# Package 'torch'

May 22, 2024

Type Package

Title Tensors and Neural Networks with 'GPU' Acceleration

**Version** 0.13.0

**Description** Provides functionality to define and train neural networks similar to 'PyTorch' by Paszke et al (2019) <doi:10.48550/arXiv.1912.01703> but written entirely in R using the 'libtorch' library. Also supports low-level tensor operations and 'GPU' acceleration.

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URL https://torch.mlverse.org/docs, https://github.com/mlverse/torch

BugReports https://github.com/mlverse/torch/issues

**Encoding** UTF-8

**SystemRequirements** LibTorch (https://pytorch.org/); Only x86\_64 platforms are currently supported except for ARM system running macOS.

Config/build/copy-method copy

LinkingTo Rcpp

**Imports** Rcpp, R6, withr, rlang, methods, utils, stats, bit64, magrittr, tools, coro (>= 1.0.2), callr, cli (>= 3.0.0), glue, ellipsis, desc, safetensors (>= 0.1.1), jsonlite

RoxygenNote 7.3.1

**Suggests** testthat (>= 3.0.0), covr, knitr (>= 1.36), rmarkdown, palmerpenguins, mvtnorm, numDeriv, katex

VignetteBuilder knitr

Collate 'R7.R' 'RcppExports.R' 'autocast.R' 'tensor.R' 'autograd.R' 'backends.R' 'call\_torch\_function.R' 'codegen-utils.R' 'compare.R' 'compat-purrr.R' 'compilation\_unit.R' 'conditions.R' 'contition.R' 'creation-ops.R' 'cuda.R' 'device.R' 'dimname\_list.R' 'utils.R' 'distributions-constraints.R' 'distributions-utils.R' 'distributions-exp-family.R' 'distributions.R' 'distributions-bernoulli.R' 'distributions-categorical.R' 'distributions-gamma.R'

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'with-indices.R' 'wrapers.R'
NeedsCompilation yes
Author Daniel Falbel [aut, cre, cph], Javier Luraschi [aut], Dmitriy Selivanov [ctb], Athos Damiani [ctb], Christophe Regouby [ctb], Krzysztof Joachimiak [ctb], Hamada S. Badr [ctb], Sebastian Fischer [ctb], RStudio [cph]
Maintainer Daniel Falbel <daniel@rstudio.com></daniel@rstudio.com>
Repository CRAN
<b>Date/Publication</b> 2024-05-21 22:30:03 UTC
R topics documented:

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as_array				
with_enable_	anomaly			
Index 584				
as_array	Converts to array			
Description				
Converts to array				
Usage				
as_array(x)				
Arguments				
х	object to be converted into an array			
AutogradContext	Class representing the context.			

# Description

Class representing the context.

Class representing the context.

## **Public fields**

ptr (Dev related) pointer to the context c++ object.

# **Active bindings**

needs\_input\_grad boolean listing arguments of forward and whether they require\_grad. saved\_variables list of objects that were saved for backward via save\_for\_backward.

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#### Methods

#### **Public methods:**

```
AutogradContext$new()
```

- AutogradContext\$save\_for\_backward()
- AutogradContext\$mark\_non\_differentiable()
- AutogradContext\$mark\_dirty()
- AutogradContext\$clone()

**Method** new(): (Dev related) Initializes the context. Not user related.

```
Usage:
AutogradContext$new(
   ptr,
   env,
   argument_names = NULL,
   argument_needs_grad = NULL
)

Arguments:
ptr pointer to the c++ object
env environment that encloses both forward and backward
argument_names names of forward arguments
argument_needs_grad whether each argument in forward needs grad.
```

**Method** save\_for\_backward(): Saves given objects for a future call to backward().

This should be called at most once, and only from inside the forward() method.

Later, saved objects can be accessed through the saved\_variables attribute. Before returning them to the user, a check is made to ensure they weren't used in any in-place operation that modified their content.

Arguments can also be any kind of R object.

```
Usage:
AutogradContext$save_for_backward(...)
Arguments:
```

... any kind of R object that will be saved for the backward pass. It's common to pass named arguments.

**Method** mark\_non\_differentiable(): Marks outputs as non-differentiable.

This should be called at most once, only from inside the forward() method, and all arguments should be outputs.

This will mark outputs as not requiring gradients, increasing the efficiency of backward computation. You still need to accept a gradient for each output in backward(), but it's always going to be a zero tensor with the same shape as the shape of a corresponding output.

This is used e.g. for indices returned from a max Function.

```
Usage:
AutogradContext$mark_non_differentiable(...)
```

autograd\_backward 19

```
Arguments:
```

```
... non-differentiable outputs.
```

Method mark\_dirty(): Marks given tensors as modified in an in-place operation.

This should be called at most once, only from inside the forward() method, and all arguments should be inputs.

Every tensor that's been modified in-place in a call to forward() should be given to this function, to ensure correctness of our checks. It doesn't matter whether the function is called before or after modification.

```
Usage:
AutogradContext$mark_dirty(...)
Arguments:
... tensors that are modified in-place.
```

**Method** clone(): The objects of this class are cloneable with this method.

Usage:

AutogradContext\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

autograd\_backward

Computes the sum of gradients of given tensors w.r.t. graph leaves.

### **Description**

The graph is differentiated using the chain rule. If any of tensors are non-scalar (i.e. their data has more than one element) and require gradient, then the Jacobian-vector product would be computed, in this case the function additionally requires specifying grad\_tensors. It should be a sequence of matching length, that contains the "vector" in the Jacobian-vector product, usually the gradient of the differentiated function w.r.t. corresponding tensors (None is an acceptable value for all tensors that don't need gradient tensors).

#### Usage

```
autograd_backward(
  tensors,
  grad_tensors = NULL,
  retain_graph = create_graph,
  create_graph = FALSE
)
```

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## Arguments

tensors (list of Tensor) – Tensors of which the derivative will be computed.

grad\_tensors (list of (Tensor or NULL)) The "vector" in the Jacobian-vector product, usually gradients w. value would be acceptable for all grad\_tensors, then this argument is optional.

retain\_graph (bool, optional) – If FALSE, the graph used to compute the grad will be freed. Note that in nearly all cases setting this option to TRUE is not needed and often can be worked around in a much more efficient way. Defaults to the value of create\_graph.

create\_graph (bool, optional) – If TRUE, graph of the derivative will be constructed, allowing to compute higher order derivative products. Defaults to FALSE.

#### **Details**

This function accumulates gradients in the leaves - you might need to zero them before calling it.

## **Examples**

```
if (torch_is_installed()) {
x <- torch_tensor(1, requires_grad = TRUE)
y <- 2 * x

a <- torch_tensor(1, requires_grad = TRUE)
b <- 3 * a

autograd_backward(list(y, b))
}</pre>
```

autograd\_function

Records operation history and defines formulas for differentiating ops.

#### **Description**

Every operation performed on Tensor's creates a new function object, that performs the computation, and records that it happened. The history is retained in the form of a DAG of functions, with edges denoting data dependencies (input <- output). Then, when backward is called, the graph is processed in the topological ordering, by calling backward() methods of each Function object, and passing returned gradients on to next Function's.

#### **Usage**

```
autograd_function(forward, backward)
```

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## Arguments

forward

Performs the operation. It must accept a context ctx as the first argument, followed by any number of arguments (tensors or other types). The context can be used to store tensors that can be then retrieved during the backward pass. See AutogradContext for more information about context methods.

backward

Defines a formula for differentiating the operation. It must accept a context ctx as the first argument, followed by as many outputs ad forward() returned (as a list()). The names of the arguments don't matter and they are passed in the order in which they were returned by forward(). The function should return a named list, where each argument is the gradient w.r.t the given output, and each element in the returned list should be the gradient w.r.t. the corresponding input. The context can be used to retrieve tensors saved during the forward pass. It also has an attribute ctx\$needs\_input\_grad as a named list of booleans representing whether each input needs gradient. E.g., backward() will have ctx\$needs\_input\_grad\$input = TRUE if the input argument to forward() needs gradient computated w.r.t. the output. See AutogradContext for more information about context methods.

## **Examples**

```
if (torch_is_installed()) {
exp2 <- autograd_function(
  forward = function(ctx, i) {
    result <- i$exp()
    ctx$save_for_backward(result = result)
    result
},
backward = function(ctx, grad_output) {
    list(i = grad_output * ctx$saved_variable$result)
}
)
}</pre>
```

autograd\_grad

Computes and returns the sum of gradients of outputs w.r.t. the inputs.

#### **Description**

grad\_outputs should be a list of length matching output containing the "vector" in Jacobian-vector product, usually the pre-computed gradients w.r.t. each of the outputs. If an output doesn't require\_grad, then the gradient can be None).

#### Usage

```
autograd_grad(
  outputs,
```

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```
inputs,
  grad_outputs = NULL,
  retain_graph = create_graph,
  create_graph = FALSE,
  allow_unused = FALSE
)
```

#### **Arguments**

outputs (sequence of Tensor) – outputs of the differentiated function.

inputs (sequence of Tensor) – Inputs w.r.t. which the gradient will be returned (and not

accumulated into .grad).

grad\_outputs (sequence of Tensor) - The "vector" in the Jacobian-vector product. Usually

gradients w.r.t. each output. None values can be specified for scalar Tensors or ones that don't require grad. If a None value would be acceptable for all

grad\_tensors, then this argument is optional. Default: None.

retain\_graph (bool, optional) - If FALSE, the graph used to compute the grad will be freed.

Note that in nearly all cases setting this option to TRUE is not needed and often can be worked around in a much more efficient way. Defaults to the value of

create\_graph.

 $create\_graph \qquad (bool, optional) - If \, \mathsf{TRUE}, \, \, \mathsf{graph} \, \, \mathsf{of} \, \, \mathsf{the} \, \, \mathsf{derivative} \, \, \mathsf{will} \, \, \mathsf{be} \, \, \mathsf{constructed}, \, \, \mathsf{allowing} \, \, \mathsf{to} \, \, \mathsf{compute} \, \, \mathsf{log} \,$ 

allow\_unused (bool, optional) – If FALSE, specifying inputs that were not used when computing

outputs (and therefore their grad is always zero) is an error. Defaults to FALSE

## Details

If only\_inputs is TRUE, the function will only return a list of gradients w.r.t the specified inputs. If it's FALSE, then gradient w.r.t. all remaining leaves will still be computed, and will be accumulated into their . grad attribute.

#### **Examples**

```
if (torch_is_installed()) {
w <- torch_tensor(0.5, requires_grad = TRUE)
b <- torch_tensor(0.9, requires_grad = TRUE)
x <- torch_tensor(runif(100))
y <- 2 * x + 1
loss <- (y - (w * x + b))^2
loss <- loss$mean()

o <- autograd_grad(loss, list(w, b))
o
}</pre>
```

```
\verb"autograd_set_grad_mode"
```

Set grad mode

# Description

Sets or disables gradient history.

# Usage

```
autograd_set_grad_mode(enabled)
```

# Arguments

enabled

bool wether to enable or disable the gradient recording.

```
backends_cudnn_is_available
```

CuDNN is available

# Description

CuDNN is available

# Usage

```
backends_cudnn_is_available()
```

backends\_cudnn\_version

CuDNN version

# Description

CuDNN version

# Usage

backends\_cudnn\_version()

backends\_mkldnn\_is\_available

MKLDNN is available

# Description

MKLDNN is available

# Usage

backends\_mkldnn\_is\_available()

## Value

Returns whether LibTorch is built with MKL-DNN support.

backends\_mkl\_is\_available

MKL is available

# Description

MKL is available

## Usage

backends\_mkl\_is\_available()

# Value

Returns whether LibTorch is built with MKL support.

backends\_mps\_is\_available

MPS is available

# Description

MPS is available

## Usage

backends\_mps\_is\_available()

#### Value

Returns whether LibTorch is built with MPS support.

backends\_openmp\_is\_available

OpenMP is available

#### **Description**

OpenMP is available

#### Usage

backends\_openmp\_is\_available()

#### Value

Returns whether LibTorch is built with OpenMP support.

broadcast\_all

Given a list of values (possibly containing numbers), returns a list where each value is broadcasted based on the following rules:

## Description

Raises value\_error: if any of the values is not a numeric instance, a torch.\*Tensor instance, or an instance implementing **torch\_function** TODO: add has\_torch\_function((v,)) See: https://github.com/pytorch/pytorch/blob/m

#### Usage

broadcast\_all(values)

### **Arguments**

values List of:

- torch.\*Tensor instances are broadcasted as per\_broadcasting-semantics.
- numeric instances (scalars) are upcast to tensors having the same size and type as the first tensor passed to values. If all the values are scalars, then they are upcasted to scalar Tensors. values (list of numeric, torch.\*Tensor or objects implementing torch\_function)

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clone\_module

Clone a torch module.

# Description

Clones a module.

## Usage

```
clone_module(module, deep = FALSE, ..., replace_values = TRUE)
```

## **Arguments**

```
module
(nn_module)
The module to clone

deep
(logical(1))
Whether to create a deep clone.
...
(any)
Additional parameters, currently unused.

replace_values
(logical(1))
Whether to replace parameters and buffers with the cloned values.
```

## **Examples**

```
if (torch_is_installed()) {
  clone_module(nn_linear(1, 1), deep = TRUE)
# is the same as
  nn_linear(1, 1)$clone(deep = TRUE)
}
```

Constraint

Abstract base class for constraints.

## **Description**

Abstract base class for constraints.

Abstract base class for constraints.

## **Details**

A constraint object represents a region over which a variable is valid, e.g. within which a variable can be optimized.

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## Methods

#### **Public methods:**

- Constraint\$check()
- Constraint\$print()
- Constraint\$clone()

**Method** check(): Returns a byte tensor of sample\_shape + batch\_shape indicating whether each event in value satisfies this constraint.

Usage:

Constraint\$check(value)

Arguments:

value each event in value will be checked.

Method print(): Define the print method for constraints,

Usage:

Constraint\$print()

**Method** clone(): The objects of this class are cloneable with this method.

Usage:

Constraint\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

contrib\_sort\_vertices Contrib sort vertices

# Description

Based on the implementation from Rotated\_IoU

## Usage

```
contrib_sort_vertices(vertices, mask, num_valid)
```

## Arguments

vertices A Tensor with the vertices.

mask A tensors containing the masks.

num\_valid A integer tensors.

#### **Details**

All tensors should be on a CUDA device so this function can be used.

#### Note

This function does not make part of the official torch API.

#### **Examples**

```
if (torch_is_installed()) {
  if (cuda_is_available()) {
    v <- torch_randn(8, 1024, 24, 2)$cuda()
    mean <- torch_mean(v, dim = 2, keepdim = TRUE)
    v <- v - mean
    m <- (torch_rand(8, 1024, 24) > 0.8)$cuda()
    nv <- torch_sum(m$to(dtype = torch_int()), dim = -1)$to(dtype = torch_int())$cuda()
    result <- contrib_sort_vertices(v, m, nv)
}
</pre>
```

#### **Description**

A gradient scaler instance is used to perform dynamic gradient scaling to avoid gradient underflow when training with mixed precision.

### Usage

```
cuda_amp_grad_scaler(
  init_scale = 2^16,
  growth_factor = 2,
  backoff_factor = 0.5,
  growth_interval = 2000,
  enabled = TRUE
)
```

## **Arguments**

```
init_scale a numeric value indicating the initial scale factor.

growth_factor a numeric value indicating the growth factor.

backoff_factor a numeric value indicating the backoff factor.

growth_interval a numeric value indicating the growth interval.

enabled a logical value indicating whether the gradient scaler should be enabled.
```

#### Value

A gradient scaler object.

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cuda\_current\_device

Returns the index of a currently selected device.

# Description

Returns the index of a currently selected device.

## Usage

```
cuda_current_device()
```

cuda\_device\_count

Returns the number of GPUs available.

# Description

Returns the number of GPUs available.

#### Usage

```
cuda_device_count()
```

cuda\_empty\_cache

Empty cache

# Description

Releases all unoccupied cached memory currently held by the caching allocator so that those can be used in other GPU application and visible in nvidia-smi.

## Usage

```
cuda_empty_cache()
```

#### Note

cuda\_empty\_cache() doesn't increase the amount of GPU memory available for torch. However, it may help reduce fragmentation of GPU memory in certain cases. See Memory management article for more details about GPU memory management.

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```
cuda_get_device_capability
```

Returns the major and minor CUDA capability of device

#### **Description**

Returns the major and minor CUDA capability of device

## Usage

```
cuda_get_device_capability(device = cuda_current_device())
```

# Arguments

device

Integer value of the CUDA device to return capabilities of.

cuda\_is\_available

Returns a bool indicating if CUDA is currently available.

## **Description**

Returns a bool indicating if CUDA is currently available.

## Usage

```
cuda_is_available()
```

cuda\_memory\_stats

Returns a dictionary of CUDA memory allocator statistics for a given device.

## **Description**

The return value of this function is a dictionary of statistics, each of which is a non-negative integer.

# Usage

```
cuda_memory_stats(device = cuda_current_device())
cuda_memory_summary(device = cuda_current_device())
```

### **Arguments**

device

Integer value of the CUDA device to return capabilities of.

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#### **Core statistics**

• "allocated.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": number of allocation requests received by the memory allocator.

- "allocated\_bytes.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": amount of allocated memory.
- "segment.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": number of reserved segments from cudaMalloc().
- "reserved\_bytes.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": amount of reserved memory.
- "active.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": number of active memory blocks.
- "active\_bytes.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": amount of active memory.
- "inactive\_split.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": number of inactive, non-releasable memory blocks.
- "inactive\_split\_bytes.{all,large\_pool,small\_pool}.{current,peak,allocated,freed}": amount of inactive, non-releasable memory.

For these core statistics, values are broken down as follows.

#### Pool type:

- all: combined statistics across all memory pools.
- large\_pool: statistics for the large allocation pool (as of October 2019, for size >= 1MB allocations).
- small\_pool: statistics for the small allocation pool (as of October 2019, for size < 1MB allocations).

## Metric type:

- current: current value of this metric.
- peak: maximum value of this metric.
- allocated: historical total increase in this metric.
- freed: historical total decrease in this metric.

#### **Additional metrics**

- "num\_alloc\_retries": number of failed cudaMalloc calls that result in a cache flush and retry.
- "num\_ooms": number of out-of-memory errors thrown.

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cuda\_runtime\_version

Returns the CUDA runtime version

## **Description**

Returns the CUDA runtime version

# Usage

```
cuda_runtime_version()
```

cuda\_synchronize

Waits for all kernels in all streams on a CUDA device to complete.

# Description

Waits for all kernels in all streams on a CUDA device to complete.

# Usage

```
cuda_synchronize(device = NULL)
```

## Arguments

device

device for which to synchronize. It uses the current device given by cuda\_current\_device() if no device is specified.

dataloader

Data loader. Combines a dataset and a sampler, and provides singleor multi-process iterators over the dataset.

# Description

Data loader. Combines a dataset and a sampler, and provides single- or multi-process iterators over the dataset.

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## Usage

```
dataloader(
  dataset,
  batch_size = 1,
  shuffle = FALSE,
  sampler = NULL,
  batch_sampler = NULL,
  num_workers = 0,
  collate_fn = NULL,
  pin_memory = FALSE,
  drop_last = FALSE,
  timeout = -1,
  worker_init_fn = NULL,
  worker_globals = NULL,
  worker_packages = NULL
)
```

# Arguments

dataset	(Dataset): dataset from which to load the data.
batch_size	(int, optional): how many samples per batch to load (default: 1).
shuffle	(bool, optional): set to TRUE to have the data reshuffled at every epoch (default: $\ensuremath{FALSE}\xspace).$
sampler	(Sampler, optional): defines the strategy to draw samples from the dataset. If specified, shuffle must be False. Custom samplers can be created with sampler().
batch_sampler	(Sampler, optional): like sampler, but returns a batch of indices at a time. Mutually exclusive with batch_size, shuffle, sampler, and drop_last. Custom samplers can be created with sampler().
num_workers	(int, optional): how many subprocesses to use for data loading. 0 means that the data will be loaded in the main process. (default: 0)
collate_fn	(callable, optional): merges a list of samples to form a mini-batch.
pin_memory	(bool, optional): If TRUE, the data loader will copy tensors into CUDA pinned memory before returning them. If your data elements are a custom type, or your collate_fn returns a batch that is a custom type see the example below.
drop_last	(bool, optional): set to TRUE to drop the last incomplete batch, if the dataset size is not divisible by the batch size. If FALSE and the size of dataset is not divisible by the batch size, then the last batch will be smaller. (default: $FALSE$ )
timeout	(numeric, optional): if positive, the timeout value for collecting a batch from workers1 means no timeout. (default: $-1$ )
worker_init_fn	(callable, optional): If not NULL, this will be called on each worker subprocess with the worker id (an int in $[1, num\_workers]$ ) as input, after seeding and before data loading. (default: NULL)

worker\_globals (list or character vector, optional) only used when num\_workers > 0. If a charac-

ter vector, then objects with those names are copied from the global environment

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to the workers. If a named list, then this list is copied and attached to the worker global environment. Notice that the objects are copied only once at the worker initialization.

worker\_packages

(character vector, optional) Only used if num\_workers > 0 optional character vector naming packages that should be loaded in each worker.

#### Parallel data loading

When using num\_workers > 0 data loading will happen in parallel for each worker. Note that batches are taken in parallel and not observations.

The worker initialization process happens in the following order:

• num\_workers R sessions are initialized.

Then in each worker we perform the following actions:

- the torch library is loaded.
- a random seed is set both using set.seed() and using torch\_manual\_seed.
- packages passed to the worker\_packages argument are loaded.
- objects passed trough the worker\_globals parameters are copied into the global environment.
- the worker\_init function is ran with an id argument.
- the dataset fetcher is copied to the worker.

## See Also

```
dataset(), sampler()
```

## Description

Creates an iterator from a DataLoader

## Usage

```
dataloader_make_iter(dataloader)
```

## **Arguments**

dataloader a dataloader object.

dataloader\_next 35

dataloader\_next

Get the next element of a dataloader iterator

## Description

Get the next element of a dataloader iterator

## Usage

```
dataloader_next(iter, completed = NULL)
```

## **Arguments**

iter a DataLoader iter created with dataloader\_make\_iter. completed the returned value when the iterator is exhausted.

dataset

Helper function to create an function that generates R6 instances of class dataset

## **Description**

All datasets that represent a map from keys to data samples should subclass this class. All subclasses should overwrite the .getitem() method, which supports fetching a data sample for a given key. Subclasses could also optionally overwrite .length(), which is expected to return the size of the dataset (e.g. number of samples) used by many sampler implementations and the default options of dataloader().

## Usage

```
dataset(
  name = NULL,
  inherit = Dataset,
  ...,
  private = NULL,
  active = NULL,
  parent_env = parent.frame()
)
```

#### **Arguments**

name a name for the dataset. It it's also used as the class for it.
inherit you can optionally inherit from a dataset when creating a new dataset.

... public methods for the dataset class

private passed to R6::R6Class().
active passed to R6::R6Class().

parent\_env An environment to use as the parent of newly-created objects.

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#### Value

The output is a function f with class dataset\_generator. Calling f() creates a new instance of the R6 class dataset. The R6 class is stored in the enclosing environment of f and can also be accessed through fs attribute Dataset.

#### Get a batch of observations

By default datasets are iterated by returning each observation/item individually. Often it's possible to have an optimized implementation to take a batch of observations (eg, subsetting a tensor by multiple indexes at once is faster than subsetting once for each index), in this case you can implement a .getbatch method that will be used instead of .getitem when getting a batch of observations within the dataloader. .getbatch must work for batches of size larger or equal to 1 and care must be taken so it doesn't drop the batch dimension when it's queried with a length 1 batch index - for instance by using drop=FALSE. .getitem() is expected to not include the batch dimension as it's added by the dataloader. For more on this see the the vignette("loading-data").

#### Note

dataloader() by default constructs a index sampler that yields integral indices. To make it work with a map-style dataset with non-integral indices/keys, a custom sampler must be provided.

 $dataset\_subset$ 

Dataset Subset

## **Description**

Subset of a dataset at specified indices.

## Usage

```
dataset_subset(dataset, indices)
```

## **Arguments**

dataset (Dataset): The whole Dataset

indices (sequence): Indices in the whole set selected for subset

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Distribution

Generic R6 class representing distributions

### **Description**

Distribution is the abstract base class for probability distributions. Note: in Python, adding torch. Size objects works as concatenation Try for example: torch.Size((2, 1)) + torch.Size((1,))

#### **Public fields**

```
.validate_args whether to validate arguments
has_rsample whether has an rsample
has_enumerate_support whether has enumerate support
```

### **Active bindings**

batch\_shape Returns the shape over which parameters are batched.

event\_shape Returns the shape of a single sample (without batching). Returns a dictionary from argument names to torch\_Constraint objects that should be satisfied by each argument of this distribution. Args that are not tensors need not appear in this dict.

support Returns a torch\_Constraint object representing this distribution's support.

mean Returns the mean on of the distribution

variance Returns the variance of the distribution

stddev Returns the standard deviation of the distribution TODO: consider different message

#### Methods

#### **Public methods:**

- Distribution\$new()
- Distribution\$expand()
- Distribution\$sample()
- Distribution\$rsample()
- Distribution\$log\_prob()
- Distribution\$cdf()
- Distribution\$icdf()
- Distribution\$enumerate\_support()
- Distribution\$entropy()
- Distribution\$perplexity()
- Distribution\$.extended\_shape()
- Distribution\$.validate\_sample()
- Distribution\$print()
- Distribution\$clone()

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**Method** new(): Initializes a distribution class.

Usage.

Distribution\$new(batch\_shape = NULL, event\_shape = NULL, validate\_args = NULL)

Arguments:

batch\_shape the shape over which parameters are batched.

event\_shape the shape of a single sample (without batching).

validate\_args whether to validate the arguments or not. Validation can be time consuming so you might want to disable it.

**Method** expand(): Returns a new distribution instance (or populates an existing instance provided by a derived class) with batch dimensions expanded to batch\_shape. This method calls expand on the distribution's parameters. As such, this does not allocate new memory for the expanded distribution instance. Additionally, this does not repeat any args checking or parameter broadcasting in initialize, when an instance is first created.

Usage:

Distribution\$expand(batch\_shape, .instance = NULL)

Arguments:

batch\_shape the desired expanded size.

. instance new instance provided by subclasses that need to override expand.

**Method** sample(): Generates a sample\_shape shaped sample or sample\_shape shaped batch of samples if the distribution parameters are batched.

Usage:

Distribution\$sample(sample\_shape = NULL)

Arguments:

sample\_shape the shape you want to sample.

**Method** rsample(): Generates a sample\_shape shaped reparameterized sample or sample\_shape shaped batch of reparameterized samples if the distribution parameters are batched.

Usage:

Distribution\$rsample(sample\_shape = NULL)

Arguments:

sample\_shape the shape you want to sample.

**Method** log\_prob(): Returns the log of the probability density/mass function evaluated at value.

Usage:

Distribution\$log\_prob(value)

Arguments:

value values to evaluate the density on.

**Method** cdf(): Returns the cumulative density/mass function evaluated at value.

Usage:

Distribution\$cdf(value) Arguments: value values to evaluate the density on. **Method** icdf(): Returns the inverse cumulative density/mass function evaluated at value. @description Returns tensor containing all values supported by a discrete distribution. The result will enumerate over dimension 0, so the shape of the result will be (cardinality,) + batch\_shape + event\_shape (whe = ()for univariate distributions). Note that this enumerates over all batched tensors in lock-steplist(c( (0), (1,1),...). With expand=FALSE, enumeration happens along dim 0, but with the remaining batch dimension c(1), ...). Usage: Distribution\$icdf(value) Arguments: value values to evaluate the density on. Method enumerate\_support(): Usage: Distribution\$enumerate\_support(expand = TRUE) Arguments: expand (bool): whether to expand the support over the batch dims to match the distribution's batch\_shape. Returns: Tensor iterating over dimension 0. **Method** entropy(): Returns entropy of distribution, batched over batch\_shape. Usage: Distribution\$entropy() *Returns:* Tensor of shape batch\_shape. Method perplexity(): Returns perplexity of distribution, batched over batch\_shape. Usage: Distribution\$perplexity() *Returns:* Tensor of shape batch\_shape. **Method** .extended\_shape(): Returns the size of the sample returned by the distribution, given a sample\_shape. Note, that the batch and event shapes of a distribution instance are fixed at the time of construction. If this is empty, the returned shape is upcast to (1,). Distribution\$.extended\_shape(sample\_shape = NULL) Arguments: sample\_shape (torch\_Size): the size of the sample to be drawn.

**Method** .validate\_sample(): Argument validation for distribution methods such as log\_prob, cdf and icdf. The rightmost dimensions of a value to be scored via these methods must agree with the distribution's batch and event shapes.

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```
Usage:
Distribution$.validate_sample(value)
Arguments:
value (Tensor): the tensor whose log probability is to be computed by the log_prob method.

Method print(): Prints the distribution instance.

Usage:
Distribution$print()

Method clone(): The objects of this class are cloneable with this method.

Usage:
Distribution$clone(deep = FALSE)

Arguments:
deep Whether to make a deep clone.
```

distr\_bernoulli

Creates a Bernoulli distribution parameterized by probs or logits (but not both). Samples are binary (0 or 1). They take the value 1 with probability p and 0 with probability 1 - p.

#### **Description**

Creates a Bernoulli distribution parameterized by probs or logits (but not both). Samples are binary (0 or 1). They take the value 1 with probability p and 0 with probability 1 - p.

# Usage

```
distr_bernoulli(probs = NULL, logits = NULL, validate_args = NULL)
```

# **Arguments**

```
probs (numeric or torch_tensor): the probability of sampling 1 logits (numeric or torch_tensor): the log-odds of sampling 1 validate_args whether to validate arguments or not.
```

### See Also

Distribution for details on the available methods.

```
Other distributions: distr_chi2(), distr_gamma(), distr_multivariate_normal(), distr_normal(), distr_poisson()
```

```
if (torch_is_installed()) {
m <- distr_bernoulli(0.3)
m$sample() # 30% chance 1; 70% chance 0
}</pre>
```

distr\_categorical 41

distr_categorical	Creates a categorical distribution parameterized by either probs or
	logits (but not both).

# **Description**

Creates a categorical distribution parameterized by either probs or logits (but not both).

#### Usage

```
distr_categorical(probs = NULL, logits = NULL, validate_args = NULL)
```

### **Arguments**

```
probs (Tensor): event probabilities
logits (Tensor): event log probabilities (unnormalized)
validate_args Additional arguments
```

#### Note

It is equivalent to the distribution that torch\_multinomial() samples from.

```
Samples are integers from \{0, \dots, K-1\} where K is probs$size(-1).
```

If probs is 1-dimensional with length-K, each element is the relative probability of sampling the class at that index.

If probs is N-dimensional, the first N-1 dimensions are treated as a batch of relative probability vectors.

The probs argument must be non-negative, finite and have a non-zero sum, and it will be normalized to sum to 1 along the last dimension. attr:probs will return this normalized value. The logits argument will be interpreted as unnormalized log probabilities and can therefore be any real number. It will likewise be normalized so that the resulting probabilities sum to 1 along the last dimension. attr:logits will return this normalized value.

```
See also: torch_multinomial()
```

```
if (torch_is_installed()) {
m <- distr_categorical(torch_tensor(c(0.25, 0.25, 0.25, 0.25)))
m$sample() # equal probability of 1,2,3,4
}</pre>
```

distr\_gamma

distr_chi2	Creates a Chi2 distribution parameterized by shape parameter df. This is exactly equivalent to distr_gamma(alpha=0.5*df, beta=0.5)
	Deta-0.5)

# **Description**

Creates a Chi2 distribution parameterized by shape parameter df. This is exactly equivalent to distr\_gamma(alpha=0.5\*df, beta=0.5)

# Usage

```
distr_chi2(df, validate_args = NULL)
```

# **Arguments**

```
df (float or torch_tensor): shape parameter of the distribution validate_args whether to validate arguments or not.
```

#### See Also

Distribution for details on the available methods.

```
Other distributions: distr_bernoulli(), distr_gamma(), distr_multivariate_normal(), distr_normal(), distr_poisson()
```

# **Examples**

```
if (torch_is_installed()) {
m <- distr_chi2(torch_tensor(1.0))
m$sample() # Chi2 distributed with shape df=1
torch_tensor(0.1046)
}</pre>
```

# Description

Creates a Gamma distribution parameterized by shape concentration and rate.

# Usage

```
distr_gamma(concentration, rate, validate_args = NULL)
```

### Arguments

```
concentration (float or Tensor): shape parameter of the distribution (often referred to as alpha)
rate (float or Tensor): rate = 1 / scale of the distribution (often referred to as beta)
validate_args whether to validate arguments or not.
```

#### See Also

Distribution for details on the available methods.

```
Other distributions: distr_bernoulli(), distr_chi2(), distr_multivariate_normal(), distr_normal(), distr_poisson()
```

### **Examples**

```
if (torch_is_installed()) {
m <- distr_gamma(torch_tensor(1.0), torch_tensor(1.0))
m$sample() # Gamma distributed with concentration=1 and rate=1
}</pre>
```

```
distr_mixture_same_family
```

Mixture of components in the same family

### **Description**

The MixtureSameFamily distribution implements a (batch of) mixture distribution where all component are from different parameterizations of the same distribution type. It is parameterized by a Categorical selecting distribution" (over k component) and a component distribution, i.e., a Distribution with a rightmost batch shape (equal to [k]) which indexes each (batch of) component.

### Usage

```
distr_mixture_same_family(
  mixture_distribution,
  component_distribution,
  validate_args = NULL
)
```

### Arguments

```
mixture_distribution
```

torch\_distributions.Categorical-like instance. Manages the probability of selecting component. The number of categories must match the rightmost batch dimension of the component\_distribution. Must have either scalar batch\_shape or batch\_shape matching component\_distribution.batch\_shape[:-1]

```
component_distribution torch_distributions.Distribution-like instance. Right-most batch dimension indexes component.
```

# Examples

validate\_args

```
if (torch_is_installed()) {
# Construct Gaussian Mixture Model in 1D consisting of 5 equally
# weighted normal distributions
mix <- distr_categorical(torch_ones(5))
comp <- distr_normal(torch_randn(5), torch_rand(5))
gmm <- distr_mixture_same_family(mix, comp)
}</pre>
```

Additional arguments

```
distr_multivariate_normal
```

Gaussian distribution

# **Description**

Creates a multivariate normal (also called Gaussian) distribution parameterized by a mean vector and a covariance matrix.

#### **Usage**

```
distr_multivariate_normal(
  loc,
  covariance_matrix = NULL,
  precision_matrix = NULL,
  scale_tril = NULL,
  validate_args = NULL
)
```

# **Arguments**

```
loc (Tensor): mean of the distribution

covariance_matrix

(Tensor): positive-definite covariance matrix

precision_matrix

(Tensor): positive-definite precision matrix

scale_tril (Tensor): lower-triangular factor of covariance, with positive-valued diagonal validate_args

Bool wether to validate the arguments or not.
```

distr\_normal 45

#### **Details**

The multivariate normal distribution can be parameterized either in terms of a positive definite covariance matrix  $\Sigma$  or a positive definite precision matrix  $\Sigma^{-1}$  or a lower-triangular matrix  $\mathbf{L}$  with positive-valued diagonal entries, such that  $\Sigma = \mathbf{L}\mathbf{L}^{\top}$ . This triangular matrix can be obtained via e.g. Cholesky decomposition of the covariance.

#### Note

Only one of covariance\_matrix or precision\_matrix or scale\_tril can be specified. Using scale\_tril will be more efficient: all computations internally are based on scale\_tril. If covariance\_matrix or precision\_matrix is passed instead, it is only used to compute the corresponding lower triangular matrices using a Cholesky decomposition.

#### See Also

Distribution for details on the available methods.

Other distributions: distr\_bernoulli(), distr\_chi2(), distr\_gamma(), distr\_normal(), distr\_poisson()

#### **Examples**

```
if (torch_is_installed()) {
m <- distr_multivariate_normal(torch_zeros(2), torch_eye(2))
m$sample() # normally distributed with mean=`[0,0]` and covariance_matrix=`I`
}</pre>
```

distr\_normal

Creates a normal (also called Gaussian) distribution parameterized by loc and scale.

#### **Description**

Creates a normal (also called Gaussian) distribution parameterized by loc and scale.

#### Usage

```
distr_normal(loc, scale, validate_args = NULL)
```

#### **Arguments**

loc (float or Tensor): mean of the distribution (often referred to as mu)

scale (float or Tensor): standard deviation of the distribution (often referred to as

sigma)

validate\_args Additional arguments

### Value

Object of torch\_Normal class

distr\_poisson

#### See Also

Distribution for details on the available methods.

```
Other distributions: distr_bernoulli(), distr_chi2(), distr_gamma(), distr_multivariate_normal(), distr_poisson()
```

# **Examples**

```
if (torch_is_installed()) {
m <- distr_normal(loc = 0, scale = 1)
m$sample() # normally distributed with loc=0 and scale=1
}</pre>
```

distr\_poisson

Creates a Poisson distribution parameterized by rate, the rate parameter.

# **Description**

Samples are nonnegative integers, with a pmf given by

$$rate^{k} \frac{e^{-rate}}{k!}$$

#### Usage

```
distr_poisson(rate, validate_args = NULL)
```

# **Arguments**

```
rate (numeric, torch_tensor): the rate parameter validate_args whether to validate arguments or not.
```

#### See Also

Distribution for details on the available methods.

```
Other distributions: distr_bernoulli(), distr_chi2(), distr_gamma(), distr_multivariate_normal(), distr_normal()
```

```
if (torch_is_installed()) {
m <- distr_poisson(torch_tensor(4))
m$sample()
}</pre>
```

enumerate 47

enumerate

Enumerate an iterator

# Description

Enumerate an iterator

# Usage

```
enumerate(x, ...)
```

# Arguments

x the generator to enumerate.

... passed to specific methods.

# Description

Enumerate an iterator

# Usage

```
## S3 method for class 'dataloader'
enumerate(x, max_len = 1e+06, ...)
```

# Arguments

x the generator to enumerate.

max\_len maximum number of iterations.

... passed to specific methods.

48 get\_install\_libs\_url

```
get_install_libs_url Install Torch from files
```

### **Description**

List the Torch and Lantern libraries URLs to download as local files in order to proceed with install\_torch\_from\_file().

Installs Torch and its dependencies from files.

# Usage

```
get_install_libs_url(version = NA, type = NA)
install_torch_from_file(version = NA, type = NA, libtorch, liblantern, ...)
```

#### **Arguments**

version	Not used
type	Not used. This function is deprecated.
libtorch	The installation archive file to use for Torch. Shall be a "file://" URL scheme.
liblantern	The installation archive file to use for Lantern. Shall be a "file://" URL scheme.
	other parameters to be passed to "install_torch()"

### **Details**

When "install\_torch()" initiated download is not possible, but installation archive files are present on local filesystem, "install\_torch\_from\_file()" can be used as a workaround to installation issue. "libtorch" is the archive containing all torch modules, and "liblantern" is the C interface to libtorch that is used for the R package. Both are highly dependent, and should be checked through "get\_install\_libs\_url()"

```
if (torch_is_installed()) {
## Not run:
# on a linux CPU platform
get_install_libs_url()
# then after making both files available into /tmp/
Sys.setenv(TORCH_URL="/tmp/libtorch-v1.13.1.zip")
Sys.setenv(LANTERN_URL="/tmp/lantern-0.9.1.9001+cpu+arm64-Darwin.zip")
torch::install_torch()
## End(Not run)
}
```

install\_torch 49

#### **Description**

Installs Torch and its dependencies.

# Usage

```
install_torch(reinstall = FALSE, ..., .inform_restart = TRUE)
```

### **Arguments**

reinstall Re-install Torch even if its already installed?
... Currently unused.

.inform\_restart

if TRUE and running in an interactive() session, after installation it will print a message to inform the user that the session must be restarted for torch to work correctly.

#### **Details**

This function is mainly controlled by environment variables that can be used to override the defaults:

- TORCH\_HOME: the installation path. By default dependencies are installed within the package directory. Eg what's given by system.file(package="torch").
- TORCH\_URL: A URL, path to a ZIP file or a directory containing a LibTorch version. Files will be installed/copied to the TORCH\_HOME directory.
- LANTERN\_URL: Same as TORCH\_URL but for the Lantern library.
- TORCH\_INSTALL\_DEBUG: Setting it to 1, shows debug log messages during installation.
- PRECXX11ABI: Setting it to 1 will will trigger the installation of a Pre-cxx11 ABI installation
  of LibTorch. This can be useful in environments with older versions of GLIBC like CentOS7
  and older Debian/Ubuntu versions.
- LANTERN\_BASE\_URL: The base URL for lantern files. This allows passing a directory where lantern binaries are located. The filename is then constructed as usual.
- TORCH\_COMMIT\_SHA: torch repository commit sha to be used when querying lantern uploads. Set it to 'none' to avoid looking for build for that commit and use the latest build for the branch.
- CUDA: We try to automatically detect the CUDA version installed in your system, but you might want to manually set it here. You can also disable CUDA installation by setting it to 'cpu'.
- TORCH\_R\_VERSION: The R torch version. It's unlikely that you need to change it, but it can be useful if you don't have the R package installed, but want to install the dependencies.

is\_nn\_buffer

The TORCH\_INSTALL environment variable can be set to 0 to prevent auto-installing torch and TORCH\_LOAD set to 0 to avoid loading dependencies automatically. These environment variables are meant for advanced use cases and troubleshooting only. When timeout error occurs during library archive download, or length of downloaded files differ from reported length, an increase of the timeout value should help.

is\_dataloader

Checks if the object is a dataloader

# Description

Checks if the object is a dataloader

# Usage

```
is_dataloader(x)
```

# Arguments

Х

object to check

is\_nn\_buffer

Checks if the object is a nn\_buffer

# **Description**

Checks if the object is a nn\_buffer

# Usage

```
is_nn_buffer(x)
```

#### **Arguments**

x object to check

is\_nn\_module 51

is\_nn\_module

Checks if the object is an nn\_module

# Description

Checks if the object is an nn\_module

# Usage

```
is_nn_module(x)
```

# **Arguments**

Х

object to check

 $is\_nn\_parameter$ 

Checks if an object is a nn\_parameter

# Description

Checks if an object is a nn\_parameter

# Usage

```
is_nn_parameter(x)
```

# Arguments

х

the object to check

is\_optimizer

Checks if the object is a torch optimizer

# Description

Checks if the object is a torch optimizer

# Usage

```
is_optimizer(x)
```

# **Arguments**

Х

object to check

is\_torch\_layout

is\_torch\_device

Checks if object is a device

# Description

Checks if object is a device

# Usage

```
is_torch_device(x)
```

# **Arguments**

Х

object to check

is\_torch\_dtype

Check if object is a torch data type

# Description

Check if object is a torch data type

# Usage

```
is_torch_dtype(x)
```

# **Arguments**

Χ

object to check.

is\_torch\_layout

Check if an object is a torch layout.

# Description

Check if an object is a torch layout.

# Usage

```
is_torch_layout(x)
```

# **Arguments**

Х

object to check

```
is_torch_memory_format
```

Check if an object is a memory format

# Description

Check if an object is a memory format

# Usage

```
is_torch_memory_format(x)
```

#### **Arguments**

Х

object to check

is\_torch\_qscheme

Checks if an object is a QScheme

# Description

Checks if an object is a QScheme

# Usage

```
is_torch_qscheme(x)
```

# **Arguments**

Х

object to check

 $is\_undefined\_tensor$ 

Checks if a tensor is undefined

# Description

Checks if a tensor is undefined

# Usage

```
is_undefined_tensor(x)
```

# **Arguments**

Х

tensor to check

54 iterable\_dataset

 $iterable\_dataset$ 

Creates an iterable dataset

# **Description**

Creates an iterable dataset

# Usage

```
iterable_dataset(
  name,
  inherit = IterableDataset,
  ...,
  private = NULL,
  active = NULL,
  parent_env = parent.frame()
)
```

# Arguments

```
name a name for the dataset. It it's also used as the class for it.

you can optionally inherit from a dataset when creating a new dataset.

public methods for the dataset class

private passed to R6::R6Class().

active passed to R6::R6Class().

An environment to use as the parent of newly-created objects.
```

```
if (torch_is_installed()) {
ids <- iterable_dataset(</pre>
 name = "hello",
 initialize = function(n = 5) {
   self$n <- n
   self$i <- 0
 },
 .iter = function() {
   i <- 0
   function() {
      i <<- i + 1
      if (i > self n)  {
        coro::exhausted()
      } else {
      }
   }
 }
```

jit\_compile 55

```
coro::collect(ids()$.iter())
}
```

jit\_compile

Compile TorchScript code into a graph

# Description

See the TorchScript language reference for documentation on how to write TorchScript code.

# Usage

```
jit_compile(source)
```

# **Arguments**

source

valid TorchScript source code.

# **Examples**

```
if (torch_is_installed()) {
comp <- jit_compile("
def fn (x):
    return torch.abs(x)

def foo (x):
    return torch.sum(x)

")

comp$fn(torch_tensor(-1))
comp$foo(torch_randn(10))
}</pre>
```

jit\_load

 $Loads\ a\ script\_function\ or\ script\_module\ previously\ saved\ with\ jit\_save$ 

# **Description**

Loads a script\_function or script\_module previously saved with jit\_save

# Usage

```
jit_load(path, ...)
```

jit\_ops

# **Arguments**

```
path a path to a script_function or script_module serialized with jit_save().
... currently unused.
```

jit\_ops

Enable idiomatic access to JIT operators from R.

# Description

Call JIT operators directly from R, keeping the familiar argument types and argument order. Note, however, that:

- all arguments are required (no defaults)
- axis numbering (as well as position numbers overall) starts from 0
- scalars have to be wrapped in jit\_scalar()

# Usage

```
jit_ops
```

#### **Format**

An object of class torch\_ops of length 0.

```
if (torch_is_installed()) {
t1 <- torch::torch_rand(4, 5)
t2 <- torch::torch_ones(5, 4)
# same as torch::torch_matmul(t1, t2)
jit_ops$aten$matmul(t1, t2)

# same as torch_split(torch::torch_arange(0, 3), 2, 1)
jit_ops$aten$split(torch::torch_arange(0, 3), torch::jit_scalar(2L), torch::jit_scalar(0L))
}</pre>
```

jit\_save 57

jit\_save

Saves a script\_function to a path

# **Description**

Saves a script\_function to a path

#### Usage

```
jit_save(obj, path, ...)
```

# **Arguments**

```
obj An script_function to save

path The path to save the serialized function.

... currently unused
```

# **Examples**

```
if (torch_is_installed()) {
fn <- function(x) {
   torch_relu(x)
}

input <- torch_tensor(c(-1, 0, 1))
tr_fn <- jit_trace(fn, input)

tmp <- tempfile("tst", fileext = "pt")
jit_save(tr_fn, tmp)
}</pre>
```

jit\_save\_for\_mobile

Saves a  $script\_function\ or\ script\_module\ in\ bytecode\ form,\ to\ be$  loaded on a mobile device

# **Description**

Saves a script\_function or script\_module in bytecode form, to be loaded on a mobile device

# Usage

```
jit_save_for_mobile(obj, path, ...)
```

jit\_trace

# **Arguments**

```
obj An script_function or script_module to save
path The path to save the serialized function.
... currently unused
```

# **Examples**

```
if (torch_is_installed()) {
fn <- function(x) {
   torch_relu(x)
}
input <- torch_tensor(c(-1, 0, 1))
tr_fn <- jit_trace(fn, input)

tmp <- tempfile("tst", fileext = "pt")
jit_save_for_mobile(tr_fn, tmp)
}</pre>
```

jit\_scalar

Adds the 'jit\_scalar' class to the input

# **Description**

Allows disambiguating length 1 vectors from scalars when passing them to the jit.

# Usage

```
jit_scalar(x)
```

# Arguments

х

a length 1 R vector.

jit\_trace

*Trace a function and return an executable* script\_function.

# Description

Using jit\_trace, you can turn an existing R function into a TorchScript script\_function. You must provide example inputs, and we run the function, recording the operations performed on all the tensors.

### Usage

```
jit_trace(func, ..., strict = TRUE)
```

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### Arguments

An R function that will be run with example\_inputs. func arguments and return values must be tensors or (possibly nested) lists that contain tensors. Can also be a nn\_module(), in such case jit\_trace\_module() is used to trace that module.

... example inputs that will be passed to the function while tracing. The resulting trace can be run with inputs of different types and shapes assuming the traced operations support those types and shapes. example\_inputs may also be a single Tensor in which case it is automatically wrapped in a list. Note that ... can not be named, and the order is respected.

strict run the tracer in a strict mode or not (default: TRUE). Only turn this off when

run the tracer in a strict mode or not (default: TRUE). Only turn this off when you want the tracer to record your mutable container types (currently list/dict) and you are sure that the container you are using in your problem is a constant

structure and does not get used as control flow (if, for) conditions.

#### **Details**

The resulting recording of a standalone function produces a script\_function. In the future we will also support tracing nn\_modules.

#### Value

An script\_function if func is a function and script\_module if func is a nn\_module().

#### Warning

Tracing only correctly records functions and modules which are not data dependent (e.g., do not have conditionals on data in tensors) and do not have any untracked external dependencies (e.g., perform input/output or access global variables). Tracing only records operations done when the given function is run on the given tensors. Therefore, the returned script\_function will always run the same traced graph on any input. This has some important implications when your module is expected to run different sets of operations, depending on the input and/or the module state. For example,

- Tracing will not record any control-flow like if-statements or loops. When this control-flow is constant across your module, this is fine and it often inlines the control-flow decisions. But sometimes the control-flow is actually part of the model itself. For instance, a recurrent network is a loop over the (possibly dynamic) length of an input sequence.
- In the returned script\_function, operations that have different behaviors in training and
  eval modes will always behave as if it is in the mode it was in during tracing, no matter which
  mode the script\_function is in.

In cases like these, tracing would not be appropriate and scripting is a better choice. If you trace such models, you may silently get incorrect results on subsequent invocations of the model. The tracer will try to emit warnings when doing something that may cause an incorrect trace to be produced.

#### Note

Scripting is not yet supported in R.

jit\_trace\_module

#### **Examples**

```
if (torch_is_installed()) {
  fn <- function(x) {
    torch_relu(x)
}
input <- torch_tensor(c(-1, 0, 1))
tr_fn <- jit_trace(fn, input)
tr_fn(input)
}</pre>
```

jit\_trace\_module

Trace a module

# **Description**

Trace a module and return an executable ScriptModule that will be optimized using just-in-time compilation. When a module is passed to jit\_trace(), only the forward method is run and traced. With jit\_trace\_module(), you can specify a named list of method names to example inputs to trace (see the inputs) argument below.

#### Usage

```
jit_trace_module(mod, ..., strict = TRUE)
```

# **Arguments**

mod	A torch nn_module() containing methods whose names are specified in inputs.  The given methods will be compiled as a part of a single ScriptModule.
• • •	A named list containing sample inputs indexed by method names in mod. The
	inputs will be passed to methods whose names correspond to inputs keys while
	<pre>tracing. list('forward'=example_forward_input, 'method2'=example_method2_input).</pre>
strict	run the tracer in a strict mode or not (default: TRUE). Only turn this off when
	you want the tracer to record your mutable container types (currently list/dict)
	and you are sure that the container you are using in your problem is a constant
	structure and does not get used as control flow (if, for) conditions.

#### **Details**

See jit\_trace for more information on tracing.

```
if (torch_is_installed()) {
linear <- nn_linear(10, 1)
tr_linear <- jit_trace_module(linear, forward = list(torch_randn(10, 10)))

x <- torch_randn(10, 10)
torch_allclose(linear(x), tr_linear(x))
}</pre>
```

jit\_tuple 61

# **Description**

Allows specifying that an output or input must be considered a jit tuple and instead of a list or dictionary when tracing.

# Usage

```
jit_tuple(x)
```

# **Arguments**

x the list object that will be converted to a tuple.

linalg_cholesky	Computes the Cholesky decomposition of a complex Hermitian or real
	symmetric positive-definite matrix.

# Description

Letting . be . or . , the **Cholesky decomposition** of a complex Hermitian or real symmetric positive-definite matrix . is defined as

# Usage

```
linalg_cholesky(A)
```

### **Arguments**

A (Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions consisting of symmetric or Hermitian positive-definite matrices.

### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is a lower triangular matrix and . is the conjugate transpose when . is complex, and the transpose when . is real-valued.

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

62 linalg\_cholesky\_ex

#### See Also

• linalg\_cholesky\_ex() for a version of this operation that skips the (slow) error checking by default and instead returns the debug information. This makes it a faster way to check if a matrix is positive-definite. linalg\_eigh() for a different decomposition of a Hermitian matrix. The eigenvalue decomposition gives more information about the matrix but it slower to compute than the Cholesky decomposition.

```
Other linalg: linalg_cholesky_ex(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

#### **Examples**

```
if (torch_is_installed()) {
a <- torch_eye(10)
linalg_cholesky(a)
}</pre>
```

linalg\_cholesky\_ex

Computes the Cholesky decomposition of a complex Hermitian or real symmetric positive-definite matrix.

#### **Description**

This function skips the (slow) error checking and error message construction of linalg\_cholesky(), instead directly returning the LAPACK error codes as part of a named tuple (L, info). This makes this function a faster way to check if a matrix is positive-definite, and it provides an opportunity to handle decomposition errors more gracefully or performantly than linalg\_cholesky() does. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions. If A is not a Hermitian positive-definite matrix, or if it's a batch of matrices and one or more of them is not a Hermitian positive-definite matrix, then info stores a positive integer for the corresponding matrix. The positive integer indicates the order of the leading minor that is not positive-definite, and the decomposition could not be completed. info filled with zeros indicates that the decomposition was successful. If check\_errors=TRUE and info contains positive integers, then a RuntimeError is thrown.

#### **Usage**

```
linalg_cholesky_ex(A, check_errors = FALSE)
```

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# Arguments

A (Tensor): the Hermitian n \times n matrix or the batch of such matrices of size (\*, n, n) where \* is one or more batch dimensions.

check\_errors (bool, optional): controls whether to check the content of infos. Default: FALSE.

#### Note

If A is on a CUDA device, this function may synchronize that device with the CPU. This function is "experimental" and it may change in a future PyTorch release.

#### See Also

```
linalg_cholesky() is a NumPy compatible variant that always checks for errors.

Other linalg: linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_solve(), linalg_tensorsolve(), linalg_vector_norm()
```

# **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(2, 2)
out <- linalg_cholesky_ex(A)
out
}</pre>
```

linalg\_cond

Computes the condition number of a matrix with respect to a matrix norm.

# Description

```
Letting . be . or . , the condition number . of a matrix . is defined as
```

#### **Usage**

```
linalg\_cond(A, p = NULL)
```

# **Arguments**

```
A (Tensor): tensor of shape (*, m, n) where * is zero or more batch dimensions for p in (2, -2), and of shape (*, n, n) where every matrix is invertible for p in ('fro', 'nuc', inf, -inf, 1, -1).

p (int, inf, -inf, 'fro', 'nuc', optional): the type of the matrix norm to use in the computations (see above). Default: NULL
```

64 linalg\_cond

#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

The condition number of A measures the numerical stability of the linear system AX = B with respect to a matrix norm.

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

p defines the matrix norm that is computed. See the table in 'Details' to find the supported norms.

For p is one of ('fro', 'nuc', inf, -inf, 1, -1), this function uses linalg\_norm() and linalg\_inv().

As such, in this case, the matrix (or every matrix in the batch) A has to be square and invertible.

For p in (2, -2), this function can be computed in terms of the singular values.

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

In these cases, it is computed using linalg\_svd(). For these norms, the matrix (or every matrix in the batch) A may have any shape.

```
matrix norm
р
NULL
        2-norm (largest singular value)
'fro'
        Frobenius norm
'nuc'
        nuclear norm
Inf
        max(sum(abs(x), dim=2))
-Inf
        min(sum(abs(x), dim=2))
1
        max(sum(abs(x), dim=1))
-1
        min(sum(abs(x), dim=1))
        largest singular value
-2
        smallest singular value
```

#### Value

A real-valued tensor, even when A is complex.

#### Note

When inputs are on a CUDA device, this function synchronizes that device with the CPU if if p is one of ('fro', 'nuc', inf, -inf, 1, -1).

```
if (torch_is_installed()) {
a <- torch_tensor(rbind(c(1., 0, -1), c(0, 1, 0), c(1, 0, 1)))
linalg_cond(a)
linalg_cond(a, "fro")
}</pre>
```

linalg\_det 65

linalg\_det

Computes the determinant of a square matrix.

#### **Description**

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

### Usage

```
linalg_det(A)
```

#### **Arguments**

Α

(Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions.

#### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangulardlinalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

#### **Examples**

```
if (torch_is_installed()) {
a <- torch_randn(3, 3)
linalg_det(a)

a <- torch_randn(3, 3, 3)
linalg_det(a)
}</pre>
```

linalg\_eig

Computes the eigenvalue decomposition of a square matrix if it exists.

# **Description**

Letting . be . or . , the **eigenvalue decomposition** of a square matrix . (if it exists) is defined as

# Usage

```
linalg_eig(A)
```

66 linalg\_eig

#### **Arguments**

Α

(Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions consisting of diagonalizable matrices.

#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

This decomposition exists if and only if . is diagonalizable\_. This is the case when all its eigenvalues are different. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

#### Value

A list (eigenvalues, eigenvectors) which corresponds to . and . above. eigenvalues and eigenvectors will always be complex-valued, even when A is real. The eigenvectors will be given by the columns of eigenvectors.

#### Warning

- This function assumes that A is diagonalizable\_ (for example, when all the eigenvalues are different). If it is not diagonalizable, the returned eigenvalues will be correct but . .
- The eigenvectors of a matrix are not unique, nor are they continuous with respect to A. Due to this lack of uniqueness, different hardware and software may compute different eigenvectors. This non-uniqueness is caused by the fact that multiplying an eigenvector by a non-zero number produces another set of valid eigenvectors of the matrix. In this implmentation, the returned eigenvectors are normalized to have norm 1 and largest real component.
- Gradients computed using V will only be finite when A does not have repeated eigenvalues.
   Furthermore, if the distance between any two eigenvalues is close to zero, the gradient will be numerically unstable, as it depends on the eigenvalues . through the computation of . .

#### Note

The eigenvalues and eigenvectors of a real matrix may be complex.

#### See Also

- linalg\_eigvals() computes only the eigenvalues. Unlike linalg\_eig(), the gradients of linalg\_eigvals() are always numerically stable.
- linalg\_eigh() for a (faster) function that computes the eigenvalue decomposition for Hermitian and symmetric matrices.
- linalg\_svd() for a function that computes another type of spectral decomposition that works on matrices of any shape.
- linalg\_qr() for another (much faster) decomposition that works on matrices of any shape.

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvals(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(),
```

linalg\_eigh 67

```
linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(),
linalg_vector_norm()
```

#### **Examples**

```
if (torch_is_installed()) {
a <- torch_randn(2, 2)
wv <- linalg_eig(a)
}</pre>
```

linalg\_eigh

Computes the eigenvalue decomposition of a complex Hermitian or real symmetric matrix.

