.

#### **Details**

### Examples::

```
 \begin{array}{l} {\rm random\_pdf\_gamma(sample=[[1,2,3,4,5]], alpha=[5], beta=[1])=[[0.01532831, 0.09022352, 0.16803136, 0.19536681, 0.17546739]]} \\ {\rm sample=[[1,2,3,4,5], [2,3,4,5,6], [3,4,5,6,7]]} \\ {\rm random\_pdf\_gamma(sample=sample, alpha=[5,6,7], beta=[1,1,1])=[[0.01532831, 0.09022352, 0.16803136, 0.19536681, 0.17546739], [0.03608941, 0.10081882, 0.15629345, 0.17546739, 0.16062315], [0.05040941, 0.10419563, 0.14622283, 0.16062315, 0.14900276]]} \\ {\rm Defined\ in\ src/operator/random/pdf\_op.cc:L303} \end{array}
```

#### Value

out The result mx.ndarray

```
mx.nd.random.pdf.generalized.negative.binomial
```

Computes the value of the PDF of \*sample\* of generalized negative binomial distributions with parameters \*mu\* (mean) and \*alpha\* (dispersion). This can be understood as a reparameterization of the negative binomial, where \*k\* = \*1 / alpha\* and  $*p* = *1 / (mu \* alpha + 1)*$ .

### Description

\*mu\* and \*alpha\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*mu\* and \*alpha\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*mu\* and \*alpha\* at index \*i\*.

### Arguments

sample	NDArray-or-Symbol Samples from the distributions.
mu	NDArray-or-Symbol Means of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
alpha	NDArray-or-Symbol Alpha (dispersion) parameters of the distributions.

#### **Details**

## Examples::

random\_pdf\_generalized\_negative\_binomial(sample=[[1, 2, 3, 4]], alpha=[1], mu=[1]) = [[0.25, 0.125, 0.0625, 0.03125]]

sample = [[1,2,3,4], [1,2,3,4]] random\_pdf\_generalized\_negative\_binomial(sample=sample, alpha=[1, 0.6666], mu=[1, 1.5]) = [[0.25, 0.125, 0.0625, 0.03125], [0.26517063, 0.16573331, 0.09667706, 0.05437994]]

Defined in src/operator/random/pdf\_op.cc:L314

#### Value

out The result mx.ndarray

mx.nd.random.pdf.negative.binomial

Computes the value of the PDF of samples of negative binomial distributions with parameters \*k\* (failure limit) and \*p\* (failure probability).

# Description

\*k\* and \*p\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*k\* and \*p\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*k\* and \*p\* at index \*i\*.

## **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
k	NDArray-or-Symbol Limits of unsuccessful experiments.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
р	NDArray-or-Symbol Failure probabilities in each experiment.

#### **Details**

### Examples::

```
 \begin{array}{l} random\_pdf\_negative\_binomial(sample=[[1,2,3,4]], k=[1], p=a[0.5]) = [[0.25, 0.125, 0.0625, 0.03125]] \\ \# Note that k may be real-valued sample = [[1,2,3,4], [1,2,3,4]] \\ random\_pdf\_negative\_binomial(sample=sample, k=[1, 1.5], p=[0.5, 0.5]) = [[0.25, 0.125, 0.0625, 0.03125], [0.26516506, 0.16572815, 0.09667476, 0.05437956]] \\ \end{array}
```

Defined in src/operator/random/pdf\_op.cc:L310

### Value

out The result mx.ndarray

mx.nd.random.pdf.normal

Computes the value of the PDF of \*sample\* of normal distributions with parameters \*mu\* (mean) and \*sigma\* (standard deviation).

# Description

\*mu\* and \*sigma\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*mu\* and \*sigma\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*mu\* and \*sigma\* at index \*i\*.

## Arguments

sample	NDArray-or-Symbol Samples from the distributions.
mu	NDArray-or-Symbol Means of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
sigma	NDArray-or-Symbol Standard deviations of the distributions.

## **Details**

# Examples::

```
sample = [[-2, -1, 0, 1, 2]] random_pdf_normal(sample=sample, mu=[0], sigma=[1]) = [[0.05399097, 0.24197073, 0.3989423, 0.24197073, 0.05399097]]
```

random\_pdf\_normal(sample=sample\*2, mu=[0,0], sigma=[1,2]) = [[0.05399097, 0.24197073, 0.3989423, 0.24197073, 0.05399097], [0.12098537, 0.17603266, 0.19947115, 0.17603266, 0.12098537]]

Defined in src/operator/random/pdf\_op.cc:L300

## Value

mx.nd.random.pdf.poisson

Computes the value of the PDF of \*sample\* of Poisson distributions with parameters \*lam\* (rate).

### **Description**

The shape of \*lam\* must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*lam\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the value of \*lam\* at index \*i\*.

### **Arguments**

sample NDArray-or-Symbol Samples from the distributions.

lam NDArray-or-Symbol Lambda (rate) parameters of the distributions.

is.log boolean, optional, default=0 If set, compute the density of the log-probability

instead of the probability.

#### **Details**

## Examples::

```
\begin{split} & \text{random\_pdf\_poisson}(\text{sample=}[[0,1,2,3]], \text{lam=}[1]) = [[0.36787945, 0.36787945, 0.18393973, 0.06131324]] \\ & \text{sample=}[[0,1,2,3], [0,1,2,3], [0,1,2,3]] \end{split}
```

random\_pdf\_poisson(sample=sample, lam=[1,2,3]) = [[0.36787945, 0.36787945, 0.18393973, 0.06131324], [0.13533528, 0.27067056, 0.27067056, 0.18044704], [0.04978707, 0.14936121, 0.22404182, 0.22404182]]

Defined in src/operator/random/pdf\_op.cc:L307

## Value

out The result mx.ndarray

mx.nd.random.pdf.uniform

Computes the value of the PDF of \*sample\* of uniform distributions on the intervals given by \*[low,high)\*.

## Description

\*low\* and \*high\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*low\* and \*high\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*low\* and \*high\* at index \*i\*.

208 mx.nd.random.poisson

#### **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
low	NDArray-or-Symbol Lower bounds of the distributions.

is.log boolean, optional, default=0 If set, compute the density of the log-probability

instead of the probability.

high NDArray-or-Symbol Upper bounds of the distributions.

### **Details**

### Examples::

```
 \begin{array}{l} {\rm random\_pdf\_uniform(sample=[[1,2,3,4]],\ low=[0],\ high=[10])=[0.1,\ 0.1,\ 0.1,\ 0.1]} \\ {\rm sample=[[[1,2,3],\ [1,2,3]],\ [[1,2,3],\ [1,2,3]]]\ low=[[0,0],\ [0,0]]\ high=[[5,10],\ [15,20]]\ random\_pdf\_uniform(sample=sample,\ low=low,\ high=high)=[[[0.2,\ 0.2,\ 0.2],\ [0.1,\ 0.1,\ 0.1]],\ [[0.06667,\ 0.06667,\ 0.06667],\ [0.05,\ 0.05,\ 0.05]]] \\ {\rm Defined\ in\ src/operator/random/pdf\_op.cc:L298} \\ \end{array}
```

### Value

out The result mx.ndarray

mx.nd.random.poisson Draw random samples from a Poisson distribution.

### **Description**

Samples are distributed according to a Poisson distribution parametrized by \*lambda\* (rate). Samples will always be returned as a floating point data type.

# Arguments

lam	float, optional, default=1 Lambda parameter (rate) of the Poisson distribution.
shape	Shape(tuple), optional, default=None Shape of the output.

Shape (tuple), optional, default—I tone Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'float16', 'float32', 'float64',optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

#### **Details**

# Example::

```
poisson(lam=4, shape=(2,2)) = [[ 5., 2.], [ 4., 6.]]
Defined in src/operator/random/sample_op.cc:L150
```

#### Value

mx.nd.random.randint 209

mx.nd.random.randint Draw random samples from a discrete uniform distribution.

#### **Description**

Samples are uniformly distributed over the half-open interval \*[low, high)\* (includes \*low\*, but excludes \*high\*).

# Arguments

low long, required Lower bound of the distribution. high long, required Upper bound of the distribution.

shape Shape(tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'int32', 'int64', optional, default='None' DType of the output in case

this can't be inferred. Defaults to int32 if not defined (dtype=None).

#### **Details**

Example::

randint(low=0, high=5, shape=(2,2)) = [[ 0, 2], [ 3, 1]] Defined in src/operator/random/sample\_op.cc:L194

## Value

out The result mx.ndarray

mx.nd.random.uniform Draw random samples from a uniform distribution.

## **Description**

.. note:: The existing alias "uniform" is deprecated.

## **Arguments**

low	float, optional, default=0 Lower bound of the distribution.
high	float, optional, default=1 Upper bound of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

210 mx.nd.ravel.multi.index

### **Details**

Samples are uniformly distributed over the half-open interval \*[low, high)\* (includes \*low\*, but excludes \*high\*).

Example::

uniform(low=0, high=1, shape=(2,2)) = [[ 0.60276335, 0.85794562], [ 0.54488319, 0.84725171]]

Defined in src/operator/random/sample\_op.cc:L96

## Value

out The result mx.ndarray

mx.nd.ravel.multi.index

Converts a batch of index arrays into an array of flat indices. The operator follows numpy conventions so a single multi index is given by a column of the input matrix. The leading dimension may be left unspecified by using -1 as placeholder.

# **Description**

Examples::

A = [[3,6,6],[4,5,1]] ravel(A, shape=(7,6)) = [22,41,37] ravel(A, shape=(-1,6)) = [22,41,37]

## Arguments

data NDArray-or-Symbol Batch of multi-indices

shape Shape(tuple), optional, default=None Shape of the array into which the multi-

indices apply.

#### **Details**

Defined in src/operator/tensor/ravel.cc:L42

## Value

mx.nd.rcbrt 211

mx.nd.rcbrt

Returns element-wise inverse cube-root value of the input.

# **Description**

```
.. math:: rcbrt(x) = 1 \land sqrt[3]x
```

# Arguments

data

NDArray-or-Symbol The input array.

### **Details**

```
Example::
```

```
rcbrt([1,8,-125]) = [1.0, 0.5, -0.2]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L323

### Value

out The result mx.ndarray

mx.nd.reciprocal

Returns the reciprocal of the argument, element-wise.

# Description

Calculates 1/x.

# **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

```
Example::
```

```
reciprocal([-2, 1, 3, 1.6, 0.2]) = [-0.5, 1.0, 0.33333334, 0.625, 5.0]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L43

# Value

212 mx.nd.repeat

mx.nd.relu

Computes rectified linear activation.

### **Description**

.. math:: max(features, 0)

## **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

The storage type of "relu" output depends upon the input storage type:

- relu(default) = default - relu(row\_sparse) = row\_sparse - relu(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L85

### Value

out The result mx.ndarray

mx.nd.repeat

Repeats elements of an array. By default, "repeat" flattens the input array into 1-D and then repeats the elements:: x = [[1, 2], [3, 4]] repeat(x, repeats=2) = [1, 1, 2, 2, 3, 3, 4, 4] The parameter "axis" specifies the axis along which to perform repeat:: repeat(x, repeats=2, x) = [[1, 1, 2, 2, 2], [3, 3, 4, 4]] repeat(x, repeats=2, x) = [[1, 2, 2], [1, 2, 2], [3, 4], [3, 4]] repeat(x, repeats=2, x) = [[1, 2, 2, 2], [3, 3, 4], [3, 4]]

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L744

# Arguments

data NDArray-or-Symbol Input data array

repeats int, required The number of repetitions for each element.

axis int or None, optional, default='None' The axis along which to repeat values.

The negative numbers are interpreted counting from the backward. By default,

use the flattened input array, and return a flat output array.

## Value

mx.nd.reset.arrays 213

mx.nd.reset.arrays
Set to zero multiple arrays

# Description

Defined in src/operator/contrib/reset\_arrays.cc:L36

# **Arguments**

data NDArray-or-Symbol[] Arrays

num.arrays int, required number of input arrays.

# Value

214 mx.nd.Reshape

mx.nd.Reshape

Reshapes the input array. .. note:: "Reshape" is deprecated, use "reshape" Given an array and a shape, this function returns a copy of the array in the new shape. The shape is a tuple of integers such as (2,3,4). The size of the new shape should be same as the size of the input array. Example:: reshape([1,2,3,4], shape=(2,2)) = [[1,2],[3,4]] Some dimensions of the shape can take special values from the set 0, -1, -2, -3, -4. The significance of each is explained below: - "0" copy this dimension from the input to the output shape. Example:: input shape = (2,3,4), shape = (4,0,2), output shape = (4,3,2) - input shape = (2,3,4), shape = (2,0,0), output shape = (2,3,4) - "-1" infers the dimension of the output shape by using the remainder of the input dimensions keeping the size of the new array same as that of the input array. At most one dimension of shape can be -1. Example:: - input shape = (2,3,4), shape = (6,1,-1), output shape = (6,1,4) - input shape = (2,3,4), shape = (3,-1,8), output shape = (3,1,8) - input shape =(2,3,4), shape=(-1,),  $output\ shape=(24,)-"-2"\ copy\ all/remainder$ of the input dimensions to the output shape. Example:: - input shape = (2,3,4), shape = (-2,), output shape = (2,3,4) - input shape = (2,3,4), shape = (2,-2), output shape = (2,3,4) - input shape = (2,3,4), shape = (-2,1,1), output shape = (2,3,4,1,1) - "-3" use the product of two consecutive dimensions of the input shape as the output dimension. Example:: - input shape = (2,3,4), shape = (-3,4), output shape = (6,4) - input shape = (2,3,4,5), shape = (-3,-3), output shape = (6,20)- input shape = (2,3,4), shape = (0,-3), output shape = (2,12) - input shape = (2,3,4), shape = (-3,-2), output shape = (6,4) - "-4" split one dimension of the input into two dimensions passed subsequent to -4 in shape (can contain -1). Example:: - input shape = (2,3,4), shape = (-4,1,2,-2), output shape = (1,2,3,4) - input shape = (2,3,4), shape = (2,-4,-1,3,-2), output shape = (2,1,3,4) If the argument 'reverse' is set to 1, then the special values are inferred from right to left. Example:: - without reverse=1, for input shape = (10,5,4), shape = (-1,0), output shape would be (40,5) - with reverse=1, output shape will be (50,4).

#### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L175

### **Arguments**

data NDArray-or-Symbol Input data to reshape.

shape Shape(tuple), optional, default=[] The target shape

reverse boolean, optional, default=0 If true then the special values are inferred from

right to left

target.shape Shape(tuple), optional, default=[] (Deprecated! Use "shape" instead.) Target

new shape. One and only one dim can be 0, in which case it will be inferred

from the rest of dims

mx.nd.reshape 215

keep.highest

boolean, optional, default=0 (Deprecated! Use "shape" instead.) Whether keep the highest dim unchanged. If set to true, then the first dim in target\_shape is ignored, and always fixed as input

#### Value

out The result mx.ndarray

mx.nd.reshape

Reshapes the input array. .. note:: "Reshape" is deprecated, use "reshape" Given an array and a shape, this function returns a copy of the array in the new shape. The shape is a tuple of integers such as (2,3,4). The size of the new shape should be same as the size of the input array. Example:: reshape([1,2,3,4], shape=(2,2)) = [[1,2],[3,4]] Some dimensions of the shape can take special values from the set 0, -1, -2, -3, -4. The significance of each is explained below: - "0" copy this dimension from the input to the output shape. Example:: input shape = (2,3,4), shape = (4,0,2), output shape = (4,3,2) - input shape = (2,3,4), shape = (2,0,0), output shape = (2,3,4) - "-1" infers the dimension of the output shape by using the remainder of the input dimensions keeping the size of the new array same as that of the input array. At most one dimension of shape can be -1. Example:: - input shape = (2,3,4), shape = (6,1,-1), output shape = (6,1,4) - input shape= (2,3,4), shape = (3,-1,8), output shape = (3,1,8) - input shape =(2,3,4), shape=(-1,), output shape=(24,) - "-2" copy all/remainder of the input dimensions to the output shape. Example:: - input shape = (2,3,4), shape = (-2,), output shape = (2,3,4) - input shape = (2,3,4), shape = (2,-2), output shape = (2,3,4) - input shape = (2,3,4), shape= (-2,1,1), output shape = (2,3,4,1,1) - "-3" use the product of two consecutive dimensions of the input shape as the output dimension. Example:: - input shape = (2,3,4), shape = (-3,4), output shape = (6,4) - input shape = (2,3,4,5), shape = (-3,-3), output shape = (6,20)- input shape = (2,3,4), shape = (0,-3), output shape = (2,12) - input shape = (2,3,4), shape = (-3,-2), output shape = (6,4) - "-4" split one dimension of the input into two dimensions passed subsequent to -4 in shape (can contain -1). Example:: - input shape = (2,3,4), shape = (-4,1,2,-2), output shape = (1,2,3,4) - input shape = (2,3,4), shape = (2,-4,-1,3,-2), output shape = (2,1,3,4) If the argument 'reverse' is set to 1, then the special values are inferred from right to left. Example:: - without reverse=1, for input shape = (10,5,4), shape = (-1,0), output shape would be (40,5) - with reverse=1, output shape will be (50,4).

### Description

Defined in src/operator/tensor/matrix\_op.cc:L175

216 mx.nd.reshape.like

### **Arguments**

data NDArray-or-Symbol Input data to reshape.

shape Shape(tuple), optional, default=[] The target shape

reverse boolean, optional, default=0 If true then the special values are inferred from

right to left

target.shape Shape(tuple), optional, default=[] (Deprecated! Use "shape" instead.) Target

new shape. One and only one dim can be 0, in which case it will be inferred

from the rest of dims

keep.highest boolean, optional, default=0 (Deprecated! Use "shape" instead.) Whether keep

the highest dim unchanged. If set to true, then the first dim in target shape is

ignored, and always fixed as input

#### Value

out The result mx.ndarray

mx.nd.reshape.like Reshape some or all dimensions of 'lhs' to have the same shape as

some or all dimensions of 'rhs'.

# Description

Returns a \*\*view\*\* of the 'lhs' array with a new shape without altering any data.

# Arguments

1hs NDArray-or-Symbol First input.

rhs NDArray-or-Symbol Second input.

lhs.begin int or None, optional, default='None' Defaults to 0. The beginning index along

which the lhs dimensions are to be reshaped. Supports negative indices.

1hs. end int or None, optional, default='None' Defaults to None. The ending index along

which the lhs dimensions are to be used for reshaping. Supports negative in-

dices.

rhs.begin int or None, optional, default='None' Defaults to 0. The beginning index along

which the rhs dimensions are to be used for reshaping. Supports negative in-

dices.

rhs.end int or None, optional, default='None' Defaults to None. The ending index along

which the rhs dimensions are to be used for reshaping. Supports negative in-

dices.

mx.nd.reverse 217

### **Details**

#### Example::

```
x = [1, 2, 3, 4, 5, 6] y = [[0, -4], [3, 2], [2, 2]] reshape_like(x, y) = [[1, 2], [3, 4], [5, 6]]
```

More precise control over how dimensions are inherited is achieved by specifying \ slices over the 'lhs' and 'rhs' array dimensions. Only the sliced 'lhs' dimensions \ are reshaped to the 'rhs' sliced dimensions, with the non-sliced 'lhs' dimensions staying the same.

### Examples::

- lhs shape = (30,7), rhs shape = (15,2,4), lhs\_begin=0, lhs\_end=1, rhs\_begin=0, rhs\_end=2, output shape = (15,2,7) - lhs shape = (3,5), rhs shape = (1,15,4), lhs\_begin=0, lhs\_end=2, rhs\_begin=1, rhs\_end=2, output shape = (15)

Negative indices are supported, and 'None' can be used for either 'lhs\_end' or 'rhs\_end' to indicate the end of the range.

#### Example::

- lhs shape = (30, 12), rhs shape = (4, 2, 2, 3), lhs\_begin=-1, lhs\_end=None, rhs\_begin=1, rhs\_end=None, output shape = (30, 2, 2, 3)

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L512

#### Value

out The result mx.ndarray

mx	. na	.rev	ver	se

Reverses the order of elements along given axis while preserving array shape. Note: reverse and flip are equivalent. We use reverse in the following examples. Examples:: x = [[0., 1., 2., 3., 4.], [5., 6., 7., 8., 9.]] reverse(x, axis=0) = [[5., 6., 7., 8., 9.], [0., 1., 2., 3., 4.]] reverse(x, axis=1) = [[4., 3., 2., 1., 0.], [9., 8., 7., 6., 5.]]

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L832

# **Arguments**

data NDArray-or-Symbol Input data array

axis Shape(tuple), required The axis which to reverse elements.

### Value

218 mx.nd.rmsprop.update

mx.nd.rint

Returns element-wise rounded value to the nearest integer of the input.

# Description

.. note:: - For input "n.5" "rint" returns "n" while "round" returns "n+1". - For input "-n.5" both "rint" and "round" returns "-n-1".

## **Arguments**

data

NDArray-or-Symbol The input array.

#### **Details**

# Example::

```
rint([-1.5, 1.5, -1.9, 1.9, 2.1]) = [-2., 1., -2., 2., 2.]
```

The storage type of "rint" output depends upon the input storage type:

- rint(default) = default - rint(row\_sparse) = row\_sparse - rint(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L799

NDArray-or-Symbol Weight

### Value

out The result mx.ndarray

mx.nd.rmsprop.update Update function for 'RMSProp' optimizer.

# Description

'RMSprop' is a variant of stochastic gradient descent where the gradients are divided by a cache which grows with the sum of squares of recent gradients?

### **Arguments**

weight

grad	NDArray-or-Symbol Gradient
n	NDArray-or-Symbol n
lr	float, required Learning rate
rho	float, optional, default=0.949999988 The decay rate of momentum estimates.
epsilon	float, optional, default=9.99999994e-09 A small constant for numerical stability.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip gradient, clip gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

clip.weights float, optional, default=-1 Clip weights to the range of [-clip\_weights, clip\_weights]

If clip\_weights <= 0, weight clipping is turned off. weights = max(min(weights,

clip\_weights), -clip\_weights).

#### **Details**

'RMSProp' is similar to 'AdaGrad', a popular variant of 'SGD' which adaptively tunes the learning rate of each parameter. 'AdaGrad' lowers the learning rate for each parameter monotonically over the course of training. While this is analytically motivated for convex optimizations, it may not be ideal for non-convex problems. 'RMSProp' deals with this heuristically by allowing the learning rates to rebound as the denominator decays over time.

Define the Root Mean Square (RMS) error criterion of the gradient as :math: 'RMS[g]\_t = \sqrtE[g^2]\_t + \epsilon', where :math: 'g' represents gradient and :math: 'E[g^2]\_t' is the decaying average over past squared gradient.

The :math:  $E[g^2]_t$  is given by:

.. math::  $E[g^2]_t = \rho * E[g^2]_{t-1} + (1-\rho) * g_t^2$ 

The update step is

.. math::  $\theta_t = \theta_t - \frac{r}{g} g_t$ 

The RMSProp code follows the version in http://www.cs.toronto.edu/~tijmen/csc321/slides/lecture\_slides\_lec6.pdf Tieleman & Hinton, 2012.

Hinton suggests the momentum term :math:'\rho' to be 0.9 and the learning rate :math:'\eta' to be 0.001.

Defined in src/operator/optimizer\_op.cc:L788

## Value

out The result mx.ndarray

mx.nd.rmspropalex.update

Update function for RMSPropAlex optimizer.

#### **Description**

'RMSPropAlex' is non-centered version of 'RMSProp'.

## **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
n	NDArray-or-Symbol n
g	NDArray-or-Symbol g
delta	NDArray-or-Symbol delta
lr	float, required Learning rate
rho	float, optional, default=0.949999988 Decay rate.
momentum	float, optional, default=0.899999976 Decay rate.
epsilon	float, optional, default=9.99999994e-09 A small constant for numerical stability.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
clip.weights	float, optional, default=-1 Clip weights to the range of [-clip_weights, clip_weights]  If clip_weights <= 0, weight clipping is turned off. weights = max(min(weights,

# **Details**

Define :math:  $E[g^2]_t$  is the decaying average over past squared gradient and :math:  $E[g]_t$  is the decaying average over past gradient.

```
.. math:: E[g^2]_t = \rho * E[g^2]_{t-1} + (1 - \rho) * g_t^2 E[g]_t = \rho * E[g]_{t-1} + (1 - \rho) * g_t \mod t = \gamma * momentum_t = \gamma *
```

The update step is

..  $math:: \theta_t + 1 = \theta_t + momentum_t$ 

clip\_weights), -clip\_weights).

The RMSPropAlex code follows the version in http://arxiv.org/pdf/1308.0850v5.pdf Eq(38) - Eq(45) by Alex Graves, 2013.

Graves suggests the momentum term :math:  $\$  to be 0.95, :math:  $\$  and the learning rate :math:  $\$  to be 0.0001.

Defined in src/operator/optimizer\_op.cc:L827

### Value

mx.nd.RNN 221

mx.nd.RNN	Applies recurrent layers to input data. Currently, vanilla RNN, LSTM and GRU are implemented, with both multi-layer and bidirectional support.

# Description

When the input data is of type float32 and the environment variables MXNET\_CUDA\_ALLOW\_TENSOR\_CORE and MXNET\_CUDA\_TENSOR\_OP\_MATH\_ALLOW\_CONVERSION are set to 1, this operator will try to use pseudo-float16 precision (float32 math with float16 I/O) precision in order to use Tensor Cores on suitable NVIDIA GPUs. This can sometimes give significant speedups.

# Arguments

	data	NDArray-or-Symbol Input data to RNN	
	parameters	NDArray-or-Symbol Vector of all RNN trainable parameters concatenated	
	state	NDArray-or-Symbol initial hidden state of the RNN	
	state.cell	NDArray-or-Symbol initial cell state for LSTM networks (only for LSTM)	
	sequence.length		
		NDArray-or-Symbol Vector of valid sequence lengths for each element in batch. (Only used if use_sequence_length kwarg is True)	
	state.size	int (non-negative), required size of the state for each layer	
	num.layers	int (non-negative), required number of stacked layers	
	bidirectional	boolean, optional, default=0 whether to use bidirectional recurrent layers	
	mode	'gru', 'lstm', 'rnn_relu', 'rnn_tanh', required the type of RNN to compute	
	p	float, optional, default=0 drop rate of the dropout on the outputs of each RNN layer, except the last layer.	
	state.outputs	boolean, optional, default=0 Whether to have the states as symbol outputs.	
projection.size			
		int or None, optional, default='None' size of project size	
lstm.state.clip.min			
		double or None, optional, default=None Minimum clip value of LSTM states. This option must be used together with lstm_state_clip_max.	
	lstm.state.clip.max		
		1 11 NI C 1 1 C 1 NI NA C 1 1 1 CI CUDA	

double or None, optional, default=None Maximum clip value of LSTM states. This option must be used together with lstm\_state\_clip\_min.

lstm.state.clip.nan

boolean, optional, default=0 Whether to stop NaN from propagating in state by clipping it to min/max. If clipping range is not specified, this option is ignored.

use.sequence.length

boolean, optional, default=0 If set to true, this layer takes in an extra input parameter 'sequence\_length' to specify variable length sequence

222 mx.nd.RNN

#### **Details**

```
**Vanilla RNN**
```

Applies a single-gate recurrent layer to input X. Two kinds of activation function are supported: ReLU and Tanh.

With ReLU activation function:

```
.. math:: h_t = relu(W_ih * x_t + b_ih + W_hh * h_(t-1) + b_hh)
```

With Tanh activtion function:

```
.. math:: h_t = \tanh(W_ih * x_t + b_ih + W_hh * h_(t-1) + b_hh)
```

Reference paper: Finding structure in time - Elman, 1988. https://crl.ucsd.edu/~elman/Papers/fsit.pdf
\*\*LSTM\*\*

Long Short-Term Memory - Hochreiter, 1997. http://www.bioinf.jku.at/publications/older/2604.pdf

With the projection size being set, LSTM could use the projection feature to reduce the parameters size and give some speedups without significant damage to the accuracy.

Long Short-Term Memory Based Recurrent Neural Network Architectures for Large Vocabulary Speech Recognition - Sak et al. 2014. https://arxiv.org/abs/1402.1128

```
**GRU**
```

Gated Recurrent Unit - Cho et al. 2014. http://arxiv.org/abs/1406.1078

The definition of GRU here is slightly different from paper but compatible with CUDNN.

Defined in src/operator/rnn.cc:L363

## Value

mx.nd.ROIPooling 223

mx.nd.ROIPooling Performs region of interest(ROI) pooling on the input array.	
---	--

# **Description**

ROI pooling is a variant of a max pooling layer, in which the output size is fixed and region of interest is a parameter. Its purpose is to perform max pooling on the inputs of non-uniform sizes to obtain fixed-size feature maps. ROI pooling is a neural-net layer mostly used in training a 'Fast R-CNN' network for object detection.

## **Arguments**

data	NDArray-or-Symbol The input array to the pooling operator, a 4D Feature maps	
rois	NDArray-or-Symbol Bounding box coordinates, a 2D array of [[batch_index, x1, y1, x2, y2]], where (x1, y1) and (x2, y2) are top left and bottom right corners of designated region of interest. 'batch_index' indicates the index of corresponding image in the input array	
pooled.size	Shape(tuple), required ROI pooling output shape (h,w)	
spatial.scale	float, required Ratio of input feature map height (or w) to raw image height (or w). Equals the reciprocal of total stride in convolutional layers	

# **Details**

This operator takes a 4D feature map as an input array and region proposals as 'rois', then it pools over sub-regions of input and produces a fixed-sized output array regardless of the ROI size.

To crop the feature map accordingly, you can resize the bounding box coordinates by changing the parameters 'rois' and 'spatial\_scale'.

The cropped feature maps are pooled by standard max pooling operation to a fixed size output indicated by a 'pooled\_size' parameter. batch\_size will change to the number of region bounding boxes after 'ROIPooling'.

The size of each region of interest doesn't have to be perfectly divisible by the number of pooling sections('pooled\_size').

#### Example::

```
x = [[[[0., 1., 2., 3., 4., 5.], [6., 7., 8., 9., 10., 11.], [12., 13., 14., 15., 16., 17.], [18., 19., 20., 21., 22., 23.], [24., 25., 26., 27., 28., 29.], [30., 31., 32., 33., 34., 35.], [36., 37., 38., 39., 40., 41.], [42., 43., 44., 45., 46., 47.]]]]
```

// region of interest i.e. bounding box coordinates. y = [[0,0,0,4,4]]

// returns array of shape (2,2) according to the given roi with max pooling. ROIPooling(x, y, (2,2), 1.0) = [[[[14., 16.], [26., 28.]]]]

// region of interest is changed due to the change in 'spacial\_scale' parameter. ROIPooling(x, y, (2,2), (0,0)) = [[[[ 7., 9.], [ 19., 21.]]]]

Defined in src/operator/roi\_pooling.cc:L225

224 mx.nd.rsqrt

# Value

out The result mx.ndarray

mx.nd.round

Returns element-wise rounded value to the nearest integer of the input.

# **Description**

Example::

# Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

```
round([-1.5, 1.5, -1.9, 1.9, 2.1]) = [-2., 2., -2., 2., 2.]
```

The storage type of "round" output depends upon the input storage type:

- round(default) = default - round(row\_sparse) = row\_sparse - round(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L778

#### Value

out The result mx.ndarray

mx.nd.rsqrt

Returns element-wise inverse square-root value of the input.

# **Description**

```
.. math:: rsqrt(x) = 1 \land sqrtx
```

# Arguments

data

NDArray-or-Symbol The input array.

## **Details**

Example::

```
rsqrt([4,9,16]) = [0.5, 0.33333334, 0.25]
```

The storage type of "rsqrt" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L221

### Value

mx.nd.sample.exponential

Concurrent sampling from multiple exponential distributions with parameters lambda (rate).

## **Description**

The parameters of the distributions are provided as an input array. Let \*[s]\* be the shape of the input array, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# **Arguments**

lam	NDArray-or-Symbol Lambda (rate) parameters of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

#### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input array, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input value at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input array.

#### Examples::

```
lam = [1.0, 8.5]
```

// Draw a single sample for each distribution sample\_exponential(lam) = [ 0.51837951, 0.09994757]

// Draw a vector containing two samples for each distribution sample\_exponential(lam, shape=(2)) = [[ 0.51837951, 0.19866663], [ 0.09994757, 0.50447971]]

Defined in src/operator/random/multisample\_op.cc:L284

#### Value

226 mx.nd.sample.gamma

mx.nd.sample.gamma	Concurrent sampling from multiple gamma distributions with parameters *alpha* (shape) and *beta* (scale).
	eters *alpha* (shape) and *beta* (scale).

# **Description**

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# **Arguments**

alpha	NDArray-or-Symbol Alpha (shape) parameters of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
beta	NDArray-or-Symbol Beta (scale) parameters of the distributions.

#### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

### Examples::

```
alpha = [ 0.0, 2.5 ] beta = [ 1.0, 0.7 ]

// Draw a single sample for each distribution sample_gamma(alpha, beta) = [ 0. , 2.25797319]

// Draw a vector containing two samples for each distribution sample_gamma(alpha, beta, shape=(2)) = [[ 0. , 0. ], [ 2.25797319, 1.70734084]]

Defined in src/operator/random/multisample_op.cc:L282
```

# Value

mx.nd.sample.generalized.negative.binomial

Concurrent sampling from multiple generalized negative binomial distributions with parameters \*mu\* (mean) and \*alpha\* (dispersion).

# Description

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# **Arguments**

mu	NDArray-or-Symbol Means of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
alpha	NDArray-or-Symbol Alpha (dispersion) parameters of the distributions.

## **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

Samples will always be returned as a floating point data type.

### Examples::

```
mu = [2.0, 2.5] alpha = [1.0, 0.1]
```

// Draw a single sample for each distribution sample\_generalized\_negative\_binomial(mu, alpha) = [0., 3.]

// Draw a vector containing two samples for each distribution sample\_generalized\_negative\_binomial(mu, alpha, shape=(2)) = [[ 0., 3.], [ 3., 1.]]

Defined in src/operator/random/multisample\_op.cc:L293

# Value

```
mx.nd.sample.multinomial
```

Concurrent sampling from multiple multinomial distributions.

# **Description**

\*data\* is an \*n\* dimensional array whose last dimension has length \*k\*, where \*k\* is the number of possible outcomes of each multinomial distribution. This operator will draw \*shape\* samples from each distribution. If shape is empty one sample will be drawn from each distribution.

### **Arguments**

data	NDArray-or-Symbol Distribution probabilities. Must sum to one on the last axis.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
get.prob	boolean, optional, default=0 Whether to also return the log probability of sampled result. This is usually used for differentiating through stochastic variables, e.g. in reinforcement learning.
dtype	'float16', 'float32', 'float64', 'int32', 'uint8',optional, default='int32' DType of the output in case this can't be inferred.

# **Details**

If \*get\_prob\* is true, a second array containing log likelihood of the drawn samples will also be returned. This is usually used for reinforcement learning where you can provide reward as head gradient for this array to estimate gradient.

Note that the input distribution must be normalized, i.e. \*data\* must sum to 1 along its last axis.

# Examples::

```
probs = [[0, 0.1, 0.2, 0.3, 0.4], [0.4, 0.3, 0.2, 0.1, 0]]

// Draw a single sample for each distribution sample_multinomial(probs) = [3, 0]

// Draw a vector containing two samples for each distribution sample_multinomial(probs, shape=(2)) = [[4, 2], [0, 0]]

// requests log likelihood sample_multinomial(probs, get_prob=True) = [2, 1], [0.2, 0.3]
```

#### Value

mx.nd.sample.negative.binomial

Concurrent sampling from multiple negative binomial distributions with parameters \*k\* (failure limit) and \*p\* (failure probability).

# **Description**

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## **Arguments**

k	NDArray-or-Symbol Limits of unsuccessful experiments.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
р	NDArray-or-Symbol Failure probabilities in each experiment.

# **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

Samples will always be returned as a floating point data type.

# Examples::

```
k = [20, 49] p = [0.4, 0.77]
```

// Draw a single sample for each distribution sample\_negative\_binomial(k, p) = [15., 16.]

// Draw a vector containing two samples for each distribution sample\_negative\_binomial(k, p, shape=(2)) = [[ 15., 50.], [ 16., 12.]]

Defined in src/operator/random/multisample\_op.cc:L289

#### Value

230 mx.nd.sample.normal

mx.nd.sample.normal	Concurrent sampling from multiple normal distributions with param-
	eters *mu* (mean) and *sigma* (standard deviation).

# **Description**

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# **Arguments**

mu	NDArray-or-Symbol Means of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
sigma	NDArray-or-Symbol Standard deviations of the distributions.

#### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

### Examples::

```
mu = [0.0, 2.5] sigma = [1.0, 3.7]
```

// Draw a single sample for each distribution sample\_normal(mu, sigma) = [-0.56410581, 0.95934606]

// Draw a vector containing two samples for each distribution sample\_normal(mu, sigma, shape=(2)) = [[-0.56410581, 0.2928229], [ 0.95934606, 4.48287058]]

Defined in src/operator/random/multisample\_op.cc:L279

# Value

mx.nd.sample.poisson 231

mx.nd.sample.poisson Concurrent sampling from multiple Poisson distributions with parameters lambda (rate).

# Description

The parameters of the distributions are provided as an input array. Let \*[s]\* be the shape of the input array, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# **Arguments**

lam	NDArray-or-Symbol Lambda (rate) parameters of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

#### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input array, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input value at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input array.

Samples will always be returned as a floating point data type.

### Examples::

```
lam = [1.0, 8.5]
```

// Draw a single sample for each distribution sample\_poisson(lam) = [0., 13.]

// Draw a vector containing two samples for each distribution sample\_poisson(lam, shape=(2)) = [[ 0, 4.], [ 13., 8.]]

Defined in src/operator/random/multisample\_op.cc:L286

## Value

232 mx.nd.sample.uniform

 $\begin{tabular}{ll} mx.nd.sample.uniform & Concurrent sampling from multiple uniform distributions on the intervals given by $$\{low,high)$*. \\ \end{tabular}$ 

# Description

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# **Arguments**

low	NDArray-or-Symbol Lower bounds of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
high	NDArray-or-Symbol Upper bounds of the distributions.

#### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

#### Examples::

```
low = [0.0, 2.5] high = [1.0, 3.7]
```

// Draw a single sample for each distribution sample\_uniform(low, high) = [ 0.40451524, 3.18687344]

// Draw a vector containing two samples for each distribution sample\_uniform(low, high, shape=(2)) = [[ 0.40451524, 0.18017688], [ 3.18687344, 3.68352246]]

Defined in src/operator/random/multisample\_op.cc:L277

## Value

mx.nd.save 233

mx.nd.save

Save an mx.nd.array object

# **Description**

Save an mx.nd.array object

# Usage

```
mx.nd.save(ndarray, filename)
```

## **Arguments**

ndarray the mx.nd.array object

filename (including the path)

# **Examples**

```
mat = mx.nd.array(1:3)
mx.nd.save(mat, 'temp.mat')
mat2 = mx.nd.load('temp.mat')
as.array(mat)
as.array(mat2[[1]])
```

mx.nd.scatter.nd

Scatters data into a new tensor according to indices.

# Description

Given 'data' with shape ' $(Y_0, ..., Y_{K-1}, X_M, ..., X_{N-1})$ ' and indices with shape ' $(M, Y_0, ..., Y_{K-1})$ ', the output will have shape ' $(X_0, X_1, ..., X_{N-1})$ ', where ' $M \le N$ '. If 'M == N', data shape should simply be ' $(Y_0, ..., Y_{K-1})$ '.

# Arguments

data	NDArray-or-Symbol data
indices	NDArray-or-Symbol indices

shape Shape(tuple), required Shape of output.

234 mx.nd.SequenceLast

### **Details**

The elements in output is defined as follows::

```
output[indices[0, y_0, ..., y_K-1], ..., indices[M-1, y_0, ..., y_K-1], x_M, ..., x_N-1] = data[y_0, ..., y_K-1, x_M, ..., x_N-1]
```

all other entries in output are 0.

.. warning::

If the indices have duplicates, the result will be non-deterministic and the gradient of 'scatter\_nd' will not be correct!!

#### Examples::

```
data = [2, 3, 0] indices = [[1, 1, 0], [0, 1, 0]] shape = (2, 2) scatter_nd(data, indices, shape) = [[0, 0], [2, 3]]
```

data = [[[1, 2], [3, 4]], [[5, 6], [7, 8]]] indices = [[0, 1], [1, 1]] shape = (2, 2, 2, 2) scatter\_nd(data, indices, shape) = [[[[0, 0], [0, 0]],

[[1, 2], [3, 4]]],

[[[0, 0], [0, 0]],

[[5, 6], [7, 8]]]]

#### Value

out The result mx.ndarray

mx.nd.SequenceLast

Takes the last element of a sequence.

# Description

This function takes an n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] and returns a (n-1)-dimensional array of the form [batch\_size, other\_feature\_dims].

### **Arguments**

data NDArray-or-Sy

NDArray-or-Symbol n-dimensional input array of the form [max\_sequence\_length,

batch\_size, other\_feature\_dims] where n>2

sequence.length

NDArray-or-Symbol vector of sequence lengths of the form [batch\_size]

use.sequence.length

boolean, optional, default=0 If set to true, this layer takes in an extra input pa-

rameter 'sequence\_length' to specify variable length sequence

axis int, optional, default='0' The sequence axis. Only values of 0 and 1 are currently

supported.

mx.nd.SequenceMask

### **Details**

Parameter 'sequence\_length' is used to handle variable-length sequences. 'sequence\_length' should be an input array of positive ints of dimension [batch\_size]. To use this parameter, set 'use\_sequence\_length' to 'True', otherwise each example in the batch is assumed to have the max sequence length.

235

.. note:: Alternatively, you can also use 'take' operator.

### Example::

```
x = [[[ 1., 2., 3.], [ 4., 5., 6.], [ 7., 8., 9.]],

[[ 10., 11., 12.], [ 13., 14., 15.], [ 16., 17., 18.]],

[[ 19., 20., 21.], [ 22., 23., 24.], [ 25., 26., 27.]]]

// returns last sequence when sequence_length parameter is not used SequenceLast(x) = [[ 19., 20., 21.], [ 22., 23., 24.], [ 25., 26., 27.]]

// sequence_length is used SequenceLast(x, sequence_length=[1,1,1], use_sequence_length=True)

= [[ 1., 2., 3.], [ 4., 5., 6.], [ 7., 8., 9.]]

// sequence_length is used SequenceLast(x, sequence_length=[1,2,3], use_sequence_length=True)

= [[ 1., 2., 3.], [ 13., 14., 15.], [ 25., 26., 27.]]

Defined in src/operator/sequence_last.cc:L106
```

#### Value

out The result mx.ndarray

supported.

mx.nd.SequenceMask

Sets all elements outside the sequence to a constant value.

# **Description**

This function takes an n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] and returns an array of the same shape.

#### **Arguments**

data	NDArray-or-Symbol n-dimensional input array of the form [max_sequence_length, batch_size, other_feature_dims] where n>2 $$	
sequence.length		
	NDArray-or-Symbol vector of sequence lengths of the form [batch_size]	
use.sequence.length		
	boolean, optional, default=0 If set to true, this layer takes in an extra input parameter 'sequence_length' to specify variable length sequence	
value	float, optional, default=0 The value to be used as a mask.	
axis	int, optional, default='0' The sequence axis. Only values of 0 and 1 are currently	

#### **Details**

Parameter 'sequence\_length' is used to handle variable-length sequences. 'sequence\_length' should be an input array of positive ints of dimension [batch\_size]. To use this parameter, set 'use\_sequence\_length' to 'True', otherwise each example in the batch is assumed to have the max sequence length and this operator works as the 'identity' operator.

# Example::

```
x = [[[ 1., 2., 3.], [ 4., 5., 6.]],

[[ 7., 8., 9.], [ 10., 11., 12.]],

[[ 13., 14., 15.], [ 16., 17., 18.]]]

// Batch 1 B1 = [[ 1., 2., 3.], [ 7., 8., 9.], [ 13., 14., 15.]]

// Batch 2 B2 = [[ 4., 5., 6.], [ 10., 11., 12.], [ 16., 17., 18.]]
```

// works as identity operator when sequence\_length parameter is not used SequenceMask(x) = [[[1, 2., 3.], [4., 5., 6.]],

```
[[ 7., 8., 9.], [ 10., 11., 12.]],
[[ 13., 14., 15.], [ 16., 17., 18.]]]
```

// sequence\_length [1,1] means 1 of each batch will be kept // and other rows are masked with default mask value = 0 SequenceMask(x, sequence\_length=[1,1], use\_sequence\_length=True) = [[[ 1., 2., 3.], [ 4., 5., 6.]],

```
[[0., 0., 0.], [0., 0., 0.]],
[[0., 0., 0.], [0., 0., 0.]]]
```

// sequence\_length [2,3] means 2 of batch B1 and 3 of batch B2 will be kept // and other rows are masked with value = 1 SequenceMask(x, sequence\_length=[2,3], use\_sequence\_length=True, value=1) = [[[1., 2., 3.], [4., 5., 6.]],

```
[[ 7., 8., 9.], [ 10., 11., 12.]],
[[ 1., 1., 1.], [ 16., 17., 18.]]]
```

Defined in src/operator/sequence\_mask.cc:L186

#### Value

out The result mx.ndarray

mx.nd.SequenceReverse Reverses the elements of each sequence.

# **Description**

This function takes an n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] and returns an array of the same shape.

## **Arguments**

data NDArray-or-Symbol n-dimensional input array of the form [max\_sequence\_length, batch\_size, other dims] where n>2

sequence.length

NDArray-or-Symbol vector of sequence lengths of the form [batch\_size]

use.sequence.length

boolean, optional, default=0 If set to true, this layer takes in an extra input parameter 'sequence\_length' to specify variable length sequence

axis int, optional, default='0' The sequence axis. Only 0 is currently supported.

#### **Details**

Parameter 'sequence\_length' is used to handle variable-length sequences. 'sequence\_length' should be an input array of positive ints of dimension [batch\_size]. To use this parameter, set 'use\_sequence\_length' to 'True', otherwise each example in the batch is assumed to have the max sequence length.

## Example::

```
x = [[[1., 2., 3.], [4., 5., 6.]],
```

[[ 7., 8., 9.], [ 10., 11., 12.]],

[[ 13., 14., 15.], [ 16., 17., 18.]]]

// Batch 1 B1 = [[ 1., 2., 3.], [ 7., 8., 9.], [ 13., 14., 15.]]

// Batch 2 B2 = 
$$[[4., 5., 6.], [10., 11., 12.], [16., 17., 18.]]$$

// returns reverse sequence when sequence\_length parameter is not used SequenceReverse(x) = [[[ 13., 14., 15.], [ 16., 17., 18.]],

[[7., 8., 9.], [10., 11., 12.]],

// sequence\_length [2,2] means 2 rows of // both batch B1 and B2 will be reversed. SequenceReverse(x, sequence\_length=[2,2], use\_sequence\_length=True) = [[[7., 8., 9.], [10., 11., 12.]],

// sequence\_length [2,3] means 2 of batch B2 and 3 of batch B3 // will be reversed. SequenceReverse(x, sequence\_length=[2,3], use\_sequence\_length=True) = [[[7., 8., 9.], [16., 17., 18.]],

Defined in src/operator/sequence\_reverse.cc:L122

### Value

## **Description**

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

## **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient mom NDArray-or-Symbol Momentum 1r float, required Learning rate

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row\_sparse and both weight and momentum have the same stype

## **Details**

.. math::

 $v_1 = \alpha v_t - 1 - \alpha v_t - 1$ 

v = momentum \* v - learning\_rate \* gradient weight += v

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

However, if grad's storage type is "row\_sparse", "lazy\_update" is True and weight's storage type is the same as momentum's storage type, only the row slices whose indices appear in grad.indices are updated (for both weight and momentum)::

for row in gradient.indices:  $v[row] = momentum[row] * v[row] - learning_rate * gradient[row] weight[row] += v[row]$ 

Defined in src/operator/optimizer\_op.cc:L556

## Value

mx.nd.sgd.update 239

mx.nd.sgd.update	Update function for Stochastic Gradient Descent (SGD) optimizer.

# Description

It updates the weights using::

# Arguments

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
lr	float, required Learning rate
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
lazy.update	boolean, optional, default=1 If true, lazy updates are applied if gradient's stype is row_sparse.

# **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
```

However, if gradient is of "row\_sparse" storage type and "lazy\_update" is True, only the row slices whose indices appear in grad.indices are updated::

for row in gradient.indices: weight[row] = weight[row] - learning\_rate \* (gradient[row] + wd \* weight[row])

Defined in src/operator/optimizer\_op.cc:L515

# Value

240 mx.nd.shuffle

mx.nd.shape.array

Returns a 1D int64 array containing the shape of data.

# Description

Example::

# **Arguments**

data

NDArray-or-Symbol Input Array.

# **Details**

```
shape_array([[1,2,3,4], [5,6,7,8]]) = [2,4]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L574

### Value

out The result mx.ndarray

mx.nd.shuffle

Randomly shuffle the elements.

# Description

This shuffles the array along the first axis. The order of the elements in each subarray does not change. For example, if a 2D array is given, the order of the rows randomly changes, but the order of the elements in each row does not change.

# **Arguments**

data

NDArray-or-Symbol Data to be shuffled.

### Value

mx.nd.sigmoid 241

mx.nd.sigmoid

Computes sigmoid of x element-wise.

# **Description**

```
.. math:: y = 1 / (1 + \exp(-x))
```

# **Arguments**

data

NDArray-or-Symbol The input array.

### **Details**

The storage type of "sigmoid" output is always dense
Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L119

#### Value

out The result mx.ndarray

mx.nd.sign

Returns element-wise sign of the input.

# Description

Example::

# Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

```
sign([-2, 0, 3]) = [-1, 0, 1]
```

The storage type of "sign" output depends upon the input storage type:

-  $sign(default) = default - sign(row\_sparse) = row\_sparse - sign(csr) = csr$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L759

# Value

242 mx.nd.signum.update

# **Description**

.. math::

### **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient lr float, required Learning rate

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

# **Details**

```
\begin{split} g\_t &= \Lambda J(W_t-1) \land W_t = W_t-1 - \epsilon_t \land (g_t) \end{split} It updates the weights using:: weight = weight - learning_rate * sign(gradient)
```

.. note:: - sparse ndarray not supported for this optimizer yet.

Defined in src/operator/optimizer\_op.cc:L63

#### Value

out The result mx.ndarray

mx.nd.signum.update SIGN momentUM (Signum) optimizer.

### Description

.. math::

mx.nd.sin 243

#### **Arguments**

weight NDArray-or-Symbol Weight
grad NDArray-or-Symbol Gradient
mom NDArray-or-Symbol Momentum
lr float, required Learning rate

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

wd.1h float, optional, default=0 The amount of weight decay that does not go into gra-

dient/momentum calculationsotherwise do weight decay algorithmically only.

#### **Details**

 $g_t = \Lambda J(W_t-1) \ m_t = \beta m_t-1 + (1 - \beta g_t \ W_t = W_t-1 - \beta g_t \ W_t = W_t-1 - \beta g_t \ W_t$ 

It updates the weights using:: state = momentum \* state + (1-momentum) \* gradient weight = weight - learning\_rate \* sign(state)

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

.. note:: - sparse ndarray not supported for this optimizer yet.

Defined in src/operator/optimizer\_op.cc:L92

#### Value

out The result mx.ndarray

mx.nd.sin

Computes the element-wise sine of the input array.

#### **Description**

The input should be in radians (:math: '2\pi' rad equals 360 degrees).

### **Arguments**

data NDArray-or-Symbol The input array.

244 mx.nd.size.array

# **Details**

```
.. math:: \sin([0, \pi/4, \pi/2]) = [0, 0.707, 1]
```

The storage type of "sin" output depends upon the input storage type:

- sin(default) = default - sin(row\_sparse) = row\_sparse - sin(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L47

### Value

out The result mx.ndarray

mx.nd.sinh

Returns the hyperbolic sine of the input array, computed element-wise.

# Description

```
.. math:: sinh(x) = 0.5 \times (exp(x) - exp(-x))
```

# **Arguments**

data

NDArray-or-Symbol The input array.

# **Details**

The storage type of "sinh" output depends upon the input storage type:

- sinh(default) = default - sinh(row\_sparse) = row\_sparse - sinh(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L371

#### Value

out The result mx.ndarray

mx.nd.size.array

Returns a 1D int64 array containing the size of data.

# Description

Example::

# **Arguments**

data

NDArray-or-Symbol Input Array.

mx.nd.slice.axis 245

# **Details**

```
size\_array([[1,2,3,4], [5,6,7,8]]) = [8]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L625

### Value

out The result mx.ndarray

mx.nd.slice.axis	Slices along a given axis. Returns an array slice along a given 'axis'
	starting from the 'begin' index to the 'end' index. Examples:: $x = [[$
	1., 2., 3., 4.], [5., 6., 7., 8.], [9., 10., 11., 12.]] slice_axis(x, axis=0,
	$begin=1, end=3) = [[5., 6., 7., 8.], [9., 10., 11., 12.]] slice_axis(x, 0.)$
	$axis=1, begin=0, end=2) = [[1., 2.], [5., 6.], [9., 10.]] slice_axis(x, x, x)$
	axis=1, begin=-3, end=-1) = [[2, 3, ], [6, 7, ], [10, 11, ]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L571

# Arguments

data	NDArray-or-Symbol Source input
axis	int, required Axis along which to be sliced, supports negative indexes.
begin	int, required The beginning index along the axis to be sliced, supports negative indexes.
end	int or None, required The ending index along the axis to be sliced, supports negative indexes.

# Value

246 mx.nd.slice.like

mx.nd.slice.like

Slices a region of the array like the shape of another array. This function is similar to "slice", however, the 'begin' are always '0's and 'end' of specific axes are inferred from the second input 'shape\_like'. Given the second 'shape\_like' input of "shape= $(d_0, d_1, ..., d_{n-1})$ ", a "slice like" operator with default empty 'axes', it performs the following operation: " out =  $slice(input, begin=(0, 0, ..., 0), end=(d_0, 0, ..., 0)$  $d_1, ..., d_{n-1}$ )". When 'axes' is not empty, it is used to speficy which axes are being sliced. Given a 4-d input data, "slice\_like" operator with "axes=(0, 2, -1)" will perform the following operation: " out =  $slice(input, begin=(0, 0, 0, 0), end=(d_0, None, d_2, d_3))$ ". Note that it is allowed to have first and second input with different dimensions, however, you have to make sure the 'axes' are specified and not exceeding the dimension limits. For example, given 'input 1' with "shape=(2,3,4,5)" and 'input\_2' with "shape=(1,2,3)", it is not allowed to use: "out = slice\_like(a, b)" because ndim of 'input\_1' is 4, and ndim of 'input\_2' is 3. The following is allowed in this situation: "  $out = slice\_like(a, b, axes=(0, 2))$ " Example:: x = [[1., 2., 3., 4.],[5., 6., 7., 8.], [9., 10., 11., 12.]] y = [[0., 0., 0.], [0., 0., 0.]] $slice\_like(x, y) = [[1., 2., 3.] [5., 6., 7.]] slice\_like(x, y, axes=(0, 1))$ = [[1., 2., 3.] [5., 6., 7.]] slice\_like(x, y, axes=(0)) = [[1., 2., 3., 4.]][5., 6., 7., 8.] slice\_like(x, y, axes=(-1)) = [[1., 2., 3.] [5., 6., 7.]9., 10., 11.]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L625

#### **Arguments**

data NDArray-or-Symbol Source input

shape.like NDArray-or-Symbol Shape like input

axes Shape(tuple), optional, default=[] List of axes on which input data will be sliced

according to the corresponding size of the second input. By default will slice on

all axes. Negative axes are supported.

#### Value

mx.nd.SliceChannel 247

mx.nd.SliceChannel

Splits an array along a particular axis into multiple sub-arrays.

### **Description**

.. note:: "SliceChannel" is deprecated. Use "split" instead.

### **Arguments**

data NDArray-or-Symbol The input

num.outputs int, required Number of splits. Note that this should evenly divide the length of

the 'axis'.

axis int, optional, default='1' Axis along which to split.

squeeze.axis boolean, optional, default=0 If true, Removes the axis with length 1 from the

shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "true" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis'

can be set to "true" only if "input.shape[axis] == num\_outputs".

#### **Details**

\*\*Note\*\* that 'num\_outputs' should evenly divide the length of the axis along which to split the array.

# Example::

```
x = [[[ 1.] [ 2.]] [[ 3.] [ 4.]] [[ 5.] [ 6.]]] x.shape = (3, 2, 1)
y = split(x, axis=1, num_outputs=2) // a list of 2 arrays with shape (3, 1, 1) y = [[[ 1.]] [[ 3.]] [[ 5.]]]
[[[ 2.]] [[ 4.]] [[ 6.]]]
y[0].shape = (3, 1, 1)
z = split(x, axis=0, num_outputs=3) // a list of 3 arrays with shape (1, 2, 1) z = [[[ 1.] [ 2.]]]
[[[ 3.] [ 4.]]]
```

[[[5.] [6.]]]z[0].shape = (1, 2, 1)

'squeeze\_axis=1' removes the axis with length 1 from the shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "1" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis' can be set to true only if "input.shape[axis] == num\_outputs".

### Example::

```
z = split(x, axis=0, num\_outputs=3, squeeze\_axis=1) // a list of 3 arrays with shape (2, 1) z = [[1.] [2.]]
```

[[ 3.] [ 4.]]

[[5.] [6.]] z[0].shape = (2,1)

Defined in src/operator/slice\_channel.cc:L107

248 mx.nd.Softmax

# Value

out The result mx.ndarray

mx.nd.smooth.l1

Calculate Smooth L1 Loss(lhs, scalar) by summing

# **Description**

.. math::

# Arguments

data

NDArray-or-Symbol source input

scalar

float scalar input

### **Details**

 $f(x) = \text{logincases (\sigma x)^2/2,\& \textif } x < 1/\text{sigma^2} |x|-0.5/\text{sigma^2,\& \textotherwise \end-cases}$ 

where :math: 'x' is an element of the tensor \*lhs\* and :math: '\sigma' is the scalar.

Example::

 $smooth_{11}([1, 2, 3, 4]) = [0.5, 1.5, 2.5, 3.5] smooth_{11}([1, 2, 3, 4], scalar=1) = [0.5, 1.5, 2.5, 3.5]$ 

Defined in src/operator/tensor/elemwise\_binary\_scalar\_op\_extended.cc:L108

## Value

out The result mx.ndarray

mx.nd.Softmax

Computes the gradient of cross entropy loss with respect to softmax output.

# **Description**

- This operator computes the gradient in two steps. The cross entropy loss does not actually need to be computed.

mx.nd.Softmax 249

#### **Arguments**

data NDArray-or-Symbol Input array.

label NDArray-or-Symbol Ground truth label.

grad. scale float, optional, default=1 Scales the gradient by a float factor.

ignore.label float, optional, default=-1 The instances whose 'labels' == 'ignore label' will

be ignored during backward, if 'use\_ignore' is set to "true").

multi.output boolean, optional, default=0 If set to "true", the softmax function will be com-

puted along axis "1". This is applied when the shape of input array differs from

the shape of label array.

use.ignore boolean, optional, default=0 If set to "true", the 'ignore\_label' value will not

contribute to the backward gradient.

preserve.shape boolean, optional, default=0 If set to "true", the softmax function will be com-

puted along the last axis ("-1").

normalization 'batch', 'null', 'valid', optional, default='null' Normalizes the gradient.

out.grad boolean, optional, default=0 Multiplies gradient with output gradient element-

wise.

smooth.alpha float, optional, default=0 Constant for computing a label smoothed version of

cross-entropyfor the backwards pass. This constant gets subtracted from theonehot encoding of the gold label and distributed uniformly to all other labels.

#### **Details**

- Applies softmax function on the input array. - Computes and returns the gradient of cross entropy loss w.r.t. the softmax output.

- The softmax function, cross entropy loss and gradient is given by:
- Softmax Function:
- .. math::  $\text{textsoftmax}(x)_i = \frac{x_i}{x_j} \exp(x_j)$
- Cross Entropy Function:
- .. math:: \textCE(label, output) = \sum\_i \textlabel\_i \log(\textoutput\_i)
- The gradient of cross entropy loss w.r.t softmax output:
- .. math:: \textgradient = \textoutput \textlabel
- During forward propagation, the softmax function is computed for each instance in the input array.

For general \*N\*-D input arrays with shape :math:  $(d_1, d_2, ..., d_n)$ . The size is :math:  $s=d_1 \cdot d_2 \cdot d_2 \cdot d_n$ . We can use the parameters 'preserve\_shape' and 'multi\_output' to specify the way to compute softmax:

- By default, 'preserve\_shape' is "false". This operator will reshape the input array into a 2-D array with shape :math: ' $(d_1, \frac{1}{n})$ ' and then compute the softmax function for each row in the reshaped array, and afterwards reshape it back to the original shape :math: ' $(d_1, d_2, ..., d_n)$ '.
- If 'preserve\_shape' is "true", the softmax function will be computed along the last axis ('axis' = "-1"). If 'multi\_output' is "true", the softmax function will be computed along the second axis ('axis' = "1").

250 mx.nd.softmax

- During backward propagation, the gradient of cross-entropy loss w.r.t softmax output array is computed. The provided label can be a one-hot label array or a probability label array.

- If the parameter 'use\_ignore' is "true", 'ignore\_label' can specify input instances with a particular label to be ignored during backward propagation. \*\*This has no effect when softmax 'output' has same shape as 'label'\*\*.

### Example::

 $\begin{array}{l} data = [[1,2,3,4],[2,2,2,2],[3,3,3,3],[4,4,4,4]] \ label = [1,0,2,3] \ ignore\_label = 1 \ SoftmaxOutput(data=data, label = label, multi\_output=true, use\_ignore=true, lignore\_label=ignore\_label) \ \# \ forward \ softmax \ output \ [[0.0320586\ 0.08714432\ 0.23688284\ 0.64391428]\ [0.25\ 0.25\ 0.25\ 0.25\ 0.25\ ]\ [0.25\ 0.$ 

- The parameter 'grad\_scale' can be used to rescale the gradient, which is often used to give each loss function different weights.
- This operator also supports various ways to normalize the gradient by 'normalization', The 'normalization' is applied if softmax output has different shape than the labels. The 'normalization' mode can be set to the followings:
- "'null'": do nothing. "'batch'": divide the gradient by the batch size. "'valid'": divide the gradient by the number of instances which are not ignored.

Defined in src/operator/softmax\_output.cc:L231

#### Value

out The result mx.ndarray

## **Description**

The resulting array contains elements in the range (0,1) and the elements along the given axis sum up to 1.

### **Arguments**

data	NDArray-or-Symbol The input array.
length	NDArray-or-Symbol The length array.
axis	int, optional, default='-1' The axis along which to compute softmax.
temperature	double or None, optional, default=None Temperature parameter in softmax
dtype	None, 'float16', 'float32', 'float64',optional, default='None' DType of the output in case this can't be inferred. Defaults to the same as input's dtype if not defined (dtype=None).
use.length	boolean or None, optional, default=0 Whether to use the length input as a mask over the data input.

### **Details**

```
.. math:: softmax(\mathbfz/t)_j = \frace^z_j/t\sum_k=1^K e^z_k/t for :math: 'j = 1, ..., K' t is the temperature parameter in softmax function. By default, t equals 1.0 Example:: x = [[\ 1.\ 1.\ 1.]\ [\ 1.\ 1.\ 1.]] softmax(x,axis=0) = [[\ 0.5\ 0.5\ 0.5]\ [\ 0.5\ 0.5\ 0.5]] softmax(x,axis=1) = [[\ 0.33333334, 0.3333334, 0.3333334], [\ 0.33333334, 0.33333334], [\ 0.33333334, 0.33333334]] Defined in src/operator/nn/softmax.cc:L134
```

#### Value

out The result mx.ndarray

```
mx.nd.softmax.cross.entropy
```

Calculate cross entropy of softmax output and one-hot label.

# Description

- This operator computes the cross entropy in two steps: - Applies softmax function on the input array. - Computes and returns the cross entropy loss between the softmax output and the labels.

## **Arguments**

data NDArray-or-Symbol Input data label NDArray-or-Symbol Input label

#### **Details**

- The softmax function and cross entropy loss is given by:
- Softmax Function:
- .. math::  $\text{textsoftmax}(x)_i = \frac{x_i}{\exp(x_i)} \sup_j \exp(x_j)$
- Cross Entropy Function:
- .. math:: \textCE(label, output) = \sum\_i \textlabel\_i \log(\textoutput\_i)

### Example::

```
Example:  x = [[1, 2, 3], [11, 7, 5]]   label = [2, 0]   softmax(x) = [[0.09003057, 0.24472848, 0.66524094], [0.97962922, 0.01794253, 0.00242826]]   softmax\_cross\_entropy(data, label) = -\log(0.66524084) - \log(0.97962922) = 0.4281871  Defined in src/operator/loss\_binary\_op.cc:L59
```

252 mx.nd.SoftmaxActivation

### Value

out The result mx.ndarray

mx.nd.SoftmaxActivation

Applies softmax activation to input. This is intended for internal layers.

# Description

.. note::

### **Arguments**

data NDArray-or-Symbol The input array.

mode 'channel', 'instance', optional, default='instance' Specifies how to compute the

softmax. If set to "instance", it computes softmax for each instance. If set to "channel", It computes cross channel softmax for each position of each instance.

## **Details**

This operator has been deprecated, please use 'softmax'.

If 'mode' = "instance", this operator will compute a softmax for each instance in the batch. This is the default mode.

If 'mode' = "channel", this operator will compute a k-class softmax at each position of each instance, where 'k' = "num\_channel". This mode can only be used when the input array has at least 3 dimensions. This can be used for 'fully convolutional network', 'image segmentation', etc.

## Example::

»> input\_array = mx.nd.array([[3., 0.5, -0.5, 2., 7.], »> [2., -4, 7., 3., 0.2]]) »> softmax\_act =
mx.nd.SoftmaxActivation(input\_array) »> print softmax\_act.asnumpy() [[ 1.78322066e-02 1.46375655e03 5.38485940e-04 6.56010211e-03 9.73605454e-01] [ 6.56221947e-03 5.95310994e-04 9.73919690e01 1.78379621e-02 1.08472735e-03]]

Defined in src/operator/nn/softmax\_activation.cc:L59

### Value

mx.nd.SoftmaxOutput 253

mx.nd.SoftmaxOutput	Computes the gradient of cross entropy loss with respect to softmax output.
---------------------	---

# Description

- This operator computes the gradient in two steps. The cross entropy loss does not actually need to be computed.

## **Arguments**

data	NDArray-or-Symbol Input array.
label	NDArray-or-Symbol Ground truth label.
grad.scale	float, optional, default=1 Scales the gradient by a float factor.
ignore.label	float, optional, default=-1 The instances whose 'labels' == 'ignore_label' will be ignored during backward, if 'use_ignore' is set to "true").
multi.output	boolean, optional, default=0 If set to "true", the softmax function will be computed along axis "1". This is applied when the shape of input array differs from the shape of label array.
use.ignore	boolean, optional, default=0 If set to "true", the 'ignore_label' value will not contribute to the backward gradient.
preserve.shape	boolean, optional, default=0 If set to "true", the softmax function will be computed along the last axis ("-1").
normalization	'batch', 'null', 'valid', optional, default='null' Normalizes the gradient.
out.grad	boolean, optional, default=0 Multiplies gradient with output gradient elementwise.
smooth.alpha	float, optional, default=0 Constant for computing a label smoothed version of cross-entropyfor the backwards pass. This constant gets subtracted from theonehot encoding of the gold label and distributed uniformly toall other labels.

### **Details**

- Applies softmax function on the input array. Computes and returns the gradient of cross entropy loss w.r.t. the softmax output.
- The softmax function, cross entropy loss and gradient is given by:
- Softmax Function:
- .. math::  $\textsoftmax(x)_i = \textsoftmax(x_i) \textsoftmax(x_j)$
- Cross Entropy Function:
- .. math:: \textCE(label, output) = \sum\_i \textlabel\_i \log(\textoutput\_i)
- The gradient of cross entropy loss w.r.t softmax output:
- .. math:: \textgradient = \textoutput \textlabel

254 mx.nd.softmin

- During forward propagation, the softmax function is computed for each instance in the input array. For general \*N\*-D input arrays with shape :math: '(d\_1, d\_2, ..., d\_n)'. The size is :math: 's=d\_1 \cdot d\_2 \cdot \cdot \cdot d\_n'. We can use the parameters 'preserve\_shape' and 'multi\_output' to specify the way to compute softmax:

- By default, 'preserve\_shape' is "false". This operator will reshape the input array into a 2-D array with shape :math: '(d\_1, \fracsd\_1)' and then compute the softmax function for each row in the reshaped array, and afterwards reshape it back to the original shape :math: '(d\_1, d\_2, ..., d\_n)'.
   If 'preserve\_shape' is "true", the softmax function will be computed along the last axis ('axis' = "-1"). If 'multi\_output' is "true", the softmax function will be computed along the second axis ('axis' = "1").
- During backward propagation, the gradient of cross-entropy loss w.r.t softmax output array is computed. The provided label can be a one-hot label array or a probability label array.
- If the parameter 'use\_ignore' is "true", 'ignore\_label' can specify input instances with a particular label to be ignored during backward propagation. \*\*This has no effect when softmax 'output' has same shape as 'label'\*\*.

#### Example::

- The parameter 'grad\_scale' can be used to rescale the gradient, which is often used to give each loss function different weights.
- This operator also supports various ways to normalize the gradient by 'normalization', The 'normalization' is applied if softmax output has different shape than the labels. The 'normalization' mode can be set to the followings:
- "'null'": do nothing. "'batch'": divide the gradient by the batch size. "'valid'": divide the gradient by the number of instances which are not ignored.

Defined in src/operator/softmax output.cc:L231

### Value

out The result mx.ndarray

mx.nd.softmin

Applies the softmin function.

### **Description**

The resulting array contains elements in the range (0,1) and the elements along the given axis sum up to 1.

mx.nd.softsign 255

### Arguments

dtype

data NDArray-or-Symbol The input array.

axis int, optional, default='-1' The axis along which to compute softmax.

 $temperature \qquad double \ or \ None, \ optional, \ default = None \ Temperature \ parameter \ in \ softmax$ 

None, 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to the same as input's dtype if not

defined (dtype=None).

use.length boolean or None, optional, default=0 Whether to use the length input as a mask

over the data input.

#### **Details**

```
.. math:: softmin(\mathbfz/t)_j = \frace^-z_j/t\sum_k=1^K e^-z_k/t
```

for :math: 'j = 1, ..., K'

t is the temperature parameter in softmax function. By default, t equals 1.0

Example::

$$x = [[1. 2. 3.] [3. 2. 1.]]$$

softmin(x,axis=0) = [[0.88079703, 0.5, 0.11920292], [0.11920292, 0.5, 0.88079703]]

softmin(x,axis=1) = [[0.66524094, 0.24472848, 0.09003057], [0.09003057, 0.24472848, 0.66524094]]

Defined in src/operator/nn/softmin.cc:L57

#### Value

out The result mx.ndarray

mx.nd.softsign

Computes softsign of x element-wise.

#### **Description**

```
.. math:: y = x / (1 + abs(x))
```

### **Arguments**

data NDArray-or-Symbol The input array.

### **Details**

The storage type of "softsign" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L191

## Value

256 mx.nd.space.to.depth

mx.nd.sort

Returns a sorted copy of an input array along the given axis.

## **Description**

Examples::

### **Arguments**

data NDArray-or-Symbol The input array

axis int or None, optional, default='-1' Axis along which to choose sort the input

tensor. If not given, the flattened array is used. Default is -1.

is.ascend boolean, optional, default=1 Whether to sort in ascending or descending order.

#### Details

```
x = [[1, 4], [3, 1]]
// sorts along the last axis sort(x) = [[1., 4.], [1., 3.]]
// flattens and then sorts sort(x, axis=None) = [1., 1., 3., 4.]
// sorts along the first axis sort(x, axis=0) = [[1., 1.], [3., 4.]]
// in a descend order sort(x, is_ascend=0) = [[4., 1.], [3., 1.]]
Defined in src/operator/tensor/ordering_op.cc:L133
```

#### Value

out The result mx.ndarray

mx.nd.space.to.depth

*Rearranges*(permutes) blocks of spatial data into SpaceToDepth depth. ONNXSimilar tooperator: https://github.com/onnx/onnx/blob/master/docs/Operators.md#SpaceToDepth The output is a new tensor where the values from height and width dimension are moved to the depth dimension. The reverse of this operation is "depth\_to\_space". .. math:: \begingather\* x \prime = reshape(x, [N, C, H / block size, block size, W / block size, $block\_size$ ]) \ x \prime \prime =  $transpose(x \prime, [0, 3, 5, 1,$ 2, 4])  $\ y = reshape(x \rangle prime \rangle prime, [N, C * (block size ^ 2), H / (block size ^ 2)$  $block \subseteq W / block \subseteq Size]) \endgather* where :math: `x` is an input$ tensor with default layout as :math: '[N, C, H, W]': [batch, channels, height, width] and :math: 'y' is the output tensor of layout :math: '[N,  $C * (block \ size \ 2), H/block \ size, W/block \ size] `Example:: x =$ [[[0, 6, 1, 7, 2, 8], [12, 18, 13, 19, 14, 20], [3, 9, 4, 10, 5, 11], [15, 21, 16, 22, 17, 23]]]  $space\_to\_depth(x, 2) = [[[[0, 1, 2], [3, 4, 5]],$ [[6, 7, 8], [9, 10, 11]], [[12, 13, 14], [15, 16, 17]], [[18, 19, 20], [21, 22, 23]]]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L1019

# Arguments

data NDArray-or-Symbol Input ndarray

block.size int, required Blocks of [block\_size. block\_size] are moved

### Value

out The result mx.ndarray

mx.nd.SpatialTransformer

Applies a spatial transformer to input feature map.

## **Description**

Applies a spatial transformer to input feature map.

## **Arguments**

data NDArray-or-Symbol Input data to the SpatialTransformerOp.

loc NDArray-or-Symbol localisation net, the output dim should be 6 when trans-

form\_type is affine. You shold initialize the weight and bias with identity tran-

form.

target.shape Shape(tuple), optional, default=[0,0] output shape(h, w) of spatial transformer:

(y, x)

transform.type 'affine', required transformation type

sampler.type 'bilinear', required sampling type

cudnn.off boolean or None, optional, default=None whether to turn cudnn off

### Value

258 mx.nd.split

mx.nd.split	Splits an array along a particular axis into multiple sub-arrays.

### **Description**

.. note:: "SliceChannel" is deprecated. Use "split" instead.

### **Arguments**

data NDArray-or-Symbol The input

num.outputs int, required Number of splits. Note that this should evenly divide the length of

the 'axis'.

axis int, optional, default='1' Axis along which to split.

squeeze.axis boolean, optional, default=0 If true, Removes the axis with length 1 from the

shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "true" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis'

can be set to "true" only if "input.shape[axis] == num\_outputs".

#### **Details**

\*\*Note\*\* that 'num\_outputs' should evenly divide the length of the axis along which to split the array.

## Example::

```
 \begin{aligned} x &= & [[[\ 1.]\ [\ 2.]]\ [[\ 3.]\ [\ 4.]]\ [[\ 5.]\ [\ 6.]]] \ x.shape = (3,\,2,\,1) \\ y &= & \text{split}(x,\,\text{axis}=1,\,\text{num\_outputs}=2)\,\text{//}\ a \ \text{list of 2 arrays with shape } (3,\,1,\,1)\,y = [[[\ 1.]]\ [[\ 3.]]\ [[\ 5.]]] \\ [[[\ 2.]]\ [[\ 4.]]\ [[\ 6.]]] \\ y &= & \text{split}(x,\,\text{axis}=0,\,\text{num\_outputs}=3)\,\text{//}\ a \ \text{list of 3 arrays with shape } (1,\,2,\,1)\,z = [[[\ 1.]\ [\ 2.]]] \\ [[[\ 3.]\ [\ 4.]]] \\ [[[\ 5.]\ [\ 6.]]] \\ z &= & \text{[0].shape} = (1,\,2,\,1) \end{aligned}
```

'squeeze\_axis=1' removes the axis with length 1 from the shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "1" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis' can be set to true only if "input.shape[axis] == num\_outputs".

### Example::

```
z = split(x, axis=0, num_outputs=3, squeeze_axis=1) // a list of 3 arrays with shape (2, 1) z = [[ 1.]
[ 2.]]
[[ 3.] [ 4.]]
[[ 5.] [ 6.]] z[0].shape = (2,1)
```

Defined in src/operator/slice\_channel.cc:L107

mx.nd.sqrt 259

### Value

out The result mx.ndarray

mx.nd.sqrt

Returns element-wise square-root value of the input.

# Description

```
.. math:: \text{textrmsqrt}(x) = \text{sqrt}x
```

### **Arguments**

data

NDArray-or-Symbol The input array.

## **Details**

Example::

$$sqrt([4, 9, 16]) = [2, 3, 4]$$

The storage type of "sqrt" output depends upon the input storage type:

- sqrt(default) = default - sqrt(row\_sparse) = row\_sparse - sqrt(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L170

### Value

out The result mx.ndarray

mx.nd.square

Returns element-wise squared value of the input.

# Description

```
.. math:: square(x) = x^2
```

### **Arguments**

data

NDArray-or-Symbol The input array.

#### **Details**

Example::

$$square([2, 3, 4]) = [4, 9, 16]$$

The storage type of "square" output depends upon the input storage type:

- square(default) = default - square(row\_sparse) = row\_sparse - square(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L119

260 mx.nd.stack

#### Value

out The result mx.ndarray

mx.nd.squeeze

Remove single-dimensional entries from the shape of an array. Same behavior of defining the output tensor shape as numpy.squeeze for the most of cases. See the following note for exception. Examples:: data = [[0], [1], [2]]] squeeze(data) = [0, 1, 2] squeeze(data, axis=0) = [[0], [1], [2]] squeeze(data, axis=2) = [[0, 1, 2]] squeeze(data, axis=(0, 2)) = [0, 1, 2]. Note:: The output of this operator will keep at least one dimension not removed. For example, squeeze([[[4]]]) = [4], while in numpy.squeeze, the output will become a scalar.

#### **Description**

Remove single-dimensional entries from the shape of an array. Same behavior of defining the output tensor shape as numpy.squeeze for the most of cases. See the following note for exception. Examples:: data = [[[0], [1], [2]]] squeeze(data) = [0, 1, 2] squeeze(data, axis=0) = [[0], [1], [2]] squeeze(data, axis=0) = [[0, 1, 2]] squeeze(data, axis=0, 2) = [0, 1, 2]. Note:: The output of this operator will keep at least one dimension not removed. For example, squeeze([[[4]]]) = [4], while in numpy.squeeze, the output will become a scalar.

#### Arguments

data NDArray-or-Symbol data to squeeze

axis Shape or None, optional, default=None Selects a subset of the single-dimensional

entries in the shape. If an axis is selected with shape entry greater than one, an

error is raised.

#### Value

out The result mx.ndarray

mx.nd.stack

Join a sequence of arrays along a new axis. The axis parameter specifies the index of the new axis in the dimensions of the result. For example, if axis=0 it will be the first dimension and if axis=-1 it will be the last dimension. Examples:: x = [1, 2] y = [3, 4] stack(x, y) = [[1, 2], [3, 4]] stack(x, y), axis=1) = [[1, 3], [2, 4]]

### **Description**

Join a sequence of arrays along a new axis. The axis parameter specifies the index of the new axis in the dimensions of the result. For example, if axis=0 it will be the first dimension and if axis=-1 it will be the last dimension. Examples:: x = [1, 2] y = [3, 4] stack(x, y) = [[1, 2], [3, 4]] stack(x, y), axis=1) = [[1, 3], [2, 4]]

mx.nd.stop.gradient 261

## Arguments

data NDArray-or-Symbol[] List of arrays to stack

axis int, optional, default='0' The axis in the result array along which the input arrays

are stacked.

num.args int, required Number of inputs to be stacked.

#### Value

out The result mx.ndarray

mx.nd.stop.gradient Stops gr

Stops gradient computation.

## **Description**

Stops the accumulated gradient of the inputs from flowing through this operator in the backward direction. In other words, this operator prevents the contribution of its inputs to be taken into account for computing gradients.

# Arguments

data

NDArray-or-Symbol The input array.

#### **Details**

# Example::

```
v1 = [1, 2] v2 = [0, 1] a = Variable('a') b = Variable('b') b_stop_grad = stop_gradient(3 * b) loss = MakeLoss(b_stop_grad + a)
```

```
executor = loss.simple_bind(ctx=cpu(), a=(1,2), b=(1,2)) executor.forward(is_train=True, a=v1, b=v2) executor.outputs [ 1. 5.]
```

```
executor.backward() executor.grad_arrays [ 0. 0.] [ 1. 1.]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L326

#### Value

262 mx.nd.sum

mx.nd.sum	Computes the sum of array elements over given axes.	

# Description

.. Note::

## **Arguments**

data	NDArray-or-Symbol The input
axis	Shape or None, optional, default=None The axis or axes along which to perform the reduction.
	The default, 'axis=()', will compute over all elements into a scalar array with shape '(1,)'.
	If 'axis' is int, a reduction is performed on a particular axis.
	If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in the tuple.
	If 'exclude' is true, reduction will be performed on the axes that are NOT in axis instead.
	Negative values means indexing from right to left.
keepdims	boolean, optional, default=0 If this is set to 'True', the reduced axes are left in the result as dimension with size one.
exclude	boolean, optional, default=0 Whether to perform reduction on axis that are NOT

# **Details**

'sum' and 'sum\_axis' are equivalent. For ndarray of csr storage type summation along axis 0 and axis 1 is supported. Setting keepdims or exclude to True will cause a fallback to dense operator.

# Example::

```
data = [[[1, 2], [2, 3], [1, 3]], [[1, 4], [4, 3], [5, 2]], [[7, 1], [7, 2], [7, 3]]] sum(data, axis=1) [[ 4. 8.] [ 10. 9.] [ 21. 6.]] sum(data, axis=[1,2]) [ 12. 19. 27.] data = [[1, 2, 0], [3, 0, 1], [4, 1, 0]] csr = cast_storage(data, 'csr') sum(csr, axis=0) [ 8. 3. 1.] sum(csr, axis=1) [ 3. 4. 5.] Defined in src/operator/tensor/broadcast_reduce_sum_value.cc:L67
```

in axis instead.

#### Value

mx.nd.sum.axis 263

# Description

.. Note::

### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

### Details

'sum' and 'sum\_axis' are equivalent. For ndarray of csr storage type summation along axis 0 and axis 1 is supported. Setting keepdims or exclude to True will cause a fallback to dense operator.

### Example::

```
data = [[[1, 2], [2, 3], [1, 3]], [[1, 4], [4, 3], [5, 2]], [[7, 1], [7, 2], [7, 3]]] sum(data, axis=1) [[ 4. 8.] [ 10. 9.] [ 21. 6.]] sum(data, axis=[1,2]) [ 12. 19. 27.] data = [[1, 2, 0], [3, 0, 1], [4, 1, 0]] csr = cast_storage(data, 'csr') sum(csr, axis=0) [ 8. 3. 1.] sum(csr, axis=1) [ 3. 4. 5.] Defined in src/operator/tensor/broadcast_reduce_sum_value.cc:L67
```

#### Value

264 mx.nd.swapaxes

mx.nd.SVMOutput	Computes support vector machine based transformation of the input.

### **Description**

This tutorial demonstrates using SVM as output layer for classification instead of softmax: https://github.com/dmlc/mxnet/tre

## **Arguments**

data NDArray-or-Symbol Input data for SVM transformation.

NDArray-or-Symbol Class label for the input data.

margin float, optional, default=1 The loss function penalizes outputs that lie outside this margin. Default margin is 1.

regularization.coefficient

float, optional, default=1 Regularization parameter for the SVM. This balances

the tradeoff between coefficient size and error.

use.linear boolean, optional, default=0 Whether to use L1-SVM objective. L2-SVM ob-

jective is used by default.

### Value

out The result mx.ndarray

mx.nd.swapaxes Interchanges two axes of an array.

# Description

Examples::

# **Arguments**

data NDArray-or-Symbol Input array.
dim1 int, optional, default='0' the first axis to be swapped.
dim2 int, optional, default='0' the second axis to be swapped.

## **Details**

```
x = [[1, 2, 3]]) swapaxes(x, 0, 1) = [[1], [2], [3]]

x = [[[0, 1], [2, 3]], [[4, 5], [6, 7]]] // (2,2,2) array

swapaxes(x, 0, 2) = [[[0, 4], [2, 6]], [[1, 5], [3, 7]]]

Defined in src/operator/swapaxis.cc:L70
```

### Value

mx.nd.SwapAxis 265

ray.	Interchanges two axes	mx.nd.SwapAxis
------	-----------------------	----------------

# Description

Examples::

# Arguments

data NDArray-or-Symbol Input array.

dim1 int, optional, default='0' the first axis to be swapped.
dim2 int, optional, default='0' the second axis to be swapped.

# **Details**

```
x = [[1, 2, 3]]) swapaxes(x, 0, 1) = [[1], [2], [3]]

x = [[[0, 1], [2, 3]], [[4, 5], [6, 7]]] // (2,2,2) array

swapaxes(x, 0, 2) = [[[0, 4], [2, 6]], [[1, 5], [3, 7]]]

Defined in src/operator/swapaxis.cc:L70
```

### Value

out The result mx.ndarray

mx.nd.take	Takes elements from an input array along the given axis.	

# Description

This function slices the input array along a particular axis with the provided indices.

# Arguments

a	NDArray-or-Symbol The input array.
indices	NDArray-or-Symbol The indices of the values to be extracted.
axis	int, optional, default='0' The axis of input array to be taken. For input tensor of rank $r$ , it could be in the range of $[-r, r-1]$
mode	'clip', 'raise', 'wrap',optional, default='clip' Specify how out-of-bound indices bahave. Default is "clip". "clip" means clip to the range. So, if all indices mentioned are too large, they are replaced by the index that addresses the last element along an axis. "wrap" means to wrap around. "raise" means to raise an error when index out of range.

266 mx.nd.tan

#### **Details**

Given data tensor of rank  $r \ge 1$ , and indices tensor of rank q, gather entries of the axis dimension of data (by default outer-most one as axis=0) indexed by indices, and concatenates them in an output tensor of rank q + (r - 1).

```
Examples::
```

```
x = [4. 5. 6.]
```

// Trivial case, take the second element along the first axis.

$$take(x, [1]) = [5.]$$

// The other trivial case, axis=-1, take the third element along the first axis

$$take(x, [3], axis=-1, mode='clip') = [6.]$$

$$x = [[1., 2.], [3., 4.], [5., 6.]]$$

// In this case we will get rows 0 and 1, then 1 and 2. Along axis 0

$$take(x, [[0,1],[1,2]]) = [[[1., 2.], [3., 4.]],$$

// In this case we will get rows 0 and 1, then 1 and 2 (calculated by wrapping around). // Along axis 1

```
take(x, [[0, 3], [-1, -2]], axis=1, mode='wrap') = [[[ 1. 2.] [ 2. 1.]]
```

[[ 3. 4.] [ 4. 3.]]

[[ 5. 6.] [ 6. 5.]]]

The storage type of "take" output depends upon the input storage type:

- take(default, default) = default - take(csr, default, axis=0) = csr

Defined in src/operator/tensor/indexing\_op.cc:L782

#### Value

out The result mx.ndarray

mx.nd.tan

Computes the element-wise tangent of the input array.

#### **Description**

The input should be in radians (:math: '2\pi' rad equals 360 degrees).

### **Arguments**

data

NDArray-or-Symbol The input array.

mx.nd.tanh 267

## **Details**

```
.. math:: tan([0, \pi/4, \pi/2]) = [0, 1, -inf]
```

The storage type of "tan" output depends upon the input storage type:

 $-\tan(default) = default - \tan(row\_sparse) = row\_sparse - \tan(csr) = csr$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L140

#### Value

out The result mx.ndarray

mx.nd.tanh

Returns the hyperbolic tangent of the input array, computed elementwise.

# Description

```
.. math:: tanh(x) = sinh(x) / cosh(x)
```

## **Arguments**

data

NDArray-or-Symbol The input array.

## **Details**

The storage type of "tanh" output depends upon the input storage type:

- tanh(default) = default - tanh(row\_sparse) = row\_sparse - tanh(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L451

#### Value

268 mx.nd.topk

mx.nd.tile

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L796

### Arguments

data NDArray-or-Symbol Input data array

reps Shape(tuple), required The number of times for repeating the tensor a. Each dim

size of reps must be a positive integer. If reps has length d, the result will have dimension of  $\max(d, a.ndim)$ ; If a.ndim < d, a is promoted to be d-dimensional by prepending new axes. If a.ndim > d, reps is promoted to a.ndim by pre-

pending 1's to it.

#### Value

out The result mx.ndarray

mx.nd.topk

Returns the indices of the top \*k\* elements in an input array along the given axis (by default). If ret\_type is set to 'value' returns the value of top \*k\* elements (instead of indices). In case of ret\_type = 'both', both value and index would be returned. The returned elements will be sorted.

### Description

Examples::

mx.nd.transpose 269

### **Arguments**

data	NDArray-or-Symbol The input array
axis	int or None, optional, default='-1' Axis along which to choose the top k indices. If not given, the flattened array is used. Default is -1.
k	int, optional, default='1' Number of top elements to select, should be always smaller than or equal to the element number in the given axis. A global sort is performed if set $k < 1$ .
ret.typ	'both', 'indices', 'mask', 'value',optional, default='indices' The return type. "value" means to return the top k values, "indices" means to return the indices of the top k values, "mask" means to return a mask array containing 0 and 1. 1 means the top k values. "both" means to return a list of both values and indices of top k elements.
is.ascend	boolean, optional, default=0 Whether to choose k largest or k smallest elements. Top K largest elements will be chosen if set to false.
dtype	'float16', 'float32', 'float64', 'int32', 'int64', 'uint8', optional, default='float32' DType of the output indices when ret_typ is "indices" or "both". An error will be raised if the selected data type cannot precisely represent the indices.

#### **Details**

```
 x = [[\ 0.3,\ 0.2,\ 0.4],\ [\ 0.1,\ 0.3,\ 0.2]]  // returns an index of the largest element on last axis topk(x) = [[\ 2.], [\ 1.]] 
 // returns the value of top-2 largest elements on last axis topk(x, ret_typ='value', k=2) = [[\ 0.4,\ 0.3], [\ 0.3,\ 0.2]] 
 // returns the value of top-2 smallest elements on last axis topk(x, ret_typ='value', k=2, is_ascend=1) = [[\ 0.2,\ 0.3], [\ 0.1,\ 0.2]] 
 // returns the value of top-2 largest elements on axis 0 topk(x, axis=0, ret_typ='value', k=2) = [[\ 0.3,\ 0.3,\ 0.4], [\ 0.1,\ 0.2,\ 0.2]] 
 // flattens and then returns list of both values and indices topk(x, ret_typ='both', k=2) = [[[\ 0.4,\ 0.3], [\ 0.3,\ 0.2]], [[\ 2.,\ 0.], [\ 1.,\ 2.]]] 
 Defined in src/operator/tensor/ordering_op.cc:L68
```

#### Value

mx.nd.transpose	Permutes the dimensions of an array. Examples:: $x = [[1, 2], [3, 4]]$ $transpose(x) = [[1., 3.], [2., 4.]]$ $x = [[[1., 2.], [3., 4.]], [[5., 6.],$
	[7., 8.]] $transpose(x) = [[[1., 5.], [3., 7.]], [[2., 6.], [4., 8.]]]$ $transpose(x, axes=(1,0,2)) = [[[1., 2.], [5., 6.]], [[3., 4.], [7., 8.]]]$

270 mx.nd.trunc

## **Description**

Defined in src/operator/tensor/matrix\_op.cc:L328

## Arguments

data NDArray-or-Symbol Source input

axes Shape(tuple), optional, default=[] Target axis order. By default the axes will be

inverted.

## Value

out The result mx.ndarray

mx.nd.trunc

Return the element-wise truncated value of the input.

# Description

The truncated value of the scalar x is the nearest integer i which is closer to zero than x is. In short, the fractional part of the signed number x is discarded.

## **Arguments**

data

NDArray-or-Symbol The input array.

#### **Details**

Example::

```
trunc([-2.1, -1.9, 1.5, 1.9, 2.1]) = [-2., -1., 1., 1., 2.]
```

The storage type of "trunc" output depends upon the input storage type:

- trunc(default) = default - trunc(row\_sparse) = row\_sparse - trunc(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L857

#### Value

mx.nd.uniform 271

mx.nd.uniform Draw random samples from a uniform distribution.	mx.nd.uniform	Draw random samples from a uniform distribution.		
--	---------------	--	--	--

### **Description**

.. note:: The existing alias "uniform" is deprecated.

# **Arguments**

low float, optional, default=0 Lower bound of the distribution.

high float, optional, default=1 Upper bound of the distribution.

Shape (tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'float16', 'float32', 'float64',optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

#### **Details**

Samples are uniformly distributed over the half-open interval \*[low, high)\* (includes \*low\*, but excludes \*high\*).

Example::

uniform(low=0, high=1, shape=(2,2)) = [[ 0.60276335, 0.85794562], [ 0.54488319, 0.84725171]] Defined in src/operator/random/sample\_op.cc:L96

#### Value

out The result mx.ndarray

mx.nd.unravel.index Converts an array of flat indices into a batch of index arrays. The

operator follows numpy conventions so a single multi index is given by a column of the output matrix. The leading dimension may be left

unspecified by using -1 as placeholder.

## Description

Examples::

### **Arguments**

data NDArray-or-Symbol Array of flat indices

shape Shape(tuple), optional, default=None Shape of the array into which the multi-

indices apply.

272 mx.nd.UpSampling

#### **Details**

```
A = [22,41,37] \text{ unravel\_index}(A, \text{shape=}(7,6)) = [[3,6,6], [4,5,1]] \text{ unravel\_index}(A, \text{shape=}(-1,6)) = [[3,6,6], [4,5,1]]
```

 $B = [[22,41,37],[10,11,15]] \text{ unravel\_index}(B, \text{shape}=(7,6)) = [[[3,6,6],[1,1,2]], [[4,5,1],[4,5,3]]] \text{ unravel\_index}(B, \text{shape}=(-1,6)) = [[[3,6,6],[1,1,2]], [[4,5,1],[4,5,3]]]$ 

Defined in src/operator/tensor/ravel.cc:L76

### Value

out The result mx.ndarray

mx.nd.UpSampling

Upsamples the given input data.

## **Description**

Two algorithms ("sample\_type") are available for upsampling:

### **Arguments**

data NDArray-or-Symbol[] Array of tensors to upsample. For bilinear upsampling,

there should be 2 inputs - 1 data and 1 weight.

scale int, required Up sampling scale

num.filter int, optional, default='0' Input filter. Only used by bilinear sample\_type.Since

bilinear upsampling uses deconvolution, num filters is set to the number of

channels.

sample.type 'bilinear', 'nearest', required upsampling method

multi.input.mode

'concat', 'sum', optional, default='concat' How to handle multiple input. concat means concatenate upsampled images along the channel dimension. sum means

add all images together, only available for nearest neighbor upsampling.

num.args int, required Number of inputs to be upsampled. For nearest neighbor upsam-

pling, this can be 1-N; the size of output will be(scale\*h\_0,scale\*w\_0) and all other inputs will be upsampled to the same size. For bilinear upsampling this

must be 2; 1 input and 1 weight.

workspace long (non-negative), optional, default=512 Tmp workspace for deconvolution

(MB)

mx.nd.where 273

#### **Details**

```
- Nearest Neighbor - Bilinear
```

\*\*Nearest Neighbor Upsampling\*\*

Input data is expected to be NCHW.

Example::

```
x = [[[[1. \ 1. \ 1.] \ [1. \ 1. \ 1.] \ [1. \ 1. \ 1.]]]]
```

```
UpSampling(x, scale=2, sample_type='nearest') = [[[[1. 1. 1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ]]]]
```

Uses 'deconvolution' algorithm under the hood. You need provide both input data and the kernel.

Input data is expected to be NCHW.

'num\_filter' is expected to be same as the number of channels.

Example::

```
 \begin{aligned} \mathbf{x} &= [[[[1. \ 1. \ 1.] \ [1. \ 1. \ 1.]]]] \\ \mathbf{w} &= [[[[1. \ 1. \ 1. \ 1.] \ [1. \ 1. \ 1.]]]] \end{aligned}
```

```
UpSampling(x, w, scale=2, sample_type='bilinear', num_filter=1) = [[[[1. 2. 2. 2. 2. 2. 1.] [2. 4. 4. 4. 2.] [2. 4. 4. 4. 2.] [2. 4. 4. 4. 2.] [1. 2. 2. 2. 2. 1.]]]]
```

Defined in src/operator/nn/upsampling.cc:L173

#### Value

out The result mx.ndarray

mx.nd.where

*Return the elements, either from x or y, depending on the condition.* 

## **Description**

Given three ndarrays, condition, x, and y, return an ndarray with the elements from x or y, depending on the elements from condition are true or false. x and y must have the same shape. If condition has the same shape as x, each element in the output array is from x if the corresponding element in the condition is true, and from y if false.

### **Arguments**

condition NDArray-or-Symbol condition array

x NDArray-or-Symbol

y NDArray-or-Symbol

<sup>\*\*</sup>Bilinear Upsampling\*\*

274 mx.nd.zeros

### **Details**

If condition does not have the same shape as x, it must be a 1D array whose size is the same as x's first dimension size. Each row of the output array is from x's row if the corresponding element from condition is true, and from y's row if false.

Note that all non-zero values are interpreted as "True" in condition.

## Examples::

```
x = [[1, 2], [3, 4]] y = [[5, 6], [7, 8]] cond = [[0, 1], [-1, 0]]
where(cond, x, y) = [[5, 2], [3, 8]]
csr_cond = cast_storage(cond, 'csr')
where(csr_cond, x, y) = [[5, 2], [3, 8]]
Defined in src/operator/tensor/control_flow_op.cc:L57
```

#### Value

out The result mx.ndarray

mx.nd.zeros

Generate an mx.nd.array object with zeros

## **Description**

Generate an mx.nd.array object with zeros

### Usage

```
mx.nd.zeros(shape, ctx = NULL)
```

## **Arguments**

shape the dimension of the mx.nd.array

ctx optional The context device of the array. mx.ctx.default() will be used in default.

## **Examples**

```
mat = mx.nd.zeros(10)
as.array(mat)
mat2 = mx.nd.zeros(c(5,5))
as.array(mat)
mat3 = mx.nd.zeroes(c(3,3,3))
as.array(mat3)
```

mx.nd.zeros.like 275

mx.nd.zeros.like

Return an array of zeros with the same shape, type and storage type as the input array.

# Description

The storage type of "zeros\_like" output depends on the storage type of the input

## **Arguments**

data

NDArray-or-Symbol The input

### **Details**

```
- zeros_like(row_sparse) = row_sparse - zeros_like(csr) = csr - zeros_like(default) = default Examples:: x = [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] zeros_like(x) = [[\ 0.,\ 0.,\ 0.],\ [\ 0.,\ 0.,\ 0.]]
```

### Value

out The result mx.ndarray

mx.opt.adadelta

Create an AdaDelta optimizer with respective parameters.

# Description

AdaDelta optimizer as described in Zeiler, M. D. (2012). \*ADADELTA: An adaptive learning rate method.\* http://arxiv.org/abs/1212.5701

### Usage

```
mx.opt.adadelta(
  rho = 0.9,
  epsilon = 1e-05,
  wd = 0,
  rescale.grad = 1,
  clip_gradient = -1
)
```

276 mx.opt.adagrad

## **Arguments**

rho float, default=0.90 Decay rate for both squared gradients and delta x.

epsilon float, default=1e-5 The constant as described in the thesis.

wd float, default=0.0 L2 regularization coefficient add to all the weights.

rescale.grad float, default=1 rescaling factor of gradient.

clip\_gradient float, default=-1 (no clipping if < 0) clip gradient in range [-clip\_gradient, clip\_gradient].

mx.opt.adagrad Create an AdaGrad optimizer with respective parameters. AdaGrad

optimizer of Duchi et al., 2011,

# Description

This code follows the version in http://arxiv.org/pdf/1212.5701v1.pdf Eq(5) by Matthew D. Zeiler, 2012. AdaGrad will help the network to converge faster in some cases.

## Usage

```
mx.opt.adagrad(
  learning.rate = 0.05,
  epsilon = 1e-08,
  wd = 0,
  rescale.grad = 1,
  clip_gradient = -1,
  lr_scheduler = NULL
)
```

## **Arguments**

learning.rate float, default=0.05 Step size.

epsilon float, default=1e-8

wd float, default=0.0 L2 regularization coefficient add to all the weights.

rescale.grad float, default=1.0 rescaling factor of gradient.

clip\_gradient float, default=-1.0 (no clipping if < 0) clip gradient in range [-clip\_gradient,

clip\_gradient].

lr\_scheduler function, optional The learning rate scheduler.

mx.opt.adam 277

mx.opt.adam	Create an Adam optimizer with respective parameters. Adam optimizer as described in [King2014].

## Description

[King2014] Diederik Kingma, Jimmy Ba, Adam: A Method for Stochastic Optimization, http://arxiv.org/abs/1412.6980

#### **Usage**

```
mx.opt.adam(
  learning.rate = 0.001,
  beta1 = 0.9,
  beta2 = 0.999,
  epsilon = 1e-08,
  wd = 0,
  rescale.grad = 1,
  clip_gradient = -1,
  lr_scheduler = NULL
)
```

#### **Arguments**

learning.rate float, default=1e-3 The initial learning rate.

beta1 float, default=0.9 Exponential decay rate for the first moment estimates.

beta2 float, default=0.999 Exponential decay rate for the second moment estimates.

epsilon float, default=1e-8

wd float, default=0.0 L2 regularization coefficient add to all the weights.

rescale.grad float, default=1.0 rescaling factor of gradient.

clip\_gradient float, optional, default=-1 (no clipping if < 0) clip gradient in range [-clip\_gradient,

clip\_gradient].

lr\_scheduler function, optional The learning rate scheduler.

mx.opt.create Create an optimizer by name and parameters

## **Description**

Create an optimizer by name and parameters

### Usage

```
mx.opt.create(name, ...)
```

278 mx.opt.nag

## **Arguments**

name The name of the optimizer
... Additional arguments

mx.opt.get.updater

Get an updater closure that can take list of weight and gradient and return updated list of weight.

## **Description**

Get an updater closure that can take list of weight and gradient and return updated list of weight.

# Usage

```
mx.opt.get.updater(optimizer, weights, ctx)
```

## **Arguments**

optimizer

The optimizer

weights

The weights to be optimized

mx.opt.nag

Create a Nesterov Accelerated SGD( NAG) optimizer.

# Description

NAG optimizer is described in Aleksandar Botev. et al (2016). \*NAG: A Nesterov accelerated SGD.\* https://arxiv.org/pdf/1607.01981.pdf

## Usage

```
mx.opt.nag(
  learning.rate = 0.01,
  momentum = 0,
  wd = 0,
  rescale.grad = 1,
  clip_gradient = -1,
  lr_scheduler = NULL
)
```

mx.opt.rmsprop 279

## Arguments

learning.rate float, default=0.01 The initial learning rate.

momentum float, default=0 The momentum value

wd float, default=0.0 L2 regularization coefficient added to all the weights.

rescale.grad float, default=1.0 rescaling factor of gradient.

clip\_gradient float, optional, default=-1 (no clipping if < 0) clip gradient in range [-clip\_gradient,

clip\_gradient].

1r\_scheduler function, optional The learning rate scheduler.

mx.opt.rmsprop Create an RMSProp optimizer with respective parameters. Refer-

ence: Tieleman T, Hinton G. Lecture 6.5-rmsprop: Divide the gradient by a running average of its recent magnitude[J]. COURSERA: Neural Networks for Machine Learning, 2012, 4(2). The code follows: http://arxiv.org/pdf/1308.0850v5.pdf Eq(38) - Eq(45) by Alex Graves,

2013.

### Description

Create an RMSProp optimizer with respective parameters. Reference: Tieleman T, Hinton G. Lecture 6.5-rmsprop: Divide the gradient by a running average of its recent magnitude[J]. COURSERA: Neural Networks for Machine Learning, 2012, 4(2). The code follows: http://arxiv.org/pdf/1308.0850v5.pdf Eq(38) - Eq(45) by Alex Graves, 2013.

### Usage

```
mx.opt.rmsprop(
  learning.rate = 0.002,
  centered = TRUE,
  rho = 0.95,
  momentum = 0.9,
  epsilon = 1e-04,
  wd = 0,
  rescale.grad = 1,
  clip_gradient = -1,
  lr_scheduler = NULL
)
```

## **Arguments**

learning.rate float, default=0.002 The initial learning rate.

rho float, default=0.95 decay factor of moving average for gradient, gradient^2.

momentum float, default=0.9 "momentum" factor.

epsilon float, default=1e-4

280 mx.opt.sgd

wd float, default=0.0 L2 regularization coefficient add to all the weights.

rescale.grad float, default=1.0 rescaling factor of gradient.

clip\_gradient float, optional, default=-1 (no clipping if < 0) clip gradient in range [-clip\_gradient, clip\_gradient].

lr\_scheduler function, optional The learning rate scheduler.

## Description

Create an SGD optimizer with respective parameters. Perform SGD with momentum update

# Usage

```
mx.opt.sgd(
  learning.rate = 0.01,
  momentum = 0,
  wd = 0,
  rescale.grad = 1,
  clip_gradient = -1,
  lr_scheduler = NULL
)
```

# Arguments

learning.rate float, default=0.01 The initial learning rate.

momentum float, default=0 The momentum value

wd float, default=0.0 L2 regularization coefficient add to all the weights.

rescale.grad float, default=1.0 rescaling factor of gradient.

clip\_gradient float, optional, default=-1 (no clipping if < 0) clip gradient in range [-clip\_gradient,

clip\_gradient].

lr\_scheduler function, optional The learning rate scheduler.

mx.profiler.config 281

mx.profiler.config

Set up the configuration of profiler.

## **Description**

Set up the configuration of profiler.

### Usage

```
mx.profiler.config(params)
```

### **Arguments**

flags

list of key/value pair tuples. Indicates configuration parameters profile\_symbolic : boolean, whether to profile symbolic operators profile\_imperative : boolean, whether to profile imperative operators profile\_memory : boolean, whether to profile memory usage profile\_api : boolean, whether to profile the C API file\_name : string, output file for profile data continuous\_dump : boolean, whether to periodically dump profiling data to file dump\_period : float, seconds between profile data dumps

mx.profiler.state

Set up the profiler state to record operator.

# Description

Set up the profiler state to record operator.

### Usage

```
mx.profiler.state(state = MX.PROF.STATE$STOP)
```

## **Arguments**

state Indicting whether to run the profiler, can be 'MX.PROF.STATE\$RUN' or 'MX.PROF.STATE\$STOP'.

Default is 'MX.PROF.STATE\$STOP'.

filename The name of output trace file. Default is 'profile.json'

282 mx.runif

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IIIX .	rno	I III

Generate nomal distribution with mean and sd.

## **Description**

Generate nomal distribution with mean and sd.

## Usage

```
mx.rnorm(shape, mean = 0, sd = 1, ctx = NULL)
```

## **Arguments**

shape Dimension, The shape(dimension) of the result.

mean numeric, The mean of distribution. sd numeric, The standard deviations.

ctx, optional The context device of the array. mx.ctx.default() will be used in default.

## **Examples**

```
mx.set.seed(0)
as.array(mx.runif(2))
# 0.5488135 0.5928446
mx.set.seed(0)
as.array(mx.rnorm(2))
# 2.212206 1.163079
```

mx.runif

Generate uniform distribution in [low, high) with specified shape.

## **Description**

Generate uniform distribution in [low, high) with specified shape.

# Usage

```
mx.runif(shape, min = 0, max = 1, ctx = NULL)
```

## **Arguments**

shape	Dimension, The shape(dimension) of the result.
min	numeric, The lower bound of distribution.
max	numeric, The upper bound of distribution.
ctx,	optional The context device of the array. mx.ctx.default() will be used in default.

mx.serialize 283

### **Examples**

```
mx.set.seed(0)
as.array(mx.runif(2))
# 0.5488135 0.5928446
mx.set.seed(0)
as.array(mx.rnorm(2))
# 2.212206 1.163079
```

mx.serialize

Serialize MXNet model into RData-compatiable format.

# Description

Serialize MXNet model into RData-compatiable format.

## Usage

```
mx.serialize(model)
```

### **Arguments**

model

The mxnet model

mx.set.seed

*Set the seed used by mxnet device-specific random number generators.* 

# Description

Set the seed used by mxnet device-specific random number generators.

### Usage

```
mx.set.seed(seed)
```

### **Arguments**

seed

the seed value to the device random number generators.

#### **Details**

We have a specific reason why mx.set.seed is introduced, instead of simply use set.seed.

The reason that is that most of mxnet random number generator can run on different devices, such as GPU. We need to use massively parallel PRNG on GPU to get fast random number generations. It can also be quite costly to seed these PRNGs. So we introduced mx.set.seed for mxnet specific device random numbers.

284 mx.symbol.abs

### **Examples**

```
mx.set.seed(0)
as.array(mx.runif(2))
# 0.5488135 0.5928446
mx.set.seed(0)
as.array(mx.rnorm(2))
# 2.212206 1.163079
```

mx.simple.bind

Simple bind the symbol to executor, with information from input shapes.

# Description

Simple bind the symbol to executor, with information from input shapes.

### Usage

```
mx.simple.bind(symbol, ctx, grad.req = "null", fixed.param = NULL, ...)
```

mx.symbol.abs

abs:Returns element-wise absolute value of the input.

# Description

Example::

# Usage

```
mx.symbol.abs(...)
```

### **Arguments**

data NI

NDArray-or-Symbol The input array.

name

string, optional Name of the resulting symbol.

### **Details**

```
abs([-2, 0, 3]) = [2, 0, 3]
```

The storage type of "abs" output depends upon the input storage type:

- abs(default) = default - abs(row\_sparse) = row\_sparse - abs(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L721

mx.symbol.Activation 285

### Value

out The result mx.symbol

mx.symbol.Activation Activation: Applies an activation function element-wise to the input.

## **Description**

The following activation functions are supported:

# Usage

```
mx.symbol.Activation(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

act.type 'relu', 'sigmoid', 'softrelu', 'softsign', 'tanh', required Activation function to be

applied.

name string, optional Name of the resulting symbol.

#### **Details**

```
- 'relu': Rectified Linear Unit, :math: 'y = max(x, 0)' - 'sigmoid': :math: 'y = \frac11 + exp(-x)' - 'tanh': Hyperbolic tangent, :math: 'y = \fracexp(x) - exp(-x)exp(x) + exp(-x)' - 'softrelu': Soft ReLU, or SoftPlus, :math: 'y = \log(1 + \exp(x))' - 'softsign': :math: 'y = \fracx1 + abs(x)'
```

Defined in src/operator/nn/activation.cc:L165

#### Value

out The result mx.symbol

mx.symbol.adam\_update adam\_update:Update function for Adam optimizer. Adam is seen as a generalization of AdaGrad.

### **Description**

Adam update consists of the following steps, where g represents gradient and m, v are 1st and 2nd order moment estimates (mean and variance).

### Usage

```
mx.symbol.adam_update(...)
```

### **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
mean	NDArray-or-Symbol Moving mean
var	NDArray-or-Symbol Moving variance
lr	float, required Learning rate
beta1	float, optional, default=0.899999976 The decay rate for the 1st moment estimates.
beta2	float, optional, default=0.999000013 The decay rate for the 2nd moment estimates.
epsilon	float, optional, default=9.9999994e-09 A small constant for numerical stability.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
lazy.update	boolean, optional, default=1 If true, lazy updates are applied if gradient's stype is row_sparse and all of w, m and v have the same stype

## **Details**

```
.. math::
```

name

```
 g_t = \adjust{0.05} \ J(W_t-1) \ m_t = \beta_1 \ m_t-1 + (1 - \beta_1) \ g_t \ v_t = \beta_2 \ v_t-1 + (1 - \beta_2) \ g_t^2 \ W_t = W_t-1 - \adjust{0.05} \ m_t \ g_t + \end{0.05}
```

string, optional Name of the resulting symbol.

It updates the weights using::

```
m = beta1*m + (1-beta1)*grad v = beta2*v + (1-beta2)*(grad**2) w += - learning_rate * m / (sqrt(v) + epsilon)
```

However, if grad's storage type is "row\_sparse", "lazy\_update" is True and the storage type of weight is the same as those of m and v, only the row slices whose indices appear in grad.indices are updated (for w, m and v)::

for row in grad.indices:  $m[row] = beta1*m[row] + (1-beta1)*grad[row] v[row] = beta2*v[row] + (1-beta2)*(grad[row]**2) w[row] += - learning_rate * m[row] / (sqrt(v[row]) + epsilon)$ 

Defined in src/operator/optimizer\_op.cc:L679

### Value

out The result mx.symbol

mx.symbol.add\_n 287

mx.symbol.add\_n

add\_n:Adds all input arguments element-wise.

### **Description**

```
.. math:: add\_n(a_1, a_2, ..., a_n) = a_1 + a_2 + ... + a_n
```

## Usage

```
mx.symbol.add_n(...)
```

### **Arguments**

args NDArray-or-Symbol[] Positional input arguments name string, optional Name of the resulting symbol.

#### **Details**

"add\_n" is potentially more efficient than calling "add" by 'n' times.

The storage type of "add\_n" output depends on storage types of inputs

- add\_n(row\_sparse, row\_sparse, ..) = row\_sparse - add\_n(default, csr, default) = default - add\_n(any input combinations longer than 4 (>4) with at least one default type) = default - otherwise, "add\_n" falls all inputs back to default storage and generates default storage

Defined in src/operator/tensor/elemwise\_sum.cc:L155

#### Value

out The result mx.symbol

```
mx.symbol.all\_finite all_finite: Check if all the float numbers in the array are finite (used for AMP)
```

# Description

Defined in src/operator/contrib/all\_finite.cc:L101

### Usage

```
mx.symbol.all_finite(...)
```

### **Arguments**

data NDArray Array

init.output boolean, optional, default=1 Initialize output to 1. string, optional Name of the resulting symbol.

### Value

out The result mx.symbol

mx.symbol.amp\_cast

amp\_cast:Cast function between low precision float/FP32 used by AMP.

### **Description**

It casts only between low precision float/FP32 and does not do anything for other types.

## Usage

```
mx.symbol.amp_cast(...)
```

### **Arguments**

data NDArray-or-Symbol The input.

dtype 'bfloat16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', required

Output data type.

name string, optional Name of the resulting symbol.

### **Details**

Defined in src/operator/tensor/amp\_cast.cc:L121

## Value

out The result mx.symbol

```
mx.symbol.amp_multicast
```

amp\_multicast: Cast function used by AMP, that casts its inputs to the common widest type.

# Description

It casts only between low precision float/FP32 and does not do anything for other types.

## Usage

```
mx.symbol.amp_multicast(...)
```

mx.symbol.arccos 289

# **Arguments**

data NDArray-or-Symbol[] Weights

num.outputs int, required Number of input/output pairs to be casted to the widest type.

cast.narrow boolean, optional, default=0 Whether to cast to the narrowest type

name string, optional Name of the resulting symbol.

### **Details**

Defined in src/operator/tensor/amp\_cast.cc:L165

#### Value

out The result mx.symbol

mx.symbol.arccos

arccos:Returns element-wise inverse cosine of the input array.

# Description

The input should be in range '[-1, 1]'. The output is in the closed interval :math: '[0, \pi]'

# Usage

```
mx.symbol.arccos(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

```
.. math:: arccos([-1, -.707, 0, .707, 1]) = [\pi, 3\pi/4, \pi/2, \pi/4, 0]
```

The storage type of "arccos" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L233

# Value

290 mx.symbol.arcsin

mx.symbol.arccosh

arccosh:Returns the element-wise inverse hyperbolic cosine of the input array, \computed element-wise.