# **Description**

Letting . be . or . , the **eigenvalue decomposition** of a complex Hermitian or real symmetric matrix . is defined as

#### Usage

```
linalg_eigh(A, UPLO = "L")
```

# **Arguments**

A (Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions consisting of symmetric or Hermitian matrices.

UPLO ('L', 'U', optional): controls whether to use the upper or lower triangular part of A in the computations. Default: 'L'.

### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is the conjugate transpose when . is complex, and the transpose when . is real-valued. . is orthogonal in the real case and unitary in the complex case.

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

A is assumed to be Hermitian (resp. symmetric), but this is not checked internally, instead:

- If UPLO\ = 'L' (default), only the lower triangular part of the matrix is used in the computation.
- If UPLO\ = 'U', only the upper triangular part of the matrix is used. The eigenvalues are returned in ascending order.

68 linalg\_eigh

#### Value

A list (eigenvalues, eigenvectors) which corresponds to . and . above. eigenvalues will always be real-valued, even when A is complex.

It will also be ordered in ascending order. eigenvectors will have the same dtype as A and will contain the eigenvectors as its columns.

# Warning

- The eigenvectors of a symmetric matrix are not unique, nor are they continuous with respect to A. Due to this lack of uniqueness, different hardware and software may compute different eigenvectors. This non-uniqueness is caused by the fact that multiplying an eigenvector by -1 in the real case or by . in the complex case produces another set of valid eigenvectors of the matrix. This non-uniqueness problem is even worse when the matrix has repeated eigenvalues. In this case, one may multiply the associated eigenvectors spanning the subspace by a rotation matrix and the resulting eigenvectors will be valid eigenvectors.
- Gradients computed using the eigenvectors tensor will only be finite when A has unique
  eigenvalues. Furthermore, if the distance between any two eigvalues is close to zero, the gradient will be numerically unstable, as it depends on the eigenvalues . through the computation
  of . .

#### Note

The eigenvalues of real symmetric or complex Hermitian matrices are always real.

#### See Also

- linalg\_eigvalsh() computes only the eigenvalues values of a Hermitian matrix. Unlike linalg\_eigh(), the gradients of linalg\_eigvalsh() are always numerically stable.
- linalg\_cholesky() for a different decomposition of a Hermitian matrix. The Cholesky decomposition gives less information about the matrix but is much faster to compute than the eigenvalue decomposition.
- linalg\_eig() for a (slower) function that computes the eigenvalue decomposition of a not necessarily Hermitian square matrix.
- linalg\_svd() for a (slower) function that computes the more general SVD decomposition of matrices of any shape.
- linalg\_qr() for another (much faster) decomposition that works on general matrices.

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigvalsh(), linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

```
if (torch_is_installed()) {
a <- torch_randn(2, 2)</pre>
```

linalg\_eigvals 69

```
linalg_eigh(a)
}
```

linalg\_eigvals

Computes the eigenvalues of a square matrix.

### **Description**

Letting . be . or . , the **eigenvalues** of a square matrix . are defined as the roots (counted with multiplicity) of the polynomial p of degree n given by

### Usage

```
linalg_eigvals(A)
```

### Arguments

Α

(Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions.

#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is the n-dimensional identity matrix. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

#### Note

The eigenvalues of a real matrix may be complex, as the roots of a real polynomial may be complex. The eigenvalues of a matrix are always well-defined, even when the matrix is not diagonalizable.

#### See Also

```
linalg_eig() computes the full eigenvalue decomposition.
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(),
```

```
linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(),
linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(),
linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(),
linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(),
```

linalg\_vector\_norm()

```
if (torch_is_installed()) {
a <- torch_randn(2, 2)
w <- linalg_eigvals(a)
}</pre>
```

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linalg_eigvalsh	Computes the eigenvalues of a complex Hermitian or real symmetric matrix.
-----------------	---

### **Description**

Letting . be . or . , the **eigenvalues** of a complex Hermitian or real symmetric matrix . are defined as the roots (counted with multiplicity) of the polynomial p of degree n given by

#### Usage

```
linalg_eigvalsh(A, UPLO = "L")
```

#### **Arguments**

A	(Tensor): tensor of shape $(*, n, n)$ where $*$ is zero or more batch dimensions consisting of symmetric or Hermitian matrices.
UPLO	('L', 'U', optional): controls whether to use the upper or lower triangular part of A in the computations. Default: 'L'.

#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is the n-dimensional identity matrix.

The eigenvalues of a real symmetric or complex Hermitian matrix are always real. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions. The eigenvalues are returned in ascending order.

A is assumed to be Hermitian (resp. symmetric), but this is not checked internally, instead:

- If UPLO\ = 'L' (default), only the lower triangular part of the matrix is used in the computation.
- If UPLO\ = 'U', only the upper triangular part of the matrix is used.

#### Value

A real-valued tensor cointaining the eigenvalues even when A is complex. The eigenvalues are returned in ascending order.

### See Also

• linalg\_eigh() computes the full eigenvalue decomposition.

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_gr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

### **Examples**

```
if (torch_is_installed()) {
a <- torch_randn(2, 2)
linalg_eigvalsh(a)
}</pre>
```

linalg\_householder\_product

Computes the first n columns of a product of Householder matrices.

# **Description**

Letting . be . or . , for a matrix . with columns . with . and a vector . with . , this function computes the first . columns of the matrix

# Usage

```
linalg_householder_product(A, tau)
```

### Arguments

```
A (Tensor): tensor of shape (*, m, n) where * is zero or more batch dimensions. tau (Tensor): tensor of shape (*, k) where * is zero or more batch dimensions.
```

### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is the m-dimensional identity matrix and . is the conjugate transpose when . is complex, and the transpose when . is real-valued. See Representation of Orthogonal or Unitary Matrices for further details.

Supports inputs of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if the inputs are batches of matrices then the output has the same batch dimensions.

#### Note

This function only uses the values strictly below the main diagonal of A. The other values are ignored.

#### See Also

• torch\_geqrf() can be used together with this function to form the Q from the linalg\_qr() decomposition.

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torch\_ormqr() is a related function that computes the matrix multiplication of a product
of Householder matrices with another matrix. However, that function is not supported by
autograd.

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_gr(), linalg_solve(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorinv(), linalg_vector_norm()
```

#### **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(2, 2)
h_tau <- torch_geqrf(A)
Q <- linalg_householder_product(h_tau[[1]], h_tau[[2]])
torch_allclose(Q, linalg_qr(A)[[1]])
}</pre>
```

linalg\_inv

Computes the inverse of a square matrix if it exists.

#### **Description**

Throws a runtime\_error if the matrix is not invertible.

#### Usage

```
linalg_inv(A)
```

### **Arguments**

Α

(Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions consisting of invertible matrices.

### **Details**

Letting . be . or . , for a matrix . , its **inverse matrix** . (if it exists) is defined as

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is the n-dimensional identity matrix.

The inverse matrix exists if and only if . is invertible. In this case, the inverse is unique. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

Consider using linalg\_solve() if possible for multiplying a matrix on the left by the inverse, as linalg\_solve(A, B) == A\$inv() %\*% B It is always prefered to use linalg\_solve() when possible, as it is faster and more numerically stable than computing the inverse explicitly.

linalg\_inv\_ex 73

#### See Also

```
linalg_pinv() computes the pseudoinverse (Moore-Penrose inverse) of matrices of any shape.
linalg_solve() computes A$inv() %*% B with a numerically stable algorithm.

Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(),
linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_lstsq(),
linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(),
linalg_norm(), linalg_pinv(), linalg_ar(), linalg_slogdet(), linalg_solve_triangular(),
linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(),
linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(4, 4)
linalg_inv(A)
}</pre>
```

linalg\_inv\_ex

Computes the inverse of a square matrix if it is invertible.

## Description

Returns a namedtuple (inverse, info). inverse contains the result of inverting A and info stores the LAPACK error codes. If A is not an invertible matrix, or if it's a batch of matrices and one or more of them is not an invertible matrix, then info stores a positive integer for the corresponding matrix. The positive integer indicates the diagonal element of the LU decomposition of the input matrix that is exactly zero. info filled with zeros indicates that the inversion was successful. If check\_errors=TRUE and info contains positive integers, then a RuntimeError is thrown. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

#### Usage

```
linalg_inv_ex(A, check_errors = FALSE)
```

## **Arguments**

```
A (Tensor): tensor of shape (*, n, n) where * is zero or more batch dimensions consisting of square matrices.

check_errors (bool, optional): controls whether to check the content of info. Default: FALSE.
```

#### Note

If A is on a CUDA device then this function may synchronize that device with the CPU.

This function is "experimental" and it may change in a future PyTorch release.

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### See Also

```
linalg_inv() is a NumPy compatible variant that always checks for errors.
```

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

# Examples

```
if (torch_is_installed()) {
A <- torch_randn(3, 3)
out <- linalg_inv_ex(A)
}</pre>
```

linalg\_lstsq

Computes a solution to the least squares problem of a system of linear equations.

## **Description**

Letting . be . or . , the **least squares problem** for a linear system . with . is defined as

## Usage

```
linalg_lstsq(A, B, rcond = NULL, ..., driver = NULL)
```

## **Arguments**

A	(Tensor): lhs tensor of shape $(*, m, n)$ where $*$ is zero or more batch dimensions.
В	(Tensor): rhs tensor of shape $(*, m, k)$ where $*$ is zero or more batch dimensions.
rcond	(float, optional): used to determine the effective rank of A. If $rcond = NULL$ , $rcond$ is set to the machine precision of the dtype of A times $max(m, n)$ . Default: $NULL$ .
	currently unused.
driver	(str, optional): name of the LAPACK/MAGMA method to be used. If NULL, 'gelsy' is used for CPU inputs and 'gels' for CUDA inputs. Default: NULL.

linalg\_lstsq 75

#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . denotes the Frobenius norm. Supports inputs of float, double, cfloat and cdouble dtypes.

Also supports batches of matrices, and if the inputs are batches of matrices then the output has the same batch dimensions. driver chooses the LAPACK/MAGMA function that will be used.

For CPU inputs the valid values are 'gels', 'gelsy', 'gelsd, 'gelss'. For CUDA input, the only valid driver is 'gels', which assumes that A is full-rank.

To choose the best driver on CPU consider:

- If A is well-conditioned (its condition number is not too large), or you do not mind some precision loss.
- For a general matrix: 'gelsy' (QR with pivoting) (default)
- If A is full-rank: 'gels' (QR)
- If A is not well-conditioned.
- 'gelsd' (tridiagonal reduction and SVD)
- But if you run into memory issues: 'gelss' (full SVD).

See also the full description of these drivers

rcond is used to determine the effective rank of the matrices in A when driver is one of ('gelsy', 'gelsd', 'gelss'). In this case, if . are the singular values of A in decreasing order, . will be rounded down to zero if . . If rcond = NULL (default), rcond is set to the machine precision of the dtype of A.

This function returns the solution to the problem and some extra information in a list of four tensors (solution, residuals, rank, singular\_values). For inputs A, B of shape (\*, m, n), (\*, m, k) respectively, it cointains

- solution: the least squares solution. It has shape (\*, n, k).
- residuals: the squared residuals of the solutions, that is, . . It has shape equal to the batch dimensions of A. It is computed when m > n and every matrix in A is full-rank, otherwise, it is an empty tensor. If A is a batch of matrices and any matrix in the batch is not full rank, then an empty tensor is returned. This behavior may change in a future PyTorch release.
- rank: tensor of ranks of the matrices in A. It has shape equal to the batch dimensions of A. It is computed when driver is one of ('gelsy', 'gelsd', 'gelss'), otherwise it is an empty tensor.
- singular\_values: tensor of singular values of the matrices in A. It has shape (\*, min(m, n)). It is computed when driver is one of ('gelsd', 'gelss'), otherwise it is an empty tensor.

#### Value

A list (solution, residuals, rank, singular\_values).

#### Warning

The default value of rcond may change in a future PyTorch release. It is therefore recommended to use a fixed value to avoid potential breaking changes.

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#### Note

This function computes X = A\$pinverse() %\*% B in a faster and more numerically stable way than performing the computations separately.

### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

### **Examples**

```
if (torch_is_installed()) {
A <- torch_tensor(rbind(c(10, 2, 3), c(3, 10, 5), c(5, 6, 12)))$unsqueeze(1) # shape (1, 3, 3)
B <- torch_stack(list(
    rbind(c(2, 5, 1), c(3, 2, 1), c(5, 1, 9)),
    rbind(c(4, 2, 9), c(2, 0, 3), c(2, 5, 3))
), dim = 1) # shape (2, 3, 3)
X <- linalg_lstsq(A, B)$solution # A is broadcasted to shape (2, 3, 3)
}</pre>
```

linalg\_matrix\_norm

Computes a matrix norm.

### **Description**

If A is complex valued, it computes the norm of A\$abs() Support input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices: the norm will be computed over the dimensions specified by the 2-tuple dim and the other dimensions will be treated as batch dimensions. The output will have the same batch dimensions.

#### **Usage**

```
linalg_matrix_norm(
   A,
   ord = "fro",
   dim = c(-2, -1),
   keepdim = FALSE,
   dtype = NULL
)
```

linalg\_matrix\_norm 77

## Arguments

A	(Tensor): tensor with two or more dimensions. By default its shape is interpreted as (*, m, n) where * is zero or more batch dimensions, but this behavior can be controlled using dim.
ord	(int, inf, -inf, 'fro', 'nuc', optional): order of norm. Default: 'fro'
dim	(int, Tupleint, optional): dimensions over which to compute the vector or matrix norm. See above for the behavior when dim=NULL. Default: NULL
keepdim	(bool, optional): If set to TRUE, the reduced dimensions are retained in the result as dimensions with size one. Default: FALSE
dtype	dtype (torch_dtype, optional): If specified, the input tensor is cast to dtype before performing the operation, and the returned tensor's type will be dtype. Default: NULL

### **Details**

ord defines the norm that is computed. The following norms are supported:

ord	norm for matrices	norm for vectors
NULL (default)	Frobenius norm	2-norm (see below)
"fro"	Frobenius norm	<ul><li>not supported –</li></ul>
"nuc"	nuclear norm	<ul><li>not supported –</li></ul>
Inf	max(sum(abs(x), dim=2))	<pre>max(abs(x))</pre>
-Inf	min(sum(abs(x), dim=2))	min(abs(x))
0	<ul><li>not supported –</li></ul>	sum(x != 0)
1	max(sum(abs(x), dim=1))	as below
-1	<pre>min(sum(abs(x), dim=1))</pre>	as below
2	largest singular value	as below
-2	smallest singular value	as below
other int or float	<ul><li>not supported –</li></ul>	sum(abs(x)^{ord})^{(1 / ord)}

## See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

# **Examples**

```
if (torch_is_installed()) {
a <- torch_arange(0, 8, dtype = torch_float())$reshape(c(3, 3))
linalg_matrix_norm(a)
linalg_matrix_norm(a, ord = -1)
b <- a$expand(c(2, -1, -1))
linalg_matrix_norm(b)
linalg_matrix_norm(b, dim = c(1, 3))</pre>
```

linalg\_matrix\_power

}

linalg\_matrix\_power

Computes the n-th power of a square matrix for an integer n.

## **Description**

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

## Usage

```
linalg_matrix_power(A, n)
```

## **Arguments**

```
A (Tensor): tensor of shape (*, m, m) where * is zero or more batch dimensions.

n (int): the exponent.
```

### **Details**

If n=0, it returns the identity matrix (or batch) of the same shape as A. If n is negative, it returns the inverse of each matrix (if invertible) raised to the power of abs(n).

## See Also

```
linalg_solve() computes A$inverse() %*% B with a numerically stable algorithm.
```

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(3, 3)
linalg_matrix_power(A, 0)
}</pre>
```

linalg\_matrix\_rank 79

linalg\_matrix\_rank

Computes the numerical rank of a matrix.

## **Description**

The matrix rank is computed as the number of singular values (or eigenvalues in absolute value when hermitian = TRUE) that are greater than the specified tol threshold.

## Usage

```
linalg_matrix_rank(
   A,
   ...,
   atol = NULL,
   rtol = NULL,
   tol = NULL,
   hermitian = FALSE
)
```

## **Arguments**

Α	(Tensor): tensor of shape ( $\star$ , m, n) where $\star$ is zero or more batch dimensions.
	Not currently used.
atol	the absolute tolerance value. When NULL it's considered to be zero.
rtol	the relative tolerance value. See above for the value it takes when NULL.
tol	(float, Tensor, optional): the tolerance value. See above for the value it takes when NULL. Default: NULL.
hermitian	(bool, optional): indicates whether A is Hermitian if complex or symmetric if real. Default: FALSE.

## Details

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

If hermitian = TRUE, A is assumed to be Hermitian if complex or symmetric if real, but this is not checked internally. Instead, just the lower triangular part of the matrix is used in the computations.

If tol is not specified and A is a matrix of dimensions (m, n), the tolerance is set to be

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is the largest singular value (or eigenvalue in absolute value when hermitian = TRUE), and . is the epsilon value for the dtype of A (see torch\_finfo()).

If A is a batch of matrices, tol is computed this way for every element of the batch.

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### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

### **Examples**

```
if (torch_is_installed()) {
a <- torch_eye(10)
linalg_matrix_rank(a)
}</pre>
```

linalg\_multi\_dot

Efficiently multiplies two or more matrices

## Description

Efficiently multiplies two or more matrices by reordering the multiplications so that the fewest arithmetic operations are performed.

### Usage

```
linalg_multi_dot(tensors)
```

#### **Arguments**

tensors

(Sequence[Tensor]): two or more tensors to multiply. The first and last tensors may be 1D or 2D. Every other tensor must be 2D.

#### **Details**

Supports inputs of float, double, cfloat and cdouble dtypes. This function does not support batched inputs.

Every tensor in tensors must be 2D, except for the first and last which may be 1D. If the first tensor is a 1D vector of shape (n,) it is treated as a row vector of shape (1, n), similarly if the last tensor is a 1D vector of shape (n,) it is treated as a column vector of shape (n, 1).

If the first and last tensors are matrices, the output will be a matrix. However, if either is a 1D vector, then the output will be a 1D vector.

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#### Note

This function is implemented by chaining torch\_mm() calls after computing the optimal matrix multiplication order.

The cost of multiplying two matrices with shapes (a, b) and (b, c) is a \* b \* c. Given matrices A, B, C with shapes (10, 100), (100, 5), (5, 50) respectively, we can calculate the cost of different multiplication orders as follows:

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

In this case, multiplying A and B first followed by C is 10 times faster.

#### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_power(), linalg_matrix_rank(), linalg_norm(), linalg_pinv(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
linalg_multi_dot(list(torch_tensor(c(1, 2)), torch_tensor(c(2, 3))))
}
```

linalg\_norm

Computes a vector or matrix norm.

## **Description**

If A is complex valued, it computes the norm of A\$abs() Supports input of float, double, cfloat and cdouble dtypes. Whether this function computes a vector or matrix norm is determined as follows:

## Usage

```
linalg_norm(A, ord = NULL, dim = NULL, keepdim = FALSE, dtype = NULL)
```

### **Arguments**

A	(Tensor): tensor of shape (*, n) or (*, m, n) where * is zero or more batch dimensions
ord	(int, float, inf, -inf, 'fro', 'nuc', optional): order of norm. Default: NULL
dim	(int, Tupleint, optional): dimensions over which to compute the vector or matrix norm. See above for the behavior when dim=NULL. Default: NULL
keepdim	(bool, optional): If set to TRUE, the reduced dimensions are retained in the result as dimensions with size one. Default: FALSE

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dtype

dtype (torch\_dtype, optional): If specified, the input tensor is cast to dtype before performing the operation, and the returned tensor's type will be dtype. Default: NULL

#### **Details**

- If dim is an int, the vector norm will be computed.
- If dim is a 2-tuple, the matrix norm will be computed.
- If dim=NULL and ord=NULL, A will be flattened to 1D and the 2-norm of the resulting vector will be computed.
- If dim=NULL and ord!=NULL, A must be 1D or 2D.

ord defines the norm that is computed. The following norms are supported:

```
norm for matrices
ord
                                                 norm for vectors
NULL (default)
                    Frobenius norm
                                                 2-norm (see below)
"fro"
                    Frobenius norm
                                                 - not supported -
"nuc"
                    nuclear norm
                                                 - not supported -
Inf
                    max(sum(abs(x), dim=2))
                                                 max(abs(x))
-Inf
                    min(sum(abs(x), dim=2))
                                                 min(abs(x))
0
                    not supported –
                                                 sum(x != 0)
1
                    max(sum(abs(x), dim=1))
                                                 as below
                                                 as below
-1
                    min(sum(abs(x), dim=1))
2
                    largest singular value
                                                 as below
-2
                    smallest singular value
                                                 as below
other int or float
                    - not supported -
                                                 sum(abs(x)^{ord})^{(1 / ord)}
```

#### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
a <- torch_arange(0, 8, dtype = torch_float()) - 4
a
b <- a$reshape(c(3, 3))
b

linalg_norm(a)
linalg_norm(b)
}</pre>
```

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linalg_pinv	Computes the pseudoinverse (Moore-Penrose inverse) of a matrix.

## **Description**

The pseudoinverse may be defined algebraically\_but it is more computationally convenient to understand it through the SVD\_Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

### Usage

```
linalg_pinv(A, rcond = NULL, hermitian = FALSE, atol = NULL, rtol = NULL)
```

## **Arguments**

Α	(Tensor): tensor of shape $(*, m, n)$ where $*$ is zero or more batch dimensions.
rcond	(float or Tensor, optional): the tolerance value to determine when is a singular value zero If it is a torch_Tensor, its shape must be broadcastable to that of the singular values of A as returned by linalg_svd(). Alias for rtol. Default: 0.
hermitian	(bool, optional): indicates whether ${\sf A}$ is Hermitian if complex or symmetric if real. Default: FALSE.
atol	the absolute tolerance value. When NULL it's considered to be zero.
rtol	the relative tolerance value. See above for the value it takes when NULL.

#### **Details**

If hermitian= TRUE, A is assumed to be Hermitian if complex or symmetric if real, but this is not checked internally. Instead, just the lower triangular part of the matrix is used in the computations. The singular values (or the norm of the eigenvalues when hermitian= TRUE) that are below the specified roond threshold are treated as zero and discarded in the computation.

### Note

This function uses linalg\_svd() if hermitian= FALSE and linalg\_eigh() if hermitian= TRUE. For CUDA inputs, this function synchronizes that device with the CPU.

Consider using linalg\_lstsq() if possible for multiplying a matrix on the left by the pseudoinverse, as linalg\_lstsq(A, B)\$solution == A\$pinv() %\*% B

It is always prefered to use linalg\_lstsq() when possible, as it is faster and more numerically stable than computing the pseudoinverse explicitly.

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### See Also

- linalg\_inv() computes the inverse of a square matrix.
- linalg\_lstsq() computes A\$pinv() %\*% B with a numerically stable algorithm.

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(3, 5)
linalg_pinv(A)
}</pre>
```

linalg\_qr

Computes the QR decomposition of a matrix.

### **Description**

```
Letting . be . or . , the full QR decomposition of a matrix . is defined as
```

#### Usage

```
linalg_qr(A, mode = "reduced")
```

#### **Arguments**

```
A (Tensor): tensor of shape (*, m, n) where * is zero or more batch dimensions.

mode (str, optional): one of 'reduced', 'complete', 'r'. Controls the shape of the returned tensors. Default: 'reduced'.
```

#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . is orthogonal in the real case and unitary in the complex case, and . is upper triangular. When m > n (tall matrix), as R is upper triangular, its last m - n rows are zero. In this case, we can drop the last m - n columns of Q to form the **reduced QR decomposition**:

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

The reduced QR decomposition agrees with the full QR decomposition when  $n \ge m$  (wide matrix). Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions. The parameter mode chooses between the full and reduced QR decomposition.

```
If A has shape (*, m, n), denoting k = min(m, n)
```

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- mode = 'reduced' (default): Returns (Q, R) of shapes (\*, m, k), (\*, k, n) respectively.
- mode = 'complete': Returns (Q, R) of shapes (\*, m, m), (\*, m, n) respectively.
- mode = 'r': Computes only the reduced R. Returns (Q, R) with Q empty and R of shape (\*, k, n).

#### Value

```
A list (Q, R).
```

#### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_slogdet(), linalg_solve_triangular(), linalg_solve(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(rbind(c(12., -51, 4), c(6, 167, -68), c(-4, 24, -41)))
qr <- linalg_qr(a)

torch_mm(qr[[1]], qr[[2]])$round()
torch_mm(qr[[1]]$t(), qr[[1]])$round()
}</pre>
```

 $linalg\_slogdet$ 

Computes the sign and natural logarithm of the absolute value of the determinant of a square matrix.

## **Description**

For complex A, it returns the angle and the natural logarithm of the modulus of the determinant, that is, a logarithmic polar decomposition of the determinant. Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

# Usage

```
linalg_slogdet(A)
```

## Arguments

A (Tensor): tensor of shape (\*, n, n) where \* is zero or more batch dimensions.

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#### Value

A list (sign, logabsdet). logabsdet will always be real-valued, even when A is complex. sign will have the same dtype as A.

#### Notes

- The determinant can be recovered as sign \* exp(logabsdet).
- When a matrix has a determinant of zero, it returns (0, -Inf).

#### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_solve_triangular(), linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

### **Examples**

```
if (torch_is_installed()) {
a <- torch_randn(3, 3)
linalg_slogdet(a)
}</pre>
```

linalg\_solve

Computes the solution of a square system of linear equations with a unique solution.

# Description

Letting . be . or . , this function computes the solution . of the **linear system** associated to . , which is defined as

## Usage

```
linalg_solve(A, B)
```

#### **Arguments**

```
A (Tensor): tensor of shape (*, n, n) where * is zero or more batch dimensions.
```

B (Tensor): right-hand side tensor of shape (\*, n) or (\*, n, k) or (n,) or (n, k) according to the rules described above

#### **Details**

$$AX = B$$

This system of linear equations has one solution if and only if . is invertible\_. This function assumes that . is invertible. Supports inputs of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if the inputs are batches of matrices then the output has the same batch dimensions.

Letting \* be zero or more batch dimensions,

- If A has shape (\*, n, n) and B has shape (\*, n) (a batch of vectors) or shape (\*, n, k) (a batch of matrices or "multiple right-hand sides"), this function returns X of shape (\*, n) or (\*, n, k) respectively.
- Otherwise, if A has shape (\*, n, n) and B has shape (n,) or (n, k), B is broadcasted to have shape (\*, n) or (\*, n, k) respectively.

This function then returns the solution of the resulting batch of systems of linear equations.

#### Note

This function computes X = A\$inverse() @ B in a faster and more numerically stable way than performing the computations separately.

#### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangularelinalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

## **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(3, 3)
b <- torch_randn(3)
x <- linalg_solve(A, b)
torch_allclose(torch_matmul(A, x), b)
}</pre>
```

linalg\_solve\_triangular

Triangular solve

# Description

Triangular solve

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### Usage

```
linalg_solve_triangular(A, B, ..., upper, left = TRUE, unitriangular = FALSE)
```

### **Arguments**

A tensor of shape (\*, n, n) or (\*, k, k) if left=TRUE) where \* is zero or more

if TRUE, the diagonal elements of A are assumed to be all equal to 1.

batch dimensions.

B right-hand side tensor of shape (\*, n, k)

... Currently ignored.

upper whether A is an upper or lower triangular matrix.

left wheter to solve the system AX=B or XA=B

#### See Also

unitriangular

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve(), linalg_svdvalsh(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

linalg\_svd

Computes the singular value decomposition (SVD) of a matrix.

# Description

```
Letting . be . or . , the full SVD of a matrix . , if k = min(m, n), is defined as
```

#### **Usage**

```
linalg_svd(A, full_matrices = TRUE)
```

#### **Arguments**

A (Tensor): tensor of shape (\*, m, n) where \* is zero or more batch dimensions.

full\_matrices (bool, optional): controls whether to compute the full or reduced SVD, and

consequently, the shape of the returned tensors U and V. Default: TRUE.

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#### **Details**

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . , . is the conjugate transpose when . is complex, and the transpose when . is real-valued.

The matrices . , . (and thus . ) are orthogonal in the real case, and unitary in the complex case. When m > n (resp. m < n) we can drop the last m - n (resp. n - m) columns of U (resp. V) to form the **reduced SVD**:

Equation not displayed. Please find it in 'https://torch.mlverse.org/docs'

where . .

In this case, . and . also have orthonormal columns. Supports input of float, double, cfloat and cdouble dtypes.

Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions.

The returned decomposition is a named tuple (U, S, V) which corresponds to . . . . . above.

The singular values are returned in descending order. The parameter full\_matrices chooses between the full (default) and reduced SVD.

#### Value

A list (U, S, V) which corresponds to . , . , . above. S will always be real-valued, even when A is complex. It will also be ordered in descending order. U and V will have the same dtype as A. The left / right singular vectors will be given by the columns of U and the rows of V respectively.

### Warnings

The returned tensors U and V are not unique, nor are they continuous with respect to A. Due to this lack of uniqueness, different hardware and software may compute different singular vectors. This non-uniqueness is caused by the fact that multiplying any pair of singular vectors . by -1 in the real case or by . in the complex case produces another two valid singular vectors of the matrix. This non-uniqueness problem is even worse when the matrix has repeated singular values. In this case, one may multiply the associated singular vectors of U and V spanning the subspace by a rotation matrix and the resulting vectors will span the same subspace.

Gradients computed using U or V will only be finite when A does not have zero as a singular value or repeated singular values. Furthermore, if the distance between any two singular values is close to zero, the gradient will be numerically unstable, as it depends on the singular values . through the computation of . . The gradient will also be numerically unstable when A has small singular values, as it also depends on the computation of . .

#### Note

When full\_matrices=TRUE, the gradients with respect to U[..., :, min(m, n):] and Vh[..., min(m, n):, :] will be ignored, as those vectors can be arbitrary bases of the corresponding subspaces.

#### See Also

• linalg\_svdvals() computes only the singular values. Unlike linalg\_svd(), the gradients of linalg\_svdvals() are always numerically stable.

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- linalg\_eig() for a function that computes another type of spectral decomposition of a matrix. The eigendecomposition works just on on square matrices.
- linalg\_eigh() for a (faster) function that computes the eigenvalue decomposition for Hermitian and symmetric matrices.
- linalg\_qr() for another (much faster) decomposition that works on general matrices.

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvals(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_invex(), linalg_invex(), linalg_invex(), linalg_invex(), linalg_invex(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(linalg_solve(), linalg_svdvals(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

#### **Examples**

```
if (torch_is_installed()) {
a <- torch_randn(5, 3)
linalg_svd(a, full_matrices = FALSE)
}</pre>
```

linalg\_svdvals

Computes the singular values of a matrix.

### **Description**

Supports input of float, double, cfloat and cdouble dtypes. Also supports batches of matrices, and if A is a batch of matrices then the output has the same batch dimensions. The singular values are returned in descending order.

### Usage

```
linalg_svdvals(A)
```

#### **Arguments**

A (Tensor): tensor of shape (\*, m, n) where \* is zero or more batch dimensions.

## Value

A real-valued tensor, even when A is complex.

#### See Also

```
linalg_svd() computes the full singular value decomposition.
```

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(linalg_solve(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve(), linalg_vector_norm()
```

linalg\_tensorinv 91

#### **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(5, 3)
S <- linalg_svdvals(A)
S
}</pre>
```

linalg\_tensorinv

Computes the multiplicative inverse of torch\_tensordot()

## **Description**

If m is the product of the first ind dimensions of A and n is the product of the rest of the dimensions, this function expects m and n to be equal. If this is the case, it computes a tensor X such that tensordot(A, X, ind) is the identity matrix in dimension m.

## Usage

```
linalg_tensorinv(A, ind = 3L)
```

## Arguments

```
A (Tensor): tensor to invert.

ind (int): index at which to compute the inverse of torch_tensordot(). Default:

3.
```

#### **Details**

Supports input of float, double, cfloat and cdouble dtypes.

### Note

```
Consider using linalg_tensorsolve() if possible for multiplying a tensor on the left by the tensor inverse as linalg_tensorsolve(A, B) == torch_tensordot(linalg_tensorinv(A), B))
```

It is always prefered to use linalg\_tensorsolve() when possible, as it is faster and more numerically stable than computing the pseudoinverse explicitly.

# See Also

• linalg\_tensorsolve() computes torch\_tensordot(linalg\_tensorinv(A), B)).

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_inv_ex(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorsolve(), linalg_vector_norm()
```

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### **Examples**

```
if (torch_is_installed()) {
A <- torch_eye(4 * 6)$reshape(c(4, 6, 8, 3))
Ainv <- linalg_tensorinv(A, ind = 3)
Ainv$shape
B <- torch_randn(4, 6)
torch_allclose(torch_tensordot(Ainv, B), linalg_tensorsolve(A, B))
A <- torch_randn(4, 4)
Atensorinv <- linalg_tensorinv(A, 2)
Ainv <- linalg_inv(A)
torch_allclose(Atensorinv, Ainv)
}</pre>
```

linalg\_tensorsolve

Computes the solution X to the system torch\_tensordot(A, X) = B.

### **Description**

If m is the product of the first B\ .ndim dimensions of A and n is the product of the rest of the dimensions, this function expects m and n to be equal. The returned tensor x satisfies tensordot(A, x, dims=x\$ndim) == B.

### Usage

```
linalg_tensorsolve(A, B, dims = NULL)
```

#### **Arguments**

```
A (Tensor): tensor to solve for.

B (Tensor): the solution

dims (Tupleint, optional): dimensions of A to be moved. If NULL, no dimensions are moved. Default: NULL.
```

#### **Details**

```
If dims is specified, A will be reshaped as A = movedim(A, dims, seq(len(dims) - A$ndim + 1, 0))
```

Supports inputs of float, double, cfloat and cdouble dtypes.

### See Also

• linalg\_tensorinv() computes the multiplicative inverse of torch\_tensordot().

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_vector_norm()
```

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### **Examples**

```
if (torch_is_installed()) {
A <- torch_eye(2 * 3 * 4)$reshape(c(2 * 3, 4, 2, 3, 4))
B <- torch_randn(2 * 3, 4)
X <- linalg_tensorsolve(A, B)
X$shape
torch_allclose(torch_tensordot(A, X, dims = X$ndim), B)

A <- torch_randn(6, 4, 4, 3, 2)
B <- torch_randn(4, 3, 2)
X <- linalg_tensorsolve(A, B, dims = c(1, 3))
A <- A$permute(c(2, 4, 5, 1, 3))
torch_allclose(torch_tensordot(A, X, dims = X$ndim), B, atol = 1e-6)
}</pre>
```

linalg\_vector\_norm

Computes a vector norm.

# Description

If A is complex valued, it computes the norm of A\$abs() Supports input of float, double, cfloat and cdouble dtypes. This function does not necessarily treat multidimensonal A as a batch of vectors, instead:

## Usage

```
linalg_vector_norm(A, ord = 2, dim = NULL, keepdim = FALSE, dtype = NULL)
```

### **Arguments**

A	(Tensor): tensor, flattened by default, but this behavior can be controlled using dim.
ord	(int, float, inf, -inf, 'fro', 'nuc', optional): order of norm. Default: 2
dim	(int, Tupleint, optional): dimensions over which to compute the vector or matrix norm. See above for the behavior when dim=NULL. Default: NULL
keepdim	(bool, optional): If set to TRUE, the reduced dimensions are retained in the result as dimensions with size one. Default: FALSE
dtype	dtype (torch_dtype, optional): If specified, the input tensor is cast to dtype before performing the operation, and the returned tensor's type will be dtype. Default: NULL

#### **Details**

• If dim=NULL, A will be flattened before the norm is computed.

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• If dim is an int or a tuple, the norm will be computed over these dimensions and the other dimensions will be treated as batch dimensions.

This behavior is for consistency with linalg\_norm().

ord defines the norm that is computed. The following norms are supported:

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```
ord
                     norm for matrices
                                                 norm for vectors
NULL (default)
                     Frobenius norm
                                                 2-norm (see below)
"fro"
                     Frobenius norm
                                                 - not supported -
"nuc"
                     nuclear norm
                                                 - not supported -
Inf
                    max(sum(abs(x), dim=2))
                                                 max(abs(x))
-Inf
                    min(sum(abs(x), dim=2))
                                                 min(abs(x))
                     - not supported -
                                                 sum(x != 0)
                     max(sum(abs(x), dim=1))
                                                 as below
1
                     min(sum(abs(x), dim=1))
                                                 as below
-1
2
                     largest singular value
                                                 as below
-2
                     smallest singular value
                                                 as below
                    - not supported -
other int or float
                                                 sum(abs(x)^{ord})^{(1/ord)}
```

### See Also

```
Other linalg: linalg_cholesky_ex(), linalg_cholesky(), linalg_det(), linalg_eigh(), linalg_eigvalsh(), linalg_eigvalsh(), linalg_eigvals(), linalg_eig(), linalg_householder_product(), linalg_inv_ex(), linalg_inv(), linalg_lstsq(), linalg_matrix_norm(), linalg_matrix_power(), linalg_matrix_rank(), linalg_multi_dot(), linalg_norm(), linalg_pinv(), linalg_qr(), linalg_slogdet(), linalg_solve_triangular(linalg_solve(), linalg_svdvals(), linalg_svd(), linalg_tensorinv(), linalg_tensorsolve()
```

### **Examples**

```
if (torch_is_installed()) {
a <- torch_arange(0, 8, dtype = torch_float()) - 4
a
b <- a$reshape(c(3, 3))
b

linalg_vector_norm(a, ord = 3.5)
linalg_vector_norm(b, ord = 3.5)
}</pre>
```

load\_state\_dict

Load a state dict file

### **Description**

This function should only be used to load models saved in python. For it to work correctly you need to use torch.save with the flag: \_use\_new\_zipfile\_serialization=True and also remove all nn.Parameter classes from the tensors in the dict.

## Usage

```
load_state_dict(path, ..., legacy_stream = FALSE)
```

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## **Arguments**

```
path to the state dict file
... additional arguments that are currently not used.

legacy_stream if TRUE then the state dict is loaded using a a legacy way of handling streams.
```

### **Details**

The above might change with development of this in pytorch's C++ api.

### Value

a named list of tensors.

local\_autocast

Autocast context manager

## **Description**

Allow regions of your code to run in mixed precision. In these regions, ops run in an op-specific dtype chosen by autocast to improve performance while maintaining accuracy.

## Usage

```
local_autocast(
  device_type,
  dtype = NULL,
  enabled = TRUE,
  cache_enabled = NULL,
  ...,
  .env = parent.frame()
)

with_autocast(
  code,
  ...,
  device_type,
  dtype = NULL,
  enabled = TRUE,
  cache_enabled = NULL
)

set_autocast(device_type, dtype = NULL, enabled = TRUE, cache_enabled = NULL)
unset_autocast(context)
```

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## Arguments

device\_type a character string indicating whether to use 'cuda' or 'cpu' device
dtype a torch data type indicating whether to use torch\_float16() or torch\_bfloat16().
enabled a logical value indicating whether autocasting should be enabled in the region.
Default: TRUE

cache\_enabled a logical value indicating whether the weight cache inside autocast should be enabled.

... currently unused.
.env The environment to use for scoping.
code code to be executed with no gradient recording.

Returned by set\_autocast and should be passed when unsetting it.

#### **Details**

context

When entering an autocast-enabled region, Tensors may be any type. You should not call half() or bfloat16() on your model(s) or inputs when using autocasting.

autocast should only be enabled during the forward pass(es) of your network, including the loss computation(s). Backward passes under autocast are not recommended. Backward ops run in the same type that autocast used for corresponding forward ops.

#### **Functions**

- with\_autocast(): A with context for automatic mixed precision.
- set\_autocast(): Set the autocast context. For advanced users only.
- unset\_autocast(): Unset the autocast context.

#### See Also

cuda\_amp\_grad\_scaler() to perform dynamic gradient scaling.

## **Examples**

```
if (torch_is_installed()) {
  x <- torch_randn(5, 5, dtype = torch_float32())
  y <- torch_randn(5, 5, dtype = torch_float32())

foo <- function(x, y) {
  local_autocast(device = "cpu")
  z <- torch_mm(x, y)
  w <- torch_mm(z, x)
  w
}

out <- foo(x, y)
}</pre>
```

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local\_device

Device contexts

## **Description**

Device contexts

### Usage

```
local_device(device, ..., .env = parent.frame())
with_device(code, ..., device)
```

## **Arguments**

device A torch device to be used by default when creating new tensors.

... currently unused.

. env The environment to use for scoping.

code The code to be evaluated in the modified environment.

## **Functions**

• with\_device(): Modifies the default device for the selected context.

lr\_cosine\_annealing

Set the learning rate of each parameter group using a cosine annealing schedule

## **Description**

Set the learning rate of each parameter group using a cosine annealing schedule

### Usage

```
lr_cosine_annealing(
  optimizer,
  T_max,
  eta_min = 0,
  last_epoch = -1,
  verbose = FALSE
)
```

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#### **Arguments**

optimizer	(Optimizer): Wrapped optimizer.
T_max	Maximum number of iterations
eta_min	Minimum learning rate. Default: 0.

last\_epoch The index of the last epoch

verbose (bool): If TRUE, prints a message to stdout for each update. Default: FALSE.

1r\_lambda Sets the learning rate of each parameter group to the initial lr times a

given function. When last\_epoch=-1, sets initial lr as lr.

# Description

Sets the learning rate of each parameter group to the initial lr times a given function. When last\_epoch=-1, sets initial lr as lr.

# Usage

```
lr_lambda(optimizer, lr_lambda, last_epoch = -1, verbose = FALSE)
```

### **Arguments**

optimizer (Optimizer): Wrapped optimizer.

1r\_lambda (function or list): A function which computes a multiplicative factor given an

integer parameter epoch, or a list of such functions, one for each group in opti-

mizer.param\_groups.

last\_epoch (int): The index of last epoch. Default: -1.

verbose (bool): If TRUE, prints a message to stdout for each update. Default: FALSE.

#### **Examples**

```
if (torch_is_installed()) {
# Assuming optimizer has two groups.
lambda1 <- function(epoch) epoch %/% 30
lambda2 <- function(epoch) 0.95^epoch
## Not run:
scheduler <- lr_lambda(optimizer, lr_lambda = list(lambda1, lambda2))
for (epoch in 1:100) {
    train(...)
    validate(...)
    scheduler$step()
}
## End(Not run)</pre>
```

lr\_one\_cycle

lr_multiplicative	Multiply the learning rate of each parameter group by the factor given
	in the specified function. When last_epoch=-1, sets initial lr as lr.

## **Description**

Multiply the learning rate of each parameter group by the factor given in the specified function. When last\_epoch=-1, sets initial lr as lr.

## Usage

```
lr_multiplicative(optimizer, lr_lambda, last_epoch = -1, verbose = FALSE)
```

## **Arguments**

optimizer (Optimizer): Wrapped optimizer.

1r\_lambda (function or list): A function which computes a multiplicative factor given an integer parameter epoch, or a list of such functions, one for each group in optimizer.param\_groups.

last\_epoch (int): The index of last epoch. Default: -1.

verbose (bool): If TRUE, prints a message to stdout for each update. Default: FALSE.

### **Examples**

```
if (torch_is_installed()) {
## Not run:
lmbda <- function(epoch) 0.95
scheduler <- lr_multiplicative(optimizer, lr_lambda = lmbda)
for (epoch in 1:100) {
    train(...)
    validate(...)
    scheduler$step()
}
## End(Not run)
}</pre>
```

lr\_one\_cycle

Once cycle learning rate

### **Description**

Sets the learning rate of each parameter group according to the 1 cycle learning rate policy. The 1 cycle policy anneals the learning rate from an initial learning rate to some maximum learning rate and then from that maximum learning rate to some minimum learning rate much lower than the initial learning rate.

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### **Usage**

```
lr_one_cycle(
  optimizer,
 max_lr,
  total_steps = NULL,
  epochs = NULL,
  steps_per_epoch = NULL,
  pct_start = 0.3,
  anneal_strategy = "cos",
  cycle_momentum = TRUE,
  base_momentum = 0.85,
 max_momentum = 0.95,
  div_factor = 25,
  final_div_factor = 10000,
  last_epoch = -1,
  verbose = FALSE
)
```

### **Arguments**

optimizer (Optimizer): Wrapped optimizer.

max\_lr (float or list): Upper learning rate boundaries in the cycle for each parameter

group.

total\_steps (int): The total number of steps in the cycle. Note that if a value is not provided

here, then it must be inferred by providing a value for epochs and steps per epoch.

Default: NULL

epochs (int): The number of epochs to train for. This is used along with steps\_per\_epoch

in order to infer the total number of steps in the cycle if a value for total\_steps is

not provided. Default: NULL

steps\_per\_epoch

(int): The number of steps per epoch to train for. This is used along with epochs in order to infer the total number of steps in the cycle if a value for total\_steps is

not provided. Default: NULL

pct\_start (float): The percentage of the cycle (in number of steps) spent increasing the

learning rate. Default: 0.3

anneal\_strategy

(str): {'cos', 'linear'} Specifies the annealing strategy: "cos" for cosine anneal-

ing, "linear" for linear annealing. Default: 'cos'

cycle\_momentum (bool): If TRUE, momentum is cycled inversely to learning rate between 'base\_momentum'

and 'max momentum'. Default: TRUE

base\_momentum (float or list): Lower momentum boundaries in the cycle for each parameter

group. Note that momentum is cycled inversely to learning rate; at the peak of a cycle, momentum is 'base\_momentum' and learning rate is 'max\_lr'. Default:

0.85

max\_momentum (float or list): Upper momentum boundaries in the cycle for each parameter

group. Functionally, it defines the cycle amplitude (max\_momentum - base\_momentum).

lr\_one\_cycle

Note that momentum is cycled inversely to learning rate; at the start of a cycle, momentum is 'max\_momentum' and learning rate is 'base\_lr' Default: 0.95

div\_factor (float): Determines the initial learning rate via initial\_lr = max\_lr/div\_factor Default: 25

. .

final\_div\_factor

(float): Determines the minimum learning rate via min\_lr = initial\_lr/final\_div\_factor

Default: 1e4

last\_epoch

(int): The index of the last batch. This parameter is used when resuming a training job. Since step() should be invoked after each batch instead of after each epoch, this number represents the total number of *batches* computed, not the total number of epochs computed. When last\_epoch=-1, the schedule is storted from the bacinning. Default: 1

started from the beginning. Default: -1

verbose

(bool): If TRUE, prints a message to stdout for each update. Default: FALSE.

#### **Details**

This policy was initially described in the paper Super-Convergence: Very Fast Training of Neural Networks Using Large Learning Rates.

The 1 cycle learning rate policy changes the learning rate after every batch. step should be called after a batch has been used for training. This scheduler is not chainable.

Note also that the total number of steps in the cycle can be determined in one of two ways (listed in order of precedence):

- A value for total\_steps is explicitly provided.
- A number of epochs (epochs) and a number of steps per epoch (steps\_per\_epoch) are provided.

In this case, the number of total steps is inferred by total steps = epochs \* steps per epoch

You must either provide a value for total\_steps or provide a value for both epochs and steps\_per\_epoch.

### **Examples**

```
if (torch_is_installed()) {
## Not run:
data_loader <- dataloader(...)</pre>
optimizer <- optim_sgd(model$parameters, lr = 0.1, momentum = 0.9)
scheduler <- lr_one_cycle(optimizer,</pre>
 max_lr = 0.01, steps_per_epoch = length(data_loader),
 epochs = 10
)
for (i in 1:epochs) {
 coro::loop(for (batch in data_loader) {
    train_batch(...)
    scheduler$step()
 })
}
## End(Not run)
}
```

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lr\_reduce\_on\_plateau Reduce learning rate on plateau

# Description

Reduce learning rate when a metric has stopped improving. Models often benefit from reducing the learning rate by a factor of 2-10 once learning stagnates. This scheduler reads a metrics quantity and if no improvement is seen for a 'patience' number of epochs, the learning rate is reduced.

# Usage

```
lr_reduce_on_plateau(
  optimizer,
  mode = "min",
  factor = 0.1,
  patience = 10,
  threshold = 1e-04,
  threshold_mode = "rel",
  cooldown = 0,
  min_lr = 0,
  eps = 1e-08,
  verbose = FALSE
)
```

## **Arguments**

optimizer	(Optimizer): Wrapped optimizer.
mode	(str): One of min, max. In min mode, lr will be reduced when the quantity monitored has stopped decreasing; in max mode it will be reduced when the quantity monitored has stopped increasing. Default: 'min'.
factor	(float): Factor by which the learning rate will be reduced. new_lr <- lr * factor. Default: 0.1.
patience	(int): Number of epochs with no improvement after which learning rate will be reduced. For example, if patience = 2, then we will ignore the first 2 epochs with no improvement, and will only decrease the LR after the 3rd epoch if the loss still hasn't improved then. Default: 10.
threshold	(float):Threshold for measuring the new optimum, to only focus on significant changes. Default: 1e-4.
threshold_mode	(str): One of rel, abs. In rel mode, dynamic_threshold <- best * (1 + threshold) in 'max' mode or best * (1 - threshold) in min mode. In abs mode, dynamic_threshold <- best + threshold in max mode or best - threshold in min mode. Default: 'rel'.
cooldown	(int): Number of epochs to wait before resuming normal operation after lr has

been reduced. Default: 0.

lr\_scheduler

min\_lr (float or list): A scalar or a list of scalars. A lower bound on the learning rate of all param groups or each group respectively. Default: 0.

eps (float): Minimal decay applied to lr. If the difference between new and old lr is smaller than eps, the update is ignored. Default: 1e-8.

verbose (bool): If TRUE, prints a message to stdout for each update. Default: FALSE.

### **Examples**

```
if (torch_is_installed()) {
## Not run:
optimizer <- optim_sgd(model$parameters(), lr=0.1, momentum=0.9)
scheduler <- lr_reduce_on_plateau(optimizer, 'min')
for (epoch in 1:10) {
   train(...)
   val_loss <- validate(...)
   # note that step should be called after validate
   scheduler$step(val_loss)
}
## End(Not run)
}</pre>
```

lr\_scheduler

Creates learning rate schedulers

## Description

Creates learning rate schedulers

## Usage

```
lr_scheduler(
  classname = NULL,
  inherit = LRScheduler,
   ...,
  parent_env = parent.frame()
)
```

### **Arguments**

classname optional name for the learning rate scheduler

inherit an optional learning rate scheduler to inherit from

... named list of methods. You must implement the get\_lr() method that doesn't take any argument and returns learning rates for each param\_group in the optimizer.

parent\_env passed to R6::R6Class().

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lr\_step

Step learning rate decay

## **Description**

Decays the learning rate of each parameter group by gamma every step\_size epochs. Notice that such decay can happen simultaneously with other changes to the learning rate from outside this scheduler. When last\_epoch=-1, sets initial lr as lr.

### Usage

```
lr_step(optimizer, step_size, gamma = 0.1, last_epoch = -1)
```

## Arguments

```
optimizer (Optimizer): Wrapped optimizer.

step_size (int): Period of learning rate decay.

gamma (float): Multiplicative factor of learning rate decay. Default: 0.1.

last_epoch (int): The index of last epoch. Default: -1.
```

## **Examples**

```
{\it nnf\_adaptive\_avg\_pool1d} \\ Adaptive\_avg\_pool1d
```

# Description

Applies a 1D adaptive average pooling over an input signal composed of several input planes.

# Usage

```
nnf_adaptive_avg_pool1d(input, output_size)
```

# Arguments

input input tensor of shape (minibatch , in\_channels , iW)

output\_size the target output size (single integer)

```
nnf\_adaptive\_avg\_pool2d \\ Adaptive\_avg\_pool2d
```

# Description

Applies a 2D adaptive average pooling over an input signal composed of several input planes.

# Usage

```
nnf_adaptive_avg_pool2d(input, output_size)
```

# Arguments

input input tensor (minibatch, in\_channels, iH, iW)

output\_size the target output size (single integer or double-integer tuple)

```
nnf_adaptive_avg_pool3d
```

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```
nnf_adaptive_avg_pool3d
```

Adaptive\_avg\_pool3d

## **Description**

Applies a 3D adaptive average pooling over an input signal composed of several input planes.

# Usage

```
nnf_adaptive_avg_pool3d(input, output_size)
```

## **Arguments**

input input tensor (minibatch, in\_channels, iT \* iH, iW)

output\_size the target output size (single integer or triple-integer tuple)

nnf\_adaptive\_max\_pool1d

Adaptive\_max\_pool1d

# **Description**

Applies a 1D adaptive max pooling over an input signal composed of several input planes.

# Usage

```
nnf_adaptive_max_pool1d(input, output_size, return_indices = FALSE)
```

## **Arguments**

input input tensor of shape (minibatch, in\_channels, iW)

output\_size the target output size (single integer)

return\_indices whether to return pooling indices. Default: FALSE

```
nnf\_adaptive\_max\_pool2d \\ Adaptive\_max\_pool2d
```

# Description

Applies a 2D adaptive max pooling over an input signal composed of several input planes.

## Usage

```
nnf_adaptive_max_pool2d(input, output_size, return_indices = FALSE)
```

## **Arguments**

```
input input tensor (minibatch, in_channels, iH, iW)
```

output\_size the target output size (single integer or double-integer tuple)

return\_indices whether to return pooling indices. Default: FALSE

```
nnf\_adaptive\_max\_pool3d \\ Adaptive\_max\_pool3d
```

## **Description**

Applies a 3D adaptive max pooling over an input signal composed of several input planes.

## Usage

```
nnf_adaptive_max_pool3d(input, output_size, return_indices = FALSE)
```

### **Arguments**

```
input input tensor (minibatch, in_channels, iT * iH, iW)
```

output\_size the target output size (single integer or triple-integer tuple)

return\_indices whether to return pooling indices. Default:FALSE

nnf\_affine\_grid 109

Affine_grid
-------------

#### **Description**

Generates a 2D or 3D flow field (sampling grid), given a batch of affine matrices theta.

## Usage

```
nnf_affine_grid(theta, size, align_corners = FALSE)
```

## **Arguments**

theta (Tensor) input batch of affine matrices with shape  $(N \times 2 \times 3)$  for 2D or  $(N \times 2 \times 3)$ 

 $3 \times 4$ ) for 3D

size (torch.Size) the target output image size.  $(N \times C \times H \times W \text{ for 2D or } N \times C \times H \times W \text{ for 2D$ 

 $D \times H \times W$  for 3D) Example: torch.Size((32, 3, 24, 24))

align\_corners (bool, optional) if True, consider -1 and 1 to refer to the centers of the cor-

ner pixels rather than the image corners. Refer to nnf\_grid\_sample() for a more complete description. A grid generated by nnf\_affine\_grid() should be passed to nnf\_grid\_sample() with the same setting for this option. Default:

False

#### Note

 $This function is often used in conjunction with \verb|nnf_grid_sample()| to build Spatial Transformer Networks\_|$ 

#### **Description**

Applies alpha dropout to the input.

## Usage

```
nnf_alpha_dropout(input, p = 0.5, training = FALSE, inplace = FALSE)
```

#### **Arguments**

input the input tensor

p probability of an element to be zeroed. Default: 0.5

training apply dropout if is TRUE. Default: TRUE

inplace If set to TRUE, will do this operation in-place. Default: FALSE

nnf\_avg\_pool2d

nnf\_avg\_pool1d

Avg\_pool1d

## **Description**

Applies a 1D average pooling over an input signal composed of several input planes.

## Usage

```
nnf_avg_pool1d(
   input,
   kernel_size,
   stride = NULL,
   padding = 0,
   ceil_mode = FALSE,
   count_include_pad = TRUE
)
```

## **Arguments**

input input tensor of shape (minibatch, in\_channels, iW)

kernel\_size the size of the window. Can be a single number or a tuple (kW,).

stride the stride of the window. Can be a single number or a tuple (sW,). Default:

kernel\_size

padding implicit zero paddings on both sides of the input. Can be a single number or a

tuple (padW,). Default: 0

ceil\_mode when True, will use ceil instead of floor to compute the output shape. Default:

**FALSE** 

count\_include\_pad

when True, will include the zero-padding in the averaging calculation. Default:

TRUE

nnf\_avg\_pool2d

Avg\_pool2d

# Description

Applies 2D average-pooling operation in kH\*kW regions by step size sH\*sW steps. The number of output features is equal to the number of input planes.

nnf\_avg\_pool3d

#### Usage

```
nnf_avg_pool2d(
  input,
  kernel_size,
  stride = NULL,
  padding = 0,
  ceil_mode = FALSE,
  count_include_pad = TRUE,
  divisor_override = NULL
)
```

#### **Arguments**

input input tensor (minibatch, in\_channels, iH, iW)

kernel\_size size of the pooling region. Can be a single number or a tuple (kH, kW)

stride stride of the pooling operation. Can be a single number or a tuple (sH, sW).

Default: kernel\_size

padding implicit zero paddings on both sides of the input. Can be a single number or a

tuple (padH, padW). Default: 0

ceil\_mode when True, will use ceil instead of floor in the formula to compute the output

shape. Default: FALSE

count\_include\_pad

when True, will include the zero-padding in the averaging calculation. Default:

TRUE

divisor\_override

if specified, it will be used as divisor, otherwise size of the pooling region will

be used. Default: NULL

nnf\_avg\_pool3d

Avg\_pool3d

## **Description**

Applies 3D average-pooling operation in kT \* kH \* kW regions by step size sT \* sH \* sW steps. The number of output features is equal to  $\left|\frac{\text{input planes}}{sT}\right|$ .

#### Usage

```
nnf_avg_pool3d(
   input,
   kernel_size,
   stride = NULL,
   padding = 0,
   ceil_mode = FALSE,
   count_include_pad = TRUE,
   divisor_override = NULL
)
```

nnf\_batch\_norm

#### **Arguments**

NA if specified, it will be used as divisor, otherwise size of the pooling region

will be used. Default: NULL

nnf\_batch\_norm Batch\_norm

#### **Description**

Applies Batch Normalization for each channel across a batch of data.