# **Description**

The storage type of "arccosh" output is always dense

# Usage

```
mx.symbol.arccosh(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L535

### Value

out The result mx.symbol

mx.symbol.arcsin

arcsin:Returns element-wise inverse sine of the input array.

# **Description**

The input should be in the range '[-1, 1]'. The output is in the closed interval of [:math:'-\pi/2', :math:'\pi/2'].

### Usage

```
mx.symbol.arcsin(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.arcsinh 291

# **Details**

```
.. math:: \arcsin([-1, -.707, 0, .707, 1]) = [-\pi/2, -\pi/4, 0, \pi/4, \pi/2]
```

The storage type of "arcsin" output depends upon the input storage type:

- arcsin(default) = default - arcsin(row\_sparse) = row\_sparse - arcsin(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L187

### Value

out The result mx.symbol

mx.symbol.arcsinh

arcsinh:Returns the element-wise inverse hyperbolic sine of the input array, \computed element-wise.

# Description

The storage type of "arcsinh" output depends upon the input storage type:

# Usage

```
mx.symbol.arcsinh(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

# **Details**

```
- arcsinh(default) = default - arcsinh(row_sparse) = row_sparse - arcsinh(csr) = csr
Defined in src/operator/tensor/elemwise_unary_op_trig.cc:L494
```

### Value

292 mx.symbol.arctanh

mx.symbol.arctan

arctan:Returns element-wise inverse tangent of the input array.

## **Description**

The output is in the closed interval :math: '[-\pi/2, \pi/2]'

# Usage

```
mx.symbol.arctan(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

# **Details**

```
.. math:: \arctan([-1, 0, 1]) = [-\pi/4, 0, \pi/4]
```

The storage type of "arctan" output depends upon the input storage type:

- arctan(default) = default - arctan(row\_sparse) = row\_sparse - arctan(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L282

### Value

out The result mx.symbol

mx.symbol.arctanh

arctanh: Returns the element-wise inverse hyperbolic tangent of the input array, \ computed element-wise.

# Description

The storage type of "arctanh" output depends upon the input storage type:

# Usage

```
mx.symbol.arctanh(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.argmax 293

### **Details**

```
- arctanh(default) = default - arctanh(row_sparse) = row_sparse - arctanh(csr) = csr
Defined in src/operator/tensor/elemwise_unary_op_trig.cc:L579
```

### Value

out The result mx.symbol

mx.symbol.argmax

argmax:Returns indices of the maximum values along an axis.

### **Description**

In the case of multiple occurrences of maximum values, the indices corresponding to the first occurrence are returned.

## Usage

```
mx.symbol.argmax(...)
```

### **Arguments**

data NDArray-or-Symbol The input

axis int or None, optional, default='None' The axis along which to perform the re-

duction. Negative values means indexing from right to left. "Requires axis to be

set as int, because global reduction is not supported yet."

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axis is left in the

result as dimension with size one.

name string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
x = [[0., 1., 2.], [3., 4., 5.]]
```

// argmax along axis 0 argmax(x, axis=0) = [1., 1., 1.]

// argmax along axis 1 argmax(x, axis=1) = [2., 2.]

// argmax along axis 1 keeping same dims as an input array argmax(x, axis=1, keepdims=True) = [[ 2.], [ 2.]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L52

#### Value

294 mx.symbol.argmin

```
mx.symbol.argmax_channel
```

argmax\_channel:Returns argmax indices of each channel from the input array.

# **Description**

The result will be an NDArray of shape (num\_channel,).

# Usage

```
mx.symbol.argmax_channel(...)
```

## **Arguments**

data NDArray-or-Symbol The input array

name string, optional Name of the resulting symbol.

### **Details**

In case of multiple occurrences of the maximum values, the indices corresponding to the first occurrence are returned.

Examples::

```
x = [[0., 1., 2.], [3., 4., 5.]]

argmax\_channel(x) = [2., 2.]
```

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L97

#### Value

out The result mx.symbol

mx.symbol.argmin

argmin:Returns indices of the minimum values along an axis.

# Description

In the case of multiple occurrences of minimum values, the indices corresponding to the first occurrence are returned.

# Usage

```
mx.symbol.argmin(...)
```

mx.symbol.argsort 295

### **Arguments**

data NDArray-or-Symbol The input

axis int or None, optional, default='None' The axis along which to perform the re-

duction. Negative values means indexing from right to left. "Requires axis to be

set as int, because global reduction is not supported yet."

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axis is left in the

result as dimension with size one.

name string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
x = [[0., 1., 2.], [3., 4., 5.]]
```

// argmin along axis 0 argmin(x, axis=0) = [0., 0., 0.]

// argmin along axis 1 argmin(x, axis=1) = [0., 0.]

// argmin along axis 1 keeping same dims as an input array argmin(x, axis=1, keepdims=True) = [[ 0.], [ 0.]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L77

# Value

out The result mx.symbol

mx.symbol.argsort argsort:Returns the indices that would sort an input array along the

given axis.

# **Description**

This function performs sorting along the given axis and returns an array of indices having same shape as an input array that index data in sorted order.

## Usage

```
mx.symbol.argsort(...)
```

### **Arguments**

data NDArray-or-Symbol	The input array
------------------------	-----------------

axis int or None, optional, default='-1' Axis along which to sort the input tensor. If

not given, the flattened array is used. Default is -1.

is ascend boolean, optional, default=1 Whether to sort in ascending or descending order.

dtype 'float16', 'float32', 'float64', 'int32', 'int64', 'uint8', optional, default='float32'

DType of the output indices. It is only valid when ret\_typ is "indices" or "both". An error will be raised if the selected data type cannot precisely represent the

indices.

name string, optional Name of the resulting symbol.

#### **Details**

## Examples::

```
x = [[ 0.3, 0.2, 0.4], [ 0.1, 0.3, 0.2]]

// sort along axis -1 argsort(x) = [[ 1., 0., 2.], [ 0., 2., 1.]]

// sort along axis 0 argsort(x, axis=0) = [[ 1., 0., 1.] [ 0., 1., 0.]]

// flatten and then sort argsort(x, axis=None) = [ 3., 1., 5., 0., 4., 2.]

Defined in src/operator/tensor/ordering_op.cc:L185
```

### Value

out The result mx.symbol

mx.symbol.BatchNorm

BatchNorm:Batch normalization.

# **Description**

Normalizes a data batch by mean and variance, and applies a scale "gamma" as well as offset "beta".

# Usage

```
mx.symbol.BatchNorm(...)
```

### **Arguments**

data NDArray-or-Symbol Input data to batch normalization

gamma NDArray-or-Symbol gamma array beta NDArray-or-Symbol beta array

moving.mean NDArray-or-Symbol running mean of input moving.var NDArray-or-Symbol running variance of input

eps double, optional, default=0.001000000474974513 Epsilon to prevent div 0.

Must be no less than CUDNN\_BN\_MIN\_EPSILON defined in cudnn.h when

using cudnn (usually 1e-5)

momentum float, optional, default=0.899999976 Momentum for moving average

fix.gamma boolean, optional, default=1 Fix gamma while training

use.global.stats

boolean, optional, default=0 Whether use global moving statistics instead of local batch-norm. This will force change batch-norm into a scale shift operator.

297

output.mean.var

boolean, optional, default=0 Output the mean and inverse std

axis int, optional, default='1' Specify which shape axis the channel is specified

cudnn.off boolean, optional, default=0 Do not select CUDNN operator, if available

min.calib.range

float or None, optional, default=None The minimum scalar value in the form of float32 obtained through calibration. If present, it will be used to by quantized batch norm op to calculate primitive scale.Note: this calib\_range is to calib bn output.

max.calib.range

float or None, optional, default=None The maximum scalar value in the form of float32 obtained through calibration. If present, it will be used to by quantized batch norm op to calculate primitive scale.Note: this calib\_range is to calib bn output.

name string, optional Name of the resulting symbol.

#### **Details**

Assume the input has more than one dimension and we normalize along axis 1. We first compute the mean and variance along this axis:

.. math::

```
data\_mean[i] = mean(data[:,i,:,...]) \setminus data\_var[i] = var(data[:,i,:,...])
```

Then compute the normalized output, which has the same shape as input, as following:

.. math::

```
out[:,i,:,...] = \frac{(i,i,:,...] - data_mean[i] \cdot qrtdata_var[i] + epsilon * gamma[i] + beta[i]}
```

Both \*mean\* and \*var\* returns a scalar by treating the input as a vector.

Assume the input has size \*k\* on axis 1, then both "gamma" and "beta" have shape \*(k,)\*. If "output\_mean\_var" is set to be true, then outputs both "data\_mean" and the inverse of "data\_var", which are needed for the backward pass. Note that gradient of these two outputs are blocked.

Besides the inputs and the outputs, this operator accepts two auxiliary states, "moving\_mean" and "moving\_var", which are \*k\*-length vectors. They are global statistics for the whole dataset, which are updated by::

moving\_mean = moving\_mean \* momentum + data\_mean \* (1 - momentum) moving\_var = moving\_var \* momentum + data\_var \* (1 - momentum)

If "use\_global\_stats" is set to be true, then "moving\_mean" and "moving\_var" are used instead of "data mean" and "data var" to compute the output. It is often used during inference.

The parameter "axis" specifies which axis of the input shape denotes the 'channel' (separately normalized groups). The default is 1. Specifying -1 sets the channel axis to be the last item in the input shape.

Both "gamma" and "beta" are learnable parameters. But if "fix\_gamma" is true, then set "gamma" to 1 and its gradient to 0.

.. Note:: When "fix\_gamma" is set to True, no sparse support is provided. If "fix\_gamma is" set to False, the sparse tensors will fallback.

Defined in src/operator/nn/batch\_norm.cc:L547

### Value

out The result mx.symbol

```
mx.symbol.BatchNorm_v1
```

BatchNorm\_v1:Batch normalization.

# **Description**

This operator is DEPRECATED. Perform BatchNorm on the input.

### Usage

```
mx.symbol.BatchNorm_v1(...)
```

### **Arguments**

data NDArray-or-Symbol Input data to batch normalization

gamma NDArray-or-Symbol gamma array beta NDArray-or-Symbol beta array

eps float, optional, default=0.00100000005 Epsilon to prevent div 0

momentum float, optional, default=0.899999976 Momentum for moving average

fix.gamma boolean, optional, default=1 Fix gamma while training

use.global.stats

boolean, optional, default=0 Whether use global moving statistics instead of

local batch-norm. This will force change batch-norm into a scale shift operator.

output.mean.var

boolean, optional, default=0 Output All,normal mean and var

name string, optional Name of the resulting symbol.

#### **Details**

Normalizes a data batch by mean and variance, and applies a scale "gamma" as well as offset "beta".

Assume the input has more than one dimension and we normalize along axis 1. We first compute the mean and variance along this axis:

```
.. math::
```

```
data\underline{nean[i]} = mean(data[:,i,:,...]) \setminus data\underline{var[i]} = var(data[:,i,:,...])
```

Then compute the normalized output, which has the same shape as input, as following:

mx.symbol.batch\_dot 299

```
.. math::
```

```
out[:,i,:,...] = \fracdata[:,i,:,...] - data\_mean[i]\sqrtdata\_var[i]+\epsilon * gamma[i] + beta[i]
```

Both \*mean\* and \*var\* returns a scalar by treating the input as a vector.

Assume the input has size \*k\* on axis 1, then both "gamma" and "beta" have shape \*(k,)\*. If "output\_mean\_var" is set to be true, then outputs both "data\_mean" and "data\_var" as well, which are needed for the backward pass.

Besides the inputs and the outputs, this operator accepts two auxiliary states, "moving\_mean" and "moving\_var", which are \*k\*-length vectors. They are global statistics for the whole dataset, which are updated by::

moving\_mean = moving\_mean \* momentum + data\_mean \* (1 - momentum) moving\_var = moving\_var \* momentum + data\_var \* (1 - momentum)

If "use\_global\_stats" is set to be true, then "moving\_mean" and "moving\_var" are used instead of "data\_mean" and "data\_var" to compute the output. It is often used during inference.

Both "gamma" and "beta" are learnable parameters. But if "fix\_gamma" is true, then set "gamma" to 1 and its gradient to 0.

There's no sparse support for this operator, and it will exhibit problematic behavior if used with sparse tensors.

Defined in src/operator/batch\_norm\_v1.cc:L95

#### Value

out The result mx.symbol

mx.symbol.batch\_dot

batch dot:Batchwise dot product.

### **Description**

"batch\_dot" is used to compute dot product of "x" and "y" when "x" and "y" are data in batch, namely N-D ( $N \ge 3$ ) arrays in shape of '( $B0, ..., B_i, ..., ...$ ).

#### Usage

```
mx.symbol.batch_dot(...)
```

### **Arguments**

1hs NDArray-or-Symbol The first input rhs NDArray-or-Symbol The second input

transpose.a boolean, optional, default=0 If true then transpose the first input before dot. boolean, optional, default=0 If true then transpose the second input before dot. forward.stype None, 'csr', 'default', 'row\_sparse', optional, default='None' The desired stor-

age type of the forward output given by user, if the combination of input storage types and this hint does not matchany implemented ones, the dot operator will perform fallback operationand still produce an output of the desired storage type.

name string, optional Name of the resulting symbol.

### **Details**

```
For example, given "x" with shape '(B_0, ..., B_i, N, M)' and "y" with shape '(B_0, ..., B_i, M, K)', the result array will have shape '(B_0, ..., B_i, N, K)', which is computed by:: batch\_dot(x,y)[b_0, ..., b_i, :, :] = dot(x[b_0, ..., b_i, :, :], y[b_0, ..., b_i, :, :])
```

Defined in src/operator/tensor/dot.cc:L127

#### Value

out The result mx.symbol

mx.symbol.batch\_take batch take: Takes elements from a data batch.

# Description

```
.. note:: 'batch_take' is deprecated. Use 'pick' instead.
```

### Usage

```
mx.symbol.batch_take(...)
```

# **Arguments**

a NDArray-or-Symbol The input array indices NDArray-or-Symbol The index array

name string, optional Name of the resulting symbol.

### **Details**

Given an input array of shape "(d0, d1)" and indices of shape "(i0,)", the result will be an output array of shape "(i0,)" with::

```
output[i] = input[i, indices[i]]
```

Examples::

```
x = [[1., 2.], [3., 4.], [5., 6.]]
```

// takes elements with specified indices batch\_take(x, [0,1,0]) = [1. 4. 5.]

Defined in src/operator/tensor/indexing\_op.cc:L841

### Value

```
mx.symbol.BilinearSampler
```

BilinearSampler: Applies bilinear sampling to input feature map.

### **Description**

Bilinear Sampling is the key of [NIPS2015] \"Spatial Transformer Networks\". The usage of the operator is very similar to remap function in OpenCV, except that the operator has the backward pass.

### Usage

```
mx.symbol.BilinearSampler(...)
```

### **Arguments**

data NDArray-or-Symbol Input data to the BilinearsamplerOp.

grid NDArray-or-Symbol Input grid to the BilinearsamplerOp.grid has two channels:

x\_src, y\_src

cudnn.off boolean or None, optional, default=None whether to turn cudnn off

name string, optional Name of the resulting symbol.

### Details

Given :math: 'data' and :math: 'grid', then the output is computed by

```
.. math:: x\_src = grid[batch, 0, y\_dst, x\_dst] \setminus y\_src = grid[batch, 1, y\_dst, x\_dst] \setminus output[batch, channel, y\_dst, x\_dst] = G(data[batch, channel, y\_src, x\_src)
```

:math:' $x_dst'$ , :math:' $y_dst'$  enumerate all spatial locations in :math:'output', and :math:'G()' denotes the bilinear interpolation kernel. The out-boundary points will be padded with zeros. The shape of the output will be (data.shape[0], data.shape[1], grid.shape[2], grid.shape[3]).

The operator assumes that :math:'data' has 'NCHW' layout and :math:'grid' has been normalized to [-1, 1].

BilinearSampler often cooperates with GridGenerator which generates sampling grids for BilinearSampler. GridGenerator supports two kinds of transformation: "affine" and "warp". If users want to design a CustomOp to manipulate :math: 'grid', please firstly refer to the code of GridGenerator.

#### Example 1::

```
## Zoom out data two times data = array([[[[1, 4, 3, 6], [1, 8, 8, 9], [0, 4, 1, 5], [1, 0, 1, 3]]]])
affine_matrix = array([[2, 0, 0], [0, 2, 0]])
affine_matrix = reshape(affine_matrix, shape=(1, 6))
grid = GridGenerator(data=affine_matrix, transform_type='affine', target_shape=(4, 4))
out = BilinearSampler(data, grid)
out [[[[0, 0, 0, 0], [0, 3.5, 6.5, 0], [0, 1.25, 2.5, 0], [0, 0, 0, 0]]]
```

```
Example 2::
## shift data horizontally by -1 pixel
data = array([[[[1, 4, 3, 6], [1, 8, 8, 9], [0, 4, 1, 5], [1, 0, 1, 3]]]])
warp_maxtrix = array([[[[1, 1, 1, 1], [1, 1, 1, 1], [1, 1, 1, 1], [1, 1, 1, 1]], [[0, 0, 0, 0], [0, 0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0, 0], [0, 0], [0, 0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0, 0], [0,
```

grid = GridGenerator(data=warp\_matrix, transform\_type='warp') out = BilinearSampler(data, grid) out [[[[ 4, 3, 6, 0], [ 8, 8, 9, 0], [ 4, 1, 5, 0], [ 0, 1, 3, 0]]]

Defined in src/operator/bilinear sampler.cc:L256

#### Value

out The result mx.symbol

0, 0, 0], [0, 0, 0, 0]]]])

mx.symbol.BlockGrad

BlockGrad:Stops gradient computation.

# Description

Stops the accumulated gradient of the inputs from flowing through this operator in the backward direction. In other words, this operator prevents the contribution of its inputs to be taken into account for computing gradients.

### Usage

```
mx.symbol.BlockGrad(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

# **Details**

# Example::

```
v1 = [1, 2] \ v2 = [0, 1] \ a = Variable('a') \ b = Variable('b') \ b\_stop\_grad = stop\_gradient(3*b) \ loss = MakeLoss(b\_stop\_grad + a)
```

```
executor = loss.simple_bind(ctx=cpu(), a=(1,2), b=(1,2)) executor.forward(is_train=True, a=v1, b=v2) executor.outputs [ 1. 5.]
```

executor.backward() executor.grad\_arrays [ 0. 0.] [ 1. 1.]

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L326

#### Value

```
mx.symbol.broadcast_add
```

broadcast\_add:Returns element-wise sum of the input arrays with broadcasting.

# Description

'broadcast\_plus' is an alias to the function 'broadcast\_add'.

# Usage

```
mx.symbol.broadcast_add(...)
```

# Arguments

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function
name	string, optional Name of the resulting symbol.

## **Details**

```
Example::
```

```
\begin{split} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_add(x,y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 2.,\ 2.,\ 2.]] \\ broadcast\_plus(x,y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 2.,\ 2.,\ 2.]] \\ Supported sparse operations: \\ broadcast\_add(csr,\ dense(1D)) &= dense \ broadcast\_add(dense(1D),\ csr) &= dense \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L58 \end{split}
```

### Value

```
mx.symbol.broadcast_axes
```

broadcast\_axes:Broadcasts the input array over particular axes.

# **Description**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

# Usage

```
mx.symbol.broadcast_axes(...)
```

# Arguments

data	NDArray-or-Symbol The input
axis	Shape(tuple), optional, default=[] The axes to perform the broadcasting.
size	Shape(tuple), optional, default=[] Target sizes of the broadcasting axes.
name	string, optional Name of the resulting symbol.

### **Details**

'broadcast\_axes' is an alias to the function 'broadcast\_axis'.

# Example::

```
// given x of shape (1,2,1) x = [[[ 1.], [ 2.]]]
```

// broadcast x on on axis 2 broadcast\_axis(x, axis=2, size=3) = [[[ 1., 1., 1.], [ 2., 2., 2.]]] // broadcast x on on axes 0 and 2 broadcast\_axis(x, axis=(0,2), size=(2,3)) = [[[ 1., 1., 1.], [ 2., 2., 2.]], [[ 1., 1., 1.], [ 2., 2., 2.]]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L93

# Value

```
mx.symbol.broadcast_axis
```

broadcast\_axis:Broadcasts the input array over particular axes.

# **Description**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

# Usage

```
mx.symbol.broadcast_axis(...)
```

# Arguments

data	NDArray-or-Symbol The input
axis	Shape(tuple), optional, default=[] The axes to perform the broadcasting.
size	Shape(tuple), optional, default=[] Target sizes of the broadcasting axes.
name	string, optional Name of the resulting symbol.

### **Details**

'broadcast\_axes' is an alias to the function 'broadcast\_axis'.

# Example::

```
// given x of shape (1,2,1) x = [[[ 1.], [ 2.]]]
```

```
// broadcast x on on axis 2 broadcast_axis(x, axis=2, size=3) = [[[ 1., 1., 1.], [ 2., 2., 2.]]] // broadcast x on on axes 0 and 2 broadcast_axis(x, axis=(0,2), size=(2,3)) = [[[ 1., 1., 1.], [ 2., 2., 2.]], [[ 1., 1., 1.], [ 2., 2., 2.]]]
```

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L93

# Value

mx.symbol.broadcast\_div

broadcast\_div:Returns element-wise division of the input arrays with broadcasting.

# **Description**

Example::

### Usage

```
mx.symbol.broadcast_div(...)
```

# **Arguments**

1hsNDArray-or-Symbol First input to the functionrhsNDArray-or-Symbol Second input to the functionnamestring, optional Name of the resulting symbol.

#### **Details**

# Value

out The result mx.symbol

```
mx.symbol.broadcast_equal
```

broadcast\_equal:Returns the result of element-wise \*\*equal to\*\* (==) comparison operation with broadcasting.

# **Description**

Example::

### Usage

```
mx.symbol.broadcast_equal(...)
```

## **Arguments**

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function
name	string, optional Name of the resulting symbol.

#### **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_equal(x,y) &= [[\ 0.,\ 0.,\ 0.],\ [\ 1.,\ 1.,\ 1.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L46 \end{aligned}
```

### Value

out The result mx.symbol

```
mx.symbol.broadcast_greater
```

 $broadcast\_greater:Returns$  the result of element-wise \*\*greater than\*\*(>) comparison operation with broadcasting.

# **Description**

Example::

### Usage

```
mx.symbol.broadcast_greater(...)
```

# Arguments

1hs NDArray-or-Symbol First input to the functionrhs NDArray-or-Symbol Second input to the functionname string, optional Name of the resulting symbol.

#### **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_greater(x,\ y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 0.,\ 0.,\ 0.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L82 \end{aligned}
```

# Value

```
mx.symbol.broadcast_greater_equal
```

broadcast\_greater\_equal:Returns the result of element-wise \*\*greater than or equal to\*\*(>=) comparison operation with broadcasting.

# **Description**

Example::

# Usage

```
\verb|mx.symbol.broadcast_greater_equal(...)|
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

### **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_greater\_equal(x,y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L100 \end{aligned}
```

### Value

out The result mx.symbol

```
mx.symbol.broadcast_hypot
```

broadcast\_hypot: Returns the hypotenuse of a right angled triangle, given its "legs" with broadcasting.

# Description

It is equivalent to doing :math:  $\frac{1^2 + x_2^2}{\cdot}$ .

# Usage

```
mx.symbol.broadcast_hypot(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

### **Details**

```
Example::
```

```
x = [[ 3., 3., 3.]]

y = [[ 4.], [ 4.]]

broadcast_hypot(x, y) = [[ 5., 5., 5.], [ 5., 5., 5.]]

z = [[ 0.], [ 4.]]

broadcast_hypot(x, z) = [[ 3., 3., 3.], [ 5., 5., 5.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L158

#### Value

out The result mx.symbol

```
mx.symbol.broadcast_lesser
```

broadcast\_lesser:Returns the result of element-wise \*\*lesser than\*\* (<) comparison operation with broadcasting.

# **Description**

Example::

## Usage

```
mx.symbol.broadcast_lesser(...)
```

### **Arguments**

1hsNDArray-or-Symbol First input to the functionrhsNDArray-or-Symbol Second input to the functionnamestring, optional Name of the resulting symbol.

# **Details**

```
x = [[1., 1., 1.], [1., 1., 1.]]

y = [[0.], [1.]]

broadcast_lesser(x, y) = [[0., 0., 0.], [0., 0., 0.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L118

# Value

out The result mx.symbol

```
mx.symbol.broadcast_lesser_equal
```

broadcast\_lesser\_equal:Returns the result of element-wise \*\*lesser than or equal to\*\*(<=) comparison operation with broadcasting.

# Description

Example::

# Usage

```
mx.symbol.broadcast_lesser_equal(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

### **Details**

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]

y = [[ 0.], [ 1.]]

broadcast_lesser_equal(x, y) = [[ 0., 0., 0.], [ 1., 1., 1.]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_logic.cc:L136
```

## Value

```
mx.symbol.broadcast_like
```

broadcast\_like:Broadcasts lhs to have the same shape as rhs.

# Description

Broadcasting is a mechanism that allows NDArrays to perform arithmetic operations with arrays of different shapes efficiently without creating multiple copies of arrays. Also see, 'Broadcasting <a href="https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html">https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html</a> '\_ for more explanation.

### Usage

```
mx.symbol.broadcast_like(...)
```

# Arguments

lhs	NDArray-or-Symbol First input.
rhs	NDArray-or-Symbol Second input.
lhs.axes	Shape or None, optional, default=None Axes to perform broadcast on in the first input array
rhs.axes	Shape or None, optional, default=None Axes to copy from the second input array
name	string, optional Name of the resulting symbol.

### **Details**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

For example::

```
broadcast_like([[1,2,3]], [[5,6,7],[7,8,9]]) = [[ 1., 2., 3.], [ 1., 2., 3.]])
broadcast_like([9], [1,2,3,4,5], lhs_axes=(0,), rhs_axes=(-1,)) = [9,9,9,9,9]
Defined in src/operator/tensor/broadcast_reduce_op_value.cc:L180
```

### Value

mx.symbol.broadcast\_logical\_and

broadcast\_logical\_and:Returns the result of element-wise \*\*logical and\*\* with broadcasting.

# Description

Example::

# Usage

```
mx.symbol.broadcast_logical_and(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

# **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_logical\_and(x,\ y) &= [[\ 0.,\ 0.,\ 0.],\ [\ 1.,\ 1.,\ 1.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L154 \end{aligned}
```

# Value

out The result mx.symbol

```
mx.symbol.broadcast_logical_or
```

broadcast\_logical\_or:Returns the result of element-wise \*\*logical or\*\* with broadcasting.

# Description

Example::

# Usage

```
mx.symbol.broadcast_logical_or(...)
```

## **Arguments**

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function
name	string, optional Name of the resulting symbol.

#### **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 0.],\ [\ 1.,\ 1.,\ 0.]] \\ y &= [[\ 1.],\ [\ 0.]] \\ broadcast\_logical\_or(x,\ y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 0.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L172 \end{aligned}
```

### Value

out The result mx.symbol

```
mx.symbol.broadcast_logical_xor
```

broadcast\_logical\_xor:Returns the result of element-wise \*\*logical xor\*\* with broadcasting.

# **Description**

Example::

### Usage

```
mx.symbol.broadcast_logical_xor(...)
```

# Arguments

1hsNDArray-or-Symbol First input to the functionrhsNDArray-or-Symbol Second input to the functionnamestring, optional Name of the resulting symbol.

#### **Details**

```
x = [[1., 1., 0.], [1., 1., 0.]]

y = [[1.], [0.]]

broadcast_logical_xor(x, y) = [[0., 0., 1.], [1., 1., 0.]]

Defined in src/operator/tensor/elemwise_binary_broadcast_op_logic.cc:L190
```

# Value

```
mx.symbol.broadcast_maximum
```

broadcast\_maximum:Returns element-wise maximum of the input arrays with broadcasting.

# **Description**

This function compares two input arrays and returns a new array having the element-wise maxima.

# Usage

```
mx.symbol.broadcast_maximum(...)
```

### **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

#### **Details**

```
Example::
```

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
broadcast_maximum(x, y) = [[ 1., 1., 1.], [ 1., 1., 1.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L81

### Value

out The result mx.symbol

```
mx.symbol.broadcast_minimum
```

broadcast\_minimum:Returns element-wise minimum of the input arrays with broadcasting.

### **Description**

This function compares two input arrays and returns a new array having the element-wise minima.

### Usage

```
mx.symbol.broadcast_minimum(...)
```

## **Arguments**

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function
name	string, optional Name of the resulting symbol.

### **Details**

# Example::

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
broadcast_maximum(x, y) = [[ 0., 0., 0.], [ 1., 1., 1.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L117

### Value

out The result mx.symbol

```
mx.symbol.broadcast_minus
```

broadcast\_minus:Returns element-wise difference of the input arrays with broadcasting.

# Description

'broadcast\_minus' is an alias to the function 'broadcast\_sub'.

# Usage

```
mx.symbol.broadcast_minus(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

# Details

# Example::

```
x = [[1., 1., 1.], [1., 1., 1.]]

y = [[0.], [1.]]

broadcast_sub(x, y) = [[1., 1., 1.], [0., 0., 0.]]

broadcast_minus(x, y) = [[1., 1., 1.], [0., 0., 0.]]
```

Supported sparse operations:

broadcast\_sub/minus(csr, dense(1D)) = dense broadcast\_sub/minus(dense(1D), csr) = dense

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L106

#### Value

out The result mx.symbol

```
mx.symbol.broadcast_mod
```

broadcast\_mod:Returns element-wise modulo of the input arrays with broadcasting.

# Description

Example::

### Usage

```
mx.symbol.broadcast_mod(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

# **Details**

```
x = [[8., 8., 8.], [8., 8., 8.]]

y = [[2.], [3.]]

broadcast_mod(x, y) = [[0., 0., 0.], [2., 2., 2.]]
```

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L222

### Value

mx.symbol.broadcast\_mul

broadcast\_mul:Returns element-wise product of the input arrays with broadcasting.

# **Description**

Example::

### Usage

```
mx.symbol.broadcast_mul(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

#### **Details**

# Value

out The result mx.symbol

```
mx.symbol.broadcast_not_equal
```

 $broadcast\_not\_equal:Returns$  the result of element-wise \*\*not equal  $to^{**}$  (!=) comparison operation with broadcasting.

# **Description**

Example::

# Usage

```
mx.symbol.broadcast_not_equal(...)
```

# Arguments

lhs	NDArray-or-Symbol First input to the function
rhs	NDArray-or-Symbol Second input to the function
name	string, optional Name of the resulting symbol.

### **Details**

```
 \begin{aligned} x &= [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] \\ y &= [[\ 0.],\ [\ 1.]] \\ broadcast\_not\_equal(x,\ y) &= [[\ 1.,\ 1.,\ 1.],\ [\ 0.,\ 0.,\ 0.]] \\ Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_logic.cc:L64 \end{aligned}
```

### Value

out The result mx.symbol

```
mx.symbol.broadcast_plus
```

broadcast\_plus:Returns element-wise sum of the input arrays with broadcasting.

# **Description**

'broadcast\_plus' is an alias to the function 'broadcast\_add'.

### Usage

```
mx.symbol.broadcast_plus(...)
```

# Arguments

1hsNDArray-or-Symbol First input to the functionrhsNDArray-or-Symbol Second input to the functionnamestring, optional Name of the resulting symbol.

#### **Details**

```
Example::
```

```
x = [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] y = [[\ 0.],\ [\ 1.]] broadcast_add(x, y) = [[\ 1.,\ 1.,\ 1.],\ [\ 2.,\ 2.,\ 2.]] broadcast_plus(x, y) = [[\ 1.,\ 1.,\ 1.],\ [\ 2.,\ 2.,\ 2.]]
Supported sparse operations: broadcast_add(csr, dense(1D)) = dense broadcast_add(dense(1D), csr) = dense Defined in src/operator/tensor/elemwise_binary_broadcast_op_basic.cc:L58
```

# Value

out The result mx.symbol

mx.symbol.broadcast\_power

broadcast\_power:Returns result of first array elements raised to powers from second array, element-wise with broadcasting.

# Description

Example::

# Usage

```
mx.symbol.broadcast_power(...)
```

# **Arguments**

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

### **Details**

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]
y = [[ 0.], [ 1.]]
broadcast_power(x, y) = [[ 2., 2., 2.], [ 4., 4., 4.]]
```

 $Defined\ in\ src/operator/tensor/elemwise\_binary\_broadcast\_op\_extended.cc:L45$ 

## Value

```
mx.symbol.broadcast_sub
```

broadcast\_sub:Returns element-wise difference of the input arrays with broadcasting.

# Description

'broadcast\_minus' is an alias to the function 'broadcast\_sub'.

# Usage

```
mx.symbol.broadcast_sub(...)
```

# Arguments

1hs NDArray-or-Symbol First input to the function
 rhs NDArray-or-Symbol Second input to the function
 name string, optional Name of the resulting symbol.

### **Details**

```
Example::
```

```
x = [[ 1., 1., 1.], [ 1., 1., 1.]]

y = [[ 0.], [ 1.]]

broadcast_sub(x, y) = [[ 1., 1., 1.], [ 0., 0., 0.]]

broadcast_minus(x, y) = [[ 1., 1., 1.], [ 0., 0., 0.]]
```

Supported sparse operations:

 $broadcast\_sub/minus(csr, dense(1D)) = dense\ broadcast\_sub/minus(dense(1D), csr) = dense$ 

Defined in src/operator/tensor/elemwise\_binary\_broadcast\_op\_basic.cc:L106

# Value

```
mx.symbol.broadcast_to
```

broadcast\_to:Broadcasts the input array to a new shape.

# **Description**

Broadcasting is a mechanism that allows NDArrays to perform arithmetic operations with arrays of different shapes efficiently without creating multiple copies of arrays. Also see, 'Broadcasting <a href="https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html">https://docs.scipy.org/doc/numpy/user/basics.broadcasting.html</a> '\_ for more explanation.

## Usage

```
mx.symbol.broadcast_to(...)
```

# **Arguments**

data	NDArray-	or-Symbol	The input

shape Shape(tuple), optional, default=[] The shape of the desired array. We can set the

dim to zero if it's same as the original. E.g 'A = broadcast\_to(B, shape=(10, 0,

0))' has the same meaning as 'A = broadcast\_axis(B, axis=0, size=10)'.

name string, optional Name of the resulting symbol.

# **Details**

Broadcasting is allowed on axes with size 1, such as from '(2,1,3,1)' to '(2,8,3,9)'. Elements will be duplicated on the broadcasted axes.

For example::

```
broadcast_to([[1,2,3]], shape=(2,3)) = [[1., 2., 3.], [1., 2., 3.]])
```

The dimension which you do not want to change can also be kept as '0' which means copy the original value. So with 'shape=(2,0)', we will obtain the same result as in the above example.

Defined in src/operator/tensor/broadcast\_reduce\_op\_value.cc:L117

## Value

322 mx.symbol.cast

mx.symbol.Cast

Cast: Casts all elements of the input to a new type.

### **Description**

```
.. note:: "Cast" is deprecated. Use "cast" instead.
```

# Usage

```
mx.symbol.Cast(...)
```

## **Arguments**

data NDArray-or-Symbol The input.

dtype 'bfloat16', 'bool', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8',

required Output data type.

name string, optional Name of the resulting symbol.

#### **Details**

## Example::

```
cast([0.9, 1.3], dtype='int32') = [0, 1] cast([1e20, 11.1], dtype='float16') = [inf, 11.09375] cast([300, 11.1, 10.9, -1, -3], dtype='uint8') = [44, 11, 10, 255, 253]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L665

### Value

out The result mx.symbol

mx.symbol.cast

cast: Casts all elements of the input to a new type.

# Description

```
.. note:: "Cast" is deprecated. Use "cast" instead.
```

### Usage

```
mx.symbol.cast(...)
```

### **Arguments**

data NDArray-or-Symbol The input.

dtype 'bfloat16', 'bool', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8',

required Output data type.

name string, optional Name of the resulting symbol.

#### **Details**

```
Example::
```

```
cast([0.9, 1.3], dtype='int32') = [0, 1] cast([1e20, 11.1], dtype='float16') = [inf, 11.09375] cast([300, 11.1, 10.9, -1, -3], dtype='uint8') = [44, 11, 10, 255, 253]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L665

#### Value

out The result mx.symbol

```
mx.symbol.cast_storage
```

cast\_storage: Casts tensor storage type to the new type.

### **Description**

When an NDArray with default storage type is cast to csr or row\_sparse storage, the result is compact, which means:

### Usage

```
mx.symbol.cast_storage(...)
```

## **Arguments**

data NDArray-or-Symbol The input.

stype 'csr', 'default', 'row\_sparse', required Output storage type.

name string, optional Name of the resulting symbol.

### **Details**

- for csr, zero values will not be retained - for row\_sparse, row slices of all zeros will not be retained The storage type of "cast\_storage" output depends on stype parameter:

```
- cast_storage(csr, 'default') = default - cast_storage(row_sparse, 'default') = default - cast_storage(default, 'csr') = csr - cast_storage(default, 'row_sparse') = row_sparse - cast_storage(csr, 'csr') = csr - cast_storage(row_sparse, 'row_sparse') = row_sparse
```

### Example::

```
dense = [[ 0., 1., 0.], [ 2., 0., 3.], [ 0., 0., 0.], [ 0., 0., 0.]]
```

```
# cast to row_sparse storage type rsp = cast_storage(dense, 'row_sparse') rsp.indices = [0, 1] rsp.values = [[ 0., 1., 0.], [ 2., 0., 3.]]
```

```
# cast to csr storage type csr = cast_storage(dense, 'csr') csr.indices = [1, 0, 2] csr.values = [1, 2, 3] csr.indptr = [0, 1, 3, 3, 3]
```

Defined in src/operator/tensor/cast\_storage.cc:L71

324 mx.symbol.ceil

# Value

out The result mx.symbol

mx.symbol.cbrt

cbrt:Returns element-wise cube-root value of the input.

# **Description**

```
.. math:: cbrt(x) = \sqrt{3}x
```

# Usage

```
mx.symbol.cbrt(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

# **Details**

```
Example::
```

```
cbrt([1, 8, -125]) = [1, 2, -5]
```

The storage type of "cbrt" output depends upon the input storage type:

-  $cbrt(default) = default - cbrt(row\_sparse) = row\_sparse - cbrt(csr) = csr$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L270

## Value

out The result mx.symbol

mx.symbol.ceil

ceil:Returns element-wise ceiling of the input.

# Description

The ceil of the scalar x is the smallest integer i, such that  $i \ge x$ .

# Usage

```
mx.symbol.ceil(...)
```

#### Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

#### Example::

```
ceil([-2.1, -1.9, 1.5, 1.9, 2.1]) = [-2., -1., 2., 2., 3.]
```

The storage type of "ceil" output depends upon the input storage type:

- ceil(default) = default - ceil(row\_sparse) = row\_sparse - ceil(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L818

#### Value

out The result mx.symbol

```
mx.symbol.choose_element_0index
```

choose\_element\_0index:Picks elements from an input array according to the input indices along the given axis.

#### **Description**

Given an input array of shape "(d0, d1)" and indices of shape "(i0,)", the result will be an output array of shape "(i0,)" with::

### Usage

```
mx.symbol.choose_element_0index(...)
```

# Arguments

data	NDArray-or-Symbol The input array
index	NDArray-or-Symbol The index array
axis	int or None, optional, default='-1' int or None. The axis to picking the elements. Negative values means indexing from right to left. If is 'None', the elements in the index w.r.t the flattened input will be picked.

keepdims boolean, optional, default=0 If true, the axis where we pick the elements is left

in the result as dimension with size one.

mode 'clip', 'wrap', optional, default='clip' Specify how out-of-bound indices behave.

Default is "clip". "clip" means clip to the range. So, if all indices mentioned are too large, they are replaced by the index that addresses the last element along an

axis. "wrap" means to wrap around.

name string, optional Name of the resulting symbol.

326 mx.symbol.clip

#### **Details**

```
output[i] = input[i, indices[i]]
```

By default, if any index mentioned is too large, it is replaced by the index that addresses the last element along an axis (the 'clip' mode).

This function supports n-dimensional input and (n-1)-dimensional indices arrays.

Examples::

$$x = [[1., 2.], [3., 4.], [5., 6.]]$$

// picks elements with specified indices along axis 0 pick(x, y=[0,1], 0) = [1., 4.]

// picks elements with specified indices along axis 1 pick(x, y=[0,1,0], 1) = [1., 4., 5.]

// picks elements with specified indices along axis 1 using 'wrap' mode // to place indicies that would normally be out of bounds pick(x, y=[2,-1,-2], 1, mode='wrap') = [1,,4,,5]

$$y = [[1.], [0.], [2.]]$$

// picks elements with specified indices along axis 1 and dims are maintained pick(x, y, 1, keep-dims=True) = [[2.], [3.], [6.]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L151

#### Value

out The result mx.symbol

mx.symbol.clip

clip:Clips (limits) the values in an array. Given an interval, values outside the interval are clipped to the interval edges. Clipping "x" between 'a\_min' and 'a\_max' would be:: .. math:: clip(x, a\_min, a\_max) = \max(\min(x, a\_max), a\_min)) Example:: x = [0, 1, 2, 3, 4, 5, 6, 7, 8, 9] clip(x,1,8) = [1., 1., 2., 3., 4., 5., 6., 7., 8., 8.] The storage type of "clip" output depends on storage types of inputs and the a\_min, a\_max \ parameter values: - clip(default) = default - clip(row\_sparse, a\_min <= 0, a\_max >= 0) = row\_sparse - clip(csr, a\_min <= 0, a\_max >= 0) = csr - clip(row\_sparse, a\_min < 0, a\_max < 0) = default - clip(row\_sparse, a\_min > 0, a\_max > 0) = default - clip(csr, a\_min < 0, a\_max < 0) = csr - clip(csr, a\_min > 0, a\_max > 0) = csr

#### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L677

### Usage

```
mx.symbol.clip(...)
```

mx.symbol.col2im 327

#### **Arguments**

data	NDArray-or-Symbol Input array.
a.min	float, required Minimum value
a.max	float, required Maximum value
name	string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.col2im	col2im:Combining the output column matrix of im2col back to image
	array.

# Description

Like :class: '~mxnet.ndarray.im2col', this operator is also used in the vanilla convolution implementation. Despite the name, col2im is not the reverse operation of im2col. Since there may be overlaps between neighbouring sliding blocks, the column elements cannot be directly put back into image. Instead, they are accumulated (i.e., summed) in the input image just like the gradient computation, so col2im is the gradient of im2col and vice versa.

### Usage

```
mx.symbol.col2im(...)
```

# Arguments

data	NDArray-or-Symbol Input array to combine sliding blocks.
output.size	Shape(tuple), required The spatial dimension of image array: (w,), (h, w) or (d, h, w).
kernel	Shape(tuple), required Sliding kernel size: (w,), (h, w) or (d, h, w).
stride	Shape(tuple), optional, default=[] The stride between adjacent sliding blocks in spatial dimension: $(w,)$ , $(h, w)$ or $(d, h, w)$ . Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] The spacing between adjacent kernel points: $(w,)$ , $(h, w)$ or $(d, h, w)$ . Defaults to 1 for each dimension.
pad	Shape(tuple), optional, default=[] The zero-value padding size on both sides of spatial dimension: $(w,)$ , $(h, w)$ or $(d, h, w)$ . Defaults to no padding.
name	string, optional Name of the resulting symbol.

328 mx.symbol.Concat

### **Details**

Using the notation in im2col, given an input column array of shape :math: '(N, C \times \prod(\textkernel), W)', this operator accumulates the column elements into output array of shape :math: '(N, C, \textout-put\_size[0], \textoutput\_size[1], ...)'. Only 1-D, 2-D and 3-D of spatial dimension is supported in this operator.

Defined in src/operator/nn/im2col.cc:L182

#### Value

out The result mx.symbol

mx.symbol.Concat

Perform an feature concat on channel dim (dim 1) over all the inputs.

### **Description**

Perform an feature concat on channel dim (dim 1) over all the inputs.

#### Usage

```
mx.symbol.Concat(data, num.args, dim = NULL, name = NULL)
```

# Arguments

data list, required List of tensors to concatenate

num. args int, required Number of inputs to be concated.

dim int, optional, default='1' the dimension to be concated.

name string, optional Name of the resulting symbol.

#### Value

mx.symbol.concat 329

mΥ	SVMho	l.concat

Perform an feature concat on channel dim (dim 1) over all the inputs.

# Description

Perform an feature concat on channel dim (dim 1) over all the inputs.

# Usage

```
mx.symbol.concat(data, num.args, dim = NULL, name = NULL)
```

### **Arguments**

data list, required List of tensors to concatenate num.args int, required Number of inputs to be concated.

dim int, optional, default='1' the dimension to be concated.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

```
mx.symbol.Convolution Convolution: Compute *N*-D convolution on *(N+2)*-D input.
```

### **Description**

In the 2-D convolution, given input data with shape \*(batch\_size, channel, height, width)\*, the output is computed by

# Usage

```
mx.symbol.Convolution(...)
```

### **Arguments**

data	NDArray-or-Symbol Input data to the ConvolutionOp.
weight	NDArray-or-Symbol Weight matrix.
bias	NDArray-or-Symbol Bias parameter.
kernel	Shape(tuple), required Convolution kernel size: (w,), (h, w) or (d, h, w)
stride	Shape(tuple), optional, default=[] Convolution stride: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] Convolution dilate: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.

pad Shape(tuple), optional, default=[] Zero pad for convolution: (w,), (h, w) or (d,

h, w). Defaults to no padding.

num.filter int (non-negative), required Convolution filter(channel) number num.group int (non-negative), optional, default=1 Number of group partitions.

workspace long (non-negative), optional, default=1024 Maximum temporary workspace al-

lowed (MB) in convolution. This parameter has two usages. When CUDNN is not used, it determines the effective batch size of the convolution kernel. When CUDNN is used, it controls the maximum temporary storage used for tuning the

best CUDNN kernel when 'limited\_workspace' strategy is used.

no.bias boolean, optional, default=0 Whether to disable bias parameter.

cudnn.tune None, 'fastest', 'limited\_workspace', 'off',optional, default='None' Whether to

pick convolution algo by running performance test.

cudnn.off boolean, optional, default=0 Turn off cudnn for this layer.

layout None, 'NCDHW', 'NCHW', 'NDHWC', 'NHWC', optional, default='None'

Set layout for input, output and weight. Empty for default layout: NCW for 1d, NCHW for 2d and NCDHW for 3d.NHWC and NDHWC are only supported on

GPU.

name string, optional Name of the resulting symbol.

#### **Details**

.. math::

 $out[n,i,:,:] = bias[i] + \sum_{j=0}^{n} data[n,j,:,:] \operatorname{star weight}[i,j,:,:]$ 

where :math: '\star' is the 2-D cross-correlation operator.

For general 2-D convolution, the shapes are

- \*\*data\*\*: \*(batch\_size, channel, height, width)\* - \*\*weight\*\*: \*(num\_filter, channel, kernel[0], kernel[1])\* - \*\*bias\*\*: \*(num\_filter,)\* - \*\*out\*\*: \*(batch\_size, num\_filter, out\_height, out\_width)\*.

Define::

f(x,k,p,s,d) = floor((x+2\*p-d\*(k-1)-1)/s)+1

then we have::

out\_height=f(height, kernel[0], pad[0], stride[0], dilate[0]) out\_width=f(width, kernel[1], pad[1], stride[1], dilate[1])

If "no\_bias" is set to be true, then the "bias" term is ignored.

The default data "layout" is \*NCHW\*, namely \*(batch\_size, channel, height, width)\*. We can choose other layouts such as \*NWC\*.

If "num\_group" is larger than 1, denoted by \*g\*, then split the input "data" evenly into \*g\* parts along the channel axis, and also evenly split "weight" along the first dimension. Next compute the convolution on the \*i\*-th part of the data with the \*i\*-th weight part. The output is obtained by concatenating all the \*g\* results.

1-D convolution does not have \*height\* dimension but only \*width\* in space.

```
- **data**: *(batch_size, channel, width)* - **weight**: *(num_filter, channel, kernel[0])* - **bias**: *(num_filter,)* - **out**: *(batch_size, num_filter, out_width)*.
```

- 3-D convolution adds an additional \*depth\* dimension besides \*height\* and \*width\*. The shapes are
- \*\*data\*\*: \*(batch\_size, channel, depth, height, width)\* \*\*weight\*\*: \*(num\_filter, channel, kernel[0], kernel[1], kernel[2])\* \*\*bias\*\*: \*(num\_filter,)\* \*\*out\*\*: \*(batch\_size, num\_filter, out\_depth, out\_height, out\_width)\*.

Both "weight" and "bias" are learnable parameters.

There are other options to tune the performance.

- \*\*cudnn\_tune\*\*: enable this option leads to higher startup time but may give faster speed. Options are
- \*\*off\*\*: no tuning \*\*limited\_workspace\*\*:run test and pick the fastest algorithm that doesn't exceed workspace limit. \*\*fastest\*\*: pick the fastest algorithm and ignore workspace limit. \*\*None\*\* (default): the behavior is determined by environment variable "MXNET\_CUDNN\_AUTOTUNE\_DEFAULT". 0 for off, 1 for limited workspace (default), 2 for fastest.
- \*\*workspace\*\*: A large number leads to more (GPU) memory usage but may improve the performance.

Defined in src/operator/nn/convolution.cc:L469

#### Value

out The result mx.symbol

```
mx.symbol.Convolution_v1
```

Convolution\_v1:This operator is DEPRECATED. Apply convolution to input then add a bias.

#### Description

Convolution\_v1:This operator is DEPRECATED. Apply convolution to input then add a bias.

#### Usage

```
mx.symbol.Convolution_v1(...)
```

#### **Arguments**

data	NDArray-or-Symbol Input data to the Convolution V1Op.
weight	NDArray-or-Symbol Weight matrix.
bias	NDArray-or-Symbol Bias parameter.
kernel	Shape(tuple), required convolution kernel size: (h, w) or (d, h, w)
stride	Shape(tuple), optional, default=[] convolution stride: (h, w) or (d, h, w)
dilate	Shape(tuple), optional, default=[] convolution dilate: (h, w) or (d, h, w)
pad	Shape(tuple), optional, default=[] pad for convolution: (h, w) or (d, h, w)

num.filter int (non-negative), required convolution filter(channel) number

int (non-negative), optional, default=1 Number of group partitions. Equivalent num.group

to slicing input into num group partitions, apply convolution on each, then con-

catenate the results

long (non-negative), optional, default=1024 Maximum temporary workspace alworkspace

> lowed for convolution (MB). This parameter determines the effective batch size of the convolution kernel, which may be smaller than the given batch size. Also, the workspace will be automatically enlarged to make sure that we can run the

kernel with batch\_size=1

no.bias boolean, optional, default=0 Whether to disable bias parameter.

None, 'fastest', 'limited\_workspace', 'off',optional, default='None' Whether to cudnn.tune

> pick convolution algo by running performance test. Leads to higher startup time but may give faster speed. Options are: 'off': no tuning 'limited\_workspace': run test and pick the fastest algorithm that doesn't exceed workspace limit. 'fastest': pick the fastest algorithm and ignore workspace limit. If set to None

(default), behavior is determined by environment variable MXNET CUDNN AUTOTUNE DEFAULT:

0 for off, 1 for limited workspace (default), 2 for fastest.

cudnn.off boolean, optional, default=0 Turn off cudnn for this layer.

None, 'NCDHW', 'NCHW', 'NDHWC', 'NHWC', optional, default='None' Set layout

layout for input, output and weight. Empty for default layout: NCHW for 2d

and NCDHW for 3d.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.Correlation Correlation: Applies correlation to inputs.

#### **Description**

The correlation layer performs multiplicative patch comparisons between two feature maps.

#### **Usage**

```
mx.symbol.Correlation(...)
```

### Arguments

data1 NDArray-or-Symbol Input data1 to the correlation. NDArray-or-Symbol Input data2 to the correlation. data2

kernel.size int (non-negative), optional, default=1 kernel size for Correlation must be an

odd number

max.displacement

int (non-negative), optional, default=1 Max displacement of Correlation

stride1 int (non-negative), optional, default=1 stride1 quantize data1 globally

stride2 int (non-negative), optional, default=1 stride2 quantize data2 within the neigh-

borhood centered around data1

pad.size int (non-negative), optional, default=0 pad for Correlation

is.multiply boolean, optional, default=1 operation type is either multiplication or subduction

name string, optional Name of the resulting symbol.

#### **Details**

Given two multi-channel feature maps :math:'f\_1, f\_2', with :math:'w', :math:'h', and :math:'c' being their width, height, and number of channels, the correlation layer lets the network compare each patch from :math:'f\_1' with each patch from :math:'f\_2'.

For now we consider only a single comparison of two patches. The 'correlation' of two patches centered at :math: 'x\_1' in the first map and :math: 'x\_2' in the second map is then defined as:

.. math::

$$c(x_1, x_2) = \sum_{e} (-k,k) < f_1(x_1 + e), f_2(x_2 + e) > c$$

for a square patch of size :math: 'K:=2k+1'.

Note that the equation above is identical to one step of a convolution in neural networks, but instead of convolving data with a filter, it convolves data with other data. For this reason, it has no training weights.

Computing :math:  $c(x_1, x_2)$  involves :math:  $k^2$  multiplications. Comparing all patch combinations involves :math:  $k^2$  such computations.

Given a maximum displacement :math:'d', for each location :math:' $x_1$ ' it computes correlations :math:' $c(x_1, x_2)$ ' only in a neighborhood of size :math:'D:=2d+1', by limiting the range of :math:' $x_2$ '. We use strides :math:' $x_1$ ,  $x_2$ ', to quantize :math:' $x_1$ ' globally and to quantize :math:' $x_2$ ' within the neighborhood centered around :math:' $x_1$ '.

The final output is defined by the following expression:

```
.. math:: out[n, q, i, j] = c(x_i, j, x_q)
```

where :math:'i' and :math:'j' enumerate spatial locations in :math:'f\_1', and :math:'q' denotes the :math:'q^th' neighborhood of :math:'x\_i,j'.

Defined in src/operator/correlation.cc:L198

#### Value

334 mx.symbol.cosh

mx.symbol.cos

cos: Computes the element-wise cosine of the input array.

# Description

The input should be in radians (:math:'2\pi' rad equals 360 degrees).

#### Usage

```
mx.symbol.cos(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

```
.. math:: cos([0, \pi/4, \pi/2]) = [1, 0.707, 0]
```

The storage type of "cos" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L90

### Value

out The result mx.symbol

mx.symbol.cosh

cosh:Returns the hyperbolic cosine of the input array, computed element-wise.

### **Description**

```
.. math:: cosh(x) = 0.5 \times (exp(x) + exp(-x))
```

### Usage

```
mx.symbol.cosh(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.Crop 335

### **Details**

The storage type of "cosh" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L409

### Value

out The result mx.symbol

mx.symbol.Crop Crop:

# Description

.. note:: 'Crop' is deprecated. Use 'slice' instead.

#### Usage

```
mx.symbol.Crop(...)
```

# Arguments

data	Symbol or Symbol[] Tensor or List of Tensors, the second input will be used as crop_like shape reference
num.args	int, required Number of inputs for crop, if equals one, then we will use the h_wfor crop height and width, else if equals two, then we will use the height and width of the second input symbol, we name crop_like here
offset	Shape(tuple), optional, default=[0,0] crop offset coordinate: (y, x)
h.w	Shape(tuple), optional, default=[0,0] crop height and width: (h, w)
center.crop	boolean, optional, default=0 If set to true, then it will use be the center_crop,or it will crop using the shape of crop_like

#### **Details**

name

Crop the 2nd and 3rd dim of input data, with the corresponding size of h\_w or with width and height of the second input symbol, i.e., with one input, we need h\_w to specify the crop height and width, otherwise the second input symbol's size will be used

string, optional Name of the resulting symbol.

Defined in src/operator/crop.cc:L50

#### Value

336 mx.symbol.crop

mx.symbol.crop

crop:Slices a region of the array. .. note:: "crop" is deprecated. Use "slice" instead. This function returns a sliced array between the indices given by 'begin' and 'end' with the corresponding 'step'. For an input array of "shape= $(d_0, d_1, ..., d_{n-1})$ ", slice operation with "begin= $(b_0, b_1...b_m-1)$ ", "end= $(e_0, e_1, ..., e_m-1)$ ", and " $step=(s_0, s_1, ..., s_m-1)$ ", where  $m \le n$ , results in an array with the shape "( $|e\ 0-b\ 0|/|s\ 0|$ , ...,  $|e\ m-1-b\ m-1|/|s\ m-1|$ ,  $d\ m$ , ...,  $d\ n-1$ 1)". The resulting array's \*k\*-th dimension contains elements from the \*k\*-th dimension of the input array starting from index "b k" (inclusive) with step "s\_k" until reaching "e\_k" (exclusive). If the \*k\*-th elements are 'None' in the sequence of 'begin', 'end', and 'step', the following rule will be used to set default values. If 's\_k' is 'None', set  $s_k=1$ . If  $s_k>0$ , set  $b_k=0$ ,  $e_k=d_k$ ; else, set  $b_k=d_k-1$ , 'e\_k=-1'. The storage type of "slice" output depends on storage types of inputs - slice(csr) = csr - otherwise, "slice" generates output with default storage .. note:: When input data storage type is csr, it only supports step=(), or step=(None,), or step=(1,) to generate a csr output. For other step parameter values, it falls back to slicing a dense tensor. Example:: x = [[1., 2., 3., 4.], [5., 6., 7., 8.], [9., 10., 11.]12.]] slice(x, begin=(0,1), end=(2,4)) = [[2., 3., 4.], [6., 7., 8.]]slice(x, begin=(None, 0), end=(None, 3), step=(-1, 2)) = [[9., 11.],[5., 7.], [1., 3.]]

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L482

### Usage

```
mx.symbol.crop(...)
```

#### **Arguments**

data	NDArray-or-Symbol Source input
begin	Shape(tuple), required starting indices for the slice operation, supports negative indices.
end	Shape(tuple), required ending indices for the slice operation, supports negative indices.
step	Shape(tuple), optional, default=[] step for the slice operation, supports negative values.
name	string, optional Name of the resulting symbol.

#### Value

mx.symbol.CTCLoss 337

mx.symbol.CTCLoss

CTCLoss: Connectionist Temporal Classification Loss.

#### **Description**

.. note:: The existing alias "contrib\_CTCLoss" is deprecated.

### Usage

```
mx.symbol.CTCLoss(...)
```

#### **Arguments**

data NDArray-or-Symbol Input ndarray

label NDArray-or-Symbol Ground-truth labels for the loss.

data.lengths NDArray-or-Symbol Lengths of data for each of the samples. Only required

when use\_data\_lengths is true.

label.lengths NDArray-or-Symbol Lengths of labels for each of the samples. Only required

when use\_label\_lengths is true.

use.data.lengths

 $boolean, optional, default = 0 \ Whether the \ data \ lengths \ are \ decided \ by \ `data\_lengths'.$ 

If false, the lengths are equal to the max sequence length.

use.label.lengths

boolean, optional, default=0 Whether the label lengths are decided by 'label\_lengths', or derived from 'padding\_mask'. If false, the lengths are derived from the first occurrence of the value of 'padding\_mask'. The value of 'padding\_mask' is "0" when first CTC label is reserved for blank, and "-1" when last label is reserved

for blank. See 'blank label'.

blank.label 'first', 'last',optional, default='first' Set the label that is reserved for blank la-

bel.If "first", 0-th label is reserved, and label values for tokens in the vocabulary are between "1" and "alphabet\_size-1", and the padding mask is "-1". If "last", last label value "alphabet\_size-1" is reserved for blank label instead, and label values for tokens in the vocabulary are between "0" and "alphabet\_size-2", and

the padding mask is "0".

name string, optional Name of the resulting symbol.

#### Details

The shapes of the inputs and outputs:

```
- **data**: '(sequence_length, batch_size, alphabet_size)' - **label**: '(batch_size, label_sequence_length)'
- **out**: '(batch_size)'
```

The 'data' tensor consists of sequences of activation vectors (without applying softmax), with ith channel in the last dimension corresponding to i-th label for i between 0 and alphabet\_size-1 (i.e always 0-indexed). Alphabet size should include one additional value reserved for blank label.

338 mx.symbol.ctc\_loss

When 'blank\_label' is ""first"", the "0"-th channel is be reserved for activation of blank label, or otherwise if it is "last", "(alphabet\_size-1)"-th channel should be reserved for blank label.

"label" is an index matrix of integers. When 'blank\_label' is ""first"", the value 0 is then reserved for blank label, and should not be passed in this matrix. Otherwise, when 'blank\_label' is ""last"", the value '(alphabet\_size-1)' is reserved for blank label.

If a sequence of labels is shorter than \*label\_sequence\_length\*, use the special padding value at the end of the sequence to conform it to the correct length. The padding value is '0' when 'blank\_label' is ""first"", and '-1' otherwise.

For example, suppose the vocabulary is '[a, b, c]', and in one batch we have three sequences 'ba', 'cbb', and 'abac'. When 'blank\_label' is ""first"", we can index the labels as 'a': 1, 'b': 2, 'c': 3', and we reserve the 0-th channel for blank label in data tensor. The resulting 'label' tensor should be padded to be::

$$[[2, 1, 0, 0], [3, 2, 2, 0], [1, 2, 1, 3]]$$

When 'blank\_label' is ""last"", we can index the labels as "a": 0, 'b": 1, 'c": 2', and we reserve the channel index 3 for blank label in data tensor. The resulting 'label' tensor should be padded to be::

$$[[1, 0, -1, -1], [2, 1, 1, -1], [0, 1, 0, 2]]$$

"out" is a list of CTC loss values, one per example in the batch.

See \*Connectionist Temporal Classification: Labelling Unsegmented Sequence Data with Recurrent Neural Networks\*, A. Graves \*et al\*. for more information on the definition and the algorithm.

Defined in src/operator/nn/ctc\_loss.cc:L100

#### Value

out The result mx.symbol

mx.symbol.ctc\_loss

ctc loss:Connectionist Temporal Classification Loss.

#### **Description**

.. note:: The existing alias "contrib\_CTCLoss" is deprecated.

#### Usage

```
mx.symbol.ctc_loss(...)
```

#### **Arguments**

data NDArray-or-Symbol Input ndarray

label NDArray-or-Symbol Ground-truth labels for the loss.

data.lengths NDArray-or-Symbol Lengths of data for each of the samples. Only required

when use\_data\_lengths is true.

label.lengths NDArray-or-Symbol Lengths of labels for each of the samples. Only required

when use\_label\_lengths is true.

mx.symbol.ctc\_loss 339

use.data.lengths

boolean, optional, default=0 Whether the data lenghts are decided by 'data\_lengths'. If false, the lengths are equal to the max sequence length.

use.label.lengths

boolean, optional, default=0 Whether the label lengths are decided by 'label\_lengths', or derived from 'padding\_mask'. If false, the lengths are derived from the first occurrence of the value of 'padding\_mask'. The value of 'padding\_mask' is "0" when first CTC label is reserved for blank, and "-1" when last label is reserved for blank. See 'blank\_label'.

blank.label

'first', 'last',optional, default='first' Set the label that is reserved for blank label.If "first", 0-th label is reserved, and label values for tokens in the vocabulary are between "1" and "alphabet\_size-1", and the padding mask is "-1". If "last", last label value "alphabet\_size-1" is reserved for blank label instead, and label values for tokens in the vocabulary are between "0" and "alphabet\_size-2", and the padding mask is "0".

name

string, optional Name of the resulting symbol.

#### **Details**

The shapes of the inputs and outputs:

- \*\*data\*\*: '(sequence\_length, batch\_size, alphabet\_size)' - \*\*label\*\*: '(batch\_size, label\_sequence\_length)' - \*\*out\*\*: '(batch\_size)'

The 'data' tensor consists of sequences of activation vectors (without applying softmax), with ith channel in the last dimension corresponding to i-th label for i between 0 and alphabet\_size-1 (i.e always 0-indexed). Alphabet size should include one additional value reserved for blank label. When 'blank\_label' is ""first"", the "0"-th channel is be reserved for activation of blank label, or otherwise if it is "last", "(alphabet\_size-1)"-th channel should be reserved for blank label.

"label" is an index matrix of integers. When 'blank\_label' is ""first"", the value 0 is then reserved for blank label, and should not be passed in this matrix. Otherwise, when 'blank\_label' is ""last"", the value '(alphabet\_size-1)' is reserved for blank label.

If a sequence of labels is shorter than \*label\_sequence\_length\*, use the special padding value at the end of the sequence to conform it to the correct length. The padding value is '0' when 'blank\_label' is ""first"", and '-1' otherwise.

For example, suppose the vocabulary is '[a, b, c]', and in one batch we have three sequences 'ba', 'cbb', and 'abac'. When 'blank\_label' is ""first"", we can index the labels as 'a': 1, 'b': 2, 'c': 3', and we reserve the 0-th channel for blank label in data tensor. The resulting 'label' tensor should be padded to be::

[[2, 1, 0, 0], [3, 2, 2, 0], [1, 2, 1, 3]]

When 'blank\_label' is ""last"", we can index the labels as 'a': 0, 'b': 1, 'c': 2', and we reserve the channel index 3 for blank label in data tensor. The resulting 'label' tensor should be padded to be::

[[1, 0, -1, -1], [2, 1, 1, -1], [0, 1, 0, 2]]

"out" is a list of CTC loss values, one per example in the batch.

See \*Connectionist Temporal Classification: Labelling Unsegmented Sequence Data with Recurrent Neural Networks\*, A. Graves \*et al\*. for more information on the definition and the algorithm.

Defined in src/operator/nn/ctc\_loss.cc:L100

340 mx.symbol.Custom

#### Value

out The result mx.symbol

mx.symbol.cumsum

cumsum: Return the cumulative sum of the elements along a given axis.

#### **Description**

Defined in src/operator/numpy/np\_cumsum.cc:L70

### Usage

```
mx.symbol.cumsum(...)
```

#### **Arguments**

a	NDArray-o	r-Symbol I	nput ndarray
---	-----------	------------	--------------

axis int or None, optional, default='None' Axis along which the cumulative sum is

computed. The default (None) is to compute the cumsum over the flattened

array.

dtype None, 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', optional, default='None'

Type of the returned array and of the accumulator in which the elements are summed. If dtype is not specified, it defaults to the dtype of a, unless a has an integer dtype with a precision less than that of the default platform integer. In

that case, the default platform integer is used.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.Custom Custom: Apply a custom operator implemented in a frontend language (like Python).

### **Description**

Custom operators should override required methods like 'forward' and 'backward'. The custom operator must be registered before it can be used. Please check the tutorial here: https://mxnet.incubator.apache.org/api/faq/new\_

#### Usage

```
mx.symbol.Custom(...)
```

### **Arguments**

data NDArray-or-Symbol[] Input data for the custom operator.

op. type string Name of the custom operator. This is the name that is passed to 'mx.operator.register'

to register the operator.

name string, optional Name of the resulting symbol.

#### **Details**

Defined in src/operator/custom/custom.cc:L547

#### Value

out The result mx.symbol

mx.symbol.Deconvolution

Deconvolution: Computes 1D or 2D transposed convolution (aka fractionally strided convolution) of the input tensor. This operation can be seen as the gradient of Convolution operation with respect to its input. Convolution usually reduces the size of the input. Transposed convolution works the other way, going from a smaller input to a larger output while preserving the connectivity pattern.

### **Description**

Deconvolution:Computes 1D or 2D transposed convolution (aka fractionally strided convolution) of the input tensor. This operation can be seen as the gradient of Convolution operation with respect to its input. Convolution usually reduces the size of the input. Transposed convolution works the other way, going from a smaller input to a larger output while preserving the connectivity pattern.

#### **Usage**

```
mx.symbol.Deconvolution(...)
```

#### Arguments

data	NDArray-or-Symbol Input tensor to the deconvolution operation.
weight	NDArray-or-Symbol Weights representing the kernel.
bias	NDArray-or-Symbol Bias added to the result after the deconvolution operation.
kernel	Shape(tuple), required Deconvolution kernel size: (w,), (h, w) or (d, h, w). This is same as the kernel size used for the corresponding convolution
stride	Shape(tuple), optional, default=[] The stride used for the corresponding convolution: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] Dilation factor for each dimension of the input: (w,), (h, w) or (d, h, w). Defaults to 1 for each dimension.

342 mx.symbol.degrees

pad	Shape(tuple), optional, default=[] The amount of implicit zero padding added during convolution for each dimension of the input: (w,), (h, w) or (d, h, w). "(kernel-1)/2" is usually a good choice. If 'target_shape' is set, 'pad' will be ignored and a padding that will generate the target shape will be used. Defaults to no padding.
adj	Shape(tuple), optional, default=[] Adjustment for output shape: (w,), (h, w) or (d, h, w). If 'target_shape' is set, 'adj' will be ignored and computed accordingly.
target.shape	Shape(tuple), optional, default=[] Shape of the output tensor: $(w,)$ , $(h, w)$ or $(d, h, w)$ .
num.filter	int (non-negative), required Number of output filters.
num.group	int (non-negative), optional, default=1 Number of groups partition.
workspace	long (non-negative), optional, default=512 Maximum temporary workspace allowed (MB) in deconvolution. This parameter has two usages. When CUDNN is not used, it determines the effective batch size of the deconvolution kernel. When CUDNN is used, it controls the maximum temporary storage used for tuning the best CUDNN kernel when 'limited_workspace' strategy is used.
no.bias	boolean, optional, default=1 Whether to disable bias parameter.
cudnn.tune	None, 'fastest', 'limited_workspace', 'off',optional, default='None' Whether to pick convolution algorithm by running performance test.
cudnn.off	boolean, optional, default=0 Turn off cudnn for this layer.
layout	None, 'NCDHW', 'NCHW', 'NCW', 'NDHWC', 'NHWC', optional, default='None' Set layout for input, output and weight. Empty for default layout, NCW for 1d, NCHW for 2d and NCDHW for 3d.NHWC and NDHWC are only supported on GPU.

### Value

name

out The result mx.symbol

 ${\tt mx.symbol.degrees}: Converts\ each\ element\ of\ the\ input\ array\ from\ radians\ to\ degrees.$ 

string, optional Name of the resulting symbol.

# Description

```
.. math:: degrees([0, \pi/2, \pi/2, 2\pi/2, 2\pi/2]) = [0, 90, 180, 270, 360]
```

# Usage

```
mx.symbol.degrees(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

The storage type of "degrees" output depends upon the input storage type:
- degrees(default) = default - degrees(row\_sparse) = row\_sparse - degrees(csr) = csr
Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L332

#### Value

out The result mx.symbol

mx.symbol.depth\_to\_space

depth\_to\_space:Rearranges(permutes) data from depth into blocks of spatial data. Similar to ONNX DepthToSpace operator: https://github.com/onnx/onnx/blob/master/docs/Operators.md#DepthToSpace. The output is a new tensor where the values from depth dimension are moved in spatial blocks to height and width dimension. The reverse of this operation is "space\_to\_depth". .. math:: \begingather\* x \prime =  $reshape(x, [N, block\_size, block\_size, C / (block\_size ^ 2), H *$  $block \leq v + block \leq v + bloc$ [0, 3, 4, 1, 5, 2]\\\\\\y = reshape(x\\\prime\\prime, [N, C/(block\\\_size^\) 2),  $H * block \_size$ ,  $W * block \_size$ ]) \endgather\* where :math: 'x' is an input tensor with default layout as :math: '[N, C, H, W]': [batch, channels, height, width] and :math: 'y' is the output tensor of layout :math: '[N, C / ( $block \le ^ 2$ ),  $H * block \le W * block \le '$ Example:: x = [[[0, 1, 2], [3, 4, 5]], [[6, 7, 8], [9, 10, 11]], [[12, 13, 1]]14], [15, 16, 17]], [[18, 19, 20], [21, 22, 23]]]] depth\_to\_space(x, 2) = [[[0, 6, 1, 7, 2, 8], [12, 18, 13, 19, 14, 20], [3, 9, 4, 10, 5, 11], [15, 15]]21, 16, 22, 17, 23]]]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L972

#### Usage

```
mx.symbol.depth_to_space(...)
```

#### **Arguments**

data NDArray-or-Symbol Input ndarray

block.size int, required Blocks of [block\_size. block\_size] are moved

name string, optional Name of the resulting symbol.

344 mx.symbol.diag

#### Value

out The result mx.symbol

mx.symbol.diag

diag:Extracts a diagonal or constructs a diagonal array.

### **Description**

"diag"'s behavior depends on the input array dimensions:

#### Usage

```
mx.symbol.diag(...)
```

### **Arguments**

data	NDArray-or-Symbol Input ndarray
k	int, optional, default='0' Diagonal in question. The default is 0. Use k>0 for diagonals above the main diagonal, and k<0 for diagonals below the main diagonal. If input has shape (S0 S1) k must be between -S0 and S1
axis1	int, optional, default='0' The first axis of the sub-arrays of interest. Ignored when the input is a 1-D array.
axis2	int, optional, default='1' The second axis of the sub-arrays of interest. Ignored when the input is a 1-D array.
name	string, optional Name of the resulting symbol.

#### **Details**

- 1-D arrays: constructs a 2-D array with the input as its diagonal, all other elements are zero. - N-D arrays: extracts the diagonals of the sub-arrays with axes specified by "axis1" and "axis2". The output shape would be decided by removing the axes numbered "axis1" and "axis2" from the input shape and appending to the result a new axis with the size of the diagonals in question.

For example, when the input shape is '(2, 3, 4, 5)', "axis1" and "axis2" are 0 and 2 respectively and "k" is 0, the resulting shape would be '(3, 5, 2)'.

#### Examples::

```
x = [[1, 2, 3], [4, 5, 6]]
diag(x) = [1, 5]
diag(x, k=1) = [2, 6]
diag(x, k=-1) = [4]
x = [1, 2, 3]
diag(x) = [[1, 0, 0], [0, 2, 0], [0, 0, 3]]
diag(x, k=1) = [[0, 1, 0], [0, 0, 2], [0, 0, 0]]
```

mx.symbol.digamma 345

```
\begin{aligned} & \text{diag}(x, k=-1) = [[0, 0, 0], [1, 0, 0], [0, 2, 0]] \\ & x = [[[1, 2], [3, 4]], \\ & [[5, 6], [7, 8]]] \\ & \text{diag}(x) = [[1, 7], [2, 8]] \\ & \text{diag}(x, k=1) = [[3], [4]] \\ & \text{diag}(x, \text{axis1}=-2, \text{axis2}=-1) = [[1, 4], [5, 8]] \\ & \text{Defined in src/operator/tensor/diag\_op.cc:L87} \end{aligned}
```

#### Value

out The result mx.symbol

 ${\tt mx.symbol.digamma}$ 

 $\label{linear} \textit{digamma:Returns element-wise log derivative of the gamma function} \\ \land \textit{of the input.}$ 

# Description

The storage type of "digamma" output is always dense

# Usage

```
mx.symbol.digamma(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

# Value

346 mx.symbol.dot

mx.symbol.dot	dot:Dot product of two arrays.	

#### **Description**

"dot"'s behavior depends on the input array dimensions:

### **Usage**

```
mx.symbol.dot(...)
```

#### **Arguments**

lhs NDArray-or-Symbol The first input NDArray-or-Symbol The second input rhs boolean, optional, default=0 If true then transpose the first input before dot. transpose.a transpose.b boolean, optional, default=0 If true then transpose the second input before dot. forward.stype None, 'csr', 'default', 'row\_sparse',optional, default='None' The desired storage type of the forward output given by user, if the combination of input storage types and this hint does not matchany implemented ones, the dot operator will

perform fallback operation and still produce an output of the desired storage type.

string, optional Name of the resulting symbol. name

#### **Details**

- 1-D arrays: inner product of vectors - 2-D arrays: matrix multiplication - N-D arrays: a sum product over the last axis of the first input and the first axis of the second input

For example, given 3-D "x" with shape '(n,m,k)' and "y" with shape '(k,r,s)', the result array will have shape '(n,m,r,s)'. It is computed by::

```
dot(x,y)[i,j,a,b] = sum(x[i,j,:]*y[:,a,b])
```

#### Example::

```
x = reshape([0,1,2,3,4,5,6,7], shape=(2,2,2)) y = reshape([7,6,5,4,3,2,1,0], shape=(2,2,2)) dot(x,y)[0,0,1,1]
= 0 \text{ sum}(x[0,0,:]*y[:,1,1]) = 0
```

The storage type of "dot" output depends on storage types of inputs, transpose option and forward\_stype option for output storage type. Implemented sparse operations include:

- dot(default, default, transpose a=True/False, transpose b=True/False) = default - dot(csr, default, transpose\_a=True) = default - dot(csr, default, transpose\_a=True) = row\_sparse - dot(csr, default) = default - dot(csr, row sparse) = default - dot(default, csr) = csr (CPU only) - dot(default, csr, forward\_stype='default') = default - dot(default, csr, transpose\_b=True, forward\_stype='default') = default

If the combination of input storage types and forward\_stype does not match any of the above patterns, "dot" will fallback and generate output with default storage.

.. Note::

mx.symbol.Dropout 347

If the storage type of the lhs is "csr", the storage type of gradient w.r.t rhs will be "row\_sparse". Only a subset of optimizers support sparse gradients, including SGD, AdaGrad and Adam. Note that by default lazy updates is turned on, which may perform differently from standard updates. For more details, please check the Optimization API at: https://mxnet.incubator.apache.org/api/python/optimization/optimization.html Defined in src/operator/tensor/dot.cc:L77

#### Value

out The result mx.symbol

mx.symbol.Dropout	Dropout: Applies dropout operation to input array.	
mx.symbol.Dropout	Dropout: Applies dropout operation to input array.	

#### **Description**

- During training, each element of the input is set to zero with probability p. The whole array is rescaled by :math:  $\frac{1}{1-p}$  to keep the expected sum of the input unchanged.

#### Usage

```
mx.symbol.Dropout(...)
```

#### **Arguments**

data	NDArray-or-Symbol Input array to which dropout will be applied.
р	float, optional, default=0.5 Fraction of the input that gets dropped out during training time.
mode	'always', 'training',optional, default='training' Whether to only turn on dropout during training or to also turn on for inference.
axes	Shape(tuple), optional, default=[] Axes for variational dropout kernel.
cudnn.off	boolean or None, optional, default=0 Whether to turn off cudnn in dropout operator. This option is ignored if axes is specified.
name	string, optional Name of the resulting symbol.

#### **Details**

- During testing, this operator does not change the input if mode is 'training'. If mode is 'always', the same computation as during training will be applied.

#### Example::

```
random.seed(998) input_array = array([[3., 0.5, -0.5, 2., 7.], [2., -0.4, 7., 3., 0.2]]) a = symbol.Variable('a') dropout = symbol.Dropout(a, p = 0.2) executor = dropout.simple_bind(a = input_array.shape)
```

```
## If training executor.forward(is_train = True, a = input_array) executor.outputs [[ 3.75 0.625 -0. 2.5 8.75 ] [ 2.5 -0.5 8.75 3.75 0. ]]
```

```
## If testing executor.forward(is_train = False, a = input_array) executor.outputs [[ 3. 0.5 -0.5 2. 7. ] [ 2. -0.4 7. 3. 0.2 ]]
```

Defined in src/operator/nn/dropout.cc:L96

#### Value

out The result mx.symbol

```
mx.symbol.ElementWiseSum
```

ElementWiseSum:Adds all input arguments element-wise.

# Description

```
.. math:: add\_n(a_1, a_2, ..., a_n) = a_1 + a_2 + ... + a_n
```

### Usage

```
mx.symbol.ElementWiseSum(...)
```

### **Arguments**

args NDArray-or-Symbol[] Positional input arguments

name string, optional Name of the resulting symbol.

#### **Details**

"add\_n" is potentially more efficient than calling "add" by 'n' times.

The storage type of "add\_n" output depends on storage types of inputs

- add\_n(row\_sparse, row\_sparse, ..) = row\_sparse - add\_n(default, csr, default) = default - add\_n(any input combinations longer than 4 (>4) with at least one default type) = default - otherwise, "add\_n" falls all inputs back to default storage and generates default storage

Defined in src/operator/tensor/elemwise\_sum.cc:L155

### Value

```
mx.symbol.elemwise_add
```

elemwise\_add:Adds arguments element-wise.

# Description

The storage type of "elemwise\_add" output depends on storage types of inputs

#### Usage

```
mx.symbol.elemwise_add(...)
```

#### Arguments

1hs NDArray-or-Symbol first inputrhs NDArray-or-Symbol second input

name string, optional Name of the resulting symbol.

#### **Details**

- elemwise\_add(row\_sparse, row\_sparse) = row\_sparse - elemwise\_add(csr, csr) = csr - elemwise\_add(default, csr) = default - elemwise\_add(csr, default) = default - elemwise\_add(default, rsp) = default - elemwise\_add(rsp, default) = default - otherwise, "elemwise\_add" generates output with default storage

#### Value

out The result mx.symbol

```
mx.symbol.elemwise_div
```

elemwise\_div:Divides arguments element-wise.

# Description

The storage type of "elemwise\_div" output is always dense

### Usage

```
mx.symbol.elemwise_div(...)
```

#### **Arguments**

1hs NDArray-or-Symbol first inputrhs NDArray-or-Symbol second input

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

```
mx.symbol.elemwise_mul
```

elemwise\_mul:Multiplies arguments element-wise.

### **Description**

The storage type of "elemwise\_mul" output depends on storage types of inputs

#### Usage

```
mx.symbol.elemwise_mul(...)
```

### Arguments

1hs NDArray-or-Symbol first inputrhs NDArray-or-Symbol second input

name string, optional Name of the resulting symbol.

#### **Details**

- elemwise\_mul(default, default) = default - elemwise\_mul(row\_sparse, row\_sparse) = row\_sparse - elemwise\_mul(default, row\_sparse) = row\_sparse - elemwise\_mul(row\_sparse, default) = row\_sparse - elemwise\_mul(csr, csr) = csr - otherwise, "elemwise\_mul" generates output with default storage

#### Value

out The result mx.symbol

```
mx.symbol.elemwise_sub
```

elemwise\_sub:Subtracts arguments element-wise.

# Description

The storage type of "elemwise\_sub" output depends on storage types of inputs

### Usage

```
mx.symbol.elemwise_sub(...)
```

#### **Arguments**

lhs	NDArray-or-Symbol first input
rhs	NDArray-or-Symbol second input

name string, optional Name of the resulting symbol.

#### **Details**

- elemwise\_sub(row\_sparse, row\_sparse) = row\_sparse - elemwise\_sub(csr, csr) = csr - elemwise\_sub(default, csr) = default - elemwise\_sub(csr, default) = default - elemwise\_sub(default, rsp) = default - elemwise\_sub(rsp, default) = default - otherwise, "elemwise\_sub" generates output with default storage

#### Value

out The result mx.symbol

# Description

This operator maps words to real-valued vectors in a high-dimensional space, called word embeddings. These embeddings can capture semantic and syntactic properties of the words. For example, it has been noted that in the learned embedding spaces, similar words tend to be close to each other and dissimilar words far apart.

#### Usage

```
mx.symbol.Embedding(...)
```

# **Arguments**

data	NDArray-or-Syn	nbol The input arra	ry to the embedding	g operator.

weight NDArray-or-Symbol The embedding weight matrix.
input.dim int, required Vocabulary size of the input indices.
output.dim int, required Dimension of the embedding vectors.

dtype 'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional,

default='float32' Data type of weight.

sparse.grad boolean, optional, default=0 Compute row sparse gradient in the backward cal-

culation. If set to True, the grad's storage type is row\_sparse.

name string, optional Name of the resulting symbol.

352 mx.symbol.erf

#### **Details**

For an input array of shape (d1, ..., dK), the shape of an output array is (d1, ..., dK, output\_dim). All the input values should be integers in the range [0, input\_dim).

If the input\_dim is ip0 and output\_dim is op0, then shape of the embedding weight matrix must be (ip0, op0).

When "sparse\_grad" is False, if any index mentioned is too large, it is replaced by the index that addresses the last vector in an embedding matrix. When "sparse\_grad" is True, an error will be raised if invalid indices are found.

### Examples::

 $input\_dim = 4 output\_dim = 5$ 

// Each row in weight matrix y represents a word. So, y = (w0,w1,w2,w3) y = [[0., 1., 2., 3., 4.], [5., 6., 7., 8., 9.], [10., 11., 12., 13., 14.], [15., 16., 17., 18., 19.]]

// Input array x represents n-grams(2-gram). So, x = [(w1, w3), (w0, w2)] x = [[1., 3.], [0., 2.]]

// Mapped input x to its vector representation y. Embedding(x, y, 4, 5) = [[[ 5., 6., 7., 8., 9.], [ 15., 16., 17., 18., 19.]],

```
[[ 0., 1., 2., 3., 4.], [ 10., 11., 12., 13., 14.]]]
```

The storage type of weight can be either row\_sparse or default.

.. Note:

If "sparse\_grad" is set to True, the storage type of gradient w.r.t weights will be "row\_sparse". Only a subset of optimizers support sparse gradients, including SGD, AdaGrad and Adam. Note that by default lazy updates is turned on, which may perform differently from standard updates. For more details, please check the Optimization API at: https://mxnet.incubator.apache.org/api/python/optimization/optimization.html Defined in src/operator/tensor/indexing\_op.cc:L603

# Value

out The result mx.symbol

mx.symbol.erf

erf:Returns element-wise gauss error function of the input.

### Description

Example::

#### **Usage**

```
mx.symbol.erf(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.erfinv 353

### **Details**

```
erf([0, -1., 10.]) = [0., -0.8427, 1.]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L887

### Value

out The result mx.symbol

mx.symbol.erfinv

erfinv:Returns element-wise inverse gauss error function of the input.

# Description

Example::

# Usage

```
mx.symbol.erfinv(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

```
erfinv([0, 0.5., -1.]) = [0., 0.4769, -inf]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L909

#### Value

mx.symbol.exp

exp:Returns element-wise exponential value of the input.

# Description

```
.. math:: exp(x) = e^x \cdot approx 2.718^x
```

### Usage

```
mx.symbol.exp(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

Example::

```
\exp([0, 1, 2]) = [1., 2.71828175, 7.38905621]
```

The storage type of "exp" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L64

#### Value

out The result mx.symbol

mx.symbol.expand\_dims  $expand\_dims:Inserts$  a new axis of size 1 into the array shape For example, given "x" with shape "(2,3,4)", then " $expand\_dims(x, axis=1)$ " will return a new array with shape "(2,1,3,4)".

### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L395

#### Usage

```
mx.symbol.expand_dims(...)
```

mx.symbol.expm1 355

### **Arguments**

data NDArray-or-Symbol Source input

axis int, required Position where new axis is to be inserted. Suppose that the in-

put 'NDArray''s dimension is 'ndim', the range of the inserted axis is '[-ndim,

ndim]'

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.expm1

expm1: Returns "exp(x) - 1" computed element-wise on the input.

# Description

This function provides greater precision than "exp(x) - 1" for small values of "x".

### Usage

```
mx.symbol.expm1(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### Details

The storage type of "expm1" output depends upon the input storage type:

- expm1(default) = default - expm1(row\_sparse) = row\_sparse - expm1(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L244

#### Value

356 mx.symbol.fix

```
mx.symbol.fill_element_0index
```

fill\_element\_0index: Fill one element of each line(row for python, column for R/Julia) in lhs according to index indicated by rhs and values indicated by mhs. This function assume rhs uses 0-based index.

#### **Description**

fill\_element\_0index:Fill one element of each line(row for python, column for R/Julia) in lhs according to index indicated by rhs and values indicated by mhs. This function assume rhs uses 0-based index.

#### Usage

```
mx.symbol.fill_element_0index(...)
```

### Arguments

1hsNDArray Left operand to the function.mhsNDArray Middle operand to the function.rhsNDArray Right operand to the function.namestring, optional Name of the resulting symbol.

### Value

out The result mx.symbol

 ${\sf mx.symbol.fix}$ 

fix:Returns element-wise rounded value to the nearest  $\setminus$  integer towards zero of the input.

### **Description**

Example::

# Usage

```
mx.symbol.fix(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.Flatten 357

#### **Details**

```
fix([-2.1, -1.9, 1.9, 2.1]) = [-2., -1., 1., 2.]
```

The storage type of "fix" output depends upon the input storage type:

- fix(default) = default - fix(row\_sparse) = row\_sparse - fix(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L875

#### Value

out The result mx.symbol

mx.symbol.Flatten

Flatten: Flattens the input array into a 2-D array by collapsing the higher dimensions. .. note:: 'Flatten' is deprecated. Use 'flatten' instead. For an input array with shape "(d1, d2, ..., dk)", 'flatten' operation reshapes the input array into an output array of shape "(d1, d2\*...\*dk)". Note that the behavior of this function is different from numpy.ndarray.flatten, which behaves similar to mxnet.ndarray.reshape((-1,1)). Example:: x = [[1,2,3], [4,5,6], [7,8,9]], [1,2,3], [4,5,6], [7,8,9]]], flatten(x = [1,2,3], [4,3,6], [7,3], [4,3,6], [4,3

### Description

Defined in src/operator/tensor/matrix\_op.cc:L250

### Usage

```
mx.symbol.Flatten(...)
```

#### **Arguments**

data NDArray-or-Symbol Input array.

name string, optional Name of the resulting symbol.

#### Value

358 mx.symbol.flip

mx.symbol.flatten

flatten:Flattens the input array into a 2-D array by collapsing the higher dimensions. .. note:: 'Flatten' is deprecated. Use 'flatten' instead. For an input array with shape "(d1, d2, ..., dk)", 'flatten' operation reshapes the input array into an output array of shape "(d1, d2\*...\*dk)". Note that the behavior of this function is different from numpy.ndarray.flatten, which behaves similar to mxnet.ndarray.reshape((-1,)). Example:: x = [[1,2,3], [4,5,6], [7,8,9]], [1,2,3], [4,5,6], [7,8,9]], flatten(x) = [1,2,3], [1,2,3], [1,2,3], [1,2,3], [1,3

#### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L250

#### Usage

```
mx.symbol.flatten(...)
```

### **Arguments**

data NDArray-or-Symbol Input array.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

 ${\tt mx.symbol.flip}$ 

flip:Reverses the order of elements along given axis while preserving array shape. Note: reverse and flip are equivalent. We use reverse in the following examples. Examples:: x = [[0., 1., 2., 3., 4.], [5., 6., 7., 8., 9.]] reverse(x, axis=0) = [[5., 6., 7., 8., 9.], [0., 1., 2., 3., 4.]] reverse(x, axis=1) = [[4., 3., 2., 1., 0.], [9., 8., 7., 6., 5.]]

### Description

Defined in src/operator/tensor/matrix\_op.cc:L832

#### Usage

```
mx.symbol.flip(...)
```

mx.symbol.floor 359

### **Arguments**

data NDArray-or-Symbol Input data array

axis Shape(tuple), required The axis which to reverse elements.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.floor

floor:Returns element-wise floor of the input.

# Description

The floor of the scalar x is the largest integer i, such that  $i \le x$ .

### Usage

```
mx.symbol.floor(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

#### Example::

```
floor([-2.1, -1.9, 1.5, 1.9, 2.1]) = [-3., -2., 1., 1., 2.]
```

The storage type of "floor" output depends upon the input storage type:

- floor(default) = default - floor(row\_sparse) = row\_sparse - floor(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L837

### Value

mx.symbol.ftml\_update ftml\_update:The FTML optimizer described in \*FTML - Follow the Moving Leader in Deep Learning\*, available at http://proceedings.mlr.press/v70/zheng17a/zheng17a.pdf.

# Description

.. math::

# Usage

```
mx.symbol.ftml_update(...)
```

# Arguments

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
d	NDArray-or-Symbol Internal state "d_t"
V	NDArray-or-Symbol Internal state "v_t"
Z	NDArray-or-Symbol Internal state "z_t"
lr	float, required Learning rate.
beta1	float, optional, default=0.600000024 Generally close to 0.5.
beta2	float, optional, default=0.999000013 Generally close to 1.
epsilon	double, optional, default=9.9999999392252903e-09 Epsilon to prevent div 0.
t	int, required Number of update.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.grad	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
name	string, optional Name of the resulting symbol.

### **Details**

```
 g_t = \addit J(W_t-1) \ v_t = \beta_2 \ v_t-1 + (1 - \beta_2) \ g_t^2 \ d_t = \frac \ 1 - \beta_1^t \ \eta_t \ \frac \ v_t \ 1 - \beta_2^t + \ensuremath{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\mbox{\s\s\mbox{\\mbox{\s\mbox{\s\mbox{\s\mbox{\s\mbox{\s\mbox{\s\s\mbox{\mbox{\mbox{\mbox{\s\mbox{\m\s\s\s\m\s\s\s\mbox{\s\s\s\s\s\s\s\s\s\s\s\s\s\s\s\s\s\
```

### Value

mx.symbol.ftrl\_update ftrl\_update:Update function for Ftrl optimizer. Referenced from \*Ad Click Prediction: a View from the Trenches\*, available at http://dl.acm.org/citation.cfm?id=2488200.

### **Description**

It updates the weights using::

### Usage

```
mx.symbol.ftrl_update(...)
```

### **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
Z	NDArray-or-Symbol z
n	NDArray-or-Symbol Square of grad
lr	float, required Learning rate
lamda1	float, optional, default=0.00999999978 The L1 regularization coefficient.
beta	float, optional, default=1 Per-Coordinate Learning Rate beta.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
name	string, optional Name of the resulting symbol.

#### **Details**

```
rescaled\_grad = clip(grad * rescale\_grad, clip\_gradient) \ z += rescaled\_grad - (sqrt(n + rescaled\_grad **2) - sqrt(n)) * weight / learning\_rate n += rescaled\_grad **2 w = (sign(z) * lamda1 - z) / ((beta + sqrt(n)) / learning\_rate + wd) * (abs(z) > lamda1)
```

If w, z and n are all of "row\_sparse" storage type, only the row slices whose indices appear in grad.indices are updated (for w, z and n)::

for row in grad.indices: rescaled\_grad[row] = clip(grad[row] \* rescale\_grad, clip\_gradient) z[row] += rescaled\_grad[row] - (sqrt(n[row] + rescaled\_grad[row]\*\*2) - sqrt(n[row])) \* weight[row] / learning\_rate n[row] += rescaled\_grad[row]\*\*2 w[row] = (sign(z[row]) \* lamda1 - z[row]) / ((beta + sqrt(n[row])) / learning\_rate + wd) \* (abs(z[row]) > lamda1)

Defined in src/operator/optimizer\_op.cc:L867

#### Value

out The result mx.symbol

```
mx.symbol.FullyConnected
```

FullyConnected:Applies a linear transformation: :math:  $Y = XW^T + b^*$ .

## **Description**

If "flatten" is set to be true, then the shapes are:

### Usage

```
mx.symbol.FullyConnected(...)
```

### **Arguments**

data NDArray-or-Symbol Input data.

weight NDArray-or-Symbol Weight matrix.

bias NDArray-or-Symbol Bias parameter.

num. hidden int, required Number of hidden nodes of the output.

no.bias boolean, optional, default=0 Whether to disable bias parameter.

flatten boolean, optional, default=1 Whether to collapse all but the first axis of the input

data tensor.

name string, optional Name of the resulting symbol.

### **Details**

```
- **data**: '(batch_size, x1, x2, ..., xn)' - **weight**: '(num_hidden, x1 * x2 * ... * xn)' - **bias**: '(num_hidden,)' - **out**: '(batch_size, num_hidden)'
```

If "flatten" is set to be false, then the shapes are:

```
- **data**: '(x1, x2, ..., xn, input_dim)' - **weight**: '(num_hidden, input_dim)' - **bias**: '(num_hidden,)' - **out**: '(x1, x2, ..., xn, num_hidden)'
```

The learnable parameters include both "weight" and "bias".

If "no\_bias" is set to be true, then the "bias" term is ignored.

.. Note:

The sparse support for FullyConnected is limited to forward evaluation with 'row\_sparse' weight and bias, where the length of 'weight.indices' and 'bias.indices' must be equal to 'num\_hidden'. This could be useful for model inference with 'row\_sparse' weights trained with importance sampling or noise contrastive estimation.

To compute linear transformation with 'csr' sparse data, sparse.dot is recommended instead of sparse.FullyConnected.

Defined in src/operator/nn/fully\_connected.cc:L287

mx.symbol.gamma 363

### Value

out The result mx.symbol

mx.symbol.gamma

gamma:Returns the gamma function (extension of the factorial function \ to the reals), computed element-wise on the input array.

## Description

The storage type of "gamma" output is always dense

### Usage

```
mx.symbol.gamma(...)
```

## Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### Value

out The result mx.symbol

mx.symbol.gammaln

gammaln: Returns element-wise log of the absolute value of the gamma function \ of the input.

### **Description**

The storage type of "gammaln" output is always dense

### Usage

```
mx.symbol.gammaln(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## Value

 $mx.symbol.gather\_nd$ 

gather\_nd:Gather elements or slices from 'data' and store to a tensor whose shape is defined by 'indices'.

## **Description**

Given 'data' with shape ' $(X_0, X_1, ..., X_{N-1})$ ' and indices with shape ' $(M, Y_0, ..., Y_{K-1})$ ', the output will have shape ' $(Y_0, ..., Y_{K-1}, X_M, ..., X_{N-1})$ ', where ' $M \le N$ '. If 'M = N', output shape will simply be ' $(Y_0, ..., Y_{K-1})$ '.

### Usage

```
mx.symbol.gather_nd(...)
```

### **Arguments**

data NDArray-or-Symbol data indices NDArray-or-Symbol indices

name string, optional Name of the resulting symbol.

### **Details**

The elements in output is defined as follows::

```
output[y_0, ..., y_K-1, x_M, ..., x_N-1] = data[indices[0, y_0, ..., y_K-1], ..., indices[M-1, y_0, ..., y_K-1], x_M, ..., x_N-1]
```

Examples::

```
data = [[0, 1], [2, 3]] indices = [[1, 1, 0], [0, 1, 0]] gather_nd(data, indices) = [2, 3, 0]
data = [[[1, 2], [3, 4]], [[5, 6], [7, 8]]] indices = [[0, 1], [1, 0]] gather_nd(data, indices) = [[3, 4], [5, 6]]
```

### Value

out The result mx.symbol

mx.symbol.GridGenerator

*GridGenerator: Generates 2D sampling grid for bilinear sampling.* 

### **Description**

GridGenerator:Generates 2D sampling grid for bilinear sampling.

mx.symbol.Group 365

## Usage

```
mx.symbol.GridGenerator(...)
```

## Arguments

data NDArray-or-Symbol Input data to the function.

transform.type 'affine', 'warp', required The type of transformation. For 'affine', input data

should be an affine matrix of size (batch, 6). For 'warp', input data should be an

optical flow of size (batch, 2, h, w).

target.shape Shape(tuple), optional, default=[0,0] Specifies the output shape (H, W). This is

required if transformation type is 'affine'. If transformation type is 'warp', this

parameter is ignored.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.Group

Create a symbol that groups symbols together.

# Description

Create a symbol that groups symbols together.

### Usage

```
mx.symbol.Group(...)
```

### **Arguments**

kwarg

Variable length of symbols or list of symbol.

### Value

The result symbol

mx.symbol.GroupNorm

GroupNorm:Group normalization.

### Description

The input channels are separated into "num\_groups" groups, each containing "num\_channels / num\_groups" channels. The mean and standard-deviation are calculated separately over the each group.

## Usage

```
mx.symbol.GroupNorm(...)
```

## **Arguments**

data NDArray-or-Symbol Input data gamma NDArray-or-Symbol gamma array beta NDArray-or-Symbol beta array

num. groups int, optional, default='1' Total number of groups.

eps float, optional, default=9.99999975e-06 An 'epsilon' parameter to prevent divi-

sion by 0.

output.mean.var

boolean, optional, default=0 Output the mean and std calculated along the given

axis.

name string, optional Name of the resulting symbol.

### **Details**

```
.. math::
```

 $\label{eq:data} $$ data = data.reshape((N, num\_groups, C // num\_groups, ...)) out = \frac{a - mean(data, axis)}{qata} + beta $$ axis) + exists for the sum of the sum o$ 

Both "gamma" and "beta" are learnable parameters.

Defined in src/operator/nn/group\_norm.cc:L77

### Value

```
mx.symbol.hard_sigmoid
```

hard\_sigmoid:Computes hard sigmoid of x element-wise.

### **Description**

```
.. math:: y = max(0, min(1, alpha * x + beta))
```

## Usage

```
mx.symbol.hard_sigmoid(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

alpha float, optional, default=0.200000003 Slope of hard sigmoid

float, optional, default=0.5 Bias of hard sigmoid.

string, optional Name of the resulting symbol.

### **Details**

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L161

### Value

out The result mx.symbol

```
mx.symbol.identity ia
```

identity:Returns a copy of the input.

## Description

From:src/operator/tensor/elemwise\_unary\_op\_basic.cc:244

## Usage

```
mx.symbol.identity(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### Value

368 mx.symbol.im2col

```
mx.symbol.IdentityAttachKLSparseReg
```

IdentityAttachKLSparseReg:Apply a sparse regularization to the output a sigmoid activation function.

### **Description**

IdentityAttachKLSparseReg:Apply a sparse regularization to the output a sigmoid activation function.

### Usage

```
mx.symbol.IdentityAttachKLSparseReg(...)
```

### **Arguments**

data NDArray-or-Symbol Input data.

sparseness.target

float, optional, default=0.100000001 The sparseness target

penalty float, optional, default=0.00100000005 The tradeoff parameter for the sparse-

ness penalty

momentum float, optional, default=0.899999976 The momentum for running average

name string, optional Name of the resulting symbol.

## Value

out The result mx.symbol

mx.symbol.im2col

im2col:Extract sliding blocks from input array.

## **Description**

This operator is used in vanilla convolution implementation to transform the sliding blocks on image to column matrix, then the convolution operation can be computed by matrix multiplication between column and convolution weight. Due to the close relation between im2col and convolution, the concept of \*\*kernel\*\*, \*\*stride\*\*, \*\*dilate\*\* and \*\*pad\*\* in this operator are inherited from convolution operation.

## Usage

```
mx.symbol.im2col(...)
```

mx.symbol.infer.shape 369

### **Arguments**

data	NDArray-or-Symbol Input array to extract sliding blocks.
kernel	Shape(tuple), required Sliding kernel size: (w,), (h, w) or (d, h, w).
stride	Shape(tuple), optional, default=[] The stride between adjacent sliding blocks in spatial dimension: $(w,)$ , $(h, w)$ or $(d, h, w)$ . Defaults to 1 for each dimension.
dilate	Shape(tuple), optional, default=[] The spacing between adjacent kernel points: $(w,)$ , $(h, w)$ or $(d, h, w)$ . Defaults to 1 for each dimension.
pad	Shape(tuple), optional, default=[] The zero-value padding size on both sides of spatial dimension: $(w_i)$ , $(h_i, w_i)$ or $(d_i, h_i, w_i)$ . Defaults to no padding.
name	string, optional Name of the resulting symbol.

#### **Details**

Given the input data of shape :math: '(N, C, \*)', where :math: 'N' is the batch size, :math: 'C' is the channel size, and :math: '\*' is the arbitrary spatial dimension, the output column array is always with shape :math: '(N, C \times \prod(\textkernel), W)', where :math: 'C \times \prod(\textkernel)' is the block size, and :math: 'W' is the block number which is the spatial size of the convolution output with same input parameters. Only 1-D, 2-D and 3-D of spatial dimension is supported in this operator.

Defined in src/operator/nn/im2col.cc:L100

### Value

out The result mx.symbol

mx.symbol.infer.shape Inference the shape of arguments, outputs, and auxiliary states.

## Description

Inference the shape of arguments, outputs, and auxiliary states.

### Usage

```
mx.symbol.infer.shape(symbol, ...)
```

# Arguments

symbol The mx.symbol object

```
mx.symbol.InstanceNorm
```

InstanceNorm:Applies instance normalization to the n-dimensional input array.

### Description

This operator takes an n-dimensional input array where (n>2) and normalizes the input using the following formula:

### Usage

```
mx.symbol.InstanceNorm(...)
```

### **Arguments**

data	NDArray-or-Symbol An n-dimensional input array $(n > 2)$ of the form [batch, channel, spatial_dim1, spatial_dim2,].
gamma	NDArray-or-Symbol A vector of length 'channel', which multiplies the normalized input.
beta	NDArray-or-Symbol A vector of length 'channel', which is added to the product of the normalized input and the weight.
eps	float, optional, default=0.00100000005 An 'epsilon' parameter to prevent division by 0.
name	string, optional Name of the resulting symbol.

## **Details**

```
.. math::
```

```
out = \fracx - mean[data] \sqrtVar[data] + \epsilon * gamma + beta
```

This layer is similar to batch normalization layer ('BatchNorm') with two differences: first, the normalization is carried out per example (instance), not over a batch. Second, the same normalization is applied both at test and train time. This operation is also known as 'contrast normalization'.

If the input data is of shape [batch, channel, spacial\_dim1, spacial\_dim2, ...], 'gamma' and 'beta' parameters must be vectors of shape [channel].

This implementation is based on this paper [1]\_

.. [1] Instance Normalization: The Missing Ingredient for Fast Stylization, D. Ulyanov, A. Vedaldi, V. Lempitsky, 2016 (arXiv:1607.08022v2).

#### Examples::

```
// Input of shape (2,1,2) x = [[[ 1.1, 2.2]], [[ 3.3, 4.4]]]
// gamma parameter of length 1 gamma = [1.5]
// beta parameter of length 1 beta = [0.5]
```

mx.symbol.khatri\_rao 371

// Instance normalization is calculated with the above formula InstanceNorm(x,gamma,beta) = [[[-0.997527, 1.99752665]], [[-0.99752653, 1.99752724]]]

Defined in src/operator/instance\_norm.cc:L95

#### Value

out The result mx.symbol

mx.symbol.khatri\_rao khatri\_rao:Computes the Khatri-Rao product of the input matrices.

### **Description**

Given a collection of :math:'n' input matrices,

## Usage

```
mx.symbol.khatri_rao(...)
```

### **Arguments**

args NDArray-or-Symbol[] Positional input matrices name string, optional Name of the resulting symbol.

### **Details**

```
.. math:: A_1 \in M_1 \in M, ..., A_n \in \mathbb{N}, the (column-wise) Khatri-Rao product is defined as the matrix,
```

.. math::  $X = A \mid \text{otimes } \cdot A \mid \text{n } \mid \text{mathbbR}^{(M)} \mid \text{cdots } M \mid \text{n} \mid \text{times } N$ ,

where the :math:'k' th column is equal to the column-wise outer product :math:' $A_1_k$  \otimes \cdots \otimes  $A_n_k$ ' where :math:' $A_i_k$ ' is the kth column of the ith matrix.

### Example::

```
»> A = mx.nd.array([[1, -1], »> [2, -3]]) »> B = mx.nd.array([[1, 4], »> [2, 5], »> [3, 6]]) »> C = mx.nd.khatri_rao(A, B) »> print(C.asnumpy()) [[ 1. -4.] [ 2. -5.] [ 3. -6.] [ 2. -12.] [ 4. -15.] [ 6. -18.]]
```

Defined in src/operator/contrib/krprod.cc:L108

### Value

```
mx.symbol.L2Normalization
```

L2Normalization:Normalize the input array using the L2 norm.

#### Description

For 1-D NDArray, it computes::

### Usage

```
mx.symbol.L2Normalization(...)
```

## **Arguments**

data NDArray-or-Symbol Input array to normalize.

eps float, optional, default=1.00000001e-10 A small constant for numerical stability.

"channel", 'instance', 'spatial', optional, default='instance' Specify the dimen-

sion along which to compute L2 norm.

name string, optional Name of the resulting symbol.

#### **Details**

```
out = data / sqrt(sum(data ** 2) + eps)
```

For N-D NDArray, if the input array has shape (N, N, ..., N),

with "mode" = "instance", it normalizes each instance in the multidimensional array by its L2 norm.::

```
for i in 0...N out[i,:,:,...,:] = data[i,:,:,...,:] / sqrt(sum(data[i,:,:,...,:] ** 2) + eps)
```

with "mode" = "channel", it normalizes each channel in the array by its L2 norm.::

```
for i in 0...N out[:,i,:,...,:] = data[:,i,:,...,:] / sqrt(sum(data[:,i,:,...,:] ** 2) + eps)
```

with "mode" = "spatial", it normalizes the cross channel norm for each position in the array by its L2 norm.::

for dim in 2...N for i in 0...N out[....,i,...] = take(out, indices=i, axis=dim) / sqrt(sum(take(out, indices=i, axis=dim) \*\* 2) + eps) -dim-

## Example::

```
x = [[[1,2], [3,4]], [[2,2], [5,6]]]
```

L2Normalization(x, mode='instance') =[[[ 0.18257418 0.36514837] [ 0.54772252 0.73029673]] [[ 0.24077171 0.24077171] [ 0.60192931 0.72231513]]]

L2Normalization(x, mode='channel') =[[[ 0.31622776 0.44721359] [ 0.94868326 0.89442718]] [[ 0.37139067 0.31622776] [ 0.92847669 0.94868326]]]

L2Normalization(x, mode='spatial') =[[[ 0.44721359 0.89442718] [ 0.60000002 0.80000001]] [[ 0.70710677 0.70710677] [ 0.6401844 0.76822126]]]

Defined in src/operator/l2\_normalization.cc:L196

## Value

out The result mx.symbol

```
mx.symbol.lamb_update_phase1
```

 $lamb\_update\_phase1:Phase\ I\ of\ lamb\ update\ it\ performs\ the\ following\ operations\ and\ returns\ g:.$ 

# Description

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

NDArray-or-Symbol Weight

## Usage

```
mx.symbol.lamb_update_phase1(...)
```

# Arguments

weight

J	, , ,
grad	NDArray-or-Symbol Gradient
mean	NDArray-or-Symbol Moving mean
var	NDArray-or-Symbol Moving variance
beta1	float, optional, default=0.899999976 The decay rate for the 1st moment estimates.
beta2	float, optional, default=0.999000013 The decay rate for the 2nd moment estimates.
epsilon	float, optional, default=9.9999997e-07 A small constant for numerical stability.
t	int, required Index update count.
bias.correction	1
	boolean, optional, default=1 Whether to use bias correction.
wd	float, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
name	string, optional Name of the resulting symbol.

```
.. math:: \begingather* grad = grad * rescale_grad if (grad < -clip_gradient) then grad = -clip_gradient if (grad > clip_gradient) then grad = clip_gradient

mean = beta1 * mean + (1 - beta1) * grad; variance = beta2 * variance + (1 - beta2) * grad ^ 2;

if (bias_correction) then mean_hat = mean_hat_specific = beta1 \text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{
```

if (bias\_correction) then mean\_hat = mean / (1. - beta1^t); var\_hat = var / (1 - beta2^t); g = mean\_hat / (var\_hat^(1/2) + epsilon) + wd \* weight; else g = mean / (var\_data^(1/2) + epsilon) + wd \* weight; \endgather\*

Defined in src/operator/optimizer\_op.cc:L944

### Value

out The result mx.symbol

```
mx.symbol.lamb_update_phase2
```

lamb\_update\_phase2:Phase II of lamb update it performs the following operations and updates grad.

### **Description**

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

#### Usage

```
mx.symbol.lamb_update_phase2(...)
```

#### **Arguments**

weight	NDArray-or-Symbol Weight
g	NDArray-or-Symbol Output of lamb_update_phase 1
r1	NDArray-or-Symbol r1
r2	NDArray-or-Symbol r2
lr	float, required Learning rate
lower.bound	float, optional, default=-1 Lower limit of norm of weight. If lower_bound <= 0, Lower limit is not set
upper.bound	float, optional, default=-1 Upper limit of norm of weight. If upper_bound <= 0, Upper limit is not set
name	string, optional Name of the resulting symbol.

### **Details**

```
.. math:: \begingather* if (lower_bound >= 0) then r1 = max(r1, lower_bound) if (upper_bound >= 0) then r1 = max(r1, upper_bound) if (r1 == 0 \text{ or } r2 == 0) then lr = lr \text{ else } lr = lr * (r1/r2) \text{ weight} = \text{weight} - lr * g \endgather* Defined in src/operator/optimizer_op.cc:L983
```

mx.symbol.LayerNorm 375

### Value

out The result mx.symbol

mx.symbol.LayerNorm LayerNorm:Layer normalization.

## Description

Normalizes the channels of the input tensor by mean and variance, and applies a scale "gamma" as well as offset "beta".

### Usage

```
mx.symbol.LayerNorm(...)
```

### **Arguments**

data	NDArray-or-Symbol Input data to layer normalization	
gamma	NDArray-or-Symbol gamma array	
beta	NDArray-or-Symbol beta array	
axis	int, optional, default='-1' The axis to perform layer normalization. Usually, this should be be axis of the channel dimension. Negative values means indexing from right to left.	
eps	float, optional, default=9.99999975e-06 An 'epsilon' parameter to prevent division by 0.	
output.mean.var		
	boolean, optional, default=0 Output the mean and std calculated along the given axis.	
name	string, optional Name of the resulting symbol.	

#### **Details**

Assume the input has more than one dimension and we normalize along axis 1. We first compute the mean and variance along this axis and then compute the normalized output, which has the same shape as input, as following:

```
.. math::
```

out = \fracdata - mean(data, axis)\sqrtvar(data, axis) + \epsilon \* gamma + beta

Both "gamma" and "beta" are learnable parameters.