#### Usage

```
nnf_batch_norm(
   input,
   running_mean,
   running_var,
   weight = NULL,
   bias = NULL,
   training = FALSE,
   momentum = 0.1,
   eps = 1e-05
)
```

## **Arguments**

input input tensor

running\_mean the running\_mean tensor
running\_var the running\_var tensor
weight the weight tensor
bias the bias tensor

training bool wether it's training. Default: FALSE

nnf\_bilinear 113

momentum	the value used for the running_mean and running_var computation. Can be
	set to None for cumulative moving average (i.e. simple average). Default: 0.1
eps	a value added to the denominator for numerical stability. Default: 1e-5

nnf\_bilinear

## **Description**

Applies a bilinear transformation to the incoming data:  $y = x_1Ax_2 + b$ 

#### Usage

```
nnf_bilinear(input1, input2, weight, bias = NULL)
```

Bilinear

# Arguments

```
input1 (N,*,H_{in1}) where H_{in1}= in1_features and * means any number of additional dimensions. All but the last dimension of the inputs should be the same. (N,*,H_{in2}) \text{ where } H_{in2}=\text{in2\_features} weight (\text{out\_features},\text{in1\_features},\text{in2\_features}) bias (\text{out\_features})
```

#### Value

output  $(N, *, H_{out})$  where  $H_{out} = \text{out\_features}$  and all but the last dimension are the same shape as the input.

```
nnf_binary_cross_entropy

**Binary_cross_entropy**
```

# Description

Function that measures the Binary Cross Entropy between the target and the output.

# Usage

```
nnf_binary_cross_entropy(
  input,
  target,
  weight = NULL,
  reduction = c("mean", "sum", "none")
)
```

#### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor (N,\*), same shape as the input

weight (tensor) weight for each value.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

nnf\_binary\_cross\_entropy\_with\_logits

Binary\_cross\_entropy\_with\_logits

## **Description**

Function that measures Binary Cross Entropy between target and output logits.

#### Usage

```
nnf_binary_cross_entropy_with_logits(
  input,
  target,
  weight = NULL,
  reduction = c("mean", "sum", "none"),
  pos_weight = NULL
)
```

## **Arguments**

input Tensor of arbitrary shape

target Tensor of the same shape as input

weight (Tensor, optional) a manual rescaling weight if provided it's repeated to match

input tensor shape.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

pos\_weight (Tensor, optional) a weight of positive examples. Must be a vector with length

equal to the number of classes.

nnf\_celu 115

nnf\_celu

Celu

#### **Description**

```
Applies element-wise, CELU(x) = max(0, x) + min(0, \alpha * (exp(x\alpha) - 1)).
```

# Usage

```
nnf_celu(input, alpha = 1, inplace = FALSE)
nnf_celu_(input, alpha = 1)
```

## **Arguments**

input (N,\*) tensor, where \* means, any number of additional dimensions

alpha the alpha value for the CELU formulation. Default: 1.0 inplace can optionally do the operation in-place. Default: FALSE

nnf\_contrib\_sparsemax Sparsemax

# Description

Applies the SparseMax activation.

# Usage

```
nnf_contrib_sparsemax(input, dim = -1)
```

# Arguments

input the input tensor

dim The dimension over which to apply the sparsemax function. (-1)

## **Details**

The SparseMax activation is described in 'From Softmax to Sparsemax: A Sparse Model of Attention and Multi-Label Classification' The implementation is based on aced125/sparsemax

nnf\_conv2d

# Description

Applies a 1D convolution over an input signal composed of several input planes.

# Usage

```
nnf_conv1d(
  input,
  weight,
  bias = NULL,
  stride = 1,
  padding = 0,
  dilation = 1,
  groups = 1
)
```

# Arguments

input	input tensor of shape (minibatch, in_channels , iW)
weight	filters of shape (out_channels, in_channels/groups , kW)
bias	optional bias of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a one-element tuple (sW,). Default: $1$
padding	implicit paddings on both sides of the input. Can be a single number or a one-element tuple (padW,). Default: $0$
dilation	the spacing between kernel elements. Can be a single number or a one-element tuple (dW,). Default: $1$
groups	split input into groups, in_channels should be divisible by the number of groups. Default: 1

_conv2d Conv2d

# Description

Applies a 2D convolution over an input image composed of several input planes.

nnf\_conv3d

# Usage

```
nnf_conv2d(
  input,
  weight,
  bias = NULL,
  stride = 1,
  padding = 0,
  dilation = 1,
  groups = 1
)
```

# Arguments

input	input tensor of shape (minibatch, in_channels, iH, iW)
weight	filters of shape (out_channels , in_channels/groups, $kH$ , $kW$ )
bias	optional bias tensor of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a tuple (sH, sW). Default: 1
padding	implicit paddings on both sides of the input. Can be a single number or a tuple (padH, padW). Default: $0$
dilation	the spacing between kernel elements. Can be a single number or a tuple (dH, dW). Default: $1$
groups	split input into groups, in_channels should be divisible by the number of groups. Default: 1

nnf\_conv3d Conv3d

# Description

Applies a 3D convolution over an input image composed of several input planes.

# Usage

```
nnf_conv3d(
  input,
  weight,
  bias = NULL,
  stride = 1,
  padding = 0,
  dilation = 1,
  groups = 1
)
```

nnf\_conv\_tbc

## **Arguments**

input	input tensor of shape (minibatch, in_channels , iT , iH , iW)
weight	filters of shape (out_channels , in_channels/groups, $kT$ , $kH$ , $kW)$
bias	optional bias tensor of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a tuple (sT, sH, sW). Default: 1
padding	implicit paddings on both sides of the input. Can be a single number or a tuple (padT, padH, padW). Default: $0$
dilation	the spacing between kernel elements. Can be a single number or a tuple (dT, dH, dW). Default: $1$
groups	split input into groups, in_channels should be divisible by the number of groups. Default: 1

nnf_conv_tbc
--------------

# Description

Applies a 1-dimensional sequence convolution over an input sequence. Input and output dimensions are (Time, Batch, Channels) - hence TBC.

# Usage

```
nnf_conv_tbc(input, weight, bias, pad = 0)
```

input	input tensor of shape (sequence length $\times  batch \times \text{in\_channels})$
weight	$filter\ of\ shape\ (kernel\ width \times in\_channels \times out\_channels)$
bias	bias of shape (out_channels)
pad	number of timesteps to pad. Default: 0

nnf\_conv\_transpose1d 119

# Description

Applies a 1D transposed convolution operator over an input signal composed of several input planes, sometimes also called "deconvolution".

# Usage

```
nnf_conv_transpose1d(
  input,
  weight,
  bias = NULL,
  stride = 1,
  padding = 0,
  output_padding = 0,
  groups = 1,
  dilation = 1
)
```

input	input tensor of shape (minibatch, in_channels, iW)
weight	filters of shape (out_channels, in_channels/groups , $kW)$
bias	optional bias of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a one-element tuple (sW,). Default: $1$
padding	implicit paddings on both sides of the input. Can be a single number or a one-element tuple (padW,). Default: $\bf 0$
output_padding	padding applied to the output
groups	split input into groups, in_channels should be divisible by the number of groups. Default: $\boldsymbol{1}$
dilation	the spacing between kernel elements. Can be a single number or a one-element tuple (dW,). Default: $1$

```
nnf_conv_transpose2d Conv_transpose2d
```

# Description

Applies a 2D transposed convolution operator over an input image composed of several input planes, sometimes also called "deconvolution".

# Usage

```
nnf_conv_transpose2d(
   input,
   weight,
   bias = NULL,
   stride = 1,
   padding = 0,
   output_padding = 0,
   groups = 1,
   dilation = 1
)
```

input	input tensor of shape (minibatch, in_channels, iH, iW)
weight	filters of shape (out_channels , in_channels/groups, $kH$ , $kW)$
bias	optional bias tensor of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a tuple (sH, sW). Default: $1$
padding	implicit paddings on both sides of the input. Can be a single number or a tuple (padH, padW). Default: $0$
output_padding	padding applied to the output
groups	split input into groups, in_channels should be divisible by the number of groups. Default: $\boldsymbol{1}$
dilation	the spacing between kernel elements. Can be a single number or a tuple (dH, dW). Default: $1$

nnf\_conv\_transpose3d 121

```
nnf_conv_transpose3d Conv_transpose3d
```

# Description

Applies a 3D transposed convolution operator over an input image composed of several input planes, sometimes also called "deconvolution"

# Usage

```
nnf_conv_transpose3d(
  input,
  weight,
  bias = NULL,
  stride = 1,
  padding = 0,
  output_padding = 0,
  groups = 1,
  dilation = 1
)
```

input	input tensor of shape (minibatch, in_channels , iT , iH , iW)
weight	filters of shape (out_channels , in_channels/groups, $kT$ , $kH$ , $kW)$
bias	optional bias tensor of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a tuple (sT, sH, sW). Default: 1
padding	implicit paddings on both sides of the input. Can be a single number or a tuple (padT, padH, padW). Default: $0$
output_padding	padding applied to the output
groups	split input into groups, in_channels should be divisible by the number of groups. Default: 1
dilation	the spacing between kernel elements. Can be a single number or a tuple ( $dT$ , $dH$ , $dW$ ). Default: 1

122 nnf\_cosine\_similarity

```
nnf_cosine_embedding_loss
```

Cosine\_embedding\_loss

## Description

Creates a criterion that measures the loss given input tensors  $x_1$ ,  $x_2$  and a Tensor label y with values 1 or -1. This is used for measuring whether two inputs are similar or dissimilar, using the cosine distance, and is typically used for learning nonlinear embeddings or semi-supervised learning.

#### Usage

```
nnf_cosine_embedding_loss(
  input1,
  input2,
  target,
  margin = 0,
  reduction = c("mean", "sum", "none")
)
```

#### **Arguments**

input1 the input x\_1 tensor input2 the input x\_2 tensor target the target tensor

margin Should be a number from -1 to 1, 0 to 0.5 is suggested. If margin is missing,

the default value is 0.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

```
nnf_cosine_similarity Cosine_similarity
```

#### **Description**

Returns cosine similarity between x1 and x2, computed along dim.

#### Usage

```
nnf_cosine_similarity(x1, x2, dim = 2, eps = 1e-08)
```

nnf\_cross\_entropy 123

# Arguments

x1	(Tensor) First input.
x2	(Tensor) Second input (of size matching x1).
dim	(int, optional) Dimension of vectors. Default: 2
eps	(float, optional) Small value to avoid division by zero. Default: 1e-8

# **Details**

$$\text{similarity} = \frac{x_1 \cdot x_2}{\max(\|x_1\|_2 \cdot \|x_2\|_2, \epsilon)}$$

nnf\_cross\_entropy

Cross\_entropy

# Description

This criterion combines log\_softmax and nll\_loss in a single function.

# Usage

```
nnf_cross_entropy(
  input,
  target,
  weight = NULL,
  ignore_index = -100,
  reduction = c("mean", "sum", "none")
)
```

input	(Tensor) $(N,C)$ where C = number of classes or $(N,C,H,W)$ in case of 2D Loss, or $(N,C,d_1,d_2,,d_K)$ where $K\geq 1$ in the case of K-dimensional loss.
target	(Tensor) $(N)$ where each value is $0 \le \text{targets}[i] \le C-1$ , or $(N,d_1,d_2,,d_K)$ where $K \ge 1$ for K-dimensional loss.
weight	(Tensor, optional) a manual rescaling weight given to each class. If given, has to be a Tensor of size C
ignore_index	(int, optional) Specifies a target value that is ignored and does not contribute to the input gradient.
reduction	(string, optional) – Specifies the reduction to apply to the output: 'none' I 'mean' I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed. Default: 'mean'

nnf\_ctc\_loss

nnf\_ctc\_loss

Ctc\_loss

## **Description**

The Connectionist Temporal Classification loss.

# Usage

```
nnf_ctc_loss(
  log_probs,
  targets,
  input_lengths,
  target_lengths,
  blank = 0,
  reduction = c("mean", "sum", "none"),
  zero_infinity = FALSE
)
```

#### **Arguments**

log\_probs (T, N, C) where C = number of characters in alphabet including blank, T =

input length, and N = batch size. The logarithmized probabilities of the outputs

(e.g. obtained with nnf\_log\_softmax).

targets (N,S) or (sum(target\_lengths)). Targets cannot be blank. In the second

form, the targets are assumed to be concatenated.

input\_lengths (N). Lengths of the inputs (must each be  $\leq T$ )

 $target_lengths$  (N). Lengths of the targets

blank (int, optional) Blank label. Default 0.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

zero\_infinity (bool, optional) Whether to zero infinite losses and the associated gradients.

Default: FALSE Infinite losses mainly occur when the inputs are too short to be

aligned to the targets.

nnf\_dropout 125

|--|--|--|

## **Description**

During training, randomly zeroes some of the elements of the input tensor with probability p using samples from a Bernoulli distribution.

## Usage

```
nnf_dropout(input, p = 0.5, training = TRUE, inplace = FALSE)
```

#### **Arguments**

input	the input tensor
Input	the input tensor

p probability of an element to be zeroed. Default: 0.5

training apply dropout if is TRUE. Default: TRUE

inplace If set to TRUE, will do this operation in-place. Default: FALSE

## **Description**

Randomly zero out entire channels (a channel is a 2D feature map, e.g., the j-th channel of the i-th sample in the batched input is a 2D tensor input[i,j]) of the input tensor). Each channel will be zeroed out independently on every forward call with probability p using samples from a Bernoulli distribution.

## Usage

```
nnf_dropout2d(input, p = 0.5, training = TRUE, inplace = FALSE)
```

#### **Arguments**

r

p probability of a channel to be zeroed. Default: 0.5

training apply dropout if is TRUE. Default: TRUE.

inplace If set to TRUE, will do this operation in-place. Default: FALSE

nnf\_elu

nnf_dropout3d	Dropout3d
---------------	-----------

## Description

Randomly zero out entire channels (a channel is a 3D feature map, e.g., the j-th channel of the i-th sample in the batched input is a 3D tensor input[i,j]) of the input tensor). Each channel will be zeroed out independently on every forward call with probability p using samples from a Bernoulli distribution.

## Usage

```
nnf_dropout3d(input, p = 0.5, training = TRUE, inplace = FALSE)
```

## **Arguments**

input the input tensor

p probability of a channel to be zeroed. Default: 0.5

training apply dropout if is TRUE. Default: TRUE.

inplace If set to TRUE, will do this operation in-place. Default: FALSE

|--|

#### **Description**

Applies element-wise,

```
ELU(x) = max(0, x) + min(0, \alpha * (exp(x) - 1))
```

# Usage

```
nnf_elu(input, alpha = 1, inplace = FALSE)
nnf_elu_(input, alpha = 1)
```

#### **Arguments**

input	(N *)	tensor	where *	means	anv	number	of:	additional	dimensio	ns
IIIDUL	(14, /	tenson,	WIICIC	micans,	anv	Humber '	$\sigma$	auunuonai	uninchisto.	пэ

alpha the alpha value for the ELU formulation. Default: 1.0 inplace can optionally do the operation in-place. Default: FALSE

nnf\_embedding 127

## **Examples**

```
if (torch_is_installed()) {
x <- torch_randn(2, 2)
y <- nnf_elu(x, alpha = 1)
nnf_elu_(x, alpha = 1)
torch_equal(x, y)
}</pre>
```

nnf\_embedding

Embedding

# Description

A simple lookup table that looks up embeddings in a fixed dictionary and size.

## Usage

```
nnf_embedding(
  input,
  weight,
  padding_idx = NULL,
  max_norm = NULL,
  norm_type = 2,
  scale_grad_by_freq = FALSE,
  sparse = FALSE
)
```

input	(LongTensor) Tensor containing indices into the embedding matrix
weight	(Tensor) The embedding matrix with number of rows equal to the maximum possible index + 1, and number of columns equal to the embedding size
padding_idx	(int, optional) If given, pads the output with the embedding vector at padding_idx (initialized to zeros) whenever it encounters the index.
max_norm	(float, optional) If given, each embedding vector with norm larger than max_norm is renormalized to have norm max_norm. Note: this will modify weight inplace.
norm_type	(float, optional) The p of the p-norm to compute for the max_norm option. Default 2.
scale_grad_by_f	Freq
	(boolean, optional) If given, this will scale gradients by the inverse of frequency of the words in the mini-batch. Default FALSE.
sparse	(bool, optional) If TRUE, gradient w.r.t. weight will be a sparse tensor. See Notes under nn_embedding for more details regarding sparse gradients.

nnf\_embedding\_bag

## **Details**

This module is often used to retrieve word embeddings using indices. The input to the module is a list of indices, and the embedding matrix, and the output is the corresponding word embeddings.

nnf\_embedding\_bag

Embedding\_bag

## Description

Computes sums, means or maxes of bags of embeddings, without instantiating the intermediate embeddings.

# Usage

```
nnf_embedding_bag(
  input,
  weight,
  offsets = NULL,
  max_norm = NULL,
  norm_type = 2,
  scale_grad_by_freq = FALSE,
  mode = "mean",
  sparse = FALSE,
  per_sample_weights = NULL,
  include_last_offset = FALSE,
  padding_idx = NULL
)
```

#### **Arguments**

input	(LongTensor) Tensor containing bags of indices into the embedding matrix		
weight	(Tensor) The embedding matrix with number of rows equal to the maximum possible index + 1, and number of columns equal to the embedding size		
offsets	(LongTensor, optional) Only used when input is 1D. offsets determines the starting index position of each bag (sequence) in input.		
max_norm	(float, optional) If given, each embedding vector with norm larger than max_norm is renormalized to have norm max_norm. Note: this will modify weight inplace.		
norm_type	(float, optional) The p in the p-norm to compute for the max_norm option. Default 2.		
scale_grad_by_freq			
	(boolean, optional) if given, this will scale gradients by the inverse of frequency		

when mode="max".

of the words in the mini-batch. Default FALSE. Note: this option is not supported

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mode (string, optional) "sum", "mean" or "max". Specifies the way to reduce the bag.

Default: 'mean'

sparse (bool, optional) if TRUE, gradient w.r.t. weight will be a sparse tensor. See

Notes under nn\_embedding for more details regarding sparse gradients. Note:

this option is not supported when mode="max".

per\_sample\_weights

(Tensor, optional) a tensor of float / double weights, or NULL to indicate all weights should be taken to be 1. If specified, per\_sample\_weights must have exactly the same shape as input and is treated as having the same offsets, if

those are not NULL.

include\_last\_offset

(bool, optional) if TRUE, the size of offsets is equal to the number of bags + 1.

padding\_idx

(int, optional) If given, pads the output with the embedding vector at  $padding\_idx$ 

(initialized to zeros) whenever it encounters the index.

nnf\_fold

Fold

#### **Description**

Combines an array of sliding local blocks into a large containing tensor.

## Usage

```
nnf_fold(
  input,
  output_size,
  kernel_size,
  dilation = 1,
  padding = 0,
  stride = 1
)
```

## **Arguments**

input the input tensor

 $output\_size \qquad \text{the shape of the spatial dimensions of the output (i.e., output \$sizes()[-c(1,2)])}$ 

kernel\_size the size of the sliding blocks

dilation a parameter that controls the stride of elements within the neighborhood. De-

fault: 1

padding implicit zero padding to be added on both sides of input. Default: 0 stride the stride of the sliding blocks in the input spatial dimensions. Default: 1

## Warning

Currently, only 4-D output tensors (batched image-like tensors) are supported.

# Description

Applies 2D fractional max pooling over an input signal composed of several input planes.

## Usage

```
nnf_fractional_max_pool2d(
   input,
   kernel_size,
   output_size = NULL,
   output_ratio = NULL,
   return_indices = FALSE,
   random_samples = NULL
)
```

## **Arguments**

input	the input tensor
kernel_size	the size of the window to take a max over. Can be a single number $k$ (for a square kernel of $k \ast k$ ) or a tuple (kH, kW)
output_size	the target output size of the image of the form $oH * oW$ . Can be a tuple (oH, oW) or a single number $oH$ for a square image $oH * oH$
output_ratio	If one wants to have an output size as a ratio of the input size, this option can be given. This has to be a number or tuple in the range $(0,1)$
return_indices	if True, will return the indices along with the outputs.
random_samples	optional random samples.

#### **Details**

Fractional MaxPooling is described in detail in the paper Fractional MaxPooling\_ by Ben Graham

The max-pooling operation is applied in kH \* kW regions by a stochastic step size determined by the target output size. The number of output features is equal to the number of input planes.

# Description

Applies 3D fractional max pooling over an input signal composed of several input planes.

## Usage

```
nnf_fractional_max_pool3d(
  input,
  kernel_size,
  output_size = NULL,
  output_ratio = NULL,
  return_indices = FALSE,
  random_samples = NULL
)
```

## Arguments

input	the input tensor
kernel_size	the size of the window to take a max over. Can be a single number $k$ (for a square kernel of $k*k*k$ ) or a tuple (kT, kH, kW)
output_size	the target output size of the form $oT*oH*oW$ . Can be a tuple (oT, oH, oW) or a single number $oH$ for a cubic output $oH*oH*oH$
output_ratio	If one wants to have an output size as a ratio of the input size, this option can be given. This has to be a number or tuple in the range $(0, 1)$
return_indices	if True, will return the indices along with the outputs.
random_samples	undocumented argument.

#### **Details**

Fractional MaxPooling is described in detail in the paper Fractional MaxPooling\_ by Ben Graham

The max-pooling operation is applied in kT \* kH \* kW regions by a stochastic step size determined by the target output size. The number of output features is equal to the number of input planes.

nnf\_glu

nnf\_gelu

Gelu

## **Description**

Gelu

#### Usage

```
nnf_gelu(input, approximate = "none")
```

## **Arguments**

input (N,\*) tensor, where \* means, any number of additional dimensions

approximate By default it's none, and applies element-wise x\*pnorm(x), if 'tanh', then GELU

is estimated. See GELU for more info.

# gelu(input) -> Tensor

Applies element-wise the function  $GELU(x) = x * \Phi(x)$ 

where  $\Phi(x)$  is the Cumulative Distribution Function for Gaussian Distribution.

See Gaussian Error Linear Units (GELUs).

nnf\_glu

Glu

# Description

The gated linear unit. Computes:

# Usage

```
nnf_glu(input, dim = -1)
```

## **Arguments**

input (Tensor) input tensor

dim (int) dimension on which to split the input. Default: -1

#### **Details**

$$GLU(a,b) = a \otimes \sigma(b)$$

where input is split in half along dim to form a and b,  $\sigma$  is the sigmoid function and  $\otimes$  is the element-wise product between matrices.

See Language Modeling with Gated Convolutional Networks.

nnf\_grid\_sample 133

#### **Description**

Given an input and a flow-field grid, computes the output using input values and pixel locations from grid.

## Usage

```
nnf_grid_sample(
  input,
  grid,
  mode = c("bilinear", "nearest"),
  padding_mode = c("zeros", "border", "reflection"),
  align_corners = FALSE
)
```

## **Arguments**

input	(Tensor) input of shape $(N, C, H_{in}, W_{in})$ (4-D case) or $(N, C, D_{in}, H_{in}, W_{in})$ (5-D case)
grid	(Tensor) flow-field of shape $(N,H_{\hbox{out}},W_{\hbox{out}},2)$ (4-D case) or $(N,D_{\hbox{out}},H_{\hbox{out}},W_{\hbox{out}},3)$ (5-D case)
mode	(str) interpolation mode to calculate output values 'bilinear'   'nearest'.  Default: 'bilinear'
padding_mode	(str) padding mode for outside grid values 'zeros' 'border' 'reflection'.  Default: 'zeros'
align_corners	(bool, optional) Geometrically, we consider the pixels of the input as squares rather than points. If set to True, the extrema (-1 and 1) are considered as referring to the center points of the input's corner pixels. If set to False, they are instead considered as referring to the corner points of the input's corner pixels, making the sampling more resolution agnostic. This option parallels the align_corners option in nnf_interpolate(), and so whichever option is used here should also be used there to resize the input image before grid sampling. Default: False

## **Details**

Currently, only spatial (4-D) and volumetric (5-D) input are supported.

In the spatial (4-D) case, for input with shape  $(N, C, H_{\rm in}, W_{\rm in})$  and grid with shape  $(N, H_{\rm out}, W_{\rm out}, 2)$ , the output will have shape  $(N, C, H_{\rm out}, W_{\rm out})$ .

For each output location output[n, :, h, w], the size-2 vector grid[n, h, w] specifies input pixel locations x and y, which are used to interpolate the output value output[n, :, h, w]. In the case of 5D inputs, grid[n, d, h, w] specifies the x, y, z pixel locations for interpolating

nnf\_group\_norm

output[n, :, d, h, w]. mode argument specifies nearest or bilinear interpolation method to sample the input pixels.

grid specifies the sampling pixel locations normalized by the input spatial dimensions. Therefore, it should have most values in the range of [-1, 1]. For example, values x = -1, y = -1 is the left-top pixel of input, and values x = 1, y = 1 is the right-bottom pixel of input.

If grid has values outside the range of [-1, 1], the corresponding outputs are handled as defined by padding\_mode. Options are

- padding\_mode="zeros": use 0 for out-of-bound grid locations,
- padding\_mode="border": use border values for out-of-bound grid locations,
- padding\_mode="reflection": use values at locations reflected by the border for out-of-bound grid locations. For location far away from the border, it will keep being reflected until becoming in bound, e.g., (normalized) pixel location x = -3.5 reflects by border -1 and becomes x' = 1.5, then reflects by border 1 and becomes x'' = -0.5.

#### Note

This function is often used in conjunction with nnf\_affine\_grid() to build Spatial Transformer Networks\_ .

nnf\_group\_norm Group\_norm

#### Description

Applies Group Normalization for last certain number of dimensions.

#### Usage

```
nnf_group_norm(input, num_groups, weight = NULL, bias = NULL, eps = 1e-05)
```

## Arguments

input the input tensor

num\_groups number of groups to separate the channels into

weight the weight tensor
bias the bias tensor

eps a value added to the denominator for numerical stability. Default: 1e-5

nnf\_gumbel\_softmax 135

nnf_gumbel_softmax	Gumbel_softmax
--------------------	----------------

## **Description**

Samples from the Gumbel-Softmax distribution and optionally discretizes.

#### Usage

```
nnf_gumbel_softmax(logits, tau = 1, hard = FALSE, dim = -1)
```

#### **Arguments**

logits [..., num\_features] unnormalized log probabilities

tau non-negative scalar temperature

hard if True, the returned samples will be discretized as one-hot vectors, but will be

differentiated as if it is the soft sample in autograd

dim (int) A dimension along which softmax will be computed. Default: -1.

nnf\_hardshrink *Hardshrink* 

#### **Description**

Applies the hard shrinkage function element-wise

## Usage

```
nnf_hardshrink(input, lambd = 0.5)
```

# Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions

lambd the lambda value for the Hardshrink formulation. Default: 0.5

nnf\_hardswish

nnf\_hardsigmoid

Hardsigmoid

## **Description**

Applies the element-wise function  $\operatorname{Hardsigmoid}(x) = \frac{ReLU6(x+3)}{6}$ 

## Usage

```
nnf_hardsigmoid(input, inplace = FALSE)
```

#### **Arguments**

input (N,\*) tensor, where \* means, any number of additional dimensions inplace NA If set to True, will do this operation in-place. Default: False

nnf\_hardswish

Hardswish

## **Description**

Applies the hardswish function, element-wise, as described in the paper: Searching for MobileNetV3.

## Usage

```
nnf_hardswish(input, inplace = FALSE)
```

## **Arguments**

input (N,\*) tensor, where \* means, any number of additional dimensions

inplace can optionally do the operation in-place. Default: FALSE

#### **Details**

$$\operatorname{Hardswish}(x) = \left\{ \begin{array}{ll} 0 & \text{if } x \leq -3, \\ x & \text{if } x \geq +3, \\ x \cdot (x+3)/6 & \text{otherwise} \end{array} \right.$$

nnf\_hardtanh 137

## Description

Applies the HardTanh function element-wise.

## Usage

```
nnf_hardtanh(input, min_val = -1, max_val = 1, inplace = FALSE)
nnf_hardtanh_(input, min_val = -1, max_val = 1)
```

## Arguments

input	(N,*) tensor, where * means, any number of additional dimensions
min_val	minimum value of the linear region range. Default: -1
max_val	maximum value of the linear region range. Default: 1
inplace	can optionally do the operation in-place. Default: FALSE

```
nnf_hinge_embedding_loss
```

Hinge\_embedding\_loss

#### **Description**

Measures the loss given an input tensor xx and a labels tensor yy (containing 1 or -1). This is usually used for measuring whether two inputs are similar or dissimilar, e.g. using the L1 pairwise distance as xx, and is typically used for learning nonlinear embeddings or semi-supervised learning.

## Usage

```
nnf_hinge_embedding_loss(input, target, margin = 1, reduction = "mean")
```

#### **Arguments**

input	tensor (N,*)	where **	means, and	v number	of additional	dimensions

target tensor (N,\*), same shape as the input

margin Has a default value of 1.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

nnf\_interpolate

nnf\_instance\_norm

Instance\_norm

# Description

Applies Instance Normalization for each channel in each data sample in a batch.

## Usage

```
nnf_instance_norm(
  input,
  running_mean = NULL,
  running_var = NULL,
  weight = NULL,
  bias = NULL,
  use_input_stats = TRUE,
  momentum = 0.1,
  eps = 1e-05
)
```

## **Arguments**

input the input tensor

running\_mean the running\_mean tensor the running var tensor

weight the weight tensor bias the bias tensor

use\_input\_stats

whether to use input stats

momentum a double for the momentum

eps an eps double for numerical stability

nnf\_interpolate

Interpolate

## **Description**

Down/up samples the input to either the given size or the given scale\_factor

nnf\_interpolate 139

#### Usage

```
nnf_interpolate(
  input,
  size = NULL,
  scale_factor = NULL,
  mode = "nearest",
  align_corners = FALSE,
  recompute_scale_factor = NULL
)
```

#### **Arguments**

input (Tensor) the input tensor

size (int or Tuple[int] or Tuple[int, int] or Tuple[int, int, int]) output spa-

tial size.

scale\_factor (float or Tuple[float]) multiplier for spatial size. Has to match input size if it

is a tuple.

mode (str) algorithm used for upsampling: 'nearest' | 'linear' | 'bilinear' | 'bicubic' |

'trilinear' | 'area' Default: 'nearest'

align\_corners (bool, optional) Geometrically, we consider the pixels of the input and output

as squares rather than points. If set to TRUE, the input and output tensors are aligned by the center points of their corner pixels, preserving the values at the corner pixels. If set to False, the input and output tensors are aligned by the corner points of their corner pixels, and the interpolation uses edge value padding for out-of-boundary values, making this operation *independent* of input size when scale\_factor is kept the same. This only has an effect when mode

is 'linear', 'bilinear', 'bicubic' or 'trilinear'. Default: False

recompute\_scale\_factor

(bool, optional) recompute the scale\_factor for use in the interpolation calculation. When scale\_factor is passed as a parameter, it is used to compute the output\_size. If recompute\_scale\_factor is "'True" or not specified, a new scale\_factor will be computed based on the output and input sizes for use in the interpolation computation (i.e. the computation will be identical to if the computed 'output\_size' were passed-in explicitly). Otherwise, the passed-in 'scale\_factor' will be used in the interpolation computation. Note that when 'scale\_factor' is floating-point, the recomputed scale\_factor may differ from the one passed in due to rounding and precision issues.

#### **Details**

The algorithm used for interpolation is determined by mode.

Currently temporal, spatial and volumetric sampling are supported, i.e. expected inputs are 3-D, 4-D or 5-D in shape.

The input dimensions are interpreted in the form:  $mini-batch \ x \ channels \ x \ [optional depth] \ x \ [optional height] \ x \ value of the modes available for resizing are: nearest, linear (3D-only), bilinear, bicubic (4D-only), trilinear (5D-only), area$ 

nnf\_11\_loss

$nnf_{-}$	レ1	div
HILL	KΙ	arv

 $Kl\_div$ 

## Description

The Kullback-Leibler divergence Loss.

## Usage

```
nnf_kl_div(input, target, reduction = "mean")
```

## **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor(N,\*), same shape as the input

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

nnf\_l1\_loss

L1\_loss

## **Description**

Function that takes the mean element-wise absolute value difference.

## Usage

```
nnf_l1_loss(input, target, reduction = "mean")
```

## **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor (N,\*), same shape as the input

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

nnf\_layer\_norm 141

nnf\_layer\_norm

Layer\_norm

#### Description

Applies Layer Normalization for last certain number of dimensions.

## Usage

```
nnf_layer_norm(
   input,
   normalized_shape,
   weight = NULL,
   bias = NULL,
   eps = 1e-05
)
```

## **Arguments**

input the input tensor

normalized\_shape

input shape from an expected input of size. If a single integer is used, it is treated as a singleton list, and this module will normalize over the last dimension which

is expected to be of that specific size.

weight the weight tensor bias the bias tensor

eps a value added to the denominator for numerical stability. Default: 1e-5

nnf\_leaky\_relu

Leaky\_relu

# Description

```
Applies element-wise, LeakyReLU(x) = max(0, x) + negative_slope * min(0, x)
```

## Usage

```
nnf_leaky_relu(input, negative_slope = 0.01, inplace = FALSE)
```

## **Arguments**

```
input (N,*) tensor, where * means, any number of additional dimensions
```

negative\_slope Controls the angle of the negative slope. Default: 1e-2 inplace can optionally do the operation in-place. Default: FALSE

nnf\_linear

Linear

## **Description**

Applies a linear transformation to the incoming data:  $y = xA^T + b$ .

# Usage

```
nnf_linear(input, weight, bias = NULL)
```

## **Arguments**

```
input (N,*,in\_features) where * means any number of additional dimensions weight (out\_features,in\_features) the weights tensor. bias optional tensor (out\_features)
```

```
nnf_local_response_norm
```

Local\_response\_norm

# Description

Applies local response normalization over an input signal composed of several input planes, where channels occupy the second dimension. Applies normalization across channels.

#### Usage

```
nnf_local_response_norm(input, size, alpha = 1e-04, beta = 0.75, k = 1)
```

input	the input tensor
size	amount of neighbouring channels used for normalization
alpha	multiplicative factor. Default: 0.0001
beta	exponent. Default: 0.75
k	additive factor. Default: 1

nnf\_logsigmoid 143

nnf\_logsigmoid

Logsigmoid

## **Description**

Applies element-wise 
$$LogSigmoid(x_i) = log(\frac{1}{1 + exp(-x_i)})$$

## Usage

```
nnf_logsigmoid(input)
```

#### **Arguments**

input

(N,\*) tensor, where \* means, any number of additional dimensions

nnf\_log\_softmax

Log\_softmax

## **Description**

Applies a softmax followed by a logarithm.

#### Usage

```
nnf_log_softmax(input, dim = NULL, dtype = NULL)
```

#### **Arguments**

input (Tensor) input

dim (int) A dimension along which log\_softmax will be computed.

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

#### **Details**

While mathematically equivalent to log(softmax(x)), doing these two operations separately is slower, and numerically unstable. This function uses an alternative formulation to compute the output and gradient correctly.

nnf\_lp\_pool2d

#### **Description**

Applies a 1D power-average pooling over an input signal composed of several input planes. If the sum of all inputs to the power of p is zero, the gradient is set to zero as well.

## Usage

```
nnf_lp_pool1d(input, norm_type, kernel_size, stride = NULL, ceil_mode = FALSE)
```

#### **Arguments**

input the input tensor

norm\_type if inf than one gets max pooling if 0 you get sum pooling (proportional to the

avg pooling)

kernel\_size a single int, the size of the window

stride a single int, the stride of the window. Default value is kernel\_size ceil\_mode when True, will use ceil instead of floor to compute the output shape

 $nnf_lp_pool2d$   $Lp_pool2d$ 

## **Description**

Applies a 2D power-average pooling over an input signal composed of several input planes. If the sum of all inputs to the power of p is zero, the gradient is set to zero as well.

#### Usage

```
nnf_lp_pool2d(input, norm_type, kernel_size, stride = NULL, ceil_mode = FALSE)
```

#### Arguments

input the input tensor

norm\_type if inf than one gets max pooling if 0 you get sum pooling (proportional to the

avg pooling)

kernel\_size a single int, the size of the window

stride a single int, the stride of the window. Default value is kernel\_size ceil\_mode when True, will use ceil instead of floor to compute the output shape

### **Description**

Creates a criterion that measures the loss given inputs x1, x2, two 1D mini-batch Tensors, and a label 1D mini-batch tensor y (containing 1 or -1).

## Usage

```
nnf_margin_ranking_loss(input1, input2, target, margin = 0, reduction = "mean")
```

## Arguments

input1 the first tensor
input2 the second input tensor
target the target tensor
margin Has a default value of 00.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

# Description

Applies a 1D max pooling over an input signal composed of several input planes.

```
nnf_max_pool1d(
  input,
  kernel_size,
  stride = NULL,
  padding = 0,
  dilation = 1,
  ceil_mode = FALSE,
  return_indices = FALSE)
```

nnf\_max\_pool2d

#### **Arguments**

input input tensor of shape (minibatch, in\_channels, iW) kernel\_size the size of the window. Can be a single number or a tuple (kW,). stride the stride of the window. Can be a single number or a tuple (sW,). Default: kernel\_size padding implicit zero paddings on both sides of the input. Can be a single number or a tuple (padW,). Default: 0 dilation controls the spacing between the kernel points; also known as the à trous algorithm. ceil\_mode when True, will use ceil instead of floor to compute the output shape. Default: **FALSE** return\_indices whether to return the indices where the max occurs.

#### **Description**

Applies a 2D max pooling over an input signal composed of several input planes.

#### Usage

```
nnf_max_pool2d(
  input,
  kernel_size,
  stride = kernel_size,
  padding = 0,
  dilation = 1,
  ceil_mode = FALSE,
  return_indices = FALSE)
```

# Arguments

input input tensor (minibatch, in\_channels, iH, iW)

kernel\_size size of the pooling region. Can be a single number or a tuple (kH, kW)

stride stride of the pooling operation. Can be a single number or a tuple (sH, sW).

Default: kernel\_size

padding implicit zero paddings on both sides of the input. Can be a single number or a

tuple (padH, padW). Default: 0

dilation controls the spacing between the kernel points; also known as the à trous algo-

rithm.

ceil\_mode when True, will use ceil instead of floor in the formula to compute the output

shape. Default: FALSE

return\_indices whether to return the indices where the max occurs.

nnf\_max\_pool3d 147

# Description

Applies a 3D max pooling over an input signal composed of several input planes.

# Usage

```
nnf_max_pool3d(
  input,
  kernel_size,
  stride = NULL,
  padding = 0,
  dilation = 1,
  ceil_mode = FALSE,
  return_indices = FALSE
)
```

# Arguments

input	input tensor (minibatch, in_channels , iT * iH , iW)
kernel_size	size of the pooling region. Can be a single number or a tuple (kT, kH, kW)
stride	stride of the pooling operation. Can be a single number or a tuple (sT, sH, sW). Default: $kernel_size$
padding	implicit zero paddings on both sides of the input. Can be a single number or a tuple (padT, padH, padW), Default: $\bf 0$
dilation	controls the spacing between the kernel points; also known as the à trous algorithm.
ceil_mode	when True, will use $\operatorname{ceil}$ instead of floor in the formula to compute the output shape
return_indices	whether to return the indices where the max occurs.

nnf_max_unpool1d	Max_unpool1d
------------------	--------------

# Description

Computes a partial inverse of MaxPool1d.

148 nnf\_max\_unpool2d

#### Usage

```
nnf_max_unpool1d(
   input,
   indices,
   kernel_size,
   stride = NULL,
   padding = 0,
   output_size = NULL)
```

### Arguments

input the input Tensor to invert

indices the indices given out by max pool kernel\_size Size of the max pooling window.

stride Stride of the max pooling window. It is set to kernel\_size by default.

padding Padding that was added to the input

output\_size the targeted output size

### **Description**

Computes a partial inverse of MaxPool2d.

## Usage

```
nnf_max_unpool2d(
  input,
  indices,
  kernel_size,
  stride = NULL,
  padding = 0,
  output_size = NULL)
```

### **Arguments**

input the input Tensor to invert

indices the indices given out by max pool kernel\_size Size of the max pooling window.

stride Stride of the max pooling window. It is set to kernel\_size by default.

padding Padding that was added to the input

output\_size the targeted output size

nnf\_max\_unpool3d 149

## Description

Computes a partial inverse of MaxPool3d.

### Usage

```
nnf_max_unpool3d(
   input,
   indices,
   kernel_size,
   stride = NULL,
   padding = 0,
   output_size = NULL)
```

### **Arguments**

input the input Tensor to invert

indices the indices given out by max pool kernel\_size Size of the max pooling window.

stride Stride of the max pooling window. It is set to kernel\_size by default.

padding Padding that was added to the input

output\_size the targeted output size

#### **Description**

Measures the element-wise mean squared error.

### Usage

```
nnf_mse_loss(input, target, reduction = "mean")
```

### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor(N,\*), same shape as the input

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

### **Description**

Creates a criterion that optimizes a multi-class multi-classification hinge loss (margin-based loss) between input x (a 2D mini-batch Tensor) and output y (which is a 2D Tensor of target class indices).

### Usage

```
nnf_multilabel_margin_loss(input, target, reduction = "mean")
```

### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor (N,\*), same shape as the input

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

```
nnf\_multilabel\_soft\_margin\_loss \\ \textit{Multilabel\_soft\_margin\_loss}
```

### **Description**

Creates a criterion that optimizes a multi-label one-versus-all loss based on max-entropy, between input x and target y of size (N, C).

```
nnf_multilabel_soft_margin_loss(
  input,
  target,
  weight = NULL,
  reduction = "mean"
)
```

#### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor (N,\*), same shape as the input weight weight tensor to apply on the loss.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

#### Note

It takes a one hot encoded target vector as input.

#### Description

Allows the model to jointly attend to information from different representation subspaces. See reference: Attention Is All You Need

```
nnf_multi_head_attention_forward(
  query,
  key,
  value,
  embed_dim_to_check,
  num_heads,
  in_proj_weight,
  in_proj_bias,
 bias_k,
 bias_v,
  add_zero_attn,
  dropout_p,
  out_proj_weight,
  out_proj_bias,
  training = TRUE,
  key_padding_mask = NULL,
  need_weights = TRUE,
  attn_mask = NULL,
  avg_weights = TRUE,
  use_separate_proj_weight = FALSE,
  q_proj_weight = NULL,
  k_proj_weight = NULL,
```

```
v_proj_weight = NULL,
static_k = NULL,
static_v = NULL,
batch_first = FALSE
)
```

#### **Arguments**

query (L, N, E) where L is the target sequence length, N is the batch size, E is the em-

bedding dimension. If batch\_first is TRUE, the first two dimensions are trans-

posed.

key (S, N, E), where S is the source sequence length, N is the batch size, E is the

embedding dimension. If batch\_first is TRUE, the first two dimensions are trans-

posed.

value (S, N, E) where S is the source sequence length, N is the batch size, E is the

embedding dimension. If batch\_first is TRUE, the first two dimensions are trans-

posed.

embed\_dim\_to\_check

total dimension of the model.

num\_heads parallel attention heads.
in\_proj\_weight input projection weight.
in\_proj\_bias input projection bias.

bias\_k bias of the key and value sequences to be added at dim=0.

bias\_v currently undocumented.

add\_zero\_attn add a new batch of zeros to the key and value sequences at dim=1.

dropout\_p probability of an element to be zeroed.

out\_proj\_weight

the output projection weight.

out\_proj\_bias output projection bias. training apply dropout if is TRUE.

key\_padding\_mask

(N,S) where N is the batch size, S is the source sequence length. If a ByteTensor is provided, the non-zero positions will be ignored while the position with the zero positions will be unchanged. If a BoolTensor is provided, the positions with the value of True will be ignored while the position with the value of False

will be unchanged.

need\_weights output attn\_output\_weights.

attn\_mask 2D mask (L, S) where L

2D mask (L,S) where L is the target sequence length, S is the source sequence length. 3D mask  $(N*num_heads,L,S)$  where N is the batch size, L is the target sequence length, S is the source sequence length. attn\_mask ensure that position i is allowed to attend the unmasked positions. If a ByteTensor is provided, the non-zero positions are not allowed to attend while the zero positions will be unchanged. If a BoolTensor is provided, positions with True is not allowed to attend while False values will be unchanged. If a FloatTensor is provided, it will be added to the attention weight.

avg\_weights Logical; whether to average attn\_output\_weights over the attention heads before

outputting them. This doesn't change the returned value of attn\_output; it only

affects the returned attention weight matrix.

use\_separate\_proj\_weight

the function accept the proj. weights for query, key, and value in different forms. If false, in\_proj\_weight will be used, which is a combination of q\_proj\_weight,

k\_proj\_weight, v\_proj\_weight.

q\_proj\_weight input projection weight and bias.

k\_proj\_weight currently undocumented.v\_proj\_weight currently undocumented.

static\_k static key and value used for attention operators.

static\_v currently undocumented.

batch\_first Logical; whether to expect query, key, and value to have batch as their first

parameter, and to return output with batch first.

## Description

Creates a criterion that optimizes a multi-class classification hinge loss (margin-based loss) between input x (a 2D mini-batch Tensor) and output y (which is a 1D tensor of target class indices,  $\emptyset \le y \le x$ .

#### Usage

```
nnf_multi_margin_loss(
  input,
  target,
  p = 1,
  margin = 1,
  weight = NULL,
  reduction = "mean"
)
```

#### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor (N,\*), same shape as the input

p Has a default value of 1. 1 and 2 are the only supported values.

margin Has a default value of 1.

weight a manual rescaling weight given to each class. If given, it has to be a Tensor of

size C. Otherwise, it is treated as if having all ones.

nnf\_nll\_loss

reduction

(string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed. Default: 'mean'

nnf\_nll\_loss

 $Nll\_loss$ 

## Description

The negative log likelihood loss.

## Usage

```
nnf_nll_loss(
  input,
  target,
  weight = NULL,
  ignore_index = -100,
  reduction = "mean"
)
```

## Arguments

input	$(N,C)$ where C = number of classes or $(N,C,H,W)$ in case of 2D Loss, or $(N,C,d_1,d_2,,d_K)$ where $K\geq 1$ in the case of K-dimensional loss.
target	$(N)$ where each value is $0 \leq \text{targets}[i] \leq C-1,$ or $(N,d_1,d_2,,d_K)$ where $K \geq 1$ for K-dimensional loss.
weight	(Tensor, optional) a manual rescaling weight given to each class. If given, has to be a Tensor of size ${\sf C}$
ignore_index	(int, optional) Specifies a target value that is ignored and does not contribute to the input gradient.
reduction	(string, optional) – Specifies the reduction to apply to the output: 'none' I 'mean' I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed. Default: 'mean'

nnf\_normalize 155

## Description

Performs  $L_p$  normalization of inputs over specified dimension.

### Usage

```
nnf_normalize(input, p = 2, dim = 2, eps = 1e-12, out = NULL)
```

## Arguments

input	input tensor of any shape
р	(float) the exponent value in the norm formulation. Default: 2
dim	(int) the dimension to reduce. Default: 1
eps	(float) small value to avoid division by zero. Default: 1e-12
out	(Tensor, optional) the output tensor. If out is used, this operation won't be differentiable.

#### **Details**

For a tensor input of sizes  $(n_0, ..., n_{dim}, ..., n_k)$ , each  $n_{dim}$  -element vector v along dimension dim is transformed as

$$v = \frac{v}{\max(\|v\|_p, \epsilon)}.$$

With the default arguments it uses the Euclidean norm over vectors along dimension 1 for normalization.

## Description

Takes LongTensor with index values of shape (\*) and returns a tensor of shape (\*, num\_classes) that have zeros everywhere except where the index of last dimension matches the corresponding value of the input tensor, in which case it will be 1.

```
nnf_one_hot(tensor, num_classes = -1)
```

nnf\_pad

#### Arguments

tensor (LongTensor) class values of any shape.

num\_classes (int) Total number of classes. If set to -1, the number of classes will be inferred

as one greater than the largest class value in the input tensor.

#### **Details**

One-hot on Wikipedia: https://en.wikipedia.org/wiki/One-hot

nnf\_pad Pad

### Description

Pads tensor.

#### Usage

```
nnf_pad(input, pad, mode = "constant", value = NULL)
```

#### **Arguments**

input (Tensor) N-dimensional tensor

pad (tuple) m-elements tuple, where  $\frac{m}{2} \le$  input dimensions and m is even.

mode 'constant', 'reflect', 'replicate' or 'circular'. Default: 'constant'

value fill value for 'constant' padding. Default: 0.

#### **Padding size**

The padding size by which to pad some dimensions of input are described starting from the last dimension and moving forward.  $\left\lfloor \frac{\operatorname{len}(\operatorname{pad})}{2} \right\rfloor$  dimensions of input will be padded. For example, to pad only the last dimension of the input tensor, then pad has the form (padding\_left, padding\_right); to pad the last 2 dimensions of the input tensor, then use (padding\_left, padding\_right, padding\_top, padding\_bottom); to pad the last 3 dimensions, use (padding\_left, padding\_right, padding\_bottom padding\_front, padding\_back).

## Padding mode

See nn\_constant\_pad\_2d, nn\_reflection\_pad\_2d, and nn\_replication\_pad\_2d for concrete examples on how each of the padding modes works. Constant padding is implemented for arbitrary dimensions. tensor, or the last 2 dimensions of 4D input tensor, or the last dimension of 3D input tensor. Reflect padding is only implemented for padding the last 2 dimensions of 4D input tensor, or the last dimension of 3D input tensor.

nnf\_pairwise\_distance 157

```
nnf_pairwise_distance
```

## Description

Computes the batchwise pairwise distance between vectors using the p-norm.

### Usage

```
nnf_pairwise_distance(x1, x2, p = 2, eps = 1e-06, keepdim = FALSE)
```

### **Arguments**

x1	(Tensor) First input.
x2	(Tensor) Second input (of size matching x1).
р	the norm degree. Default: 2
eps	(float, optional) Small value to avoid division by zero. Default: 1e-8
keepdim	Determines whether or not to keep the vector dimension. Default: False

### **Description**

Computes the p-norm distance between every pair of row vectors in the input. This is identical to the upper triangular portion, excluding the diagonal, of torch\_norm(input[:, None] - input, dim=2, p=p). This function will be faster if the rows are contiguous.

### Usage

```
nnf_pdist(input, p = 2)
```

## **Arguments**

```
input input tensor of shape N \times M.

p value for the p-norm distance to calculate between each vector pair \in [0, \infty].
```

#### **Details**

If input has shape  $N \times M$  then the output will have shape  $\frac{1}{2}N(N-1)$ .

nnf\_poisson\_nll\_loss

#### Description

Rearranges elements in a tensor of shape  $(*, C \times r^2, H, W)$  to a tensor of shape  $(*, C, H \times r, W \times r)$ .

### Usage

```
nnf_pixel_shuffle(input, upscale_factor)
```

## Arguments

```
input (Tensor) the input tensor upscale_factor (int) factor to increase spatial resolution by
```

## Description

Poisson negative log likelihood loss.

#### Usage

```
nnf_poisson_nll_loss(
  input,
  target,
  log_input = TRUE,
  full = FALSE,
  eps = 1e-08,
  reduction = "mean"
)
```

#### Arguments

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor(N,\*), same shape as the input

log\_input if TRUE the loss is computed as exp(input) — target \* input, if FALSE then loss is

input - target  $* \log(input + eps)$ . Default: TRUE.

full whether to compute full loss, i. e. to add the Stirling approximation term. De-

fault: FALSE.

eps (float, optional) Small value to avoid evaluation of log(0) when  $log_input=FALSE$ .

Default: 1e-8

nnf\_prelu 159

reduction

(string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed. Default: 'mean'

nnf\_prelu

Prelu

## Description

Applies element-wise the function PReLU(x) = max(0, x) + weight \* min(0, x) where weight is a learnable parameter.

# Usage

```
nnf_prelu(input, weight)
```

# Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions weight (Tensor) the learnable weights

nnf\_relu

Relu

## Description

Applies the rectified linear unit function element-wise.

## Usage

```
nnf_relu(input, inplace = FALSE)
nnf_relu_(input)
```

### **Arguments**

input (N,\*) tensor, where \* means, any number of additional dimensions

inplace can optionally do the operation in-place. Default: FALSE

nnf\_rrelu

nnf_relu6	Relu6
-----------	-------

# Description

```
Applies the element-wise function ReLU6(x) = min(max(0, x), 6).
```

## Usage

```
nnf_relu6(input, inplace = FALSE)
```

# Arguments

input	(N,*) tensor, where * means, any number of additional dimensions
inplace	can optionally do the operation in-place. Default: FALSE

## Description

Randomized leaky ReLU.

## Usage

```
nnf_rrelu(input, lower = 1/8, upper = 1/3, training = FALSE, inplace = FALSE)
nnf_rrelu_(input, lower = 1/8, upper = 1/3, training = FALSE)
```

## Arguments

input	(N,*) tensor, where * means, any number of additional dimensions
lower	lower bound of the uniform distribution. Default: 1/8
upper	upper bound of the uniform distribution. Default: 1/3
training	bool wether it's a training pass. DEfault: FALSE
inplace	can optionally do the operation in-place. Default: FALSE

nnf\_selu 161

nnf\_selu

Selu

### **Description**

Applies element-wise,

$$SELU(x) = scale * (max(0, x) + min(0, \alpha * (exp(x) - 1)))$$

, with  $\alpha=1.6732632423543772848170429916717$  and scale=1.0507009873554804934193349852946.

### Usage

```
nnf_selu(input, inplace = FALSE)
nnf_selu_(input)
```

## Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions inplace can optionally do the operation in-place. Default: FALSE

## **Examples**

```
if (torch_is_installed()) {
x <- torch_randn(2, 2)
y <- nnf_selu(x)
nnf_selu_(x)
torch_equal(x, y)
}</pre>
```

nnf\_sigmoid

Sigmoid

### **Description**

```
Applies element-wise Sigmoid(x_i) = \frac{1}{1 + exp(-x_i)}
```

#### Usage

```
nnf_sigmoid(input)
```

## Arguments

input

(N,\*) tensor, where \* means, any number of additional dimensions

nnf\_smooth\_11\_loss

### **Description**

Applies the Sigmoid Linear Unit (SiLU) function, element-wise. See nn\_silu() for more information.

#### Usage

```
nnf_silu(input, inplace = FALSE)
```

# Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions

inplace can optionally do the operation in-place. Default: FALSE

#### See Also

nn\_silu().

### **Description**

Function that uses a squared term if the absolute element-wise error falls below 1 and an L1 term otherwise.

#### Usage

```
nnf_smooth_l1_loss(input, target, reduction = "mean")
```

### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor(N,\*), same shape as the input

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

nnf\_softmax 163

nnf\_softmax Softmax

## Description

Applies a softmax function.

## Usage

```
nnf_softmax(input, dim, dtype = NULL)
```

#### Arguments

input (Tensor) input

dim (int) A dimension along which softmax will be computed.

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

#### **Details**

Softmax is defined as:

$$Softmax(x_i) = exp(x_i) / \sum_j exp(x_j)$$

It is applied to all slices along dim, and will re-scale them so that the elements lie in the range [0, 1] and sum to 1.

nnf\_softmin Softmin

## Description

Applies a softmin function.

```
nnf_softmin(input, dim, dtype = NULL)
```

nnf\_softplus

#### **Arguments**

input (Tensor) input

dim (int) A dimension along which softmin will be computed (so every slice along

dim will sum to 1).

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

### **Details**

Note that

$$Softmin(x) = Softmax(-x)$$

•

See nnf\_softmax definition for mathematical formula.

nnf\_softplus Softplus

## Description

Applies element-wise, the function  $Softplus(x) = 1/\beta * log(1 + exp(\beta * x)).$ 

### Usage

```
nnf_softplus(input, beta = 1, threshold = 20)
```

# Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions

beta the beta value for the Softplus formulation. Default: 1

threshold values above this revert to a linear function. Default: 20

## **Details**

For numerical stability the implementation reverts to the linear function when  $input*\beta > threshold$ .

nnf\_softshrink 165

nnf\_softshrink

Softshrink

### **Description**

Applies the soft shrinkage function elementwise

## Usage

```
nnf_softshrink(input, lambd = 0.5)
```

## Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions

lambd the lambda (must be no less than zero) value for the Softshrink formulation.

Default: 0.5

nnf\_softsign

Softsign

## Description

Applies element-wise, the function SoftSign(x) = x/(1 + |x|)

## Usage

```
nnf_softsign(input)
```

## Arguments

input

(N,\*) tensor, where \* means, any number of additional dimensions

nnf\_tanhshrink

### **Description**

Creates a criterion that optimizes a two-class classification logistic loss between input tensor x and target tensor y (containing 1 or -1).

## Usage

```
nnf_soft_margin_loss(input, target, reduction = "mean")
```

#### **Arguments**

input tensor (N,\*) where \*\* means, any number of additional dimensions

target tensor (N,\*), same shape as the input

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

### **Description**

```
Applies element-wise, Tanhshrink(x) = x - Tanh(x)
```

### Usage

```
nnf_tanhshrink(input)
```

### **Arguments**

input (N,\*) tensor, where \* means, any number of additional dimensions

nnf\_threshold 167

nnf\_threshold

Threshold

### **Description**

Thresholds each element of the input Tensor.

## Usage

```
nnf_threshold(input, threshold, value, inplace = FALSE)
nnf_threshold_(input, threshold, value)
```

## Arguments

input (N,\*) tensor, where \* means, any number of additional dimensions

threshold The value to threshold at value The value to replace with

inplace can optionally do the operation in-place. Default: FALSE

```
nnf_triplet_margin_loss
```

Triplet\_margin\_loss

## Description

Creates a criterion that measures the triplet loss given an input tensors x1, x2, x3 and a margin with a value greater than 0. This is used for measuring a relative similarity between samples. A triplet is composed by a, p and n (i.e., anchor, positive examples and negative examples respectively). The shapes of all input tensors should be (N, D).

```
nnf_triplet_margin_loss(
  anchor,
  positive,
  negative,
  margin = 1,
  p = 2,
  eps = 1e-06,
  swap = FALSE,
  reduction = "mean"
)
```

## Arguments

anchor the anchor input tensor
positive the positive input tensor
negative the negative input tensor

margin Default: 1.

p The norm degree for pairwise distance. Default: 2. eps (float, optional) Small value to avoid division by zero.

swap The distance swap is described in detail in the paper Learning shallow convolu-

tional feature descriptors with triplet losses by V. Balntas, E. Riba et al. Default:

FALSE.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

```
nnf_triplet_margin_with_distance_loss

*Triplet margin with distance loss*
```

## Description

```
See nn_triplet_margin_with_distance_loss()
```

#### **Usage**

```
nnf_triplet_margin_with_distance_loss(
   anchor,
   positive,
   negative,
   distance_function = NULL,
   margin = 1,
   swap = FALSE,
   reduction = "mean"
)
```

## Arguments

anchor the anchor input tensor positive the positive input tensor negative the negative input tensor

distance\_function

(callable, optional): A nonnegative, real-valued function that quantifies the closeness of two tensors. If not specified, nn\_pairwise\_distance() will be used.