Unlike BatchNorm and InstanceNorm, the \*mean\* and \*var\* are computed along the channel dimension.

Assume the input has size  $k^*$  on axis 1, then both "gamma" and "beta" have shape k,k. If "output\_mean\_var" is set to be true, then outputs both "data\_mean" and "data\_std". Note that no gradient will be passed through these two outputs.

The parameter "axis" specifies which axis of the input shape denotes the 'channel' (separately normalized groups). The default is -1, which sets the channel axis to be the last item in the input shape.

Defined in src/operator/nn/layer\_norm.cc:L158

### Value

out The result mx.symbol

mx.symbol.LeakyReLU LeakyReLU:Applies Leaky rectified linear unit activation elementwise to the input.

### **Description**

Leaky ReLUs attempt to fix the "dying ReLU" problem by allowing a small 'slope' when the input is negative and has a slope of one when input is positive.

### Usage

```
mx.symbol.LeakyReLU(...)
```

### **Arguments**

data	NDArray-or-Symbol Input data to activation function.
gamma	NDArray-or-Symbol Input data to activation function.
act.type	'elu', 'gelu', 'leaky', 'prelu', 'rrelu', 'selu',optional, default='leaky' Activation function to be applied.
slope	float, optional, default=0.25 Init slope for the activation. (For leaky and elu only)
lower.bound	float, optional, default=0.125 Lower bound of random slope. (For rrelu only)
upper.bound	float, optional, default= $0.333999991$ Upper bound of random slope. (For rrelu only)
name	string, optional Name of the resulting symbol.

### **Details**

The following modified ReLU Activation functions are supported:

- \*elu\*: Exponential Linear Unit. 'y = x > 0 ? x : slope \* (exp(x)-1)' - \*selu\*: Scaled Exponential Linear Unit. 'y = lambda \* (x > 0 ? x : alpha \* (exp(x) - 1))' where \*lambda = 1.0507009873554804934193349852946\* and \*alpha = 1.6732632423543772848170429916717\*. - \*leaky\*: Leaky ReLU. 'y = x > 0 ? x : slope \* x' - \*prelu\*: Parametric ReLU. This is same as \*leaky\* except that 'slope' is learnt during training. - \*rrelu\*: Randomized ReLU. same as \*leaky\* but the 'slope' is uniformly and randomly chosen from \*[lower\_bound, upper\_bound)\* for training, while fixed to be \*(lower\_bound+upper\_bound)/2\* for inference.

Defined in src/operator/leaky\_relu.cc:L161

mx.symbol.linalg\_det

### Value

out The result mx.symbol

```
mx.symbol.linalg_det:Compute the determinant of a matrix. Input is a tensor *A* of dimension *n >= 2*.
```

377

## Description

```
If *n=2*, *A* is a square matrix. We compute:
```

### Usage

```
mx.symbol.linalg_det(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of square matrix

name string, optional Name of the resulting symbol.

## **Details**

```
*out* = *det(A)*
```

If \*n>2\*, \*det\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only. .. note:: There is no gradient backwarded when A is non-invertible (which is equivalent to det(A) = 0) because zero is rarely hit upon in float point computation and the Jacobi's formula on determinant gradient is not computationally efficient when A is non-invertible.

## Examples::

```
Single matrix determinant A = [[1., 4.], [2., 3.]] det(A) = [-5.]
```

Batch matrix determinant A = [[[1., 4.], [2., 3.]], [[2., 3.], [1., 4.]]] det(A) = [-5., 5.]

Defined in src/operator/tensor/la\_op.cc:L975

### Value

```
mx.symbol.linalg_extractdiag
```

linalg\_extractdiag:Extracts the diagonal entries of a square matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

### **Description**

If \*n=2\*, then \*A\* represents a single square matrix which diagonal elements get extracted as a 1-dimensional tensor.

### Usage

```
mx.symbol.linalg_extractdiag(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of square matrices

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

name string, optional Name of the resulting symbol.

### **Details**

If \*n>2\*, then \*A\* represents a batch of square matrices on the trailing two dimensions. The extracted diagonals are returned as an \*n-1\*-dimensional tensor.

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix diagonal extraction A = [[1.0, 2.0], [3.0, 4.0]]

extractdiag(A) = [1.0, 4.0]

extractdiag(A, 1) = [2.0]

Batch matrix diagonal extraction A = [[[1.0, 2.0], [3.0, 4.0]], [[5.0, 6.0], [7.0, 8.0]]]

extractdiag(A) = [[1.0, 4.0], [5.0, 8.0]]

Defined in src/operator/tensor/la\_op.cc:L495

### Value

```
mx.symbol.linalg_extracttrian
```

linalg\_extracttrian:Extracts a triangular sub-matrix from a square matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

### **Description**

If \*n=2\*, then \*A\* represents a single square matrix from which a triangular sub-matrix is extracted as a 1-dimensional tensor.

#### Usage

```
mx.symbol.linalg_extracttrian(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of square matrices

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

lower boolean, optional, default=1 Refer to the lower triangular matrix if lower=true,

refer to the upper otherwise. Only relevant when offset=0

name string, optional Name of the resulting symbol.

#### **Details**

If \*n>2\*, then \*A\* represents a batch of square matrices on the trailing two dimensions. The extracted triangular sub-matrices are returned as an \*n-1\*-dimensional tensor.

The \*offset\* and \*lower\* parameters determine the triangle to be extracted:

- When \*offset = 0\* either the lower or upper triangle with respect to the main diagonal is extracted depending on the value of parameter \*lower\*. When \*offset = k > 0\* the upper triangle with respect to the k-th diagonal above the main diagonal is extracted. When \*offset = k < 0\* the lower triangle with respect to the k-th diagonal below the main diagonal is extracted.
- .. note:: The operator supports float32 and float64 data types only.

#### Examples::

```
Single triagonal extraction A = [[1.0, 2.0], [3.0, 4.0]]
```

extracttrian(A) = [1.0, 3.0, 4.0] extracttrian(A, lower=False) = [1.0, 2.0, 4.0] extracttrian(A, 1) = [2.0] extracttrian(A, -1) = [3.0]

Batch triagonal extraction A = [[[1.0, 2.0], [3.0, 4.0]], [[5.0, 6.0], [7.0, 8.0]]]

extracttrian(A) = [[1.0, 3.0, 4.0], [5.0, 7.0, 8.0]]

Defined in src/operator/tensor/la\_op.cc:L605

#### Value

mx.symbol.linalg\_gelqf

linalg\_gelqf:LQ factorization for general matrix. Input is a tensor \*A\* of dimension  $*n \ge 2*$ .

### Description

If \*n=2\*, we compute the LQ factorization (LAPACK \*gelqf\*, followed by \*orglq\*). \*A\* must have shape \*(x, y)\* with \*x <= y\*, and must have full rank \*=x\*. The LQ factorization consists of \*L\* with shape \*(x, x)\* and \*Q\* with shape \*(x, y)\*, so that:

### Usage

```
mx.symbol.linalg_gelqf(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of input matrices to be factorized

name string, optional Name of the resulting symbol.

### **Details**

```
*A* = *L* \ *O*
```

Here, \*L\* is lower triangular (upper triangle equal to zero) with nonzero diagonal, and \*Q\* is row-orthonormal, meaning that

is equal to the identity matrix of shape \*(x, x)\*.

If \*n>2\*, \*gelqf\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

### Examples::

Single LQ factorization A = [[1., 2., 3.], [4., 5., 6.]] Q, L = gelqf(A) Q = [[-0.26726124, -0.53452248, -0.80178373], [0.87287156, 0.21821789, -0.43643578]] L = [[-3.74165739, 0.], [-8.55235974, 1.96396101]]

Batch LQ factorization A = [[[1, 2., 3.], [4., 5., 6.]], [[7., 8., 9.], [10., 11., 12.]]] Q, L = gelqf(A) Q = [[[-0.26726124, -0.53452248, -0.80178373], [0.87287156, 0.21821789, -0.43643578]], [[-0.50257071, -0.57436653, -0.64616234], [0.7620735, 0.05862104, -0.64483142]]] L = [[[-3.74165739, 0.], [-8.55235974, 1.96396101]], [[-13.92838828, 0.], [-19.09768702, 0.52758934]]]

Defined in src/operator/tensor/la\_op.cc:L798

### Value

mx.symbol.linalg\_gemm linalg\_gemm:Performs general matrix multiplication and accumulation. Input are tensors \*A\*, \*B\*, \*C\*, each of dimension \*n >= 2\* and having the same shape on the leading \*n-2\* dimensions.

### **Description**

If \*n=2\*, the BLAS3 function \*gemm\* is performed:

#### **Usage**

```
mx.symbol.linalg_gemm(...)
```

### **Arguments**

A	NDArray-or-Symbol Tensor of input matrices
В	NDArray-or-Symbol Tensor of input matrices
С	NDArray-or-Symbol Tensor of input matrices
transpose.a	boolean, optional, default=0 Multiply with transposed of first input (A).
transpose.b	boolean, optional, default=0 Multiply with transposed of second input (B).
alpha	double, optional, default=1 Scalar factor multiplied with A*B.
beta	double, optional, default=1 Scalar factor multiplied with C.
axis	int, optional, default='-2' Axis corresponding to the matrix rows.
name	string, optional Name of the resulting symbol.

#### **Details**

```
*out* = *alpha* \* *op* \ (*A*) \* *op* \ (*B*) + *beta* \* *C*
```

Here, \*alpha\* and \*beta\* are scalar parameters, and \*op()\* is either the identity or matrix transposition (depending on \*transpose\_a\*, \*transpose\_b\*).

If \*n>2\*, \*gemm\* is performed separately for a batch of matrices. The column indices of the matrices are given by the last dimensions of the tensors, the row indices by the axis specified with the \*axis\* parameter. By default, the trailing two dimensions will be used for matrix encoding.

For a non-default axis parameter, the operation performed is equivalent to a series of swapaxes/gemm/swapaxes calls. For example let \*A\*, \*B\*, \*C\* be 5 dimensional tensors. Then gemm(\*A\*, \*B\*, \*C\*, axis=1) is equivalent to the following without the overhead of the additional swapaxis operations::

```
A1 = \text{swapaxes}(A, \text{dim}1=1, \text{dim}2=3) B1 = \text{swapaxes}(B, \text{dim}1=1, \text{dim}2=3) C = \text{swapaxes}(C, \text{dim}1=1, \text{d
```

When the input data is of type float32 and the environment variables MXNET\_CUDA\_ALLOW\_TENSOR\_CORE and MXNET\_CUDA\_TENSOR\_OP\_MATH\_ALLOW\_CONVERSION are set to 1, this operator will try to use pseudo-float16 precision (float32 math with float16 I/O) precision in order to use Tensor Cores on suitable NVIDIA GPUs. This can sometimes give significant speedups.

.. note:: The operator supports float32 and float64 data types only.

#### Examples::

Single matrix multiply-add  $A = [[1.0, 1.0], [1.0, 1.0]] B = [[1.0, 1.0], [1.0, 1.0], [1.0, 1.0]] C = [[1.0, 1.0, 1.0], [1.0, 1.0], [1.0, 1.0]] gemm(A, B, C, transpose_b=True, alpha=2.0, beta=10.0) = [[14.0, 14.0, 14.0], [14.0, 14.0, 14.0]]$ 

Batch matrix multiply-add  $A = [[[1.0, 1.0]], [[0.1, 0.1]]] B = [[[1.0, 1.0]], [[0.1, 0.1]]] C = [[[10.0]], [[0.01]]] gemm(A, B, C, transpose_b=True, alpha=2.0, beta=10.0) = [[[104.0]], [[0.14]]]$ 

Defined in src/operator/tensor/la\_op.cc:L89

#### Value

out The result mx.symbol

```
mx.symbol.linalg_gemm2
```

linalg\_gemm2:Performs general matrix multiplication. Input are tensors \*A\*, \*B\*, each of dimension \*n >= 2\* and having the same shape on the leading \*n-2\* dimensions.

### **Description**

If \*n=2\*, the BLAS3 function \*gemm\* is performed:

### Usage

```
mx.symbol.linalg_gemm2(...)
```

## **Arguments**

A NDArray-or-Symbol Tensor of input matrices

B NDArray-or-Symbol Tensor of input matrices

transpose.a boolean, optional, default=0 Multiply with transposed of first input (A). transpose.b boolean, optional, default=0 Multiply with transposed of second input (B).

alpha double, optional, default=1 Scalar factor multiplied with A\*B.

axis int, optional, default='-2' Axis corresponding to the matrix row indices.

name string, optional Name of the resulting symbol.

#### **Details**

```
*out* = *alpha* \ **op* \ (*A*) \ **op* \ (*B*)
```

Here \*alpha\* is a scalar parameter and \*op()\* is either the identity or the matrix transposition (depending on \*transpose\_a\*, \*transpose\_b\*).

If \*n>2\*, \*gemm\* is performed separately for a batch of matrices. The column indices of the matrices are given by the last dimensions of the tensors, the row indices by the axis specified with the \*axis\* parameter. By default, the trailing two dimensions will be used for matrix encoding.

For a non-default axis parameter, the operation performed is equivalent to a series of swapaxes/gemm/swapaxes calls. For example let \*A\*, \*B\* be 5 dimensional tensors. Then gemm(\*A\*, \*B\*, axis=1) is equivalent to the following without the overhead of the additional swapaxis operations::

```
A1 = swapaxes(A, dim1=1, dim2=3) B1 = swapaxes(B, dim1=1, dim2=3) C = gemm2(A1, B1) C = swapaxis(C, dim1=1, dim2=3)
```

When the input data is of type float32 and the environment variables MXNET\_CUDA\_ALLOW\_TENSOR\_CORE and MXNET\_CUDA\_TENSOR\_OP\_MATH\_ALLOW\_CONVERSION are set to 1, this operator will try to use pseudo-float16 precision (float32 math with float16 I/O) precision in order to use Tensor Cores on suitable NVIDIA GPUs. This can sometimes give significant speedups.

.. note:: The operator supports float32 and float64 data types only.

#### Examples::

```
Single matrix multiply A = [[1.0, 1.0], [1.0, 1.0]] B = [[1.0, 1.0], [1.0, 1.0], [1.0, 1.0]] gemm2(A, B, transpose_b=True, alpha=2.0) = [[4.0, 4.0, 4.0], [4.0, 4.0, 4.0]]
```

Batch matrix multiply  $A = [[[1.0, 1.0]], [[0.1, 0.1]]] B = [[[1.0, 1.0]], [[0.1, 0.1]]] gemm2(A, B, transpose_b=True, alpha=<math>2.0$ ) = [[[4.0]], [[0.04]]]

Defined in src/operator/tensor/la\_op.cc:L163

#### Value

out The result mx.symbol

```
mx.symbol.linalg_inverse
```

linalg\_inverse: Compute the inverse of a matrix. Input is a tensor  $A^*$  of dimension  $n \ge 2$ .

### Description

```
If *n=2*, *A* is a square matrix. We compute:
```

### Usage

```
mx.symbol.linalg_inverse(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of square matrix

name string, optional Name of the resulting symbol.

```
*out* = *A* \ :sup: `-1`
```

If \*n>2\*, \*inverse\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

Examples::

```
Single matrix inverse A = [[1., 4.], [2., 3.]] inverse(A) = [[-0.6, 0.8], [0.4, -0.2]]
```

Batch matrix inverse A = [[[1., 4.], [2., 3.]], [[1., 3.], [2., 4.]]] inverse(A) = [[[-0.6, 0.8], [0.4, -0.2]], [[-2., 1.5], [1., -0.5]]]

Defined in src/operator/tensor/la\_op.cc:L920

#### Value

out The result mx.symbol

```
mx.symbol.linalg_makediag
```

linalg\_makediag:Constructs a square matrix with the input as diagonal. Input is a tensor \*A\* of dimension \*n >= 1\*.

### **Description**

If \*n=1\*, then \*A\* represents the diagonal entries of a single square matrix. This matrix will be returned as a 2-dimensional tensor. If \*n>1\*, then \*A\* represents a batch of diagonals of square matrices. The batch of diagonal matrices will be returned as an \*n+1\*-dimensional tensor.

### Usage

```
mx.symbol.linalg_makediag(...)
```

### **Arguments**

Α	NDArray-or-	Symbol Tensor	of diagonal	entries

offset int, optional, default='0' Offset of the diagonal versus the main diagonal. 0

corresponds to the main diagonal, a negative/positive value to diagonals be-

low/above the main diagonal.

name string, optional Name of the resulting symbol.

```
.. note:: The operator supports float32 and float64 data types only. Examples:: Single diagonal matrix construction A = [1.0, 2.0] makediag(A) = [[1.0, 0.0], [0.0, 2.0]] makediag(A, 1) = [[0.0, 1.0, 0.0], [0.0, 0.0, 2.0], [0.0, 0.0, 0.0]] Batch diagonal matrix construction A = [[1.0, 2.0], [3.0, 4.0]] makediag(A) = [[[1.0, 0.0], [0.0, 2.0]], [[3.0, 0.0], [0.0, 4.0]]] Defined in src/operator/tensor/la_op.cc:L547
```

#### Value

out The result mx.symbol

```
mx.symbol.linalg_maketrian
```

linalg\_maketrian:Constructs a square matrix with the input representing a specific triangular sub-matrix. This is basically the inverse of \*linalg.extracttrian\*. Input is a tensor \*A\* of dimension \*n >= 1\*.

# Description

If \*n=1\*, then \*A\* represents the entries of a triangular matrix which is lower triangular if \*off-set<0\* or \*offset=0\*, \*lower=true\*. The resulting matrix is derived by first constructing the square matrix with the entries outside the triangle set to zero and then adding \*offset\*-times an additional diagonal with zero entries to the square matrix.

### Usage

```
mx.symbol.linalg_maketrian(...)
```

## **Arguments**

Α	NDArray-or-Symbol Tensor of triangular matrices stored as vectors
offset	int, optional, default='0' Offset of the diagonal versus the main diagonal. 0 corresponds to the main diagonal, a negative/positive value to diagonals below/above the main diagonal.
lower	boolean, optional, default=1 Refer to the lower triangular matrix if lower=true, refer to the upper otherwise. Only relevant when offset=0
name	string, optional Name of the resulting symbol.

If \*n>1\*, then \*A\* represents a batch of triangular sub-matrices. The batch of corresponding square matrices is returned as an \*n+1\*-dimensional tensor.

.. note:: The operator supports float32 and float64 data types only.

#### Examples::

```
Single matrix construction A = [1.0, 2.0, 3.0]
```

```
maketrian(A) = [[1.0, 0.0], [2.0, 3.0]]
```

maketrian(A, lower=false) = [[1.0, 2.0], [0.0, 3.0]]

maketrian(A, offset=1) = [[0.0, 1.0, 2.0], [0.0, 0.0, 3.0], [0.0, 0.0, 0.0]] maketrian(A, offset=-1) = [[0.0, 0.0, 0.0], [1.0, 0.0, 0.0], [2.0, 3.0, 0.0]]

Batch matrix construction A = [[1.0, 2.0, 3.0], [4.0, 5.0, 6.0]]

```
maketrian(A) = [[[1.0, 0.0], [2.0, 3.0]], [[4.0, 0.0], [5.0, 6.0]]]
```

maketrian(A, offset=1) = [[[0.0, 1.0, 2.0], [0.0, 0.0, 3.0], [0.0, 0.0, 0.0]], [[0.0, 4.0, 5.0], [0.0, 0.0, 6.0], [0.0, 0.0, 0.0]]]

Defined in src/operator/tensor/la\_op.cc:L673

### Value

out The result mx.symbol

```
mx.symbol.linalg_potrf
```

linalg\_potrf: Performs Cholesky factorization of a symmetric positivedefinite matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

### Description

If \*n=2\*, the Cholesky factor \*B\* of the symmetric, positive definite matrix \*A\* is computed. \*B\* is triangular (entries of upper or lower triangle are all zero), has positive diagonal entries, and:

### Usage

```
mx.symbol.linalg_potrf(...)
```

## **Arguments**

A NDArray-or-Symbol Tensor of input matrices to be decomposed

name string, optional Name of the resulting symbol.

```
*A* = *B* \* *B*\ :sup:'T' if *lower* = *true* *A* = *B*\ :sup:'T' \* *B* if *lower* = *false*
```

If \*n>2\*, \*potrf\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

#### Examples::

Single matrix factorization A = [[4.0, 1.0], [1.0, 4.25]] potrf(A) = [[2.0, 0], [0.5, 2.0]]

Batch matrix factorization A = [[[4.0, 1.0], [1.0, 4.25]], [[16.0, 4.0], [4.0, 17.0]]] potrf(A) = [[[2.0, 0], [0.5, 2.0]], [[4.0, 0], [1.0, 4.0]]]

Defined in src/operator/tensor/la\_op.cc:L214

#### Value

out The result mx.symbol

```
mx.symbol.linalg_potri
```

linalg\_potri:Performs matrix inversion from a Cholesky factorization. Input is a tensor \*A\* of dimension \*n >= 2\*.

### **Description**

If \*n=2\*, \*A\* is a triangular matrix (entries of upper or lower triangle are all zero) with positive diagonal. We compute:

#### Usage

```
mx.symbol.linalg_potri(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of lower triangular matrices

name string, optional Name of the resulting symbol.

#### Details

```
*out* = *A*\ :sup: '-T' \* *A*\ :sup: '-1' if *lower* = *true* *out* = *A*\ :sup: '-1' \* *A*\ :sup: '-T' if *lower* = *false*
```

In other words, if \*A\* is the Cholesky factor of a symmetric positive definite matrix \*B\* (obtained by \*potrf\*), then

```
*out* = *B*\ :sup:'-1'
```

If \*n>2\*, \*potri\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

.. note:: Use this operator only if you are certain you need the inverse of \*B\*, and cannot use the Cholesky factor \*A\* (\*potrf\*), together with backsubstitution (\*trsm\*). The latter is numerically much safer, and also cheaper.

#### Examples::

Single matrix inverse A = [[2.0, 0], [0.5, 2.0]] potri(A) = [[0.26563, -0.0625], [-0.0625, 0.25]]Batch matrix inverse A = [[[2.0, 0], [0.5, 2.0]], [[4.0, 0], [1.0, 4.0]]] potri(A) = [[[0.26563, -0.0625], [-0.0625, 0.25]], [[0.06641, -0.01562], [-0.01562, 0.0625]]]

Defined in src/operator/tensor/la\_op.cc:L275

#### Value

out The result mx.symbol

```
mx.symbol.linalg_slogdet
```

linalg\_slogdet:Compute the sign and log of the determinant of a matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

### Description

If \*n=2\*, \*A\* is a square matrix. We compute:

### Usage

```
mx.symbol.linalg_slogdet(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of square matrix name string, optional Name of the resulting symbol.

#### **Details**

```
*sign* = *sign(det(A))* *logabsdet* = *log(abs(det(A)))*
```

If \*n>2\*, \*slogdet\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only. .. note:: The gradient is not properly defined on sign, so the gradient of it is not backwarded. .. note:: No gradient is backwarded when A is non-invertible. Please see the docs of operator det for detail.

#### Examples::

Single matrix signed log determinant A = [[2., 3.], [1., 4.]] sign, logabsdet = slogdet(A) sign = [1.] logabsdet = [1.609438]

Batch matrix signed log determinant A = [[[2., 3.], [1., 4.]], [[1., 2.], [2., 4.]], [[1., 2.], [4., 3.]]] sign, logabsdet = slogdet(A) sign = [1., 0., -1.] logabsdet = [1.609438, -inf, 1.609438]

Defined in src/operator/tensor/la\_op.cc:L1034

### Value

out The result mx.symbol

```
mx.symbol.linalg_sumlogdiag
```

linalg\_sumlogdiag:Computes the sum of the logarithms of the diagonal elements of a square matrix. Input is a tensor \*A\* of dimension \*n >= 2\*.

# Description

If \*n=2\*, \*A\* must be square with positive diagonal entries. We sum the natural logarithms of the diagonal elements, the result has shape (1,).

### Usage

```
mx.symbol.linalg_sumlogdiag(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of square matrices

name string, optional Name of the resulting symbol.

### **Details**

If \*n>2\*, \*sumlogdiag\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

Examples::

Single matrix reduction A = [[1.0, 1.0], [1.0, 7.0]] sumlogdiag(A) = [1.9459]

Batch matrix reduction A = [[[1.0, 1.0], [1.0, 7.0]], [[3.0, 0], [0, 17.0]]] sumlogdiag(A) = [1.9459, 3.9318]

Defined in src/operator/tensor/la\_op.cc:L445

## Value

mx.symbol.linalg\_syrk linalg\_syrk:Multiplication of matrix with its transpose. Input is a tensor \*A\* of dimension \*n >= 2\*.

### **Description**

If \*n=2\*, the operator performs the BLAS3 function \*syrk\*:

### Usage

```
mx.symbol.linalg_syrk(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of input matrices

transpose boolean, optional, default=0 Use transpose of input matrix.

alpha double, optional, default=1 Scalar factor to be applied to the result.

name string, optional Name of the resulting symbol.

#### **Details**

```
*out* = *alpha* \* *A* \* *A*\ :sup: 'T' if *transpose=False*, or *out* = *alpha* \* *A*\ :sup: 'T' \ \* *A* if *transpose=True*.
```

If \*n>2\*, \*syrk\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

### Examples::

```
Single matrix multiply A = [[1., 2., 3.], [4., 5., 6.]] syrk(A, alpha=1., transpose=False) = [[14., 32.], [32., 77.]] syrk(A, alpha=1., transpose=True) = [[17., 22., 27.], [22., 29., 36.], [27., 36., 45.]]
```

Batch matrix multiply A = [[[1., 1.]], [[0.1, 0.1]]] syrk(A, alpha=2., transpose=False) = [[[4.]], [[0.04]]]

Defined in src/operator/tensor/la\_op.cc:L730

### Value

mx.symbol.linalg\_trmm linalg\_trmm:Performs multiplication with a lower triangular matrix. Input are tensors \*A\*, \*B\*, each of dimension \*n >= 2\* and having the same shape on the leading \*n-2\* dimensions.

### Description

If \*n=2\*, \*A\* must be triangular. The operator performs the BLAS3 function \*trmm\*:

### Usage

```
mx.symbol.linalg_trmm(...)
```

#### **Arguments**

A NDArray-or-Symbol Tensor of lower triangular matrices

B NDArray-or-Symbol Tensor of matrices

transpose boolean, optional, default=0 Use transposed of the triangular matrix

rightside boolean, optional, default=0 Multiply triangular matrix from the right to non-

triangular one.

lower boolean, optional, default=1 True if the triangular matrix is lower triangular,

false if it is upper triangular.

alpha double, optional, default=1 Scalar factor to be applied to the result.

name string, optional Name of the resulting symbol.

#### **Details**

```
*out* = *alpha* \* *op*\ (*A*) \* *B*
if *rightside=False*, or
*out* = *alpha* \* *B* \* *op*\ (*A*)
```

if \*rightside=True\*. Here, \*alpha\* is a scalar parameter, and \*op()\* is either the identity or the matrix transposition (depending on \*transpose\*).

If \*n>2\*, \*trmm\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

### Examples::

Single triangular matrix multiply A = [[1.0, 0], [1.0, 1.0]] B = [[1.0, 1.0, 1.0], [1.0, 1.0]] trmm(A, B, alpha=2.0) = [[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]]

Batch triangular matrix multiply A = [[[1.0, 0], [1.0, 1.0]], [[1.0, 0], [1.0, 1.0]]] B = [[[1.0, 1.0, 1.0], [1.0, 1.0]], [[0.5, 0.5, 0.5], [0.5, 0.5, 0.5]]] trmm(A, B, alpha=2.0) = [[[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]], [[1.0, 1.0, 1.0], [2.0, 2.0, 2.0]]]

Defined in src/operator/tensor/la\_op.cc:L333

#### Value

out The result mx.symbol

mx.symbol.linalg\_trsm linalg\_trsm:Solves matrix equation involving a lower triangular matrix. Input are tensors \*A\*, \*B\*, each of dimension \*n >= 2\* and having the same shape on the leading \*n-2\* dimensions.

### Description

If \*n=2\*, \*A\* must be triangular. The operator performs the BLAS3 function \*trsm\*, solving for \*out\* in:

#### Usage

```
mx.symbol.linalg_trsm(...)
```

### **Arguments**

A NDArray-or-Symbol Tensor of lower triangular matrices

B NDArray-or-Symbol Tensor of matrices

transpose boolean, optional, default=0 Use transposed of the triangular matrix

rightside boolean, optional, default=0 Multiply triangular matrix from the right to non-

triangular one.

lower boolean, optional, default=1 True if the triangular matrix is lower triangular,

false if it is upper triangular.

alpha double, optional, default=1 Scalar factor to be applied to the result.

name string, optional Name of the resulting symbol.

#### **Details**

```
*op*\(*A*)\* *out* = *alpha* \* *B*
if *rightside=False*, or
```

 $*out* \ **op* \ (*A*) = *alpha* \ **B*$ 

if \*rightside=True\*. Here, \*alpha\* is a scalar parameter, and \*op()\* is either the identity or the matrix transposition (depending on \*transpose\*).

If \*n>2\*, \*trsm\* is performed separately on the trailing two dimensions for all inputs (batch mode).

.. note:: The operator supports float32 and float64 data types only.

### Examples::

Single matrix solve A = [[1.0, 0], [1.0, 1.0]] B = [[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]] trsm(A, B, alpha=0.5) = [[1.0, 1.0, 1.0], [1.0, 1.0, 1.0]]

Batch matrix solve A = [[[1.0, 0], [1.0, 1.0]], [[1.0, 0], [1.0, 1.0]]] B = [[[2.0, 2.0, 2.0], [4.0, 4.0, 4.0]], [[4.0, 4.0, 4.0], [8.0, 8.0, 8.0]]] trsm(A, B, alpha=0.5) = [[[1.0, 1.0, 1.0], [1.0, 1.0, 1.0]], [[2.0, 2.0, 2.0], [2.0, 2.0, 2.0]]]

Defined in src/operator/tensor/la\_op.cc:L396

#### Value

out The result mx.symbol

mx.symbol.LinearRegressionOutput

LinearRegressionOutput:Computes and optimizes for squared loss during backward propagation. Just outputs "data" during forward propagation.

### **Description**

If :math: '\haty\_i' is the predicted value of the i-th sample, and :math: 'y\_i' is the corresponding target value, then the squared loss estimated over :math: 'n' samples is defined as

### Usage

```
mx.symbol.LinearRegressionOutput(...)
```

### **Arguments**

data NDArray-or-Symbol Input data to the function.

1abel NDArray-or-Symbol Input label to the function.

grad.scale float, optional, default=1 Scale the gradient by a float factor

name string, optional Name of the resulting symbol.

### Details

 $: math: `\text{textSquaredLoss}(\text{textbfY}, \text{hat}\text{textbfY}) = \frac{1n \sum_{i=0}^{n-1} \text{textbfy}_i - \text{hat}\text{textbfy}_i - \text{hat}\text{textbfy}_i$ 

.. note:: Use the LinearRegressionOutput as the final output layer of a net.

The storage type of "label" can be "default" or "csr"

 $- Linear Regression Output (default, \ default) = default - Linear Regression Output (default, \ csr) = default$ 

By default, gradients of this loss function are scaled by factor '1/m', where m is the number of regression outputs of a training example. The parameter 'grad\_scale' can be used to change this scale to 'grad\_scale/m'.

Defined in src/operator/regression\_output.cc:L92

### Value

394 mx.symbol.load.json

 ${\tt mx.symbol.load}$ 

Load an mx.symbol object

## Description

Load an mx.symbol object

## Usage

```
mx.symbol.load(file.name)
```

## Arguments

filename

the filename (including the path)

# **Examples**

```
data = mx.symbol.Variable('data')
mx.symbol.save(data, 'temp.symbol')
data2 = mx.symbol.load('temp.symbol')
```

mx.symbol.load.json

Load an mx.symbol object from a json string

# Description

Load an mx.symbol object from a json string

## Arguments

str

the json str represent a mx.symbol

mx.symbol.log 395

mx.symbol.log

log:Returns element-wise Natural logarithmic value of the input.

### **Description**

The natural logarithm is logarithm in base  $e^*$ , so that "log(exp(x)) = x"

## Usage

```
mx.symbol.log(...)
```

## Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

The storage type of "log" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L77

### Value

out The result mx.symbol

mx.symbol.log10

log10:Returns element-wise Base-10 logarithmic value of the input.

## **Description**

```
10**log10(x) = x
```

### Usage

```
mx.symbol.log10(...)
```

### Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

The storage type of "log10" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L94

396 mx.symbol.log2

### Value

out The result mx.symbol

mx.symbol.log1p

log1p:Returns element-wise "log(1 + x)" value of the input.

### **Description**

This function is more accurate than "log(1 + x)" for small "x" so that :math: '1+x\approx 1'

## Usage

```
mx.symbol.log1p(...)
```

## Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

The storage type of "log1p" output depends upon the input storage type:

 $-\log 1p(\text{default}) = \text{default} - \log 1p(\text{row\_sparse}) = \text{row\_sparse} - \log 1p(\text{csr}) = \text{csr}$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L199

#### Value

out The result mx.symbol

mx.symbol.log2

log2:Returns element-wise Base-2 logarithmic value of the input.

### **Description**

$$2**\log 2(x) = x$$

### Usage

```
mx.symbol.log2(...)
```

## Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.logical\_not

397

## **Details**

The storage type of "log2" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_logexp.cc:L106

### Value

out The result mx.symbol

mx.symbol.logical\_not logical\_not:Returns the result of logical NOT (!) function

## **Description**

```
Example: logical_not([-2., 0., 1.]) = [0., 1., 0.]
```

## Usage

```
mx.symbol.logical_not(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### Value

out The result mx.symbol

```
\verb|mx.symbol.LogisticRegressionOutput|\\
```

LogisticRegressionOutput:Applies a logistic function to the input.

# **Description**

The logistic function, also known as the sigmoid function, is computed as :math: '\frac11+exp(-\textbfx)'.

```
mx.symbol.LogisticRegressionOutput(...)
```

## Arguments

data NDArray-or-Symbol Input data to the function.

1abel NDArray-or-Symbol Input label to the function.

grad. scale float, optional, default=1 Scale the gradient by a float factor

name string, optional Name of the resulting symbol.

### **Details**

Commonly, the sigmoid is used to squash the real-valued output of a linear model :math: 'wTx+b' into the [0,1] range so that it can be interpreted as a probability. It is suitable for binary classification or probability prediction tasks.

.. note:: Use the LogisticRegressionOutput as the final output layer of a net.

The storage type of "label" can be "default" or "csr"

LogisticRegressionOutput(default, default) = default - LogisticRegressionOutput(default, csr) = default

The loss function used is the Binary Cross Entropy Loss:

```
:math: -(y\log(p) + (1 - y)\log(1 - p))
```

Where 'y' is the ground truth probability of positive outcome for a given example, and 'p' the probability predicted by the model. By default, gradients of this loss function are scaled by factor '1/m', where m is the number of regression outputs of a training example. The parameter 'grad\_scale' can be used to change this scale to 'grad\_scale/m'.

Defined in src/operator/regression\_output.cc:L152

#### Value

out The result mx.symbol

### Description

Examples::

```
mx.symbol.log_softmax(...)
```

mx.symbol.LRN399

## **Arguments**

data	NDArray-or-Symbol The input array.
axis	int, optional, default='-1' The axis along which to compute softmax.
temperature	double or None, optional, default=None Temperature parameter in softmax
dtype	None, 'float16', 'float32', 'float64',optional, default='None' DType of the output in case this can't be inferred. Defaults to the same as input's dtype if not defined (dtype=None).
use.length	boolean or None, optional, default=0 Whether to use the length input as a mask

over the data input.

string, optional Name of the resulting symbol. name

### **Details**

```
\gg x = mx.nd.array([1, 2, .1]) \gg mx.nd.log_softmax(x).asnumpy() array([-1.41702998, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.41702995, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.4170295, -0.41702005, -0.4170205, -0.4170205, -0.4170205, -0.4170205, -0.4170200
  -2.31702995], dtype=float32)
  >> x = mx.nd.array([[1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, 1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1]]) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() array([[-1, 2, .1], [.1, 2, .1])) >> mx.nd.log_softmax(x, axis=0).asnumpy() >> mx.nd.log_softmax(x, axis=0).asnumpy() >> mx.nd.log_softmax(x,
0.34115392, -0.69314718, -1.24115396], [-1.24115396, -0.69314718, -0.34115392]], dtype=float32)
```

### Value

out The result mx.symbol

mx.symbol.LRN LRN:Applies local response normalization to the input.

# **Description**

The local response normalization layer performs "lateral inhibition" by normalizing over local input regions.

## Usage

```
mx.symbol.LRN(...)
```

## **Arguments**

data	NDArray-or-Symbol Input data to LRN
alpha	float, optional, default=9.99999975e-05 The variance scaling parameter :math: '\alpha' in the LRN expression.
beta	float, optional, default=0.75 The power parameter :math: '\beta' in the LRN expression.
knorm	float, optional, default=2 The parameter :math: 'k' in the LRN expression.
nsize	int (non-negative), required normalization window width in elements.
name	string, optional Name of the resulting symbol.

#### **Details**

If :math: 'a\_x,y^i' is the activity of a neuron computed by applying kernel :math: 'i' at position :math: '(x, y)' and then applying the ReLU nonlinearity, the response-normalized activity :math: 'b\_x,y^i' is given by the expression:

```
.. math:: b_x,y^i = \frac{x,y^i}Bigg(k + \frac{sum_j=max(0, i-\frac{2)^min(N-1, i+\frac{2)}{min(N-1, i+\frac{
```

where the sum runs over :math: 'n' "adjacent" kernel maps at the same spatial position, and :math: 'N' is the total number of kernels in the layer.

Defined in src/operator/nn/lrn.cc:L158

#### Value

out The result mx.symbol

mx.symbol.MAERegressionOutput

MAERegressionOutput:Computes mean absolute error of the input.

### **Description**

MAE is a risk metric corresponding to the expected value of the absolute error.

## Usage

```
mx.symbol.MAERegressionOutput(...)
```

### **Arguments**

data NDArray-or-Symbol Input data to the function.

1abel NDArray-or-Symbol Input label to the function.

grad.scale float, optional, default=1 Scale the gradient by a float factor

name string, optional Name of the resulting symbol.

#### **Details**

If :math: '\haty\_i' is the predicted value of the i-th sample, and :math: 'y\_i' is the corresponding target value, then the mean absolute error (MAE) estimated over :math: 'n' samples is defined as :math: '\textMAE(\textbfY, \hat\textbfY) = \frac1n \sum\_i=0^n-1 \lVert \textbfy\_i - \hat\textbfy\_i \rVert 1'

.. note:: Use the MAERegressionOutput as the final output layer of a net.

The storage type of "label" can be "default" or "csr"

- MAERegressionOutput(default, default) = default - MAERegressionOutput(default, csr) = default By default, gradients of this loss function are scaled by factor '1/m', where m is the number of regression outputs of a training example. The parameter 'grad\_scale' can be used to change this scale to 'grad scale/m'.

Defined in src/operator/regression\_output.cc:L120

mx.symbol.MakeLoss 401

#### Value

out The result mx.symbol

mx.symbol.MakeLoss

MakeLoss:Make your own loss function in network construction.

# **Description**

This operator accepts a customized loss function symbol as a terminal loss and the symbol should be an operator with no backward dependency. The output of this function is the gradient of loss with respect to the input data.

#### Usage

```
mx.symbol.MakeLoss(...)
```

#### **Arguments**

data NDArray-or-Symbol Input array.

grad.scale float, optional, default=1 Gradient scale as a supplement to unary and binary

operators

valid.thresh float, optional, default=0 clip each element in the array to 0 when it is less than

"valid\_thresh". This is used when "normalization" is set to "valid".

normalization 'batch', 'null', 'valid', optional, default='null' If this is set to null, the output

gradient will not be normalized. If this is set to batch, the output gradient will be divided by the batch size. If this is set to valid, the output gradient will be

divided by the number of valid input elements.

name string, optional Name of the resulting symbol.

### **Details**

For example, if you are a making a cross entropy loss function. Assume "out" is the predicted output and "label" is the true label, then the cross entropy can be defined as::

```
cross\_entropy = label * log(out) + (1 - label) * log(1 - out) loss = MakeLoss(cross\_entropy)
```

We will need to use "MakeLoss" when we are creating our own loss function or we want to combine multiple loss functions. Also we may want to stop some variables' gradients from backpropagation. See more detail in "BlockGrad" or "stop\_gradient".

In addition, we can give a scale to the loss by setting "grad\_scale", so that the gradient of the loss will be rescaled in the backpropagation.

.. note:: This operator should be used as a Symbol instead of NDArray.

Defined in src/operator/make\_loss.cc:L71

## Value

402 mx.symbol.max

mx.symbol.make\_loss

make\_loss:Make your own loss function in network construction.

#### **Description**

This operator accepts a customized loss function symbol as a terminal loss and the symbol should be an operator with no backward dependency. The output of this function is the gradient of loss with respect to the input data.

## Usage

```
mx.symbol.make_loss(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

For example, if you are a making a cross entropy loss function. Assume "out" is the predicted output and "label" is the true label, then the cross entropy can be defined as::

```
cross\_entropy = label * log(out) + (1 - label) * log(1 - out) loss = make\_loss(cross\_entropy)
```

We will need to use "make\_loss" when we are creating our own loss function or we want to combine multiple loss functions. Also we may want to stop some variables' gradients from backpropagation. See more detail in "BlockGrad" or "stop\_gradient".

The storage type of "make\_loss" output depends upon the input storage type:

- make\_loss(default) = default - make\_loss(row\_sparse) = row\_sparse

Defined in src/operator/tensor/elemwise unary op basic.cc:L359

### Value

out The result mx.symbol

mx.symbol.max

max: Computes the max of array elements over given axes.

### **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L32

```
mx.symbol.max(...)
```

mx.symbol.max\_axis 403

#### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

 $mx.symbol.max_axis$   $max_a$ 

max\_axis:Computes the max of array elements over given axes.

# **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L32

### Usage

```
mx.symbol.max_axis(...)
```

### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

404 mx.symbol.mean

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.mean m

mean: Computes the mean of array elements over given axes.

## **Description**

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L84

### Usage

```
mx.symbol.mean(...)
```

### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

## Value

mx.symbol.moments 405

mx.symbol.moments

moments: Calculate the mean and variance of 'data'.

# **Description**

The mean and variance are calculated by aggregating the contents of data across axes. If x is 1-D and axes = [0] this is just the mean and variance of a vector.

#### Usage

```
mx.symbol.moments(...)
```

#### **Arguments**

data NDArray-or-Symbol Input ndarray

axes Shape or None, optional, default=None Array of ints. Axes along which to

compute mean and variance.

keepdims boolean, optional, default=0 produce moments with the same dimensionality as

the input.

name string, optional Name of the resulting symbol.

### **Details**

#### Example:

```
x = [[1, 2, 3], [4, 5, 6]] mean, var = moments(data=x, axes=[0]) mean = [2.5, 3.5, 4.5] var = [2.25, 2.25, 2.25] mean, var = moments(data=x, axes=[1]) mean = [2.0, 5.0] var = [0.66666667] mean, var = moments(data=x, axis=[0, 1]) mean = [3.5] var = [2.9166667]
```

Defined in src/operator/nn/moments.cc:L54

#### Value

out The result mx.symbol

```
mx.symbol.mp_lamb_update_phase1
```

mp\_lamb\_update\_phase1:Mixed Precision version of Phase I of lamb update it performs the following operations and returns g:.

## Description

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

```
mx.symbol.mp_lamb_update_phase1(...)
```

### **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient

mean NDArray-or-Symbol Moving mean

var NDArray-or-Symbol Moving variance

weight32 NDArray-or-Symbol Weight32

beta1 float, optional, default=0.899999976 The decay rate for the 1st moment esti-

mates.

beta2 float, optional, default=0.999000013 The decay rate for the 2nd moment esti-

mates.

epsilon float, optional, default=9.99999997e-07 A small constant for numerical stability.

t int, required Index update count.

bias.correction

boolean, optional, default=1 Whether to use bias correction.

wd float, required Weight decay augments the objective function with a regulariza-

tion term that penalizes large weights. The penalty scales with the square of the

magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

name string, optional Name of the resulting symbol.

### **Details**

.. math:: \begingather\* grad32 = grad(float16) \* rescale\_grad if (grad < -clip\_gradient) then grad = -clip\_gradient if (grad > clip\_gradient) then grad = clip\_gradient

mean = beta1 \* mean + (1 - beta1) \* grad; variance = beta2 \* variance + (1. - beta2) \* grad ^ 2;

if (bias\_correction) then mean\_hat = mean / (1. - beta1^t); var\_hat = var / (1 - beta2^t); g = mean\_hat / (var\_hat^(1/2) + epsilon) + wd \* weight32; else g = mean / (var\_data^(1/2) + epsilon) + wd \* weight32; \endgather\*

Defined in src/operator/optimizer\_op.cc:L1024

## Value

```
mx.symbol.mp_lamb_update_phase2
```

mp\_lamb\_update\_phase2:Mixed Precision version Phase II of lamb update it performs the following operations and updates grad.

# Description

Link to paper: https://arxiv.org/pdf/1904.00962.pdf

## Usage

```
mx.symbol.mp_lamb_update_phase2(...)
```

## **Arguments**

weight	NDArray-or-Symbol Weight
g	NDArray-or-Symbol Output of mp_lamb_update_phase 1
r1	NDArray-or-Symbol r1
r2	NDArray-or-Symbol r2
weight32	NDArray-or-Symbol Weight32
lr	float, required Learning rate
lower.bound	float, optional, default=-1 Lower limit of norm of weight. If lower_bound <= 0, Lower limit is not set
upper.bound	float, optional, default=-1 Upper limit of norm of weight. If upper_bound <= 0, Upper limit is not set
name	string, optional Name of the resulting symbol.

### **Details**

```
.. math:: \begingather* if (lower_bound >= 0) then r1 = max(r1, lower_bound) if (upper_bound >= 0) then r1 = max(r1, upper_bound) if (r1 == 0 \text{ or } r2 == 0) then lr = lr \text{ else } lr = lr * (r1/r2) \text{ weight32} = weight32 - lr * g weight(float16) = weight32 \endgather*

Defined in src/operator/optimizer_op.cc:L1066
```

# Value

mx.symbol.mp\_nag\_mom\_update

mp\_nag\_mom\_update:Update function for multi-precision Nesterov Accelerated Gradient( NAG) optimizer.

## **Description**

Defined in src/operator/optimizer\_op.cc:L736

## Usage

```
mx.symbol.mp_nag_mom_update(...)
```

## **Arguments**

weight NDArray-or-Symbol Weight NDArray-or-Symbol Gradient grad NDArray-or-Symbol Momentum mom NDArray-or-Symbol Weight32 weight32 lr float, required Learning rate momentum float, optional, default=0 The decay rate of momentum estimates at each epoch. float, optional, default=0 Weight decay augments the objective function with a wd regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight. rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad. clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

clip\_gradient), -clip\_gradient).

string, optional Name of the resulting symbol.

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

## Value

name

```
mx.symbol.mp_sgd_mom_update
```

mp\_sgd\_mom\_update:Updater function for multi-precision sgd optimizer

### **Description**

mp\_sgd\_mom\_update:Updater function for multi-precision sgd optimizer

### Usage

```
mx.symbol.mp_sgd_mom_update(...)
```

### **Arguments**

NDArray-or-Symbol Weight weight grad NDArray-or-Symbol Gradient NDArray-or-Symbol Momentum mom weight32 NDArray-or-Symbol Weight32 lr float, required Learning rate float, optional, default=0 The decay rate of momentum estimates at each epoch. momentum float, optional, default=0 Weight decay augments the objective function with a wd regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight. rescale.grad float, optional, default=1 Rescale gradient to grad = rescale grad\*grad. clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient] If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip\_gradient), -clip\_gradient). lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype is row\_sparse and both weight and momentum have the same stype name string, optional Name of the resulting symbol.

## Value

mx.symbol.mp\_sgd\_update

mp\_sgd\_update:Updater function for multi-precision sgd optimizer

### **Description**

mp\_sgd\_update:Updater function for multi-precision sgd optimizer

#### Usage

```
mx.symbol.mp_sgd_update(...)
```

## **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol gradient weight32 NDArray-or-Symbol Weight32 lr float, required Learning rate

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row\_sparse.

name string, optional Name of the resulting symbol.

## Value

out The result mx.symbol

mx.symbol.multi\_all\_finite

multi\_all\_finite:Check if all the float numbers in all the arrays are finite (used for AMP)

## Description

Defined in src/operator/contrib/all\_finite.cc:L133

mx.symbol.multi\_lars 411

### Usage

```
mx.symbol.multi_all_finite(...)
```

### **Arguments**

data NDArray-or-Symbol[] Arrays

num.arrays int, optional, default='1' Number of arrays.

init.output boolean, optional, default=1 Initialize output to 1.

name string, optional Name of the resulting symbol.

### Value

out The result mx.symbol

 $\verb|mx.symbol.multi_lars| \textit{multi_lars}: \textit{Compute the LARS coefficients of multiple weights and} \\$ 

grads from their sums of square"

# Description

Defined in src/operator/contrib/multi\_lars.cc:L37

# Usage

```
mx.symbol.multi_lars(...)
```

# **Arguments**

1rs NDArray-or-Symbol Learning rates to scale by LARS coefficient

weights.sum.sq NDArray-or-Symbol sum of square of weights arrays grads.sum.sq NDArray-or-Symbol sum of square of gradients arrays

wds NDArray-or-Symbol weight decays

eta float, required LARS eta eps float, required LARS eps

rescale.grad float, optional, default=1 Gradient rescaling factor name string, optional Name of the resulting symbol.

## Value

```
mx.symbol.multi_mp_sgd_mom_update
```

multi\_mp\_sgd\_mom\_update:Momentum update function for multiprecision Stochastic Gradient Descent (SGD) optimizer.

## **Description**

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

# Usage

```
mx.symbol.multi_mp_sgd_mom_update(...)
```

## **Arguments**

	data	NDArray-or-Symbol[] Weights
	lrs	tuple of <float>, required Learning rates.</float>
		tuple of <float>, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.</float>
	momentum	float, optional, default=0 The decay rate of momentum estimates at each epoch.
	rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
	clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
	num.weights	int, optional, default='1' Number of updated weights.

## **Details**

name

```
.. math:: v_1 = \alpha V_0 \times V_t = \gamma V_t - \alpha V_t
```

string, optional Name of the resulting symbol.

# Value

```
\verb|mx.symbol.multi_mp_sgd_update| \\
```

multi\_mp\_sgd\_update:Update function for multi-precision Stochastic Gradient Descent (SDG) optimizer.

# Description

It updates the weights using::

# Usage

```
mx.symbol.multi_mp_sgd_update(...)
```

# Arguments

data	NDArray-or-Symbol[] Weights	
lrs	tuple of <float>, required Learning rates.</float>	
wds	tuple of <float>, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.</float>	
rescale.grad float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.		
clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient clip_gradient]. If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(gradient), -clip_gradient).		
num.weights	int, optional, default='1' Number of updated weights.	
name	string, optional Name of the resulting symbol.	

# **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
Defined in src/operator/optimizer_op.cc:L408
```

# Value

```
mx.symbol.multi_sgd_mom_update
```

multi\_sgd\_mom\_update:Momentum update function for Stochastic Gradient Descent (SGD) optimizer.

## **Description**

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

### Usage

```
mx.symbol.multi_sgd_mom_update(...)
```

#### **Arguments**

data	NDArray-or-Symbol[]	Weights,	gradients and momentum
------	---------------------	----------	------------------------

1rs tuple of <float>, required Learning rates.

wds tuple of <float>, required Weight decay augments the objective function with

a regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

name string, optional Name of the resulting symbol.

#### **Details**

```
.. math::
```

```
v_1 = \alpha \ T(W_0) \ v_t = \gamma \ v_{-1} - \alpha \ T(W_{t-1}) \ W_t = W_{t-1} + v_t
```

It updates the weights using::

```
v = momentum * v - learning_rate * gradient weight += v
```

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

Defined in src/operator/optimizer\_op.cc:L365

#### Value

```
{\tt mx.symbol.multi\_sgd\_update}
```

multi\_sgd\_update:Update function for Stochastic Gradient Descent (SDG) optimizer.

# Description

It updates the weights using::

# Usage

```
mx.symbol.multi_sgd_update(...)
```

# Arguments

data	NDArray-or-Symbol[] Weights	
lrs	tuple of <float>, required Learning rates.</float>	
wds	tuple of <float>, required Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.</float>	
rescale.grad float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.		
clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient clip_gradient]. If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(gradient), -clip_gradient).		
num.weights	int, optional, default='1' Number of updated weights.	
name	string, optional Name of the resulting symbol.	

# **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
Defined in src/operator/optimizer_op.cc:L320
```

# Value

```
mx.symbol.multi_sum_sq
```

multi\_sum\_sq:Compute the sums of squares of multiple arrays

## **Description**

Defined in src/operator/contrib/multi\_sum\_sq.cc:L36

## Usage

```
mx.symbol.multi_sum_sq(...)
```

# **Arguments**

data NDArray-or-Symbol[] Arrays

num.arrays int, required number of input arrays.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

```
mx.symbol.nag_mom_update
```

nag\_mom\_update:Update function for Nesterov Accelerated Gradient( NAG) optimizer. It updates the weights using the following formula,

### **Description**

```
.. math:: v_t = \gamma v_{t-1} + \epsilon * \Lambda J(W_{t-1} - \gamma v_t) W_t = W_{t-1} - v_t
```

### Usage

```
mx.symbol.nag_mom_update(...)
```

# **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient mom NDArray-or-Symbol Momentum 1r float, required Learning rate

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

mx.symbol.nanprod 417

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

name string, optional Name of the resulting symbol.

### **Details**

Where :math: '\eta' is the learning rate of the optimizer :math: '\gamma' is the decay rate of the momentum estimate :math: '\v\_t' is the update vector at time step 't' :math: '\W\_t' is the weight vector at time step 't'

Defined in src/operator/optimizer\_op.cc:L717

#### Value

out The result mx.symbol

mx.symbol.nanprod nanprod:Computes the product of array elements over given axes treating Not a Numbers ("NaN") as one.

### Description

nanprod:Computes the product of array elements over given axes treating Not a Numbers ("NaN") as one.

### Usage

```
mx.symbol.nanprod(...)
```

### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

418 mx.symbol.nansum

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

### **Details**

Defined in src/operator/tensor/broadcast\_reduce\_prod\_value.cc:L47

#### Value

out The result mx.symbol

mx.symbol.nansum nansum: Computes the sum of array elements over given axes treating

Not a Numbers ("NaN") as zero.

## **Description**

nansum:Computes the sum of array elements over given axes treating Not a Numbers ("NaN") as zero.

### Usage

```
mx.symbol.nansum(...)
```

## **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

mx.symbol.negative 419

### **Details**

Defined in src/operator/tensor/broadcast\_reduce\_sum\_value.cc:L102

### Value

out The result mx.symbol

mx.symbol.negative

negative: Numerical negative of the argument, element-wise.

## **Description**

The storage type of "negative" output depends upon the input storage type:

### Usage

```
mx.symbol.negative(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

```
- negative(default) = default - negative(row_sparse) = row_sparse - negative(csr) = csr
```

## Value

out The result mx.symbol

mx.symbol.norm

norm: Computes the norm on an NDArray.

# Description

This operator computes the norm on an NDArray with the specified axis, depending on the value of the ord parameter. By default, it computes the L2 norm on the entire array. Currently only ord=2 supports sparse ndarrays.

```
mx.symbol.norm(...)
```

420 mx.symbol.normal

## **Arguments**

data	NDArray-or-Symbol The input
ord	int, optional, default='2' Order of the norm. Currently ord=1 and ord=2 is supported.
axis	Shape or None, optional, default=None The axis or axes along which to perform the reduction. The default, 'axis=()', will compute over all elements into a scalar array with shape '(1,)'. If 'axis' is int, a reduction is performed on a particular axis. If 'axis' is a 2-tuple, it specifies the axes that hold 2-D matrices, and the matrix norms of these matrices are computed.
out.dtype	None, 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', optional, default='None' The data type of the output.
keepdims	boolean, optional, default=0 If this is set to 'True', the reduced axis is left in the result as dimension with size one.
name	string, optional Name of the resulting symbol.

## **Details**

```
Examples::
```

```
x = [[[1, 2], [3, 4]], [[2, 2], [5, 6]]]

norm(x, ord=2, axis=1) = [[3.1622777 4.472136] [5.3851647 6.3245554]]

norm(x, ord=1, axis=1) = [[4., 6.], [7., 8.]]

rsp = x.cast_storage('row_sparse')

norm(rsp) = [5.47722578]

csr = x.cast_storage('csr')

norm(csr) = [5.47722578]

Defined in src/operator/tensor/broadcast_reduce_norm_value.cc:L89
```

### Value

out The result mx.symbol

mx.symbol.normal

normal:Draw random samples from a normal (Gaussian) distribution.

# Description

.. note:: The existing alias "normal" is deprecated.

```
mx.symbol.normal(...)
```

mx.symbol.ones\_like 421

## **Arguments**

loc	float, optional, default=0 Mean of the distribution.
scale	float, optional, default=1 Standard deviation of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

### **Details**

Samples are distributed according to a normal distribution parametrized by \*loc\* (mean) and \*scale\* (standard deviation).

Example::

```
normal(loc=0, scale=1, shape=(2,2)) = [[ 1.89171135, -1.16881478], [-1.23474145, 1.55807114]] Defined in src/operator/random/sample_op.cc:L113
```

## Value

out The result mx.symbol

 ${\tt mx.symbol.ones\_like}$  ones\_like: Return an array of ones with the same shape and type as the input array.

# Description

Examples::

# Usage

```
mx.symbol.ones_like(...)
```

# Arguments

data	NDArray-or-Symbol	The input
------	-------------------	-----------

name string, optional Name of the resulting symbol.

# **Details**

```
x = [[0., 0., 0.], [0., 0., 0.]]
ones_like(x) = [[1., 1., 1.], [1., 1., 1.]]
```

422 mx.symbol.one\_hot

## Value

out The result mx.symbol

mx.symbol.one\_hot

one\_hot:Returns a one-hot array.

# **Description**

The locations represented by 'indices' take value 'on\_value', while all other locations take value 'off value'.

## Usage

```
mx.symbol.one_hot(...)
```

## **Arguments**

indices	NDArray-or-Symbol array of locations where to set on_value
depth	int, required Depth of the one hot dimension.
ueptn	int, required Depth of the one not difficultion.
on.value	double, optional, default=1 The value assigned to the locations represented by indices.
off.value	double, optional, default=0 The value assigned to the locations not represented by indices.
dtype	'bfloat16', 'float16', 'float32', 'float64', 'int32', 'int64', 'int8', 'uint8', optional, default='float32' DType of the output
name	string, optional Name of the resulting symbol.

## **Details**

```
'one_hot' operation with 'indices' of shape "(i0, i1)" and 'depth' of "d" would result in an output
array of shape "(i0, i1, d)" with::
output[i,j,:] = off_value output[i,j,indices[i,j]] = on_value
Examples::
one_hot([1,0,2,0], 3) = [[ 0. 1. 0.] [ 1. 0. 0.] [ 0. 0. 1.] [ 1. 0. 0.]]
one_hot([1,0,2,0], 3, on_value=8, off_value=1, dtype='int32') = [[1 8 1] [8 1 1] [1 1 8] [8 1 1]]
one_hot([[1,0],[1,0],[2,0]], 3) = [[[0.1.0.][1.0.0.]]
[[ 0. 1. 0.] [ 1. 0. 0.]]
[[ 0. 0. 1.] [ 1. 0. 0.]]]
Defined in src/operator/tensor/indexing_op.cc:L888
```

### Value

mx.symbol.Pad 423

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Pad: Pads an input array with a constant or edge values of the array.

### **Description**

```
.. note:: 'Pad' is deprecated. Use 'pad' instead.
```

#### Usage

```
mx.symbol.Pad(...)
```

## **Arguments**

data NDArray-or-Symbol An n-dimensional input array.

mode 'constant', 'edge', 'reflect', required Padding type to use. "constant" pads with

'constant\_value' "edge" pads using the edge values of the input array "reflect"

pads by reflecting values with respect to the edges.

pad.width Shape(tuple), required Widths of the padding regions applied to the edges of

each axis. It is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". It should be of length "2\*N" where "N" is the number of dimensions of the array. This is equivalent to pad\_width in

numpy.pad, but flattened.

constant.value double, optional, default=0 The value used for padding when 'mode' is "con-

stant".

name string, optional Name of the resulting symbol.

# Details

.. note:: Current implementation only supports 4D and 5D input arrays with padding applied only on axes 1, 2 and 3. Expects axes 4 and 5 in 'pad\_width' to be zero.

This operation pads an input array with either a 'constant\_value' or edge values along each axis of the input array. The amount of padding is specified by 'pad\_width'.

'pad\_width' is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". The 'pad\_width' should be of length "2\*N" where "N" is the number of dimensions of the array.

For dimension "N" of the input array, "before\_N" and "after\_N" indicates how many values to add before and after the elements of the array along dimension "N". The widths of the higher two dimensions "before\_1", "after\_1", "before\_2", "after\_2" must be 0.

#### Example::

```
x = [[[[ 1. 2. 3.] [ 4. 5. 6.]]
[[ 7. 8. 9.] [ 10. 11. 12.]]]
[[[ 11. 12. 13.] [ 14. 15. 16.]]
[[ 17. 18. 19.] [ 20. 21. 22.]]]]
```

424 mx.symbol.pad

```
pad(x,mode="edge", pad_width=(0,0,0,0,1,1,1,1)) =
[[[[ 1. 1. 2. 3. 3.] [ 1. 1. 2. 3. 3.] [ 4. 4. 5. 6. 6.] [ 4. 4. 5. 6. 6.]]
[[ 7. 7. 8. 9. 9.] [ 7. 7. 8. 9. 9.] [ 10. 10. 11. 12. 12.] [ 10. 10. 11. 12. 12.]]]
[[[ 11. 11. 12. 13. 13.] [ 11. 11. 12. 13. 13.] [ 14. 14. 15. 16. 16.] [ 14. 14. 15. 16. 16.]]
[[ 17. 17. 18. 19. 19.] [ 17. 17. 18. 19. 19.] [ 20. 20. 21. 22. 22.] [ 20. 20. 21. 22. 22.]]]]
pad(x, mode="constant", constant_value=0, pad_width=(0,0,0,0,1,1,1,1)) =
[[[ 0. 0. 0. 0. 0.] [ 0. 1. 2. 3. 0.] [ 0. 4. 5. 6. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 7. 8. 9. 0.] [ 0. 10. 11. 12. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 11. 12. 13. 0.] [ 0. 14. 15. 16. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 17. 18. 19. 0.] [ 0. 20. 21. 22. 0.] [ 0. 0. 0. 0. 0.]]]]
Defined in src/operator/pad.cc:L766
```

#### Value

out The result mx.symbol

mx.symbol.pad

pad:Pads an input array with a constant or edge values of the array.

# **Description**

.. note:: 'Pad' is deprecated. Use 'pad' instead.

#### **Usage**

```
mx.symbol.pad(...)
```

#### **Arguments**

data	NDArray-or-Symbol An n-dimensional inp	ut array.
------	--	-----------

mode 'constant', 'edge', 'reflect', required Padding type to use. "constant" pads with

'constant\_value' "edge" pads using the edge values of the input array "reflect"

pads by reflecting values with respect to the edges.

pad.width Shape(tuple), required Widths of the padding regions applied to the edges of

each axis. It is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". It should be of length "2\*N" where "N" is the number of dimensions of the array. This is equivalent to pad\_width in

numpy.pad, but flattened.

constant.value double, optional, default=0 The value used for padding when 'mode' is "con-

stant".

name string, optional Name of the resulting symbol.

mx.symbol.pick 425

#### **Details**

.. note:: Current implementation only supports 4D and 5D input arrays with padding applied only on axes 1, 2 and 3. Expects axes 4 and 5 in 'pad\_width' to be zero.

This operation pads an input array with either a 'constant\_value' or edge values along each axis of the input array. The amount of padding is specified by 'pad\_width'.

'pad\_width' is a tuple of integer padding widths for each axis of the format "(before\_1, after\_1, ..., before\_N, after\_N)". The 'pad\_width' should be of length "2\*N" where "N" is the number of dimensions of the array.

For dimension "N" of the input array, "before\_N" and "after\_N" indicates how many values to add before and after the elements of the array along dimension "N". The widths of the higher two dimensions "before\_1", "after\_1", "before\_2", "after\_2" must be 0.

## Example::

```
x = [[[[ 1. 2. 3.] [ 4. 5. 6.]]
[[ 7. 8. 9.] [ 10. 11. 12.]]]
[[[ 11. 12. 13.] [ 14. 15. 16.]]
[[ 17. 18. 19.] [ 20. 21. 22.]]]]
pad(x,mode="edge", pad_width=(0,0,0,1,1,1,1)) =
[[[[ 1. 1. 2. 3. 3.] [ 1. 1. 2. 3. 3.] [ 4. 4. 5. 6. 6.] [ 4. 4. 5. 6. 6.]]
[[ 7. 7. 8. 9. 9.] [ 7. 7. 8. 9. 9.] [ 10. 10. 11. 12. 12.] [ 10. 10. 11. 12. 12.]]]
[[[ 11. 11. 12. 13. 13.] [ 11. 11. 12. 13. 13.] [ 14. 14. 15. 16. 16.] [ 14. 14. 15. 16. 16.]]
[[ 17. 17. 18. 19. 19.] [ 17. 17. 18. 19. 19.] [ 20. 20. 21. 22. 22.] [ 20. 20. 21. 22. 22.]]]]
pad(x, mode="constant", constant_value=0, pad_width=(0,0,0,0,1,1,1,1)) =
[[[ 0. 0. 0. 0. 0.] [ 0. 1. 2. 3. 0.] [ 0. 4. 5. 6. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 11. 12. 13. 0.] [ 0. 14. 15. 16. 0.] [ 0. 0. 0. 0. 0.]]
[[ 0. 0. 0. 0. 0.] [ 0. 17. 18. 19. 0.] [ 0. 20. 21. 22. 0.] [ 0. 0. 0. 0. 0.]]]]
Defined in src/operator/pad.cc:L766
```

#### Value

out The result mx.symbol

mx.symbol.pick

pick: Picks elements from an input array according to the input indices along the given axis.

#### **Description**

Given an input array of shape "(d0, d1)" and indices of shape "(i0,)", the result will be an output array of shape "(i0,)" with::

426 mx.symbol.pick

## Usage

```
mx.symbol.pick(...)
```

# **Arguments**

data NDArray-or-Symbol The input array index NDArray-or-Symbol The index array

axis int or None, optional, default='-1' int or None. The axis to picking the elements.

Negative values means indexing from right to left. If is 'None', the elements in

the index w.r.t the flattened input will be picked.

keepdims boolean, optional, default=0 If true, the axis where we pick the elements is left

in the result as dimension with size one.

mode 'clip', 'wrap', optional, default='clip' Specify how out-of-bound indices behave.

Default is "clip". "clip" means clip to the range. So, if all indices mentioned are too large, they are replaced by the index that addresses the last element along an

axis. "wrap" means to wrap around.

name string, optional Name of the resulting symbol.

#### **Details**

output[i] = input[i, indices[i]]

By default, if any index mentioned is too large, it is replaced by the index that addresses the last element along an axis (the 'clip' mode).

This function supports n-dimensional input and (n-1)-dimensional indices arrays.

Examples::

x = [[1., 2.], [3., 4.], [5., 6.]]

// picks elements with specified indices along axis 0 pick(x, y=[0,1], 0) = [1., 4.]

// picks elements with specified indices along axis 1 pick(x, y=[0,1,0], 1) = [1., 4., 5.]

// picks elements with specified indices along axis 1 using 'wrap' mode // to place indicies that would normally be out of bounds pick(x, y=[2,-1,-2], 1, mode='wrap') = [1., 4., 5.]

y = [[1.], [0.], [2.]]

// picks elements with specified indices along axis 1 and dims are maintained pick(x, y, 1, keep-dims=True) = [[2.], [3.], [6.]]

Defined in src/operator/tensor/broadcast\_reduce\_op\_index.cc:L151

#### Value

mx.symbol.Pooling 427

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Pooling:Performs pooling on the input.

### **Description**

The shapes for 1-D pooling are

### Usage

```
mx.symbol.Pooling(...)
```

# **Arguments**

data NDArray-or-Symbol Input data to the pooling operator.

kernel Shape(tuple), optional, default=[] Pooling kernel size: (y, x) or (d, y, x)

pool.type 'avg', 'lp', 'max', 'sum', optional, default='max' Pooling type to be applied.

global.pool boolean, optional, default=0 Ignore kernel size, do global pooling based on cur-

rent input feature map.

cudnn.off boolean, optional, default=0 Turn off cudnn pooling and use MXNet pooling

operator.

pooling.convention

'full', 'same', 'valid', optional, default='valid' Pooling convention to be applied.

stride Shape(tuple), optional, default=[] Stride: for pooling (y, x) or (d, y, x). Defaults

to 1 for each dimension.

pad Shape(tuple), optional, default=[] Pad for pooling: (y, x) or (d, y, x). Defaults to

no padding.

p.value int or None, optional, default='None' Value of p for Lp pooling, can be 1 or 2,

required for Lp Pooling.

count.include.pad

boolean or None, optional, default=None Only used for AvgPool, specify whether to count padding elements for averagecalculation. For example, with a 5\*5 kernel on a 3\*3 corner of a image, the sum of the 9 valid elements will be divided by 25 if this is set to true, or it will be divided by 9 if this is set to false. Defaults

to true.

layout None, 'NCDHW', 'NCHW', 'NCW', 'NDHWC', 'NHWC', 'NWC', optional,

default='None' Set layout for input and output. Empty for default layout: NCW

for 1d, NCHW for 2d and NCDHW for 3d.

name string, optional Name of the resulting symbol.

#### **Details**

- \*\*data\*\* and \*\*out\*\*: \*(batch\_size, channel, width)\* (NCW layout) or \*(batch\_size, width, channel)\* (NWC layout),

The shapes for 2-D pooling are

- \*\*data\*\* and \*\*out\*\*: \*(batch\_size, channel, height, width)\* (NCHW layout) or \*(batch\_size, height, width, channel)\* (NHWC layout),

out\_height = f(height, kernel[0], pad[0], stride[0]) out\_width = f(width, kernel[1], pad[1], stride[1])

The definition of \*f\* depends on "pooling\_convention", which has two options:

- \*\*valid\*\* (default)::

f(x, k, p, s) = floor((x+2\*p-k)/s)+1

- \*\*full\*\*, which is compatible with Caffe::

$$f(x, k, p, s) = ceil((x+2*p-k)/s)+1$$

When "global\_pool" is set to be true, then global pooling is performed. It will reset "kernel=(height, width)" and set the appropriate padding to 0.

Three pooling options are supported by "pool\_type":

- \*\*avg\*\*: average pooling - \*\*max\*\*: max pooling - \*\*sum\*\*: sum pooling - \*\*lp\*\*: Lp pooling

For 3-D pooling, an additional \*depth\* dimension is added before \*height\*. Namely the input data and output will have shape \*(batch\_size, channel, depth, height, width)\* (NCDHW layout) or \*(batch\_size, depth, height, width, channel)\* (NDHWC layout).

Notes on Lp pooling:

Lp pooling was first introduced by this paper: https://arxiv.org/pdf/1204.3968.pdf. L-1 pooling is simply sum pooling, while L-inf pooling is simply max pooling. We can see that Lp pooling stands between those two, in practice the most common value for p is 2.

For each window "X", the mathematical expression for Lp pooling is:

```
:math: f(X) = \sqrt{p}\sum_x^X x^p
```

Defined in src/operator/nn/pooling.cc:L414

#### Value

out The result mx.symbol

mx.symbol.Pooling\_v1 Pooling\_v1:This operator is DEPRECATED. Perform pooling on the input.

## **Description**

The shapes for 2-D pooling is

```
mx.symbol.Pooling_v1(...)
```

## **Arguments**

data NDArray-or-Symbol Input data to the pooling operator.

kernel Shape(tuple), optional, default=[] pooling kernel size: (y, x) or (d, y, x)

pool.type 'avg', 'max', 'sum',optional, default='max' Pooling type to be applied.

global.pool boolean, optional, default=0 Ignore kernel size, do global pooling based on cur-

rent input feature map.

pooling.convention

'full', 'valid', optional, default='valid' Pooling convention to be applied.

stride Shape(tuple), optional, default=[] stride: for pooling (y, x) or (d, y, x)

Shape(tuple), optional, default=[] pad for pooling: (y, x) or (d, y, x)

name string, optional Name of the resulting symbol.

#### **Details**

- \*\*data\*\*: \*(batch\_size, channel, height, width)\* - \*\*out\*\*: \*(batch\_size, num\_filter, out\_height, out\_width)\*, with::

 $out\_height = f(height, kernel[0], pad[0], stride[0]) out\_width = f(width, kernel[1], pad[1], stride[1])$ 

The definition of \*f\* depends on "pooling convention", which has two options:

- \*\*valid\*\* (default)::

f(x, k, p, s) = floor((x+2\*p-k)/s)+1

- \*\*full\*\*, which is compatible with Caffe::

f(x, k, p, s) = ceil((x+2\*p-k)/s)+1

But "global\_pool" is set to be true, then do a global pooling, namely reset "kernel=(height, width)".

Three pooling options are supported by "pool\_type":

- \*\*avg\*\*: average pooling - \*\*max\*\*: max pooling - \*\*sum\*\*: sum pooling

1-D pooling is special case of 2-D pooling with \*weight=1\* and \*kernel[1]=1\*.

For 3-D pooling, an additional \*depth\* dimension is added before \*height\*. Namely the input data will have shape \*(batch\_size, channel, depth, height, width)\*.

Defined in src/operator/pooling\_v1.cc:L104

#### Value

```
mx.symbol.preloaded_multi_mp_sgd_mom_update
```

preloaded\_multi\_mp\_sgd\_mom\_update:Momentum update function for multi-precision Stochastic Gradient Descent (SGD) optimizer.

# Description

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

### Usage

```
mx.symbol.preloaded_multi_mp_sgd_mom_update(...)
```

# **Arguments**

data	NDArray-or-Symbol[] Weights, gradients, momentums, learning rates and weight

decays

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

name string, optional Name of the resulting symbol.

#### **Details**

```
.. math::
```

```
v_1 = \alpha V_1 - \alpha V_2 - \alpha V_1 - \alpha V_1 - \alpha V_2 - \alpha V_1 - \alpha V_2 - \alpha V_2
```

It updates the weights using::

v = momentum \* v - learning\_rate \* gradient weight += v

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

Defined in src/operator/contrib/preloaded\_multi\_sgd.cc:L200

### Value

mx.symbol.preloaded\_multi\_mp\_sgd\_update

preloaded\_multi\_mp\_sgd\_update:Update function for multi-precision Stochastic Gradient Descent (SDG) optimizer.

### **Description**

It updates the weights using::

## Usage

```
mx.symbol.preloaded_multi_mp_sgd_update(...)
```

## **Arguments**

data NDArray-or-Symbol[] Weights, gradients, learning rates and weight decays rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

name string, optional Name of the resulting symbol.

## Details

```
weight = weight - learning_rate * (gradient + wd * weight)

Defined in src/operator/contrib/preloaded_multi_sgd.cc:L140
```

#### Value

out The result mx.symbol

```
mx.symbol.preloaded_multi_sgd_mom_update
```

preloaded\_multi\_sgd\_mom\_update:Momentum update function for Stochastic Gradient Descent (SGD) optimizer.

## Description

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

```
mx.symbol.preloaded_multi_sgd_mom_update(...)
```

## Arguments

data NDArray-or-Symbol[] Weights, gradients, momentum, learning rates and weight

decays

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

name string, optional Name of the resulting symbol.

#### **Details**

.. math::

 $v_1 = \alpha V_t - 1 - \alpha V_t - \Omega V_t = \gamma V_t - 1 - \alpha V_t$ 

It updates the weights using::

v = momentum \* v - learning\_rate \* gradient weight += v

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

Defined in src/operator/contrib/preloaded\_multi\_sgd.cc:L91

### Value

out The result mx.symbol

mx.symbol.preloaded\_multi\_sgd\_update

preloaded\_multi\_sgd\_update:Update function for Stochastic Gradient

Descent (SDG) optimizer.

## Description

It updates the weights using::

# Usage

```
mx.symbol.preloaded_multi_sgd_update(...)
```

## **Arguments**

data NDArray-or-Symbol[] Weights, gradients, learning rates and weight decays rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip gradient, clip gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

num.weights int, optional, default='1' Number of updated weights.

name string, optional Name of the resulting symbol.

mx.symbol.prod 433

# **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
Defined in src/operator/contrib/preloaded_multi_sgd.cc:L42
```

## Value

out The result mx.symbol

mx.symbol.prod

prod:Computes the product of array elements over given axes.

# Description

Defined in src/operator/tensor/./broadcast\_reduce\_op.h:L31

# Usage

```
mx.symbol.prod(...)
```

# Arguments

data	NDArray-or-Symbol The input
axis	Shape or None, optional, default=None The axis or axes along which to perform the reduction.
	The default, 'axis=()', will compute over all elements into a scalar array with shape '(1,)'.
	If 'axis' is int, a reduction is performed on a particular axis.
	If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in the tuple.
	If 'exclude' is true, reduction will be performed on the axes that are NOT in axis instead.
	Negative values means indexing from right to left.
keepdims	boolean, optional, default=0 If this is set to 'True', the reduced axes are left in the result as dimension with size one.
exclude	boolean, optional, default=0 Whether to perform reduction on axis that are NOT in axis instead.
name	string, optional Name of the resulting symbol.

## Value

mx.symbol.radians

radians: Converts each element of the input array from degrees to radians.

## **Description**

## Usage

```
mx.symbol.radians(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

The storage type of "radians" output depends upon the input storage type:

- radians(default) = default - radians(row\_sparse) = row\_sparse - radians(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L351

## Value

out The result mx.symbol

```
mx.symbol.random_exponential
```

random\_exponential:Draw random samples from an exponential distribution.

# Description

Samples are distributed according to an exponential distribution parametrized by \*lambda\* (rate).

### Usage

```
mx.symbol.random_exponential(...)
```

### **Arguments**

lam	float, optional, default=1 Lambda parameter (rate) of the exponential distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64',optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

### **Details**

## Example::

```
exponential(lam=4, shape=(2,2)) = \hbox{\tt [[\ 0.0097189\ ,\ 0.08999364],\ [\ 0.04146638,\ 0.31715935]]} Defined in src/operator/random/sample_op.cc:L137
```

# Value

out The result mx.symbol

```
mx.symbol.random_gamma
```

 $random\_gamma: Draw\ random\ samples\ from\ a\ gamma\ distribution.$ 

# Description

Samples are distributed according to a gamma distribution parametrized by \*alpha\* (shape) and \*beta\* (scale).

## Usage

```
mx.symbol.random_gamma(...)
```

### **Arguments**

alpha	float, optional, default=1 Alpha parameter (shape) of the gamma distribution.
beta	float, optional, default=1 Beta parameter (scale) of the gamma distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64',optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

### **Details**

Example::

gamma(alpha=9, beta=0.5, shape=(2,2)) = [[ 7.10486984, 3.37695289], [ 3.91697288, 3.65933681]] Defined in src/operator/random/sample\_op.cc:L125

#### Value

out The result mx.symbol

```
mx.symbol.random_generalized_negative_binomial
```

random\_generalized\_negative\_binomial:Draw random samples from a generalized negative binomial distribution.

## **Description**

Samples are distributed according to a generalized negative binomial distribution parametrized by \*mu\* (mean) and \*alpha\* (dispersion). \*alpha\* is defined as \*1/k\* where \*k\* is the failure limit of the number of unsuccessful experiments (generalized to real numbers). Samples will always be returned as a floating point data type.

## Usage

```
mx.symbol.random_generalized_negative_binomial(...)
```

# Arguments

mu	float, optional, default=1 Mean of the negative binomial distribution.
alpha	float, optional, default=1 Alpha (dispersion) parameter of the negative binomial distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

## Details

```
Example::
```

```
generalized_negative_binomial(mu=2.0, alpha=0.3, shape=(2,2)) = [[ 2., 1.], [ 6., 4.]] Defined in src/operator/random/sample_op.cc:L179
```

### Value

```
mx.symbol.random_negative_binomial
```

random\_negative\_binomial:Draw random samples from a negative binomial distribution.

# Description

Samples are distributed according to a negative binomial distribution parametrized by \*k\* (limit of unsuccessful experiments) and \*p\* (failure probability in each experiment). Samples will always be returned as a floating point data type.

### Usage

```
mx.symbol.random_negative_binomial(...)
```

## **Arguments**

k	int, optional, default='1' Limit of unsuccessful experiments.
р	float, optional, default=1 Failure probability in each experiment.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

## **Details**

```
Example::
```

```
negative\_binomial(k=3, p=0.4, shape=(2,2)) = [[\ 4.,\ 7.],\ [\ 2.,\ 5.]]
```

Defined in src/operator/random/sample\_op.cc:L164

## Value

```
mx.symbol.random_normal
```

random\_normal:Draw random samples from a normal (Gaussian) distribution.

# Description

.. note:: The existing alias "normal" is deprecated.

## Usage

```
mx.symbol.random_normal(...)
```

# Arguments

loc	float, optional, default=0 Mean of the distribution.
scale	float, optional, default=1 Standard deviation of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

### **Details**

Samples are distributed according to a normal distribution parametrized by \*loc\* (mean) and \*scale\* (standard deviation).

## Example::

```
normal(loc=0, scale=1, shape=(2,2)) = [[1.89171135, -1.16881478], [-1.23474145, 1.55807114]]
Defined in src/operator/random/sample_op.cc:L113
```

## Value

```
mx.symbol.random_pdf_dirichlet
```

random\_pdf\_dirichlet:Computes the value of the PDF of \*sample\* of Dirichlet distributions with parameter \*alpha\*.

## **Description**

The shape of \*alpha\* must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*alpha\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the value of \*alpha\* at index \*i\*.

### Usage

```
mx.symbol.random_pdf_dirichlet(...)
```

## Arguments

sample	NDArray-or-Symbol Samples from the distributions.
alpha	NDArray-or-Symbol Concentration parameters of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
name	string, optional Name of the resulting symbol.

## **Details**

## Examples::

```
 \begin{array}{l} {\rm random\_pdf\_dirichlet(sample=[[1,2],[2,3],[3,4]], \, alpha=[2.5,\,2.5])=[38.413498,\,199.60245,\,564.56085]} \\ {\rm sample=[[[1,2,3],\,[10,\,20,\,30],\,[100,\,200,\,300]],\,[[0.1,\,0.2,\,0.3],\,[0.01,\,0.02,\,0.03],\,[0.001,\,0.002,\,0.003]]]} \\ {\rm random\_pdf\_dirichlet(sample=sample,\,\,alpha=[0.1,\,0.4,\,0.9])=[[2.3257459e-02,\,5.8420084e-04,\,1.4674458e-05],\,[9.2589635e-01,\,3.6860607e+01,\,1.4674468e+03]]} \\ {\rm Defined\ in\ src/operator/random/pdf\_op.cc:L316} \\ \end{array}
```

### Value

```
mx.symbol.random_pdf_exponential
```

random\_pdf\_exponential:Computes the value of the PDF of \*sample\* of exponential distributions with parameters \*lam\* (rate).

## **Description**

The shape of \*lam\* must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*lam\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the value of \*lam\* at index \*i\*.

## Usage

```
mx.symbol.random_pdf_exponential(...)
```

### **Arguments**

sample NDArray-or-Symbol Samples from the distributions.

lam NDArray-or-Symbol Lambda (rate) parameters of the distributions.

is.log boolean, optional, default=0 If set, compute the density of the log-probability

instead of the probability.

name string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
random_pdf_exponential(sample=[[1, 2, 3]], lam=[1]) = [[0.36787945, 0.13533528, 0.04978707]] sample = [[1,2,3], [1,2,3], [1,2,3]]
```

 $\begin{array}{l} random\_pdf\_exponential(sample=sample, lam=[1,0.5,0.25]) = [[0.36787945, 0.13533528, 0.04978707], \\ [0.30326533, 0.18393973, 0.11156508], [0.1947002, 0.15163267, 0.11809164]] \end{array}$ 

Defined in src/operator/random/pdf\_op.cc:L305

### Value

```
mx.symbol.random_pdf_gamma
```

random\_pdf\_gamma:Computes the value of the PDF of \*sample\* of gamma distributions with parameters \*alpha\* (shape) and \*beta\* (rate).

## Description

\*alpha\* and \*beta\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*alpha\* and \*beta\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*alpha\* and \*beta\* at index \*i\*.

## Usage

```
mx.symbol.random_pdf_gamma(...)
```

### **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
alpha	NDArray-or-Symbol Alpha (shape) parameters of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
beta	NDArray-or-Symbol Beta (scale) parameters of the distributions.
name	string, optional Name of the resulting symbol.

## **Details**

### Examples::

```
random_pdf_gamma(sample=[[1,2,3,4,5]], alpha=[5], beta=[1]) = [[0.01532831, 0.09022352, 0.16803136, 0.19536681, 0.17546739]] sample = [[1, 2, 3, 4, 5], [2, 3, 4, 5, 6], [3, 4, 5, 6, 7]] random_pdf_gamma(sample=sample, alpha=[5,6,7], beta=[1,1,1]) = [[0.01532831, 0.09022352, 0.16803136, 0.19536681, 0.17546739], [0.03608941, 0.10081882, 0.15629345, 0.17546739, 0.16062315], [0.05040941, 0.10419563, 0.14622283, 0.16062315, 0.14900276]] Defined in src/operator/random/pdf_op.cc:L303
```

### Value

```
mx.symbol.random_pdf_generalized_negative_binomial
```

random\_pdf\_generalized\_negative\_binomial:Computes the value of the PDF of \*sample\* of generalized negative binomial distributions with parameters \*mu\* (mean) and \*alpha\* (dispersion). This can be understood as a reparameterization of the negative binomial, where \*k\* = \*1 / alpha\*and \*p\* = \*1 / (mu \* alpha + 1)\*.

## **Description**

\*mu\* and \*alpha\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*mu\* and \*alpha\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*mu\* and \*alpha\* at index \*i\*.

## Usage

```
mx.symbol.random_pdf_generalized_negative_binomial(...)
```

### **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
mu	NDArray-or-Symbol Means of the distributions.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
alpha	NDArray-or-Symbol Alpha (dispersion) parameters of the distributions.
name	string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
random\_pdf\_generalized\_negative\_binomial(sample=[[1, 2, 3, 4]], alpha=[1], mu=[1]) = [[0.25, 0.125, 0.0625, 0.03125]]
```

 $sample = \hbox{\tt [[1,2,3,4], [1,2,3,4]]} \ random\_pdf\_generalized\_negative\_binomial(sample=sample, alpha=\hbox{\tt [1,0.6666]}, mu=\hbox{\tt [1,1.5]}) = \hbox{\tt [[0.25,0.125,0.0625,0.03125]}, \hbox{\tt [0.26517063,0.16573331,0.09667706,0.05437994]}]$ 

Defined in src/operator/random/pdf\_op.cc:L314

### Value

```
mx.symbol.random_pdf_negative_binomial
```

random\_pdf\_negative\_binomial:Computes the value of the PDF of samples of negative binomial distributions with parameters \*k\* (failure limit) and \*p\* (failure probability).

### **Description**

\*k\* and \*p\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*k\* and \*p\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*k\* and \*p\* at index \*i\*.

# Usage

```
mx.symbol.random_pdf_negative_binomial(...)
```

## **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
k	NDArray-or-Symbol Limits of unsuccessful experiments.
is.log	boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.
p	NDArray-or-Symbol Failure probabilities in each experiment.
name	string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
\begin{aligned} & \text{random\_pdf\_negative\_binomial}(\text{sample=}[[1,2,3,4]], \text{k=}[1], \text{p=a}[0.5]) = [[0.25, 0.125, 0.0625, 0.03125]] \\ & \text{# Note that k may be real-valued sample} = [[1,2,3,4], [1,2,3,4]] \\ & \text{random\_pdf\_negative\_binomial}(\text{sample=sample}, \text{k=}[1, 1.5], \text{p=}[0.5, 0.5]) = [[0.25, 0.125, 0.0625, 0.03125], [0.26516506, 0.16572815, 0.09667476, 0.05437956]] \end{aligned}
```

Defined in src/operator/random/pdf\_op.cc:L310

#### Value

```
mx.symbol.random_pdf_normal
```

random\_pdf\_normal:Computes the value of the PDF of \*sample\* of normal distributions with parameters \*mu\* (mean) and \*sigma\* (standard deviation).

### **Description**

\*mu\* and \*sigma\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*mu\* and \*sigma\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*mu\* and \*sigma\* at index \*i\*.

### Usage

```
mx.symbol.random_pdf_normal(...)
```

## **Arguments**

sample NDArray-or-Symbol Samples from the distributions.

mu NDArray-or-Symbol Means of the distributions.

is.log boolean, optional, default=0 If set, compute the density of the log-probability instead of the probability.

sigma NDArray-or-Symbol Standard deviations of the distributions.

name string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
sample = [[-2, -1, 0, 1, 2]] random_pdf_normal(sample=sample, mu=[0], sigma=[1]) = [[0.05399097, 0.24197073, 0.3989423, 0.24197073, 0.05399097]]
```

random\_pdf\_normal(sample=sample\*2, mu=[0,0], sigma=[1,2]) = [[0.05399097, 0.24197073, 0.3989423, 0.24197073, 0.05399097], [0.12098537, 0.17603266, 0.19947115, 0.17603266, 0.12098537]]

Defined in src/operator/random/pdf\_op.cc:L300

#### Value

```
mx.symbol.random_pdf_poisson
```

random\_pdf\_poisson: Computes the value of the PDF of \*sample\* of Poisson distributions with parameters \*lam\* (rate).

## **Description**

The shape of \*lam\* must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*lam\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the value of \*lam\* at index \*i\*.

## Usage

```
mx.symbol.random_pdf_poisson(...)
```

### **Arguments**

sample NDArray-or-Symbol Samples from the distributions.

lam NDArray-or-Symbol Lambda (rate) parameters of the distributions.

is.log boolean, optional, default=0 If set, compute the density of the log-probability

instead of the probability.

name string, optional Name of the resulting symbol.

#### **Details**

### Examples::

```
\begin{split} & \text{random\_pdf\_poisson}(\text{sample=}[[0,1,2,3]], \text{lam=}[1]) = [[0.36787945, 0.36787945, 0.18393973, 0.06131324]] \\ & \text{sample} = [[0,1,2,3], [0,1,2,3], [0,1,2,3]] \end{split}
```

random\_pdf\_poisson(sample=sample, lam=[1,2,3]) = [[0.36787945, 0.36787945, 0.18393973, 0.06131324], [0.13533528, 0.27067056, 0.27067056, 0.18044704], [0.04978707, 0.14936121, 0.22404182, 0.22404182]]

Defined in src/operator/random/pdf\_op.cc:L307

### Value

```
mx.symbol.random_pdf_uniform
```

random\_pdf\_uniform: Computes the value of the PDF of \*sample\* of uniform distributions on the intervals given by \*[low,high)\*.

## Description

\*low\* and \*high\* must have the same shape, which must match the leftmost subshape of \*sample\*. That is, \*sample\* can have the same shape as \*low\* and \*high\*, in which case the output contains one density per distribution, or \*sample\* can be a tensor of tensors with that shape, in which case the output is a tensor of densities such that the densities at index \*i\* in the output are given by the samples at index \*i\* in \*sample\* parameterized by the values of \*low\* and \*high\* at index \*i\*.

### Usage

```
mx.symbol.random_pdf_uniform(...)
```

## **Arguments**

sample	NDArray-or-Symbol Samples from the distributions.
low	NDArray-or-Symbol Lower bounds of the distributions.
is.log	boolean, optional, default= $0$ If set, compute the density of the log-probability instead of the probability.
high	NDArray-or-Symbol Upper bounds of the distributions.
name	string, optional Name of the resulting symbol.

### **Details**

### Examples::

```
random_pdf_uniform(sample=[[1,2,3,4]], low=[0], high=[10]) = [0.1, 0.1, 0.1, 0.1] sample = [[[1, 2, 3], [1, 2, 3]], [[1, 2, 3], [1, 2, 3]]] low = [[0, 0], [0, 0]] high = [[ 5, 10], [15, 20]] random_pdf_uniform(sample=sample, low=low, high=high) = [[[0.2, 0.2, 0.2], [0.1, 0.1, 0.1]], [[0.06667, 0.06667, 0.06667], [0.05, 0.05, 0.05]]]

Defined in src/operator/random/pdf_op.cc:L298
```

## Value

mx.symbol.random\_poisson

random\_poisson:Draw random samples from a Poisson distribution.

## **Description**

Samples are distributed according to a Poisson distribution parametrized by \*lambda\* (rate). Samples will always be returned as a floating point data type.

## Usage

```
mx.symbol.random_poisson(...)
```

# Arguments

lam	float, optional, default=1 Lambda parameter (rate) of the Poisson distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

# **Details**

```
Example::
```

```
poisson(lam=4, shape=(2,2)) = [[ 5., 2.], [ 4., 6.]]
Defined in src/operator/random/sample_op.cc:L150
```

## Value

out The result mx.symbol

```
mx.symbol.random_randint
```

random\_randint:Draw random samples from a discrete uniform distribution.

# Description

Samples are uniformly distributed over the half-open interval \*[low, high)\* (includes \*low\*, but excludes \*high\*).

### Usage

```
mx.symbol.random_randint(...)
```

#### **Arguments**

low long, required Lower bound of the distribution. high long, required Upper bound of the distribution.

shape Shape(tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'int32', 'int64',optional, default='None' DType of the output in case

this can't be inferred. Defaults to int32 if not defined (dtype=None).

name string, optional Name of the resulting symbol.

### **Details**

Example::

randint(low=0, high=5, shape=(2,2)) = [[ 0, 2], [ 3, 1]] Defined in src/operator/random/sample\_op.cc:L194

### Value

out The result mx.symbol

```
mx.symbol.random_uniform
```

random\_uniform:Draw random samples from a uniform distribution.

## **Description**

.. note:: The existing alias "uniform" is deprecated.

### Usage

```
mx.symbol.random_uniform(...)
```

## **Arguments**

low	float, optional, default=0 Lower bound of the distribution.
high	float, optional, default=1 Upper bound of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.

ctx string, optional, default="Context of output, in format [cpulgpulcpu\_pinned](n).

Only used for imperative calls.

dtype 'None', 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

name string, optional Name of the resulting symbol.

### **Details**

Samples are uniformly distributed over the half-open interval \*[low, high)\* (includes \*low\*, but excludes \*high\*).

Example::

uniform(low=0, high=1, shape=(2,2)) = [[ 0.60276335, 0.85794562], [ 0.54488319, 0.84725171]] Defined in src/operator/random/sample\_op.cc:L96

## Value

out The result mx.symbol

```
mx.symbol.ravel_multi_index
```

ravel\_multi\_index:Converts a batch of index arrays into an array of flat indices. The operator follows numpy conventions so a single multi index is given by a column of the input matrix. The leading dimension may be left unspecified by using -1 as placeholder.

# Description

Examples::

```
A = [[3,6,6],[4,5,1]] \text{ ravel}(A, \text{shape}=(7,6)) = [22,41,37] \text{ ravel}(A, \text{shape}=(-1,6)) = [22,41,37]
```

## Usage

```
mx.symbol.ravel_multi_index(...)
```

## **Arguments**

data NDArray-or-Symbol Batch of multi-indices

shape Shape(tuple), optional, default=None Shape of the array into which the multi-

indices apply.

name string, optional Name of the resulting symbol.

### **Details**

Defined in src/operator/tensor/ravel.cc:L42

### Value

450 mx.symbol.reciprocal

mx.symbol.rcbrt

rcbrt:Returns element-wise inverse cube-root value of the input.

## Description

```
.. math:: rcbrt(x) = 1 \land sqrt[3]x
```

## Usage

```
mx.symbol.rcbrt(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

```
Example::
```

```
rcbrt([1,8,-125]) = [1.0, 0.5, -0.2]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L323

## Value

out The result mx.symbol

mx.symbol.reciprocal

reciprocal: Returns the reciprocal of the argument, element-wise.

# Description

Calculates 1/x.

## Usage

```
mx.symbol.reciprocal(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.relu 451

## **Details**

```
Example::
```

```
reciprocal([-2, 1, 3, 1.6, 0.2]) = [-0.5, 1.0, 0.33333334, 0.625, 5.0]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L43

## Value

out The result mx.symbol

mx.symbol.relu

relu:Computes rectified linear activation.

## **Description**

```
.. math:: max(features, 0)
```

### Usage

```
mx.symbol.relu(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

The storage type of "relu" output depends upon the input storage type:

```
- relu(default) = default - relu(row_sparse) = row_sparse - relu(csr) = csr
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L85

# Value

mx.symbol.repeat

repeat:Repeats elements of an array. By default, "repeat" flattens the input array into 1-D and then repeats the elements:: x = [[1, 2], [3, 4]] repeat(x, repeats=2) = [1, 1, 2, 2, 3, 3, 4, 4] The parameter "axis" specifies the axis along which to perform repeat:: repeat(x, repeats=2, axis=1) = [[1, 1, 2, 2, 1, [3, 3, 4, 4]] repeat(x, repeats=2, axis=0) = [[1, 2, 1, [1, 2, 2, 1, [3, 4, 4]]] repeat(x, repeats=2, axis=-1) = [[1, 1, 2, 2, 2, 1, [3, 3, 4, 4, 4]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L744

### Usage

```
mx.symbol.repeat(...)
```

## **Arguments**

data NDArray-or-Symbol Input data array

repeats int, required The number of repetitions for each element.

axis int or None, optional, default='None' The axis along which to repeat values.

The negative numbers are interpreted counting from the backward. By default,

use the flattened input array, and return a flat output array.

name string, optional Name of the resulting symbol.

## Value

out The result mx.symbol

```
mx.symbol.reset_arrays
```

reset\_arrays:Set to zero multiple arrays

## **Description**

Defined in src/operator/contrib/reset\_arrays.cc:L36

## Usage

```
mx.symbol.reset_arrays(...)
```

mx.symbol.Reshape 453

#### **Arguments**

data NDArray-or-Symbol[] Arrays

num.arrays int, required number of input arrays.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.Reshape

Reshape: Reshapes the input array. .. note:: "Reshape" is deprecated, use "reshape" Given an array and a shape, this function returns a copy of the array in the new shape. The shape is a tuple of integers such as (2,3,4). The size of the new shape should be same as the size of the input array. Example:: reshape([1,2,3,4], shape=(2,2)) = [[1,2],[3,4]] Some dimensions of the shape can take special values from the set 0, -1, -2, -3, -4. The significance of each is explained below: - "0" copy this dimension from the input to the output shape. Example:: input shape = (2,3,4), shape = (4,0,2), output shape = (4,3,2) - input shape = (2,3,4), shape = (2,0,0), output shape = (2,3,4) - "-1" infers the dimension of the output shape by using the remainder of the input dimensions keeping the size of the new array same as that of the input array. At most one dimension of shape can be -1. Example:: - input shape = (2,3,4), shape = (6,1,-1), output shape = (6,1,4) - input shape= (2,3,4), shape = (3,-1,8), output shape = (3,1,8) - input shape =(2,3,4), shape=(-1,), output shape=(24,) - "-2" copy all/remainder of the input dimensions to the output shape. Example:: - input shape = (2,3,4), shape = (-2,), output shape = (2,3,4) - input shape = (2,3,4), shape = (2,-2), output shape = (2,3,4) - input shape = (2,3,4), shape = (-2,1,1), output shape = (2,3,4,1,1) - "-3" use the product of two consecutive dimensions of the input shape as the output dimension. Example:: - input shape = (2,3,4), shape = (-3,4), output shape = (6,4) - input shape = (2,3,4,5), shape = (-3,-3), output shape = (6,20)- input shape = (2,3,4), shape = (0,-3), output shape = (2,12) - input shape = (2,3,4), shape = (-3,-2), output shape = (6,4) - "-4" split one dimension of the input into two dimensions passed subsequent to -4 in shape (can contain -1). Example:: - input shape = (2,3,4), shape = (-4,1,2,-2), output shape = (1,2,3,4) - input shape = (2,3,4), shape = (2,-4,-1,3,-2), output shape = (2,1,3,4) If the argument 'reverse' is set to 1, then the special values are inferred from right to left. Example:: - without reverse=1, for input shape = (10,5,4), shape = (-1,0), output shape would be (40,5) - with reverse=1, output shape will be (50,4).

# Description

Defined in src/operator/tensor/matrix\_op.cc:L175

454 mx.symbol.reshape

## Usage

```
mx.symbol.Reshape(...)
```

## **Arguments**

data	NDArray-or-Symbol Input data to reshape.

shape Shape(tuple), optional, default=[] The target shape

reverse boolean, optional, default=0 If true then the special values are inferred from

right to left

target.shape Shape(tuple), optional, default=[] (Deprecated! Use "shape" instead.) Target

new shape. One and only one dim can be 0, in which case it will be inferred

from the rest of dims

keep.highest boolean, optional, default=0 (Deprecated! Use "shape" instead.) Whether keep

the highest dim unchanged. If set to true, then the first dim in target\_shape is

ignored, and always fixed as input

name string, optional Name of the resulting symbol.

### Value

mx.symbol.reshape 455

mx.symbol.reshape

reshape: Reshapes the input array. .. note:: "Reshape" is deprecated, use "reshape" Given an array and a shape, this function returns a copy of the array in the new shape. The shape is a tuple of integers such as (2,3,4). The size of the new shape should be same as the size of the input array. Example:: reshape([1,2,3,4], shape=(2,2)) = [[1,2],[3,4]] Some dimensions of the shape can take special values from the set 0, -1, -2, -3, -4. The significance of each is explained below: - "0" copy this dimension from the input to the output shape. Example:: input shape = (2,3,4), shape = (4,0,2), output shape = (4,3,2) - input shape = (2,3,4), shape = (2,0,0), output shape = (2,3,4) - "-1" infers the dimension of the output shape by using the remainder of the input dimensions keeping the size of the new array same as that of the input array. At most one dimension of shape can be -1. Example:: - input shape = (2,3,4), shape = (6,1,-1), output shape = (6,1,4) - input shape = (2,3,4), shape = (3,-1,8), output shape = (3,1,8) - input shape =(2,3,4), shape=(-1,),  $output\ shape=(24,)$  - "-2"  $copy\ all/remainder$ of the input dimensions to the output shape. Example:: - input shape = (2,3,4), shape = (-2,), output shape = (2,3,4) - input shape = (2,3,4), shape = (2,-2), output shape = (2,3,4) - input shape = (2,3,4), shape = (-2,1,1), output shape = (2,3,4,1,1) - "-3" use the product of two consecutive dimensions of the input shape as the output dimension. Example:: - input shape = (2,3,4), shape = (-3,4), output shape = (6,4) - input shape = (2,3,4,5), shape = (-3,-3), output shape = (6,20)- input shape = (2,3,4), shape = (0,-3), output shape = (2,12) - input shape = (2,3,4), shape = (-3,-2), output shape = (6,4) - "-4" split one dimension of the input into two dimensions passed subsequent to -4 in shape (can contain -1). Example:: - input shape = (2,3,4), shape = (-4,1,2,-2), output shape = (1,2,3,4) - input shape = (2,3,4), shape = (2,-4,-1,3,-2), output shape = (2,1,3,4) If the argument 'reverse' is set to 1, then the special values are inferred from right to left. Example:: - without reverse=1, for input shape = (10,5,4), shape = (-1,0), output shape would be (40,5) - with reverse=1, output shape will be (50,4).

### Description

Defined in src/operator/tensor/matrix\_op.cc:L175

#### Usage

```
mx.symbol.reshape(...)
```

## **Arguments**

data NDArray-or-Symbol Input data to reshape.

shape Shape(tuple), optional, default=[] The target shape

reverse boolean, optional, default=0 If true then the special values are inferred from

right to left

target.shape Shape(tuple), optional, default=[] (Deprecated! Use "shape" instead.) Target

new shape. One and only one dim can be 0, in which case it will be inferred

from the rest of dims

keep.highest boolean, optional, default=0 (Deprecated! Use "shape" instead.) Whether keep

the highest dim unchanged. If set to true, then the first dim in target\_shape is

ignored, and always fixed as input

name string, optional Name of the resulting symbol.

### Value

out The result mx.symbol

mx.symbol.reshape\_like

reshape\_like:Reshape some or all dimensions of 'lhs' to have the same shape as some or all dimensions of 'rhs'.

## **Description**

Returns a \*\*view\*\* of the 'lhs' array with a new shape without altering any data.

## Usage

```
mx.symbol.reshape_like(...)
```

## **Arguments**

1hs NDArray-or-Symbol First input.rhs NDArray-or-Symbol Second input.

1hs.begin int or None, optional, default='None' Defaults to 0. The beginning index along

which the lhs dimensions are to be reshaped. Supports negative indices.

1hs.end int or None, optional, default='None' Defaults to None. The ending index along

which the lhs dimensions are to be used for reshaping. Supports negative in-

dices.

rhs.begin int or None, optional, default='None' Defaults to 0. The beginning index along

which the rhs dimensions are to be used for reshaping. Supports negative in-

dices.

rhs. end int or None, optional, default='None' Defaults to None. The ending index along

which the rhs dimensions are to be used for reshaping. Supports negative in-

dices.

name string, optional Name of the resulting symbol.

mx.symbol.reverse 457

#### **Details**

#### Example::

```
x = [1, 2, 3, 4, 5, 6] y = [[0, -4], [3, 2], [2, 2]] reshape_like(x, y) = [[1, 2], [3, 4], [5, 6]]
```

More precise control over how dimensions are inherited is achieved by specifying \ slices over the 'lhs' and 'rhs' array dimensions. Only the sliced 'lhs' dimensions \ are reshaped to the 'rhs' sliced dimensions, with the non-sliced 'lhs' dimensions staying the same.

### Examples::

- lhs shape = (30,7), rhs shape = (15,2,4), lhs\_begin=0, lhs\_end=1, rhs\_begin=0, rhs\_end=2, output shape = (15,2,7) - lhs shape = (3,5), rhs shape = (1,15,4), lhs\_begin=0, lhs\_end=2, rhs\_begin=1, rhs\_end=2, output shape = (15)

Negative indices are supported, and 'None' can be used for either 'lhs\_end' or 'rhs\_end' to indicate the end of the range.

### Example::

- lhs shape = (30, 12), rhs shape = (4, 2, 2, 3), lhs\_begin=-1, lhs\_end=None, rhs\_begin=1, rhs\_end=None, output shape = (30, 2, 2, 3)

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L512

#### Value

out The result mx.symbol

reverse(x, axis=1) = [[4., 3., 2., 1., 0.], [9., 8., 7., 6., 5.]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L832

## Usage

```
mx.symbol.reverse(...)
```

#### **Arguments**

data NDArray-or-Symbol Input data array

axis Shape(tuple), required The axis which to reverse elements.

name string, optional Name of the resulting symbol.

### Value

mx.symbol.rint

rint:Returns element-wise rounded value to the nearest integer of the input.

## **Description**

```
.. note:: - For input "n.5" "rint" returns "n" while "round" returns "n+1". - For input "-n.5" both "rint" and "round" returns "-n-1".
```

### Usage

```
mx.symbol.rint(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

# Example::

```
rint([-1.5, 1.5, -1.9, 1.9, 2.1]) = [-2., 1., -2., 2., 2.]
```

The storage type of "rint" output depends upon the input storage type:

- rint(default) = default - rint(row\_sparse) = row\_sparse - rint(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L799

### Value

out The result mx.symbol

```
mx.symbol.rmspropalex_update
```

 $rmspropalex\_update: Update\ function\ for\ RMSPropAlex\ optimizer.$ 

## **Description**

'RMSPropAlex' is non-centered version of 'RMSProp'.

## Usage

```
mx.symbol.rmspropalex_update(...)
```

### **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
n	NDArray-or-Symbol n
g	NDArray-or-Symbol g
delta	NDArray-or-Symbol delta
lr	float, required Learning rate
rho	float, optional, default=0.949999988 Decay rate.
momentum	float, optional, default=0.899999976 Decay rate.
epsilon	float, optional, default=9.99999994e-09 A small constant for numerical stability.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
clip.weights	float, optional, default=-1 Clip weights to the range of [-clip_weights, clip_weights] If clip_weights <= 0, weight clipping is turned off. weights = max(min(weights, clip_weights), -clip_weights).
name	string, optional Name of the resulting symbol.

## **Details**

Define :math: 'E[g^2]\_t' is the decaying average over past squared gradient and :math: 'E[g]\_t' is the decaying average over past gradient.

```
.. \ math:: \ E[g^2]_t = \ * \ E[g^2]_{t-1} + (1 - \ ) * \ g_t^2 \ E[g]_t = \ * \ E[g]_{t-1} + (1 - \ ) * \ g_t \ momentum_t = \ * \ momentum_{t-1} - \ F[g^2]_t - \ E[g]_t^2 + \ epsilon \ g_t \ * \ Momentum_t = \ Momentum_t - \ M
```

The update step is

..  $math:: \theta_t = \theta_t + momentum_t$ 

The RMSPropAlex code follows the version in http://arxiv.org/pdf/1308.0850v5.pdf Eq(38) - Eq(45) by Alex Graves, 2013.

Graves suggests the momentum term :math:  $\hline 0.95$ , :math:  $\arrown a^{\prime}$  to be 0.9 and the learning rate :math:  $\end{0.0001}$ .

Defined in src/operator/optimizer\_op.cc:L827

### Value

mx.symbol.rmsprop\_update

rmsprop\_update:Update function for 'RMSProp' optimizer.

## **Description**

'RMSprop' is a variant of stochastic gradient descent where the gradients are divided by a cache which grows with the sum of squares of recent gradients?

### Usage

```
mx.symbol.rmsprop_update(...)
```

## **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
n	NDArray-or-Symbol n
lr	float, required Learning rate
rho	float, optional, default=0.949999988 The decay rate of momentum estimates.
epsilon	float, optional, default=9.99999994e-09 A small constant for numerical stability.
wd	float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight.
rescale.grad	float, optional, default=1 Rescale gradient to grad = rescale_grad*grad.
clip.gradient	float, optional, default=-1 Clip gradient to the range of [-clip_gradient, clip_gradient] If clip_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip_gradient), -clip_gradient).
clip.weights	float, optional, default=-1 Clip weights to the range of [-clip_weights, clip_weights] If clip_weights <= 0, weight clipping is turned off. weights = max(min(weights, clip_weights), -clip_weights).
name	string, optional Name of the resulting symbol.

#### **Details**

'RMSProp' is similar to 'AdaGrad', a popular variant of 'SGD' which adaptively tunes the learning rate of each parameter. 'AdaGrad' lowers the learning rate for each parameter monotonically over the course of training. While this is analytically motivated for convex optimizations, it may not be ideal for non-convex problems. 'RMSProp' deals with this heuristically by allowing the learning rates to rebound as the denominator decays over time.

Define the Root Mean Square (RMS) error criterion of the gradient as :math: 'RMS[g]\_t = \sqrtE[g^2]\_t + \epsilon', where :math: 'g' represents gradient and :math: 'E[g^2]\_t' is the decaying average over past squared gradient.

mx.symbol.RNN 461

```
The :math: E[g^2]_t is given by:
```

```
.. math:: E[g^2]_t = \ E[g^2]_{t-1} + (1-\ g_t^2)
```

The update step is

```
.. math:: \theta_t = \theta_t - \frac{r}{g} g_t
```

The RMSProp code follows the version in http://www.cs.toronto.edu/~tijmen/csc321/slides/lecture\_slides\_lec6.pdf Tieleman & Hinton, 2012.

Hinton suggests the momentum term :math: '\rho' to be 0.9 and the learning rate :math: '\eta' to be 0.001.

Defined in src/operator/optimizer\_op.cc:L788

#### Value

out The result mx.symbol

mx.symbol.RNN	RNN:Applies recurrent layers to input data. Currently, vanilla RNN, LSTM and GRU are implemented, with both multi-layer and bidirectional support
	tional support.

## Description

When the input data is of type float32 and the environment variables MXNET\_CUDA\_ALLOW\_TENSOR\_CORE and MXNET\_CUDA\_TENSOR\_OP\_MATH\_ALLOW\_CONVERSION are set to 1, this operator will try to use pseudo-float16 precision (float32 math with float16 I/O) precision in order to use Tensor Cores on suitable NVIDIA GPUs. This can sometimes give significant speedups.

## Usage

```
mx.symbol.RNN(...)
```

### **Arguments**

data	NDArray-or-Symbol Input data to RNN
parameters	NDArray-or-Symbol Vector of all RNN trainable parameters concatenated
state	NDArray-or-Symbol initial hidden state of the RNN
state.cell	NDArray-or-Symbol initial cell state for LSTM networks (only for LSTM)
sequence.lengt	h
	NDArray-or-Symbol Vector of valid sequence lengths for each element in batch. (Only used if use_sequence_length kwarg is True)
state.size	int (non-negative), required size of the state for each layer
num.layers	int (non-negative), required number of stacked layers
bidirectional	boolean, optional, default=0 whether to use bidirectional recurrent layers
mode	'gru', 'lstm', 'rnn_relu', 'rnn_tanh', required the type of RNN to compute

462 mx.symbol.RNN

p float, optional, default=0 drop rate of the dropout on the outputs of each RNN layer, except the last layer.

state.outputs boolean, optional, default=0 Whether to have the states as symbol outputs. projection.size

int or None, optional, default='None' size of project size

lstm.state.clip.min

double or None, optional, default=None Minimum clip value of LSTM states. This option must be used together with lstm\_state\_clip\_max.

lstm.state.clip.max

double or None, optional, default=None Maximum clip value of LSTM states. This option must be used together with lstm\_state\_clip\_min.

lstm.state.clip.nan

boolean, optional, default=0 Whether to stop NaN from propagating in state by clipping it to min/max. If clipping range is not specified, this option is ignored.

use.sequence.length

boolean, optional, default=0 If set to true, this layer takes in an extra input parameter 'sequence\_length' to specify variable length sequence

name string, optional Name of the resulting symbol.

#### **Details**

\*\*Vanilla RNN\*\*

Applies a single-gate recurrent layer to input X. Two kinds of activation function are supported: ReLU and Tanh.

With ReLU activation function:

```
.. math:: h_t = relu(W_ih * x_t + b_ih + W_hh * h_(t-1) + b_hh)
```

With Tanh activtion function:

```
.. math:: h_t = \tanh(W_ih * x_t + b_ih + W_hh * h_(t-1) + b_hh)
```

Reference paper: Finding structure in time - Elman, 1988. https://crl.ucsd.edu/~elman/Papers/fsit.pdf \*\*LSTM\*\*

Long Short-Term Memory - Hochreiter, 1997. http://www.bioinf.jku.at/publications/older/2604.pdf

With the projection size being set, LSTM could use the projection feature to reduce the parameters size and give some speedups without significant damage to the accuracy.

Long Short-Term Memory Based Recurrent Neural Network Architectures for Large Vocabulary Speech Recognition - Sak et al. 2014. https://arxiv.org/abs/1402.1128

```
.. math:: \beginarrayll i_t = \mathrmsigmoid(W_ii x_t + b_ii + W_ri r_(t-1) + b_ri) \ f_t = \mathrmsigmoid(W_if x_t + b_if + W_rf r_(t-1) + b_rf) \ g_t = \tanh(W_ig x_t + b_ig + W_rc r_(t-1) + b_rg) \ o_t = \mathrmsigmoid(W_io x_t + b_o + W_ro r_(t-1) + b_ro) \ c_t = f_t * c_(t-1) + i_t * g_t \ h_t = o_t * \tanh(c_t) r_t = W_hr h_t \ h_t = o_t * (t-1) + t_t + t_t
```

```
**GRU**
```

Gated Recurrent Unit - Cho et al. 2014. http://arxiv.org/abs/1406.1078

The definition of GRU here is slightly different from paper but compatible with CUDNN.