Default: None

nnf\_unfold 169

margin Default: 1.

swap The distance swap is described in detail in the paper Learning shallow convolu-

tional feature descriptors with triplet losses by V. Balntas, E. Riba et al. Default:

FALSE.

reduction (string, optional) – Specifies the reduction to apply to the output: 'none' I'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be

summed. Default: 'mean'

nnf\_unfold Unfold

## **Description**

Extracts sliding local blocks from an batched input tensor.

## Usage

```
nnf_unfold(input, kernel_size, dilation = 1, padding = 0, stride = 1)
```

### **Arguments**

input the input tensor

kernel\_size the size of the sliding blocks

dilation a parameter that controls the stride of elements within the neighborhood. De-

fault: 1

padding implicit zero padding to be added on both sides of input. Default: 0

stride the stride of the sliding blocks in the input spatial dimensions. Default: 1

#### Warning

More than one element of the unfolded tensor may refer to a single memory location. As a result, in-place operations (especially ones that are vectorized) may result in incorrect behavior. If you need to write to the tensor, please clone it first.

```
nn_adaptive_avg_pool1d
```

Applies a 1D adaptive average pooling over an input signal composed of several input planes.

## **Description**

The output size is H, for any input size. The number of output features is equal to the number of input planes.

#### Usage

```
nn_adaptive_avg_pool1d(output_size)
```

#### **Arguments**

```
output_size the target output size H
```

#### **Examples**

```
if (torch_is_installed()) {
# target output size of 5
m <- nn_adaptive_avg_pool1d(5)
input <- torch_randn(1, 64, 8)
output <- m(input)
}</pre>
```

nn\_adaptive\_avg\_pool2d

Applies a 2D adaptive average pooling over an input signal composed of several input planes.

### **Description**

The output is of size H x W, for any input size. The number of output features is equal to the number of input planes.

### Usage

```
nn_adaptive_avg_pool2d(output_size)
```

### **Arguments**

output\_size

the target output size of the image of the form H x W. Can be a tuple (H, W) or a single H for a square image H x H. H and W can be either a int, or NULL which means the size will be the same as that of the input.

#### **Examples**

```
if (torch_is_installed()) {
# target output size of 5x7
m <- nn_adaptive_avg_pool2d(c(5, 7))
input <- torch_randn(1, 64, 8, 9)
output <- m(input)
# target output size of 7x7 (square)
m <- nn_adaptive_avg_pool2d(7)
input <- torch_randn(1, 64, 10, 9)
output <- m(input)
}</pre>
```

nn\_adaptive\_avg\_pool3d

Applies a 3D adaptive average pooling over an input signal composed of several input planes.

### **Description**

The output is of size D x H x W, for any input size. The number of output features is equal to the number of input planes.

### Usage

```
nn_adaptive_avg_pool3d(output_size)
```

## Arguments

output\_size

the target output size of the form D x H x W. Can be a tuple (D, H, W) or a single number D for a cube D x D x D. D, H and W can be either a int, or None which means the size will be the same as that of the input.

#### **Examples**

```
if (torch_is_installed()) {
# target output size of 5x7x9
m <- nn_adaptive_avg_pool3d(c(5, 7, 9))
input <- torch_randn(1, 64, 8, 9, 10)
output <- m(input)
# target output size of 7x7x7 (cube)
m <- nn_adaptive_avg_pool3d(7)
input <- torch_randn(1, 64, 10, 9, 8)
output <- m(input)
}</pre>
```

```
nn\_adaptive\_log\_softmax\_with\_loss\\ AdaptiveLogSoftmaxWithLoss\ module
```

#### **Description**

Efficient softmax approximation as described in Efficient softmax approximation for GPUs by Edouard Grave, Armand Joulin, Moustapha Cissé, David Grangier, and Hervé Jégou

#### Usage

```
nn_adaptive_log_softmax_with_loss(
  in_features,
  n_classes,
  cutoffs,
  div_value = 4,
  head_bias = FALSE
)
```

#### **Arguments**

in\_features (int): Number of features in the input tensor
 n\_classes (int): Number of classes in the dataset
 cutoffs (Sequence): Cutoffs used to assign targets to their buckets
 div\_value (float, optional): value used as an exponent to compute sizes of the clusters. Default: 4.0
 head\_bias (bool, optional): If True, adds a bias term to the 'head' of the adaptive softmax. Default: False

#### **Details**

Adaptive softmax is an approximate strategy for training models with large output spaces. It is most effective when the label distribution is highly imbalanced, for example in natural language modelling, where the word frequency distribution approximately follows the Zipf's law.

Adaptive softmax partitions the labels into several clusters, according to their frequency. These clusters may contain different number of targets each.

Additionally, clusters containing less frequent labels assign lower dimensional embeddings to those labels, which speeds up the computation. For each minibatch, only clusters for which at least one target is present are evaluated.

The idea is that the clusters which are accessed frequently (like the first one, containing most frequent labels), should also be cheap to compute – that is, contain a small number of assigned labels. We highly recommend taking a look at the original paper for more details.

- cutoffs should be an ordered Sequence of integers sorted in the increasing order. It controls number of clusters and the partitioning of targets into clusters. For example setting cutoffs = c(10, 100, 1000) means that first 10 targets will be assigned to the 'head' of the adaptive softmax, targets 11, 12, ..., 100 will be assigned to the first cluster, and targets 101, 102, ..., 1000 will be assigned to the second cluster, while targets 1001, 1002, ..., n\_classes 1 will be assigned to the last, third cluster.
- div\_value is used to compute the size of each additional cluster, which is given as  $\left\lfloor \frac{\text{in\_features}}{\text{div\_value}^{idx}} \right\rfloor$ , where idx is the cluster index (with clusters for less frequent words having larger indices, and indices starting from 1).
- head\_bias if set to True, adds a bias term to the 'head' of the adaptive softmax. See paper for details. Set to False in the official implementation.

#### Value

NamedTuple with output and loss fields:

- output is a Tensor of size N containing computed target log probabilities for each example
- loss is a Scalar representing the computed negative log likelihood loss

#### Warning

Labels passed as inputs to this module should be sorted according to their frequency. This means that the most frequent label should be represented by the index 0, and the least frequent label should be represented by the index n\_classes - 1.

#### Shape

- input:  $(N, in\_features)$
- target: (N) where each value satisfies  $0 \le target[i] \le n_classes$
- output1: (*N*)
- output2: Scalar

## Note

This module returns a NamedTuple with output and loss fields. See further documentation for details.

To compute log-probabilities for all classes, the log\_prob method can be used.

```
nn_adaptive_max_pool1d
```

Applies a 1D adaptive max pooling over an input signal composed of several input planes.

## Description

The output size is H, for any input size. The number of output features is equal to the number of input planes.

#### Usage

```
nn_adaptive_max_pool1d(output_size, return_indices = FALSE)
```

## Arguments

```
output_size the target output size H
return_indices if TRUE, will return the indices along with the outputs. Useful to pass to nn_max_unpool1d().

Default: FALSE
```

#### **Examples**

```
if (torch_is_installed()) {
# target output size of 5
m <- nn_adaptive_max_pool1d(5)
input <- torch_randn(1, 64, 8)
output <- m(input)
}</pre>
```

nn\_adaptive\_max\_pool2d

Applies a 2D adaptive max pooling over an input signal composed of several input planes.

# Description

The output is of size H x W, for any input size. The number of output features is equal to the number of input planes.

```
nn_adaptive_max_pool2d(output_size, return_indices = FALSE)
```

## **Arguments**

output\_size the target output size of the image of the form H x W. Can be a tuple (H, W)

or a single H for a square image H x H. H and W can be either a int, or None

which means the size will be the same as that of the input.

return\_indices if TRUE, will return the indices along with the outputs. Useful to pass to nn\_max\_unpool2d().

Default: FALSE

## **Examples**

```
if (torch_is_installed()) {
# target output size of 5x7
m <- nn_adaptive_max_pool2d(c(5, 7))
input <- torch_randn(1, 64, 8, 9)
output <- m(input)
# target output size of 7x7 (square)
m <- nn_adaptive_max_pool2d(7)
input <- torch_randn(1, 64, 10, 9)
output <- m(input)
}</pre>
```

nn\_adaptive\_max\_pool3d

Applies a 3D adaptive max pooling over an input signal composed of several input planes.

### **Description**

The output is of size D x H x W, for any input size. The number of output features is equal to the number of input planes.

## Usage

```
nn_adaptive_max_pool3d(output_size, return_indices = FALSE)
```

## Arguments

output\_size the target output size of the image of the form D x H x W. Can be a tuple (D, H,

W) or a single D for a cube D x D x D. D, H and W can be either a int, or None

which means the size will be the same as that of the input.

return\_indices if TRUE, will return the indices along with the outputs. Useful to pass to nn\_max\_unpool3d().

Default: FALSE

nn\_avg\_pool1d

### **Examples**

```
if (torch_is_installed()) {
# target output size of 5x7x9
m <- nn_adaptive_max_pool3d(c(5, 7, 9))
input <- torch_randn(1, 64, 8, 9, 10)
output <- m(input)
# target output size of 7x7x7 (cube)
m <- nn_adaptive_max_pool3d(7)
input <- torch_randn(1, 64, 10, 9, 8)
output <- m(input)
}</pre>
```

nn\_avg\_pool1d

Applies a 1D average pooling over an input signal composed of several input planes.

## Description

In the simplest case, the output value of the layer with input size (N, C, L), output  $(N, C, L_{out})$  and kernel\_size k can be precisely described as:

## Usage

```
nn_avg_pool1d(
   kernel_size,
   stride = NULL,
   padding = 0,
   ceil_mode = FALSE,
   count_include_pad = TRUE
)
```

#### **Arguments**

kernel\_size the size of the window

stride the stride of the window. Default value is kernel\_size

padding implicit zero padding to be added on both sides

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape

count\_include\_pad

when TRUE, will include the zero-padding in the averaging calculation

#### **Details**

$$\operatorname{out}(N_i,C_j,l) = \frac{1}{k} \sum_{m=0}^{k-1} \operatorname{input}(N_i,C_j,\operatorname{stride} \times l + m)$$

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If padding is non-zero, then the input is implicitly zero-padded on both sides for padding number of points.

The parameters kernel\_size, stride, padding can each be an int or a one-element tuple.

### Shape

- Input:  $(N, C, L_{in})$
- Output:  $(N, C, L_{out})$ , where

$$L_{out} = \left\lfloor \frac{L_{in} + 2 \times \text{padding} - \text{kernel\_size}}{\text{stride}} + 1 \right\rfloor$$

# Examples

```
if (torch_is_installed()) {
# pool with window of size=3, stride=2
m <- nn_avg_pool1d(3, stride = 2)
m(torch_randn(1, 1, 8))
}</pre>
```

nn\_avg\_pool2d

Applies a 2D average pooling over an input signal composed of several input planes.

#### Description

In the simplest case, the output value of the layer with input size (N, C, H, W), output  $(N, C, H_{out}, W_{out})$  and kernel\_size (kH, kW) can be precisely described as:

#### Usage

```
nn_avg_pool2d(
  kernel_size,
  stride = NULL,
  padding = 0,
  ceil_mode = FALSE,
  count_include_pad = TRUE,
  divisor_override = NULL
)
```

#### **Arguments**

```
kernel_size the size of the window
stride the stride of the window. Default value is kernel_size
padding implicit zero padding to be added on both sides
```

nn\_avg\_pool2d

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape count\_include\_pad

when TRUE, will include the zero-padding in the averaging calculation divisor\_override

if specified, it will be used as divisor, otherwise kernel\_size will be used

#### **Details**

$$out(N_i, C_j, h, w) = \frac{1}{kH * kW} \sum_{m=0}^{kH-1} \sum_{n=0}^{kW-1} input(N_i, C_j, stride[0] \times h + m, stride[1] \times w + n)$$

If padding is non-zero, then the input is implicitly zero-padded on both sides for padding number of points.

The parameters kernel\_size, stride, padding can either be:

- a single int in which case the same value is used for the height and width dimension
- a tuple of two ints in which case, the first int is used for the height dimension, and the second int for the width dimension

### Shape

- Input:  $(N, C, H_{in}, W_{in})$
- Output:  $(N, C, H_{out}, W_{out})$ , where

$$H_{out} = \left\lfloor \frac{H_{in} + 2 \times \text{padding}[0] - \text{kernel\_size}[0]}{\text{stride}[0]} + 1 \right\rfloor$$

$$W_{out} = \left\lfloor \frac{W_{in} + 2 \times \mathsf{padding}[1] - \mathsf{kernel\_size}[1]}{\mathsf{stride}[1]} + 1 \right\rfloor$$

### **Examples**

```
if (torch_is_installed()) {
# pool of square window of size=3, stride=2
m <- nn_avg_pool2d(3, stride = 2)
# pool of non-square window
m <- nn_avg_pool2d(c(3, 2), stride = c(2, 1))
input <- torch_randn(20, 16, 50, 32)
output <- m(input)
}</pre>
```

nn\_avg\_pool3d

nn_avg_pool3d	Applies a 3D average pooling over an input signal composed of several input planes.
---------------	---

#### **Description**

In the simplest case, the output value of the layer with input size (N, C, D, H, W), output  $(N, C, D_{out}, H_{out}, W_{out})$  and kernel\_size (kD, kH, kW) can be precisely described as:

## Usage

```
nn_avg_pool3d(
  kernel_size,
  stride = NULL,
  padding = 0,
  ceil_mode = FALSE,
  count_include_pad = TRUE,
  divisor_override = NULL
)
```

#### **Arguments**

```
kernel_size the size of the window

stride the stride of the window. Default value is kernel_size

padding implicit zero padding to be added on all three sides

ceil_mode when TRUE, will use ceil instead of floor to compute the output shape

count_include_pad

when TRUE, will include the zero-padding in the averaging calculation

divisor_override

if specified, it will be used as divisor, otherwise kernel_size will be used
```

## **Details**

```
\operatorname{out}(N_i, C_j, d, h, w) = \underbrace{\sum_{k=0}^{kD-1} \sum_{m=0}^{kH-1} \sum_{n=0}^{kW-1}}_{\substack{input(N_i, C_j, \text{stride}[0] \times d+k, \text{stride}[1] \times h+m, \text{stride}[2] \times w+n)}}_{\substack{kD \times kW \times kW}}
```

If padding is non-zero, then the input is implicitly zero-padded on all three sides for padding number of points.

The parameters kernel\_size, stride can either be:

- a single int in which case the same value is used for the depth, height and width dimension
- a tuple of three ints in which case, the first int is used for the depth dimension, the second int for the height dimension and the third int for the width dimension

nn\_batch\_norm1d

## Shape

- Input:  $(N, C, D_{in}, H_{in}, W_{in})$
- Output:  $(N, C, D_{out}, H_{out}, W_{out})$ , where

$$\begin{split} D_{out} &= \left\lfloor \frac{D_{in} + 2 \times \text{padding}[0] - \text{kernel\_size}[0]}{\text{stride}[0]} + 1 \right\rfloor \\ H_{out} &= \left\lfloor \frac{H_{in} + 2 \times \text{padding}[1] - \text{kernel\_size}[1]}{\text{stride}[1]} + 1 \right\rfloor \\ W_{out} &= \left\lfloor \frac{W_{in} + 2 \times \text{padding}[2] - \text{kernel\_size}[2]}{\text{stride}[2]} + 1 \right\rfloor \end{split}$$

#### **Examples**

```
if (torch_is_installed()) {

# pool of square window of size=3, stride=2

m <- nn_avg_pool3d(3, stride = 2)

# pool of non-square window

m <- nn_avg_pool3d(c(3, 2, 2), stride = c(2, 1, 2))
input <- torch_randn(20, 16, 50, 44, 31)
output <- m(input)
}</pre>
```

nn\_batch\_norm1d

BatchNorm1D module

#### Description

Applies Batch Normalization over a 2D or 3D input (a mini-batch of 1D inputs with optional additional channel dimension) as described in the paper Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift

```
nn_batch_norm1d(
  num_features,
  eps = 1e-05,
  momentum = 0.1,
  affine = TRUE,
  track_running_stats = TRUE
)
```

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# Arguments

num\_features C from an expected input of size (N, C, L) or L from input of size (N, L) eps a value added to the denominator for numerical stability. Default: 1e-5

momentum the value used for the running\_mean and running\_var computation. Can be set

to NULL for cumulative moving average (i.e. simple average). Default: 0.1

affine a boolean value that when set to TRUE, this module has learnable affine parame-

ters. Default: TRUE

track\_running\_stats

a boolean value that when set to TRUE, this module tracks the running mean and variance, and when set to FALSE, this module does not track such statistics and always uses batch statistics in both training and eval modes. Default: TRUE

#### **Details**

$$y = \frac{x - \mathbf{E}[x]}{\sqrt{\mathbf{Var}[x] + \epsilon}} * \gamma + \beta$$

The mean and standard-deviation are calculated per-dimension over the mini-batches and  $\gamma$  and  $\beta$  are learnable parameter vectors of size C (where C is the input size). By default, the elements of  $\gamma$  are set to 1 and the elements of  $\beta$  are set to 0.

Also by default, during training this layer keeps running estimates of its computed mean and variance, which are then used for normalization during evaluation. The running estimates are kept with a default :attr:momentum of 0.1. If track\_running\_stats is set to FALSE, this layer then does not keep running estimates, and batch statistics are instead used during evaluation time as well.

## Note

This momentum argument is different from one used in optimizer classes and the conventional notion of momentum. Mathematically, the update rule for running statistics here is  $\hat{x}_{\text{new}} = (1 - \text{momentum}) \times \hat{x} + \text{momentum} \times x_t$ , where  $\hat{x}$  is the estimated statistic and  $x_t$  is the new observed value.

Because the Batch Normalization is done over the C dimension, computing statistics on (N, L) slices, it's common terminology to call this Temporal Batch Normalization.

# Shape

- Input: (N, C) or (N, C, L)
- Output: (N, C) or (N, C, L) (same shape as input)

# **Examples**

```
if (torch_is_installed()) {
# With Learnable Parameters
m <- nn_batch_norm1d(100)
# Without Learnable Parameters
m <- nn_batch_norm1d(100, affine = FALSE)
input <- torch_randn(20, 100)</pre>
```

nn\_batch\_norm2d

```
output <- m(input)
}</pre>
```

nn\_batch\_norm2d

BatchNorm2D

### **Description**

Applies Batch Normalization over a 4D input (a mini-batch of 2D inputs additional channel dimension) as described in the paper Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift.

# Usage

```
nn_batch_norm2d(
  num_features,
  eps = 1e-05,
  momentum = 0.1,
  affine = TRUE,
  track_running_stats = TRUE
)
```

### **Arguments**

num\_features C from an expected input of size (N, C, H, W)

eps a value added to the denominator for numerical stability. Default: 1e-5

momentum the value used for the running\_mean and running\_var computation. Can be set

to None for cumulative moving average (i.e. simple average). Default: 0.1

affine a boolean value that when set to TRUE, this module has learnable affine parame-

ters. Default: TRUE

track\_running\_stats

a boolean value that when set to TRUE, this module tracks the running mean and variance, and when set to FALSE, this module does not track such statistics and uses batch statistics instead in both training and eval modes if the running mean and variance are None. Default: TRUE

#### **Details**

$$y = \frac{x - \mathbf{E}[x]}{\sqrt{\mathbf{Var}[x] + \epsilon}} * \gamma + \beta$$

The mean and standard-deviation are calculated per-dimension over the mini-batches and  $\gamma$  and  $\beta$  are learnable parameter vectors of size C (where C is the input size). By default, the elements of  $\gamma$  are set to 1 and the elements of  $\beta$  are set to 0. The standard-deviation is calculated via the biased estimator, equivalent to torch\_var(input, unbiased=FALSE). Also by default, during training

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this layer keeps running estimates of its computed mean and variance, which are then used for normalization during evaluation. The running estimates are kept with a default momentum of 0.1.

If track\_running\_stats is set to FALSE, this layer then does not keep running estimates, and batch statistics are instead used during evaluation time as well.

# Shape

```
Input: (N, C, H, W)
Output: (N, C, H, W) (same shape as input)
```

#### Note

This momentum argument is different from one used in optimizer classes and the conventional notion of momentum. Mathematically, the update rule for running statistics here is  $\hat{x}_{\text{new}} = (1 - \text{momentum}) \times \hat{x} + \text{momentum} \times x_t$ , where  $\hat{x}$  is the estimated statistic and  $x_t$  is the new observed value. Because the Batch Normalization is done over the C dimension, computing statistics on (N, H, W) slices, it's common terminology to call this Spatial Batch Normalization.

### **Examples**

```
if (torch_is_installed()) {
# With Learnable Parameters
m <- nn_batch_norm2d(100)
# Without Learnable Parameters
m <- nn_batch_norm2d(100, affine = FALSE)
input <- torch_randn(20, 100, 35, 45)
output <- m(input)
}</pre>
```

nn\_batch\_norm3d

BatchNorm3D

# **Description**

Applies Batch Normalization over a 5D input (a mini-batch of 3D inputs with additional channel dimension) as described in the paper Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift.

# Usage

```
nn_batch_norm3d(
  num_features,
  eps = 1e-05,
  momentum = 0.1,
  affine = TRUE,
  track_running_stats = TRUE
)
```

nn\_batch\_norm3d

#### **Arguments**

num\_features C from an expected input of size (N, C, D, H, W)

eps a value added to the denominator for numerical stability. Default: 1e-5

momentum the value used for the running\_mean and running\_var computation. Can be set

to None for cumulative moving average (i.e. simple average). Default: 0.1

affine a boolean value that when set to TRUE, this module has learnable affine parame-

ters. Default: TRUE

track\_running\_stats

a boolean value that when set to TRUE, this module tracks the running mean and variance, and when set to FALSE, this module does not track such statistics and uses batch statistics instead in both training and eval modes if the running mean and variance are None. Default: TRUE

#### **Details**

$$y = \frac{x - \mathbf{E}[x]}{\sqrt{\mathbf{Var}[x] + \epsilon}} * \gamma + \beta$$

The mean and standard-deviation are calculated per-dimension over the mini-batches and  $\gamma$  and  $\beta$  are learnable parameter vectors of size C (where C is the input size). By default, the elements of  $\gamma$  are set to 1 and the elements of  $\beta$  are set to 0. The standard-deviation is calculated via the biased estimator, equivalent to torch\_var(input, unbiased = FALSE).

Also by default, during training this layer keeps running estimates of its computed mean and variance, which are then used for normalization during evaluation. The running estimates are kept with a default momentum of 0.1.

If track\_running\_stats is set to FALSE, this layer then does not keep running estimates, and batch statistics are instead used during evaluation time as well.

## Shape

• Input: (N, C, D, H, W)

• Output: (N, C, D, H, W) (same shape as input)

#### Note

This momentum argument is different from one used in optimizer classes and the conventional notion of momentum. Mathematically, the update rule for running statistics here is:  $\hat{x}_{\text{new}} = (1 - \text{momentum}) \times \hat{x} + \text{momentum} \times x_t$ , where  $\hat{x}$  is the estimated statistic and  $x_t$  is the new observed value.

Because the Batch Normalization is done over the C dimension, computing statistics on (N, D, H, W) slices, it's common terminology to call this Volumetric Batch Normalization or Spatio-temporal Batch Normalization.

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#### **Examples**

```
if (torch_is_installed()) {
# With Learnable Parameters
m <- nn_batch_norm3d(100)</pre>
# Without Learnable Parameters
m <- nn_batch_norm3d(100, affine = FALSE)</pre>
input <- torch_randn(20, 100, 35, 45, 55)
output <- m(input)</pre>
```

nn\_bce\_loss

Binary cross entropy loss

# **Description**

Creates a criterion that measures the Binary Cross Entropy between the target and the output:

### Usage

```
nn_bce_loss(weight = NULL, reduction = "mean")
```

# **Arguments**

weight (Tensor, optional): a manual rescaling weight given to the loss of each batch

element. If given, has to be a Tensor of size nbatch.

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

# **Details**

The unreduced (i.e. with reduction set to 'none') loss can be described as:

$$\ell(x,y) = L = \{l_1, \dots, l_N\}^{\top}, \quad l_n = -w_n [y_n \cdot \log x_n + (1 - y_n) \cdot \log(1 - x_n)]$$

where N is the batch size. If reduction is not 'none' (default 'mean'), then

$$\ell(x,y) = \left\{ \begin{array}{ll} \operatorname{mean}(L), & \text{if reduction = 'mean';} \\ \operatorname{sum}(L), & \text{if reduction = 'sum'.} \end{array} \right.$$

This is used for measuring the error of a reconstruction in for example an auto-encoder. Note that the targets y should be numbers between 0 and 1.

Notice that if  $x_n$  is either 0 or 1, one of the log terms would be mathematically undefined in the above loss equation. PyTorch chooses to set  $\log(0) = -\infty$ , since  $\lim_{x\to 0} \log(x) = -\infty$ .

However, an infinite term in the loss equation is not desirable for several reasons. For one, if either  $y_n = 0$  or  $(1 - y_n) = 0$ , then we would be multiplying 0 with infinity. Secondly, if we have an infinite loss value, then we would also have an infinite term in our gradient, since  $\lim_{x\to 0} \frac{d}{dx} \log(x) = \infty$ .

This would make BCELoss's backward method nonlinear with respect to  $x_n$ , and using it for things like linear regression would not be straight-forward. Our solution is that BCELoss clamps its log function outputs to be greater than or equal to -100. This way, we can always have a finite loss value and a linear backward method.

# Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input
- Output: scalar. If reduction is 'none', then (N,\*), same shape as input.

## **Examples**

```
if (torch_is_installed()) {
m <- nn_sigmoid()
loss <- nn_bce_loss()
input <- torch_randn(3, requires_grad = TRUE)
target <- torch_rand(3)
output <- loss(m(input), target)
output$backward()
}</pre>
```

nn\_bce\_with\_logits\_loss

BCE with logits loss

### **Description**

This loss combines a Sigmoid layer and the BCELoss in one single class. This version is more numerically stable than using a plain Sigmoid followed by a BCELoss as, by combining the operations into one layer, we take advantage of the log-sum-exp trick for numerical stability.

#### Usage

```
nn_bce_with_logits_loss(weight = NULL, reduction = "mean", pos_weight = NULL)
```

# **Arguments**

weight	(Tensor, optional): a manual rescaling weight given to the loss of each batch element. If given, has to be a Tensor of size nbatch.
reduction	(string, optional): Specifies the reduction to apply to the output: 'none'   'mean'   'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.
pos_weight	(Tensor, optional): a weight of positive examples. Must be a vector with length equal to the number of classes.

#### **Details**

The unreduced (i.e. with reduction set to 'none') loss can be described as:

$$\ell(x,y) = L = \{l_1, \dots, l_N\}^{\top}, \quad l_n = -w_n [y_n \cdot \log \sigma(x_n) + (1-y_n) \cdot \log(1-\sigma(x_n))],$$

where N is the batch size. If reduction is not 'none' (default 'mean'), then

$$\ell(x,y) = \begin{array}{ll} \mathrm{mean}(L), & \mathrm{if\ reduction = 'mean';} \\ \mathrm{sum}(L), & \mathrm{if\ reduction = 'sum'.} \end{array}$$

This is used for measuring the error of a reconstruction in for example an auto-encoder. Note that the targets t[i] should be numbers between 0 and 1. It's possible to trade off recall and precision by adding weights to positive examples. In the case of multi-label classification the loss can be described as:

$$\ell_c(x,y) = L_c = \{l_{1,c}, \dots, l_{N,c}\}^{\top}, \quad l_{n,c} = -w_{n,c} \left[p_c y_{n,c} \cdot \log \sigma(x_{n,c}) + (1 - y_{n,c}) \cdot \log(1 - \sigma(x_{n,c}))\right],$$

where c is the class number (c > 1 for multi-label binary classification,

c=1 for single-label binary classification), n is the number of the sample in the batch and  $p_c$  is the weight of the positive answer for the class c.  $p_c>1$  increases the recall,  $p_c<1$  increases the precision. For example, if a dataset contains 100 positive and 300 negative examples of a single class, then pos\_weight for the class should be equal to  $\frac{300}{100}=3$ . The loss would act as if the dataset contains  $3\times 100=300$  positive examples.

### Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input
- Output: scalar. If reduction is 'none', then (N, \*), same shape as input.

### **Examples**

```
if (torch_is_installed()) {
  loss <- nn_bce_with_logits_loss()
  input <- torch_randn(3, requires_grad = TRUE)
  target <- torch_empty(3)$random_(1, 2)
  output <- loss(input, target)
  output$backward()

target <- torch_ones(10, 64, dtype = torch_float32()) # 64 classes, batch size = 10
  output <- torch_full(c(10, 64), 1.5) # A prediction (logit)
  pos_weight <- torch_ones(64) # All weights are equal to 1
  criterion <- nn_bce_with_logits_loss(pos_weight = pos_weight)
  criterion(output, target) # -log(sigmoid(1.5))
}</pre>
```

nn\_bilinear

nn\_bilinear

Bilinear module

# **Description**

Applies a bilinear transformation to the incoming data  $y = x_1^T A x_2 + b$ 

## Usage

```
nn_bilinear(in1_features, in2_features, out_features, bias = TRUE)
```

# **Arguments**

```
in1_features size of each first input sample
in2_features size of each second input sample
out_features size of each output sample
bias If set to FALSE, the layer will not learn an additive bias. Default: TRUE
```

# Shape

- Input1:  $(N, *, H_{in1})$   $H_{in1} = \text{in1\_features}$  and \* means any number of additional dimensions. All but the last dimension of the inputs should be the same.
- Input2:  $(N, *, H_{in2})$  where  $H_{in2} = \text{in2\_features}$ .
- Output:  $(N, *, H_{out})$  where  $H_{out} = \text{out\_features}$  and all but the last dimension are the same shape as the input.

#### **Attributes**

- weight: the learnable weights of the module of shape (out\_features, in1\_features, in2\_features). The values are initialized from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$ , where  $k=\frac{1}{\text{in1}\_\text{features}}$
- bias: the learnable bias of the module of shape (out\_features). If bias is TRUE, the values are initialized from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$ , where  $k=\frac{1}{\text{in}1\_\text{features}}$

# **Examples**

```
if (torch_is_installed()) {
m <- nn_bilinear(20, 30, 50)
input1 <- torch_randn(128, 20)
input2 <- torch_randn(128, 30)
output <- m(input1, input2)
print(output$size())
}</pre>
```

nn\_buffer 189

nn\_buffer

Creates a nn\_buffer

# Description

Indicates that a tensor is a buffer in a nn\_module

# Usage

```
nn_buffer(x, persistent = TRUE)
```

# **Arguments**

x the tensor that will be converted to nn\_buffer persistent whether the buffer should be persistent or not.

nn\_celu

CELU module

# **Description**

Applies the element-wise function:

# Usage

```
nn_celu(alpha = 1, inplace = FALSE)
```

# Arguments

alpha the  $\alpha$  value for the CELU formulation. Default: 1.0 inplace can optionally do the operation in-place. Default: FALSE

### **Details**

$$CELU(x) = \max(0, x) + \min(0, \alpha * (\exp(x/\alpha) - 1))$$

More details can be found in the paper Continuously Differentiable Exponential Linear Units.

# Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

nn\_conv1d

# **Examples**

```
if (torch_is_installed()) {
m <- nn_celu()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_contrib\_sparsemax Sparsemax activation

# **Description**

Sparsemax activation module.

# Usage

```
nn\_contrib\_sparsemax(dim = -1)
```

# **Arguments**

dim

The dimension over which to apply the sparsemax function. (-1)

#### **Details**

The SparseMax activation is described in 'From Softmax to Sparsemax: A Sparse Model of Attention and Multi-Label Classification' The implementation is based on aced125/sparsemax

nn\_conv1d

Conv1D module

# **Description**

Applies a 1D convolution over an input signal composed of several input planes. In the simplest case, the output value of the layer with input size  $(N, C_{\rm in}, L)$  and output  $(N, C_{\rm out}, L_{\rm out})$  can be precisely described as:

# Usage

```
nn_conv1d(
   in_channels,
   out_channels,
   kernel_size,
   stride = 1,
   padding = 0,
   dilation = 1,
   groups = 1,
   bias = TRUE,
   padding_mode = "zeros"
)
```

nn\_conv1d

#### **Arguments**

in\_channels (int): Number of channels in the input image out\_channels (int): Number of channels produced by the convolution kernel\_size (int or tuple): Size of the convolving kernel stride (int or tuple, optional): Stride of the convolution. Default: 1 (int, tuple or str, optional) – Padding added to both sides of the input. Default: 0 padding (int or tuple, optional): Spacing between kernel elements. Default: 1 dilation (int, optional): Number of blocked connections from input channels to output groups channels. Default: 1 bias (bool, optional): If TRUE, adds a learnable bias to the output. Default: TRUE

padding\_mode (string, optional): 'zeros', 'reflect', 'replicate' or 'circular'. Default:

'zeros'

#### **Details**

$$\operatorname{out}(N_i, C_{\operatorname{Out}_j}) = \operatorname{bias}(C_{\operatorname{Out}_j}) + \sum_{k=0}^{C_{in}-1} \operatorname{weight}(C_{\operatorname{Out}_j}, k) \star \operatorname{input}(N_i, k)$$

where  $\star$  is the valid cross-correlation operator, N is a batch size, C denotes a number of channels, L is a length of signal sequence.

- stride controls the stride for the cross-correlation, a single number or a one-element tuple.
- padding controls the amount of implicit zero-paddings on both sides for padding number of points.
- dilation controls the spacing between the kernel points; also known as the à trous algorithm. It is harder to describe, but this link has a nice visualization of what dilation does.
- groups controls the connections between inputs and outputs. in\_channels and out\_channels must both be divisible by groups. For example,
  - At groups=1, all inputs are convolved to all outputs.
  - At groups=2, the operation becomes equivalent to having two conv layers side by side, each seeing half the input channels, and producing half the output channels, and both subsequently concatenated.
  - At groups= in\_channels, each input channel is convolved with its own set of filters, of size  $\left| \frac{out\_channels}{in\_channels} \right|$ .

## Note

Depending of the size of your kernel, several (of the last) columns of the input might be lost, because it is a valid cross-correlation, *and not a full* cross-correlation. It is up to the user to add proper padding.

When groups == in\_channels and out\_channels == K \* in\_channels, where K is a positive integer, this operation is also termed in literature as depthwise convolution. In other words, for an input of size  $(N, C_{in}, L_{in})$ , a depthwise convolution with a depthwise multiplier K, can be constructed by arguments  $(C_{in} = C_{in}, C_{out} = C_{in} \times K, ..., \text{groups} = C_{in})$ .

nn\_conv2d

### Shape

- Input:  $(N, C_{in}, L_{in})$
- Output:  $(N, C_{out}, L_{out})$  where

$$L_{out} = \left\lfloor \frac{L_{in} + 2 \times \text{padding} - \text{dilation} \times (\text{kernel\_size} - 1) - 1}{\text{stride}} + 1 \right\rfloor$$

### **Attributes**

- weight (Tensor): the learnable weights of the module of shape (out\_channels,  $\frac{\text{in\_channels}}{\text{groups}}$ , kernel\_size). The values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\text{in}}*\text{kernel\_size}}$
- bias (Tensor): the learnable bias of the module of shape (out\_channels). If bias is TRUE, then the values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\mbox{in}}*\mbox{kernel\_size}}$

# **Examples**

```
if (torch_is_installed()) {
m <- nn_conv1d(16, 33, 3, stride = 2)
input <- torch_randn(20, 16, 50)
output <- m(input)
}</pre>
```

nn\_conv2d

Conv2D module

# **Description**

Applies a 2D convolution over an input signal composed of several input planes.

# Usage

```
nn_conv2d(
   in_channels,
   out_channels,
   kernel_size,
   stride = 1,
   padding = 0,
   dilation = 1,
   groups = 1,
   bias = TRUE,
   padding_mode = "zeros"
)
```

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### **Arguments**

in\_channels (int): Number of channels in the input image

out\_channels (int): Number of channels produced by the convolution

kernel\_size (int or tuple): Size of the convolving kernel

stride (int or tuple, optional): Stride of the convolution. Default: 1

padding (int or tuple or string, optional): Zero-padding added to both sides of the input.

controls the amount of padding applied to the input. It can be either a string 'valid', 'same' or a tuple of ints giving the amount of implicit padding applied

on both sides. Default: 0

dilation (int or tuple, optional): Spacing between kernel elements. Default: 1

groups (int, optional): Number of blocked connections from input channels to output

channels. Default: 1

bias (bool, optional): If TRUE, adds a learnable bias to the output. Default: TRUE

padding\_mode (string, optional): 'zeros', 'reflect', 'replicate' or 'circular'. Default:

'zeros'

#### **Details**

In the simplest case, the output value of the layer with input size  $(N, C_{in}, H, W)$  and output  $(N, C_{out}, H_{out}, W_{out})$  can be precisely described as:

$$\operatorname{out}(N_i, C_{\operatorname{Out}_j}) = \operatorname{bias}(C_{\operatorname{Out}_j}) + \sum_{k=0}^{C_{\operatorname{in}}-1} \operatorname{weight}(C_{\operatorname{Out}_j}, k) \star \operatorname{input}(N_i, k)$$

where  $\star$  is the valid 2D cross-correlation operator, N is a batch size, C denotes a number of channels, H is a height of input planes in pixels, and W is width in pixels.

- stride controls the stride for the cross-correlation, a single number or a tuple.
- padding controls the amount of implicit zero-paddings on both sides for padding number of points for each dimension.
- dilation controls the spacing between the kernel points; also known as the à trous algorithm. It is harder to describe, but this link\_ has a nice visualization of what dilation does.
- groups controls the connections between inputs and outputs. in\_channels and out\_channels must both be divisible by groups. For example,
  - At groups=1, all inputs are convolved to all outputs.
  - At groups=2, the operation becomes equivalent to having two conv layers side by side, each seeing half the input channels, and producing half the output channels, and both subsequently concatenated.
  - At groups= in\_channels, each input channel is convolved with its own set of filters, of size: \( \begin{array}{c} \frac{out\_channels}{in \channels} \end{array} \).

The parameters kernel\_size, stride, padding, dilation can either be:

- a single int in which case the same value is used for the height and width dimension
- a tuple of two ints in which case, the first int is used for the height dimension, and the second int for the width dimension

nn\_conv2d

### Note

Depending of the size of your kernel, several (of the last) columns of the input might be lost, because it is a valid cross-correlation, and not a full cross-correlation. It is up to the user to add proper padding.

When groups == in\_channels and out\_channels == K \* in\_channels, where K is a positive integer, this operation is also termed in literature as depthwise convolution. In other words, for an input of size :math: (N, C\_{in}, H\_{in}, W\_{in}), a depthwise convolution with a depthwise multiplier K, can be constructed by arguments  $(in\_channels = C_{in}, out\_channels = C_{in} \times K, ..., groups = C_{in})$ .

In some circumstances when using the CUDA backend with CuDNN, this operator may select a nondeterministic algorithm to increase performance. If this is undesirable, you can try to make the operation deterministic (potentially at a performance cost) by setting backends\_cudnn\_deterministic = TRUE.

# Shape

- Input:  $(N, C_{in}, H_{in}, W_{in})$
- Output:  $(N, C_{out}, H_{out}, W_{out})$  where

$$H_{out} = \left\lfloor \frac{H_{in} + 2 \times \operatorname{padding}[0] - \operatorname{dilation}[0] \times (\operatorname{kernel\_size}[0] - 1) - 1}{\operatorname{stride}[0]} + 1 \right\rfloor$$

$$W_{out} = \left\lfloor \frac{W_{in} + 2 \times \operatorname{padding}[1] - \operatorname{dilation}[1] \times (\operatorname{kernel\_size}[1] - 1) - 1}{\operatorname{stride}[1]} + 1 \right\rfloor$$

#### **Attributes**

- weight (Tensor): the learnable weights of the module of shape (out\_channels,  $\frac{\text{in\_channels}}{\text{groups}}$ , kernel\_size[0], kernel\_size[1]). The values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k = \frac{groups}{C_{\inf}*\prod_{i=0}^{i} \text{kernel\_size}[i]}$
- bias (Tensor): the learnable bias of the module of shape (out\_channels). If bias is TRUE, then the values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\mathbf{in}}*\prod_{i=0}^{l} \text{kernel\_size}[i]}$

#### **Examples**

```
if (torch_is_installed()) {

# With square kernels and equal stride
m <- nn_conv2d(16, 33, 3, stride = 2)
# non-square kernels and unequal stride and with padding
m <- nn_conv2d(16, 33, c(3, 5), stride = c(2, 1), padding = c(4, 2))
# non-square kernels and unequal stride and with padding and dilation
m <- nn_conv2d(16, 33, c(3, 5), stride = c(2, 1), padding = c(4, 2), dilation = c(3, 1))
input <- torch_randn(20, 16, 50, 100)
output <- m(input)
}</pre>
```

nn\_conv3d 195

nn\_conv3d Conv3D module

### **Description**

Applies a 3D convolution over an input signal composed of several input planes. In the simplest case, the output value of the layer with input size  $(N, C_{in}, D, H, W)$  and output  $(N, C_{out}, D_{out}, H_{out}, W_{out})$  can be precisely described as:

### Usage

```
nn_conv3d(
   in_channels,
   out_channels,
   kernel_size,
   stride = 1,
   padding = 0,
   dilation = 1,
   groups = 1,
   bias = TRUE,
   padding_mode = "zeros"
)
```

#### **Arguments**

in\_channels (int): Number of channels in the input image out\_channels (int): Number of channels produced by the convolution kernel\_size (int or tuple): Size of the convolving kernel stride (int or tuple, optional): Stride of the convolution. Default: 1 (int, tuple or str, optional): padding added to all six sides of the input. Default: padding dilation (int or tuple, optional): Spacing between kernel elements. Default: 1 (int, optional): Number of blocked connections from input channels to output groups channels. Default: 1 bias (bool, optional): If TRUE, adds a learnable bias to the output. Default: TRUE padding\_mode (string, optional): 'zeros', 'reflect', 'replicate' or 'circular'. Default: 'zeros'

# **Details**

$$out(N_i, C_{out_j}) = bias(C_{out_j}) + \sum_{k=0}^{C_{in}-1} weight(C_{out_j}, k) \star input(N_i, k)$$

where  $\star$  is the valid 3D cross-correlation operator

nn\_conv3d

- stride controls the stride for the cross-correlation.
- padding controls the amount of implicit zero-paddings on both sides for padding number of points for each dimension.
- dilation controls the spacing between the kernel points; also known as the à trous algorithm. It is harder to describe, but this link\_ has a nice visualization of what dilation does.
- groups controls the connections between inputs and outputs. in\_channels and out\_channels must both be divisible by groups. For example,
- At groups=1, all inputs are convolved to all outputs.
- At groups=2, the operation becomes equivalent to having two conv layers side by side, each seeing half the input channels, and producing half the output channels, and both subsequently concatenated.
- At groups= in\_channels, each input channel is convolved with its own set of filters, of size
   \[ \frac{out\_channels}{in\_channels} \] \[ \].

The parameters kernel\_size, stride, padding, dilation can either be:

- a single int in which case the same value is used for the depth, height and width dimension
- a tuple of three ints in which case, the first int is used for the depth dimension, the second int for the height dimension and the third int for the width dimension

# Shape

- Input:  $(N, C_{in}, D_{in}, H_{in}, W_{in})$
- Output:  $(N, C_{out}, D_{out}, H_{out}, W_{out})$  where

$$D_{out} = \left\lfloor \frac{D_{in} + 2 \times \operatorname{padding}[0] - \operatorname{dilation}[0] \times (\operatorname{kernel\_size}[0] - 1) - 1}{\operatorname{stride}[0]} + 1 \right\rfloor$$

$$H_{out} = \left\lfloor \frac{H_{in} + 2 \times \operatorname{padding}[1] - \operatorname{dilation}[1] \times (\operatorname{kernel\_size}[1] - 1) - 1}{\operatorname{stride}[1]} + 1 \right\rfloor$$

$$W_{out} = \left\lfloor \frac{W_{in} + 2 \times \operatorname{padding}[2] - \operatorname{dilation}[2] \times (\operatorname{kernel\_size}[2] - 1) - 1}{\operatorname{stride}[2]} + 1 \right\rfloor$$

#### **Attributes**

- weight (Tensor): the learnable weights of the module of shape (out\_channels,  $\frac{\text{in\_channels}}{\text{groups}}$ , kernel\_size[0], kernel\_size[1], kernel\_size[2]). The values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\mbox{in}}*\prod_{i=0}^2 \mbox{kernel\_size}[i]}$
- bias (Tensor): the learnable bias of the module of shape (out\_channels). If bias is True, then the values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\mbox{in}}*\prod_{i=0}^2 \mbox{kernel\_size}[i]}$

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#### Note

Depending of the size of your kernel, several (of the last) columns of the input might be lost, because it is a valid cross-correlation, *and not a full* cross-correlation. It is up to the user to add proper padding.

When groups == in\_channels and out\_channels == K \* in\_channels, where K is a positive integer, this operation is also termed in literature as depthwise convolution. In other words, for an input of size  $(N, C_{in}, D_{in}, H_{in}, W_{in})$ , a depthwise convolution with a depthwise multiplier K, can be constructed by arguments  $(in\_channels = C_{in}, out\_channels = C_{in} \times K, ..., groups = C_{in})$ .

In some circumstances when using the CUDA backend with CuDNN, this operator may select a nondeterministic algorithm to increase performance. If this is undesirable, you can try to make the operation deterministic (potentially at a performance cost) by setting torch.backends.cudnn.deterministic = TRUE. Please see the notes on :doc:/notes/randomness for background.

# **Examples**

```
if (torch_is_installed()) {
# With square kernels and equal stride
m <- nn_conv3d(16, 33, 3, stride = 2)
# non-square kernels and unequal stride and with padding
m <- nn_conv3d(16, 33, c(3, 5, 2), stride = c(2, 1, 1), padding = c(4, 2, 0))
input <- torch_randn(20, 16, 10, 50, 100)
output <- m(input)
}</pre>
```

#### **Description**

Applies a 1D transposed convolution operator over an input image composed of several input planes.

### Usage

```
nn_conv_transpose1d(
   in_channels,
   out_channels,
   kernel_size,
   stride = 1,
   padding = 0,
   output_padding = 0,
   groups = 1,
   bias = TRUE,
   dilation = 1,
   padding_mode = "zeros"
)
```

#### **Arguments**

in\_channels (int): Number of channels in the input image (int): Number of channels produced by the convolution out\_channels kernel\_size (int or tuple): Size of the convolving kernel (int or tuple, optional): Stride of the convolution. Default: 1 stride (int or tuple, optional): dilation \* (kernel\_size - 1) - padding zero-padding padding will be added to both sides of the input. Default: 0 (int or tuple, optional): Additional size added to one side of the output shape. output\_padding Default: 0 (int, optional): Number of blocked connections from input channels to output groups channels. Default: 1 bias (bool, optional): If True, adds a learnable bias to the output. Default: TRUE dilation (int or tuple, optional): Spacing between kernel elements. Default: 1

#### Details

padding\_mode

This module can be seen as the gradient of Conv1d with respect to its input. It is also known as a fractionally-strided convolution or a deconvolution (although it is not an actual deconvolution operation).

• stride controls the stride for the cross-correlation.

'zeros'

padding controls the amount of implicit zero-paddings on both sides for dilation \* (kernel\_size
 - 1) - padding number of points. See note below for details.

(string, optional): 'zeros', 'reflect', 'replicate' or 'circular'. Default:

- output\_padding controls the additional size added to one side of the output shape. See note below for details.
- dilation controls the spacing between the kernel points; also known as the à trous algorithm. It is harder to describe, but this link has a nice visualization of what dilation does.
- groups controls the connections between inputs and outputs. in\_channels and out\_channels must both be divisible by groups. For example,
  - At groups=1, all inputs are convolved to all outputs.
  - At groups=2, the operation becomes equivalent to having two conv layers side by side, each seeing half the input channels, and producing half the output channels, and both subsequently concatenated.
  - At groups= in\_channels, each input channel is convolved with its own set of filters (of size  $\left| \frac{out\_channels}{in\_channels} \right|$ ).

# Shape

- Input:  $(N, C_{in}, L_{in})$
- Output:  $(N, C_{out}, L_{out})$  where

 $L_{out} = (L_{in} - 1) \times \text{stride} - 2 \times \text{padding} + \text{dilation} \times (\text{kernel\_size} - 1) + \text{output\_padding} + 1$ 

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### **Attributes**

• weight (Tensor): the learnable weights of the module of shape (in\_channels,  $\frac{\text{out\_channels}}{\text{groups}}$ , kernel\_size). The values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k = \frac{groups}{C_{\text{Out}}*\text{kernel\_size}}$ 

• bias (Tensor): the learnable bias of the module of shape (out\_channels). If bias is TRUE, then the values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\text{Out}}*\text{kernel\_size}}$ 

#### Note

Depending of the size of your kernel, several (of the last) columns of the input might be lost, because it is a valid cross-correlation, *and not a full* cross-correlation. It is up to the user to add proper padding.

The padding argument effectively adds dilation \* (kernel\_size - 1) - padding amount of zero padding to both sizes of the input. This is set so that when a ~torch.nn.Conv1d and a ~torch.nn.ConvTranspose1d are initialized with same parameters, they are inverses of each other in regard to the input and output shapes. However, when stride > 1, ~torch.nn.Conv1d maps multiple input shapes to the same output shape. output\_padding is provided to resolve this ambiguity by effectively increasing the calculated output shape on one side. Note that output\_padding is only used to find output shape, but does not actually add zero-padding to output.

In some circumstances when using the CUDA backend with CuDNN, this operator may select a nondeterministic algorithm to increase performance. If this is undesirable, you can try to make the operation deterministic (potentially at a performance cost) by setting torch.backends.cudnn.deterministic = TRUE.

### **Examples**

```
if (torch_is_installed()) {
m <- nn_conv_transpose1d(32, 16, 2)
input <- torch_randn(10, 32, 2)
output <- m(input)
}</pre>
```

nn\_conv\_transpose2d

ConvTranpose2D module

# Description

Applies a 2D transposed convolution operator over an input image composed of several input planes.

# Usage

```
nn_conv_transpose2d(
  in_channels,
  out_channels,
  kernel_size,
  stride = 1,
```

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```
padding = 0,
output_padding = 0,
groups = 1,
bias = TRUE,
dilation = 1,
padding_mode = "zeros"
```

#### **Arguments**

in\_channels (int): Number of channels in the input image out\_channels (int): Number of channels produced by the convolution kernel size (int or tuple): Size of the convolving kernel stride (int or tuple, optional): Stride of the convolution. Default: 1 (int or tuple, optional): dilation \* (kernel\_size - 1) - padding zero-padding padding will be added to both sides of each dimension in the input. Default: 0 (int or tuple, optional): Additional size added to one side of each dimension in output\_padding the output shape. Default: 0 (int, optional): Number of blocked connections from input channels to output groups channels. Default: 1 bias (bool, optional): If True, adds a learnable bias to the output. Default: True (int or tuple, optional): Spacing between kernel elements. Default: 1 dilation (string, optional): 'zeros', 'reflect', 'replicate' or 'circular'. Default: padding\_mode 'zeros'

#### **Details**

This module can be seen as the gradient of Conv2d with respect to its input. It is also known as a fractionally-strided convolution or a deconvolution (although it is not an actual deconvolution operation).

- stride controls the stride for the cross-correlation.
- padding controls the amount of implicit zero-paddings on both sides for dilation \* (kernel\_size
   1) padding number of points. See note below for details.
- output\_padding controls the additional size added to one side of the output shape. See note below for details.
- dilation controls the spacing between the kernel points; also known as the à trous algorithm. It is harder to describe, but this link\_ has a nice visualization of what dilation does.
- groups controls the connections between inputs and outputs. in\_channels and out\_channels must both be divisible by groups. For example,
  - At groups=1, all inputs are convolved to all outputs.
  - At groups=2, the operation becomes equivalent to having two conv layers side by side, each seeing half the input channels, and producing half the output channels, and both subsequently concatenated.

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At groups= in\_channels, each input channel is convolved with its own set of filters (of size \( \frac{out\_channels}{in\_channels} \) ).

The parameters kernel\_size, stride, padding, output\_padding can either be:

- a single int in which case the same value is used for the height and width dimensions
- a tuple of two ints in which case, the first int is used for the height dimension, and the second int for the width dimension

# Shape

- Input:  $(N, C_{in}, H_{in}, W_{in})$
- Output:  $(N, C_{out}, H_{out}, W_{out})$  where

$$H_{out} = (H_{in} - 1) \times \text{stride}[0] - 2 \times \text{padding}[0] + \text{dilation}[0] \times (\text{kernel\_size}[0] - 1) + \text{output\_padding}[0] + 1 \times (\text{padding}[0] + 1) \times$$

$$W_{out} = (W_{in} - 1) \times \text{stride}[1] - 2 \times \text{padding}[1] + \text{dilation}[1] \times (\text{kernel\_size}[1] - 1) + \text{output\_padding}[1] + 1 \times (\text{padding}[1] + 1) \times$$

#### **Attributes**

- weight (Tensor): the learnable weights of the module of shape (in\_channels,  $\frac{\text{out\_channels}}{\text{groups}}$ , kernel\_size[0], kernel\_size[1]). The values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k = \frac{groups}{C_{\text{OUI}}*\prod_{i=0}^{1} \text{kernel\_size}[i]}$
- bias (Tensor): the learnable bias of the module of shape (out\_channels) If bias is True, then the values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\mathbf{Out}}*\Pi^1_{i=0}\operatorname{kernel\_size}[i]}$

#### Note

Depending of the size of your kernel, several (of the last) columns of the input might be lost, because it is a valid cross-correlation\_, and not a full cross-correlation. It is up to the user to add proper padding.

The padding argument effectively adds dilation \* (kernel\_size - 1) - padding amount of zero padding to both sizes of the input. This is set so that when a nn\_conv2d and a nn\_conv\_transpose2d are initialized with same parameters, they are inverses of each other in regard to the input and output shapes. However, when stride > 1, nn\_conv2d maps multiple input shapes to the same output shape. output\_padding is provided to resolve this ambiguity by effectively increasing the calculated output shape on one side. Note that output\_padding is only used to find output shape, but does not actually add zero-padding to output.

In some circumstances when using the CUDA backend with CuDNN, this operator may select a nondeterministic algorithm to increase performance. If this is undesirable, you can try to make the operation deterministic (potentially at a performance cost) by setting torch.backends.cudnn.deterministic = TRUE.

202 nn\_conv\_transpose3d

### **Examples**

```
if (torch_is_installed()) {
# With square kernels and equal stride
m <- nn_conv_transpose2d(16, 33, 3, stride = 2)</pre>
# non-square kernels and unequal stride and with padding
m < -nn_{conv_{transpose}} (16, 33, c(3, 5), stride = c(2, 1), padding = c(4, 2))
input <- torch_randn(20, 16, 50, 100)
output <- m(input)</pre>
# exact output size can be also specified as an argument
input <- torch_randn(1, 16, 12, 12)</pre>
downsample <- nn_conv2d(16, 16, 3, stride = 2, padding = 1)
upsample <- nn_conv_transpose2d(16, 16, 3, stride = 2, padding = 1)</pre>
h <- downsample(input)</pre>
h$size()
output <- upsample(h, output_size = input$size())</pre>
output$size()
}
```

nn\_conv\_transpose3d

ConvTranpose3D module

# **Description**

Applies a 3D transposed convolution operator over an input image composed of several input planes.

### Usage

```
nn_conv_transpose3d(
   in_channels,
   out_channels,
   kernel_size,
   stride = 1,
   padding = 0,
   output_padding = 0,
   groups = 1,
   bias = TRUE,
   dilation = 1,
   padding_mode = "zeros"
)
```

# **Arguments**

```
    in_channels (int): Number of channels in the input image
    out_channels (int): Number of channels produced by the convolution
    kernel_size (int or tuple): Size of the convolving kernel
    stride (int or tuple, optional): Stride of the convolution. Default: 1
```

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padding	(int or tuple, optional): dilation * (kernel_size - 1) - padding zero-padding will be added to both sides of each dimension in the input. Default: 0 output_padding (int or tuple, optional): Additional size added to one side of each dimension in the output shape. Default: 0
output_padding	(int or tuple, optional): Additional size added to one side of each dimension in the output shape. Default: $0$
groups	(int, optional): Number of blocked connections from input channels to output channels. Default: 1
bias	(bool, optional): If True, adds a learnable bias to the output. Default: True
dilation	(int or tuple, optional): Spacing between kernel elements. Default: 1
padding_mode	(string, optional): 'zeros', 'reflect', 'replicate' or 'circular'. Default: 'zeros'

#### **Details**

The transposed convolution operator multiplies each input value element-wise by a learnable kernel, and sums over the outputs from all input feature planes.

This module can be seen as the gradient of Conv3d with respect to its input. It is also known as a fractionally-strided convolution or a deconvolution (although it is not an actual deconvolution operation).

- stride controls the stride for the cross-correlation.
- padding controls the amount of implicit zero-paddings on both sides for dilation \* (kernel\_size
   1) padding number of points. See note below for details.
- output\_padding controls the additional size added to one side of the output shape. See note below for details.
- dilation controls the spacing between the kernel points; also known as the à trous algorithm. It is harder to describe, but this link\_ has a nice visualization of what dilation does.
- groups controls the connections between inputs and outputs. in\_channels and out\_channels must both be divisible by groups. For example,
  - At groups=1, all inputs are convolved to all outputs.
  - At groups=2, the operation becomes equivalent to having two conv layers side by side, each seeing half the input channels, and producing half the output channels, and both subsequently concatenated.
  - At groups= in\_channels, each input channel is convolved with its own set of filters (of size | out\_channels | ).

The parameters kernel\_size, stride, padding, output\_padding can either be:

- a single int in which case the same value is used for the depth, height and width dimensions
- a tuple of three ints in which case, the first int is used for the depth dimension, the second int for the height dimension and the third int for the width dimension

# Shape

```
• Input: (N, C_{in}, D_{in}, H_{in}, W_{in})

• Output: (N, C_{out}, D_{out}, H_{out}, W_{out}) where D_{out} = (D_{in}-1) \times \text{stride}[0] - 2 \times \text{padding}[0] + \text{dilation}[0] \times (\text{kernel\_size}[0]-1) + \text{output\_padding}[0] + 1 H_{out} = (H_{in}-1) \times \text{stride}[1] - 2 \times \text{padding}[1] + \text{dilation}[1] \times (\text{kernel\_size}[1]-1) + \text{output\_padding}[1] + 1 W_{out} = (W_{in}-1) \times \text{stride}[2] - 2 \times \text{padding}[2] + \text{dilation}[2] \times (\text{kernel\_size}[2]-1) + \text{output\_padding}[2] + 1
```

#### Attributes

- weight (Tensor): the learnable weights of the module of shape (in\_channels,  $\frac{\text{out\_channels}}{\text{groups}}$ , kernel\_size[0], kernel\_size[1], kernel\_size[2]). The values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\text{Out}}*\prod_{i=0}^2 \text{kernel\_size}[i]}$
- bias (Tensor): the learnable bias of the module of shape (out\_channels) If bias is True, then the values of these weights are sampled from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{groups}{C_{\mathrm{Out}}*\Pi_{i=0}^2 \text{ kernel\_size}[i]}$

#### Note

Depending of the size of your kernel, several (of the last) columns of the input might be lost, because it is a valid cross-correlation, *and not a full* cross-correlation. It is up to the user to add proper padding.

The padding argument effectively adds dilation \* (kernel\_size - 1) - padding amount of zero padding to both sizes of the input. This is set so that when a ~torch.nn.Conv3d and a ~torch.nn.ConvTranspose3d are initialized with same parameters, they are inverses of each other in regard to the input and output shapes. However, when stride > 1, ~torch.nn.Conv3d maps multiple input shapes to the same output shape. output\_padding is provided to resolve this ambiguity by effectively increasing the calculated output shape on one side. Note that output\_padding is only used to find output shape, but does not actually add zero-padding to output.

In some circumstances when using the CUDA backend with CuDNN, this operator may select a nondeterministic algorithm to increase performance. If this is undesirable, you can try to make the operation deterministic (potentially at a performance cost) by setting torch.backends.cudnn.deterministic = TRUE.

# Examples

```
if (torch_is_installed()) {
## Not run:

# With square kernels and equal stride
m <- nn_conv_transpose3d(16, 33, 3, stride = 2)
# non-square kernels and unequal stride and with padding
m <- nn_conv_transpose3d(16, 33, c(3, 5, 2), stride = c(2, 1, 1), padding = c(0, 4, 2))
input <- torch_randn(20, 16, 10, 50, 100)
output <- m(input)

## End(Not run)
}</pre>
```

nn\_cosine\_embedding\_loss

Cosine embedding loss

# **Description**

Creates a criterion that measures the loss given input tensors  $x_1$ ,  $x_2$  and a Tensor label y with values 1 or -1. This is used for measuring whether two inputs are similar or dissimilar, using the cosine distance, and is typically used for learning nonlinear embeddings or semi-supervised learning. The loss function for each sample is:

# Usage

```
nn_cosine_embedding_loss(margin = 0, reduction = "mean")
```

### **Arguments**

margin (float, optional): Should be a number from -1 to 1, 0 to 0.5 is suggested. If

margin is missing, the default value is 0.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

### **Details**

$$\label{eq:loss} \text{loss}(x,y) = \begin{array}{ll} 1 - \cos(x_1,x_2), & \text{if } y = 1 \\ \max(0,\cos(x_1,x_2) - \text{margin}), & \text{if } y = -1 \end{array}$$

# **Description**

This criterion combines nn\_log\_softmax() and nn\_nll\_loss() in one single class. It is useful when training a classification problem with C classes.

## Usage

```
nn_cross_entropy_loss(weight = NULL, ignore_index = -100, reduction = "mean")
```

#### **Arguments**

weight (Tensor, optional): a manual rescaling weight given to each class. If given, has

to be a Tensor of size C

ignore\_index (int, optional): Specifies a target value that is ignored and does not contribute

to the input gradient. When size\_average is TRUE, the loss is averaged over

non-ignored targets.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

#### **Details**

If provided, the optional argument weight should be a 1D Tensor assigning weight to each of the classes.

This is particularly useful when you have an unbalanced training set. The input is expected to contain raw, unnormalized scores for each class. input has to be a Tensor of size either (minibatch, C) or  $(minibatch, C, d_1, d_2, ..., d_K)$  with  $K \ge 1$  for the K-dimensional case (described later).

This criterion expects a class index in the range [0, C-1] as the target for each value of a 1D tensor of size minibatch; if ignore\_index is specified, this criterion also accepts this class index (this index may not necessarily be in the class range).

The loss can be described as:

$$loss(x, class) = -\log\left(\frac{\exp(x[class])}{\sum_{j} \exp(x[j])}\right) = -x[class] + \log\left(\sum_{j} \exp(x[j])\right)$$

or in the case of the weight argument being specified:

$$loss(x, class) = weight[class] \left( -x[class] + log \left( \sum_{j} exp(x[j]) \right) \right)$$

The losses are averaged across observations for each minibatch. Can also be used for higher dimension inputs, such as 2D images, by providing an input of size  $(minibatch, C, d_1, d_2, ..., d_K)$  with  $K \ge 1$ , where K is the number of dimensions, and a target of appropriate shape (see below).

# Shape

- Input: (N, C) where C = number of classes, or  $(N, C, d_1, d_2, ..., d_K)$  with  $K \ge 1$  in the case of K-dimensional loss.
- Target: (N) where each value is  $0 \le \operatorname{targets}[i] \le C 1$ , or  $(N, d_1, d_2, ..., d_K)$  with  $K \ge 1$  in the case of K-dimensional loss.
- Output: scalar. If reduction is 'none', then the same size as the target: (N), or  $(N, d_1, d_2, ..., d_K)$  with  $K \ge 1$  in the case of K-dimensional loss.

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### **Examples**

```
if (torch_is_installed()) {
  loss <- nn_cross_entropy_loss()
  input <- torch_randn(3, 5, requires_grad = TRUE)
  target <- torch_randint(low = 1, high = 5, size = 3, dtype = torch_long())
  output <- loss(input, target)
  output$backward()
}</pre>
```

nn\_ctc\_loss

The Connectionist Temporal Classification loss.

# **Description**

Calculates loss between a continuous (unsegmented) time series and a target sequence. CTCLoss sums over the probability of possible alignments of input to target, producing a loss value which is differentiable with respect to each input node. The alignment of input to target is assumed to be "many-to-one", which limits the length of the target sequence such that it must be  $\leq$  the input length.

# Usage

```
nn_ctc_loss(blank = 0, reduction = "mean", zero_infinity = FALSE)
```

# **Arguments**

blank (int, optional): blank label. Default 0.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the output losses will be divided by the target lengths and then the mean over the batch is taken. Default:

'mean'

zero\_infinity (bool, optional): Whether to zero infinite losses and the associated gradients.

Default: FALSE Infinite losses mainly occur when the inputs are too short to be

aligned to the targets.

## Shape

- Log\_probs: Tensor of size (T, N, C), where T = input length, N = batch size, and C = number of classes (including blank). The logarithmized probabilities of the outputs (e.g. obtained with [nnf)log\_softmax()]).
- Targets: Tensor of size (N,S) or  $(\text{sum}(\text{target\_lengths}))$ , where N= batch size and S= max target length, if shape is (N,S). It represent the target sequences. Each element in the target sequence is a class index. And the target index cannot be blank (default=0). In the (N,S) form, targets are padded to the length of the longest sequence, and stacked. In the  $(\text{sum}(\text{target\_lengths}))$  form, the targets are assumed to be un-padded and concatenated within 1 dimension.

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Input\_lengths: Tuple or tensor of size (N), where N = batch size. It represent the lengths
of the inputs (must each be ≤ T). And the lengths are specified for each sequence to achieve
masking under the assumption that sequences are padded to equal lengths.

- Target\_lengths: Tuple or tensor of size (N), where N= batch size. It represent lengths of the targets. Lengths are specified for each sequence to achieve masking under the assumption that sequences are padded to equal lengths. If target shape is (N,S), target\_lengths are effectively the stop index  $s_n$  for each target sequence, such that target\_n = targets[n,0:s\_n] for each target in a batch. Lengths must each be  $\leq S$  If the targets are given as a 1d tensor that is the concatenation of individual targets, the target\_lengths must add up to the total length of the tensor.
- Output: scalar. If reduction is 'none', then (N), where N =batch size.

[nnf)log\_softmax()]: R:nnf)log\_softmax() [n,0:s\_n]: R:n,0:s\_n

#### Note

In order to use CuDNN, the following must be satisfied: targets must be in concatenated format, all input\_lengths must be T. blank = 0, target\_lengths  $\leq 256$ , the integer arguments must be of The regular implementation uses the (more common in PyTorch) torch\_long dtype. dtype torch\_int32.

In some circumstances when using the CUDA backend with CuDNN, this operator may select a nondeterministic algorithm to increase performance. If this is undesirable, you can try to make the operation deterministic (potentially at a performance cost) by setting torch.backends.cudnn.deterministic = TRUE.

### References

A. Graves et al.: Connectionist Temporal Classification: Labelling Unsegmented Sequence Data with Recurrent Neural Networks: https://www.cs.toronto.edu/~graves/icml\_2006.pdf

# **Examples**

```
if (torch_is_installed()) {
# Target are to be padded
T <- 50 # Input sequence length
C <- 20 # Number of classes (including blank)
N <- 16 # Batch size
S <- 30 # Target sequence length of longest target in batch (padding length)
S_min <- 10 # Minimum target length, for demonstration purposes
# Initialize random batch of input vectors, for *size = (T,N,C)
input <- torch_randn(T, N, C)$log_softmax(2)$detach()$requires_grad_()
# Initialize random batch of targets (0 = blank, 1:C = classes)
target <- torch_randint(low = 1, high = C, size = c(N, S), dtype = torch_long())
input_lengths <- torch_full(size = c(N), fill_value = TRUE, dtype = torch_long())
target_lengths <- torch_randint(low = S_min, high = S, size = c(N), dtype = torch_long())
ctc_loss <- nn_ctc_loss()
loss <- ctc_loss(input, target, input_lengths, target_lengths)</pre>
```

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```
loss$backward()
# Target are to be un-padded
T <- 50 # Input sequence length
C <- 20 # Number of classes (including blank)
N <- 16 # Batch size
# Initialize random batch of input vectors, for *size = (T,N,C)
input <- torch_randn(T, N, C)$log_softmax(2)$detach()$requires_grad_()</pre>
input_lengths <- torch_full(size = c(N), fill_value = TRUE, dtype = torch_long())
# Initialize random batch of targets (0 = blank, 1:C = classes)
target_lengths < -torch_randint(low = 1, high = T, size = c(N), dtype = torch_long())
target <- torch_randint(</pre>
 low = 1, high = C, size = as.integer(sum(target_lengths)),
 dtype = torch_long()
)
ctc_loss <- nn_ctc_loss()</pre>
loss <- ctc_loss(input, target, input_lengths, target_lengths)</pre>
loss$backward()
}
```

nn\_dropout

Dropout module

### **Description**

During training, randomly zeroes some of the elements of the input tensor with probability p using samples from a Bernoulli distribution. Each channel will be zeroed out independently on every forward call.

# Usage

```
nn_dropout(p = 0.5, inplace = FALSE)
```

# **Arguments**

p probability of an element to be zeroed. Default: 0.5 inplace If set to TRUE, will do this operation in-place. Default: FALSE.

## **Details**

This has proven to be an effective technique for regularization and preventing the co-adaptation of neurons as described in the paper Improving neural networks by preventing co-adaptation of feature detectors.

Furthermore, the outputs are scaled by a factor of :math:\frac{1}{1-p} during training. This means that during evaluation the module simply computes an identity function.

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### Shape

- Input: (\*). Input can be of any shape
- Output: (\*). Output is of the same shape as input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_dropout(p = 0.2)
input <- torch_randn(20, 16)
output <- m(input)
}</pre>
```

nn\_dropout2d

Dropout2D module

# **Description**

Randomly zero out entire channels (a channel is a 2D feature map, e.g., the j-th channel of the i-th sample in the batched input is a 2D tensor input[i, j]).

### Usage

```
nn_dropout2d(p = 0.5, inplace = FALSE)
```

### **Arguments**

```
p (float, optional): probability of an element to be zero-ed.
inplace (bool, optional): If set to TRUE, will do this operation in-place
```

#### **Details**

Each channel will be zeroed out independently on every forward call with probability p using samples from a Bernoulli distribution. Usually the input comes from nn conv2d modules.

As described in the paper Efficient Object Localization Using Convolutional Networks , if adjacent pixels within feature maps are strongly correlated (as is normally the case in early convolution layers) then i.i.d. dropout will not regularize the activations and will otherwise just result in an effective learning rate decrease. In this case, nn\_dropout2d will help promote independence between feature maps and should be used instead.

### Shape

- Input: (N, C, H, W)
- Output: (N, C, H, W) (same shape as input)

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# **Examples**

```
if (torch_is_installed()) {
m <- nn_dropout2d(p = 0.2)
input <- torch_randn(20, 16, 32, 32)
output <- m(input)
}</pre>
```

nn\_dropout3d

Dropout3D module

## Description

Randomly zero out entire channels (a channel is a 3D feature map, e.g., the j-th channel of the i-th sample in the batched input is a 3D tensor input[i, j]).

# Usage

```
nn_dropout3d(p = 0.5, inplace = FALSE)
```

# **Arguments**

```
p (float, optional): probability of an element to be zeroed.
inplace (bool, optional): If set to TRUE, will do this operation in-place
```

#### Details

Each channel will be zeroed out independently on every forward call with probability p using samples from a Bernoulli distribution. Usually the input comes from nn\_conv2d modules.

As described in the paper Efficient Object Localization Using Convolutional Networks, if adjacent pixels within feature maps are strongly correlated (as is normally the case in early convolution layers) then i.i.d. dropout will not regularize the activations and will otherwise just result in an effective learning rate decrease.

In this case, nn\_dropout3d will help promote independence between feature maps and should be used instead.

# Shape

```
Input: (N, C, D, H, W)
Output: (N, C, D, H, W) (same shape as input)
```

## **Examples**

```
if (torch_is_installed()) {
m <- nn_dropout3d(p = 0.2)
input <- torch_randn(20, 16, 4, 32, 32)
output <- m(input)
}</pre>
```

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nn\_elu

ELU module

# **Description**

Applies the element-wise function:

# Usage

```
nn_elu(alpha = 1, inplace = FALSE)
```

# Arguments

alpha the  $\alpha$  value for the ELU formulation. Default: 1.0 inplace can optionally do the operation in-place. Default: FALSE

### **Details**

$$ELU(x) = max(0, x) + min(0, \alpha * (exp(x) - 1))$$

# Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_elu()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_embedding

Embedding module

# **Description**

A simple lookup table that stores embeddings of a fixed dictionary and size. This module is often used to store word embeddings and retrieve them using indices. The input to the module is a list of indices, and the output is the corresponding word embeddings.

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# Usage

```
nn_embedding(
  num_embeddings,
  embedding_dim,
  padding_idx = NULL,
  max_norm = NULL,
  norm_type = 2,
  scale_grad_by_freq = FALSE,
  sparse = FALSE,
  .weight = NULL
)
```

#### **Arguments**

```
num_embeddings
                  (int): size of the dictionary of embeddings
embedding_dim
                   (int): the size of each embedding vector
padding_idx
                   (int, optional): If given, pads the output with the embedding vector at padding_idx
                   (initialized to zeros) whenever it encounters the index.
                   (float, optional): If given, each embedding vector with norm larger than max_norm
max_norm
                   is renormalized to have norm max_norm.
                   (float, optional): The p of the p-norm to compute for the max_norm option. De-
norm_type
scale_grad_by_freq
                   (boolean, optional): If given, this will scale gradients by the inverse of frequency
                   of the words in the mini-batch. Default False.
                   (bool, optional): If True, gradient w.r.t. weight matrix will be a sparse tensor.
sparse
.weight
                   (Tensor) embeddings weights (in case you want to set it manually)
                   See Notes for more details regarding sparse gradients.
```

#### Attributes

• weight (Tensor): the learnable weights of the module of shape (num\_embeddings, embedding\_dim) initialized from  $\mathcal{N}(0,1)$ 

#### Shape

- Input: (\*), LongTensor of arbitrary shape containing the indices to extract
- Output: (\*, H), where \* is the input shape and  $H = \text{embedding\_dim}$

#### Note

Keep in mind that only a limited number of optimizers support sparse gradients: currently it's optim.SGD (CUDA and CPU), optim.SparseAdam (CUDA and CPU) and optim.Adagrad (CPU)

With padding\_idx set, the embedding vector at padding\_idx is initialized to all zeros. However, note that this vector can be modified afterwards, e.g., using a customized initialization method, and thus changing the vector used to pad the output. The gradient for this vector from nn\_embedding is always zero.

214 nn\_embedding\_bag

### **Examples**

```
if (torch_is_installed()) {
# an Embedding module containing 10 tensors of size 3
embedding <- nn_embedding(10, 3)
# a batch of 2 samples of 4 indices each
input <- torch_tensor(rbind(c(1, 2, 4, 5), c(4, 3, 2, 9)), dtype = torch_long())
embedding(input)
# example with padding_idx
embedding <- nn_embedding(10, 3, padding_idx = 1)
input <- torch_tensor(matrix(c(1, 3, 1, 6), nrow = 1), dtype = torch_long())
embedding(input)
}</pre>
```

nn\_embedding\_bag

Embedding bag module

# **Description**

Computes sums, means or maxes of bags of embeddings, without instantiating the intermediate embeddings.

#### Usage

```
nn_embedding_bag(
  num_embeddings,
  embedding_dim,
  max_norm = NULL,
  norm_type = 2,
  scale_grad_by_freq = FALSE,
  mode = "mean",
  sparse = FALSE,
  include_last_offset = FALSE,
  padding_idx = NULL,
  .weight = NULL
)
```

#### **Arguments**

```
num_embeddings (int): size of the dictionary of embeddings
embedding_dim (int): the size of each embedding vector

max_norm (float, optional): If given, each embedding vector with norm larger than max_norm
is renormalized to have norm max_norm.

norm_type (float, optional): The p of the p-norm to compute for the max_norm option. Default 2

scale_grad_by_freq
(boolean, optional): If given, this will scale gradients by the inverse of frequency
```

(boolean, optional): If given, this will scale gradients by the inverse of frequency of the words in the mini-batch. Default False.

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(string, optional): "sum", "mean" or "max". Specifies the way to reduce the bag. mode "sum" computes the weighted sum, taking per\_sample\_weights into consideration. "mean" computes the average of the values in the bag, "max" computes the max value over each bag. (bool, optional): If True, gradient w.r.t. weight matrix will be a sparse tensor. sparse See Notes for more details regarding sparse gradients. include\_last\_offset (bool, optional): if True, offsets has one additional element, where the last element is equivalent to the size of indices. This matches the CSR format. (int, optional): If given, pads the output with the embedding vector at padding\_idx padding\_idx (initialized to zeros) whenever it encounters the index. (Tensor, optional) embeddings weights (in case you want to set it manually) .weight

#### **Attributes**

• weight (Tensor): the learnable weights of the module of shape (num\_embeddings, embedding dim) initialized from  $\mathcal{N}(0,1)$ 

# **Examples**

```
if (torch_is_installed()) {
# an EmbeddingBag module containing 10 tensors of size 3
embedding_sum <- nn_embedding_bag(10, 3, mode = 'sum')</pre>
# a batch of 2 samples of 4 indices each
input \leftarrow torch_tensor(c(1, 2, 4, 5, 4, 3, 2, 9), dtype = torch_long())
offsets \leftarrow torch_tensor(c(0, 4), dtype = torch_long())
embedding_sum(input, offsets)
# example with padding_idx
embedding_sum <- nn_embedding_bag(10, 3, mode = 'sum', padding_idx = 1)</pre>
input <- torch_tensor(c(2, 2, 2, 2, 4, 3, 2, 9), dtype = torch_long())
offsets <- torch_tensor(c(0, 4), dtype = torch_long())
embedding_sum(input, offsets)
# An EmbeddingBag can be loaded from an Embedding like so
embedding <- nn_embedding(10, 3, padding_idx = 2)</pre>
embedding_sum <- nn_embedding_bag$from_pretrained(embedding$weight,</pre>
                                                    padding_idx = embedding$padding_idx,
                                                    mode='sum')
}
```

nn\_flatten

Flattens a contiguous range of dims into a tensor.

### **Description**

For use with nn\_sequential.

# Usage

```
nn_flatten(start_dim = 2, end_dim = -1)
```

# Arguments

```
start_dim first dim to flatten (default = 2).
end_dim last dim to flatten (default = -1).
```

# Shape

- Input: (\*, S\_start,..., S\_i, ..., S\_end, \*), where S\_i is the size at dimension i and \* means any number of dimensions including none.
- Output: (\*, S\_start\*...\*S\_i\*...S\_end, \*).

# See Also

nn\_unflatten

### **Examples**

```
if (torch_is_installed()) {
input <- torch_randn(32, 1, 5, 5)
m <- nn_flatten()
m(input)
}</pre>
```

```
nn_fractional_max_pool2d
```

Applies a 2D fractional max pooling over an input signal composed of several input planes.

# Description

Fractional MaxPooling is described in detail in the paper Fractional MaxPooling by Ben Graham

# Usage

```
nn_fractional_max_pool2d(
  kernel_size,
  output_size = NULL,
  output_ratio = NULL,
  return_indices = FALSE
)
```

# **Arguments**

the size of the window to take a max over. Can be a single number k (for a square kernel of k x k) or a tuple (kh, kw)

output\_size the target output size of the image of the form oH x oW. Can be a tuple (oH, oW) or a single number oH for a square image oH x oH

```
output_ratio If one wants to have an output size as a ratio of the input size, this option can be given. This has to be a number or tuple in the range (0, 1)

return_indices if TRUE, will return the indices along with the outputs. Useful to pass to nn_max_unpool2d().

Default: FALSE
```

#### **Details**

The max-pooling operation is applied in  $kH \times kW$  regions by a stochastic step size determined by the target output size. The number of output features is equal to the number of input planes.

## **Examples**

```
if (torch_is_installed()) {
# pool of square window of size=3, and target output size 13x12
m <- nn_fractional_max_pool2d(3, output_size = c(13, 12))
# pool of square window and target output size being half of input image size
m <- nn_fractional_max_pool2d(3, output_ratio = c(0.5, 0.5))
input <- torch_randn(20, 16, 50, 32)
output <- m(input)
}</pre>
```

nn\_fractional\_max\_pool3d

Applies a 3D fractional max pooling over an input signal composed of several input planes.

### Description

Fractional MaxPooling is described in detail in the paper Fractional MaxPooling by Ben Graham

## Usage

```
nn_fractional_max_pool3d(
  kernel_size,
  output_size = NULL,
  output_ratio = NULL,
  return_indices = FALSE
)
```

#### **Arguments**

kernel_size	the size of the window to take a max over. Can be a single number k (for a square kernel of k x k x k) or a tuple (kt x kh x kw)
output_size	the target output size of the image of the form oT $\times$ oH $\times$ oW. Can be a tuple (oT, oH, oW) or a single number oH for a square image oH $\times$ oH
output_ratio	If one wants to have an output size as a ratio of the input size, this option can be given. This has to be a number or tuple in the range $(0, 1)$
return_indices	if TRUE, will return the indices along with the outputs. Useful to pass to $nn_max_unpool3d()$ . Default: FALSE

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### **Details**

The max-pooling operation is applied in kTxkHxkW regions by a stochastic step size determined by the target output size. The number of output features is equal to the number of input planes.

### **Examples**

```
if (torch_is_installed()) {
# pool of cubic window of size=3, and target output size 13x12x11
m <- nn_fractional_max_pool3d(3, output_size = c(13, 12, 11))
# pool of cubic window and target output size being half of input size
m <- nn_fractional_max_pool3d(3, output_ratio = c(0.5, 0.5, 0.5))
input <- torch_randn(20, 16, 50, 32, 16)
output <- m(input)
}</pre>
```

nn\_gelu

GELU module

### **Description**

Applies the Gaussian Error Linear Units function:

$$GELU(x) = x * \Phi(x)$$

#### Usage

```
nn_gelu(approximate = "none")
```

### **Arguments**

```
approximate the gelu approximation algorithm to use: 'none' or 'tanh'. Default: 'none'.
```

#### **Details**

where  $\Phi(x)$  is the Cumulative Distribution Function for Gaussian Distribution.

# Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

```
if (torch_is_installed()) {
m <- nn_gelu()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

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nn\_glu

GLU module

## **Description**

Applies the gated linear unit function  $GLU(a,b) = a \otimes \sigma(b)$  where a is the first half of the input matrices and b is the second half.

### Usage

```
nn_glu(dim = -1)
```

# Arguments

dim

(int): the dimension on which to split the input. Default: -1

## Shape

- Input:  $(*_1, N, *_2)$  where \* means, any number of additional dimensions
- Output:  $(*_1, M, *_2)$  where M = N/2

## **Examples**

```
if (torch_is_installed()) {
m <- nn_glu()
input <- torch_randn(4, 2)
output <- m(input)
}</pre>
```

nn\_group\_norm

Group normalization

# Description

Applies Group Normalization over a mini-batch of inputs as described in the paper Group Normalization.

## Usage

```
nn_group_norm(num_groups, num_channels, eps = 1e-05, affine = TRUE)
```

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### Arguments

num\_groups (int): number of groups to separate the channels into

num\_channels (int): number of channels expected in input

eps a value added to the denominator for numerical stability. Default: 1e-5

affine a boolean value that when set to TRUE, this module has learnable per-channel

affine parameters initialized to ones (for weights) and zeros (for biases). Default:

TRUE.

#### **Details**

$$y = \frac{x - \mathbf{E}[x]}{\sqrt{\mathbf{Var}[x] + \epsilon}} * \gamma + \beta$$

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.  $\gamma$  and  $\beta$  are learnable per-channel affine transform parameter vectors of size num\_channels if affine is TRUE. The standard-deviation is calculated via the biased estimator, equivalent to torch\_var(input, unbiased=FALSE).

### Shape

- Input: (N, C, \*) where  $C = \text{num\_channels}$
- Output: (N, C, \*) (same shape as input)

### Note

This layer uses statistics computed from input data in both training and evaluation modes.

```
if (torch_is_installed()) {
  input <- torch_randn(20, 6, 10, 10)
# Separate 6 channels into 3 groups
m <- nn_group_norm(3, 6)
# Separate 6 channels into 6 groups (equivalent with [nn_instance_morm])
m <- nn_group_norm(6, 6)
# Put all 6 channels into a single group (equivalent with [nn_layer_norm])
m <- nn_group_norm(1, 6)
# Activating the module
output <- m(input)
}</pre>
```

nn\_gru 221

nn\_gru Applies a multi-layer gated recurrent unit (GRU) RNN to an input sequence.

### **Description**

For each element in the input sequence, each layer computes the following function:

### Usage

```
nn_gru(
   input_size,
   hidden_size,
   num_layers = 1,
   bias = TRUE,
   batch_first = FALSE,
   dropout = 0,
   bidirectional = FALSE,
   ...
)
```

### Arguments

input\_size The number of expected features in the input x hidden\_size The number of features in the hidden state h Number of recurrent layers. E.g., setting num\_layers=2 would mean stacking num\_layers two GRUs together to form a stacked GRU, with the second GRU taking in outputs of the first GRU and computing the final results. Default: 1 bias If FALSE, then the layer does not use bias weights b\_ih and b\_hh. Default: TRUE batch\_first If TRUE, then the input and output tensors are provided as (batch, seq, feature). Default: FALSE If non-zero, introduces a Dropout layer on the outputs of each GRU layer except dropout the last layer, with dropout probability equal to dropout. Default: 0 If TRUE, becomes a bidirectional GRU. Default: FALSE bidirectional currently unused. . . .

#### **Details**

$$\begin{split} r_t &= \sigma(W_{ir}x_t + b_{ir} + W_{hr}h_{(t-1)} + b_{hr}) \\ z_t &= \sigma(W_{iz}x_t + b_{iz} + W_{hz}h_{(t-1)} + b_{hz}) \\ n_t &= \tanh(W_{in}x_t + b_{in} + r_t(W_{hn}h_{(t-1)} + b_{hn})) \\ h_t &= (1 - z_t)n_t + z_th_{(t-1)} \end{split}$$

where  $h_t$  is the hidden state at time t,  $x_t$  is the input at time t,  $h_{(t-1)}$  is the hidden state of the previous layer at time t-1 or the initial hidden state at time 0, and  $r_t$ ,  $z_t$ ,  $n_t$  are the reset, update, and new gates, respectively.  $\sigma$  is the sigmoid function.

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### **Inputs**

Inputs: input, h\_0

• **input** of shape (seq\_len, batch, input\_size): tensor containing the features of the input sequence. The input can also be a packed variable length sequence. See nn\_utils\_rnn\_pack\_padded\_sequence() for details.

• h\_0 of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the initial hidden state for each element in the batch. Defaults to zero if not provided.

### **Outputs**

Outputs: output, h\_n

- output of shape (seq\_len, batch, num\_directions \* hidden\_size): tensor containing the output features h\_t from the last layer of the GRU, for each t. If a PackedSequence has been given as the input, the output will also be a packed sequence. For the unpacked case, the directions can be separated using output\$view(c(seq\_len, batch, num\_directions, hidden\_size)), with forward and backward being direction 0 and 1 respectively. Similarly, the directions can be separated in the packed case.
- h\_n of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the hidden state for t = seq\_len Like *output*, the layers can be separated using h\_n\$view(num\_layers, num\_directions, batch, hidden\_size).

#### **Attributes**

- weight\_ih\_1[k]: the learnable input-hidden weights of the  $k^{th}$  layer (W\_irlW\_izlW\_in), of shape (3\*hidden\_size x input\_size)
- weight\_hh\_1[k]: the learnable hidden-hidden weights of the k<sup>th</sup> layer (W\_hrlW\_hzlW\_hn), of shape (3\*hidden\_size x hidden\_size)
- bias\_ih\_1[k] : the learnable input-hidden bias of the  $k^{th}$  layer (b\_irlb\_izlb\_in), of shape (3\*hidden\_size)
- bias\_hh\_1[k] : the learnable hidden-hidden bias of the  $k^{th}$  layer (b\_hrlb\_hzlb\_hn), of shape (3\*hidden\_size)

#### Note

All the weights and biases are initialized from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{1}{\text{hidden size}}$ 

```
if (torch_is_installed()) {
rnn <- nn_gru(10, 20, 2)
input <- torch_randn(5, 3, 10)
h0 <- torch_randn(2, 3, 20)
output <- rnn(input, h0)
}</pre>
```

nn\_hardshrink 223

nn\_hardshrink

Hardshwink module

## **Description**

Applies the hard shrinkage function element-wise:

# Usage

```
nn_hardshrink(lambd = 0.5)
```

### **Arguments**

lambd

the  $\lambda$  value for the Hardshrink formulation. Default: 0.5

## **Details**

$$\mbox{HardShrink}(x) = \left\{ \begin{array}{ll} x, & \mbox{if } x > \lambda \\ x, & \mbox{if } x < -\lambda \\ 0, & \mbox{otherwise} \end{array} \right.$$

## Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N,\*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_hardshrink()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_hardsigmoid

Hardsigmoid module

# Description

Applies the element-wise function:

## Usage

```
nn_hardsigmoid()
```

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## **Details**

$$\operatorname{Hardsigmoid}(x) = \left\{ \begin{array}{ll} 0 & \text{if } x \leq -3, \\ 1 & \text{if } x \geq +3, \\ x/6 + 1/2 & \text{otherwise} \end{array} \right.$$

## Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N,\*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_hardsigmoid()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_hardswish

Hardswish module

### **Description**

Applies the hardswish function, element-wise, as described in the paper: Searching for MobileNetV3

## Usage

```
nn_hardswish()
```

#### **Details**

$$\operatorname{Hardswish}(x) = \left\{ \begin{array}{ll} 0 & \text{if } x \leq -3, \\ x & \text{if } x \geq +3, \\ x \cdot (x+3)/6 & \text{otherwise} \end{array} \right.$$

## Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

nn\_hardtanh 225

### **Examples**

```
if (torch_is_installed()) {
## Not run:
m <- nn_hardswish()
input <- torch_randn(2)
output <- m(input)
## End(Not run)
}</pre>
```

nn\_hardtanh

Hardtanh module

### **Description**

Applies the HardTanh function element-wise HardTanh is defined as:

### Usage

```
nn_hardtanh(min_val = -1, max_val = 1, inplace = FALSE)
```

# Arguments

min_val	minimum value of the linear region range. Default: -1
max_val	maximum value of the linear region range. Default: 1
inplace	can optionally do the operation in-place. Default: FALSE

## **Details**

$$\operatorname{HardTanh}(x) = \left\{ \begin{array}{ll} 1 & \text{if } x > 1 \\ -1 & \text{if } x < -1 \\ x & \text{otherwise} \end{array} \right.$$

The range of the linear region :math:[-1, 1] can be adjusted using min\_val and max\_val.

## Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N,\*), same shape as the input

```
if (torch_is_installed()) {
m <- nn_hardtanh(-2, 2)
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_hinge\_embedding\_loss

Hinge embedding loss

### **Description**

Measures the loss given an input tensor x and a labels tensor y (containing 1 or -1).

### Usage

```
nn_hinge_embedding_loss(margin = 1, reduction = "mean")
```

### **Arguments**

margin (float, optional): Has a default value of 1.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

#### **Details**

This is usually used for measuring whether two inputs are similar or dissimilar, e.g. using the L1 pairwise distance as x, and is typically used for learning nonlinear embeddings or semi-supervised learning. The loss function for n-th sample in the mini-batch is

$$l_n = \begin{cases} x_n, & \text{if } y_n = 1, \\ \max\{0, \Delta - x_n\}, & \text{if } y_n = -1, \end{cases}$$

and the total loss functions is

$$\ell(x,y) = \begin{array}{ll} \mathrm{mean}(L), & \mathrm{if\ reduction = 'mean';} \\ \mathrm{sum}(L), & \mathrm{if\ reduction = 'sum'.} \end{array}$$

where  $L = \{l_1, ..., l_N\}^{\top}$ .

# Shape

- Input: (\*) where \* means, any number of dimensions. The sum operation operates over all the elements.
- Target: (\*), same shape as the input
- Output: scalar. If reduction is 'none', then same shape as the input

nn\_identity 227

nn\_identity

Identity module

# Description

A placeholder identity operator that is argument-insensitive.

# Usage

```
nn_identity(...)
```

## **Arguments**

```
... any arguments (unused)
```

# **Examples**

```
if (torch_is_installed()) {
m <- nn_identity(54, unused_argument1 = 0.1, unused_argument2 = FALSE)
input <- torch_randn(128, 20)
output <- m(input)
print(output$size())
}</pre>
```

```
nn_init_calculate_gain
```

Calculate gain

# Description

Return the recommended gain value for the given nonlinearity function.

# Usage

```
nn_init_calculate_gain(nonlinearity, param = NULL)
```

# Arguments

```
nonlinearity the non-linear function
param optional parameter for the non-linear function
```

nn\_init\_dirac\_

nn\_init\_constant\_

Constant initialization

## **Description**

Fills the input Tensor with the value val.

## Usage

```
nn_init_constant_(tensor, val)
```

## **Arguments**

```
tensor an n-dimensional Tensor
val the value to fill the tensor with
```

# **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_constant_(w, 0.3)
}</pre>
```

nn\_init\_dirac\_

Dirac initialization

## **Description**

Fills the {3, 4, 5}-dimensional input Tensor with the Dirac delta function. Preserves the identity of the inputs in Convolutional layers, where as many input channels are preserved as possible. In case of groups>1, each group of channels preserves identity.

## Usage

```
nn_init_dirac_(tensor, groups = 1)
```

## Arguments

```
tensor  a \; \{3,4,5\} \text{-dimensional torch.Tensor}
```

groups (optional) number of groups in the conv layer (default: 1)

nn\_init\_eye\_ 229

### **Examples**

```
if (torch_is_installed()) {
## Not run:
w <- torch_empty(3, 16, 5, 5)
nn_init_dirac_(w)
## End(Not run)
}</pre>
```

nn\_init\_eye\_

Eye initialization

## **Description**

Fills the 2-dimensional input Tensor with the identity matrix. Preserves the identity of the inputs in Linear layers, where as many inputs are preserved as possible.

### Usage

```
nn_init_eye_(tensor)
```

## Arguments

tensor

a 2-dimensional torch tensor.

# **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_eye_(w)
}</pre>
```

nn\_init\_kaiming\_normal\_

Kaiming normal initialization

# Description

Fills the input Tensor with values according to the method described in Delving deep into rectifiers: Surpassing humanite He, K. et al. (2015), using a normal distribution.

### Usage

```
nn_init_kaiming_normal_(
  tensor,
  a = 0,
  mode = "fan_in",
  nonlinearity = "leaky_relu"
)
```

## Arguments

an n-dimensional torch. Tensor

the negative slope of the rectifier used after this layer (only used with 'leaky\_relu')

mode either 'fan\_in' (default) or 'fan\_out'. Choosing 'fan\_in' preserves the magnitude of the variance of the weights in the forward pass. Choosing 'fan\_out' preserves the magnitudes in the backwards pass.

nonlinearity the non-linear function. recommended to use only with 'relu' or 'leaky\_relu' (default).

# Examples

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_kaiming_normal_(w, mode = "fan_in", nonlinearity = "leaky_relu")
}</pre>
```

nn\_init\_kaiming\_uniform\_

Kaiming uniform initialization

# Description

Fills the input Tensor with values according to the method described in Delving deep into rectifiers: Surpassing humanum - He, K. et al. (2015), using a uniform distribution.

## Usage

```
nn_init_kaiming_uniform_(
  tensor,
  a = 0,
  mode = "fan_in",
  nonlinearity = "leaky_relu"
)
```

nn\_init\_normal\_ 231

## **Arguments**

tensor an n-dimensional torch. Tensor

a the negative slope of the rectifier used after this layer (only used with 'leaky\_relu')

mode either 'fan\_in' (default) or 'fan\_out'. Choosing 'fan\_in' preserves the magnitude

of the variance of the weights in the forward pass. Choosing 'fan\_out' preserves

the magnitudes in the backwards pass.

nonlinearity the non-linear function. recommended to use only with 'relu' or 'leaky\_relu'

(default).

### **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_kaiming_uniform_(w, mode = "fan_in", nonlinearity = "leaky_relu")
}</pre>
```

nn\_init\_normal\_

Normal initialization

# Description

Fills the input Tensor with values drawn from the normal distribution

### Usage

```
nn_init_normal_(tensor, mean = 0, std = 1)
```

# **Arguments**

tensor an n-dimensional Tensor

mean the mean of the normal distribution

std the standard deviation of the normal distribution

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_normal_(w)
}</pre>
```

232 nn\_init\_orthogonal\_

nn\_init\_ones\_

Ones initialization

## Description

Fills the input Tensor with the scalar value 1

# Usage

```
nn_init_ones_(tensor)
```

## **Arguments**

tensor

an n-dimensional Tensor

# **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_ones_(w)
}</pre>
```

nn\_init\_orthogonal\_

Orthogonal initialization

# Description

Fills the input Tensor with a (semi) orthogonal matrix, as described in Exact solutions to the nonlinear dynamics of 1 - Saxe, A. et al. (2013). The input tensor must have at least 2 dimensions, and for tensors with more than 2 dimensions the trailing dimensions are flattened.

## Usage

```
nn_init_orthogonal_(tensor, gain = 1)
```

# Arguments

tensor an n-dimensional Tensor gain optional scaling factor

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_orthogonal_(w)
}</pre>
```

nn\_init\_sparse\_ 233

nn\_init\_sparse\_

Sparse initialization

## **Description**

Fills the 2D input Tensor as a sparse matrix, where the non-zero elements will be drawn from the normal distribution as described in Deep learning via Hessian-free optimization - Martens, J. (2010).

# Usage

```
nn_init_sparse_(tensor, sparsity, std = 0.01)
```

## **Arguments**

tensor an n-dimensional Tensor

sparsity The fraction of elements in each column to be set to zero

std the standard deviation of the normal distribution used to generate the non-zero

values

## **Examples**

```
if (torch_is_installed()) {
## Not run:
w <- torch_empty(3, 5)
nn_init_sparse_(w, sparsity = 0.1)
## End(Not run)
}</pre>
```

nn\_init\_trunc\_normal\_ Truncated normal initialization

# Description

Fills the input Tensor with values drawn from a truncated normal distribution.

### Usage

```
nn_init_trunc_normal_(tensor, mean = 0, std = 1, a = -2, b = 2)
```

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## **Arguments**

tensor	an n-dimensional Tensor
mean	the mean of the normal distribution
std	the standard deviation of the normal distribution
а	the minimum cutoff value
b	the maximum cutoff value

# Examples

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_trunc_normal_(w)
}</pre>
```

nn\_init\_uniform\_

Uniform initialization

## **Description**

Fills the input Tensor with values drawn from the uniform distribution

# Usage

```
nn_init_uniform_(tensor, a = 0, b = 1)
```

# Arguments

```
tensor an n-dimensional Tensor

a the lower bound of the uniform distribution

b the upper bound of the uniform distribution
```

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_uniform_(w)
}</pre>
```

nn\_init\_xavier\_normal\_ 235

```
nn_init_xavier_normal_
```

Xavier normal initialization

### **Description**

Fills the input Tensor with values according to the method described in Understanding the difficulty of training deep - Glorot, X. & Bengio, Y. (2010), using a normal distribution.

### Usage

```
nn_init_xavier_normal_(tensor, gain = 1)
```

## **Arguments**

```
tensor an n-dimensional Tensor
gain an optional scaling factor
```

# **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_xavier_normal_(w)
}</pre>
```

```
nn_init_xavier_uniform_
```

Xavier uniform initialization

# Description

Fills the input Tensor with values according to the method described in Understanding the difficulty of training deep - Glorot, X. & Bengio, Y. (2010), using a uniform distribution.

## Usage

```
nn_init_xavier_uniform_(tensor, gain = 1)
```

### **Arguments**

tensor an n-dimensional Tensor gain an optional scaling factor nn\_kl\_div\_loss

### **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_xavier_uniform_(w)
}</pre>
```

nn\_init\_zeros\_

Zeros initialization

### **Description**

Fills the input Tensor with the scalar value 0

### Usage

```
nn_init_zeros_(tensor)
```

### **Arguments**

tensor

an n-dimensional tensor

# **Examples**

```
if (torch_is_installed()) {
w <- torch_empty(3, 5)
nn_init_zeros_(w)
}</pre>
```

nn\_kl\_div\_loss

Kullback-Leibler divergence loss

## **Description**

The Kullback-Leibler divergence loss measure Kullback-Leibler divergence is a useful distance measure for continuous distributions and is often useful when performing direct regression over the space of (discretely sampled) continuous output distributions.

## Usage

```
nn_kl_div_loss(reduction = "mean")
```

# Arguments

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'batchmean' | 'sum' | 'mean'. 'none': no reduction will be applied. 'batchmean': the sum of the output will be divided by batchsize. 'sum': the output will be summed. 'mean': the output will be divided by the number of elements in the output. Default: 'mean'

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#### **Details**

As with nn\_nll\_loss(), the input given is expected to contain *log-probabilities* and is not restricted to a 2D Tensor.

The targets are interpreted as *probabilities* by default, but could be considered as *log-probabilities* with log\_target set to TRUE.

This criterion expects a target Tensor of the same size as the input Tensor.

The unreduced (i.e. with reduction set to 'none') loss can be described as:

$$l(x,y) = L = \{l_1, \dots, l_N\}, \quad l_n = y_n \cdot (\log y_n - x_n)$$

where the index N spans all dimensions of input and L has the same shape as input. If reduction is not 'none' (default 'mean'), then:

$$\ell(x,y) = \begin{array}{ll} \operatorname{mean}(L), & \text{if reduction} = \text{'mean'}; \\ \operatorname{sum}(L), & \text{if reduction} = \text{'sum'}. \end{array}$$

In default reduction mode 'mean', the losses are averaged for each minibatch over observations as well as over dimensions. 'batchmean' mode gives the correct KL divergence where losses are averaged over batch dimension only. 'mean' mode's behavior will be changed to the same as 'batchmean' in the next major release.

### Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input
- Output: scalar by default. If reduction is 'none', then (N,\*), the same shape as the input

#### Note

reduction = 'mean' doesn't return the true kl divergence value, please use reduction = 'batchmean' which aligns with KL math definition. In the next major release, 'mean' will be changed to be the same as 'batchmean'.

nn\_l1\_loss

L1 loss

## **Description**

Creates a criterion that measures the mean absolute error (MAE) between each element in the input x and target y.

## Usage

```
nn_l1_loss(reduction = "mean")
```

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### **Arguments**

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

### **Details**

The unreduced (i.e. with reduction set to 'none') loss can be described as:

$$\ell(x,y) = L = \{l_1, \dots, l_N\}^{\top}, \quad l_n = |x_n - y_n|,$$

where N is the batch size. If reduction is not 'none' (default 'mean'), then:

$$\ell(x,y) = \begin{array}{ll} \mathrm{mean}(L), & \mathrm{if\ reduction = 'mean';} \\ \mathrm{sum}(L), & \mathrm{if\ reduction = 'sum'.} \end{array}$$

x and y are tensors of arbitrary shapes with a total of n elements each.

The sum operation still operates over all the elements, and divides by n. The division by n can be avoided if one sets reduction = 'sum'.

## Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input
- Output: scalar. If reduction is 'none', then (N,\*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
loss <- nn_l1_loss()
input <- torch_randn(3, 5, requires_grad = TRUE)
target <- torch_randn(3, 5)
output <- loss(input, target)
output$backward()
}</pre>
```

nn\_layer\_norm

Layer normalization

### **Description**

Applies Layer Normalization over a mini-batch of inputs as described in the paper Layer Normalization

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### Usage

```
nn_layer_norm(normalized_shape, eps = 1e-05, elementwise_affine = TRUE)
```

### **Arguments**

normalized\_shape

(int or list): input shape from an expected input of size  $[*\times normalized\_shape[0] \times normalized\_shape[1] \times \ldots \times normalized\_shape[-1]]$  If a single integer is used, it is treated as a singleton list, and this module will normalize over the last dimension which is expected to be of that specific size.

eps

a value added to the denominator for numerical stability. Default: 1e-5

elementwise\_affine

a boolean value that when set to TRUE, this module has learnable per-element affine parameters initialized to ones (for weights) and zeros (for biases). Default: TRUE.

#### **Details**

$$y = \frac{x - \mathbf{E}[x]}{\sqrt{\mathbf{Var}[x] + \epsilon}} * \gamma + \beta$$

The mean and standard-deviation are calculated separately over the last certain number dimensions which have to be of the shape specified by normalized\_shape.

 $\gamma$  and  $\beta$  are learnable affine transform parameters of normalized\_shape if elementwise\_affine is TRUE.

The standard-deviation is calculated via the biased estimator, equivalent to torch\_var(input, unbiased=FALSE).

### Shape

• Input: (N,\*)

• Output: (N, \*) (same shape as input)

#### Note

Unlike Batch Normalization and Instance Normalization, which applies scalar scale and bias for each entire channel/plane with the affine option, Layer Normalization applies per-element scale and bias with elementwise\_affine.

This layer uses statistics computed from input data in both training and evaluation modes.

```
if (torch_is_installed()) {
input <- torch_randn(20, 5, 10, 10)
# With Learnable Parameters
m <- nn_layer_norm(input$size()[-1])</pre>
```

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```
# Without Learnable Parameters
m <- nn_layer_norm(input$size()[-1], elementwise_affine = FALSE)
# Normalize over last two dimensions
m <- nn_layer_norm(c(10, 10))
# Normalize over last dimension of size 10
m <- nn_layer_norm(10)
# Activating the module
output <- m(input)
}</pre>
```

nn\_leaky\_relu

LeakyReLU module

### **Description**

Applies the element-wise function:

## Usage

```
nn_leaky_relu(negative_slope = 0.01, inplace = FALSE)
```

## **Arguments**

```
negative_slope Controls the angle of the negative slope. Default: 1e-2 inplace can optionally do the operation in-place. Default: FALSE
```

# **Details**

$$LeakyReLU(x) = \max(0, x) + negative\_slope * \min(0, x)$$

or

$$\mbox{LeakyRELU}(x) = \left\{ \begin{array}{ll} x, & \mbox{if } x \geq 0 \\ \mbox{negative\_slope} \times x, & \mbox{otherwise} \end{array} \right.$$

### Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

```
if (torch_is_installed()) {
m <- nn_leaky_relu(0.1)
input <- torch_randn(2)
output <- m(input)
}</pre>
```

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nn\_linear

Linear module

## Description

Applies a linear transformation to the incoming data:  $y = xA^T + b$ 

## Usage

```
nn_linear(in_features, out_features, bias = TRUE)
```

### **Arguments**

```
in_features size of each input sample
out_features size of each output sample
bias If set to FALSE, the layer will not learn an additive bias. Default: TRUE
```

## Shape

- Input: (N, \*, H\_in) where \* means any number of additional dimensions and H\_in = in\_features.
- Output: (N, \*, H\_out) where all but the last dimension are the same shape as the input and :math:H\_out = out\_features.

### Attributes

- weight: the learnable weights of the module of shape (out\_features, in\_features). The values are initialized from  $U(-\sqrt{k},\sqrt{k})$ s, where  $k=\frac{1}{\text{in features}}$
- bias: the learnable bias of the module of shape (out\_features). If bias is TRUE, the values are initialized from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{1}{\text{in_features}}$

```
if (torch_is_installed()) {
m <- nn_linear(20, 30)
input <- torch_randn(128, 20)
output <- m(input)
print(output$size())
}</pre>
```

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nn\_log\_sigmoid

LogSigmoid module

## **Description**

Applies the element-wise function:

$$\mathsf{LogSigmoid}(x) = \log\left(\frac{1}{1 + \exp(-x)}\right)$$

## Usage

```
nn_log_sigmoid()
```

### Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_log_sigmoid()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_log\_softmax

LogSoftmax module

## **Description**

Applies the  $\log(\operatorname{Softmax}(x))$  function to an n-dimensional input Tensor. The LogSoftmax formulation can be simplified as:

# Usage

```
nn_log_softmax(dim)
```

## **Arguments**

dim

(int): A dimension along which LogSoftmax will be computed.

## **Details**

$$\operatorname{LogSoftmax}(x_i) = \log \left( \frac{\exp(x_i)}{\sum_j \exp(x_j)} \right)$$

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## Value

a Tensor of the same dimension and shape as the input with values in the range [-inf, 0)

### Shape

- Input: (\*) where \* means, any number of additional dimensions
- Output: (\*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_log_softmax(1)
input <- torch_randn(2, 3)
output <- m(input)
}</pre>
```

nn\_lp\_pool1d

Applies a 1D power-average pooling over an input signal composed of several input planes.

## **Description**

On each window, the function computed is:

## Usage

```
nn_lp_pool1d(norm_type, kernel_size, stride = NULL, ceil_mode = FALSE)
```

### **Arguments**

 $norm\_type$  if inf than one gets max pooling if 0 you get sum pooling (proportional to the

avg pooling)

kernel\_size a single int, the size of the window

stride a single int, the stride of the window. Default value is kernel\_size

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape

#### **Details**

$$f(X) = \sqrt[p]{\sum_{x \in X} x^p}$$

- At  $p = \infty$ , one gets Max Pooling
- At p = 1, one gets Sum Pooling (which is proportional to Average Pooling)

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### **Shape**

• Input:  $(N, C, L_{in})$ 

• Output:  $(N, C, L_{out})$ , where

$$L_{out} = \left\lfloor \frac{L_{in} - \text{kernel\_size}}{\text{stride}} + 1 \right\rfloor$$

### Note

If the sum to the power of p is zero, the gradient of this function is not defined. This implementation will set the gradient to zero in this case.

## **Examples**

```
if (torch_is_installed()) {
# power-2 pool of window of length 3, with stride 2.
m <- nn_lp_pool1d(2, 3, stride = 2)
input <- torch_randn(20, 16, 50)
output <- m(input)
}</pre>
```

nn\_lp\_pool2d

Applies a 2D power-average pooling over an input signal composed of several input planes.

## Description

On each window, the function computed is:

### Usage

```
nn_lp_pool2d(norm_type, kernel_size, stride = NULL, ceil_mode = FALSE)
```

### **Arguments**

norm\_type if inf than one gets max pooling if 0 you get sum pooling (proportional to the

avg pooling)

kernel\_size the size of the window

stride the stride of the window. Default value is kernel\_size

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape

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### **Details**

$$f(X) = \sqrt[p]{\sum_{x \in X} x^p}$$

- At  $p = \infty$ , one gets Max Pooling
- At p = 1, one gets Sum Pooling (which is proportional to average pooling)

The parameters kernel\_size, stride can either be:

- a single int in which case the same value is used for the height and width dimension
- a tuple of two ints in which case, the first int is used for the height dimension, and the second int for the width dimension

## Shape

• Input:  $(N, C, H_{in}, W_{in})$ 

• Output:  $(N, C, H_{out}, W_{out})$ , where

$$H_{out} = \left\lfloor \frac{H_{in} - \text{kernel\_size}[0]}{\text{stride}[0]} + 1 \right\rfloor$$

$$W_{out} = \left\lfloor \frac{W_{in} - \text{kernel\_size}[1]}{\text{stride}[1]} + 1 \right\rfloor$$

### Note

If the sum to the power of p is zero, the gradient of this function is not defined. This implementation will set the gradient to zero in this case.

```
if (torch_is_installed()) {

# power-2 pool of square window of size=3, stride=2
m <- nn_lp_pool2d(2, 3, stride = 2)
# pool of non-square window of power 1.2
m <- nn_lp_pool2d(1.2, c(3, 2), stride = c(2, 1))
input <- torch_randn(20, 16, 50, 32)
output <- m(input)
}</pre>
```

246 nn\_lstm

nn_lstm	Applies a multi-layer long short-term memory (LSTM) RNN to an input sequence.
	pui sequence.

# Description

For each element in the input sequence, each layer computes the following function:

# Usage

```
nn_lstm(
  input_size,
  hidden_size,
  num_layers = 1,
  bias = TRUE,
  batch_first = FALSE,
  dropout = 0,
  bidirectional = FALSE,
  ...
)
```

# Arguments

input_size	The number of expected features in the input x
hidden_size	The number of features in the hidden state h
num_layers	Number of recurrent layers. E.g., setting num_layers=2 would mean stacking two LSTMs together to form a stacked LSTM, with the second LSTM taking in outputs of the first LSTM and computing the final results. Default: 1
bias	If FALSE, then the layer does not use bias weights b_ih and b_hh. Default: TRUE
batch_first	If TRUE, then the input and output tensors are provided as (batch, seq, feature). Default: FALSE
dropout	If non-zero, introduces a Dropout layer on the outputs of each LSTM layer except the last layer, with dropout probability equal to dropout. Default: 0
bidirectional	If TRUE, becomes a bidirectional LSTM. Default: FALSE
	currently unused.

# **Details**

```
\begin{split} i_t &= \sigma(W_{ii}x_t + b_{ii} + W_{hi}h_{(t-1)} + b_{hi}) \\ f_t &= \sigma(W_{if}x_t + b_{if} + W_{hf}h_{(t-1)} + b_{hf}) \\ g_t &= \tanh(W_{ig}x_t + b_{ig} + W_{hg}h_{(t-1)} + b_{hg}) \\ o_t &= \sigma(W_{io}x_t + b_{io} + W_{ho}h_{(t-1)} + b_{ho}) \\ c_t &= f_t c_{(t-1)} + i_t g_t \\ h_t &= o_t \tanh(c_t) \end{split}
```

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where  $h_t$  is the hidden state at time t,  $c_t$  is the cell state at time t,  $x_t$  is the input at time t,  $h_{(t-1)}$  is the hidden state of the previous layer at time t-1 or the initial hidden state at time 0, and  $i_t$ ,  $f_t$ ,  $g_t$ ,  $o_t$  are the input, forget, cell, and output gates, respectively.  $\sigma$  is the sigmoid function.

### **Inputs**

Inputs: input, (h\_0, c\_0)

- input of shape (seq\_len, batch, input\_size): tensor containing the features of the input sequence. The input can also be a packed variable length sequence. See nn\_utils\_rnn\_pack\_padded\_sequence() or nn\_utils\_rnn\_pack\_sequence() for details.
- h\_0 of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the initial hidden state for each element in the batch.
- $c_0$  of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the initial cell state for each element in the batch.

If  $(h_0, c_0)$  is not provided, both  $h_0$  and  $c_0$  default to zero.

### **Outputs**

Outputs: output, (h\_n, c\_n)

- output of shape (seq\_len, batch, num\_directions \* hidden\_size): tensor containing the output features (h\_t) from the last layer of the LSTM, for each t. If a torch\_nn.utils.rnn.PackedSequence has been given as the input, the output will also be a packed sequence. For the unpacked case, the directions can be separated using output\$view(c(seq\_len, batch, num\_directions, hidden\_size)), with forward and backward being direction 0 and 1 respectively. Similarly, the directions can be separated in the packed case.
- h\_n of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the hidden state for t = seq\_len. Like *output*, the layers can be separated using h\_n\$view(c(num\_layers, num\_directions, batch, hidden\_size)) and similarly for c\_n.
- **c\_n** (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the cell state for t = seq\_len

#### **Attributes**

- weight\_ih\_1[k]: the learnable input-hidden weights of the k<sup>th</sup> layer (W\_ii|W\_if|W\_ig|W\_io), of shape (4\*hidden\_size x input\_size)
- weight\_hh\_1[k]: the learnable hidden-hidden weights of the k<sup>th</sup> layer (W\_hi|W\_hf|W\_hg|W\_ho), of shape (4\*hidden\_size x hidden\_size)
- bias\_ih\_l[k]: the learnable input-hidden bias of the k<sup>th</sup> layer (b\_ii|b\_if|b\_ig|b\_io), of shape (4\*hidden\_size)
- bias\_hh\_1[k]: the learnable hidden-hidden bias of the k<sup>th</sup> layer (b\_hi|b\_hf|b\_hg|b\_ho), of shape (4\*hidden\_size)

#### Note

All the weights and biases are initialized from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{1}{\text{hidden size}}$ 

### **Examples**

```
if (torch_is_installed()) {
  rnn <- nn_lstm(10, 20, 2)
  input <- torch_randn(5, 3, 10)
  h0 <- torch_randn(2, 3, 20)
  c0 <- torch_randn(2, 3, 20)
  output <- rnn(input, list(h0, c0))
}</pre>
```

nn\_margin\_ranking\_loss

Margin ranking loss

## Description

Creates a criterion that measures the loss given inputs x1, x2, two 1D mini-batch Tensors, and a label 1D mini-batch tensor y (containing 1 or -1). If y=1 then it assumed the first input should be ranked higher (have a larger value) than the second input, and vice-versa for y=-1.