```
.. math:: \beginarrayll r_t = \mathrmsigmoid(W_ir x_t + b_ir + W_hr h_(t-1) + b_hr) \ z_t = \mathrmsigmoid(W_iz x_t + b_iz + W_hz h_(t-1) + b_hz) \ n_t = \tanh(W_in x_t + b_in + r_t * (W_hn h_(t-1) + b_hn)) \ h_t = (1 - z_t) * n_t + z_t * h_(t-1) \ \endarray
```

Defined in src/operator/rnn.cc:L363

#### Value

out The result mx.symbol

 ${\tt mx.symbol.ROIPooling:} Performs\ region\ of\ interest(ROI)\ pooling\ on\ the\ input\ array.$ 

## **Description**

ROI pooling is a variant of a max pooling layer, in which the output size is fixed and region of interest is a parameter. Its purpose is to perform max pooling on the inputs of non-uniform sizes to obtain fixed-size feature maps. ROI pooling is a neural-net layer mostly used in training a 'Fast R-CNN' network for object detection.

### Usage

```
mx.symbol.ROIPooling(...)
```

### **Arguments**

data	NDArray-or-Symbol The input array to the pooling operator, a 4D Feature maps
rois	NDArray-or-Symbol Bounding box coordinates, a 2D array of [[batch_index, x1, y1, x2, y2]], where (x1, y1) and (x2, y2) are top left and bottom right corners of designated region of interest. 'batch_index' indicates the index of corresponding image in the input array
pooled.size	Shape(tuple), required ROI pooling output shape (h,w)
spatial.scale	float, required Ratio of input feature map height (or w) to raw image height (or w). Equals the reciprocal of total stride in convolutional layers
name	string, optional Name of the resulting symbol.

464 mx.symbol.round

### **Details**

This operator takes a 4D feature map as an input array and region proposals as 'rois', then it pools over sub-regions of input and produces a fixed-sized output array regardless of the ROI size.

To crop the feature map accordingly, you can resize the bounding box coordinates by changing the parameters 'rois' and 'spatial\_scale'.

The cropped feature maps are pooled by standard max pooling operation to a fixed size output indicated by a 'pooled\_size' parameter. batch\_size will change to the number of region bounding boxes after 'ROIPooling'.

The size of each region of interest doesn't have to be perfectly divisible by the number of pooling sections ('pooled\_size').

### Example::

```
x = [[[[0., 1., 2., 3., 4., 5.], [6., 7., 8., 9., 10., 11.], [12., 13., 14., 15., 16., 17.], [18., 19., 20., 21., 22., 23.], [24., 25., 26., 27., 28., 29.], [30., 31., 32., 33., 34., 35.], [36., 37., 38., 39., 40., 41.], [42., 43., 44., 45., 46., 47.]]]]
```

// region of interest i.e. bounding box coordinates. y = [[0,0,0,4,4]]

// returns array of shape (2,2) according to the given roi with max pooling. ROIPooling(x, y, (2,2), 1.0) = [[[[14., 16.], [26., 28.]]]]

// region of interest is changed due to the change in 'spacial\_scale' parameter. ROIPooling(x, y, (2,2), (2,7) = [[[[ 7., 9.], [ 19., 21.]]]]

Defined in src/operator/roi\_pooling.cc:L225

### Value

out The result mx.symbol

mx.symbol.round

round:Returns element-wise rounded value to the nearest integer of the input.

# **Description**

Example::

## Usage

```
mx.symbol.round(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

mx.symbol.rsqrt 465

## **Details**

```
round([-1.5, 1.5, -1.9, 1.9, 2.1]) = [-2., 2., -2., 2., 2.]
```

The storage type of "round" output depends upon the input storage type:

- round(default) = default - round(row\_sparse) = row\_sparse - round(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L778

#### Value

out The result mx.symbol

mx.symbol.rsqrt

rsqrt:Returns element-wise inverse square-root value of the input.

## **Description**

```
.. math:: rsqrt(x) = 1 \land sqrtx
```

## Usage

```
mx.symbol.rsqrt(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

### **Details**

Example::

```
rsqrt([4,9,16]) = [0.5, 0.33333334, 0.25]
```

The storage type of "rsqrt" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L221

# Value

```
mx.symbol.sample_exponential
```

sample\_exponential:Concurrent sampling from multiple exponential distributions with parameters lambda (rate).

# Description

The parameters of the distributions are provided as an input array. Let \*[s]\* be the shape of the input array, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## Usage

```
mx.symbol.sample_exponential(...)
```

## Arguments

lam	NDArray-or-Symbol Lambda (rate) parameters of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

#### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input array, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input value at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input array.

## Examples::

```
lam = [1.0, 8.5]
```

// Draw a single sample for each distribution sample\_exponential(lam) = [ 0.51837951, 0.09994757]

// Draw a vector containing two samples for each distribution sample\_exponential(lam, shape=(2)) = [[ 0.51837951, 0.19866663], [ 0.09994757, 0.50447971]]

Defined in src/operator/random/multisample\_op.cc:L284

### Value

```
mx.symbol.sample_gamma
```

sample\_gamma: Concurrent sampling from multiple gamma distributions with parameters \*alpha\* (shape) and \*beta\* (scale).

## Description

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## Usage

```
mx.symbol.sample_gamma(...)
```

### **Arguments**

alpha	NDArray-or-Symbol Alpha (shape) parameters of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
beta	NDArray-or-Symbol Beta (scale) parameters of the distributions.
name	string, optional Name of the resulting symbol.

## **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

### Examples::

```
alpha = [ 0.0, 2.5 ] beta = [ 1.0, 0.7 ]

// Draw a single sample for each distribution sample_gamma(alpha, beta) = [ 0. , 2.25797319]

// Draw a vector containing two samples for each distribution sample_gamma(alpha, beta, shape=(2))

= [[ 0. , 0. ], [ 2.25797319, 1.70734084]]

Defined in src/operator/random/multisample op.cc:L282
```

## Value

```
mx.symbol.sample_generalized_negative_binomial
```

sample\_generalized\_negative\_binomial:Concurrent sampling from multiple generalized negative binomial distributions with parameters \*mu\* (mean) and \*alpha\* (dispersion).

### **Description**

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## Usage

```
mx.symbol.sample_generalized_negative_binomial(...)
```

### Arguments

mu	NDArray-or-Symbol Means of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
alpha	NDArray-or-Symbol Alpha (dispersion) parameters of the distributions.
name	string, optional Name of the resulting symbol.

### **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

Samples will always be returned as a floating point data type.

## Examples::

```
mu = [2.0, 2.5] alpha = [1.0, 0.1]
```

// Draw a single sample for each distribution sample\_generalized\_negative\_binomial(mu, alpha) = [0., 3.]

// Draw a vector containing two samples for each distribution sample\_generalized\_negative\_binomial(mu, alpha, shape=(2)) = [[ 0., 3.], [ 3., 1.]]

Defined in src/operator/random/multisample\_op.cc:L293

### Value

```
mx.symbol.sample_multinomial
```

sample\_multinomial:Concurrent sampling from multiple multinomial distributions.

# **Description**

\*data\* is an \*n\* dimensional array whose last dimension has length \*k\*, where \*k\* is the number of possible outcomes of each multinomial distribution. This operator will draw \*shape\* samples from each distribution. If shape is empty one sample will be drawn from each distribution.

## Usage

```
mx.symbol.sample_multinomial(...)
```

### **Arguments**

data	NDArray-or-Symbol Distribution probabilities. Must sum to one on the last axis.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
get.prob	boolean, optional, default=0 Whether to also return the log probability of sampled result. This is usually used for differentiating through stochastic variables, e.g. in reinforcement learning.
dtype	'float16', 'float32', 'float64', 'int32', 'uint8', optional, default='int32' DType of the output in case this can't be inferred.
name	string, optional Name of the resulting symbol.

## Details

If \*get\_prob\* is true, a second array containing log likelihood of the drawn samples will also be returned. This is usually used for reinforcement learning where you can provide reward as head gradient for this array to estimate gradient.

Note that the input distribution must be normalized, i.e. \*data\* must sum to 1 along its last axis.

# Examples::

```
probs = [[0, 0.1, 0.2, 0.3, 0.4], [0.4, 0.3, 0.2, 0.1, 0]]

// Draw a single sample for each distribution sample_multinomial(probs) = [3, 0]

// Draw a vector containing two samples for each distribution sample_multinomial(probs, shape=(2)) = [[4, 2], [0, 0]]

// requests log likelihood sample_multinomial(probs, get_prob=True) = [2, 1], [0.2, 0.3]
```

### Value

```
mx.symbol.sample_negative_binomial
```

sample\_negative\_binomial:Concurrent sampling from multiple negative binomial distributions with parameters \*k\* (failure limit) and \*p\* (failure probability).

## **Description**

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## Usage

```
mx.symbol.sample_negative_binomial(...)
```

### **Arguments**

k	NDArray-or-Symbol Limits of unsuccessful experiments.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
p	NDArray-or-Symbol Failure probabilities in each experiment.

string, optional Name of the resulting symbol.

## **Details**

name

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

Samples will always be returned as a floating point data type.

## Examples::

```
k = [20, 49] p = [0.4, 0.77]

// Draw a single sample for each distribution sample_negative_binomial(k, p) = [15., 16.]

// Draw a vector containing two samples for each distribution sample_negative_binomial(k, p, shape=(2)) = [[15., 50.], [16., 12.]]
```

Defined in src/operator/random/multisample\_op.cc:L289

### Value

```
mx.symbol.sample_normal
```

sample\_normal:Concurrent sampling from multiple normal distributions with parameters \*mu\* (mean) and \*sigma\* (standard deviation).

# Description

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

# Usage

```
mx.symbol.sample_normal(...)
```

## **Arguments**

mu	NDArray-or-Symbol Means of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
sigma	NDArray-or-Symbol Standard deviations of the distributions.
name	string, optional Name of the resulting symbol.

# **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

## Examples::

```
mu = [ 0.0, 2.5 ] sigma = [ 1.0, 3.7 ]

// Draw a single sample for each distribution sample_normal(mu, sigma) = [-0.56410581, 0.95934606]

// Draw a vector containing two samples for each distribution sample_normal(mu, sigma, shape=(2))

= [[-0.56410581, 0.2928229 ], [ 0.95934606, 4.48287058]]

Defined in src/operator/random/multisample op.cc:L279
```

## Value

```
mx.symbol.sample_poisson
```

sample\_poisson:Concurrent sampling from multiple Poisson distributions with parameters lambda (rate).

## Description

The parameters of the distributions are provided as an input array. Let \*[s]\* be the shape of the input array, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## Usage

```
mx.symbol.sample_poisson(...)
```

## **Arguments**

lam	NDArray-or-Symbol Lambda (rate) parameters of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random dis-

tribution.

dtype 'None', 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to float32 if not defined (dtype=None).

name string, optional Name of the resulting symbol.

## **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input array, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input value at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input array.

Samples will always be returned as a floating point data type.

# Examples::

```
lam = [1.0, 8.5]
```

// Draw a single sample for each distribution sample\_poisson(lam) = [0., 13.]

// Draw a vector containing two samples for each distribution sample\_poisson(lam, shape=(2)) = [[ 0., 4.], [ 13., 8.]]

Defined in src/operator/random/multisample\_op.cc:L286

## Value

```
mx.symbol.sample_uniform
```

sample\_uniform: Concurrent sampling from multiple uniform distributions on the intervals given by \*[low,high)\*.

# Description

The parameters of the distributions are provided as input arrays. Let \*[s]\* be the shape of the input arrays, \*n\* be the dimension of \*[s]\*, \*[t]\* be the shape specified as the parameter of the operator, and \*m\* be the dimension of \*[t]\*. Then the output will be a \*(n+m)\*-dimensional array with shape \*[s]x[t]\*.

## Usage

```
mx.symbol.sample_uniform(...)
```

## **Arguments**

low	NDArray-or-Symbol Lower bounds of the distributions.
shape	Shape(tuple), optional, default=[] Shape to be sampled from each random distribution.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
high	NDArray-or-Symbol Upper bounds of the distributions.
name	string, optional Name of the resulting symbol.

## **Details**

For any valid \*n\*-dimensional index \*i\* with respect to the input arrays, \*output[i]\* will be an \*m\*-dimensional array that holds randomly drawn samples from the distribution which is parameterized by the input values at index \*i\*. If the shape parameter of the operator is not set, then one sample will be drawn per distribution and the output array has the same shape as the input arrays.

# Examples::

```
low = [ 0.0, 2.5 ] high = [ 1.0, 3.7 ]

// Draw a single sample for each distribution sample_uniform(low, high) = [ 0.40451524, 3.18687344]

// Draw a vector containing two samples for each distribution sample_uniform(low, high, shape=(2))

= [[ 0.40451524, 0.18017688], [ 3.18687344, 3.68352246]]

Defined in src/operator/random/multisample op.cc:L277
```

## Value

mx.symbol.save

Save an mx.symbol object

# Description

Save an mx.symbol object

## Usage

```
mx.symbol.save(symbol, filename)
```

# Arguments

symbol the mx.symbol object

filename (including the path)

## **Examples**

```
data = mx.symbol.Variable('data')
mx.symbol.save(data, 'temp.symbol')
data2 = mx.symbol.load('temp.symbol')
```

mx.symbol.scatter\_nd scatter\_nd:Scatters data into a new tensor according to indices.

# Description

Given 'data' with shape ' $(Y_0, ..., Y_{K-1}, X_M, ..., X_{N-1})$ ' and indices with shape ' $(M, Y_0, ..., Y_{K-1})$ ', the output will have shape ' $(X_0, X_1, ..., X_{N-1})$ ', where ' $M \le N$ '. If 'M == N', data shape should simply be ' $(Y_0, ..., Y_{K-1})$ '.

## Usage

```
mx.symbol.scatter_nd(...)
```

# Arguments

data NDArray-or-Symbol data indices NDArray-or-Symbol indices

shape Shape(tuple), required Shape of output.

name string, optional Name of the resulting symbol.

### **Details**

The elements in output is defined as follows::

```
output[indices[0, y_0, ..., y_K-1], ..., indices[M-1, y_0, ..., y_K-1], x_M, ..., x_N-1] = data[y_0, ..., y_K-1, x_M, ..., x_N-1]
```

all other entries in output are 0.

.. warning::

If the indices have duplicates, the result will be non-deterministic and the gradient of 'scatter\_nd' will not be correct!!

### Examples::

```
data = [2, 3, 0] indices = [[1, 1, 0], [0, 1, 0]] shape = (2, 2) scatter_nd(data, indices, shape) = [[0, 0], [2, 3]]
```

 $data = [[[1, 2], [3, 4]], [[5, 6], [7, 8]]] indices = [[0, 1], [1, 1]] shape = (2, 2, 2, 2) scatter_nd(data, indices, shape) = [[[[0, 0], [0, 0]],$ 

[[1, 2], [3, 4]]],

[[[0, 0], [0, 0]],

[[5, 6], [7, 8]]]]

### Value

out The result mx.symbol

mx.symbol.SequenceLast

SequenceLast:Takes the last element of a sequence.

# **Description**

This function takes an n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] and returns a (n-1)-dimensional array of the form [batch\_size, other\_feature\_dims].

## Usage

```
mx.symbol.SequenceLast(...)
```

## **Arguments**

data

NDArray-or-Symbol n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] where n>2

sequence.length

NDArray-or-Symbol vector of sequence lengths of the form [batch\_size]

use.sequence.length

boolean, optional, default=0 If set to true, this layer takes in an extra input parameter 'sequence\_length' to specify variable length sequence

axis int, optional, default='0' The sequence axis. Only values of 0 and 1 are currently

supported.

name string, optional Name of the resulting symbol.

### **Details**

Parameter 'sequence\_length' is used to handle variable-length sequences. 'sequence\_length' should be an input array of positive ints of dimension [batch\_size]. To use this parameter, set 'use\_sequence\_length' to 'True', otherwise each example in the batch is assumed to have the max sequence length.

.. note:: Alternatively, you can also use 'take' operator.

## Example::

```
x = [[[ 1., 2., 3.], [ 4., 5., 6.], [ 7., 8., 9.]],

[[ 10., 11., 12.], [ 13., 14., 15.], [ 16., 17., 18.]],

[[ 19., 20., 21.], [ 22., 23., 24.], [ 25., 26., 27.]]]

// returns last sequence when sequence_length parameter is not used SequenceLast(x) = [[ 19., 20., 21.], [ 22., 23., 24.], [ 25., 26., 27.]]

// sequence_length is used SequenceLast(x, sequence_length=[1,1,1], use_sequence_length=True)

= [[ 1., 2., 3.], [ 4., 5., 6.], [ 7., 8., 9.]]

// sequence_length is used SequenceLast(x, sequence_length=[1,2,3], use_sequence_length=True)

= [[ 1., 2., 3.], [ 13., 14., 15.], [ 25., 26., 27.]]

Defined in src/operator/sequence_last.cc:L106
```

### Value

out The result mx.symbol

```
mx.symbol.SequenceMask
```

SequenceMask:Sets all elements outside the sequence to a constant value.

# Description

This function takes an n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] and returns an array of the same shape.

```
mx.symbol.SequenceMask(...)
```

## **Arguments**

data NDArray-or-Symbol n-dimensional input array of the form [max\_sequence\_length,

batch\_size, other\_feature\_dims] where n>2

sequence.length

NDArray-or-Symbol vector of sequence lengths of the form [batch\_size]

use.sequence.length

boolean, optional, default=0 If set to true, this layer takes in an extra input pa-

rameter 'sequence\_length' to specify variable length sequence

value float, optional, default=0 The value to be used as a mask.

axis int, optional, default='0' The sequence axis. Only values of 0 and 1 are currently

supported.

name string, optional Name of the resulting symbol.

### **Details**

Parameter 'sequence\_length' is used to handle variable-length sequences. 'sequence\_length' should be an input array of positive ints of dimension [batch\_size]. To use this parameter, set 'use\_sequence\_length' to 'True', otherwise each example in the batch is assumed to have the max sequence length and this operator works as the 'identity' operator.

## Example::

```
x = [[[1., 2., 3.], [4., 5., 6.]],
```

[[7., 8., 9.], [10., 11., 12.]],

[[ 13., 14., 15.], [ 16., 17., 18.]]]

// Batch 1 B1 = [[ 1., 2., 3.], [ 7., 8., 9.], [ 13., 14., 15.]]

// Batch 2 B2 = [[4., 5., 6.], [10., 11., 12.], [16., 17., 18.]]

// works as identity operator when sequence\_length parameter is not used SequenceMask(x) = [[[1, 2, 3, 3, [4, 5, 6, 6]]]

[[ 7., 8., 9.], [ 10., 11., 12.]],

[[ 13., 14., 15.], [ 16., 17., 18.]]]

// sequence\_length [1,1] means 1 of each batch will be kept // and other rows are masked with default mask value = 0 SequenceMask(x, sequence\_length=[1,1], use\_sequence\_length=True) = [[[ 1., 2., 3.], [ 4., 5., 6.]],

[[0., 0., 0.], [0., 0., 0.]],

[[0., 0., 0.], [0., 0., 0.]]]

// sequence\_length [2,3] means 2 of batch B1 and 3 of batch B2 will be kept // and other rows are masked with value = 1 SequenceMask(x, sequence\_length=[2,3], use\_sequence\_length=True, value=1) = [[[1., 2., 3.], [4., 5., 6.]],

[[7., 8., 9.], [10., 11., 12.]],

[[ 1., 1., 1.], [ 16., 17., 18.]]]

Defined in src/operator/sequence\_mask.cc:L186

### Value

out The result mx.symbol

```
mx.symbol.SequenceReverse
```

SequenceReverse:Reverses the elements of each sequence.

## **Description**

This function takes an n-dimensional input array of the form [max\_sequence\_length, batch\_size, other\_feature\_dims] and returns an array of the same shape.

## Usage

```
mx.symbol.SequenceReverse(...)
```

## **Arguments**

data NDArray-or-Symbol n-dimensional input array of the form [max\_sequence\_length,

batch\_size, other dims] where n>2

sequence.length

NDArray-or-Symbol vector of sequence lengths of the form [batch\_size]

 $\verb"use.sequence.length"$ 

boolean, optional, default=0 If set to true, this layer takes in an extra input pa-

rameter 'sequence\_length' to specify variable length sequence

axis int, optional, default='0' The sequence axis. Only 0 is currently supported.

name string, optional Name of the resulting symbol.

### **Details**

Parameter 'sequence\_length' is used to handle variable-length sequences. 'sequence\_length' should be an input array of positive ints of dimension [batch\_size]. To use this parameter, set 'use\_sequence\_length' to 'True', otherwise each example in the batch is assumed to have the max sequence length.

### Example::

```
x = [[[ 1., 2., 3.], [ 4., 5., 6.]],

[[ 7., 8., 9.], [ 10., 11., 12.]],

[[ 13., 14., 15.], [ 16., 17., 18.]]]

// Batch 1 B1 = [[ 1., 2., 3.], [ 7., 8., 9.], [ 13., 14., 15.]]

// Batch 2 B2 = [[ 4., 5., 6.], [ 10., 11., 12.], [ 16., 17., 18.]]

// returns reverse sequence when sequence_length parameter is not used SequenceReverse(x) = [[[ 13., 14., 15.], [ 16., 17., 18.]],

[[ 7., 8., 9.], [ 10., 11., 12.]],

[[ 1., 2., 3.], [ 4., 5., 6.]]]
```

```
// sequence_length [2,2] means 2 rows of // both batch B1 and B2 will be reversed. SequenceReverse(x, sequence_length=[2,2], use_sequence_length=True) = [[[7., 8., 9.], [10., 11., 12.]], [[1., 2., 3.], [4., 5., 6.]],
```

// sequence\_length [2,3] means 2 of batch B2 and 3 of batch B3 // will be reversed. SequenceReverse(x, sequence\_length=[2,3], use\_sequence\_length=True) = [[[7., 8., 9.], [16., 17., 18.]],

```
[[ 1., 2., 3.], [ 10., 11., 12.]],
[[ 13., 14, 15.], [ 4., 5., 6.]]]
```

Defined in src/operator/sequence\_reverse.cc:L122

## Value

out The result mx.symbol

```
mx.symbol.sgd_mom_update
```

sgd\_mom\_update:Momentum update function for Stochastic Gradient Descent (SGD) optimizer.

# **Description**

Momentum update has better convergence rates on neural networks. Mathematically it looks like below:

# Usage

```
mx.symbol.sgd_mom_update(...)
```

# **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient mom NDArray-or-Symbol Momentum lr float, required Learning rate

momentum float, optional, default=0 The decay rate of momentum estimates at each epoch. wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row\_sparse and both weight and momentum have the same stype

name string, optional Name of the resulting symbol.

### **Details**

```
.. math:
```

 $v_1 = \alpha y_0 * \quad J(W_0) \ v_t = \gamma v_1 - \alpha y_0 * \quad J(W_{t-1}) \ W_t = W_{t-1} + v_t$  It updates the weights using::

v = momentum \* v - learning\_rate \* gradient weight += v

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

However, if grad's storage type is "row\_sparse", "lazy\_update" is True and weight's storage type is the same as momentum's storage type, only the row slices whose indices appear in grad.indices are updated (for both weight and momentum)::

for row in gradient.indices:  $v[row] = momentum[row] * v[row] - learning_rate * gradient[row] weight[row] += v[row]$ 

Defined in src/operator/optimizer\_op.cc:L556

#### Value

out The result mx.symbol

# **Description**

It updates the weights using::

### Usage

```
mx.symbol.sgd_update(...)
```

# **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient lr float, required Learning rate

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

lazy.update boolean, optional, default=1 If true, lazy updates are applied if gradient's stype

is row sparse.

name string, optional Name of the resulting symbol.

## **Details**

```
weight = weight - learning_rate * (gradient + wd * weight)
```

However, if gradient is of "row\_sparse" storage type and "lazy\_update" is True, only the row slices whose indices appear in grad.indices are updated::

for row in gradient.indices: weight[row] = weight[row] - learning\_rate \* (gradient[row] + wd \* weight[row])

Defined in src/operator/optimizer\_op.cc:L515

## Value

out The result mx.symbol

mx.symbol.shape\_array shape\_array:Returns a 1D int64 array containing the shape of data.

# **Description**

Example::

# Usage

```
mx.symbol.shape_array(...)
```

## **Arguments**

data NDArray-or-Symbol Input Array.

name string, optional Name of the resulting symbol.

# **Details**

```
shape_array([[1,2,3,4], [5,6,7,8]]) = [2,4]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L574

### Value

482 mx.symbol.sigmoid

mx.symbol.shuffle

shuffle:Randomly shuffle the elements.

## **Description**

This shuffles the array along the first axis. The order of the elements in each subarray does not change. For example, if a 2D array is given, the order of the rows randomly changes, but the order of the elements in each row does not change.

# Usage

```
mx.symbol.shuffle(...)
```

# **Arguments**

data NDArray-or-Symbol Data to be shuffled.
name string, optional Name of the resulting symbol.

### Value

out The result mx.symbol

mx.symbol.sigmoid

sigmoid:Computes sigmoid of x element-wise.

# **Description**

```
.. math:: y = 1 / (1 + \exp(-x))
```

# Usage

```
mx.symbol.sigmoid(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

The storage type of "sigmoid" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L119

# Value

mx.symbol.sign 483

mx.symbol.sign

sign:Returns element-wise sign of the input.

# Description

Example::

# Usage

```
mx.symbol.sign(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

```
sign([-2, 0, 3]) = [-1, 0, 1]
```

The storage type of "sign" output depends upon the input storage type:

- sign(default) = default - sign(row\_sparse) = row\_sparse - sign(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L759

# Value

out The result mx.symbol

```
mx.symbol.signsgd_update
```

signsgd\_update:Update function for SignSGD optimizer.

# Description

.. math::

```
mx.symbol.signsgd_update(...)
```

## **Arguments**

weight	NDArray-or-Symbol Weight
grad	NDArray-or-Symbol Gradient
lr	float, required Learning rate

wd float, optional, default=0 Weight decay augments the objective function with a

regularization term that penalizes large weights. The penalty scales with the

square of the magnitude of each weight.

rescale.grad float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad.

clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient]

If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad,

clip\_gradient), -clip\_gradient).

name string, optional Name of the resulting symbol.

## **Details**

```
\begin{split} g_t &= \Lambda J(W_{t-1}) \ W_t = W_{t-1} - \epsilon a_t \ \lambda g_t \end{split} It updates the weights using:: weight = weight - learning_rate * sign(gradient) 
... note:: - sparse ndarray not supported for this optimizer yet.
```

Defined in src/operator/optimizer\_op.cc:L63

## Value

out The result mx.symbol

# Description

.. math::

```
mx.symbol.signum_update(...)
```

mx.symbol.sin 485

## **Arguments**

weight NDArray-or-Symbol Weight grad NDArray-or-Symbol Gradient NDArray-or-Symbol Momentum mom float, required Learning rate lr float, optional, default=0 The decay rate of momentum estimates at each epoch. momentum wd float, optional, default=0 Weight decay augments the objective function with a regularization term that penalizes large weights. The penalty scales with the square of the magnitude of each weight. float, optional, default=1 Rescale gradient to grad = rescale\_grad\*grad. rescale.grad clip.gradient float, optional, default=-1 Clip gradient to the range of [-clip\_gradient, clip\_gradient] If clip\_gradient <= 0, gradient clipping is turned off. grad = max(min(grad, clip\_gradient), -clip\_gradient). wd.1h float, optional, default=0 The amount of weight decay that does not go into gradient/momentum calculationsotherwise do weight decay algorithmically only.

name string, optional Name of the resulting symbol.

### **Details**

```
g_t = \Lambda J(W_{t-1}) + (1 - \beta g_t) + (1 - \beta
```

It updates the weights using:: state = momentum \* state + (1-momentum) \* gradient weight = weight - learning\_rate \* sign(state)

Where the parameter "momentum" is the decay rate of momentum estimates at each epoch.

.. note:: - sparse ndarray not supported for this optimizer yet.

Defined in src/operator/optimizer\_op.cc:L92

# Value

out The result mx.symbol

mx.symbol.sin sin: Computes the element-wise sine of the input array.

# **Description**

The input should be in radians (:math: '2\pi' rad equals 360 degrees).

```
mx.symbol.sin(...)
```

486 mx.symbol.sinh

## Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

```
.. math:: \sin([0, \pi/4, \pi/2]) = [0, 0.707, 1]
```

The storage type of "sin" output depends upon the input storage type:

- sin(default) = default - sin(row\_sparse) = row\_sparse - sin(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L47

# Value

out The result mx.symbol

mx.symbol.sinh

sinh: Returns the hyperbolic sine of the input array, computed elementwise.

# **Description**

```
.. math:: sinh(x) = 0.5 \times (exp(x) - exp(-x))
```

## Usage

```
mx.symbol.sinh(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

The storage type of "sinh" output depends upon the input storage type:

- sinh(default) = default - sinh(row\_sparse) = row\_sparse - sinh(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L371

# Value

mx.symbol.size\_array 487

mx.symbol.size\_array size\_array:Returns a 1D int64 array containing the size of data.

# Description

Example::

# Usage

```
mx.symbol.size_array(...)
```

# Arguments

data NDArray-or-Symbol Input Array.

name string, optional Name of the resulting symbol.

## **Details**

```
size\_array([[1,2,3,4], [5,6,7,8]]) = [8]
```

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L625

# Value

488 mx.symbol.slice

mx.symbol.slice

slice:Slices a region of the array. .. note:: "crop" is deprecated. Use "slice" instead. This function returns a sliced array between the indices given by 'begin' and 'end' with the corresponding 'step'. For an input array of "shape= $(d_0, d_1, ..., d_{n-1})$ ", slice operation with "begin=(b\_0, b\_1...b\_m-1)", "end=(e\_0, e\_1, ..., e\_m-1)", and "step=(s 0, s 1, ..., s m-1)", where  $m \le n$ , results in an array with the shape "(|e\_0-b\_0|/|s\_0|, ..., |e\_m-1-b\_m-1|/|s\_m-1|, d\_m, ..., d\_n-1)". The resulting array's \*k\*-th dimension contains elements from the \*k\*-th dimension of the input array starting from index "b\_k" (inclusive) with step "s\_k" until reaching "e\_k" (exclusive). If the \*k\*-th elements are 'None' in the sequence of 'begin', 'end', and 'step', the following rule will be used to set default values. If 's\_k' is 'None', set  $s_k=1$ . If  $s_k>0$ , set  $b_k=0$ ,  $e_k=d_k$ ; else, set  $b_k=d_k-1$ , 'e\_k=-1'. The storage type of "slice" output depends on storage types of inputs - slice(csr) = csr - otherwise, "slice" generates output with default storage .. note:: When input data storage type is csr, it only supports step=(), or step=(None,), or step=(1,) to generate a csr output. For other step parameter values, it falls back to slicing a dense tensor. Example:: x = [[1., 2., 3., 4.], [5., 6., 7., 8.], [9., 10., 11.]12.]] slice(x, begin=(0,1), end=(2,4)) = [[2., 3., 4.], [6., 7., 8.]]slice(x, begin=(None, 0), end=(None, 3), step=(-1, 2)) = [[9., 11.],[5., 7.], [1., 3.]]

# Description

Defined in src/operator/tensor/matrix\_op.cc:L482

# Usage

```
mx.symbol.slice(...)
```

## **Arguments**

data	NDArray-or-Symbol Source input
begin	Shape(tuple), required starting indices for the slice operation, supports negative indices.
end	Shape(tuple), required ending indices for the slice operation, supports negative indices.
step	Shape(tuple), optional, default=[] step for the slice operation, supports negative values.
name	string, optional Name of the resulting symbol.

#### Value

```
mx.symbol.SliceChannel
```

SliceChannel:Splits an array along a particular axis into multiple subarrays.

## **Description**

.. note:: "SliceChannel" is deprecated. Use "split" instead.

## Usage

```
mx.symbol.SliceChannel(...)
```

### **Arguments**

data NDArray-or-Symbol The input

num.outputs int, required Number of splits. Note that this should evenly divide the length of

the 'axis'.

axis int, optional, default='1' Axis along which to split.

squeeze.axis boolean, optional, default=0 If true, Removes the axis with length 1 from the

shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "true" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis'

can be set to "true" only if "input.shape[axis] == num\_outputs".

name string, optional Name of the resulting symbol.

### **Details**

\*\*Note\*\* that 'num\_outputs' should evenly divide the length of the axis along which to split the array.

### Example::

```
 \begin{aligned} x &= & [[[\ 1.]\ [\ 2.]]\ [[\ 3.]\ [\ 4.]]\ [[\ 5.]\ [\ 6.]]] \ x.shape = (3,\,2,\,1) \\ y &= & \text{split}(x,\,axis=1,\,num\_outputs=2)\,\,\text{//}\ a \ \text{list of 2 arrays with shape } (3,\,1,\,1)\,\,y = [[[\ 1.]]\ [[\ 3.]]\ [[\ 5.]]] \\ [[[\ 2.]]\ [[\ 4.]]\ [[\ 6.]]] \\ y[0].shape &= (3,\,1,\,1) \\ z &= & \text{split}(x,\,axis=0,\,num\_outputs=3)\,\,\text{//}\ a \ \text{list of 3 arrays with shape } (1,\,2,\,1)\,\,z = [[[\ 1.]\ [\ 2.]]] \\ [[[\ 3.]\ [\ 4.]]] \\ [[[\ 5.]\ [\ 6.]]] \\ z[0].shape &= (1,\,2,\,1) \end{aligned}
```

'squeeze\_axis=1' removes the axis with length 1 from the shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "1" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis' can be set to true only if "input.shape[axis] == num\_outputs".

## Example::

490 mx.symbol.slice\_axis

## Value

out The result mx.symbol

```
mx.symbol.slice_axis slice_axis:Slices along a given axis. Returns an array slice along a given 'axis' starting from the 'begin' index to the 'end' index. Examples:: x = [[\ 1.,\ 2.,\ 3.,\ 4.],\ [\ 5.,\ 6.,\ 7.,\ 8.],\ [\ 9.,\ 10.,\ 11.,\ 12.]] slice_axis(x,\ axis=0,\ begin=1,\ end=3) = [[\ 5.,\ 6.,\ 7.,\ 8.],\ [\ 9.,\ 10.,\ 11.]] slice_axis(x,\ axis=1,\ begin=-3,\ end=-1) = [[\ 2.,\ 3.],\ [\ 6.,\ 7.],\ [\ 10.,\ 11.]]
```

# Description

Defined in src/operator/tensor/matrix\_op.cc:L571

# Usage

```
mx.symbol.slice_axis(...)
```

# **Arguments**

data	NDArray-or-Symbol Source input
axis	int, required Axis along which to be sliced, supports negative indexes.
begin	int, required The beginning index along the axis to be sliced, supports negative indexes.
end	int or None, required The ending index along the axis to be sliced, supports negative indexes.
name	string, optional Name of the resulting symbol.

# Value

mx.symbol.slice\_like 491

mx.symbol.slice\_like

slice\_like:Slices a region of the array like the shape of another array. This function is similar to "slice", however, the 'begin' are always '0's and 'end' of specific axes are inferred from the second input 'shape\_like'. Given the second 'shape\_like' input of "shape= $(d_0, d_0, d_0)$ ". d 1, ..., d n-1)", a "slice like" operator with default empty 'axes', it performs the following operation: "out = slice(input, begin=(0, 0, 0, 0))" ..., 0),  $end=(d_0, d_1, ..., d_{n-1}))$ ". When 'axes' is not empty, it is used to speficy which axes are being sliced. Given a 4-d input data, "slice\_like" operator with "axes=(0, 2, -1)" will perform the following operation: " out = slice(input, begin=(0, 0, 0, 0), end= $(d_0, 0, 0)$ ), end= $(d_0, 0, 0)$ None,  $d_2$ ,  $d_3$ )". Note that it is allowed to have first and second input with different dimensions, however, you have to make sure the 'axes' are specified and not exceeding the dimension limits. For example, given 'input\_1' with "shape=(2,3,4,5)" and 'input\_2' with "shape=(1,2,3)", it is not allowed to use: "out = slice like(a, b)" because ndim of 'input\_1' is 4, and ndim of 'input\_2' is 3. The following is allowed in this situation: "out =  $slice_like(a, b, axes=(0, 2))$ " Example:: x = [[1., 2., 3., 4.], [5., 6., 7., 8.], [9., 10., 11., 12.]] y =[[0., 0., 0.], [0., 0., 0.]]  $slice\_like(x, y) = [[1., 2., 3.], [5., 6., 7.]]$  $slice\_like(x, y, axes=(0, 1)) = [[1., 2., 3.] [5., 6., 7.]] slice\_like(x, y, axes=(0, 1)) = [[1., 2., 3.] [5., 4.]] slice\_like(x, y, axes=(0, 1)) = [[1., 2., 3.]] slice\_like$ axes=(0)) = [[ 1., 2., 3., 4.] [ 5., 6., 7., 8.]]  $slice_like(x, y, axes=(-1))$ = [[1., 2., 3.] [5., 6., 7.] [9., 10., 11.]]

### Description

Defined in src/operator/tensor/matrix\_op.cc:L625

## Usage

```
mx.symbol.slice_like(...)
```

## **Arguments**

data NDArray-or-Symbol Source input shape.like NDArray-or-Symbol Shape like input

axes Shape(tuple), optional, default=[] List of axes on which input data will be sliced

according to the corresponding size of the second input. By default will slice on

all axes. Negative axes are supported.

name string, optional Name of the resulting symbol.

### Value

492 mx.symbol.Softmax

mx.symbol.smooth\_11

smooth\_l1:Calculate Smooth L1 Loss(lhs, scalar) by summing

# Description

.. math::

# Usage

```
mx.symbol.smooth_l1(...)
```

# Arguments

data NDArray-or-Symbol source input

scalar float scalar input

name string, optional Name of the resulting symbol.

### **Details**

 $f(x) = \text{logincases (\sigma x)^2/2,\& \textif x < 1/sigma^2\ |x|-0.5/\sigma^2,\& \textotherwise \endoxes | \cdots | \cdo$ 

where :math: 'x' is an element of the tensor \*lhs\* and :math: '\sigma' is the scalar.

Example::

 $smooth_{11}([1, 2, 3, 4]) = [0.5, 1.5, 2.5, 3.5] smooth_{11}([1, 2, 3, 4], scalar=1) = [0.5, 1.5, 2.5, 3.5]$ 

Defined in src/operator/tensor/elemwise\_binary\_scalar\_op\_extended.cc:L108

## Value

out The result mx.symbol

mx.symbol.Softmax

Softmax: Computes the gradient of cross entropy loss with respect to softmax output.

# Description

- This operator computes the gradient in two steps. The cross entropy loss does not actually need to be computed.

```
mx.symbol.Softmax(...)
```

mx.symbol.Softmax 493

### **Arguments**

data NDArray-or-Symbol Input array. label NDArray-or-Symbol Ground truth label. grad.scale float, optional, default=1 Scales the gradient by a float factor. ignore.label float, optional, default=-1 The instances whose 'labels' == 'ignore\_label' will be ignored during backward, if 'use\_ignore' is set to "true"). boolean, optional, default=0 If set to "true", the softmax function will be commulti.output puted along axis "1". This is applied when the shape of input array differs from the shape of label array. use.ignore boolean, optional, default=0 If set to "true", the 'ignore\_label' value will not contribute to the backward gradient. boolean, optional, default=0 If set to "true", the softmax function will be compreserve.shape puted along the last axis ("-1"). 'batch', 'null', 'valid', optional, default='null' Normalizes the gradient. normalization

vaich, huir, vand ,optional, default= nun Normanzes the gradient.

out.grad boolean, optional, default=0 Multiplies gradient with output gradient element-

wise.

smooth.alpha float, optional, default=0 Constant for computing a label smoothed version of

cross-entropyfor the backwards pass. This constant gets subtracted from theonehot encoding of the gold label and distributed uniformly to all other labels.

name string, optional Name of the resulting symbol.

### **Details**

- Applies softmax function on the input array. - Computes and returns the gradient of cross entropy loss w.r.t. the softmax output.

- The softmax function, cross entropy loss and gradient is given by:
- Softmax Function:
- .. math::  $\text{textsoftmax}(x)_i = \frac{\text{fracexp}(x_i)}{\text{sum}_j \exp(x_j)}$
- Cross Entropy Function:
- .. math:: \textCE(label, output) = \sum\_i \textlabel\_i \log(\textoutput\_i)
- The gradient of cross entropy loss w.r.t softmax output:
- .. math:: \textgradient = \textoutput \textlabel
- During forward propagation, the softmax function is computed for each instance in the input array.

For general \*N\*-D input arrays with shape :math:  $(d_1, d_2, ..., d_n)$ . The size is :math:  $s=d_1 \cdot d_2 \cdot d_2 \cdot d_n$ . We can use the parameters 'preserve\_shape' and 'multi\_output' to specify the way to compute softmax:

- By default, 'preserve\_shape' is "false". This operator will reshape the input array into a 2-D array with shape :math: ' $(d_1, \frac{1}{2}, \frac{1}{2})$ ' and then compute the softmax function for each row in the reshaped array, and afterwards reshape it back to the original shape :math: ' $(d_1, d_2, ..., d_n)$ '. - If 'preserve\_shape' is "true", the softmax function will be computed along the last axis ('axis' = "-1"). - If 'multi\_output' is "true", the softmax function will be computed along the second axis ('axis' = "1").

494 mx.symbol.softmax

- During backward propagation, the gradient of cross-entropy loss w.r.t softmax output array is computed. The provided label can be a one-hot label array or a probability label array.

- If the parameter 'use\_ignore' is "true", 'ignore\_label' can specify input instances with a particular label to be ignored during backward propagation. \*\*This has no effect when softmax 'output' has same shape as 'label'\*\*.

### Example::

- The parameter 'grad\_scale' can be used to rescale the gradient, which is often used to give each loss function different weights.
- This operator also supports various ways to normalize the gradient by 'normalization', The 'normalization' is applied if softmax output has different shape than the labels. The 'normalization' mode can be set to the followings:
- "'null'": do nothing. "'batch'": divide the gradient by the batch size. "'valid'": divide the gradient by the number of instances which are not ignored.

Defined in src/operator/softmax\_output.cc:L231

### Value

out The result mx.symbol

mx.symbol.softmax

softmax: Applies the softmax function.

## **Description**

The resulting array contains elements in the range (0,1) and the elements along the given axis sum up to 1.

# Usage

```
mx.symbol.softmax(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

length NDArray-or-Symbol The length array.

axis int, optional, default='-1' The axis along which to compute softmax.

temperature double or None, optional, default=None Temperature parameter in softmax

None, 'float16', 'float32', 'float64',optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to the same as input's dtype if not

defined (dtype=None).

use.length boolean or None, optional, default=0 Whether to use the length input as a mask

over the data input.

name string, optional Name of the resulting symbol.

### **Details**

```
.. math:: softmax(\mathbfz/t)_j = \frace^z_j/t\sum_k=1^K e^z_k/t for :math: 'j = 1, ..., K' t is the temperature parameter in softmax function. By default, t equals 1.0 Example:: x = [[\ 1.\ 1.\ 1.]\ [\ 1.\ 1.\ 1.]] softmax(x,axis=0) = [[\ 0.5\ 0.5\ 0.5]\ [\ 0.5\ 0.5\ 0.5]] softmax(x,axis=1) = [[\ 0.33333334, 0.3333334, 0.3333334], [\ 0.33333334, 0.33333334]] Defined in src/operator/nn/softmax.cc:L134
```

### Value

out The result mx.symbol

```
mx.symbol.SoftmaxActivation
```

SoftmaxActivation:Applies softmax activation to input. This is intended for internal layers.

# Description

.. note::

## Usage

```
mx.symbol.SoftmaxActivation(...)
```

### **Arguments**

data NDArray-or-Symbol The input array.

mode 'channel', 'instance', optional, default='instance' Specifies how to compute the

softmax. If set to "instance", it computes softmax for each instance. If set to "channel", It computes cross channel softmax for each position of each instance.

name string, optional Name of the resulting symbol.

## **Details**

This operator has been deprecated, please use 'softmax'.

If 'mode' = "instance", this operator will compute a softmax for each instance in the batch. This is the default mode.

If 'mode' = "channel", this operator will compute a k-class softmax at each position of each instance, where 'k' = "num\_channel". This mode can only be used when the input array has at least 3 dimensions. This can be used for 'fully convolutional network', 'image segmentation', etc.

### Example::

»> input\_array = mx.nd.array([[3., 0.5, -0.5, 2., 7.], »> [2., -.4, 7., 3., 0.2]]) »> softmax\_act =
mx.nd.SoftmaxActivation(input\_array) »> print softmax\_act.asnumpy() [[ 1.78322066e-02 1.46375655e03 5.38485940e-04 6.56010211e-03 9.73605454e-01] [ 6.56221947e-03 5.95310994e-04 9.73919690e01 1.78379621e-02 1.08472735e-03]]

Defined in src/operator/nn/softmax\_activation.cc:L59

#### Value

out The result mx.symbol

```
mx.symbol.SoftmaxOutput
```

SoftmaxOutput: Computes the gradient of cross entropy loss with respect to softmax output.

## **Description**

- This operator computes the gradient in two steps. The cross entropy loss does not actually need to be computed.

# Usage

```
mx.symbol.SoftmaxOutput(...)
```

## **Arguments**

data	NDArray-or-Symbol Input array.
label	NDArray-or-Symbol Ground truth label.
grad.scale	float, optional, default=1 Scales the gradient by a float factor.
ignore.label	float, optional, default=-1 The instances whose 'labels' == 'ignore_label' will be ignored during backward, if 'use_ignore' is set to "true").
multi.output	boolean, optional, default=0 If set to "true", the softmax function will be computed along axis "1". This is applied when the shape of input array differs from the shape of label array.
use.ignore	boolean, optional, default=0 If set to "true", the 'ignore_label' value will not contribute to the backward gradient.

preserve.shape boolean, optional, default=0 If set to "true", the softmax function will be com-

puted along the last axis ("-1").

normalization 'batch', 'null', 'valid', optional, default='null' Normalizes the gradient.

out.grad boolean, optional, default=0 Multiplies gradient with output gradient element-

wise.

smooth.alpha float, optional, default=0 Constant for computing a label smoothed version of

cross-entropyfor the backwards pass. This constant gets subtracted from theonehot encoding of the gold label and distributed uniformly to all other labels.

name string, optional Name of the resulting symbol.

### **Details**

- Applies softmax function on the input array. - Computes and returns the gradient of cross entropy loss w.r.t. the softmax output.

- The softmax function, cross entropy loss and gradient is given by:
- Softmax Function:
- .. math::  $\text{textsoftmax}(x)_i = \frac{x_i}{\sup_i \exp(x_i)}$
- Cross Entropy Function:
- .. math:: \textCE(label, output) = \sum\_i \textlabel\_i \log(\textoutput\_i)
- The gradient of cross entropy loss w.r.t softmax output:
- .. math:: \textgradient = \textoutput \textlabel
- During forward propagation, the softmax function is computed for each instance in the input array.

For general  $N^*$ -D input arrays with shape :math:  $(d_1, d_2, ..., d_n)$ . The size is :math:  $s=d_1 \cdot d_2 \cdot d_1 \cdot d_2 \cdot d_n$ . We can use the parameters 'preserve\_shape' and 'multi\_output' to specify the way to compute softmax:

- By default, 'preserve\_shape' is "false". This operator will reshape the input array into a 2-D array with shape :math: '(d\_1, \fracsd\_1)' and then compute the softmax function for each row in the reshaped array, and afterwards reshape it back to the original shape :math: '(d\_1, d\_2, ..., d\_n)'.
- If 'preserve\_shape' is "true", the softmax function will be computed along the last axis ('axis' = "-1"). If 'multi\_output' is "true", the softmax function will be computed along the second axis ('axis' = "1").
- During backward propagation, the gradient of cross-entropy loss w.r.t softmax output array is computed. The provided label can be a one-hot label array or a probability label array.
- If the parameter 'use\_ignore' is "true", 'ignore\_label' can specify input instances with a particular label to be ignored during backward propagation. \*\*This has no effect when softmax 'output' has same shape as 'label'\*\*.

### Example::

data = [[1,2,3,4],[2,2,2,2],[3,3,3,3],[4,4,4,4]] label = [1,0,2,3] ignore\_label = 1 SoftmaxOutput(data=data, label = label,\ multi\_output=true, use\_ignore=true,\ ignore\_label=ignore\_label) ## forward softmax output [[ 0.0320586 0.08714432 0.23688284 0.64391428] [ 0.25 0.25 0.25 0.25 ] [ 0.25 0.25 0.25 0.25 0.25 ] [ 0.25 0.25 0.25 0.25 ] ## backward gradient output [[ 0. 0. 0. 0. 0. ] [-0.75 0.25 0.25 0.25 0.25 ] [ 0.25 0.25 0.25 0.25 -0.75 0.25] [ 0.25 0.25 -0.75 0.25] [ 0.25 0.25 -0.75 0.25] [ 0.25 0.25 -0.75]] ## notice that the first row is all 0 because label[0] is 1, which is equal to ignore\_label.

- The parameter 'grad\_scale' can be used to rescale the gradient, which is often used to give each loss function different weights.
- This operator also supports various ways to normalize the gradient by 'normalization', The 'normalization' is applied if softmax output has different shape than the labels. The 'normalization' mode can be set to the followings:
- "'null'": do nothing. "'batch'": divide the gradient by the batch size. "'valid'": divide the gradient by the number of instances which are not ignored.

Defined in src/operator/softmax\_output.cc:L231

## Value

out The result mx.symbol

```
mx.symbol.softmax_cross_entropy
```

softmax\_cross\_entropy:Calculate cross entropy of softmax output and one-hot label.

# **Description**

- This operator computes the cross entropy in two steps: - Applies softmax function on the input array. - Computes and returns the cross entropy loss between the softmax output and the labels.

### Usage

```
mx.symbol.softmax_cross_entropy(...)
```

## **Arguments**

data NDArray-or-Symbol Input data label NDArray-or-Symbol Input label

name string, optional Name of the resulting symbol.

### **Details**

- The softmax function and cross entropy loss is given by:
- Softmax Function:
- .. math::  $\text{textsoftmax}(x)_i = \frac{x_i}{\sum_i \exp(x_i)}$
- Cross Entropy Function:
- .. math:: \textCE(label, output) = \sum\_i \textlabel\_i \log(\textoutput\_i)

# Example::

```
x = [[1, 2, 3], [11, 7, 5]]
```

label = [2, 0]

softmax(x) = [[0.09003057, 0.24472848, 0.66524094], [0.97962922, 0.01794253, 0.00242826]]

 $softmax\_cross\_entropy(data, label) = -log(0.66524084) - log(0.97962922) = 0.4281871$ 

Defined in src/operator/loss\_binary\_op.cc:L59

mx.symbol.softmin 499

## Value

out The result mx.symbol

mx.symbol.softmin softmin:Applies the softmin function.

# Description

The resulting array contains elements in the range (0,1) and the elements along the given axis sum up to 1.

# Usage

```
mx.symbol.softmin(...)
```

# **Arguments**

data NDArray-or-Symbol The input array.

axis int, optional, default='-1' The axis along which to compute softmax.

temperature double or None, optional, default=None Temperature parameter in softmax

dtype None, 'float16', 'float32', 'float64', optional, default='None' DType of the out-

put in case this can't be inferred. Defaults to the same as input's dtype if not

defined (dtype=None).

use.length boolean or None, optional, default=0 Whether to use the length input as a mask

over the data input.

name string, optional Name of the resulting symbol.

# **Details**

```
.. math:: softmin(\mathbfz/t)_j = \frace^-z_j/t\sum_k=1^K e^-z_k/t for :math: 'j = 1, ..., K' t is the temperature parameter in softmax function. By default, t equals 1.0 Example:: x = [[\ 1.\ 2.\ 3.]\ [\ 3.\ 2.\ 1.]] softmin(x,axis=0) = [[\ 0.88079703, 0.5, 0.11920292], [\ 0.11920292, 0.5, 0.88079703]] softmin(x,axis=1) = [[\ 0.66524094, 0.24472848, 0.09003057], [\ 0.09003057, 0.24472848, 0.66524094]] Defined in src/operator/nn/softmin.cc:L57
```

## Value

500 mx.symbol.sort

mx.symbol.softsign

softsign: Computes softsign of x element-wise.

## **Description**

```
.. math:: y = x / (1 + abs(x))
```

# Usage

```
mx.symbol.softsign(...)
```

## **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

The storage type of "softsign" output is always dense

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L191

## Value

out The result mx.symbol

mx.symbol.sort

sort:Returns a sorted copy of an input array along the given axis.

## **Description**

Examples::

### **Usage**

```
mx.symbol.sort(...)
```

# **Arguments**

data NDArray-or-Symbol The input array

axis int or None, optional, default='-1' Axis along which to choose sort the input

tensor. If not given, the flattened array is used. Default is -1.

is.ascend boolean, optional, default=1 Whether to sort in ascending or descending order.

name string, optional Name of the resulting symbol.

### **Details**

```
x = [[ 1, 4], [ 3, 1]]
// sorts along the last axis sort(x) = [[ 1., 4.], [ 1., 3.]]
// flattens and then sorts sort(x, axis=None) = [ 1., 1., 3., 4.]
// sorts along the first axis sort(x, axis=0) = [[ 1., 1.], [ 3., 4.]]
// in a descend order sort(x, is_ascend=0) = [[ 4., 1.], [ 3., 1.]]
Defined in src/operator/tensor/ordering_op.cc:L133
```

### Value

out The result mx.symbol

mx.symbol.space\_to\_depth

space\_to\_depth:Rearranges(permutes) blocks of spatial data Similar to ONNX SpaceToDepth operator: into depth. https://github.com/onnx/onnx/blob/master/docs/Operators.md#SpaceToDepth The output is a new tensor where the values from height and width dimension are moved to the depth dimension. The reverse of this operation is "depth\_to\_space". .. math:: \begingather\* x \prime = reshape(x, [N, C, H / block | size, block | size, W / block | size, $block\_size]) \setminus x \setminus prime \setminus prime = transpose(x \setminus prime, [0, 3, 5, 1,$ 2, 4])  $\ y = reshape(x \rangle prime \rangle prime, [N, C * (block size ^ 2), H /$ block\\_size, W / block\\_size]) \endgather\* where :math: 'x' is an input tensor with default layout as :math: '[N, C, H, W]': [batch, channels, height, width] and :math: 'y' is the output tensor of layout :math: '[N,  $C * (block size ^ 2), H/block size, W/block size] Example:: x =$ [[[0, 6, 1, 7, 2, 8], [12, 18, 13, 19, 14, 20], [3, 9, 4, 10, 5, 11], [15, 21, 16, 22, 17, 23]]]  $space_to_depth(x, 2) = [[[[0, 1, 2], [3, 4, 5]],$ [[6, 7, 8], [9, 10, 11]], [[12, 13, 14], [15, 16, 17]], [[18, 19, 20], [21, 22, 23]]]]

# **Description**

Defined in src/operator/tensor/matrix\_op.cc:L1019

## Usage

```
mx.symbol.space_to_depth(...)
```

# **Arguments**

data NDArray-or-Symbol Input ndarray

block.size int, required Blocks of [block\_size. block\_size] are moved

name string, optional Name of the resulting symbol.

# Value

out The result mx.symbol

 $\verb|mx.symbol.SpatialTransformer| \\$ 

SpatialTransformer:Applies a spatial transformer to input feature map.

# **Description**

SpatialTransformer:Applies a spatial transformer to input feature map.

# Usage

```
mx.symbol.SpatialTransformer(...)
```

# Arguments

data NDArray-or-Symbol Input data to the SpatialTransformer	Op.
---	-----

loc NDArray-or-Symbol localisation net, the output dim should be 6 when trans-

form\_type is affine. You shold initialize the weight and bias with identity tran-

form.

target.shape Shape(tuple), optional, default=[0,0] output shape(h, w) of spatial transformer:

(y, x)

transform.type 'affine', required transformation type

sampler.type 'bilinear', required sampling type

cudnn.off boolean or None, optional, default=None whether to turn cudnn off

name string, optional Name of the resulting symbol.