### Usage

```
nn_margin_ranking_loss(margin = 0, reduction = "mean")
```

### **Arguments**

margin (float, optional): Has a default value of 0.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

## **Details**

The loss function for each pair of samples in the mini-batch is:

$$loss(x1, x2, y) = max(0, -y * (x1 - x2) + margin)$$

### Shape

- Input1: (N) where N is the batch size.
- Input2: (N), same shape as the Input1.
- Target: (N), same shape as the inputs.
- Output: scalar. If reduction is 'none', then (N).

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### **Examples**

```
if (torch_is_installed()) {
loss <- nn_margin_ranking_loss()
input1 <- torch_randn(3, requires_grad = TRUE)
input2 <- torch_randn(3, requires_grad = TRUE)
target <- torch_randn(3)$sign()
output <- loss(input1, input2, target)
output$backward()
}</pre>
```

nn\_max\_pool1d

MaxPool1D module

#### **Description**

Applies a 1D max pooling over an input signal composed of several input planes.

### Usage

```
nn_max_pool1d(
  kernel_size,
  stride = NULL,
  padding = 0,
  dilation = 1,
  return_indices = FALSE,
  ceil_mode = FALSE
)
```

## Arguments

kernel\_size the size of the window to take a max over

stride the stride of the window. Default value is kernel\_size

padding implicit zero padding to be added on both sides

dilation a parameter that controls the stride of elements in the window

return\_indices if TRUE, will return the max indices along with the outputs. Useful for nn\_max\_unpool1d()

later.

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape

#### **Details**

In the simplest case, the output value of the layer with input size (N,C,L) and output  $(N,C,L_{out})$  can be precisely described as:

$$out(N_i, C_j, k) = \max_{m = 0, \dots, \text{kernel\_size-}1} input(N_i, C_j, stride \times k + m)$$

If padding is non-zero, then the input is implicitly zero-padded on both sides for padding number of points. dilation controls the spacing between the kernel points. It is harder to describe, but this link has a nice visualization of what dilation does.

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### **Shape**

```
• Input: (N, C, L_{in})
```

• Output:  $(N, C, L_{out})$ , where

$$L_{out} = \left\lfloor \frac{L_{in} + 2 \times \text{padding} - \text{dilation} \times (\text{kernel\_size} - 1) - 1}{\text{stride}} + 1 \right\rfloor$$

## **Examples**

```
if (torch_is_installed()) {
# pool of size=3, stride=2
m <- nn_max_pool1d(3, stride = 2)
input <- torch_randn(20, 16, 50)
output <- m(input)
}</pre>
```

nn\_max\_pool2d

MaxPool2D module

## **Description**

Applies a 2D max pooling over an input signal composed of several input planes.

### Usage

```
nn_max_pool2d(
  kernel_size,
  stride = NULL,
  padding = 0,
  dilation = 1,
  return_indices = FALSE,
  ceil_mode = FALSE
)
```

## **Arguments**

kernel\_size the size of the window to take a max over

stride the stride of the window. Default value is kernel\_size

padding implicit zero padding to be added on both sides

dilation a parameter that controls the stride of elements in the window

return\_indices if TRUE, will return the max indices along with the outputs. Useful for nn\_max\_unpool2d()

later.

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape

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### **Details**

In the simplest case, the output value of the layer with input size (N, C, H, W), output  $(N, C, H_{out}, W_{out})$  and kernel\_size (kH, kW) can be precisely described as:

$$\begin{aligned} out(N_i, C_j, h, w) = & & \max_{m=0, \dots, kH-1} \max_{n=0, \dots, kW-1} \\ & & & \text{input}(N_i, C_j, \text{stride}[0] \times h + m, \text{stride}[1] \times w + n) \end{aligned}$$

If padding is non-zero, then the input is implicitly zero-padded on both sides for padding number of points. dilation controls the spacing between the kernel points. It is harder to describe, but this link has a nice visualization of what dilation does.

The parameters kernel\_size, stride, padding, dilation can either be:

- a single int in which case the same value is used for the height and width dimension
- a tuple of two ints in which case, the first int is used for the height dimension, and the second int for the width dimension

### Shape

- Input:  $(N, C, H_{in}, W_{in})$
- Output:  $(N, C, H_{out}, W_{out})$ , where

$$H_{out} = \left\lfloor \frac{H_{in} + 2*\operatorname{padding}[0] - \operatorname{dilation}[0] \times (\operatorname{kernel\_size}[0] - 1) - 1}{\operatorname{stride}[0]} + 1 \right\rfloor$$

$$W_{out} = \left\lfloor \frac{W_{in} + 2 * \texttt{padding[1]} - \texttt{dilation[1]} \times (\texttt{kernel\_size[1]} - 1) - 1}{\texttt{stride[1]}} + 1 \right\rfloor$$

#### **Examples**

```
if (torch_is_installed()) {
# pool of square window of size=3, stride=2
m <- nn_max_pool2d(3, stride = 2)
# pool of non-square window
m <- nn_max_pool2d(c(3, 2), stride = c(2, 1))
input <- torch_randn(20, 16, 50, 32)
output <- m(input)
}</pre>
```

nn\_max\_pool3d

Applies a 3D max pooling over an input signal composed of several input planes.

#### **Description**

In the simplest case, the output value of the layer with input size (N, C, D, H, W), output  $(N, C, D_{out}, H_{out}, W_{out})$  and kernel\_size (kD, kH, kW) can be precisely described as:

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### Usage

```
nn_max_pool3d(
  kernel_size,
  stride = NULL,
  padding = 0,
  dilation = 1,
  return_indices = FALSE,
  ceil_mode = FALSE
)
```

#### **Arguments**

kernel\_size the size of the window to take a max over
stride the stride of the window. Default value is kernel\_size
padding implicit zero padding to be added on all three sides
dilation a parameter that controls the stride of elements in the window

return indices if TRUE will return the max indices along with the outputs. Useful for torch on Max/Inpoel 3

return\_indices if TRUE, will return the max indices along with the outputs. Useful for torch\_nn.MaxUnpool3d

later

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape

#### **Details**

```
\begin{aligned} \text{out}(N_i,C_j,d,h,w) = & & \max_{k=0,\dots,kD-1} \max_{m=0,\dots,kH-1} \max_{n=0,\dots,kW-1} \\ & & \text{input}(N_i,C_j,\text{stride}[0] \times d + k,\text{stride}[1] \times h + m,\text{stride}[2] \times w + n) \end{aligned}
```

If padding is non-zero, then the input is implicitly zero-padded on both sides for padding number of points. dilation controls the spacing between the kernel points. It is harder to describe, but this link\_has a nice visualization of what dilation does. The parameters kernel\_size, stride, padding, dilation can either be:

- a single int in which case the same value is used for the depth, height and width dimension
- a tuple of three ints in which case, the first int is used for the depth dimension, the second int for the height dimension and the third int for the width dimension

#### Shape

- Input:  $(N, C, D_{in}, H_{in}, W_{in})$
- Output:  $(N, C, D_{out}, H_{out}, W_{out})$ , where

$$D_{out} = \left\lfloor \frac{D_{in} + 2 \times \operatorname{padding}[0] - \operatorname{dilation}[0] \times (\operatorname{kernel\_size}[0] - 1) - 1}{\operatorname{stride}[0]} + 1 \right\rfloor$$

$$H_{out} = \left\lfloor \frac{H_{in} + 2 \times \text{padding}[1] - \text{dilation}[1] \times (\text{kernel\_size}[1] - 1) - 1}{\text{stride}[1]} + 1 \right\rfloor$$

$$W_{out} = \left\lfloor \frac{W_{in} + 2 \times \operatorname{padding}[2] - \operatorname{dilation}[2] \times (\operatorname{kernel\_size}[2] - 1) - 1}{\operatorname{stride}[2]} + 1 \right\rfloor$$

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### **Examples**

```
if (torch_is_installed()) {
# pool of square window of size=3, stride=2
m <- nn_max_pool3d(3, stride = 2)
# pool of non-square window
m <- nn_max_pool3d(c(3, 2, 2), stride = c(2, 1, 2))
input <- torch_randn(20, 16, 50, 44, 31)
output <- m(input)
}</pre>
```

nn\_max\_unpool1d

Computes a partial inverse of MaxPool1d.

## **Description**

MaxPool1d is not fully invertible, since the non-maximal values are lost. MaxUnpool1d takes in as input the output of MaxPool1d including the indices of the maximal values and computes a partial inverse in which all non-maximal values are set to zero.

#### Usage

```
nn_max_unpool1d(kernel_size, stride = NULL, padding = 0)
```

## **Arguments**

kernel\_size (int or tuple): Size of the max pooling window.

stride (int or tuple): Stride of the max pooling window. It is set to kernel\_size by

default.

padding (int or tuple): Padding that was added to the input

## **Inputs**

- input: the input Tensor to invert
- indices: the indices given out by nn\_max\_pool1d()
- output\_size (optional): the targeted output size

### Shape

- Input:  $(N, C, H_{in})$
- Output:  $(N, C, H_{out})$ , where

$$H_{out} = (H_{in} - 1) \times \text{stride}[0] - 2 \times \text{padding}[0] + \text{kernel\_size}[0]$$

or as given by output\_size in the call operator

254 nn\_max\_unpool2d

#### Note

MaxPool1d can map several input sizes to the same output sizes. Hence, the inversion process can get ambiguous. To accommodate this, you can provide the needed output size as an additional argument output\_size in the forward call. See the Inputs and Example below.

### **Examples**

```
if (torch_is_installed()) {
pool <- nn_max_pool1d(2, stride = 2, return_indices = TRUE)
unpool <- nn_max_unpool1d(2, stride = 2)

input <- torch_tensor(array(1:8 / 1, dim = c(1, 1, 8)))
out <- pool(input)
unpool(out[[1]], out[[2]])

# Example showcasing the use of output_size
input <- torch_tensor(array(1:8 / 1, dim = c(1, 1, 8)))
out <- pool(input)
unpool(out[[1]], out[[2]], output_size = input$size())
unpool(out[[1]], out[[2]])
}</pre>
```

nn\_max\_unpool2d

Computes a partial inverse of MaxPool2d.

# Description

MaxPool2d is not fully invertible, since the non-maximal values are lost. MaxUnpool2d takes in as input the output of MaxPool2d including the indices of the maximal values and computes a partial inverse in which all non-maximal values are set to zero.

### Usage

```
nn_max_unpool2d(kernel_size, stride = NULL, padding = 0)
```

### **Arguments**

```
kernel_size (int or tuple): Size of the max pooling window.

stride (int or tuple): Stride of the max pooling window. It is set to kernel_size by default.

padding (int or tuple): Padding that was added to the input
```

### **Inputs**

- input: the input Tensor to invert
- indices: the indices given out by nn\_max\_pool2d()
- output\_size (optional): the targeted output size

nn\_max\_unpool3d 255

### Shape

```
• Input: (N,C,H_{in},W_{in})

• Output: (N,C,H_{out},W_{out}), where H_{out}=(H_{in}-1)\times \text{stride}[0]-2\times \text{padding}[0]+\text{kernel\_size}[0] W_{out}=(W_{in}-1)\times \text{stride}[1]-2\times \text{padding}[1]+\text{kernel\_size}[1] or as given by output_size in the call operator
```

#### Note

MaxPool2d can map several input sizes to the same output sizes. Hence, the inversion process can get ambiguous. To accommodate this, you can provide the needed output size as an additional argument output\_size in the forward call. See the Inputs and Example below.

## **Examples**

```
if (torch_is_installed()) {
pool <- nn_max_pool2d(2, stride = 2, return_indices = TRUE)
unpool <- nn_max_unpool2d(2, stride = 2)
input <- torch_randn(1, 1, 4, 4)
out <- pool(input)
unpool(out[[1]], out[[2]])

# specify a different output size than input size
unpool(out[[1]], out[[2]], output_size = c(1, 1, 5, 5))
}</pre>
```

nn\_max\_unpool3d

Computes a partial inverse of MaxPool3d.

# Description

MaxPool3d is not fully invertible, since the non-maximal values are lost. MaxUnpool3d takes in as input the output of MaxPool3d including the indices of the maximal values and computes a partial inverse in which all non-maximal values are set to zero.

# Usage

```
nn_max_unpool3d(kernel_size, stride = NULL, padding = 0)
```

## **Arguments**

```
kernel_size (int or tuple): Size of the max pooling window.

stride (int or tuple): Stride of the max pooling window. It is set to kernel_size by default.

padding (int or tuple): Padding that was added to the input
```

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## **Inputs**

- input: the input Tensor to invert
- indices: the indices given out by nn\_max\_pool3d()
- output\_size (optional): the targeted output size

### Shape

```
• Input: (N, C, D_{in}, H_{in}, W_{in})
```

• Output:  $(N, C, D_{out}, H_{out}, W_{out})$ , where

$$\begin{split} D_{out} &= (D_{in} - 1) \times \text{stride}[0] - 2 \times \text{padding}[0] + \text{kernel\_size}[0] \\ H_{out} &= (H_{in} - 1) \times \text{stride}[1] - 2 \times \text{padding}[1] + \text{kernel\_size}[1] \\ W_{out} &= (W_{in} - 1) \times \text{stride}[2] - 2 \times \text{padding}[2] + \text{kernel\_size}[2] \end{split}$$

or as given by output\_size in the call operator

#### Note

MaxPool3d can map several input sizes to the same output sizes. Hence, the inversion process can get ambiguous. To accommodate this, you can provide the needed output size as an additional argument output\_size in the forward call. See the Inputs section below.

## **Examples**

```
if (torch_is_installed()) {
# pool of square window of size=3, stride=2
pool <- nn_max_pool3d(3, stride = 2, return_indices = TRUE)
unpool <- nn_max_unpool3d(3, stride = 2)
out <- pool(torch_randn(20, 16, 51, 33, 15))
unpooled_output <- unpool(out[[1]], out[[2]])
unpooled_output$size()
}</pre>
```

nn\_module

Base class for all neural network modules.

#### **Description**

Your models should also subclass this class.

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### Usage

```
nn_module(
  classname = NULL,
  inherit = nn_Module,
    ...,
  private = NULL,
  active = NULL,
  parent_env = parent.frame()
)
```

#### **Arguments**

```
classname an optional name for the module inherit an optional module to inherit from ... methods implementation private passed to R6::R6Class(). active passed to R6::R6Class(). parent_env passed to R6::R6Class().
```

#### **Details**

Modules can also contain other Modules, allowing to nest them in a tree structure. You can assign the submodules as regular attributes.

You are expected to implement the initialize and the forward to create a new nn\_module.

#### **Initialize**

The initialize function will be called whenever a new instance of the nn\_module is created. We use the initialize functions to define submodules and parameters of the module. For example:

```
initialize = function(input_size, output_size) {
   self$conv1 <- nn_conv2d(input_size, output_size, 5)
   self$conv2 <- nn_conv2d(output_size, output_size, 5)
}</pre>
```

The initialize function can have any number of parameters. All objects assigned to self\$ will be available for other methods that you implement. Tensors wrapped with nn\_parameter() or nn\_buffer() and submodules are automatically tracked when assigned to self\$.

The initialize function is optional if the module you are defining doesn't have weights, submodules or buffers.

### **Forward**

The forward method is called whenever an instance of nn\_module is called. This is usually used to implement the computation that the module does with the weights ad submodules defined in the initialize function.

For example:

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```
forward = function(input) {
   input <- self$conv1(input)
   input <- nnf_relu(input)
   input <- self$conv2(input)
   input <- nnf_relu(input)
   input
}</pre>
```

The forward function can use the self\$training attribute to make different computations depending wether the model is training or not, for example if you were implementing the dropout module.

### **Cloning**

To finalize the cloning of a module, you can define a private finalize\_deep\_clone() method. This method is called on the cloned object when deep-cloning a module, after all the modules, parameters and buffers were already cloned.

## **Examples**

```
if (torch_is_installed()) {
model <- nn_module(
   initialize = function() {
    self$conv1 <- nn_conv2d(1, 20, 5)
    self$conv2 <- nn_conv2d(20, 20, 5)
},
forward = function(input) {
   input <- self$conv1(input)
   input <- nnf_relu(input)
   input <- self$conv2(input)
   input <- nnf_relu(input)
   input <- nnf_relu(input)
   input <- nnf_relu(input)
   input
}
</pre>
```

nn\_module\_dict

Container that allows named values

### **Description**

Container that allows named values

# Usage

```
nn_module_dict(dict)
```

### **Arguments**

dict

A named list of submodules that will be saved in that module.

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### See Also

```
nn_module_list()
```

## **Examples**

```
if (torch_is_installed()) {
nn_module <- nn_module(
  initialize = function() {
    self$dict <- nn_module_dict(list(
         l1 = nn_linear(10, 20),
         l2 = nn_linear(20, 10)
    ))
    },
    forward = function(x) {
        x <- self$dict$l1(x)
        self$dict$l2(x)
    }
)
</pre>
```

nn\_module\_list

Holds submodules in a list.

## **Description**

nn\_module\_list can be indexed like a regular R list, but modules it contains are properly registered, and will be visible by all nn\_module methods.

## Usage

```
nn_module_list(modules = list())
```

### **Arguments**

modules

a list of modules to add

### See Also

```
nn_module_dict()
```

# **Examples**

```
if (torch_is_installed()) {

my_module <- nn_module(
  initialize = function() {
    self$linears <- nn_module_list(lapply(1:10, function(x) nn_linear(10, 10)))
  },
  forward = function(x) {</pre>
```

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```
for (i in 1:length(self$linears)) {
    x <- self$linears[[i]](x)
    }
    x
}
</pre>
```

nn\_mse\_loss

MSE loss

### **Description**

Creates a criterion that measures the mean squared error (squared L2 norm) between each element in the input x and target y. The unreduced (i.e. with reduction set to 'none') loss can be described as:

### Usage

```
nn_mse_loss(reduction = "mean")
```

### **Arguments**

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

### **Details**

$$\ell(x,y) = L = \{l_1, \dots, l_N\}^{\top}, \quad l_n = (x_n - y_n)^2,$$

where N is the batch size. If reduction is not 'none' (default 'mean'), then:

$$\ell(x,y) = \begin{array}{ll} \operatorname{mean}(L), & \text{if reduction} = \text{'mean';} \\ \operatorname{sum}(L), & \text{if reduction} = \text{'sum'.} \end{array}$$

x and y are tensors of arbitrary shapes with a total of n elements each.

The mean operation still operates over all the elements, and divides by n. The division by n can be avoided if one sets reduction = 'sum'.

### Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input

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### **Examples**

```
if (torch_is_installed()) {
loss <- nn_mse_loss()
input <- torch_randn(3, 5, requires_grad = TRUE)
target <- torch_randn(3, 5)
output <- loss(input, target)
output$backward()
}</pre>
```

nn\_multihead\_attention

MultiHead attention

### **Description**

Allows the model to jointly attend to information from different representation subspaces. See reference: Attention Is All You Need

### Usage

```
nn_multihead_attention(
  embed_dim,
  num_heads,
  dropout = 0,
  bias = TRUE,
  add_bias_kv = FALSE,
  add_zero_attn = FALSE,
  kdim = NULL,
  vdim = NULL,
  batch_first = FALSE
)
```

### **Arguments**

embed\_dim total dimension of the model.

num\_heads parallel attention heads. Note that embed\_dim will be split across num\_heads

(i.e. each head will have dimension embed\_dim %/% num\_heads).

dropout a Dropout layer on attn\_output\_weights. Default: 0.0.

bias add bias as module parameter. Default: True.

add\_bias\_kv add bias to the key and value sequences at dim=0.

add\_zero\_attn add a new batch of zeros to the key and value sequences at dim=1.

kdim total number of features in key. Default: NULL

vdim total number of features in value. Default: NULL. Note: if kdim and vdim are

NULL, they will be set to embed\_dim such that query, key, and value have the

same number of features.

batch\_first

if TRUE then the input and output tensors are (N, S, E) instead of (S, N, E), where N is the batch size, S is the sequence length, and E is the embedding dimension.

#### **Details**

```
MultiHead(Q, K, V) = Concat(head_1, ..., head_h)W^O where head_i = Attention(QW_i^Q, KW_i^K, VW_i^V)
```

## Shape

## Inputs:

- query: (L, N, E) where L is the target sequence length, N is the batch size, E is the embedding dimension. (but see the batch\_first argument)
- key: (S, N, E), where S is the source sequence length, N is the batch size, E is the embedding dimension. (but see the batch\_first argument)
- value: (S, N, E) where S is the source sequence length, N is the batch size, E is the embedding dimension. (but see the batch\_first argument)
- key\_padding\_mask: (N, S) where N is the batch size, S is the source sequence length. If a ByteTensor is provided, the non-zero positions will be ignored while the position with the zero positions will be unchanged. If a BoolTensor is provided, the positions with the value of True will be ignored while the position with the value of False will be unchanged.
- attn\_mask: 2D mask (L,S) where L is the target sequence length, S is the source sequence length. 3D mask  $(N*num_heads,L,S)$  where N is the batch size, L is the target sequence length, S is the source sequence length. attn\_mask ensure that position i is allowed to attend the unmasked positions. If a ByteTensor is provided, the non-zero positions are not allowed to attend while the zero positions will be unchanged. If a BoolTensor is provided, positions with True are not allowed to attend while False values will be unchanged. If a FloatTensor is provided, it will be added to the attention weight.

### Outputs:

- attn\_output: (L, N, E) where L is the target sequence length, N is the batch size, E is the embedding dimension. (but see the batch\_first argument)
- attn\_output\_weights:
  - if avg\_weights is TRUE (the default), the output attention weights are averaged over the attention heads, giving a tensor of shape (N,L,S) where N is the batch size, L is the target sequence length, S is the source sequence length.
  - if avg\_weights is FALSE, the attention weight tensor is output as-is, with shape (N, H, L, S), where H is the number of attention heads.

### **Examples**

```
if (torch_is_installed()) {
## Not run:
multihead_attn <- nn_multihead_attention(embed_dim, num_heads)
out <- multihead_attn(query, key, value)</pre>
```

```
attn_output <- out[[1]]
attn_output_weights <- out[[2]]
## End(Not run)
}</pre>
```

nn\_multilabel\_margin\_loss

Multilabel margin loss

### Description

Creates a criterion that optimizes a multi-class multi-classification hinge loss (margin-based loss) between input x (a 2D mini-batch Tensor) and output y (which is a 2D Tensor of target class indices). For each sample in the mini-batch:

### Usage

```
nn_multilabel_margin_loss(reduction = "mean")
```

### **Arguments**

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

### **Details**

$$loss(x, y) = \sum_{ij} \frac{\max(0, 1 - (x[y[j]] - x[i]))}{\text{x.size}(0)}$$

where  $x \in \{0, \dots, \text{ x.size}(0) - 1\}$ ,  $\forall y \in \{0, \dots, \text{ y.size}(0) - 1\}$ ,  $\forall 0 \leq y[j] \leq \text{ x.size}(0) - 1$ , and  $i \neq y[j]$  for all i and j. y and x must have the same size.

The criterion only considers a contiguous block of non-negative targets that starts at the front. This allows for different samples to have variable amounts of target classes.

### Shape

- Input: (C) or (N, C) where N is the batch size and C is the number of classes.
- Target: (C) or (N,C), label targets padded by -1 ensuring same shape as the input.
- Output: scalar. If reduction is 'none', then (N).

### **Examples**

```
if (torch_is_installed()) {
  loss <- nn_multilabel_margin_loss()
  x <- torch_tensor(c(0.1, 0.2, 0.4, 0.8))$view(c(1, 4))
# for target y, only consider labels 4 and 1, not after label -1
  y <- torch_tensor(c(4, 1, -1, 2), dtype = torch_long())$view(c(1, 4))
  loss(x, y)
}</pre>
```

nn\_multilabel\_soft\_margin\_loss

Multi label soft margin loss

## Description

Creates a criterion that optimizes a multi-label one-versus-all loss based on max-entropy, between input x and target y of size (N, C).

### Usage

```
nn_multilabel_soft_margin_loss(weight = NULL, reduction = "mean")
```

#### **Arguments**

weight (Tensor, optional): a manual rescaling weight given to each class. If given, it

has to be a Tensor of size C. Otherwise, it is treated as if having all ones.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

### Details

For each sample in the minibatch:

$$loss(x,y) = -\frac{1}{C} * \sum_{i} y[i] * \log((1 + \exp(-x[i]))^{-1}) + (1 - y[i]) * \log\left(\frac{\exp(-x[i])}{(1 + \exp(-x[i]))}\right)$$

where  $i \in \{0, \dots, x.nElement() - 1\}, y[i] \in \{0, 1\}.$ 

## Shape

- Input: (N, C) where N is the batch size and C is the number of classes.
- Target: (N, C), label targets padded by -1 ensuring same shape as the input.
- Output: scalar. If reduction is 'none', then (N).

nn\_multi\_margin\_loss 265

## **Description**

Creates a criterion that optimizes a multi-class classification hinge loss (margin-based loss) between input x (a 2D mini-batch Tensor) and output y (which is a 1D tensor of target class indices,  $0 \le y \le x.\text{size}(1) - 1$ ):

### Usage

```
nn_multi_margin_loss(p = 1, margin = 1, weight = NULL, reduction = "mean")
```

### **Arguments**

p (int, optional): Has a default value of 1. 1 and 2 are the only supported values.

margin (float, optional): Has a default value of 1.

weight (Tensor, optional): a manual rescaling weight given to each class. If given, it

has to be a Tensor of size C. Otherwise, it is treated as if having all ones.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will

be summed.

## **Details**

For each mini-batch sample, the loss in terms of the 1D input x and scalar output y is:

$$loss(x,y) = \frac{\sum_{i} \max(0, \text{margin} - x[y] + x[i]))^{p}}{\text{x.size}(0)}$$

where  $x \in \{0, \dots, \text{ x.size}(0) - 1\}$  and  $i \neq y$ .

Optionally, you can give non-equal weighting on the classes by passing a 1D weight tensor into the constructor. The loss function then becomes:

$$loss(x,y) = \frac{\sum_{i} \max(0, w[y] * (\mathsf{margin} - x[y] + x[i]))^p)}{\mathsf{x.size}(0)}$$

266 nn\_nll\_loss

nn_nll_loss	Nll loss

#### **Description**

The negative log likelihood loss. It is useful to train a classification problem with C classes.

### Usage

```
nn_nll_loss(weight = NULL, ignore_index = -100, reduction = "mean")
```

## **Arguments**

weight (Tensor, optional): a manual rescaling weight given to each class. If given, it

has to be a Tensor of size C. Otherwise, it is treated as if having all ones.

ignore\_index (int, optional): Specifies a target value that is ignored and does not contribute to

the input gradient.

reduction (string, optional): Specifies the reduction to apply to the output: 'none' | 'mean'

I 'sum'. 'none': no reduction will be applied, 'mean': the weighted mean of

the output is taken, 'sum': the output will be summed.

#### **Details**

If provided, the optional argument weight should be a 1D Tensor assigning weight to each of the classes. This is particularly useful when you have an unbalanced training set.

The input given through a forward call is expected to contain log-probabilities of each class. input has to be a Tensor of size either (minibatch, C) or  $(minibatch, C, d_1, d_2, ..., d_K)$  with  $K \ge 1$  for the K-dimensional case (described later).

Obtaining log-probabilities in a neural network is easily achieved by adding a LogSoftmax layer in the last layer of your network.

You may use CrossEntropyLoss instead, if you prefer not to add an extra layer.

The target that this loss expects should be a class index in the range [0, C-1] where C = number of classes; if ignore\_index is specified, this loss also accepts this class index (this index may not necessarily be in the class range).

The unreduced (i.e. with reduction set to 'none') loss can be described as:

$$\ell(x,y) = L = \{l_1,\dots,l_N\}^\top, \quad l_n = -w_{y_n}x_{n,y_n}, \quad w_c = \mathrm{weight}[c] \cdot 1\{c \neq \mathrm{ignore\_index}\},$$

where x is the input, y is the target, w is the weight, and N is the batch size. If reduction is not 'none' (default 'mean'), then

$$\ell(x,y) = \begin{array}{l} \sum_{n=1}^N \frac{1}{\sum_{n=1}^N w_{y_n}} l_n, & \text{if reduction = 'mean';} \\ \sum_{n=1}^N l_n, & \text{if reduction = 'sum'.} \end{array}$$

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Can also be used for higher dimension inputs, such as 2D images, by providing an input of size  $(minibatch, C, d_1, d_2, ..., d_K)$  with  $K \ge 1$ , where K is the number of dimensions, and a target of appropriate shape (see below). In the case of images, it computes NLL loss per-pixel.

## Shape

- Input: (N,C) where C = number of classes, or  $(N,C,d_1,d_2,...,d_K)$  with  $K\geq 1$  in the case of K-dimensional loss.
- Target: (N) where each value is  $0 \le \operatorname{targets}[i] \le C 1$ , or  $(N, d_1, d_2, ..., d_K)$  with  $K \ge 1$  in the case of K-dimensional loss.
- Output: scalar.

If reduction is 'none', then the same size as the target: (N), or  $(N, d_1, d_2, ..., d_K)$  with  $K \ge 1$  in the case of K-dimensional loss.

### **Examples**

```
if (torch_is_installed()) {
m <- nn_log_softmax(dim = 2)</pre>
loss <- nn_nll_loss()</pre>
# input is of size N x C = 3 \times 5
input <- torch_randn(3, 5, requires_grad = TRUE)</pre>
# each element in target has to have 0 <= value < C</pre>
target <- torch_tensor(c(2, 1, 5), dtype = torch_long())</pre>
output <- loss(m(input), target)</pre>
output$backward()
# 2D loss example (used, for example, with image inputs)
N <- 5
C <- 4
loss <- nn_nll_loss()</pre>
# input is of size N x C x height x width
data <- torch_randn(N, 16, 10, 10)</pre>
conv <- nn_conv2d(16, C, c(3, 3))</pre>
m <- nn_log_softmax(dim = 1)</pre>
# each element in target has to have 0 <= value < C</pre>
target <- torch_empty(N, 8, 8, dtype = torch_long())$random_(1, C)</pre>
output <- loss(m(conv(data)), target)</pre>
output$backward()
}
```

### Description

Computes the batchwise pairwise distance between vectors  $v_1$ ,  $v_2$  using the p-norm:

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### Usage

```
nn_pairwise_distance(p = 2, eps = 1e-06, keepdim = FALSE)
```

### **Arguments**

p (real): the norm degree. Default: 2

eps (float, optional): Small value to avoid division by zero. Default: 1e-6

keepdim (bool, optional): Determines whether or not to keep the vector dimension. De-

fault: FALSE

### **Details**

$$||x||_p = \left(\sum_{i=1}^n |x_i|^p\right)^{1/p}.$$

## Shape

• Input1: (N, D) where D = vector dimension

• Input2: (N, D), same shape as the Input1

• Output: (N). If keepdim is TRUE, then (N, 1).

## **Examples**

```
if (torch_is_installed()) {
pdist <- nn_pairwise_distance(p = 2)
input1 <- torch_randn(100, 128)
input2 <- torch_randn(100, 128)
output <- pdist(input1, input2)
}</pre>
```

nn\_parameter

Creates an nn\_parameter

### **Description**

Indicates to nn\_module that x is a parameter

### Usage

```
nn_parameter(x, requires_grad = TRUE)
```

## **Arguments**

x the tensor that you want to indicate as parameter

requires\_grad whether this parameter should have requires\_grad = TRUE

nn\_poisson\_nll\_loss 269

# Description

Negative log likelihood loss with Poisson distribution of target. The loss can be described as:

### Usage

```
nn_poisson_nll_loss(
  log_input = TRUE,
  full = FALSE,
  eps = 1e-08,
  reduction = "mean"
)
```

# Arguments

log_input	(bool, optional): if TRUE the loss is computed as $\exp(\text{input}) - \text{target} * \text{input}$ , if FALSE the loss is input $- \text{target} * \log(\text{input} + \text{eps})$ .
full	(bool, optional): whether to compute full loss, i. e. to add the Stirling approximation term target $*\log(\text{target}) - \text{target} + 0.5*\log(2\pi \text{target})$ .
eps	(float, optional): Small value to avoid evaluation of $\log(0)$ when log_input = FALSE. Default: 1e-8
reduction	(string, optional): Specifies the reduction to apply to the output: 'none'   'mean'   'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

### **Details**

```
target \sim Poisson(input)loss(input, target) = input - target * log(input) + log(target!)
```

The last term can be omitted or approximated with Stirling formula. The approximation is used for target values more than 1. For targets less or equal to 1 zeros are added to the loss.

# Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input
- Output: scalar by default. If reduction is 'none', then (N,\*), the same shape as the input

270 nn\_prelu

## **Examples**

```
if (torch_is_installed()) {
loss <- nn_poisson_nll_loss()
log_input <- torch_randn(5, 2, requires_grad = TRUE)
target <- torch_randn(5, 2)
output <- loss(log_input, target)
output$backward()
}</pre>
```

nn\_prelu

PReLU module

### **Description**

Applies the element-wise function:

$$PReLU(x) = \max(0, x) + a * \min(0, x)$$

or

$$\operatorname{PReLU}(x) = \left\{ \begin{array}{ll} x, & \text{if } x \geq 0 \\ ax, & \text{otherwise} \end{array} \right.$$

### Usage

```
nn_prelu(num_parameters = 1, init = 0.25)
```

# **Arguments**

```
num_parameters (int): number of a to learn. Although it takes an int as input, there is only two values are legitimate: 1, or the number of channels at input. Default: 1 init (float): the initial value of a. Default: 0.25
```

### **Details**

Here a is a learnable parameter. When called without arguments, nn.prelu() uses a single parameter a across all input channels. If called with nn\_prelu(nChannels), a separate a is used for each input channel.

### Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

### Attributes

• weight (Tensor): the learnable weights of shape (num\_parameters).

nn\_prune\_head 271

## Note

weight decay should not be used when learning a for good performance.

Channel dim is the 2nd dim of input. When input has dims < 2, then there is no channel dim and the number of channels = 1.

# **Examples**

```
if (torch_is_installed()) {
m <- nn_prelu()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_prune\_head

Prune top layer(s) of a network

# Description

Prune head\_size last layers of a nn\_module in order to replace them by your own head, or in order to use the pruned module as a sequential embedding module.

## Usage

```
nn_prune_head(x, head_size)
```

### Arguments

```
x nn_network to prune
head_size number of nn_layers to prune
```

### Value

a nn\_sequential network with the top nn\_layer removed

## **Examples**

```
if (torch_is_installed()) {
  if (torch_is_installed()) {
    x <- nn_sequential(
        nn_relu(),
        nn_tanh(),
        nn_relu6(),
        nn_relu(),
        nn_linear(2,10),
        nn_batch_norm1d(10),
        nn_tanh(),
        nn_linear(10,3)
)</pre>
```

272 nn\_relu6

```
prune <- nn_prune_head(x, 3)
prune
}
</pre>
```

nn\_relu

ReLU module

# Description

Applies the rectified linear unit function element-wise

$$ReLU(x) = (x)^{+} = \max(0, x)$$

## Usage

```
nn_relu(inplace = FALSE)
```

# Arguments

inplace

can optionally do the operation in-place. Default: FALSE

# Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_relu()
input <- torch_randn(2)
m(input)
}</pre>
```

nn\_relu6

ReLu6 module

# Description

Applies the element-wise function:

## Usage

```
nn_relu6(inplace = FALSE)
```

nn\_rnn 273

## **Arguments**

inplace

can optionally do the operation in-place. Default: FALSE

## **Details**

$$ReLU6(x) = \min(\max(0, x), 6)$$

## Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_relu6()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_rnn

RNN module

## **Description**

Applies a multi-layer Elman RNN with tanh or ReLU non-linearity to an input sequence.

# Usage

```
nn_rnn(
  input_size,
  hidden_size,
  num_layers = 1,
  nonlinearity = NULL,
  bias = TRUE,
  batch_first = FALSE,
  dropout = 0,
  bidirectional = FALSE,
  ...
)
```

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#### **Arguments**

The number of expected features in the input x input\_size hidden\_size The number of features in the hidden state h Number of recurrent layers. E.g., setting num\_layers=2 would mean stacking num\_layers two RNNs together to form a stacked RNN, with the second RNN taking in outputs of the first RNN and computing the final results. Default: 1 nonlinearity The non-linearity to use. Can be either 'tanh' or 'relu'. Default: 'tanh' If FALSE, then the layer does not use bias weights b\_ih and b\_hh. Default: TRUE bias If TRUE, then the input and output tensors are provided as (batch, seq, feature). batch\_first Default: FALSE If non-zero, introduces a Dropout layer on the outputs of each RNN layer except dropout the last layer, with dropout probability equal to dropout. Default: 0 bidirectional If TRUE, becomes a bidirectional RNN. Default: FALSE other arguments that can be passed to the super class.

#### **Details**

For each element in the input sequence, each layer computes the following function:

$$h_t = \tanh(W_{ih}x_t + b_{ih} + W_{hh}h_{(t-1)} + b_{hh})$$

where  $h_t$  is the hidden state at time t,  $x_t$  is the input at time t, and  $h_{(t-1)}$  is the hidden state of the previous layer at time t-1 or the initial hidden state at time 0. If nonlinearity is 'relu', then ReLU is used instead of tanh.

### **Inputs**

- **input** of shape (seq\_len, batch, input\_size): tensor containing the features of the input sequence. The input can also be a packed variable length sequence.
- h\_0 of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the initial hidden state for each element in the batch. Defaults to zero if not provided. If the RNN is bidirectional, num\_directions should be 2, else it should be 1.

#### **Outputs**

- output of shape (seq\_len, batch, num\_directions \* hidden\_size): tensor containing the output features (h\_t) from the last layer of the RNN, for each t. If a :class:nn\_packed\_sequence has been given as the input, the output will also be a packed sequence. For the unpacked case, the directions can be separated using output\$view(seq\_len, batch, num\_directions, hidden\_size), with forward and backward being direction 0 and 1 respectively. Similarly, the directions can be separated in the packed case.
- h\_n of shape (num\_layers \* num\_directions, batch, hidden\_size): tensor containing the hidden state for t = seq\_len. Like *output*, the layers can be separated using h\_n\$view(num\_layers, num\_directions, batch, hidden\_size).

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#### Shape

- Input1:  $(L, N, H_{in})$  tensor containing input features where  $H_{in} = \text{input\_size}$  and L represents a sequence length.
- Input2:  $(S, N, H_{out})$  tensor containing the initial hidden state for each element in the batch.  $H_{out} = \text{hidden\_size Defaults to zero if not provided.}$  where  $S = \text{num\_layers*num\_directions}$  If the RNN is bidirectional, num\_directions should be 2, else it should be 1.
- Output1:  $(L, N, H_{all})$  where  $H_{all} = \text{num\_directions} * \text{hidden\_size}$
- Output2:  $(S, N, H_{out})$  tensor containing the next hidden state for each element in the batch

#### **Attributes**

- weight\_ih\_1[k]: the learnable input-hidden weights of the k-th layer, of shape (hidden\_size, input\_size) for k = 0. Otherwise, the shape is (hidden\_size, num\_directions \* hidden\_size)
- weight\_hh\_1[k]: the learnable hidden-hidden weights of the k-th layer, of shape (hidden\_size, hidden\_size)
- bias\_ih\_l[k]: the learnable input-hidden bias of the k-th layer, of shape (hidden\_size)
- bias\_hh\_1[k]: the learnable hidden-hidden bias of the k-th layer, of shape (hidden\_size)

#### Note

All the weights and biases are initialized from  $\mathcal{U}(-\sqrt{k},\sqrt{k})$  where  $k=\frac{1}{\text{hidden size}}$ 

### **Examples**

```
if (torch_is_installed()) {
  rnn <- nn_rnn(10, 20, 2)
  input <- torch_randn(5, 3, 10)
  h0 <- torch_randn(2, 3, 20)
  rnn(input, h0)
}</pre>
```

nn\_rrelu

RReLU module

### **Description**

Applies the randomized leaky rectified liner unit function, element-wise, as described in the paper:

#### Usage

```
nn_rrelu(lower = 1/8, upper = 1/3, inplace = FALSE)
```

#### **Arguments**

```
lower lower bound of the uniform distribution. Default: \frac{1}{8} upper upper bound of the uniform distribution. Default: \frac{1}{3} inplace can optionally do the operation in-place. Default: FALSE
```

276 nn\_selu

### **Details**

Empirical Evaluation of Rectified Activations in Convolutional Network.

The function is defined as:

$$RReLU(x) = \begin{cases} x & \text{if } x \ge 0 \\ ax & \text{otherwise} \end{cases}$$

where a is randomly sampled from uniform distribution  $\mathcal{U}(\text{lower}, \text{upper})$ . See: https://arxiv.org/pdf/1505.00853.pdf

# Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_rrelu(0.1, 0.3)
input <- torch_randn(2)
m(input)
}</pre>
```

nn\_selu

SELU module

# Description

Applied element-wise, as:

# Usage

```
nn_selu(inplace = FALSE)
```

## **Arguments**

inplace

(bool, optional): can optionally do the operation in-place. Default: FALSE

## **Details**

$$\mathrm{SELU}(x) = \mathrm{scale} * (\max(0, x) + \min(0, \alpha * (\exp(x) - 1)))$$

with  $\alpha = 1.6732632423543772848170429916717$  and scale = 1.0507009873554804934193349852946.

More details can be found in the paper Self-Normalizing Neural Networks.

nn\_sequential 277

## Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_selu()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_sequential

A sequential container

## **Description**

A sequential container. Modules will be added to it in the order they are passed in the constructor. See examples.

## Usage

```
nn_sequential(...)
```

## **Arguments**

. . . sequence of modules to be added

### **Examples**

```
if (torch_is_installed()) {

model <- nn_sequential(
   nn_conv2d(1, 20, 5),
   nn_relu(),
   nn_conv2d(20, 64, 5),
   nn_relu()
)
input <- torch_randn(32, 1, 28, 28)
output <- model(input)
}</pre>
```

nn\_silu

 $nn\_sigmoid$ 

Sigmoid module

# Description

Applies the element-wise function:

## Usage

```
nn_sigmoid()
```

### **Details**

$$\operatorname{Sigmoid}(x) = \sigma(x) = \frac{1}{1 + \exp(-x)}$$

# Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_sigmoid()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_silu

Applies the Sigmoid Linear Unit (SiLU) function, element-wise. The SiLU function is also known as the swish function.

# Description

Applies the Sigmoid Linear Unit (SiLU) function, element-wise. The SiLU function is also known as the swish function.

## Usage

```
nn_silu(inplace = FALSE)
```

## **Arguments**

inplace

can optionally do the operation in-place. Default: FALSE

nn\_smooth\_11\_loss 279

### **Details**

See Gaussian Error Linear Units (GELUs) where the SiLU (Sigmoid Linear Unit) was originally coined, and see Sigmoid-Weighted Linear Units for Neural Network Function Approximation in Reinforcement Learning and Swish: a Self-Gated Activation Function where the SiLU was experimented with later.

nn\_smooth\_l1\_loss

Smooth L1 loss

### **Description**

Creates a criterion that uses a squared term if the absolute element-wise error falls below 1 and an L1 term otherwise. It is less sensitive to outliers than the MSELoss and in some cases prevents exploding gradients (e.g. see Fast R-CNN paper by Ross Girshick). Also known as the Huber loss:

## Usage

```
nn_smooth_l1_loss(reduction = "mean")
```

## **Arguments**

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

#### **Details**

$$loss(x,y) = \frac{1}{n} \sum_{i} z_{i}$$

where  $z_i$  is given by:

$$z_i = \begin{array}{ll} 0.5(x_i - y_i)^2, & \text{if } |x_i - y_i| < 1 \\ |x_i - y_i| - 0.5, & \text{otherwise} \end{array}$$

x and y arbitrary shapes with a total of n elements each the sum operation still operates over all the elements, and divides by n. The division by n can be avoided if sets reduction = 'sum'.

# Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Target: (N, \*), same shape as the input
- Output: scalar. If reduction is 'none', then (N, \*), same shape as the input

280 nn\_softmax

nn\_softmax

Softmax module

## **Description**

Applies the Softmax function to an n-dimensional input Tensor rescaling them so that the elements of the n-dimensional output Tensor lie in the range [0,1] and sum to 1. Softmax is defined as:

## Usage

```
nn_softmax(dim)
```

### **Arguments**

dim

(int): A dimension along which Softmax will be computed (so every slice along dim will sum to 1).

#### **Details**

$$Softmax(x_i) = \frac{\exp(x_i)}{\sum_{j} \exp(x_j)}$$

When the input Tensor is a sparse tensor then the unspecifed values are treated as -Inf.

## Value

: a Tensor of the same dimension and shape as the input with values in the range [0, 1]

### Shape

- Input: (\*) where \* means, any number of additional dimensions
- Output: (\*), same shape as the input

#### Note

This module doesn't work directly with NLLLoss, which expects the Log to be computed between the Softmax and itself. Use LogSoftmax instead (it's faster and has better numerical properties).

### **Examples**

```
if (torch_is_installed()) {
m <- nn_softmax(1)
input <- torch_randn(2, 3)
output <- m(input)
}</pre>
```

nn\_softmax2d 281

 $nn\_softmax2d$ 

Softmax2d module

### **Description**

Applies SoftMax over features to each spatial location. When given an image of Channels x Height x Width, it will apply Softmax to each location  $(Channels, h_i, w_i)$ 

## Usage

```
nn_softmax2d()
```

#### Value

a Tensor of the same dimension and shape as the input with values in the range [0, 1]

# Shape

- Input: (N, C, H, W)
- Output: (N, C, H, W) (same shape as input)

## **Examples**

```
if (torch_is_installed()) {
m <- nn_softmax2d()
input <- torch_randn(2, 3, 12, 13)
output <- m(input)
}</pre>
```

nn\_softmin

Softmin

### **Description**

Applies the Softmin function to an n-dimensional input Tensor rescaling them so that the elements of the n-dimensional output Tensor lie in the range [0, 1] and sum to 1. Softmin is defined as:

## Usage

```
nn_softmin(dim)
```

### **Arguments**

dim

(int): A dimension along which Softmin will be computed (so every slice along dim will sum to 1).

282 nn\_softplus

### **Details**

Softmin
$$(x_i) = \frac{\exp(-x_i)}{\sum_j \exp(-x_j)}$$

#### Value

a Tensor of the same dimension and shape as the input, with values in the range [0, 1].

### Shape

- Input: (\*) where \* means, any number of additional dimensions
- Output: (\*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_softmin(dim = 1)
input <- torch_randn(2, 2)
output <- m(input)
}</pre>
```

nn\_softplus

Softplus module

# Description

Applies the element-wise function:

$$\operatorname{Softplus}(x) = \frac{1}{\beta} * \log(1 + \exp(\beta * x))$$

## Usage

```
nn_softplus(beta = 1, threshold = 20)
```

## **Arguments**

beta the  $\beta$  value for the Softplus formulation. Default: 1 threshold values above this revert to a linear function. Default: 20

## **Details**

SoftPlus is a smooth approximation to the ReLU function and can be used to constrain the output of a machine to always be positive. For numerical stability the implementation reverts to the linear function when  $input \times \beta > threshold$ .

nn\_softshrink 283

## Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_softplus()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_softshrink

Softshrink module

## **Description**

Applies the soft shrinkage function elementwise:

## Usage

```
nn_softshrink(lambd = 0.5)
```

## **Arguments**

lambd

the  $\lambda$  (must be no less than zero) value for the Softshrink formulation. Default:  $0.5\,$ 

### **Details**

$$\mbox{SoftShrinkage}(x) = \left\{ \begin{array}{ll} x - \lambda, & \mbox{if } x > \lambda \\ x + \lambda, & \mbox{if } x < -\lambda \\ 0, & \mbox{otherwise} \end{array} \right.$$

## Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_softshrink()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

284 nn\_soft\_margin\_loss

nn\_softsign

Softsign module

### **Description**

Applies the element-wise function:

$$SoftSign(x) = \frac{x}{1 + |x|}$$

### Usage

```
nn_softsign()
```

## Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_softsign()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_soft\_margin\_loss

Soft margin loss

# Description

Creates a criterion that optimizes a two-class classification logistic loss between input tensor x and target tensor y (containing 1 or -1).

## Usage

```
nn_soft_margin_loss(reduction = "mean")
```

## **Arguments**

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

nn\_tanh

## **Details**

$$loss(x, y) = \sum_{i} \frac{\log(1 + \exp(-y[i] * x[i]))}{\text{x.nelement()}}$$

## Shape

- Input: (\*) where \* means, any number of additional dimensions
- Target: (\*), same shape as the input
- Output: scalar. If reduction is 'none', then same shape as the input

nn\_tanh

Tanh module

# Description

Applies the element-wise function:

## Usage

nn\_tanh()

## **Details**

$$\mathrm{Tanh}(x) = \tanh(x) = \frac{\exp(x) - \exp(-x)}{\exp(x) + \exp(-x)}$$

### Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_tanh()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

286 nn\_threshold

nn\_tanhshrink

Tanhshrink module

## **Description**

Applies the element-wise function:

## Usage

```
nn_tanhshrink()
```

## **Details**

$$Tanhshrink(x) = x - tanh(x)$$

## Shape

- Input: (N,\*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

# **Examples**

```
if (torch_is_installed()) {
m <- nn_tanhshrink()
input <- torch_randn(2)
output <- m(input)
}</pre>
```

nn\_threshold

Threshold module

# **Description**

Thresholds each element of the input Tensor.

# Usage

```
nn_threshold(threshold, value, inplace = FALSE)
```

## **Arguments**

threshold The value to threshold at value The value to replace with

inplace can optionally do the operation in-place. Default: FALSE

### **Details**

Threshold is defined as:

$$y = \begin{cases} x, & \text{if } x > \text{threshold} \\ \text{value}, & \text{otherwise} \end{cases}$$

### Shape

- Input: (N, \*) where \* means, any number of additional dimensions
- Output: (N, \*), same shape as the input

## **Examples**

```
if (torch_is_installed()) {
m <- nn_threshold(0.1, 20)
input <- torch_randn(2)
output <- m(input)
}</pre>
```

```
nn_triplet_margin_loss
```

Triplet margin loss

## **Description**

Creates a criterion that measures the triplet loss given an input tensors x1, x2, x3 and a margin with a value greater than 0. This is used for measuring a relative similarity between samples. A triplet is composed by a, p and n (i.e., anchor, positive examples and negative examples respectively). The shapes of all input tensors should be (N, D).

### Usage

```
nn_triplet_margin_loss(
  margin = 1,
  p = 2,
  eps = 1e-06,
  swap = FALSE,
  reduction = "mean"
)
```

## **Arguments**

margin (float, optional): Default: 1.

p (int, optional): The norm degree for pairwise distance. Default: 2.

eps constant to avoid NaN's

swap (bool, optional): The distance swap is described in detail in the paper Learning shallow convolutional feature descriptors with triplet losses by V. Balntas, E. Riba et al. Default: FALSE.

reduction

(string, optional): Specifies the reduction to apply to the output: 'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output, 'sum': the output will be summed.

#### **Details**

The distance swap is described in detail in the paper Learning shallow convolutional feature descriptors with triplet losses by V. Balntas, E. Riba et al.

The loss function for each sample in the mini-batch is:

$$L(a, p, n) = \max\{d(a_i, p_i) - d(a_i, n_i) + \text{margin}, 0\}$$

where

$$d(x_i, y_i) = |\mathbf{x}_i - \mathbf{y}_i|_p$$

See also nn\_triplet\_margin\_with\_distance\_loss(), which computes the triplet margin loss for input tensors using a custom distance function.

## Shape

- Input: (N, D) where D is the vector dimension.
- Output: A Tensor of shape (N) if reduction is 'none', or a scalar otherwise.

### **Examples**

```
if (torch_is_installed()) {
  triplet_loss <- nn_triplet_margin_loss(margin = 1, p = 2)
  anchor <- torch_randn(100, 128, requires_grad = TRUE)
  positive <- torch_randn(100, 128, requires_grad = TRUE)
  negative <- torch_randn(100, 128, requires_grad = TRUE)
  output <- triplet_loss(anchor, positive, negative)
  output$backward()
}</pre>
```

```
nn_triplet_margin_with_distance_loss

*Triplet margin with distance loss*
```

## Description

Creates a criterion that measures the triplet loss given input tensors a, p, and n (representing anchor, positive, and negative examples, respectively), and a nonnegative, real-valued function ("distance function") used to compute the relationship between the anchor and positive example ("positive distance") and the anchor and negative example ("negative distance").

#### Usage

```
nn_triplet_margin_with_distance_loss(
  distance_function = NULL,
  margin = 1,
  swap = FALSE,
  reduction = "mean"
)
```

#### Arguments

distance\_function

(callable, optional): A nonnegative, real-valued function that quantifies the closeness of two tensors. If not specified, nn\_pairwise\_distance() will be used.

Default: None

margin (float, optional): A non-negative margin representing the minimum difference

between the positive and negative distances required for the loss to be 0. Larger margins penalize cases where the negative examples are not distant enough from

the anchors, relative to the positives. Default: 1.

swap (bool, optional): Whether to use the distance swap described in the paper Learn-

ing shallow convolutional feature descriptors with triplet losses by V. Balntas, E. Riba et al. If TRUE, and if the positive example is closer to the negative example than the anchor is, swaps the positive example and the anchor in the loss

computation. Default: FALSE.

reduction (string, optional): Specifies the (optional) reduction to apply to the output:

'none' | 'mean' | 'sum'. 'none': no reduction will be applied, 'mean': the sum of the output will be divided by the number of elements in the output,

'sum': the output will be summed. Default: 'mean'

## Details

The unreduced loss (i.e., with reduction set to 'none') can be described as:

$$\ell(a, p, n) = L = \{l_1, \dots, l_N\}^{\top}, \quad l_i = \max\{d(a_i, p_i) - d(a_i, n_i) + \text{margin}, 0\}$$

where N is the batch size; d is a nonnegative, real-valued function quantifying the closeness of two tensors, referred to as the distance\_function; and margin is a non-negative margin representing the minimum difference between the positive and negative distances that is required for the loss to be 0. The input tensors have N elements each and can be of any shape that the distance function can handle. If reduction is not 'none' (default 'mean'), then:

```
\ell(x,y) = \begin{array}{ll} \operatorname{mean}(L), & \text{if reduction} = \text{`mean'}; \\ \operatorname{sum}(L), & \text{if reduction} = \text{`sum'}. \end{array}
```

See also nn\_triplet\_margin\_loss(), which computes the triplet loss for input tensors using the  $l_p$  distance as the distance function.

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#### Shape

• Input: (N,\*) where \* represents any number of additional dimensions as supported by the distance function.

• Output: A Tensor of shape (N) if reduction is 'none', or a scalar otherwise.

#### **Examples**

```
if (torch_is_installed()) {
# Initialize embeddings
embedding <- nn_embedding(1000, 128)</pre>
anchor_ids <- torch_randint(1, 1000, 1, dtype = torch_long())</pre>
positive_ids <- torch_randint(1, 1000, 1, dtype = torch_long())</pre>
negative_ids <- torch_randint(1, 1000, 1, dtype = torch_long())</pre>
anchor <- embedding(anchor_ids)</pre>
positive <- embedding(positive_ids)</pre>
negative <- embedding(negative_ids)</pre>
# Built-in Distance Function
triplet_loss <- nn_triplet_margin_with_distance_loss(</pre>
  distance_function = nn_pairwise_distance()
output <- triplet_loss(anchor, positive, negative)</pre>
# Custom Distance Function
l_infinity <- function(x1, x2) {</pre>
  torch_max(torch_abs(x1 - x2), dim = 1)[[1]]
}
triplet_loss <- nn_triplet_margin_with_distance_loss(</pre>
  distance_function = l_infinity, margin = 1.5
)
output <- triplet_loss(anchor, positive, negative)</pre>
# Custom Distance Function (Lambda)
triplet_loss <- nn_triplet_margin_with_distance_loss(</pre>
  distance_function = function(x, y) {
    1 - nnf_cosine_similarity(x, y)
  }
)
output <- triplet_loss(anchor, positive, negative)</pre>
}
```

 $nn\_unflatten$ 

Unflattens a tensor dim expanding it to a desired shape. For use with [nn\_sequential.

#### **Description**

Unflattens a tensor dim expanding it to a desired shape. For use with [nn\_sequential.

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#### Usage

```
nn_unflatten(dim, unflattened_size)
```

#### **Arguments**

```
dim Dimension to be unflattened unflattened_size
```

New shape of the unflattened dimension

## **Examples**

```
if (torch_is_installed()) {
input <- torch_randn(2, 50)

m <- nn_sequential(
    nn_linear(50, 50),
    nn_unflatten(2, c(2, 5, 5))
)
output <- m(input)
output$size()
}</pre>
```

nn\_upsample

Upsample module

### **Description**

Upsamples a given multi-channel 1D (temporal), 2D (spatial) or 3D (volumetric) data. The input data is assumed to be of the form minibatch x channels x optional depth x optional height] x width. Hence, for spatial inputs, we expect a 4D Tensor and for volumetric inputs, we expect a 5D Tensor.

### Usage

```
nn_upsample(
    size = NULL,
    scale_factor = NULL,
    mode = "nearest",
    align_corners = NULL)
```

### **Arguments**

align\_corners (bool, optional): if TRUE, the corner pixels of the input and output tensors are aligned, and thus preserving the values at those pixels. This only has effect when mode is 'linear', 'bilinear', or 'trilinear'. Default: FALSE

#### **Details**

The algorithms available for upsampling are nearest neighbor and linear, bilinear, bicubic and trilinear for 3D, 4D and 5D input Tensor, respectively.

One can either give a scale\_factor or the target output size to calculate the output size. (You cannot give both, as it is ambiguous)

### **Examples**

```
if (torch_is_installed()) {
  input <- torch_arange(start = 1, end = 4, dtype = torch_float())$view(c(1, 1, 2, 2))
  nn_upsample(scale_factor = c(2), mode = "nearest")(input)
  nn_upsample(scale_factor = c(2, 2), mode = "nearest")(input)
}</pre>
```

```
nn_utils_clip_grad_norm_
```

Clips gradient norm of an iterable of parameters.

#### **Description**

The norm is computed over all gradients together, as if they were concatenated into a single vector. Gradients are modified in-place.

### Usage

```
nn_utils_clip_grad_norm_(parameters, max_norm, norm_type = 2)
```

#### **Arguments**

parameters	(IterableTensor or Tensor): an iterable of Tensors or a single Tensor that will have gradients normalized
max_norm	(float or int): max norm of the gradients
norm_type	(float or int): type of the used p-norm. Can be Inf for infinity norm.

#### Value

Total norm of the parameters (viewed as a single vector).

```
nn_utils_clip_grad_value_
```

Clips gradient of an iterable of parameters at specified value.

## Description

Gradients are modified in-place.

#### Usage

```
nn_utils_clip_grad_value_(parameters, clip_value)
```

### **Arguments**

parameters (Iterable(Tensor) or Tensor): an iterable of Tensors or a single Tensor that will

have gradients normalized

clip\_value (float or int): maximum allowed value of the gradients.

#### **Details**

The gradients are clipped in the range [-clip\_value, clip\_value]

```
nn_utils_rnn_pack_padded_sequence
```

Packs a Tensor containing padded sequences of variable length.

## **Description**

input can be of size T  $\times$  B  $\times$  \* where T is the length of the longest sequence (equal to lengths[1]), B is the batch size, and \* is any number of dimensions (including 0). If batch\_first is TRUE, B  $\times$  T  $\times$  \* input is expected.

### Usage

```
nn_utils_rnn_pack_padded_sequence(
  input,
  lengths,
  batch_first = FALSE,
  enforce_sorted = TRUE
)
```

#### Arguments

input (Tensor): padded batch of variable length sequences.

lengths (Tensor): list of sequences lengths of each batch element.

batch\_first (bool, optional): if TRUE, the input is expected in B x T x \* format.

enforce\_sorted (bool, optional): if TRUE, the input is expected to contain sequences sorted by

length in a decreasing order. If FALSE, the input will get sorted unconditionally.

Default: TRUE.

#### **Details**

For unsorted sequences, use enforce\_sorted = FALSE. If enforce\_sorted is TRUE, the sequences should be sorted by length in a decreasing order, i.e. input[,1] should be the longest sequence, and input[,B] the shortest one. enforce\_sorted = TRUE is only necessary for ONNX export.

#### Value

a PackedSequence object

#### Note

This function accepts any input that has at least two dimensions. You can apply it to pack the labels, and use the output of the RNN with them to compute the loss directly. A Tensor can be retrieved from a PackedSequence object by accessing its .data attribute.

```
nn_utils_rnn_pack_sequence
```

Packs a list of variable length Tensors

#### **Description**

sequences should be a list of Tensors of size  $L \times *$ , where L is the length of a sequence and \* is any number of trailing dimensions, including zero.

## Usage

```
nn_utils_rnn_pack_sequence(sequences, enforce_sorted = TRUE)
```

#### **Arguments**

sequences (list[Tensor]): A list of sequences of decreasing length.

enforce\_sorted (bool, optional): if TRUE, checks that the input contains sequences sorted by

length in a decreasing order. If FALSE, this condition is not checked. Default:

TRUE.

#### **Details**

For unsorted sequences, use enforce\_sorted = FALSE. If enforce\_sorted is TRUE, the sequences should be sorted in the order of decreasing length. enforce\_sorted = TRUE is only necessary for ONNX export.

#### Value

a PackedSequence object

### **Examples**

```
if (torch_is_installed()) {
x <- torch_tensor(c(1, 2, 3), dtype = torch_long())
y <- torch_tensor(c(4, 5), dtype = torch_long())
z <- torch_tensor(c(6), dtype = torch_long())
p <- nn_utils_rnn_pack_sequence(list(x, y, z))
}</pre>
```

nn\_utils\_rnn\_pad\_packed\_sequence

Pads a packed batch of variable length sequences.

### **Description**

It is an inverse operation to nn\_utils\_rnn\_pack\_padded\_sequence().

### Usage

```
nn_utils_rnn_pad_packed_sequence(
   sequence,
   batch_first = FALSE,
   padding_value = 0,
   total_length = NULL
)
```

# Arguments

```
sequence (PackedSequence): batch to pad

batch_first (bool, optional): if True, the output will be in "B x T x * format.

padding_value (float, optional): values for padded elements.

total_length (int, optional): if not NULL, the output will be padded to have length total_length.

This method will throw ValueError if total_length is less than the max sequence length in sequence.
```

#### **Details**

The returned Tensor's data will be of size T  $\times$  B  $\times$  \*, where T is the length of the longest sequence and B is the batch size. If batch\_first is TRUE, the data will be transposed into B  $\times$  T  $\times$  \* format.

#### Value

Tuple of Tensor containing the padded sequence, and a Tensor containing the list of lengths of each sequence in the batch. Batch elements will be re-ordered as they were ordered originally when the batch was passed to nn\_utils\_rnn\_pack\_padded\_sequence() or nn\_utils\_rnn\_pack\_sequence().

#### Note

total\_length is useful to implement the pack sequence -> recurrent network -> unpack sequence pattern in a nn\_module wrapped in ~torch.nn.DataParallel.

#### **Examples**

```
if (torch_is_installed()) {
  seq <- torch_tensor(rbind(c(1, 2, 0), c(3, 0, 0), c(4, 5, 6)))
  lens <- c(2, 1, 3)
  packed <- nn_utils_rnn_pack_padded_sequence(seq, lens,
      batch_first = TRUE,
      enforce_sorted = FALSE
)
  packed
  nn_utils_rnn_pad_packed_sequence(packed, batch_first = TRUE)
}</pre>
```

nn\_utils\_rnn\_pad\_sequence

Pad a list of variable length Tensors with padding\_value

### Description

pad\_sequence stacks a list of Tensors along a new dimension, and pads them to equal length. For example, if the input is list of sequences with size  $L \times *$  and if batch\_first is False, and  $T \times B \times *$  otherwise.

#### Usage

```
nn_utils_rnn_pad_sequence(sequences, batch_first = FALSE, padding_value = 0)
```

# Arguments

```
sequences (list[Tensor]): list of variable length sequences.

batch_first (bool, optional): output will be in B x T x * if TRUE, or in T x B x * otherwise padding_value (float, optional): value for padded elements. Default: 0.
```

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#### **Details**

B is batch size. It is equal to the number of elements in sequences. T is length of the longest sequence. L is length of the sequence. \* is any number of trailing dimensions, including none.

#### Value

```
Tensor of size T x B x * if batch_first is FALSE. Tensor of size B x T x * otherwise
```

#### Note

This function returns a Tensor of size  $T \times B \times *$  or  $B \times T \times *$  where T is the length of the longest sequence. This function assumes trailing dimensions and type of all the Tensors in sequences are same.

# **Examples**

```
if (torch_is_installed()) {
a <- torch_ones(25, 300)
b <- torch_ones(22, 300)
c <- torch_ones(15, 300)
nn_utils_rnn_pad_sequence(list(a, b, c))$size()
}</pre>
```

### **Description**

Applies weight normalization to a parameter in the given module.

#### **Details**

```
\left\{ w = g \left( mathbf\{v\} \right) \right\}
```

Weight normalization is a reparameterization that decouples the magnitude of a weight tensor from its direction. This replaces the parameter specified by name (e.g. 'weight') with two parameters: one specifying the magnitude (e.g. 'weight\_g') and one specifying the direction (e.g. 'weight\_v').

#### Value

The original module with the weight\_v and weight\_g paramters.

#### Methods

#### **Public methods:**

```
• nn_utils_weight_norm$new()
  • nn_utils_weight_norm$compute_weight()
  • nn_utils_weight_norm$apply()
  • nn_utils_weight_norm$call()
  • nn_utils_weight_norm$recompute()
  • nn_utils_weight_norm$remove()
  • nn_utils_weight_norm$clone()
Method new():
 Usage:
 nn_utils_weight_norm$new(name, dim)
 Arguments:
 name (str, optional): name of weight parameter
 dim (int, optional): dimension over which to compute the norm
Method compute_weight():
 Usage:
 nn_utils_weight_norm$compute_weight(module, name = NULL, dim = NULL)
 Arguments:
 module (Module): containing module
 name (str, optional): name of weight parameter
 dim (int, optional): dimension over which to compute the norm
Method apply():
 Usage:
 nn_utils_weight_norm$apply(module, name = NULL, dim = NULL)
 Arguments:
 module (Module): containing module
 name (str, optional): name of weight parameter
 dim (int, optional): dimension over which to compute the norm
Method call():
 Usage:
 nn_utils_weight_norm$call(module)
 Arguments:
 module (Module): containing module
Method recompute():
 Usage:
 nn_utils_weight_norm$recompute(module)
```

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```
Arguments:
  module (Module): containing module

Method remove():
    Usage:
    nn_utils_weight_norm$remove(module, name = NULL)

Arguments:
    module (Module): containing module
    name (str, optional): name of weight parameter

Method clone(): The objects of this class are cloneable with this method.
    Usage:
    nn_utils_weight_norm$clone(deep = FALSE)

Arguments:
    deep Whether to make a deep clone.
```

#### Note

The pytorch Weight normalization is implemented via a hook that recomputes the weight tensor from the magnitude and direction before every forward() call. Since torch for R still do not support hooks, the weight recomputation need to be done explicitly inside the forward() definition trough a call of the recompute() method. See examples.

By default, with dim = 0, the norm is computed independently per output channel/plane. To compute a norm over the entire weight tensor, use dim = NULL.

@references https://arxiv.org/abs/1602.07868

```
if (torch_is_installed()) {
    x = nn_linear(in_features = 20, out_features = 40)
    weight_norm = nn_utils_weight_norm$new(name = 'weight', dim = 2)
    weight_norm$apply(x)
    x$weight_g$size()
    x$weight_v$size()
    x$weight
# the recompute() method recomputes the weight using g and v. It must be called
# explicitly inside `forward()`.
    weight_norm$recompute(x)
}
```

300 optimizer

optimizer	Creates a custom optimizer

## Description

When implementing custom optimizers you will usually need to implement the initialize and step methods. See the example section below for a full example.

# Usage

```
optimizer(
  name = NULL,
  inherit = Optimizer,
  ...,
  private = NULL,
  active = NULL,
  parent_env = parent.frame()
)
```

### **Arguments**

name	(optional) name of the optimizer
inherit	(optional) you can inherit from other optimizers to re-use some methods.
• • •	Pass any number of fields or methods. You should at least define the initialize and step methods. See the examples section.
private	(optional) a list of private methods for the optimizer.
active	(optional) a list of active methods for the optimizer.
parent_env	used to capture the right environment to define the class. The default is fine for most situations.

# Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

```
if (torch_is_installed()) {
# In this example we will create a custom optimizer
# that's just a simplified version of the `optim_sgd` function.

optim_sgd2 <- optimizer(
  initialize = function(params, learning_rate) {</pre>
```

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```
defaults <- list(</pre>
      learning_rate = learning_rate
    super$initialize(params, defaults)
  },
  step = function() {
    with_no_grad({
      for (g in seq_along(self$param_groups)) {
        group <- self$param_groups[[g]]</pre>
        for (p in seq_along(group$params)) {
          param <- group$params[[p]]</pre>
          if (is.null(param$grad) || is_undefined_tensor(param$grad)) {
            next
          }
          param$add_(param$grad, alpha = -group$learning_rate)
        }
      }
    })
 }
x <- torch_randn(1, requires_grad = TRUE)</pre>
opt <- optim_sgd2(x, learning_rate = 0.1)</pre>
for (i in 1:100) {
  opt$zero_grad()
  y < - x^2
  y$backward()
  opt$step()
}
all.equal(x$item(), 0, tolerance = 1e-9)
```

optim\_adadelta

Adadelta optimizer

### **Description**

It has been proposed in ADADELTA: An Adaptive Learning Rate Method

# Usage

```
optim_adadelta(params, lr = 1, rho = 0.9, eps = 1e-06, weight_decay = 0)
```

## Arguments

```
params (iterable): list of parameters to optimize or list defining parameter groups 
1r (float, optional): learning rate (default: 1e-3)
```

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rho	(float, optional): coefficient used for computing a running average of squared gradients (default: 0.9)
eps	(float, optional): term added to the denominator to improve numerical stability (default: 1e-6)
weight_decay	(float, optional): weight decay (L2 penalty) (default: 0)

# Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

#### Note

According to the original paper, decaying average of the squared gradients is computed as follows:

$$E[g^2]_t = \rho E[g^2]_{t-1} + (1 - \rho)g_t^2$$

RMS of previous squared gradients up to time t:

$$RMS[g_t] = \sqrt{E[g^2]_t + \epsilon}$$

Adadelta update rule:

$$\Delta\theta_t = -\frac{RMS[\Delta\theta]_{t-1}}{RMS[g]_t}\theta_{t+1} = \theta_t + \Delta\theta_t$$

## **Examples**

```
if (torch_is_installed()) {
## Not run:
optimizer <- optim_adadelta(model$parameters, lr = 0.1)
optimizer$zero_grad()
loss_fn(model(input), target)$backward()
optimizer$step()
## End(Not run)</pre>
```

optim\_adagrad

Adagrad optimizer

#### Description

Proposed in Adaptive Subgradient Methods for Online Learning and Stochastic Optimization

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#### Usage

```
optim_adagrad(
  params,
  lr = 0.01,
  lr_decay = 0,
  weight_decay = 0,
  initial_accumulator_value = 0,
  eps = 1e-10
)
```

#### **Arguments**

params (iterable): list of parameters to optimize or list parameter groups

1r (float, optional): learning rate (default: 1e-2)

1r\_decay (float, optional): learning rate decay (default: 0)

weight\_decay (float, optional): weight decay (L2 penalty) (default: 0)

initial\_accumulator\_value

the initial value for the accumulator. (default: 0)

Adagrad is an especially good optimizer for sparse data. It individually modifies learning rate for every single parameter, dividing the original learning rate value by sum of the squares of the gradients. It causes that the rarely occurring features get greater learning rates. The main downside of this method is the fact that learning rate may be getting small too fast, so that at some point a model cannot

learn anymore.

eps (float, optional): term added to the denominator to improve numerical stability

(default: 1e-10)

#### Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

#### Note

Update rule:

$$\theta_{t+1} = \theta_t - \frac{\eta}{\sqrt{G_t + \epsilon}} \odot g_t$$

The equation above and some remarks quoted after *An overview of gradient descent optimization algorithms* by Sebastian Ruder.

304 optim\_adam

optim\_adam

Implements Adam algorithm.

### **Description**

It has been proposed in Adam: A Method for Stochastic Optimization.

## Usage

```
optim_adam(
  params,
  lr = 0.001,
  betas = c(0.9, 0.999),
  eps = 1e-08,
  weight_decay = 0,
  amsgrad = FALSE
)
```

### **Arguments**

params (iterable): iterable of parameters to optimize or dicts defining parameter groups

lr (float, optional): learning rate (default: 1e-3)

betas (Tuple[float, float], optional): coefficients used for computing running averages of gradient and its square (default: (0.9, 0.999))

eps (float, optional): term added to the denominator to improve numerical stability (default: 1e-8)

weight\_decay (float, optional): weight decay (L2 penalty) (default: 0)

amsgrad (boolean, optional): whether to use the AMSGrad variant of this algorithm from the paper On the Convergence of Adam and Beyond (default: FALSE)

#### Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

```
if (torch_is_installed()) {
## Not run:
optimizer <- optim_adam(model$parameters(), lr = 0.1)
optimizer$zero_grad()
loss_fn(model(input), target)$backward()
optimizer$step()</pre>
```

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```
## End(Not run)
}
```

optim\_adamw

Implements AdamW algorithm

# Description

For further details regarding the algorithm we refer to Decoupled Weight Decay Regularization

# Usage

```
optim_adamw(
  params,
  lr = 0.001,
  betas = c(0.9, 0.999),
  eps = 1e-08,
  weight_decay = 0.01,
  amsgrad = FALSE
)
```

# Arguments

params	(iterable): iterable of parameters to optimize or dicts defining parameter groups
lr	(float, optional): learning rate (default: 1e-3)
betas	(Tuple[float, float], optional): coefficients used for computing running averages of gradient and its square (default: $(0.9,0.999)$ )
eps	(float, optional): term added to the denominator to improve numerical stability (default: $1e-8$ )
weight_decay	(float, optional): weight decay (L2 penalty) (default: 0)
amsgrad	(boolean, optional): whether to use the AMSGrad variant of this algorithm from the paper On the Convergence of Adam and Beyond (default: FALSE)

optim_asgd	Averaged Stochastic Gradien	t Descent optimizer
-1		

# Description

Proposed in Acceleration of stochastic approximation by averaging

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#### Usage

```
optim_asgd(
  params,
  lr = 0.01,
  lambda = 1e-04,
  alpha = 0.75,
  t0 = 1e+06,
  weight_decay = 0
)
```

### **Arguments**

```
params (iterable): iterable of parameters to optimize or lists defining parameter groups

lr (float): learning rate

lambda (float, optional): decay term (default: 1e-4)

alpha (float, optional): power for eta update (default: 0.75)

t0 (float, optional): point at which to start averaging (default: 1e6)

weight_decay (float, optional): weight decay (L2 penalty) (default: 0)
```

#### Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

### **Examples**

```
if (torch_is_installed()) {
## Not run:
optimizer <- optim_asgd(model$parameters(), lr = 0.1)
optimizer$zero_grad()
loss_fn(model(input), target)$backward()
optimizer$step()
## End(Not run)
}</pre>
```

optim\_lbfgs

LBFGS optimizer

### **Description**

Implements L-BFGS algorithm, heavily inspired by minFunc

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#### Usage

```
optim_lbfgs(
  params,
  lr = 1,
  max_iter = 20,
  max_eval = NULL,
  tolerance_grad = 1e-07,
  tolerance_change = 1e-09,
  history_size = 100,
  line_search_fn = NULL
)
```

#### **Arguments**

```
params (iterable): iterable of parameters to optimize or dicts defining parameter groups

(float): learning rate (default: 1)

max_iter (int): maximal number of iterations per optimization step (default: 20)

max_eval (int): maximal number of function evaluations per optimization step (default: max_iter * 1.25).

tolerance_grad (float): termination tolerance on first order optimality (default: 1e-5).

tolerance_change (float): termination tolerance on function value/parameter changes (default: 1e-9).

history_size (int): update history size (default: 100).

line_search_fn (str): either 'strong_wolfe' or None (default: None).
```

#### **Details**

This optimizer is different from the others in that in optimizer\$step(), it needs to be passed a closure that (1) calculates the loss, (2) calls backward() on it, and (3) returns it. See example below.

#### Warning

This optimizer doesn't support per-parameter options and parameter groups (there can be only one).

Right now all parameters have to be on a single device. This will be improved in the future.

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

### Note

This is a very memory intensive optimizer (it requires additional param\_bytes \* (history\_size + 1) bytes). If it doesn't fit in memory try reducing the history size, or use a different algorithm.

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### **Examples**

```
if (torch_is_installed()) {
a <- 1
b <- 5
rosenbrock <- function(x) {</pre>
  x1 <- x[1]
  x2 <- x[2]
  (a - x1)^2 + b * (x2 - x1^2)^2
x <- torch_tensor(c(-1, 1), requires_grad = TRUE)</pre>
optimizer <- optim_lbfgs(x)</pre>
calc_loss <- function() {</pre>
  optimizer$zero_grad()
  value <- rosenbrock(x)</pre>
  value$backward()
  value
}
num_iterations <- 2</pre>
for (i in 1:num_iterations) {
  optimizer$step(calc_loss)
}
rosenbrock(x)
}
```

optim\_required

Dummy value indicating a required value.

# Description

export

### Usage

optim\_required()

optim\_rmsprop

RMSprop optimizer

# Description

Proposed by G. Hinton in his course.

optim\_rmsprop 309

#### Usage

```
optim_rmsprop(
  params,
  lr = 0.01,
  alpha = 0.99,
  eps = 1e-08,
  weight_decay = 0,
  momentum = 0,
  centered = FALSE
)
```

# **Arguments**

params (iterable): iterable of parameters to optimize or list defining parameter groups

1r (float, optional): learning rate (default: 1e-2)

alpha (float, optional): smoothing constant (default: 0.99)

eps (float, optional): term added to the denominator to improve numerical stability

(default: 1e-8)

weight\_decay optional weight decay penalty. (default: 0)
momentum (float, optional): momentum factor (default: 0)

centered (bool, optional): if TRUE, compute the centered RMSProp, the gradient is nor-

malized by an estimation of its variance weight decay (float, optional): weight

decay (L2 penalty) (default: 0)

#### Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

#### Note

The centered version first appears in Generating Sequences With Recurrent Neural Networks. The implementation here takes the square root of the gradient average before adding epsilon (note that TensorFlow interchanges these two operations). The effective learning rate is thus  $\alpha/(\sqrt{v}+\epsilon)$  where  $\alpha$  is the scheduled learning rate and v is the weighted moving average of the squared gradient.

Update rule:

$$\theta_{t+1} = \theta_t - \frac{\eta}{\sqrt{E[g^2]_t + \epsilon}} * g_t$$

310 optim\_rprop

optim_rprop	Implements the resilient backpropagation algorithm.	

### **Description**

Proposed first in RPROP - A Fast Adaptive Learning Algorithm

### Usage

```
optim_rprop(params, lr = 0.01, etas = c(0.5, 1.2), step_sizes = c(1e-06, 50))
```

# Arguments

params	(iterable): iterable of parameters to optimize or lists defining parameter groups
lr	(float, optional): learning rate (default: 1e-2)
etas	(Tuple(float, float), optional): pair of (etaminus, etaplis), that are multiplicative increase and decrease factors (default: $(0.5,1.2)$ )
step_sizes	(vector(float, float), optional): a pair of minimal and maximal allowed step sizes (default: (1e-6, 50))

# Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

```
if (torch_is_installed()) {
## Not run:
optimizer <- optim_rprop(model$parameters(), lr = 0.1)
optimizer$zero_grad()
loss_fn(model(input), target)$backward()
optimizer$step()
## End(Not run)
}</pre>
```

optim\_sgd 311

optim\_sgd SGD optimizer

### **Description**

Implements stochastic gradient descent (optionally with momentum). Nesterov momentum is based on the formula from On the importance of initialization and momentum in deep learning.

### Usage

```
optim_sgd(
  params,
  lr = optim_required(),
  momentum = 0,
  dampening = 0,
  weight_decay = 0,
  nesterov = FALSE
)
```

#### **Arguments**

params (iterable): iterable of parameters to optimize or dicts defining parameter groups

lr (float): learning rate

momentum (float, optional): momentum factor (default: 0)

dampening (float, optional): dampening for momentum (default: 0)

weight\_decay (float, optional): weight decay (L2 penalty) (default: 0)

nesterov (bool, optional): enables Nesterov momentum (default: FALSE)

#### Note

The implementation of SGD with Momentum-Nesterov subtly differs from Sutskever et. al. and implementations in some other frameworks.

Considering the specific case of Momentum, the update can be written as

$$v_{t+1} = \mu * v_t + g_{t+1},$$
  
 $p_{t+1} = p_t - \operatorname{lr} * v_{t+1},$ 

where p, g, v and  $\mu$  denote the parameters, gradient, velocity, and momentum respectively.

This is in contrast to Sutskever et. al. and other frameworks which employ an update of the form

$$v_{t+1} = \mu * v_t + \text{lr} * g_{t+1},$$
  
 $p_{t+1} = p_t - v_{t+1}.$ 

The Nesterov version is analogously modified.

312 sampler

#### Warning

If you need to move a model to GPU via \$cuda(), please do so before constructing optimizers for it. Parameters of a model after \$cuda() will be different objects from those before the call. In general, you should make sure that the objects pointed to by model parameters subject to optimization remain the same over the whole lifecycle of optimizer creation and usage.

### **Examples**

```
if (torch_is_installed()) {
## Not run:
optimizer <- optim_sgd(model$parameters(), lr = 0.1, momentum = 0.9)
optimizer$zero_grad()
loss_fn(model(input), target)$backward()
optimizer$step()
## End(Not run)
}</pre>
```

sampler

Creates a new Sampler

# Description

Samplers can be used with dataloader() when creating batches from a torch dataset().

### Usage

```
sampler(
  name = NULL,
  inherit = Sampler,
  ...,
  private = NULL,
  active = NULL,
  parent_env = parent.frame()
)
```

### Arguments

name	(optional) name of the sampler
inherit	(optional) you can inherit from other samplers to re-use some methods.
•••	Pass any number of fields or methods. You should at least define the initialize and step methods. See the examples section.
private	(optional) a list of private methods for the sampler
active	(optional) a list of active methods for the sampler.
parent_env	used to capture the right environment to define the class. The default is fine for most situations.

tensor\_dataset 313

#### **Details**

A sampler must implement the .iter and .length() methods.

- initialize takes in a data\_source. In general this is a dataset().
- .iter returns a function that returns a dataset index everytime it's called.
- .length returns the maximum number of samples that can be retrieved from that sampler.

tensor\_dataset

Dataset wrapping tensors.

### **Description**

Each sample will be retrieved by indexing tensors along the first dimension.

### Usage

```
tensor_dataset(...)
```

#### **Arguments**

tensors that have the same size of the first dimension.

threads

Number of threads

### **Description**

Get and set the numbers used by torch computations.

## Usage

```
torch_set_num_threads(num_threads)
torch_set_num_interop_threads(num_threads)
torch_get_num_interop_threads()
torch_get_num_threads()
```

#### **Arguments**

 $num\_threads \qquad number\ of\ threads\ to\ set.$ 

#### **Details**

For details see the CPU threading article in the PyTorch documentation.

314 torch\_absolute

### Note

torch\_set\_threads do not work on macOS system as it must be 1.

torch\_abs

Abs

# Description

Abs

# Usage

```
torch_abs(self)
```

# Arguments

self

(Tensor) the input tensor.

### abs(input) -> Tensor

Computes the element-wise absolute value of the given input tensor.

$$out_i = |input_i|$$

## **Examples**

```
if (torch_is_installed()) {
torch_abs(torch_tensor(c(-1, -2, 3)))
}
```

torch\_absolute

Absolute

# Description

Absolute

### Usage

```
torch_absolute(self)
```

# Arguments

self

(Tensor) the input tensor.

# absolute(input, \*, out=None) -> Tensor

Alias for torch\_abs()

torch\_acos 315

 $torch\_acos$ 

Acos

# Description

Acos

# Usage

```
torch_acos(self)
```

# Arguments

self

(Tensor) the input tensor.

# acos(input) -> Tensor

Returns a new tensor with the arccosine of the elements of input.

$$\operatorname{out}_i = \cos^{-1}(\operatorname{input}_i)$$

# Examples

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_acos(a)
}
```

 $torch\_acosh$ 

Acosh

# Description

Acosh

# Usage

```
torch_acosh(self)
```

# Arguments

self

(Tensor) the input tensor.

#### acosh(input, \*, out=None) -> Tensor

Returns a new tensor with the inverse hyperbolic cosine of the elements of input.

#### Note

The domain of the inverse hyperbolic cosine is [1, inf) and values outside this range will be mapped to NaN, except for + INF for which the output is mapped to + INF.

$$\mathsf{out}_i = \cosh^{-1}(\mathsf{input}_i)$$

### **Examples**

```
if (torch_is_installed()) {
a <- torch_randn(c(4))$uniform_(1, 2)
a
torch_acosh(a)
}</pre>
```

```
torch_adaptive_avg_pool1d
```

Adaptive\_avg\_pool1d

## Description

Adaptive\_avg\_pool1d

### Usage

```
torch_adaptive_avg_pool1d(self, output_size)
```

#### **Arguments**

self the input tensor

output\_size the target output size (single integer)

### adaptive\_avg\_pool1d(input, output\_size) -> Tensor

Applies a 1D adaptive average pooling over an input signal composed of several input planes.

See nn\_adaptive\_avg\_pool1d() for details and output shape.

torch\_add 317

torch\_add

Add

### **Description**

Add

#### Usage

```
torch_add(self, other, alpha = 1L)
```

### **Arguments**

self (Tensor) the input tensor.

other (Tensor/Number) the second input tensor/number.

alpha (Number) the scalar multiplier for other

#### add(input, other, out=NULL)

Adds the scalar other to each element of the input input and returns a new resulting tensor.

$$out = input + other$$

If input is of type FloatTensor or DoubleTensor, other must be a real number, otherwise it should be an integer.

### add(input, other, \*, alpha=1, out=NULL)

Each element of the tensor other is multiplied by the scalar alpha and added to each element of the tensor input. The resulting tensor is returned.

The shapes of input and other must be broadcastable.

$$out = input + alpha \times other$$

If other is of type FloatTensor or DoubleTensor, alpha must be a real number, otherwise it should be an integer.

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_add(a, 20)

a = torch_randn(c(4))
a
```

318 torch\_addbmm

```
b = torch_randn(c(4, 1))
b
torch_add(a, b)
}
```

torch\_addbmm

Addbmm

#### **Description**

Addbmm

#### Usage

```
torch_addbmm(self, batch1, batch2, beta = 1L, alpha = 1L)
```

### **Arguments**

self	(Tensor) matrix to be added
batch1	(Tensor) the first batch of matrices to be multiplied
batch2	(Tensor) the second batch of matrices to be multiplied
beta	(Number, optional) multiplier for input $(\beta)$
alpha	(Number, optional) multiplier for batch1 @ batch2 ( $\alpha$ )

### addbmm(input, batch1, batch2, \*, beta=1, alpha=1, out=NULL) -> Tensor

Performs a batch matrix-matrix product of matrices stored in batch1 and batch2, with a reduced add step (all matrix multiplications get accumulated along the first dimension). input is added to the final result.

batch1 and batch2 must be 3-D tensors each containing the same number of matrices.

If batch1 is a  $(b \times n \times m)$  tensor, batch2 is a  $(b \times m \times p)$  tensor, input must be broadcastable with a  $(n \times p)$  tensor and out will be a  $(n \times p)$  tensor.

$$out = \beta \text{ input} + \alpha \ (\sum_{i=0}^{b-1} \text{batch1}_i \ @ \text{ batch2}_i)$$

For inputs of type FloatTensor or DoubleTensor, arguments beta and alpha must be real numbers, otherwise they should be integers.

```
if (torch_is_installed()) {

M = torch_randn(c(3, 5))

batch1 = torch_randn(c(10, 3, 4))

batch2 = torch_randn(c(10, 4, 5))

torch_addbmm(M, batch1, batch2)
}
```

torch\_addcdiv 319

torch\_addcdiv

Addcdiv

#### Description

Addcdiv

### Usage

```
torch_addcdiv(self, tensor1, tensor2, value = 1L)
```

### **Arguments**

```
self (Tensor) the tensor to be added
tensor1 (Tensor) the numerator tensor
tensor2 (Tensor) the denominator tensor
value (Number, optional) multiplier for tensor1/tensor2
```

#### addcdiv(input, tensor1, tensor2, \*, value=1, out=NULL) -> Tensor

Performs the element-wise division of tensor1 by tensor2, multiply the result by the scalar value and add it to input.

### Warning

Integer division with addediv is deprecated, and in a future release addediv will perform a true division of tensor1 and tensor2. The current addediv behavior can be replicated using torch\_floor\_divide() for integral inputs (input + value \* tensor1 // tensor2) and torch\_div() for float inputs (input + value \* tensor1 // tensor2). The new addediv behavior can be implemented with torch\_true\_divide() (input + value \* torch.true\_divide(tensor1, tensor2).

$$\operatorname{out}_i = \operatorname{input}_i + \operatorname{value} \times \frac{\operatorname{tensor1}_i}{\operatorname{tensor2}_i}$$

The shapes of input, tensor1, and tensor2 must be broadcastable.

For inputs of type FloatTensor or DoubleTensor, value must be a real number, otherwise an integer.

```
if (torch_is_installed()) {
t = torch_randn(c(1, 3))
t1 = torch_randn(c(3, 1))
t2 = torch_randn(c(1, 3))
torch_addcdiv(t, t1, t2, 0.1)
}
```

320 torch\_addcmul

torch\_addcmul

Addcmul

## Description

Addcmul

### Usage

```
torch_addcmul(self, tensor1, tensor2, value = 1L)
```

### Arguments

```
self (Tensor) the tensor to be added

tensor1 (Tensor) the tensor to be multiplied

tensor2 (Tensor) the tensor to be multiplied

value (Number, optional) multiplier for tensor1. * tensor2
```

### addcmul(input, tensor1, tensor2, \*, value=1, out=NULL) -> Tensor

Performs the element-wise multiplication of tensor1 by tensor2, multiply the result by the scalar value and add it to input.

```
\operatorname{out}_i = \operatorname{input}_i + \operatorname{value} \times \operatorname{tensor1}_i \times \operatorname{tensor2}_i
```

The shapes of tensor, tensor1, and tensor2 must be broadcastable .

For inputs of type FloatTensor or DoubleTensor, value must be a real number, otherwise an integer.

```
if (torch_is_installed()) {
t = torch_randn(c(1, 3))
t1 = torch_randn(c(3, 1))
t2 = torch_randn(c(1, 3))
torch_addcmul(t, t1, t2, 0.1)
}
```

torch\_addmm 321

|--|

# Description

Addmm

# Usage

```
torch_addmm(self, mat1, mat2, beta = 1L, alpha = 1L)
```

#### **Arguments**

self	(Tensor) matrix to be added
mat1	(Tensor) the first matrix to be multiplied
mat2	(Tensor) the second matrix to be multiplied
beta	(Number, optional) multiplier for input $(\beta)$
alpha	(Number, optional) multiplier for $mat1@mat2$ ( $\alpha$ )

# addmm(input, mat1, mat2, \*, beta=1, alpha=1, out=NULL) -> Tensor

Performs a matrix multiplication of the matrices mat1 and mat2. The matrix input is added to the final result.

If mat1 is a  $(n \times m)$  tensor, mat2 is a  $(m \times p)$  tensor, then input must be broadcastable with a  $(n \times p)$  tensor and out will be a  $(n \times p)$  tensor.

alpha and beta are scaling factors on matrix-vector product between mat1 and mat2 and the added matrix input respectively.

```
out = \beta input + \alpha (mat1<sub>i</sub> @ mat2<sub>i</sub>)
```

For inputs of type FloatTensor or DoubleTensor, arguments beta and alpha must be real numbers, otherwise they should be integers.

```
if (torch_is_installed()) {

M = torch_randn(c(2, 3))

mat1 = torch_randn(c(2, 3))

mat2 = torch_randn(c(3, 3))

torch_addmm(M, mat1, mat2)
}
```

322 torch\_addmv

torch_addmv
-------------

Addmv

# Description

Addmv

# Usage

```
torch_addmv(self, mat, vec, beta = 1L, alpha = 1L)
```

### **Arguments**

self	(Tensor) vector to be added
mat	(Tensor) matrix to be multiplied
vec	(Tensor) vector to be multiplied
beta	(Number, optional) multiplier for input $(\beta)$
alpha	(Number, optional) multiplier for $mat@vec\left(\alpha\right)$

# addmv(input, mat, vec, \*, beta=1, alpha=1, out=NULL) -> Tensor

Performs a matrix-vector product of the matrix mat and the vector vec. The vector input is added to the final result.

If mat is a  $(n \times m)$  tensor, vec is a 1-D tensor of size m, then input must be broadcastable with a 1-D tensor of size n and out will be 1-D tensor of size n.

alpha and beta are scaling factors on matrix-vector product between mat and vec and the added tensor input respectively.

```
out = \beta input + \alpha (mat @ vec)
```

For inputs of type FloatTensor or DoubleTensor, arguments beta and alpha must be real numbers, otherwise they should be integers

```
if (torch_is_installed()) {

M = torch_randn(c(2))
mat = torch_randn(c(2, 3))
vec = torch_randn(c(3))
torch_addmv(M, mat, vec)
}
```

torch\_addr 323

torch_addr	Addr
------------	------

### **Description**

Addr

### Usage

```
torch_addr(self, vec1, vec2, beta = 1L, alpha = 1L)
```

#### **Arguments**

```
self (Tensor) matrix to be added vec1 (Tensor) the first vector of the outer product vec2 (Tensor) the second vector of the outer product beta (Number, optional) multiplier for input (\beta) alpha (Number, optional) multiplier for vec1 \otimes vec2 (\alpha)
```

#### addr(input, vec1, vec2, \*, beta=1, alpha=1, out=NULL) -> Tensor

Performs the outer-product of vectors vec1 and vec2 and adds it to the matrix input.

Optional values beta and alpha are scaling factors on the outer product between vec1 and vec2 and the added matrix input respectively.

```
out = \beta input + \alpha (vec1 \otimes vec2)
```

If vec1 is a vector of size n and vec2 is a vector of size m, then input must be broadcastable with a matrix of size  $(n \times m)$  and out will be a matrix of size  $(n \times m)$ .

For inputs of type FloatTensor or DoubleTensor, arguments beta and alpha must be real numbers, otherwise they should be integers

```
if (torch_is_installed()) {
  vec1 = torch_arange(1, 3)
  vec2 = torch_arange(1, 2)
  M = torch_zeros(c(3, 2))
  torch_addr(M, vec1, vec2)
}
```

324 torch\_allclose

### **Description**

Allclose

#### Usage

```
torch_allclose(self, other, rtol = 1e-05, atol = 1e-08, equal_nan = FALSE)
```

#### **Arguments**

self	(Tensor) first tensor to compare
other	(Tensor) second tensor to compare
rtol	(float, optional) relative tolerance. Default: 1e-05
atol	(float, optional) absolute tolerance. Default: 1e-08
equal_nan	(bool, optional) if TRUE, then two NaN s will be compared as equal. Default: $\ensuremath{FALSE}$

### allclose(input, other, rtol=1e-05, atol=1e-08, equal\_nan=False) -> bool

This function checks if all input and other satisfy the condition:

```
|input - other| \le atol + rtol \times |other|
```

elementwise, for all elements of input and other. The behaviour of this function is analogous to numpy.allclose <a href="https://docs.scipy.org/doc/numpy/reference/generated/numpy.allclose.html">numpy.allclose.html</a>.

```
if (torch_is_installed()) {

torch_allclose(torch_tensor(c(10000., 1e-07)), torch_tensor(c(10000.1, 1e-08)))

torch_allclose(torch_tensor(c(10000., 1e-08)), torch_tensor(c(10000.1, 1e-09)))

torch_allclose(torch_tensor(c(1.0, NaN)), torch_tensor(c(1.0, NaN)))

torch_allclose(torch_tensor(c(1.0, NaN)), torch_tensor(c(1.0, NaN)), equal_nan=TRUE)
}
```

torch\_amax 325

torch\_amax Amax

## Description

Amax

## Usage

```
torch_amax(self, dim = list(), keepdim = FALSE)
```

#### **Arguments**

self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

#### amax(input, dim, keepdim=FALSE, \*, out=None) -> Tensor

Returns the maximum value of each slice of the input tensor in the given dimension(s) dim.

#### Note

The difference between max/min and amax/amin is:

- amax/amin supports reducing on multiple dimensions,
- amax/amin does not return indices,
- amax/amin evenly distributes gradient between equal values, while max(dim)/min(dim) propagates gradient only to a single index in the source tensor.

If keepdim is TRUE, the output tensors are of the same size as inputexcept in the dimension(s)dimwhere the put.

```
if (torch_is_installed()) {
a <- torch_randn(c(4, 4))
a
torch_amax(a, 1)
}</pre>
```

326 torch\_amin

torch_amin	Amin
------------	------

## **Description**

Amin

#### Usage

```
torch_amin(self, dim = list(), keepdim = FALSE)
```

#### **Arguments**

```
self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

keepdim (bool) whether the output tensor has dim retained or not.
```

## amin(input, dim, keepdim=FALSE, \*, out=None) -> Tensor

Returns the minimum value of each slice of the input tensor in the given dimension(s) dim.

#### Note

The difference between max/min and amax/amin is:

- amax/amin supports reducing on multiple dimensions,
- amax/amin does not return indices,
- amax/amin evenly distributes gradient between equal values, while max(dim)/min(dim) propagates gradient only to a single index in the source tensor.

If keepdim is TRUE, the output tensors are of the same size as input except in the dimension(s) dim where they are of size 1. Otherwise, dims are squeezed (see torch\_squeeze()), resulting in the output tensors having fewer dimensions than input.

```
if (torch_is_installed()) {
a <- torch_randn(c(4, 4))
a
torch_amin(a, 1)
}</pre>
```

torch\_angle 327

torch\_angle

Angle

## Description

Angle

## Usage

```
torch_angle(self)
```

## Arguments

self

(Tensor) the input tensor.

## angle(input) -> Tensor

Computes the element-wise angle (in radians) of the given input tensor.

```
\operatorname{out}_i = \operatorname{angle}(\operatorname{input}_i)
```

## **Examples**

```
if (torch_is_installed()) {
## Not run:
torch_angle(torch_tensor(c(-1 + 1i, -2 + 2i, 3 - 3i)))*180/3.14159
## End(Not run)
}
```

torch\_arange

Arange

## Description

Arange

328 torch\_arange

#### Usage

```
torch_arange(
   start,
   end,
   step = 1L,
   dtype = NULL,
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

#### **Arguments**

start (Number) the starting value for the set of points. Default: 0.

end (Number) the ending value for the set of points

step (Number) the gap between each pair of adjacent points. Default: 1.

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type). If dtype is not given, infer the data type from the other input arguments. If any of start, end, or stop are floating-point, the dtype is inferred to be the default dtype, see ~torch.get\_default\_dtype. Otherwise, the dtype is inferred to be

torch.int64.

layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

## arange(start=0, end, step=1, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a 1-D tensor of size  $\left\lceil \frac{\text{end-start}}{\text{step}} \right\rceil$  with values from the interval [start, end) taken with common difference step beginning from start.

Note that non-integer step is subject to floating point rounding errors when comparing against end; to avoid inconsistency, we advise adding a small epsilon to end in such cases.

$$out_{i+1} = out_i + step$$

```
if (torch_is_installed()) {
torch_arange(start = 0, end = 5)
torch_arange(1, 4)
```

```
torch_arccos 329
```

```
torch_arange(1, 2.5, 0.5)
}
```

torch\_arccos

Arccos

## Description

Arccos

## Usage

```
torch_arccos(self)
```

## Arguments

self

(Tensor) the input tensor.

## arccos(input, \*, out=None) -> Tensor

Alias for torch\_acos().

torch\_arccosh

Arccosh

## Description

Arccosh

## Usage

```
torch_arccosh(self)
```

## Arguments

self

(Tensor) the input tensor.

## arccosh(input, \*, out=None) -> Tensor

Alias for torch\_acosh().

330 torch\_arcsinh

torch\_arcsin

Arcsin

## Description

Arcsin

## Usage

```
torch_arcsin(self)
```

## Arguments

self

(Tensor) the input tensor.

## arcsin(input, \*, out=None) -> Tensor

Alias for torch\_asin().

torch\_arcsinh

Arcsinh

## Description

Arcsinh

## Usage

```
torch_arcsinh(self)
```

## Arguments

self

(Tensor) the input tensor.

## arcsinh(input, \*, out=None) -> Tensor

Alias for torch\_asinh().

torch\_arctan 331

torch\_arctan

Arctan

## Description

Arctan

## Usage

```
torch_arctan(self)
```

## **Arguments**

self

(Tensor) the input tensor.

## arctan(input, \*, out=None) -> Tensor

Alias for torch\_atan().

torch\_arctanh

Arctanh

## Description

Arctanh

## Usage

```
torch_arctanh(self)
```

## Arguments

self

(Tensor) the input tensor.

## arctanh(input, \*, out=None) -> Tensor

Alias for torch\_atanh().

332 torch\_argmax

## Description

Argmax

## Arguments

self (Tensor) the input tensor.

dim (int) the dimension to reduce. If NULL, the argmax of the flattened input is returned.

keepdim (bool) whether the output tensor has dim retained or not. Ignored if dim=NULL.

## argmax(input) -> LongTensor

Returns the indices of the maximum value of all elements in the input tensor.

This is the second value returned by torch\_max. See its documentation for the exact semantics of this method.

#### argmax(input, dim, keepdim=False) -> LongTensor

Returns the indices of the maximum values of a tensor across a dimension.

This is the second value returned by torch\_max. See its documentation for the exact semantics of this method.

```
if (torch_is_installed()) {
## Not run:
a = torch_randn(c(4, 4))
a
torch_argmax(a)
## End(Not run)

a = torch_randn(c(4, 4))
a
torch_argmax(a, dim=1)
}
```

torch\_argmin 333

## Description

Argmin

## Arguments

self	(Tensor) the input tensor.
dim	(int) the dimension to reduce. If NULL, the argmin of the flattened input is returned.
keepdim	(bool) whether the output tensor has dim retained or not. Ignored if dim=NULL.

## argmin(input) -> LongTensor

Returns the indices of the minimum value of all elements in the input tensor.

This is the second value returned by torch\_min. See its documentation for the exact semantics of this method.

## argmin(input, dim, keepdim=False, out=NULL) -> LongTensor

Returns the indices of the minimum values of a tensor across a dimension.

This is the second value returned by torch\_min. See its documentation for the exact semantics of this method.

```
if (torch_is_installed()) {
a = torch_randn(c(4, 4))
a
torch_argmin(a)

a = torch_randn(c(4, 4))
a
torch_argmin(a, dim=1)
}
```

334 torch\_asin

torch\_argsort

Argsort

## **Description**

Argsort

#### Usage

```
torch_argsort(self, dim = -1L, descending = FALSE)
```

#### **Arguments**

self (Tensor) the input tensor.

dim (int, optional) the dimension to sort along

descending (bool, optional) controls the sorting order (ascending or descending)

#### argsort(input, dim=-1, descending=False) -> LongTensor

Returns the indices that sort a tensor along a given dimension in ascending order by value.

This is the second value returned by torch\_sort. See its documentation for the exact semantics of this method.

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4, 4))
a
torch_argsort(a, dim=1)
}
```

torch\_asin

Asin

## **Description**

Asin

## Usage

```
torch_asin(self)
```

## Arguments

self

(Tensor) the input tensor.

torch\_asinh 335

## asin(input, out=NULL) -> Tensor

Returns a new tensor with the arcsine of the elements of input.

$$\operatorname{out}_i = \sin^{-1}(\operatorname{input}_i)$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_asin(a)
}
```

torch\_asinh

As inh

## Description

Asinh

#### Usage

```
torch_asinh(self)
```

## Arguments

self

(Tensor) the input tensor.

## asinh(input, \*, out=None) -> Tensor

Returns a new tensor with the inverse hyperbolic sine of the elements of input.

$$\operatorname{out}_i = \sinh^{-1}(\operatorname{input}_i)$$

```
if (torch_is_installed()) {
a <- torch_randn(c(4))
a
torch_asinh(a)
}</pre>
```

336 torch\_as\_strided

### **Description**

As\_strided

#### Usage

```
torch_as_strided(self, size, stride, storage_offset = NULL)
```

## **Arguments**

```
self (Tensor) the input tensor.

size (tuple or ints) the shape of the output tensor

stride (tuple or ints) the stride of the output tensor

storage_offset (int, optional) the offset in the underlying storage of the output tensor
```

## as\_strided(input, size, stride, storage\_offset=0) -> Tensor

Create a view of an existing torch\_Tensor input with specified size, stride and storage\_offset.

#### Warning

More than one element of a created tensor may refer to a single memory location. As a result, inplace operations (especially ones that are vectorized) may result in incorrect behavior. If you need to write to the tensors, please clone them first.

```
Many PyTorch functions, which return a view of a tensor, are internally implemented with this function. Those functions, like `torch_Tensor.expand`, are easier to read and are therefore more advisable to use.
```

```
if (torch_is_installed()) {
x = torch_randn(c(3, 3))
x
t = torch_as_strided(x, list(2, 2), list(1, 2))
t
t = torch_as_strided(x, list(2, 2), list(1, 2), 1)
t
}
```

torch\_atan 337

torch\_atan

Atan

## Description

Atan

## Usage

```
torch_atan(self)
```

## Arguments

self

(Tensor) the input tensor.

## atan(input, out=NULL) -> Tensor

Returns a new tensor with the arctangent of the elements of input.

$$\operatorname{out}_i = \tan^{-1}(\operatorname{input}_i)$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_atan(a)
}
```

torch\_atan2

Atan2

## Description

Atan2

## Usage

```
torch_atan2(self, other)
```

## Arguments

self (Tensor) the first input tensor other (Tensor) the second input tensor 338 torch\_atanh

#### atan2(input, other, out=NULL) -> Tensor

Element-wise arctangent of  $\operatorname{input}_i/\operatorname{other}_i$  with consideration of the quadrant. Returns a new tensor with the signed angles in radians between vector  $(\operatorname{other}_i,\operatorname{input}_i)$  and vector (1,0). (Note that other, the second parameter, is the x-coordinate, while  $\operatorname{input}_i$ , the first parameter, is the y-coordinate.)

The shapes of input and other must be broadcastable.

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_atan2(a, torch_randn(c(4)))
}
```

torch\_atanh

Atanh

#### **Description**

Atanh

#### Usage

```
torch_atanh(self)
```

## **Arguments**

self

(Tensor) the input tensor.

#### atanh(input, \*, out=None) -> Tensor

Returns a new tensor with the inverse hyperbolic tangent of the elements of input.

## Note

The domain of the inverse hyperbolic tangent is (-1, 1) and values outside this range will be mapped to NaN, except for the values 1 and -1 for which the output is mapped to +/-INF respectively.

$$\operatorname{out}_i = \tanh^{-1}(\operatorname{input}_i)$$

torch\_atleast\_1d 339

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))$uniform_(-1, 1)
a
torch_atanh(a)
}
```

torch\_atleast\_1d

 $At least\_1d$ 

## Description

Returns a 1-dimensional view of each input tensor with zero dimensions. Input tensors with one or more dimensions are returned as-is.

## Usage

```
torch_atleast_1d(self)
```

## Arguments

self

(Tensor or list of Tensors)

```
if (torch_is_installed()) {

x <- torch_randn(c(2))
x
torch_atleast_1d(x)
x <- torch_tensor(1.)
x
torch_atleast_1d(x)
x <- torch_tensor(0.5)
y <- torch_tensor(1.)
torch_atleast_1d(list(x,y))
}</pre>
```

340 torch\_atleast\_3d

torch\_atleast\_2d

 $At least\_2d$ 

## Description

Returns a 2-dimensional view of each each input tensor with zero dimensions. Input tensors with two or more dimensions are returned as-is.

#### Usage

```
torch_atleast_2d(self)
```

#### **Arguments**

self

(Tensor or list of Tensors)

#### **Examples**

```
if (torch_is_installed()) {

x <- torch_tensor(1.)
x
torch_atleast_2d(x)
x <- torch_randn(c(2,2))
x
torch_atleast_2d(x)
x <- torch_tensor(0.5)
y <- torch_tensor(1.)
torch_atleast_2d(list(x,y))
}</pre>
```

torch\_atleast\_3d

Atleast\_3d

#### **Description**

Returns a 3-dimensional view of each each input tensor with zero dimensions. Input tensors with three or more dimensions are returned as-is.

## Usage

```
torch_atleast_3d(self)
```

#### **Arguments**

self

(Tensor or list of Tensors)

torch\_avg\_pool1d 341

## Description

Avg\_pool1d

#### Usage

```
torch_avg_pool1d(
  self,
  kernel_size,
  stride = list(),
  padding = 0L,
  ceil_mode = FALSE,
  count_include_pad = TRUE
)
```

#### **Arguments**

self input tensor of shape (minibatch, in\_channels, iW)

kernel\_size the size of the window. Can be a single number or a tuple (kW,)

stride the stride of the window. Can be a single number or a tuple (sW,). Default:

kernel\_size

padding implicit zero paddings on both sides of the input. Can be a single number or a

tuple (padW,). Default: 0

ceil\_mode when TRUE, will use ceil instead of floor to compute the output shape. De-

fault: FALSE

count\_include\_pad

when TRUE, will include the zero-padding in the averaging calculation. Default:

TRUE

## $avg\_pool1d(input, kernel\_size, stride=NULL, padding=0, ceil\_mode=FALSE, count\_include\_pad=TRUE) \\ -> Tensor$

Applies a 1D average pooling over an input signal composed of several input planes.

See nn\_avg\_pool1d() for details and output shape.

342 torch\_baddbmm

torch_baddbmm	Baddbmm

#### **Description**

Baddbmm

## Usage

```
torch_baddbmm(self, batch1, batch2, beta = 1L, alpha = 1L)
```

#### **Arguments**

self	(Tensor) the tensor to be added
batch1	(Tensor) the first batch of matrices to be multiplied
batch2	(Tensor) the second batch of matrices to be multiplied
beta	(Number, optional) multiplier for input ( $\beta$ )
alpha	(Number, optional) multiplier for batch1 @ batch2 ( $\alpha$ )

#### baddbmm(input, batch1, batch2, \*, beta=1, alpha=1, out=NULL) -> Tensor

Performs a batch matrix-matrix product of matrices in batch1 and batch2. input is added to the final result.

batch1 and batch2 must be 3-D tensors each containing the same number of matrices.

If batch1 is a  $(b \times n \times m)$  tensor, batch2 is a  $(b \times m \times p)$  tensor, then input must be broadcastable with a  $(b \times n \times p)$  tensor and out will be a  $(b \times n \times p)$  tensor. Both alpha and beta mean the same as the scaling factors used in torch\_addbmm.

```
\operatorname{out}_i = \beta \operatorname{input}_i + \alpha \operatorname{(batch1}_i \otimes \operatorname{batch2}_i)
```

For inputs of type FloatTensor or DoubleTensor, arguments beta and alpha must be real numbers, otherwise they should be integers.

```
if (torch_is_installed()) {

M = torch_randn(c(10, 3, 5))
batch1 = torch_randn(c(10, 3, 4))
batch2 = torch_randn(c(10, 4, 5))
torch_baddbmm(M, batch1, batch2)
}
```

torch\_bartlett\_window 343

torch\_bartlett\_window Bartlett window

#### **Description**

Bartlett\_window

## Usage

```
torch_bartlett_window(
  window_length,
  periodic = TRUE,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE
)
```

#### **Arguments**

window\_length (int) the size of returned window

periodic (bool, optional) If TRUE, returns a window to be used as periodic function. If

False, return a symmetric window.

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type). Only

floating point types are supported.

layout (torch.layout, optional) the desired layout of returned window tensor. Only

torch\_strided (dense layout) is supported.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

# bartlett\_window(window\_length, periodic=TRUE, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Bartlett window function.

$$w[n] = 1 - \left| \frac{2n}{N-1} - 1 \right| = \left\{ \begin{array}{ll} \frac{2n}{N-1} & \text{if } 0 \le n \le \frac{N-1}{2} \\ 2 - \frac{2n}{N-1} & \text{if } \frac{N-1}{2} < n < N \end{array} \right.,$$

where N is the full window size.

The input window\_length is a positive integer controlling the returned window size. periodic flag determines whether the returned window trims off the last duplicate value from the symmetric

344 torch\_bernoulli

window and is ready to be used as a periodic window with functions like torch\_stft. Therefore, if periodic is true, the N in above formula is in fact window\_length + 1. Also, we always have torch\_bartlett\_window(L, periodic=TRUE) equal to torch\_bartlett\_window(L + 1, periodic=False)[:-1]).

#### Note

If `window\_length` \eqn{=1}, the returned window contains a single value 1.

torch\_bernoulli

**Description** 

Bernoulli

## Usage

```
torch_bernoulli(self, p, generator = NULL)
```

Bernoulli

#### **Arguments**

self (Tensor) the input tensor of probability values for the Bernoulli distribution

p (Number) a probability value. If p is passed than it's used instead of the values in self tensor.

generator (torch.Generator, optional) a pseudorandom number generator for sampling

## bernoulli(input, \*, generator=NULL, out=NULL) -> Tensor

Draws binary random numbers (0 or 1) from a Bernoulli distribution.

The input tensor should be a tensor containing probabilities to be used for drawing the binary random number. Hence, all values in input have to be in the range:  $0 \le \text{input}_i \le 1$ .

The  $i^{th}$  element of the output tensor will draw a value 1 according to the  $i^{th}$  probability value given in input.

 $\operatorname{out}_i \sim \operatorname{Bernoulli}(p = \operatorname{input}_i)$ 

The returned out tensor only has values 0 or 1 and is of the same shape as input.

out can have integral dtype, but input must have floating point dtype.

torch\_bincount 345

#### **Examples**

```
if (torch_is_installed()) {
a = torch_empty(c(3, 3))$uniform_(0, 1) # generate a uniform random matrix with range c(0, 1)
a
torch_bernoulli(a)
a = torch_ones(c(3, 3)) # probability of drawing "1" is 1
torch_bernoulli(a)
a = torch_zeros(c(3, 3)) # probability of drawing "1" is 0
torch_bernoulli(a)
}
```

torch\_bincount

**Bincount** 

#### **Description**

Bincount

#### **Arguments**

self (Tensor) 1-d int tensor

weights (Tensor) optional, weight for each value in the input tensor. Should be of same size as input tensor.

minlength (int) optional, minimum number of bins. Should be non-negative.

#### bincount(input, weights=NULL, minlength=0) -> Tensor

Count the frequency of each value in an array of non-negative ints.

The number of bins (size 1) is one larger than the largest value in input unless input is empty, in which case the result is a tensor of size 0. If minlength is specified, the number of bins is at least minlength and if input is empty, then the result is tensor of size minlength filled with zeros. If n is the value at position i, out[n] += weights[i] if weights is specified else out[n] += 1.