## Value

mx.symbol.split 503

mx.symbol.split

split:Splits an array along a particular axis into multiple sub-arrays.

# Description

```
.. note:: "SliceChannel" is deprecated. Use "split" instead.
```

## Usage

```
mx.symbol.split(...)
```

## Arguments

data NDArray-or-Symbol The input

num.outputs int, required Number of splits. Note that this should evenly divide the length of

the 'axis'.

axis int, optional, default='1' Axis along which to split.

squeeze.axis boolean, optional, default=0 If true, Removes the axis with length 1 from the

shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "true" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis'

can be set to "true" only if "input.shape[axis] == num\_outputs".

name string, optional Name of the resulting symbol.

### **Details**

\*\*Note\*\* that 'num\_outputs' should evenly divide the length of the axis along which to split the array.

## Example::

```
 \begin{aligned} x &= \text{[[[ 1.] [ 2.]] [[ 3.] [ 4.]] [[ 5.] [ 6.]]] } x.shape &= (3, 2, 1) \\ y &= \text{split}(x, \text{axis=1}, \text{num\_outputs=2}) \text{// a list of 2 arrays with shape } (3, 1, 1) \text{ y} &= \text{[[[ 1.]] [[ 3.]] [[ 5.]]]} \\ \text{[[[ 2.]] [[ 4.]] [[ 6.]]]} \\ y &[0].\text{shape} &= (3, 1, 1) \\ z &= \text{split}(x, \text{axis=0}, \text{num\_outputs=3}) \text{// a list of 3 arrays with shape } (1, 2, 1) \text{ z} &= \text{[[[ 1.] [ 2.]]]} \\ \text{[[[ 3.] [ 4.]]]} \\ \text{[[[ 5.] [ 6.]]]} \\ z &[0].\text{shape} &= (1, 2, 1) \end{aligned}
```

'squeeze\_axis=1' removes the axis with length 1 from the shapes of the output arrays. \*\*Note\*\* that setting 'squeeze\_axis' to "1" removes axis with length 1 only along the 'axis' which it is split. Also 'squeeze\_axis' can be set to true only if "input.shape[axis] == num\_outputs".

## Example::

```
z = split(x, axis=0, num\_outputs=3, squeeze\_axis=1) // a list of 3 arrays with shape (2, 1) <math>z = [[1.] [2.]]
```

504 mx.symbol.sqrt

```
[[ 3.] [ 4.]]
[[ 5.] [ 6.]] z[0].shape = (2 ,1 )
Defined in src/operator/slice_channel.cc:L107
```

### Value

out The result mx.symbol

mx.symbol.sqrt

sqrt:Returns element-wise square-root value of the input.

# **Description**

```
.. math:: \text{textrmsqrt}(x) = \text{sqrt}x
```

# Usage

```
mx.symbol.sqrt(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

## **Details**

Example::

```
sqrt([4, 9, 16]) = [2, 3, 4]
```

The storage type of "sqrt" output depends upon the input storage type:

-  $sqrt(default) = default - sqrt(row\_sparse) = row\_sparse - sqrt(csr) = csr$ 

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L170

### Value

mx.symbol.square 505

mx.symbol.square

square: Returns element-wise squared value of the input.

## **Description**

```
.. math:: square(x) = x^2
```

#### Usage

```
mx.symbol.square(...)
```

## Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

Example::

square([2, 3, 4]) = [4, 9, 16]

The storage type of "square" output depends upon the input storage type:

- square(default) = default - square(row\_sparse) = row\_sparse - square(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_pow.cc:L119

#### Value

out The result mx.symbol

mx.symbol.squeeze

squeeze:Remove single-dimensional entries from the shape of an array. Same behavior of defining the output tensor shape as numpy.squeeze for the most of cases. See the following note for exception. Examples:: data = [[[0], [1], [2]]] squeeze(data) = [0, 1, 2] squeeze(data, axis=0) = [[0], [1], [2]] squeeze(data, axis=2) = [[0, 1, 2]] squeeze(data, axis=0) = [0, 1, 2] . Note:: The output of this operator will keep at least one dimension not removed. For example, squeeze([[[4]]]) = [4], while in numpy.squeeze, the output will become a scalar.

506 mx.symbol.stack

#### **Description**

squeeze:Remove single-dimensional entries from the shape of an array. Same behavior of defining the output tensor shape as numpy.squeeze for the most of cases. See the following note for exception. Examples:: data = [[[0], [1], [2]]] squeeze(data) = [0, 1, 2] squeeze(data, axis=0) = [[0], [1], [2]] squeeze(data, axis=0) = [[0, 1, 2]] squeeze(data, axis=0, 2) = [0, 1, 2] . Note:: The output of this operator will keep at least one dimension not removed. For example, squeeze([[[4]]]) = [4], while in numpy.squeeze, the output will become a scalar.

## Usage

```
mx.symbol.squeeze(...)
```

#### **Arguments**

data NDArray-or-Symbol data to squeeze

axis Shape or None, optional, default=None Selects a subset of the single-dimensional

entries in the shape. If an axis is selected with shape entry greater than one, an

error is raised.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.stack

stack: Join a sequence of arrays along a new axis. The axis parameter specifies the index of the new axis in the dimensions of the result. For example, if axis=0 it will be the first dimension and if axis=-1 it will be the last dimension. Examples:: x = [1, 2] y = [3, 4] stack(x, y) = [1, 2], [3, 4] stack(x, y), [4, 4]

## Description

stack: Join a sequence of arrays along a new axis. The axis parameter specifies the index of the new axis in the dimensions of the result. For example, if axis=0 it will be the first dimension and if axis=-1 it will be the last dimension. Examples:: x = [1, 2] y = [3, 4] stack(x, y) = [[1, 2], [3, 4]] stack(x, y, axis=1) = [[1, 3], [2, 4]]

#### Usage

```
mx.symbol.stack(...)
```

#### **Arguments**

data NDArray-or-Symbol[] List of arrays to stack

axis int, optional, default='0' The axis in the result array along which the input arrays

are stacked.

num.args int, required Number of inputs to be stacked.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

```
mx.symbol.stop_gradient
```

stop\_gradient:Stops gradient computation.

# Description

Stops the accumulated gradient of the inputs from flowing through this operator in the backward direction. In other words, this operator prevents the contribution of its inputs to be taken into account for computing gradients.

## Usage

```
mx.symbol.stop_gradient(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L326

#### **Details**

## Example::

```
v1 = [1, 2] v2 = [0, 1] a = Variable('a') b = Variable('b') b_stop_grad = stop_gradient(3 * b) loss = MakeLoss(b_stop_grad + a) executor = loss.simple_bind(ctx=cpu(), a=(1,2), b=(1,2)) executor.forward(is_train=True, a=v1, b=v2) executor.outputs [ 1. 5.] executor.backward() executor.grad_arrays [ 0. 0.] [ 1. 1.]
```

# Value

508 mx.symbol.sum

mx.symbol.sum

sum: Computes the sum of array elements over given axes.

#### **Description**

.. Note::

## Usage

```
mx.symbol.sum(...)
```

## **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

#### **Details**

'sum' and 'sum\_axis' are equivalent. For ndarray of csr storage type summation along axis 0 and axis 1 is supported. Setting keepdims or exclude to True will cause a fallback to dense operator.

#### Example::

```
data = [[[1, 2], [2, 3], [1, 3]], [[1, 4], [4, 3], [5, 2]], [[7, 1], [7, 2], [7, 3]]] sum(data, axis=1) [[ 4. 8.] [ 10. 9.] [ 21. 6.]] sum(data, axis=[1,2]) [ 12. 19. 27.] data = [[1, 2, 0], [3, 0, 1], [4, 1, 0]] csr = cast_storage(data, 'csr') sum(csr, axis=0) [ 8. 3. 1.] sum(csr, axis=1) [ 3. 4. 5.]
```

Defined in src/operator/tensor/broadcast\_reduce\_sum\_value.cc:L67

mx.symbol.sum\_axis 509

#### Value

out The result mx.symbol

mx.symbol.sum\_axis

sum\_axis: Computes the sum of array elements over given axes.

#### **Description**

.. Note::

#### Usage

```
mx.symbol.sum_axis(...)
```

#### **Arguments**

data NDArray-or-Symbol The input

axis Shape or None, optional, default=None The axis or axes along which to perform

the reduction.

The default, 'axis=()', will compute over all elements into a scalar array with

shape '(1,)'.

If 'axis' is int, a reduction is performed on a particular axis.

If 'axis' is a tuple of ints, a reduction is performed on all the axes specified in

the tuple.

If 'exclude' is true, reduction will be performed on the axes that are NOT in axis

instead.

Negative values means indexing from right to left.

keepdims boolean, optional, default=0 If this is set to 'True', the reduced axes are left in

the result as dimension with size one.

exclude boolean, optional, default=0 Whether to perform reduction on axis that are NOT

in axis instead.

name string, optional Name of the resulting symbol.

#### **Details**

'sum' and 'sum\_axis' are equivalent. For ndarray of csr storage type summation along axis 0 and axis 1 is supported. Setting keepdims or exclude to True will cause a fallback to dense operator.

#### Example::

```
data = [[[1, 2], [2, 3], [1, 3]], [[1, 4], [4, 3], [5, 2]], [[7, 1], [7, 2], [7, 3]]] sum(data, axis=1) [[ 4. 8.] [ 10. 9.] [ 21. 6.]] sum(data, axis=[1,2]) [ 12. 19. 27.] data = [[1, 2, 0], [3, 0, 1], [4, 1, 0]] csr = cast_storage(data, 'csr')
```

```
sum(csr, axis=0) [ 8. 3. 1.]
sum(csr, axis=1) [ 3. 4. 5.]
```

Defined in src/operator/tensor/broadcast\_reduce\_sum\_value.cc:L67

#### Value

out The result mx.symbol

 ${\tt mx.symbol.SVMOutput:} \ SVMOutput: Computes support vector machine based transformation of the input.$ 

## **Description**

This tutorial demonstrates using SVM as output layer for classification instead of softmax: https://github.com/dmlc/mxnet/tre

#### Usage

```
mx.symbol.SVMOutput(...)
```

#### **Arguments**

data NDArray-or-Symbol Input data for SVM transformation.

label NDArray-or-Symbol Class label for the input data.

margin float, optional, default=1 The loss function penalizes outputs that lie outside this

margin. Default margin is 1.

regularization.coefficient

float, optional, default=1 Regularization parameter for the SVM. This balances

the tradeoff between coefficient size and error.

use.linear boolean, optional, default=0 Whether to use L1-SVM objective. L2-SVM ob-

jective is used by default.

name string, optional Name of the resulting symbol.

## Value

mx.symbol.swapaxes 511

mx.symbol.swapaxes

swapaxes:Interchanges two axes of an array.

# Description

Examples::

#### Usage

```
mx.symbol.swapaxes(...)
```

## Arguments

data NDArray-or-Symbol Input array.

dim1 int, optional, default='0' the first axis to be swapped.dim2 int, optional, default='0' the second axis to be swapped.

name string, optional Name of the resulting symbol.

#### **Details**

```
x = [[1, 2, 3]]) swapaxes(x, 0, 1) = [[1], [2], [3]]

x = [[[0, 1], [2, 3]], [[4, 5], [6, 7]]] // (2,2,2) array

swapaxes(x, 0, 2) = [[[0, 4], [2, 6]], [[1, 5], [3, 7]]]

Defined in src/operator/swapaxis.cc:L70
```

## Value

out The result mx.symbol

mx.symbol.SwapAxis

SwapAxis:Interchanges two axes of an array.

## **Description**

Examples::

## Usage

```
mx.symbol.SwapAxis(...)
```

512 mx.symbol.take

## **Arguments**

data	NDArray-or-Symbol Input array.
dim1	int, optional, default='0' the first axis to be swapped.
dim2	int, optional, default='0' the second axis to be swapped.
name	string, optional Name of the resulting symbol.

#### **Details**

```
x = [[1, 2, 3]]) swapaxes(x, 0, 1) = [[1], [2], [3]]

x = [[[0, 1], [2, 3]], [[4, 5], [6, 7]]] // (2,2,2) array

swapaxes(x, 0, 2) = [[[0, 4], [2, 6]], [[1, 5], [3, 7]]]

Defined in src/operator/swapaxis.cc:L70
```

## Value

out The result mx.symbol

mx.symbol.take

take: Takes elements from an input array along the given axis.

# Description

This function slices the input array along a particular axis with the provided indices.

## Usage

```
mx.symbol.take(...)
```

# Arguments

a	NDArray-or-Symbol The input array.
indices	NDArray-or-Symbol The indices of the values to be extracted.
axis	int, optional, default='0' The axis of input array to be taken. For input tensor of rank $r$ , it could be in the range of $[-r, r-1]$
mode	'clip', 'raise', 'wrap',optional, default='clip' Specify how out-of-bound indices bahave. Default is "clip". "clip" means clip to the range. So, if all indices mentioned are too large, they are replaced by the index that addresses the last element along an axis. "wrap" means to wrap around. "raise" means to raise an error when index out of range.
name	string, optional Name of the resulting symbol.

mx.symbol.tan 513

#### **Details**

Given data tensor of rank  $r \ge 1$ , and indices tensor of rank q, gather entries of the axis dimension of data (by default outer-most one as axis=0) indexed by indices, and concatenates them in an output tensor of rank q + (r - 1).

```
Examples::
```

```
x = [4. 5. 6.]
```

// Trivial case, take the second element along the first axis.

$$take(x, [1]) = [5.]$$

// The other trivial case, axis=-1, take the third element along the first axis

$$take(x, [3], axis=-1, mode='clip') = [6.]$$

$$x = [[1., 2.], [3., 4.], [5., 6.]]$$

// In this case we will get rows 0 and 1, then 1 and 2. Along axis 0

$$take(x, [[0,1],[1,2]]) = [[[1., 2.], [3., 4.]],$$

// In this case we will get rows 0 and 1, then 1 and 2 (calculated by wrapping around). // Along axis 1

```
take(x, [[0, 3], [-1, -2]], axis=1, mode='wrap') = [[[ 1. 2.] [ 2. 1.]]
```

[[ 3. 4.] [ 4. 3.]]

[[ 5. 6.] [ 6. 5.]]]

The storage type of "take" output depends upon the input storage type:

- take(default, default) = default - take(csr, default, axis=0) = csr

Defined in src/operator/tensor/indexing\_op.cc:L782

#### Value

out The result mx.symbol

mx.symbol.tan

tan: Computes the element-wise tangent of the input array.

#### **Description**

The input should be in radians (:math: '2\pi' rad equals 360 degrees).

## Usage

```
mx.symbol.tan(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

514 mx.symbol.tanh

## **Details**

```
.. math:: tan([0, \pi/4, \pi/2]) = [0, 1, -inf]
```

The storage type of "tan" output depends upon the input storage type:

- tan(default) = default - tan(row\_sparse) = row\_sparse - tan(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L140

## Value

out The result mx.symbol

mx.symbol.tanh

tanh:Returns the hyperbolic tangent of the input array, computed element-wise.

# Description

```
.. math:: tanh(x) = sinh(x) / cosh(x)
```

# Usage

```
mx.symbol.tanh(...)
```

# Arguments

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

#### **Details**

The storage type of "tanh" output depends upon the input storage type:

- tanh(default) = default - tanh(row\_sparse) = row\_sparse - tanh(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_trig.cc:L451

#### Value

mx.symbol.tile 515

mx.symbol.tile

tile:Repeats the whole array multiple times. If "reps" has length \*d\*, and input array has dimension of \*n\*. There are three cases: \*n=d\*. Repeat \*i\*-th dimension of the input by "reps[i]" times:: x = [[1, 2], [3, 4]] tile(x, reps=(2,3)) = [[1, 2, 1, 2, 1, 2, 1, 2, 1, 2, 1, 3, 4,

## Description

Defined in src/operator/tensor/matrix\_op.cc:L796

#### Usage

```
mx.symbol.tile(...)
```

#### **Arguments**

data NDArray-or-Symbol Input data array

reps Shape(tuple), required The number of times for repeating the tensor a. Each dim

size of reps must be a positive integer. If reps has length d, the result will have dimension of max(d, a.ndim); If a.ndim < d, a is promoted to be d-dimensional by prepending new axes. If a.ndim > d, reps is promoted to a.ndim by pre-

pending 1's to it.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

 ${\it mx.symbol.topk}$ 

topk:Returns the indices of the top \*k\* elements in an input array along the given axis (by default). If ret\_type is set to 'value' returns the value of top \*k\* elements (instead of indices). In case of ret\_type = 'both', both value and index would be returned. The returned elements will be sorted.

516 mx.symbol.topk

## **Description**

Examples::

#### Usage

```
mx.symbol.topk(...)
```

#### **Arguments**

data	NDArray-or-Symbol The input array
axis	int or None, optional, default='-1' Axis along which to choose the top k indices. If not given, the flattened array is used. Default is -1.
k	int, optional, default='1' Number of top elements to select, should be always smaller than or equal to the element number in the given axis. A global sort is performed if set $k < 1$ .
ret.typ	'both', 'indices', 'mask', 'value',optional, default='indices' The return type. "value" means to return the top k values, "indices" means to return the indices of the top k values, "mask" means to return a mask array containing 0 and 1. 1 means the top k values. "both" means to return a list of both values and indices of top k elements.
is.ascend	boolean, optional, default=0 Whether to choose k largest or k smallest elements. Top K largest elements will be chosen if set to false.
dtype	'float16', 'float32', 'float64', 'int32', 'int64', 'uint8', optional, default='float32' DType of the output indices when ret_typ is "indices" or "both". An error will be raised if the selected data type cannot precisely represent the indices.
name	string, optional Name of the resulting symbol.

## **Details**

```
 x = [[\ 0.3, 0.2, 0.4], [\ 0.1, 0.3, 0.2]]  // returns an index of the largest element on last axis topk(x) = [[\ 2.], [\ 1.]] 
 // returns the value of top-2 largest elements on last axis topk(x, ret_typ='value', k=2) = [[\ 0.4, 0.3], [\ 0.3, 0.2]] 
 // returns the value of top-2 smallest elements on last axis topk(x, ret_typ='value', k=2, is_ascend=1) 
 = [[\ 0.2, 0.3], [\ 0.1, 0.2]] 
 // returns the value of top-2 largest elements on axis 0 topk(x, axis=0, ret_typ='value', k=2) = [[\ 0.3, 0.3, 0.4], [\ 0.1, 0.2, 0.2]] 
 // flattens and then returns list of both values and indices topk(x, ret_typ='both', k=2) = [[[\ 0.4, 0.3], [\ 0.3, 0.2]], [[\ 2., 0.], [\ 1., 2.]]] 
 Defined in src/operator/tensor/ordering_op.cc:L68
```

#### Value

mx.symbol.transpose 517

```
mx.symbol.transpose transpose: Permutes the dimensions of an array. Examples:: <math>x = [[\ 1, \ 2], \ [\ 3, \ 4]] transpose(x) = [[\ 1., \ 3.], \ [\ 2., \ 4.]] x = [[\ 1., \ 2.], \ [\ 3., \ 4.]], [\ 5., \ 6.], \ [\ 7., \ 8.]]] transpose(x) = [[[\ 1., \ 2.], \ [\ 5., \ 6.]], \ [[\ 3., \ 4.], \ [\ 7., \ 8.]]] transpose(x, \ axes=(1,0,2)) = [[[\ 1., \ 2.], \ [\ 5., \ 6.]], \ [[\ 3., \ 4.], \ [\ 7., \ 8.]]]
```

#### **Description**

Defined in src/operator/tensor/matrix\_op.cc:L328

#### Usage

```
mx.symbol.transpose(...)
```

#### **Arguments**

data NDArray-or-Symbol Source input

axes Shape(tuple), optional, default=[] Target axis order. By default the axes will be

inverted.

name string, optional Name of the resulting symbol.

#### Value

out The result mx.symbol

mx.symbol.trunc trunc:Return the element-wise truncated value of the input.

## **Description**

The truncated value of the scalar x is the nearest integer i which is closer to zero than x is. In short, the fractional part of the signed number x is discarded.

## Usage

```
mx.symbol.trunc(...)
```

#### **Arguments**

data NDArray-or-Symbol The input array.

name string, optional Name of the resulting symbol.

518 mx.symbol.uniform

## **Details**

```
Example::
```

```
trunc([-2.1, -1.9, 1.5, 1.9, 2.1]) = [-2., -1., 1., 1., 2.]
```

The storage type of "trunc" output depends upon the input storage type:

- trunc(default) = default - trunc(row\_sparse) = row\_sparse - trunc(csr) = csr

Defined in src/operator/tensor/elemwise\_unary\_op\_basic.cc:L857

#### Value

out The result mx.symbol

mx.symbol.uniform

uniform:Draw random samples from a uniform distribution.

#### **Description**

.. note:: The existing alias "uniform" is deprecated.

## Usage

```
mx.symbol.uniform(...)
```

# Arguments

low	float, optional, default=0 Lower bound of the distribution.
high	float, optional, default=1 Upper bound of the distribution.
shape	Shape(tuple), optional, default=None Shape of the output.
ctx	string, optional, default="Context of output, in format [cpulgpulcpu_pinned](n). Only used for imperative calls.
dtype	'None', 'float16', 'float32', 'float64', optional, default='None' DType of the output in case this can't be inferred. Defaults to float32 if not defined (dtype=None).
name	string, optional Name of the resulting symbol.

#### **Details**

Samples are uniformly distributed over the half-open interval \*[low, high)\* (includes \*low\*, but excludes \*high\*).

Example::

```
uniform(low=0, high=1, shape=(2,2)) = [[ 0.60276335, 0.85794562], [ 0.54488319, 0.84725171]] Defined in src/operator/random/sample_op.cc:L96
```

#### Value

mx.symbol.unravel\_index

unravel\_index:Converts an array of flat indices into a batch of index arrays. The operator follows numpy conventions so a single multi index is given by a column of the output matrix. The leading dimension may be left unspecified by using -1 as placeholder.

#### **Description**

Examples::

#### Usage

```
mx.symbol.unravel_index(...)
```

## **Arguments**

data NDArray-or-Symbol Array of flat indices

shape Shape(tuple), optional, default=None Shape of the array into which the multi-

indices apply.

name string, optional Name of the resulting symbol.

## **Details**

```
A = [22,41,37] \ unravel\_index(A, shape=(7,6)) = [[3,6,6], [4,5,1]] \ unravel\_index(A, shape=(-1,6)) \\ = [[3,6,6], [4,5,1]]
```

 $B = [[22,41,37],[10,11,15]] \ unravel\_index(B, shape=(7,6)) = [[[3,6,6],[1,1,2]], [[4,5,1],[4,5,3]]] \ unravel\_index(B, shape=(-1,6)) = [[[3,6,6],[1,1,2]], [[4,5,1],[4,5,3]]]$ 

Defined in src/operator/tensor/ravel.cc:L76

## Value

out The result mx.symbol

 $\verb|mx.symbol.UpSampling| UpSampling: Upsamples the given input data.$ 

#### **Description**

Two algorithms ("sample\_type") are available for upsampling:

#### Usage

```
mx.symbol.UpSampling(...)
```

#### **Arguments**

data NDArray-or-Symbol[] Array of tensors to upsample. For bilinear upsampling,

there should be 2 inputs - 1 data and 1 weight.

scale int, required Up sampling scale

num.filter int, optional, default='0' Input filter. Only used by bilinear sample\_type.Since

bilinear upsampling uses deconvolution, num\_filters is set to the number of

channels.

sample.type 'bilinear', 'nearest', required upsampling method

multi.input.mode

'concat', 'sum',optional, default='concat' How to handle multiple input. concat means concatenate upsampled images along the channel dimension. sum means

add all images together, only available for nearest neighbor upsampling.

num.args int, required Number of inputs to be upsampled. For nearest neighbor upsam-

pling, this can be 1-N; the size of output will be(scale\* $h_0$ ,scale\* $w_0$ ) and all other inputs will be upsampled to the same size. For bilinear upsampling this

must be 2; 1 input and 1 weight.

workspace long (non-negative), optional, default=512 Tmp workspace for deconvolution

(MB)

name string, optional Name of the resulting symbol.

#### **Details**

- Nearest Neighbor - Bilinear

\*\*Nearest Neighbor Upsampling\*\*

Input data is expected to be NCHW.

Example::

```
\mathbf{x} = [[[[1. \ 1. \ 1.] \ [1. \ 1. \ 1.] \ [1. \ 1. \ 1.]]]]
```

UpSampling(x, scale=2, sample\_type='nearest') = [[[[1. 1. 1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ] [1. 1. 1. 1. 1. ]]]]

\*\*Bilinear Upsampling\*\*

Uses 'deconvolution' algorithm under the hood. You need provide both input data and the kernel.

Input data is expected to be NCHW.

'num\_filter' is expected to be same as the number of channels.

Example::

```
\mathbf{x} = [[[[1. \ 1. \ 1.] \ [1. \ 1. \ 1.] \ [1. \ 1. \ 1.]]]]

\mathbf{w} = [[[[1. \ 1. \ 1. \ 1.] \ [1. \ 1. \ 1. \ 1.] \ [1. \ 1. \ 1. \ 1.]]]]
```

UpSampling(x, w, scale=2, sample\_type='bilinear', num\_filter=1) = [[[[1. 2. 2. 2. 2. 2. 1.] [2. 4. 4. 4. 2.] [2. 4. 4. 4. 2.] [2. 4. 4. 4. 2.] [1. 2. 2. 2. 2. 1.]]]]

Defined in src/operator/nn/upsampling.cc:L173

#### Value

mx.symbol.Variable 521

mx.symbol.Variable

Create a symbolic variable with specified name.

## **Description**

Create a symbolic variable with specified name.

#### **Arguments**

name

string The name of the result symbol.

#### Value

The result symbol

mx.symbol.where

where: Return the elements, either from x or y, depending on the condition.

# Description

Given three ndarrays, condition, x, and y, return an ndarray with the elements from x or y, depending on the elements from condition are true or false. x and y must have the same shape. If condition has the same shape as x, each element in the output array is from x if the corresponding element in the condition is true, and from y if false.

# Usage

```
mx.symbol.where(...)
```

## **Arguments**

condition NDArray-or-Symbol condition array

x NDArray-or-Symbol y NDArray-or-Symbol

name string, optional Name of the resulting symbol.

#### **Details**

If condition does not have the same shape as x, it must be a 1D array whose size is the same as x's first dimension size. Each row of the output array is from x's row if the corresponding element from condition is true, and from y's row if false.

Note that all non-zero values are interpreted as "True" in condition.

Examples::

```
x = [[1, 2], [3, 4]] y = [[5, 6], [7, 8]] cond = [[0, 1], [-1, 0]]
where(cond, x, y) = [[5, 2], [3, 8]]
csr_cond = cast_storage(cond, 'csr')
where(csr_cond, x, y) = [[5, 2], [3, 8]]
Defined in src/operator/tensor/control_flow_op.cc:L57
```

#### Value

out The result mx.symbol

```
mx.symbol.zeros_like zeros_like:Return an array of zeros with the same shape, type and storage type as the input array.
```

## Description

The storage type of "zeros\_like" output depends on the storage type of the input

#### Usage

```
mx.symbol.zeros_like(...)
```

## **Arguments**

data NDArray-or-Symbol The input

name string, optional Name of the resulting symbol.

#### **Details**

```
- zeros_like(row_sparse) = row_sparse - zeros_like(csr) = csr - zeros_like(default) = default Examples:: x = [[\ 1.,\ 1.,\ 1.],\ [\ 1.,\ 1.,\ 1.]] zeros_like(x) = [[\ 0.,\ 0.,\ 0.],\ [\ 0.,\ 0.,\ 0.]]
```

#### Value

mx.unserialize 523

mx.unserialize

Unserialize MXNet model from Robject.

## **Description**

Unserialize MXNet model from Robject.

#### Usage

```
mx.unserialize(model)
```

#### **Arguments**

model

The mxnet model loaded from RData files.

mxnet

MXNet: Flexible and Efficient GPU computing and Deep Learning.

#### **Description**

MXNet is a flexible and efficient GPU computing and deep learning framework.

#### **Details**

It enables you to write seamless tensor/matrix computation with multiple GPUs in R.

It also enables you construct and customize the state-of-art deep learning models in R, and apply them to tasks such as image classification and data science challenges.

mxnet.export

Internal function to generate mxnet\_generated.R Users do not need to call this function.

## Description

Internal function to generate mxnet\_generated.R Users do not need to call this function.

#### Usage

```
mxnet.export(path)
```

#### **Arguments**

path

The path to the root of the package.

Ops.MXNDArray

Binary operator overloading of mx.ndarray

# Description

Binary operator overloading of mx.ndarray

# Usage

```
## S3 method for class 'MXNDArray'
Ops(e1, e2)
```

# Arguments

e1

The second operand

outputs

Get the outputs of a symbol.

# Description

Get the outputs of a symbol.

# Usage

```
outputs(x)
```

# Arguments

Χ

The input symbol

predict.MXFeedForwardModel

Predict the outputs given a model and dataset.

# Description

Predict the outputs given a model and dataset.

print.MXNDArray 525

#### Usage

```
## S3 method for class 'MXFeedForwardModel'
predict(
  model,
  X,
  ctx = NULL,
  array.batch.size = 128,
  array.layout = "auto",
  allow.extra.params = FALSE
)
```

#### **Arguments**

model The MXNet Model.

X The dataset to predict.

ctx mx.cpu() or mx.gpu(). The device used to generate the prediction.

array.batch.size

The batch size used in batching. Only used when X is R's array.

array.layout

can be "auto", "colmajor", "rowmajor", (detault=auto) The layout of array. "rowmajor" is only supported for two dimensional array. For matrix, "rowmajor" means  $\dim(X) = c(\text{nexample}, \text{nfeatures})$ , "colmajor" means  $\dim(X) = c(\text{nfeatures}, \text{nexample})$  "auto" will auto detect the layout by match the feature size, and will report error when X is a square matrix to ask user to explicitly specify layout.

allow.extra.params

Whether allow extra parameters that are not needed by symbol. If this is TRUE, no error will be thrown when arg\_params or aux\_params contain extra parameters that is not needed by the executor.

print.MXNDArray

print operator overload of mx.ndarray

# Description

print operator overload of mx.ndarray

#### Usage

```
## S3 method for class 'MXNDArray'
print(nd)
```

## Arguments

nd

The mx.ndarray

rnn.graph

rnn.graph

Generate a RNN symbolic model - requires CUDA

# Description

Generate a RNN symbolic model - requires CUDA

## Usage

```
rnn.graph(
  num_rnn_layer,
  input_size = NULL,
  num\_embed = NULL,
 num_hidden,
 num_decode,
  dropout = 0,
  ignore\_label = -1,
 bidirectional = F,
  loss_output = NULL,
  config,
  cell_type,
 masking = F,
 output_last_state = F,
  rnn.state = NULL,
  rnn.state.cell = NULL,
 prefix = ""
)
```

## Arguments

num\_rnn\_layer int, number of stacked layers
input\_size int, number of levels in the data - only used for embedding
num\_embed int, default = NULL - no embedding. Dimension of the embedding vectors
num\_hidden int, size of the state in each RNN layer
num\_decode int, number of output variables in the decoding layer
dropout
config Either seq-to-one or one-to-one
cell\_type Type of RNN cell: either gru or lstm

rnn.graph.unroll 527

rnn.graph.unroll

Unroll representation of RNN running on non CUDA device

# Description

Unroll representation of RNN running on non CUDA device

# Usage

```
rnn.graph.unroll(
  num_rnn_layer,
  seq_len,
  input_size = NULL,
 num_embed = NULL,
 num_hidden,
 num_decode,
 dropout = 0,
  ignore\_label = -1,
  loss_output = NULL,
  init.state = NULL,
 config,
 cell_type = "lstm",
 masking = F,
 output_last_state = F,
 prefix = "",
 data_name = "data",
  label_name = "label"
)
```

# Arguments

num_rnn_layer	int, number of stacked layers
seq_len	int, number of time steps to unroll
input_size	int, number of levels in the data - only used for embedding
num_embed	int, default = NULL - no embedding. Dimension of the embedding vectors
num_hidden	int, size of the state in each RNN layer
num_decode	int, number of output variables in the decoding layer
dropout	
config	Either seq-to-one or one-to-one
cell_type	Type of RNN cell: either gru or lstm

# **Index**

```
*Topic datasets
                                                mx.exec.update.arg.arrays, 26
    mx.metric.accuracy, 58
                                                mx.exec.update.aux.arrays, 26
    mx.metric.logistic_acc, 58
                                                mx.exec.update.grad.arrays, 27
    mx.metric.logloss, 59
                                                mx.gpu, 27
    mx.metric.mae, 59
                                                mx.infer.rnn, 28
    mx.metric.mse, 59
                                                mx.infer.rnn.one, 28
    mx.metric.Perplexity, 60
                                                mx.infer.rnn.one.unroll, 29
    mx.metric.rmse, 60
                                                mx.init.create, 29
    mx.metric.rmsle, 60
                                                mx.init.internal.default, 30
    mx.metric.top_k_accuracy, 61
                                                mx.init.normal, 30
                                                mx.init.uniform, 30
arguments, 15
                                                mx.init.Xavier, 31
as.array.MXNDArray, 16
                                                mx.io.arrayiter, 31
as.matrix.MXNDArray, 16
                                                mx.io.bucket.iter, 32
                                                mx.io.CSVIter, 32
children, 16
                                                mx.io.extract, 34
ctx, 17
                                                mx.io.ImageDetRecordIter, 34
                                                mx.io.ImageRecordInt8Iter, 38
dim. MXNDArray, 17
                                                mx.io.ImageRecordIter, 41
graph.viz, 17
                                                mx.io.ImageRecordIter_v1, 44
                                                mx.io.ImageRecordUInt8Iter, 47
im2rec. 18
                                                mx.io.ImageRecordUInt8Iter_v1, 50
internals, 19
                                                mx.io.LibSVMIter, 53
is.mx.context, 20
                                                mx.io.MNISTIter, 55
is.mx.dataiter, 20
                                                mx.kv.create, 56
is.mx.ndarray, 20
                                                mx.lr_scheduler.FactorScheduler, 56
is.mx.symbol, 21
                                                mx.lr_scheduler.MultiFactorScheduler,
is.serialized, 21
                                                mx.metric.accuracy, 58
length.MXNDArray, 22
                                                mx.metric.custom, 58
                                                mx.metric.logistic_acc, 58
mx.apply, 22
                                                {\tt mx.metric.logloss}, {\tt 59}
mx.callback.early.stop, 23
                                                mx.metric.mae, 59
mx.callback.log.speedometer, 23
                                                mx.metric.mse, 59
mx.callback.log.train.metric, 24
                                                mx.metric.Perplexity, 60
mx.callback.save.checkpoint, 24
                                                mx.metric.rmse, 60
mx.cpu, 25
mx.ctx.default, 25
                                                mx.metric.rmsle, 60
mx.exec.backward, 25
                                                mx.metric.top_k_accuracy, 61
mx.exec.forward, 26
                                                mx.mlp, 61
```

mx.model.buckets, 62	$\verb mx.nd.broadcast.not.equal , 95$
mx.model.FeedForward.create, 63	mx.nd.broadcast.plus, 95
mx.model.init.params, 65	mx.nd.broadcast.power,96
mx.model.load, 65	mx.nd.broadcast.sub,97
mx.model.save, 66	mx.nd.broadcast.to,97
mx.nd.abs, 66	mx.nd.Cast, 98
mx.nd.Activation, 67	mx.nd.cast, 99
mx.nd.adam.update, 67	mx.nd.cast.storage, 99
mx.nd.add.n, 68	mx.nd.cbrt, 100
mx.nd.all.finite, 69	mx.nd.ceil, 101
mx.nd.amp.cast, 69	mx.nd.choose.element.0index, 101
mx.nd.amp.multicast, 70	mx.nd.clip, 102
mx.nd.arccos, 70	mx.nd.col2im, 103
mx.nd.arccosh, 71	mx.nd.Concat, 104
mx.nd.arcsin, 71	mx.nd.concat, 104
mx.nd.arcsinh, 72	mx.nd.Convolution, 105
mx.nd.arctan, 72	mx.nd.Convolution.v1, 107
mx.nd.arctanh, 73	mx.nd.copyto, 108
mx.nd.argmax, 73	mx.nd.Correlation, 109
mx.nd.argmax.channel, 74	mx.nd.cos, 110
mx.nd.argmin, 75	mx.nd.cosh, 110
mx.nd.argsort, 75	mx.nd.Crop, 111
mx.nd.array, 76	mx.nd.crop, 112
mx.nd.batch.dot, 77	mx.nd.ctc.loss, 113
mx.nd.batch.take, 78	mx.nd.CTCLoss, 114
mx.nd.BatchNorm, 78	mx.nd.cumsum, 116
mx.nd.BatchNorm.v1, 80	mx.nd.Custom, 116
mx.nd.BilinearSampler, 81	mx.nd.Deconvolution, 117
mx.nd.BlockGrad, 82	mx.nd.degrees, 118
mx.nd.broadcast.add, 83	mx.nd.depth.to.space, 119
mx.nd.broadcast.axes, 84	mx.nd.diag, 119
mx.nd.broadcast.axis, 84	mx.nd.digamma, 120
mx.nd.broadcast.div, 85	mx.nd.dot, 121
mx.nd.broadcast.equal, 86	mx.nd.Dropout, 122
mx.nd.broadcast.greater, 86	mx.nd.ElementWiseSum, 123
mx.nd.broadcast.greater.equal, 87	mx.nd.elemwise.add, 123
mx.nd.broadcast.hypot, 87	mx.nd.elemwise.div, 124
mx.nd.broadcast.lesser, 88	mx.nd.elemwise.mul, 124
mx.nd.broadcast.lesser.equal, 89	mx.nd.elemwise.sub, 125
mx.nd.broadcast.like, 89	mx.nd.Embedding, 125
mx.nd.broadcast.logical.and, 90	mx.nd.erf, 126
mx.nd.broadcast.logical.or, 91	mx.nd.erfinv, 127
mx.nd.broadcast.logical.xor, 91	mx.nd.exp, 127
mx.nd.broadcast.maximum, 92	mx.nd.exp, 127 mx.nd.expand.dims, 128
mx.nd.broadcast.minimum, 92	mx.nd.expm1, 128
mx.nd.broadcast.minus, 93	mx.nd.fill.element.0index, 129
mx.nd.broadcast.mod, 94	mx.nd.fix, 129
mx.nd.broadcast.mul, 94	mx.nd.Flatten, 130
IIIA. Hu. DI Daucast. IIIuI, 74	IIIA. 114.1 TALLEII, 130

mx.nd.flatten, 130	mx.nd.MAERegressionOutput, 166
mx.nd.flip, 131	mx.nd.make.loss, 167
mx.nd.floor, 131	mx.nd.MakeLoss, 168
mx.nd.ftml.update, 132	mx.nd.max, 169
mx.nd.ftrl.update, 133	mx.nd.max.axis, 169
mx.nd.FullyConnected, 134	mx.nd.mean, 170
mx.nd.gamma, 135	mx.nd.min, 171
mx.nd.gammaln, 135	mx.nd.min.axis, 171
mx.nd.gather.nd, 136	mx.nd.moments, 172
mx.nd.GridGenerator, 136	mx.nd.mp.lamb.update.phase1, 173
mx.nd.GroupNorm, 137	mx.nd.mp.lamb.update.phase2, 174
mx.nd.hard.sigmoid, 138	mx.nd.mp.nag.mom.update, 175
mx.nd.identity, 138	mx.nd.mp.sgd.mom.update, 175
mx.nd.IdentityAttachKLSparseReg, 139	mx.nd.mp.sgd.update, 176
mx.nd.im2col, 139	mx.nd.multi.all.finite, 177
mx.nd.InstanceNorm, 140	mx.nd.multi.lars, 177
mx.nd.khatri.rao, 141	mx.nd.multi.mp.sgd.mom.update, 178
mx.nd.L2Normalization, 142	mx.nd.multi.mp.sgd.update, 179
mx.nd.lamb.update.phase1, 143	mx.nd.multi.sgd.mom.update, 179
mx.nd.lamb.update.phase2, 144	mx.nd.multi.sgd.update, 180
mx.nd.LayerNorm, 144	mx.nd.multi.sum.sq, 181
mx.nd.LeakyReLU, 145	mx.nd.nag.mom.update, 181
mx.nd.linalg.det, 146	mx.nd.nanprod, 182
mx.nd.linalg.extractdiag, 147	mx.nd.nansum, 183
mx.nd.linalg.extracttrian, 148	mx.nd.negative, 184
mx.nd.linalg.gelqf, 149	mx.nd.norm, 184
mx.nd.linalg.gemm, 150	mx.nd.normal, 185
mx.nd.linalg.gemm2, 151	mx.nd.one.hot, 186
mx.nd.linalg.inverse, 152	mx.nd.ones, 187
mx.nd.linalg.makediag, 153	mx.nd.ones.like, 187
mx.nd.linalg.maketrian, 153	mx.nd.Pad, 188
mx.nd.linalg.potrf, 154	mx.nd.pad, 189
mx.nd.linalg.potri, 155	mx.nd.pick, 190
mx.nd.linalg.slogdet, 156	mx.nd.Pooling, 191
mx.nd.linalg.sumlogdiag, 157	mx.nd.Pooling.v1, 193
mx.nd.linalg.syrk, 157	<pre>mx.nd.preloaded.multi.mp.sgd.mom.update,</pre>
mx.nd.linalg.trmm, 158	194
mx.nd.linalg.trsm, 159	<pre>mx.nd.preloaded.multi.mp.sgd.update,</pre>
mx.nd.LinearRegressionOutput, 160	195
mx.nd.load, 161	<pre>mx.nd.preloaded.multi.sgd.mom.update,</pre>
mx.nd.log, 161	195
mx.nd.log.softmax, 162	mx.nd.preloaded.multi.sgd.update, 196
mx.nd.log10, 163	mx.nd.prod, 197
mx.nd.log1p, 163	mx.nd.radians, 197
mx.nd.log2, 164	mx.nd.random.exponential, 198
mx.nd.logical.not, 164	mx.nd.random.gamma, 199
mx.nd.LogisticRegressionOutput, 165	<pre>mx.nd.random.generalized.negative.binomial,</pre>
mx.nd.LRN, 166	199

mx.nd.random.negative.binomial, 200	mx.nd.shape.array, 240
mx.nd.random.normal, 201	mx.nd.shuffle, 240
mx.nd.random.pdf.dirichlet, 202	mx.nd.sigmoid, 241
mx.nd.random.pdf.exponential, 202	mx.nd.sign, 241
mx.nd.random.pdf.gamma, 203	mx.nd.signsgd.update, 242
mx.nd.random.pdf.generalized.negative.binomi	
204	
	mx.nd.sin, 243
mx.nd.random.pdf.negative.binomial,	mx.nd.sinh, 244
	mx.nd.size.array, 244
mx.nd.random.pdf.normal, 206	mx.nd.slice.axis, 245
mx.nd.random.pdf.poisson, 207	mx.nd.slice.like, 246
mx.nd.random.pdf.uniform, 207	mx.nd.SliceChannel, 247
mx.nd.random.poisson, 208	mx.nd.smooth.11, 248
mx.nd.random.randint, 209	mx.nd.Softmax, 248
mx.nd.random.uniform, 209	mx.nd.softmax, 250
mx.nd.ravel.multi.index, 210	mx.nd.softmax.cross.entropy, 251
mx.nd.rcbrt, 211	mx.nd.SoftmaxActivation, 252
mx.nd.reciprocal, 211	mx.nd.SoftmaxOutput, 253
mx.nd.relu,212	mx.nd.softmin, 254
mx.nd.repeat, 212	mx.nd.softsign, 255
mx.nd.reset.arrays, 213	mx.nd.sort, 256
mx.nd.Reshape, 213	mx.nd.space.to.depth, 256
mx.nd.reshape, 215	mx.nd.SpatialTransformer, 257
mx.nd.reshape.like,216	mx.nd.split, 258
mx.nd.reverse, 217	mx.nd.sqrt, 259
mx.nd.rint, 218	mx.nd.square, 259
mx.nd.rmsprop.update, 218	mx.nd.squeeze, 260
mx.nd.rmspropalex.update, 219	mx.nd.stack, 260
mx.nd.RNN, 221	mx.nd.stop.gradient, 261
mx.nd.ROIPooling, 223	mx.nd.sum, 262
mx.nd.round, 224	mx.nd.sum.axis, 263
mx.nd.rsqrt, 224	mx.nd.SVMOutput, 264
mx.nd.sample.exponential, 225	mx.nd.swapaxes, 264
mx.nd.sample.gamma, 226	mx.nd.SwapAxis, 265
<pre>mx.nd.sample.generalized.negative.binomial,</pre>	mx.nd.take, 265
227	mx.nd.tan, 266
mx.nd.sample.multinomial, 228	mx.nd.tanh, 267
mx.nd.sample.negative.binomial, 229	mx.nd.tile, 268
mx.nd.sample.normal, 230	mx.nd.topk, 268
mx.nd.sample.poisson, 231	mx.nd.transpose, 269
mx.nd.sample.uniform, 232	mx.nd.trunc, 270
mx.nd.save, 233	mx.nd.uniform, 271
mx.nd.scatter.nd, 233	mx.nd.unravel.index, 271
mx.nd.SequenceLast, 234	mx.nd.UpSampling, 272
mx.nd.SequenceMask, 235	mx.nd.where, 273
mx.nd.SequenceReverse, 236	mx.nd.zeros, 274
mx.nd.sgd.mom.update, 238	mx.nd.zeros.like, 275
mx.nd.sgd.update, 239	mx.opt.adadelta, 275
· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·

mx.opt.adagrad, 276	mx.symbol.broadcast_logical_and,312
mx.opt.adam, 277	mx.symbol.broadcast_logical_or,312
mx.opt.create, 277	mx.symbol.broadcast_logical_xor, 313
mx.opt.get.updater, 278	mx.symbol.broadcast_maximum, 314
mx.opt.nag, 278	mx.symbol.broadcast_minimum, 314
mx.opt.rmsprop, 279	mx.symbol.broadcast_minus, 315
mx.opt.sgd, 280	mx.symbol.broadcast_mod, 316
mx.profiler.config, 281	mx.symbol.broadcast_mul, 317
mx.profiler.state, 281	<pre>mx.symbol.broadcast_not_equal, 317</pre>
mx.rnorm, 282	mx.symbol.broadcast_plus, 318
mx.runif, 282	mx.symbol.broadcast_power,319
mx.serialize, 283	mx.symbol.broadcast_sub, 320
mx.set.seed, 283	<pre>mx.symbol.broadcast_to, 321</pre>
mx.simple.bind, 284	mx.symbol.Cast,322
mx.symbol.abs, 284	mx.symbol.cast, 322
mx.symbol.Activation, 285	mx.symbol.cast_storage, 323
mx.symbol.adam_update, 285	mx.symbol.cbrt, 324
mx.symbol.add_n, 287	mx.symbol.ceil, 324
mx.symbol.all_finite, 287	mx.symbol.choose_element_0index, 325
mx.symbol.amp_cast, 288	mx.symbol.clip, 326
mx.symbol.amp_multicast, 288	mx.symbol.col2im, 327
mx.symbol.arccos, 289	mx.symbol.Concat, 328
mx.symbol.arccosh, 290	mx.symbol.concat, 329
mx.symbol.arcsin, 290	mx.symbol.Convolution, 329
mx.symbol.arcsinh, 291	mx.symbol.Convolution_v1, 331
mx.symbol.arctan, 292	mx.symbol.Correlation, 332
mx.symbol.arctanh, 292	mx.symbol.cos, 334
mx.symbol.argmax, 293	mx.symbol.cosh, 334
mx.symbol.argmax_channel, 294	mx.symbol.Crop, 335
mx.symbol.argmin, 294	mx.symbol.crop, 336
mx.symbol.argsort, 295	mx.symbol.ctc_loss, 338
mx.symbol.drgsort, 299	mx.symbol.CTCLoss, 337
mx.symbol.batch_take, 300	mx.symbol.cumsum, 340
mx.symbol.BatchNorm, 296	mx.symbol.Custom, 340
mx.symbol.BatchNorm_v1, 298	mx.symbol.Deconvolution, 341
-	mx.symbol.degrees, 342
mx.symbol.BilinearSampler, 301	
mx.symbol.BlockGrad, 302	mx.symbol.depth_to_space, 343
mx.symbol.broadcast_add, 303	mx.symbol.diag, 344
mx.symbol.broadcast_axes, 304	mx.symbol.digamma, 345
mx.symbol.broadcast_axis, 305	mx.symbol.dot, 346
mx.symbol.broadcast_div, 306	mx.symbol.Dropout, 347
mx.symbol.broadcast_equal, 306	mx.symbol.ElementWiseSum, 348
mx.symbol.broadcast_greater, 307	mx.symbol.elemwise_add, 349
mx.symbol.broadcast_greater_equal, 308	mx.symbol.elemwise_div, 349
mx.symbol.broadcast_hypot, 308	mx.symbol.elemwise_mul, 350
mx.symbol.broadcast_lesser,309	mx.symbol.elemwise_sub, 350
mx.symbol.broadcast_lesser_equal, 310	mx.symbol.Embedding, 351
mx.symbol.broadcast_like, 311	mx.symbol.erf, 352

mx.symbol.erfinv, 353	mx.symbol.LinearRegressionOutput, 393
mx.symbol.exp, 354	mx.symbol.load, 394
mx.symbol.expand_dims, 354	mx.symbol.load.json, 394
mx.symbol.expm1,355	mx.symbol.log,395
<pre>mx.symbol.fill_element_0index, 356</pre>	mx.symbol.log10,395
mx.symbol.fix,356	mx.symbol.log1p,396
mx.symbol.Flatten, 357	mx.symbol.log2,396
mx.symbol.flatten, 358	mx.symbol.log_softmax,398
mx.symbol.flip, 358	mx.symbol.logical_not,397
mx.symbol.floor, 359	<pre>mx.symbol.LogisticRegressionOutput,</pre>
<pre>mx.symbol.ftml_update, 360</pre>	397
mx.symbol.ftrl_update, 361	mx.symbol.LRN, 399
mx.symbol.FullyConnected, 362	mx.symbol.MAERegressionOutput,400
mx.symbol.gamma, 363	mx.symbol.make_loss, 402
mx.symbol.gammaln, 363	mx.symbol.MakeLoss, 401
mx.symbol.gather_nd,364	mx.symbol.max, 402
mx.symbol.GridGenerator, 364	mx.symbol.max_axis, 403
mx.symbol.Group, 365	mx.symbol.mean,404
mx.symbol.GroupNorm, 366	mx.symbol.moments, 405
mx.symbol.hard_sigmoid, 367	mx.symbol.mp_lamb_update_phase1,405
mx.symbol.identity, 367	mx.symbol.mp_lamb_update_phase2,407
mx.symbol.IdentityAttachKLSparseReg,	mx.symbol.mp_nag_mom_update, 408
368	mx.symbol.mp_sgd_mom_update, 409
mx.symbol.im2col, 368	mx.symbol.mp_sgd_update, 410
mx.symbol.infer.shape, 369	mx.symbol.multi_all_finite, 410
mx.symbol.InstanceNorm, 370	mx.symbol.multi_lars, 411
mx.symbol.khatri_rao,371	mx.symbol.multi_mp_sgd_mom_update, 412
mx.symbol.L2Normalization, 372	mx.symbol.multi_mp_sgd_update, 413
mx.symbol.lamb_update_phase1,373	mx.symbol.multi_sgd_mom_update, 414
mx.symbol.lamb_update_phase2, 374	mx.symbol.multi_sgd_update, 415
mx.symbol.LayerNorm, 375	mx.symbol.multi_sum_sq, 416
mx.symbol.LeakyReLU, 376	mx.symbol.nag_mom_update, 416
mx.symbol.linalg_det, 377	mx.symbol.nanprod, 417
mx.symbol.linalg_extractdiag, 378	mx.symbol.nansum, 418
mx.symbol.linalg_extracttrian, 379	mx.symbol.negative, 419
mx.symbol.linalg_gelqf, 380	mx.symbol.norm, 419
mx.symbol.linalg_gemm, 381	mx.symbol.normal, 420
mx.symbol.linalg_gemm2, 382	mx.symbol.one_hot, 422
mx.symbol.linalg_inverse, 383	mx.symbol.ones_like, 421
mx.symbol.linalg_makediag, 384	mx.symbol.Pad, 423
mx.symbol.linalg_maketrian, 385	mx.symbol.pad, 424
mx.symbol.linalg_potrf, 386	mx.symbol.pick, 425
mx.symbol.linalg_potri, 387	mx.symbol.Pooling, 427
mx.symbol.linalg_slogdet, 388	mx.symbol.Pooling_v1, 428
mx.symbol.linalg_sumlogdiag, 389	mx.symbol.preloaded_multi_mp_sgd_mom_update
mx.symbol.linalg_syrk, 390	430
mx.symbol.linalg_trmm, 391	mx.symbol.preloaded_multi_mp_sgd_update,
mx.symbol.linalg_trmm, 391	431
IIIA. SYIIIDUI. IIIIaig_ti SIII, 394	431

```
mx.symbol.preloaded_multi_sgd_mom_update,
                                               mx.symbol.sample_negative_binomial,
mx.symbol.preloaded_multi_sgd_update,
                                               mx.symbol.sample_normal, 471
        432
                                               mx.symbol.sample_poisson, 472
mx.symbol.prod, 433
                                               mx.symbol.sample_uniform, 473
mx.symbol.radians, 434
                                               mx.symbol.save, 474
mx.symbol.random_exponential, 434
                                               mx.symbol.scatter_nd, 474
mx.symbol.random_gamma, 435
                                               mx.symbol.SequenceLast, 475
mx.symbol.random_generalized_negative_binomianlx.symbol.SequenceMask, 476
                                               mx.symbol.SequenceReverse, 478
mx.symbol.random_negative_binomial,
                                               mx.symbol.sgd_mom_update, 479
        437
                                               mx.symbol.sgd_update, 480
mx.symbol.random_normal, 438
                                               mx.symbol.shape_array, 481
mx.symbol.random_pdf_dirichlet, 439
                                               mx.symbol.shuffle, 482
mx.symbol.random_pdf_exponential, 440
                                               mx.symbol.sigmoid, 482
mx.symbol.random_pdf_gamma, 441
                                               mx.symbol.sign, 483
mx.symbol.random_pdf_generalized_negative_binomis/lmbol.signsgd_update, 483
                                               \verb|mx.symbol.signum_update|, 484|
mx.symbol.random_pdf_negative_binomial,
                                               mx.symbol.sin, 485
        443
                                               mx.symbol.sinh, 486
mx.symbol.random_pdf_normal, 444
                                               mx.symbol.size_array, 487
mx.symbol.random_pdf_poisson, 445
                                               mx.symbol.slice, 487
mx.symbol.random_pdf_uniform, 446
                                               mx.symbol.slice_axis, 490
mx.symbol.random_poisson, 447
                                               mx.symbol.slice_like, 491
mx.symbol.random_randint, 447
                                               mx.symbol.SliceChannel, 489
mx.symbol.random_uniform, 448
                                               mx.symbol.smooth_11,492
mx.symbol.ravel_multi_index, 449
                                               mx.symbol.Softmax, 492
mx.symbol.rcbrt, 450
                                               mx.symbol.softmax, 494
mx.symbol.reciprocal, 450
                                               mx.symbol.softmax_cross_entropy, 498
mx.symbol.relu,451
                                               mx.symbol.SoftmaxActivation, 495
mx.symbol.repeat, 452
                                               mx.symbol.SoftmaxOutput, 496
mx.symbol.reset_arrays, 452
                                               mx.symbol.softmin, 499
mx.symbol.Reshape, 453
                                               mx.symbol.softsign, 500
mx.symbol.reshape, 454
                                               mx.symbol.sort, 500
mx.symbol.reshape_like, 456
                                               mx.symbol.space_to_depth, 501
mx.symbol.reverse, 457
                                               mx.symbol.SpatialTransformer, 502
mx.symbol.rint, 458
                                               mx.symbol.split, 503
mx.symbol.rmsprop_update, 460
                                               mx.symbol.sqrt, 504
mx.symbol.rmspropalex_update, 458
                                               mx.symbol.square, 505
mx.symbol.RNN, 461
                                               mx.symbol.squeeze, 505
mx.symbol.ROIPooling, 463
                                               mx.symbol.stack, 506
mx.symbol.round, 464
                                               mx.symbol.stop_gradient, 507
mx.symbol.rsqrt, 465
                                               mx.symbol.sum, 508
                                               mx.symbol.sum_axis, 509
mx.symbol.sample_exponential, 466
mx.symbol.sample_gamma, 467
                                               mx.symbol.SVMOutput, 510
mx.symbol.sample_generalized_negative_binomianlx.symbol.swapaxes, 511
                                               mx.symbol.SwapAxis, 511
mx.symbol.sample_multinomial, 469
                                               mx.symbol.take, 512
```

```
mx.symbol.tan, 513
mx.symbol.tanh, 514
mx.symbol.tile, 515
mx.symbol.topk, 515
mx.symbol.transpose, 517
mx.symbol.trunc, 517
mx.symbol.uniform, 518
mx.symbol.unravel_index, 519
mx.symbol.UpSampling, 519
mx.symbol.Variable, 521
{\tt mx.symbol.where}, {\tt 521}
mx.symbol.zeros_like, 522
mx.unserialize, 523
mxnet, 523
mxnet.export, 523
Ops.MXNDArray, 524
outputs, 524
predict.MXFeedForwardModel, 524
print.MXNDArray, 525
rnn.graph, 526
rnn.graph.unroll, 527
```