```
.. include:: cuda_deterministic.rst
```

```
if (torch_is_installed()) {
input = torch_randint(1, 8, list(5), dtype=torch_int64())
weights = torch_linspace(0, 1, steps=5)
input
weights
torch_bincount(input, weights)
input$bincount(weights)
}
```

346 torch\_bitwise\_not

torch\_bitwise\_and

Bitwise\_and

#### **Description**

Bitwise\_and

#### Usage

```
torch_bitwise_and(self, other)
```

## Arguments

self NA the first input tensor other NA the second input tensor

#### bitwise\_and(input, other, out=NULL) -> Tensor

Computes the bitwise AND of input and other. The input tensor must be of integral or Boolean types. For bool tensors, it computes the logical AND.

torch\_bitwise\_not

Bitwise\_not

#### **Description**

Bitwise\_not

#### Usage

```
torch_bitwise_not(self)
```

#### **Arguments**

self

(Tensor) the input tensor.

## bitwise\_not(input, out=NULL) -> Tensor

Computes the bitwise NOT of the given input tensor. The input tensor must be of integral or Boolean types. For bool tensors, it computes the logical NOT.

torch\_bitwise\_or 347

torch\_bitwise\_or

Bitwise\_or

## Description

Bitwise\_or

#### Usage

```
torch_bitwise_or(self, other)
```

#### **Arguments**

self NA the first input tensor other NA the second input tensor

#### bitwise\_or(input, other, out=NULL) -> Tensor

Computes the bitwise OR of input and other. The input tensor must be of integral or Boolean types. For bool tensors, it computes the logical OR.

torch\_bitwise\_xor

Bitwise\_xor

#### **Description**

Bitwise\_xor

## Usage

```
torch_bitwise_xor(self, other)
```

## Arguments

self NA the first input tensor other NA the second input tensor

#### bitwise\_xor(input, other, out=NULL) -> Tensor

Computes the bitwise XOR of input and other. The input tensor must be of integral or Boolean types. For bool tensors, it computes the logical XOR.

torch\_blackman\_window Blackman\_window

#### **Description**

Blackman\_window

#### Usage

```
torch_blackman_window(
  window_length,
  periodic = TRUE,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE
)
```

#### **Arguments**

window\_length (int) the size of returned window

periodic (bool, optional) If TRUE, returns a window to be used as periodic function. If

False, return a symmetric window.

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type). Only

floating point types are supported.

layout (torch.layout, optional) the desired layout of returned window tensor. Only

torch\_strided (dense layout) is supported.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

## blackman\_window(window\_length, periodic=TRUE, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Blackman window function.

$$w[n] = 0.42 - 0.5\cos\left(\frac{2\pi n}{N-1}\right) + 0.08\cos\left(\frac{4\pi n}{N-1}\right)$$

where N is the full window size.

The input window\_length is a positive integer controlling the returned window size. periodic flag determines whether the returned window trims off the last duplicate value from the symmetric

torch\_block\_diag 349

window and is ready to be used as a periodic window with functions like torch\_stft. Therefore, if periodic is true, the N in above formula is in fact window\_length + 1. Also, we always have torch\_blackman\_window(L, periodic=TRUE) equal to torch\_blackman\_window(L + 1, periodic=False)[:-1]).

#### Note

If `window\_length` \eqn{=1}, the returned window contains a single value 1.

torch\_block\_diag

Block\_diag

#### **Description**

Create a block diagonal matrix from provided tensors.

#### Usage

```
torch_block_diag(tensors)
```

#### **Arguments**

tensors

(list of tensors) One or more tensors with 0, 1, or 2 dimensions.

## Examples

```
if (torch_is_installed()) {

A <- torch_tensor(rbind(c(0, 1), c(1, 0)))

B <- torch_tensor(rbind(c(3, 4, 5), c(6, 7, 8)))

C <- torch_tensor(7)

D <- torch_tensor(c(1, 2, 3))

E <- torch_tensor(rbind(4, 5, 6))

torch_block_diag(list(A, B, C, D, E))
}</pre>
```

torch\_bmm

Bmm

## Description

Bmm

## Usage

```
torch_bmm(self, mat2)
```

#### **Arguments**

```
self (Tensor) the first batch of matrices to be multiplied
mat2 (Tensor) the second batch of matrices to be multiplied
```

#### bmm(input, mat2, out=NULL) -> Tensor

Performs a batch matrix-matrix product of matrices stored in input and mat2.

input and mat2 must be 3-D tensors each containing the same number of matrices.

If input is a  $(b \times n \times m)$  tensor, mat 2 is a  $(b \times m \times p)$  tensor, out will be a  $(b \times n \times p)$  tensor.

$$\operatorname{out}_i = \operatorname{input}_i @ \operatorname{mat} 2_i$$

#### Note

This function does not broadcast . For broadcasting matrix products, see torch\_matmul.

#### **Examples**

```
if (torch_is_installed()) {
input = torch_randn(c(10, 3, 4))
mat2 = torch_randn(c(10, 4, 5))
res = torch_bmm(input, mat2)
res
}
```

torch\_broadcast\_tensors

Broadcast\_tensors

#### **Description**

Broadcast\_tensors

#### **Usage**

```
torch_broadcast_tensors(tensors)
```

## Arguments

tensors a list containing any number of tensors of the same type

## broadcast\_tensors(tensors) -> List of Tensors

Broadcasts the given tensors according to broadcasting-semantics.

torch\_bucketize 351

#### **Examples**

```
if (torch_is_installed()) {
x = torch_arange(0, 3)$view(c(1, 4))
y = torch_arange(0, 2)$view(c(3, 1))
out = torch_broadcast_tensors(list(x, y))
out[[1]]
}
```

torch\_bucketize

Bucketize

#### **Description**

Bucketize

#### Usage

```
torch_bucketize(self, boundaries, out_int32 = FALSE, right = FALSE)
```

#### **Arguments**

self (Tensor or Scalar) N-D tensor or a Scalar containing the search value(s).

(Tensor) 1-D tensor, must contain a monotonically increasing sequence.

(bool, optional) – indicate the output data type. torch\_int32() if True, torch\_int64()

otherwise. Default value is FALSE, i.e. default output data type is torch\_int64().

(bool, optional) – if False, return the first suitable location that is found. If

True, return the last such index. If no suitable index found, return 0 for nonnumerical value (eg. nan, inf) or the size of boundaries (one pass the last index).

In other words, if False, gets the lower bound index for each value in input from
boundaries. If True, gets the upper bound index instead. Default value is False.

## bucketize(input, boundaries, \*, out\_int32=FALSE, right=FALSE, out=None) -> Tensor

Returns the indices of the buckets to which each value in the input belongs, where the boundaries of the buckets are set by boundaries. Return a new tensor with the same size as input. If right is FALSE (default), then the left boundary is closed.

```
if (torch_is_installed()) {
boundaries <- torch_tensor(c(1, 3, 5, 7, 9))
boundaries
v <- torch_tensor(rbind(c(3, 6, 9), c(3, 6, 9)))
v
torch_bucketize(v, boundaries)
torch_bucketize(v, boundaries, right=TRUE)
}</pre>
```

352 torch\_cartesian\_prod

torch\_can\_cast

Can\_cast

## Description

Can\_cast

## Usage

```
torch_can_cast(from, to)
```

### **Arguments**

```
from (dtype) The original torch_dtype. to (dtype) The target torch_dtype.
```

#### can\_cast(from, to) -> bool

Determines if a type conversion is allowed under PyTorch casting rules described in the type promotion documentation .

#### **Examples**

```
if (torch_is_installed()) {
torch_can_cast(torch_double(), torch_float())
torch_can_cast(torch_float(), torch_int())
}
```

torch\_cartesian\_prod Cartesian\_prod

## Description

Do cartesian product of the given sequence of tensors.

## Usage

```
torch_cartesian_prod(tensors)
```

## Arguments

tensors

a list containing any number of 1 dimensional tensors.

torch\_cat 353

#### **Examples**

```
if (torch_is_installed()) {
a = c(1, 2, 3)
b = c(4, 5)
tensor_a = torch_tensor(a)
tensor_b = torch_tensor(b)
torch_cartesian_prod(list(tensor_a, tensor_b))
}
```

torch\_cat

Cat

#### **Description**

Cat

#### Usage

```
torch_cat(tensors, dim = 1L)
```

#### Arguments

tensors (sequence of Tensors) any python sequence of tensors of the same type. Nonempty tensors provided must have the same shape, except in the cat dimension. dim (int, optional) the dimension over which the tensors are concatenated

#### cat(tensors, dim=0, out=NULL) -> Tensor

Concatenates the given sequence of seq tensors in the given dimension. All tensors must either have the same shape (except in the concatenating dimension) or be empty.

```
torch_cat can be seen as an inverse operation for torch_split() and torch_chunk. torch_cat can be best understood via examples.
```

```
if (torch_is_installed()) {
x = torch_randn(c(2, 3))
x
torch_cat(list(x, x, x), 1)
torch_cat(list(x, x, x), 2)
}
```

354 torch\_ceil

torch\_cdist

Cdist

#### **Description**

Cdist

#### Usage

```
torch_cdist(x1, x2, p = 2L, compute_mode = NULL)
```

## **Arguments**

x1 (Tensor) input tensor of shape  $B \times P \times M$ . x2 (Tensor) input tensor of shape  $B \times R \times M$ .

p NA p value for the p-norm distance to calculate between each vector pair  $\in$ 

 $[0,\infty].$ 

compute\_mode NA 'use\_mm\_for\_euclid\_dist\_if\_necessary' - will use matrix multiplication ap-

proach to calculate euclidean distance (p = 2) if P > 25 or R > 25 'use\_mm\_for\_euclid\_dist'

- will always use matrix multiplication approach to calculate euclidean distance (p = 2) 'donot\_use\_mm\_for\_euclid\_dist' - will never use matrix multiplication

approach to calculate euclidean distance (p = 2) Default: use\_mm\_for\_euclid\_dist\_if\_necessary.

#### **TEST**

Computes batched the p-norm distance between each pair of the two collections of row vectors.

torch\_ceil Ceil

## Description

Ceil

## Usage

```
torch_ceil(self)
```

#### **Arguments**

self (Tensor) the input tensor.

torch\_celu 355

## ceil(input, out=NULL) -> Tensor

Returns a new tensor with the ceil of the elements of input, the smallest integer greater than or equal to each element.

$$\operatorname{out}_i = \lceil \operatorname{input}_i \rceil = \lfloor \operatorname{input}_i \rfloor + 1$$

## Examples

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_ceil(a)
}
```

torch\_celu

Celu

## Description

Celu

#### Usage

```
torch_celu(self, alpha = 1L)
```

## Arguments

self the input tensor

alpha the alpha value for the CELU formulation. Default: 1.0

## celu(input, alpha=1.) -> Tensor

See nnf\_celu() for more info.

356 torch\_chain\_matmul

torch\_celu\_

Celu\_

## Description

Celu\_

#### Usage

```
torch_celu_(self, alpha = 1L)
```

#### **Arguments**

self the input tensor

alpha the alpha value for the CELU formulation. Default: 1.0

#### celu\_(input, alpha=1.) -> Tensor

In-place version of torch\_celu().

torch\_chain\_matmul

Chain\_matmul

#### **Description**

Chain\_matmul

#### Usage

```
torch_chain_matmul(matrices)
```

## Arguments

matrices

(Tensors...) a sequence of 2 or more 2-D tensors whose product is to be determined.

#### **TEST**

Returns the matrix product of the N 2-D tensors. This product is efficiently computed using the matrix chain order algorithm which selects the order in which incurs the lowest cost in terms of arithmetic operations ([CLRS]\_). Note that since this is a function to compute the product, N needs to be greater than or equal to 2; if equal to 2 then a trivial matrix-matrix product is returned. If N is 1, then this is a no-op - the original matrix is returned as is.

torch\_channel\_shuffle 357

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3, 4))
b = torch_randn(c(4, 5))
c = torch_randn(c(5, 6))
d = torch_randn(c(6, 7))
torch_chain_matmul(list(a, b, c, d))
}
```

```
torch_channel_shuffle Channel_shuffle
```

## Description

Channel\_shuffle

## Usage

```
torch_channel_shuffle(self, groups)
```

#### **Arguments**

```
self (Tensor) the input tensor
groups (int) number of groups to divide channels in and rearrange.
```

## Divide the channels in a tensor of shape

```
math:(*, C , H, W):
```

Divide the channels in a tensor of shape (\*, C, H, W) into g groups and rearrange them as  $(*, C \stackrel{g}{,} g, H, W)$ , while keeping the original tensor shape.

```
if (torch_is_installed()) {
input <- torch_randn(c(1, 4, 2, 2))
print(input)
output <- torch_channel_shuffle(input, 2)
print(output)
}</pre>
```

358 torch\_cholesky

torch\_cholesky

Cholesky

### **Description**

Cholesky

#### Usage

```
torch_cholesky(self, upper = FALSE)
```

#### **Arguments**

self (Tensor) the input tensor A of size (\*, n, n) where \* is zero or more batch di-

mensions consisting of symmetric positive-definite matrices.

upper (bool, optional) flag that indicates whether to return a upper or lower triangular

matrix. Default: FALSE

#### cholesky(input, upper=False, out=NULL) -> Tensor

Computes the Cholesky decomposition of a symmetric positive-definite matrix A or for batches of symmetric positive-definite matrices.

If upper is TRUE, the returned matrix U is upper-triangular, and the decomposition has the form:

$$A = U^T U$$

If upper is FALSE, the returned matrix L is lower-triangular, and the decomposition has the form:

$$A = LL^T$$

If upper is TRUE, and A is a batch of symmetric positive-definite matrices, then the returned tensor will be composed of upper-triangular Cholesky factors of each of the individual matrices. Similarly, when upper is FALSE, the returned tensor will be composed of lower-triangular Cholesky factors of each of the individual matrices.

```
if (torch_is_installed()) {
a = torch_randn(c(3, 3))
a = torch_mm(a, a$t()) # make symmetric positive-definite
l = torch_cholesky(a)
a
l
torch_mm(l, l$t())
a = torch_randn(c(3, 2, 2))
## Not run:
a = torch_matmul(a, a$transpose(-1, -2)) + 1e-03 # make symmetric positive-definite
```

torch\_cholesky\_inverse 359

```
1 = torch_cholesky(a)
z = torch_matmul(1, l$transpose(-1, -2))
torch_max(torch_abs(z - a)) # Max non-zero
## End(Not run)
}
```

torch\_cholesky\_inverse

Cholesky\_inverse

#### **Description**

Cholesky\_inverse

#### Usage

```
torch_cholesky_inverse(self, upper = FALSE)
```

## Arguments

self (Tensor) the input 2-D tensor u, a upper or lower triangular Cholesky factor upper (bool, optional) whether to return a lower (default) or upper triangular matrix

## cholesky\_inverse(input, upper=False, out=NULL) -> Tensor

Computes the inverse of a symmetric positive-definite matrix A using its Cholesky factor u: returns matrix inv. The inverse is computed using LAPACK routines dpotri and spotri (and the corresponding MAGMA routines).

If upper is FALSE, u is lower triangular such that the returned tensor is

$$inv = (uu^T)^{-1}$$

If upper is TRUE or not provided, u is upper triangular such that the returned tensor is

$$inv = (u^T u)^{-1}$$

```
if (torch_is_installed()) {
## Not run:
a = torch_randn(c(3, 3))
a = torch_mm(a, a$t()) + 1e-05 * torch_eye(3) # make symmetric positive definite
u = torch_cholesky(a)
a
torch_cholesky_inverse(u)
a$inverse()
```

360 torch\_cholesky\_solve

```
## End(Not run)
}
```

## Description

Cholesky\_solve

#### Usage

```
torch_cholesky_solve(self, input2, upper = FALSE)
```

## Arguments

self	(Tensor) input matrix $b$ of size $(\ast,m,k),$ where $\ast$ is zero or more batch dimensions
input2	(Tensor) input matrix $u$ of size $(*,m,m)$ , where $*$ is zero of more batch dimensions composed of upper or lower triangular Cholesky factor
upper	(bool, optional) whether to consider the Cholesky factor as a lower or upper

## cholesky\_solve(input, input2, upper=False, out=NULL) -> Tensor

Solves a linear system of equations with a positive semidefinite matrix to be inverted given its Cholesky factor matrix u.

If upper is FALSE, u is and lower triangular and c is returned such that:

triangular matrix. Default: FALSE.

$$c = (uu^T)^{-1}b$$

If upper is TRUE or not provided, u is upper triangular and c is returned such that:

$$c = (u^T u)^{-1} b$$

 $torch\_cholesky\_solve(b, u)$  can take in 2D inputs b, u or inputs that are batches of 2D matrices. If the inputs are batches, then returns batched outputs c

```
if (torch_is_installed()) {
a = torch_randn(c(3, 3))
a = torch_mm(a, a$t()) # make symmetric positive definite
u = torch_cholesky(a)
a
b = torch_randn(c(3, 2))
```

torch\_chunk 361

```
b
torch_cholesky_solve(b, u)
torch_mm(a$inverse(), b)
}
```

torch\_chunk

Chunk

#### **Description**

Chunk

## Usage

```
torch_chunk(self, chunks, dim = 1L)
```

# **Arguments**

self (Tensor) the tensor to split

chunks (int) number of chunks to return

dim (int) dimension along which to split the tensor

# chunk(input, chunks, dim=0) -> List of Tensors

Splits a tensor into a specific number of chunks. Each chunk is a view of the input tensor.

Last chunk will be smaller if the tensor size along the given dimension dim is not divisible by chunks.

torch\_clamp

Clamp

## **Description**

Clamp

## Usage

```
torch_clamp(self, min = NULL, max = NULL)
```

#### **Arguments**

self (Tensor) the input tensor.

min (Number) lower-bound of the range to be clamped to max (Number) upper-bound of the range to be clamped to

362 torch\_clip

#### clamp(input, min, max, out=NULL) -> Tensor

Clamp all elements in input into the range [ min, max ] and return a resulting tensor:

$$y_i = \left\{ \begin{array}{ll} \min & \text{if } x_i < \min \\ x_i & \text{if } \min \leq x_i \leq \max \\ \max & \text{if } x_i > \max \end{array} \right.$$

If input is of type FloatTensor or DoubleTensor, args min and max must be real numbers, otherwise they should be integers.

#### clamp(input, \*, min, out=NULL) -> Tensor

Clamps all elements in input to be larger or equal min.

If input is of type FloatTensor or DoubleTensor, value should be a real number, otherwise it should be an integer.

#### clamp(input, \*, max, out=NULL) -> Tensor

Clamps all elements in input to be smaller or equal max.

If input is of type FloatTensor or DoubleTensor, value should be a real number, otherwise it should be an integer.

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_clamp(a, min=-0.5, max=0.5)

a = torch_randn(c(4))
a
torch_clamp(a, min=0.5)

a = torch_randn(c(4))
a
torch_clamp(a, max=0.5)
}
```

torch\_clip

Clip

## **Description**

Clip

torch\_clone 363

#### Usage

```
torch_clip(self, min = NULL, max = NULL)
```

#### **Arguments**

self (Tensor) the input tensor.

min (Number) lower-bound of the range to be clamped to
max (Number) upper-bound of the range to be clamped to

# clip(input, min, max, \*, out=None) -> Tensor

Alias for torch\_clamp().

torch\_clone

Clone

#### **Description**

Clone

#### Usage

```
torch_clone(self, memory_format = NULL)
```

# **Arguments**

```
self (Tensor) the input tensor.
memory_format a torch memory format. see torch_preserve_format().
```

# clone(input, \*, memory\_format=torch.preserve\_format) -> Tensor

Returns a copy of input.

#### Note

This function is differentiable, so gradients will flow back from the result of this operation to input. To create a tensor without an autograd relationship to input see Tensor\$detach.

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torch\_combinations

**Combinations** 

## Description

Combinations

#### Usage

```
torch_combinations(self, r = 2L, with_replacement = FALSE)
```

#### **Arguments**

```
self (Tensor) 1D vector.

r (int, optional) number of elements to combine
with_replacement
(boolean, optional) whether to allow duplication in combination
```

#### combinations(input, r=2, with\_replacement=False) -> seq

Compute combinations of length r of the given tensor. The behavior is similar to python's itertools.combinations when with\_replacement is set to False, and itertools.combinations\_with\_replacement when with\_replacement is set to TRUE.

#### **Examples**

```
if (torch_is_installed()) {
a = c(1, 2, 3)
tensor_a = torch_tensor(a)
torch_combinations(tensor_a)
torch_combinations(tensor_a, r=3)
torch_combinations(tensor_a, with_replacement=TRUE)
}
```

torch\_complex

Complex

## **Description**

Complex

#### Usage

```
torch_complex(real, imag)
```

torch\_conj 365

## Arguments

```
real (Tensor) The real part of the complex tensor. Must be float or double.
```

imag (Tensor) The imaginary part of the complex tensor. Must be same dtype as real.

# complex(real, imag, \*, out=None) -> Tensor

Constructs a complex tensor with its real part equal to real and its imaginary part equal to imag.

#### **Examples**

```
if (torch_is_installed()) {
real <- torch_tensor(c(1, 2), dtype=torch_float32())
imag <- torch_tensor(c(3, 4), dtype=torch_float32())
z <- torch_complex(real, imag)
z
z$dtype
}</pre>
```

torch\_conj

Conj

#### **Description**

Conj

#### Usage

```
torch_conj(self)
```

#### **Arguments**

self

(Tensor) the input tensor.

#### conj(input) -> Tensor

Computes the element-wise conjugate of the given input tensor.

```
out_i = conj(input_i)
```

```
if (torch_is_installed()) {
## Not run:
torch_conj(torch_tensor(c(-1 + 1i, -2 + 2i, 3 - 3i)))
## End(Not run)
}
```

366 torch\_conv1d

torch\_conv1d

Conv1d

#### Description

Conv1d

#### Usage

```
torch_conv1d(
  input,
  weight,
  bias = list(),
  stride = 1L,
  padding = 0L,
  dilation = 1L,
  groups = 1L
)
```

#### Arguments

```
input tensor of shape (minibatch, in_channels, iW)
input
                   filters of shape (out_channels, \frac{\text{in\_channels}}{\text{groups}}, kW)
weight
bias
                   optional bias of shape (out_channels). Default: NULL
                   the stride of the convolving kernel. Can be a single number or a one-element
stride
                   tuple (sW,). Default: 1
                   implicit paddings on both sides of the input. Can be a single number or a one-
padding
                   element tuple (padW,). Default: 0
dilation
                   the spacing between kernel elements. Can be a single number or a one-element
                   tuple (dW,). Default: 1
                   split input into groups, in_channels should be divisible by the number of groups.
groups
                   Default: 1
```

#### conv1d(input, weight, bias=NULL, stride=1, padding=0, dilation=1, groups=1) -> Tensor

Applies a 1D convolution over an input signal composed of several input planes.

See nn\_conv1d() for details and output shape.

```
if (torch_is_installed()) {
filters = torch_randn(c(33, 16, 3))
inputs = torch_randn(c(20, 16, 50))
nnf_conv1d(inputs, filters)
}
```

torch\_conv2d 367

torch\_conv2d Conv2d

#### **Description**

Conv2d

#### Usage

```
torch_conv2d(
  input,
  weight,
  bias = list(),
  stride = 1L,
  padding = 0L,
  dilation = 1L,
  groups = 1L
)
```

#### Arguments

```
input tensor of shape (minibatch, in_channels, iH, iW)
input
                    \text{filters of shape (out\_channels, } \\ \\ \frac{\text{in\_channels}}{\text{groups}}, kH, kW)
weight
bias
                    optional bias tensor of shape (out_channels). Default: NULL
stride
                    the stride of the convolving kernel. Can be a single number or a tuple (sH, sW).
                    Default: 1
padding
                    implicit paddings on both sides of the input. Can be a single number or a tuple
                    (padH, padW). Default: 0
dilation
                    the spacing between kernel elements. Can be a single number or a tuple (dH, dW).
                    Default: 1
                    split input into groups, in_channels should be divisible by the number of groups.
groups
                    Default: 1
```

# conv2d(input, weight, bias=NULL, stride=1, padding=0, dilation=1, groups=1) -> Tensor

Applies a 2D convolution over an input image composed of several input planes. See nn\_conv2d() for details and output shape.

```
if (torch_is_installed()) {
# With square kernels and equal stride
filters = torch_randn(c(8,4,3,3))
inputs = torch_randn(c(1,4,5,5))
nnf_conv2d(inputs, filters, padding=1)
}
```

368 torch\_conv3d

torch\_conv3d

Conv3d

#### **Description**

Conv3d

#### Usage

```
torch_conv3d(
  input,
  weight,
  bias = list(),
  stride = 1L,
  padding = 0L,
  dilation = 1L,
  groups = 1L
)
```

#### **Arguments**

```
input tensor of shape (minibatch, in_channels, iT, iH, iW)
input
                   filters of shape (out_channels, \frac{\text{in\_channels}}{\text{groups}}, kT, kH, kW)
weight
bias
                   optional bias tensor of shape (out channels). Default: NULL
stride
                   the stride of the convolving kernel. Can be a single number or a tuple (sT, sH, sW).
                   Default: 1
                   implicit paddings on both sides of the input. Can be a single number or a tuple
padding
                    (padT, padH, padW). Default: 0
dilation
                   the spacing between kernel elements. Can be a single number or a tuple (dT, dH, dW).
                   Default: 1
groups
                   split input into groups, in_channels should be divisible by the number of groups.
                   Default: 1
```

#### conv3d(input, weight, bias=NULL, stride=1, padding=0, dilation=1, groups=1) -> Tensor

Applies a 3D convolution over an input image composed of several input planes.

See nn\_conv3d() for details and output shape.

```
if (torch_is_installed()) {
# filters = torch_randn(c(33, 16, 3, 3, 3))
# inputs = torch_randn(c(20, 16, 50, 10, 20))
# nnf_conv3d(inputs, filters)
}
```

torch\_conv\_tbc 369

torch\_conv\_tbc

Conv\_tbc

#### **Description**

Conv\_tbc

#### Usage

```
torch_conv_tbc(self, weight, bias, pad = 0L)
```

#### **Arguments**

pad

self NA input tensor of shape (sequence length  $\times$   $batch \times$  in\_channels) weight NA filter of shape (kernel width  $\times$  in\_channels  $\times$  out\_channels) bias NA bias of shape (out\_channels)

NA number of timesteps to pad. Default: 0

# TEST

Applies a 1-dimensional sequence convolution over an input sequence. Input and output dimensions are (Time, Batch, Channels) - hence TBC.

```
torch_conv_transpose1d
```

Conv\_transpose1d

# Description

Conv\_transpose1d

#### Usage

```
torch_conv_transpose1d(
  input,
  weight,
  bias = list(),
  stride = 1L,
  padding = 0L,
  output_padding = 0L,
  groups = 1L,
  dilation = 1L
)
```

#### **Arguments**

input	input tensor of shape (minibatch, in_channels, $iW$ )
weight	filters of shape (in_channels, $\frac{\text{out\_channels}}{\text{groups}}, kW$ )
bias	optional bias of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a tuple ( $sW$ ,). Default: $1$
padding	dilation $\star$ (kernel_size - 1) - padding zero-padding will be added to both sides of each dimension in the input. Can be a single number or a tuple (padW,). Default: 0
output_padding	additional size added to one side of each dimension in the output shape. Can be a single number or a tuple (out_padW). Default: $0$
groups	split input into groups, in_channels should be divisible by the number of groups. Default: 1
dilation	the spacing between kernel elements. Can be a single number or a tuple $(dW,)$ . Default: 1

# $conv\_transpose1d(input, weight, bias=NULL, stride=1, padding=0, output\_padding=0, groups=1, dilation=1) -> Tensor$

Applies a 1D transposed convolution operator over an input signal composed of several input planes, sometimes also called "deconvolution".

See nn\_conv\_transpose1d() for details and output shape.

# **Examples**

```
if (torch_is_installed()) {
inputs = torch_randn(c(20, 16, 50))
weights = torch_randn(c(16, 33, 5))
nnf_conv_transpose1d(inputs, weights)
}
```

torch\_conv\_transpose2d

Conv\_transpose2d

# Description

Conv\_transpose2d

## Usage

```
torch_conv_transpose2d(
  input,
  weight,
  bias = list(),
  stride = 1L,
  padding = 0L,
  output_padding = 0L,
  groups = 1L,
  dilation = 1L
)
```

#### **Arguments**

input	input tensor of shape (minibatch, in_channels, $iH$ , $iW$ )
weight	filters of shape (in_channels, $\frac{\text{out\_channels}}{\text{groups}}, kH, kW$ )
bias	optional bias of shape (out_channels). Default: NULL
stride	the stride of the convolving kernel. Can be a single number or a tuple (sH, sW). Default: 1
padding	dilation * (kernel_size - 1) - padding zero-padding will be added to both sides of each dimension in the input. Can be a single number or a tuple (padH, padW). Default: 0
output_padding	additional size added to one side of each dimension in the output shape. Can be a single number or a tuple (out_padH, out_padW). Default: 0
groups	split input into groups, in_channels should be divisible by the number of groups. Default: 1
dilation	the spacing between kernel elements. Can be a single number or a tuple (dH, dW). Default: 1

# $conv\_transpose2d(input, weight, bias=NULL, stride=1, padding=0, output\_padding=0, groups=1, dilation=1) -> Tensor$

Applies a 2D transposed convolution operator over an input image composed of several input planes, sometimes also called "deconvolution".

See nn\_conv\_transpose2d() for details and output shape.

```
if (torch_is_installed()) {

# With square kernels and equal stride
inputs = torch_randn(c(1, 4, 5, 5))
weights = torch_randn(c(4, 8, 3, 3))
nnf_conv_transpose2d(inputs, weights, padding=1)
}
```

```
torch_conv_transpose3d
```

Conv\_transpose3d

#### **Description**

Conv\_transpose3d

#### Usage

```
torch_conv_transpose3d(
  input,
  weight,
  bias = list(),
  stride = 1L,
  padding = 0L,
  output_padding = 0L,
  groups = 1L,
  dilation = 1L
)
```

## **Arguments**

```
input tensor of shape (minibatch, in_channels, iT, iH, iW)
input
                   \text{filters of shape (in\_channels,} \frac{\text{out\_channels}}{\text{groups}}, kT, kH, kW)
weight
bias
                   optional bias of shape (out_channels). Default: NULL
                   the stride of the convolving kernel. Can be a single number or a tuple (sT, sH, sW).
stride
                   Default: 1
padding
                   dilation * (kernel_size - 1) - padding zero-padding will be added to both
                   sides of each dimension in the input. Can be a single number or a tuple (padT, padH, padW).
                   Default: 0
                  additional size added to one side of each dimension in the output shape. Can be
output_padding
                   a single number or a tuple (out_padT, out_padH, out_padW). Default: 0
                   split input into groups, in channels should be divisible by the number of groups.
groups
                   Default: 1
dilation
                   the spacing between kernel elements. Can be a single number or a tuple (dT, dH, dW).
                   Default: 1
```

# conv\_transpose3d(input, weight, bias=NULL, stride=1, padding=0, output\_padding=0, groups=1, dilation=1) -> Tensor

Applies a 3D transposed convolution operator over an input image composed of several input planes, sometimes also called "deconvolution"

See nn\_conv\_transpose3d() for details and output shape.

torch\_cos 373

# **Examples**

```
if (torch_is_installed()) {
## Not run:
inputs = torch_randn(c(20, 16, 50, 10, 20))
weights = torch_randn(c(16, 33, 3, 3, 3))
nnf_conv_transpose3d(inputs, weights)
## End(Not run)
}
```

torch\_cos

Cos

# Description

Cos

#### Usage

```
torch_cos(self)
```

# Arguments

self

(Tensor) the input tensor.

# cos(input, out=NULL) -> Tensor

Returns a new tensor with the cosine of the elements of input.

```
\operatorname{out}_i = \cos(\operatorname{input}_i)
```

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_cos(a)
}
```

torch\_cosh

Cosh

# Description

Cosh

#### Usage

```
torch_cosh(self)
```

# Arguments

self

(Tensor) the input tensor.

# cosh(input, out=NULL) -> Tensor

Returns a new tensor with the hyperbolic cosine of the elements of input.

$$\operatorname{out}_i = \cosh(\operatorname{input}_i)$$

# Examples

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_cosh(a)
}
```

torch\_cosine\_similarity

Cosine\_similarity

# Description

Cosine\_similarity

# Usage

```
torch_cosine_similarity(x1, x2, dim = 2L, eps = 1e-08)
```

torch\_count\_nonzero 375

## **Arguments**

x1	(Tensor) First input.
x2	(Tensor) Second input (of size matching x1).
dim	(int, optional) Dimension of vectors. Default: 1
eps	(float, optional) Small value to avoid division by zero. Default: 1e-8

# cosine\_similarity(x1, x2, dim=1, eps=1e-8) -> Tensor

Returns cosine similarity between x1 and x2, computed along dim.

$$\text{similarity} = \frac{x_1 \cdot x_2}{\max(\|x_1\|_2 \cdot \|x_2\|_2, \epsilon)}$$

#### **Examples**

```
if (torch_is_installed()) {
input1 = torch_randn(c(100, 128))
input2 = torch_randn(c(100, 128))
output = torch_cosine_similarity(input1, input2)
output
}
```

torch\_count\_nonzero Count\_nonzero

#### **Description**

Count\_nonzero

#### Usage

```
torch_count_nonzero(self, dim = NULL)
```

# Arguments

self (Tensor) the input tensor.

dim (int or tuple of ints, optional) Dim or tuple of dims along which to count non-zeros.

# count\_nonzero(input, dim=None) -> Tensor

Counts the number of non-zero values in the tensor input along the given dim. If no dim is specified then all non-zeros in the tensor are counted.

376 torch\_cross

#### **Examples**

```
if (torch_is_installed()) {
x <- torch_zeros(3,3)
x[torch_randn(3,3) > 0.5] = 1
x
torch_count_nonzero(x)
torch_count_nonzero(x, dim=1)
}
```

torch\_cross

Cross

#### **Description**

Cross

#### Usage

```
torch_cross(self, other, dim = NULL)
```

# Arguments

```
self (Tensor) the input tensor.

other (Tensor) the second input tensor

dim (int, optional) the dimension to take the cross-product in.
```

#### cross(input, other, dim=-1, out=NULL) -> Tensor

Returns the cross product of vectors in dimension dim of input and other.

input and other must have the same size, and the size of their dim dimension should be 3.

If dim is not given, it defaults to the first dimension found with the size 3.

```
if (torch_is_installed()) {
a = torch_randn(c(4, 3))
a
b = torch_randn(c(4, 3))
b
torch_cross(a, b, dim=2)
torch_cross(a, b)
}
```

torch\_cummax 377

torch\_cummax

**Cummax** 

# Description

Cummax

#### Usage

```
torch_cummax(self, dim)
```

## **Arguments**

self (Tensor) the input tensor.

dim (int) the dimension to do the operation over

# cummax(input, dim) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the cumulative maximum of elements of input in the dimension dim. And indices is the index location of each maximum value found in the dimension dim.

$$y_i = max(x_1, x_2, x_3, \dots, x_i)$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(10))
a
torch_cummax(a, dim=1)
}
```

torch\_cummin

Cummin

# Description

Cummin

## Usage

```
torch_cummin(self, dim)
```

378 torch\_cumprod

#### **Arguments**

self (Tensor) the input tensor.

dim (int) the dimension to do the operation over

#### cummin(input, dim) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the cumulative minimum of elements of input in the dimension dim. And indices is the index location of each maximum value found in the dimension dim.

$$y_i = min(x_1, x_2, x_3, \dots, x_i)$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(10))
a
torch_cummin(a, dim=1)
}
```

torch\_cumprod

Cumprod

## **Description**

Cumprod

# Usage

```
torch_cumprod(self, dim, dtype = NULL)
```

#### **Arguments**

self (Tensor) the input tensor.

dim (int) the dimension to do the operation over

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

#### cumprod(input, dim, out=NULL, dtype=NULL) -> Tensor

Returns the cumulative product of elements of input in the dimension dim.

For example, if input is a vector of size N, the result will also be a vector of size N, with elements.

$$y_i = x_1 \times x_2 \times x_3 \times \cdots \times x_i$$

torch\_cumsum 379

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(10))
a
torch_cumprod(a, dim=1)
}
```

torch\_cumsum

Cumsum

## **Description**

Cumsum

## Usage

```
torch_cumsum(self, dim, dtype = NULL)
```

#### Arguments

self (Tensor) the input tensor.

dim (int) the dimension to do the operation over

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

## cumsum(input, dim, out=NULL, dtype=NULL) -> Tensor

Returns the cumulative sum of elements of input in the dimension dim.

For example, if input is a vector of size N, the result will also be a vector of size N, with elements.

$$y_i = x_1 + x_2 + x_3 + \dots + x_i$$

```
if (torch_is_installed()) {
a = torch_randn(c(10))
a
torch_cumsum(a, dim=1)
}
```

380 torch\_dequantize

torch\_deg2rad

Deg2rad

#### **Description**

Deg2rad

# Usage

```
torch_deg2rad(self)
```

#### **Arguments**

self

(Tensor) the input tensor.

# deg2rad(input, \*, out=None) -> Tensor

Returns a new tensor with each of the elements of input converted from angles in degrees to radians.

# **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(rbind(c(180.0, -180.0), c(360.0, -360.0), c(90.0, -90.0)))
torch_deg2rad(a)
}</pre>
```

torch\_dequantize

Dequantize

#### **Description**

Dequantize

# Usage

```
torch_dequantize(tensor)
```

# Arguments

tensor

(Tensor) A quantized Tensor or a list oof quantized tensors

# dequantize(tensor) -> Tensor

Returns an fp32 Tensor by dequantizing a quantized Tensor

torch\_det 381

#### dequantize(tensors) -> sequence of Tensors

Given a list of quantized Tensors, dequantize them and return a list of fp32 Tensors

torch\_det Det

# **Description**

Det

#### Usage

```
torch_det(self)
```

#### **Arguments**

self

(Tensor) the input tensor of size (\*, n, n) where \* is zero or more batch dimensions.

# det(input) -> Tensor

Calculates determinant of a square matrix or batches of square matrices.

#### Note

Backward through `det` internally uses SVD results when `input` is not invertible. In this case, double backward through `det` will be unstable in when `input` doesn't have distinct singular values. See `~torch.svd` for details.

```
if (torch_is_installed()) {
A = torch_randn(c(3, 3))
torch_det(A)
A = torch_randn(c(3, 2, 2))
A
A$det()
}
```

382 torch\_diag

torch\_device

Create a Device object

#### **Description**

A torch\_device is an object representing the device on which a torch\_tensor is or will be allocated.

## Usage

```
torch_device(type, index = NULL)
```

#### Arguments

type

(character) a device type "cuda" or "cpu"

index

(integer) optional device ordinal for the device type. If the device ordinal is not present, this object will always represent the current device for the device type, even after torch\_cuda\_set\_device() is called; e.g., a torch\_tensor constructed with device 'cuda' is equivalent to 'cuda:X' where X is the result of torch outperformed device()

of  $torch\_cuda\_current\_device()$ .

A torch\_device can be constructed via a string or via a string and device ordinal

## **Examples**

```
if (torch_is_installed()) {

# Via string
torch_device("cuda:1")
torch_device("cpu")
torch_device("cuda") # current cuda device

# Via string and device ordinal
torch_device("cuda", 0)
torch_device("cpu", 0)
}
```

torch\_diag

Diag

#### **Description**

Diag

#### Usage

```
torch_diag(self, diagonal = 0L)
```

torch\_diagflat 383

## **Arguments**

self (Tensor) the input tensor.

diagonal (int, optional) the diagonal to consider

#### diag(input, diagonal=0, out=NULL) -> Tensor

• If input is a vector (1-D tensor), then returns a 2-D square tensor with the elements of input as the diagonal.

• If input is a matrix (2-D tensor), then returns a 1-D tensor with the diagonal elements of input.

The argument diagonal controls which diagonal to consider:

- If diagonal = 0, it is the main diagonal.
- If diagonal > 0, it is above the main diagonal.
- If diagonal < 0, it is below the main diagonal.

torch\_diagflat

Diagflat

#### **Description**

Diagflat

#### Usage

```
torch_diagflat(self, offset = 0L)
```

#### **Arguments**

self (Tensor) the input tensor.

offset (int, optional) the diagonal to consider. Default: 0 (main diagonal).

#### diagflat(input, offset=0) -> Tensor

- If input is a vector (1-D tensor), then returns a 2-D square tensor with the elements of input as the diagonal.
- If input is a tensor with more than one dimension, then returns a 2-D tensor with diagonal elements equal to a flattened input.

The argument offset controls which diagonal to consider:

- If offset = 0, it is the main diagonal.
- If offset > 0, it is above the main diagonal.
- If offset < 0, it is below the main diagonal.

384 torch\_diagonal

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3))
a
torch_diagflat(a)
torch_diagflat(a, 1)
a = torch_randn(c(2, 2))
a
torch_diagflat(a)
}
```

torch\_diagonal

Diagonal

#### **Description**

Diagonal

#### Usage

```
torch_diagonal(self, outdim, dim1 = 1L, dim2 = 2L, offset = 0L)
```

#### **Arguments**

```
self (Tensor) the input tensor. Must be at least 2-dimensional.

outdim dimension name if self is a named tensor.

dim1 (int, optional) first dimension with respect to which to take diagonal. Default: 0.

dim2 (int, optional) second dimension with respect to which to take diagonal. Default: 1.

offset (int, optional) which diagonal to consider. Default: 0 (main diagonal).
```

#### diagonal(input, offset=0, dim1=0, dim2=1) -> Tensor

Returns a partial view of input with the its diagonal elements with respect to dim1 and dim2 appended as a dimension at the end of the shape.

The argument offset controls which diagonal to consider:

- If offset = 0, it is the main diagonal.
- If offset > 0, it is above the main diagonal.
- If offset < 0, it is below the main diagonal.

Applying torch\_diag\_embed to the output of this function with the same arguments yields a diagonal matrix with the diagonal entries of the input. However, torch\_diag\_embed has different default dimensions, so those need to be explicitly specified.

torch\_diag\_embed 385

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3, 3))
a
torch_diagonal(a, offset = 0)
torch_diagonal(a, offset = 1)
x = torch_randn(c(2, 5, 4, 2))
torch_diagonal(x, offset=-1, dim1=1, dim2=2)
}
```

torch\_diag\_embed

Diag\_embed

#### **Description**

Diag\_embed

#### Usage

```
torch_diag_embed(self, offset = 0L, dim1 = -2L, dim2 = -1L)
```

#### **Arguments**

```
self (Tensor) the input tensor. Must be at least 1-dimensional.

offset (int, optional) which diagonal to consider. Default: 0 (main diagonal).

dim1 (int, optional) first dimension with respect to which to take diagonal. Default:
-2.

dim2 (int, optional) second dimension with respect to which to take diagonal. Default:
-1.
```

#### diag\_embed(input, offset=0, dim1=-2, dim2=-1) -> Tensor

Creates a tensor whose diagonals of certain 2D planes (specified by dim1 and dim2) are filled by input. To facilitate creating batched diagonal matrices, the 2D planes formed by the last two dimensions of the returned tensor are chosen by default.

The argument offset controls which diagonal to consider:

- If offset = 0, it is the main diagonal.
- If offset > 0, it is above the main diagonal.
- If offset < 0, it is below the main diagonal.

The size of the new matrix will be calculated to make the specified diagonal of the size of the last input dimension. Note that for offset other than 0, the order of dim1 and dim2 matters. Exchanging them is equivalent to changing the sign of offset.

Applying torch\_diagonal to the output of this function with the same arguments yields a matrix identical to input. However, torch\_diagonal has different default dimensions, so those need to be explicitly specified.

386 torch\_diff

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(2, 3))
torch_diag_embed(a)
torch_diag_embed(a, offset=1, dim1=1, dim2=3)
}
```

torch\_diff

Computes the n-th forward difference along the given dimension.

#### Description

The first-order differences are given by out[i] = input[i + 1] - input[i]. Higher-order differences are calculated by using torch\_diff() recursively.

## Usage

```
torch_diff(self, n = 1L, dim = -1L, prepend = list(), append = list())
```

#### **Arguments**

the tensor to compute the differences on
the number of times to recursively compute the difference
the dimension to compute the difference along. Default is the last dimension.

prepend values to prepend to input along dim before computing the difference. Their dimensions must be equivalent to that of input, and their shapes must match input's shape except on dim.

append values to append to input along dim before computing the difference. Their dimensions must be equivalent to that of input, and their shapes must match

input's shape except on dim.

#### Note

Only n = 1 is currently supported

```
if (torch_is_installed()) {
a <- torch_tensor(c(1,2,3))
torch_diff(a)

b <- torch_tensor(c(4, 5))
torch_diff(a, append = b)

c <- torch_tensor(rbind(c(1,2,3), c(3,4,5)))
torch_diff(c, dim = 1)</pre>
```

torch\_digamma 387

```
torch_diff(c, dim = 2)
}
```

torch\_digamma

Digamma

## **Description**

Digamma

## Usage

```
torch_digamma(self)
```

#### **Arguments**

self

(Tensor) the tensor to compute the digamma function on

#### digamma(input, out=NULL) -> Tensor

Computes the logarithmic derivative of the gamma function on input.

$$\psi(x) = \frac{d}{dx} \ln (\Gamma(x)) = \frac{\Gamma'(x)}{\Gamma(x)}$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_tensor(c(1, 0.5))
torch_digamma(a)
}
```

torch\_dist

Dist

# Description

Dist

## Usage

```
torch_dist(self, other, p = 2L)
```

388 torch\_div

#### **Arguments**

```
self (Tensor) the input tensor.

other (Tensor) the Right-hand-side input tensor

p (float, optional) the norm to be computed
```

#### dist(input, other, p=2) -> Tensor

```
Returns the p-norm of (input - other)
```

The shapes of input and other must be broadcastable.

#### **Examples**

```
if (torch_is_installed()) {
x = torch_randn(c(4))
x
y = torch_randn(c(4))
y
torch_dist(x, y, 3.5)
torch_dist(x, y, 3)
torch_dist(x, y, 0)
torch_dist(x, y, 1)
}
```

torch\_div

Div

#### **Description**

Div

# Usage

```
torch_div(self, other, rounding_mode)
```

#### **Arguments**

self (Tensor) the input tensor.

other (Number) the number to be divided to each element of input

rounding\_mode (str, optional) – Type of rounding applied to the result:

- NULL default behavior. Performs no rounding and, if both input and other are integer types, promotes the inputs to the default scalar type. Equivalent to true division in Python (the / operator) and NumPy's np.true\_divide.
- "trunc" rounds the results of the division towards zero. Equivalent to C-style integer division.
- "floor" rounds the results of the division down. Equivalent to floor division in Python (the // operator) and NumPy's np. floor\_divide.

torch\_divide 389

#### div(input, other, out=NULL) -> Tensor

Divides each element of the input input with the scalar other and returns a new resulting tensor.

Each element of the tensor input is divided by each element of the tensor other. The resulting tensor is returned.

$$\operatorname{out}_i = \frac{\operatorname{input}_i}{\operatorname{other}_i}$$

The shapes of input and other must be broadcastable. If the torch\_dtype of input and other differ, the torch\_dtype of the result tensor is determined following rules described in the type promotion documentation. If out is specified, the result must be castable to the torch\_dtype of the specified output tensor. Integral division by zero leads to undefined behavior.

#### Warning

Integer division using div is deprecated, and in a future release div will perform true division like torch\_true\_divide(). Use torch\_floor\_divide() to perform integer division, instead.

$$\operatorname{out}_i = \frac{\operatorname{input}_i}{\operatorname{other}}$$

If the torch\_dtype of input and other differ, the torch\_dtype of the result tensor is determined following rules described in the type promotion documentation. If out is specified, the result must be castable to the torch\_dtype of the specified output tensor. Integral division by zero leads to undefined behavior.

#### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(5))
a
torch_div(a, 0.5)

a = torch_randn(c(4, 4))
a
b = torch_randn(c(4))
b
torch_div(a, b)
}
```

torch\_divide

Divide

## **Description**

Divide

390 torch\_dot

#### Usage

```
torch_divide(self, other, rounding_mode)
```

# Arguments

self (Tensor) the input tensor.

other (Number) the number to be divided to each element of input

rounding\_mode (str, optional) – Type of rounding applied to the result:

- NULL default behavior. Performs no rounding and, if both input and other are integer types, promotes the inputs to the default scalar type. Equivalent to true division in Python (the / operator) and NumPy's np.true\_divide.
- "trunc" rounds the results of the division towards zero. Equivalent to C-style integer division.
- "floor" rounds the results of the division down. Equivalent to floor division in Python (the // operator) and NumPy's np. floor\_divide.

# divide(input, other, \*, out=None) -> Tensor

Alias for torch\_div().

torch\_dot Dot

# Description

Dot

#### Usage

```
torch_dot(self, tensor)
```

#### **Arguments**

self the input tensor tensor the other input tensor

#### dot(input, tensor) -> Tensor

Computes the dot product (inner product) of two tensors.

#### Note

This function does not broadcast.

torch\_dstack 391

## **Examples**

```
if (torch_is_installed()) {
  torch_dot(torch_tensor(c(2, 3)), torch_tensor(c(2, 1)))
}
```

torch\_dstack

Dstack

# Description

Dstack

# Usage

```
torch_dstack(tensors)
```

# **Arguments**

tensors

(sequence of Tensors) sequence of tensors to concatenate

#### dstack(tensors, \*, out=None) -> Tensor

Stack tensors in sequence depthwise (along third axis).

This is equivalent to concatenation along the third axis after 1-D and 2-D tensors have been reshaped by torch\_atleast\_3d().

```
if (torch_is_installed()) {
a <- torch_tensor(c(1, 2, 3))
b <- torch_tensor(c(4, 5, 6))
torch_dstack(list(a,b))
a <- torch_tensor(rbind(1,2,3))
b <- torch_tensor(rbind(4,5,6))
torch_dstack(list(a,b))
}</pre>
```

392 torch\_dtype

 $torch\_dtype$ 

Torch data types

# Description

Returns the correspondent data type.

# Usage

```
torch_float32()
torch_float()
torch_float64()
torch_double()
torch_cfloat32()
torch_chalf()
torch_cfloat()
torch_cfloat64()
torch_cdouble()
torch_cfloat128()
torch_float16()
torch_half()
torch_uint8()
torch_int8()
torch_int16()
torch_short()
torch_int32()
torch_int()
```

torch\_int64()

torch\_eig 393

torch\_long()

torch\_bool()

torch\_quint8()

torch\_qint8()

torch\_qint32()

torch\_eig

Eig

# Description

Eig

# Arguments

self (Tensor) the square matrix of shape  $(n \times n)$  for which the eigenvalues and eigen-

vectors will be computed

eigenvectors (bool) TRUE to compute both eigenvalues and eigenvectors; otherwise, only

eigenvalues will be computed

#### eig(input, eigenvectors=False, out=NULL) -> (Tensor, Tensor)

Computes the eigenvalues and eigenvectors of a real square matrix.

torch\_einsum Einsum

# **Description**

Einsum

# Usage

```
torch_einsum(equation, tensors, path = NULL)
```

394 torch\_einsum

#### **Arguments**

equation

(string) The equation is given in terms of lower case letters (indices) to be associated with each dimension of the operands and result. The left hand side lists the operands dimensions, separated by commas. There should be one index letter per tensor dimension. The right hand side follows after -> and gives the indices for the output. If the -> and right hand side are omitted, it implicitly defined as the alphabetically sorted list of all indices appearing exactly once in the left hand side. The indices not apprearing in the output are summed over after multiplying the operands entries. If an index appears several times for the same operand, a diagonal is taken. Ellipses . . . represent a fixed number of dimensions. If the right hand side is inferred, the ellipsis dimensions are at the beginning of the output.

tensors

(Tensor) The operands to compute the Einstein sum of.

path

(int) This function uses opt\_einsum to speed up computation or to consume less memory by optimizing contraction order. This optimization occurs when there are at least three inputs, since the order does not matter otherwise. Note that finding *the* optimal path is an NP-hard problem, thus, opt\_einsum relies on different heuristics to achieve near-optimal results. If opt\_einsum is not available, the default order is to contract from left to right. The path argument is used to changed that default, but it should only be set by advanced users.

## einsum(equation, \*operands) -> Tensor

This function provides a way of computing multilinear expressions (i.e. sums of products) using the Einstein summation convention.

```
if (torch_is_installed()) {
x = torch_randn(c(5))
y = torch_randn(c(4))
torch_einsum('i,j->ij', list(x, y)) # outer product
A = torch_randn(c(3,5,4))
1 = torch_randn(c(2,5))
r = torch_randn(c(2,4))
torch_einsum('bn,anm,bm->ba', list(1, A, r)) # compare torch_nn$functional$bilinear
As = torch_randn(c(3,2,5))
Bs = torch_randn(c(3,5,4))
torch_einsum('bij,bjk->bik', list(As, Bs)) # batch matrix multiplication
A = torch_randn(c(3, 3))
torch_einsum('ii->i', list(A)) # diagonal
A = torch_randn(c(4, 3, 3))
torch_einsum('...ii->...i', list(A)) # batch diagonal
A = torch_randn(c(2, 3, 4, 5))
torch_einsum('...ij->...ji', list(A))$shape # batch permute
}
```

torch\_empty 395

#### **Description**

**Empty** 

#### Usage

```
torch_empty(
    ...,
    names = NULL,
    dtype = NULL,
    layout = NULL,
    device = NULL,
    requires_grad = FALSE
)
```

#### **Arguments**

a sequence of integers defining the shape of the output tensor. . . . names optional character vector naming each dimension. (torch.dtype, optional) the desired data type of returned tensor. Default: if dtype NULL, uses a global default (see torch\_set\_default\_tensor\_type). (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided. layout device (torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types. requires\_grad (bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

# empty(\*size, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False, pin\_memory=False) -> Tensor

Returns a tensor filled with uninitialized data. The shape of the tensor is defined by the variable argument size.

```
if (torch_is_installed()) {
torch_empty(c(2, 3))
}
```

396 torch\_empty\_like

#### **Description**

Empty\_like

# Usage

```
torch_empty_like(
  input,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

## **Arguments**

input (Tensor) the size of input will determine size of the output tensor. dtype (torch.dtype, optional) the desired data type of returned Tensor. Default: if NULL, defaults to the dtype of input. (torch.layout, optional) the desired layout of returned tensor. Default: if layout NULL, defaults to the layout of input. device (torch.device, optional) the desired device of returned tensor. Default: if NULL, defaults to the device of input. requires\_grad (bool, optional) If autograd should record operations on the returned tensor. Default: FALSE. memory\_format (torch.memory\_format, optional) the desired memory format of returned Tensor. Default: torch\_preserve\_format.

# empty\_like(input, dtype=NULL, layout=NULL, device=NULL, requires\_grad=False, memory\_format=torch.preserve\_for-> Tensor

Returns an uninitialized tensor with the same size as input. torch\_empty\_like(input) is equivalent to torch\_empty(input.size(), dtype=input.dtype, layout=input.layout, device=input.device).

```
if (torch_is_installed()) {
torch_empty(list(2,3), dtype = torch_int64())
}
```

torch\_empty\_strided 397

#### **Description**

Empty\_strided

## Usage

```
torch_empty_strided(
    size,
    stride,
    dtype = NULL,
    layout = NULL,
    device = NULL,
    requires_grad = FALSE,
    pin_memory = FALSE
)
```

#### Arguments

size	(tuple of ints) the shape of the output tensor
stride	(tuple of ints) the strides of the output tensor
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch_set_default_tensor_type).
layout	(torch.layout, optional) the desired layout of returned Tensor. Default: torch_strided.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.
pin_memory	(bool, optional) If set, returned tensor would be allocated in the pinned memory. Works only for CPU tensors. Default: FALSE.

# empty\_strided(size, stride, dtype=NULL, layout=NULL, device=NULL, requires\_grad=False, pin\_memory=False) -> Tensor

Returns a tensor filled with uninitialized data. The shape and strides of the tensor is defined by the variable argument size and stride respectively. torch\_empty\_strided(size, stride) is equivalent to torch\_empty(size).as\_strided(size, stride).

## Warning

More than one element of the created tensor may refer to a single memory location. As a result, in-place operations (especially ones that are vectorized) may result in incorrect behavior. If you need to write to the tensors, please clone them first.

398 torch\_eq

## **Examples**

```
if (torch_is_installed()) {
a = torch_empty_strided(list(2, 3), list(1, 2))
a
a$stride(1)
a$size(1)
}
```

torch\_eq

Eq

# Description

Eq

## Usage

```
torch_eq(self, other)
```

# Arguments

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare Must be a ByteTensor

# $eq(input, other, out=NULL) \rightarrow Tensor$

Computes element-wise equality

The second argument can be a number or a tensor whose shape is broadcastable with the first argument.

```
if (torch_is_installed()) {
torch_eq(torch_tensor(c(1,2,3,4)), torch_tensor(c(1, 3, 2, 4)))
}
```

torch\_equal 399

torch\_equal

Equal

## **Description**

Equal

## Usage

```
torch_equal(self, other)
```

# Arguments

self the input tensor other the other input tensor

# equal(input, other) -> bool

TRUE if two tensors have the same size and elements, FALSE otherwise.

# **Examples**

```
if (torch_is_installed()) {
  torch_equal(torch_tensor(c(1, 2)), torch_tensor(c(1, 2)))
}
```

torch\_erf

Erf

# **Description**

Erf

# Usage

```
torch_erf(self)
```

# Arguments

self

(Tensor) the input tensor.

# erf(input, out=NULL) -> Tensor

Computes the error function of each element. The error function is defined as follows:

$$\operatorname{erf}(x) = \frac{2}{\sqrt{\pi}} \int_0^x e^{-t^2} dt$$

400 torch\_erfc

# **Examples**

```
if (torch_is_installed()) {
  torch_erf(torch_tensor(c(0, -1., 10.)))
}
```

torch\_erfc

Erfc

# Description

Erfc

# Usage

```
torch_erfc(self)
```

# Arguments

self

(Tensor) the input tensor.

# erfc(input, out=NULL) -> Tensor

Computes the complementary error function of each element of input. The complementary error function is defined as follows:

$$\operatorname{erfc}(x) = 1 - \frac{2}{\sqrt{\pi}} \int_0^x e^{-t^2} dt$$

```
if (torch_is_installed()) {
  torch_erfc(torch_tensor(c(0, -1., 10.)))
}
```

torch\_erfinv 401

torch\_erfinv

**Erfinv** 

## **Description**

Erfinv

# Usage

```
torch_erfinv(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# erfinv(input, out=NULL) -> Tensor

Computes the inverse error function of each element of input. The inverse error function is defined in the range (-1,1) as:

$$\operatorname{erfinv}(\operatorname{erf}(x)) = x$$

# **Examples**

```
if (torch_is_installed()) {
torch_erfinv(torch_tensor(c(0, 0.5, -1.)))
}
```

torch\_exp

Exp

# Description

Exp

## Usage

```
torch_exp(self)
```

# Arguments

self

(Tensor) the input tensor.

402 torch\_exp2

# exp(input, out=NULL) -> Tensor

Returns a new tensor with the exponential of the elements of the input tensor input.

$$y_i = e^{x_i}$$

# **Examples**

```
if (torch_is_installed()) {
  torch_exp(torch_tensor(c(0, log(2))))
}
```

torch\_exp2

Exp2

# Description

Exp2

# Usage

```
torch_exp2(self)
```

# Arguments

self

(Tensor) the input tensor.

# exp2(input, \*, out=None) -> Tensor

Computes the base two exponential function of input.

$$y_i = 2^{x_i}$$

```
if (torch_is_installed()) {
torch_exp2(torch_tensor(c(0, log2(2.), 3, 4)))
}
```

torch\_expm1 403

torch\_expm1

Expm1

# Description

Expm1

# Usage

```
torch_expm1(self)
```

# Arguments

self

(Tensor) the input tensor.

# expm1(input, out=NULL) -> Tensor

Returns a new tensor with the exponential of the elements minus 1 of input.

$$y_i = e^{x_i} - 1$$

# **Examples**

```
if (torch_is_installed()) {
  torch_expm1(torch_tensor(c(0, log(2))))
}
```

torch\_eye

Eye

# Description

Eye

# Usage

```
torch_eye(
   n,
   m = n,
   dtype = NULL,
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

404 torch\_fft\_fft

#### **Arguments**

n (int) the number of rows (int, optional) the number of columns with default being n m (torch.dtype, optional) the desired data type of returned tensor. Default: if dtype NULL, uses a global default (see torch\_set\_default\_tensor\_type). layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided. device (torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types. (bool, optional) If autograd should record operations on the returned tensor. Derequires\_grad

# eye(n, m=NULL, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a 2-D tensor with ones on the diagonal and zeros elsewhere.

fault: FALSE.

# Examples

## **Description**

Computes the one dimensional discrete Fourier transform of input.

## Usage

```
torch_fft_fft(self, n = NULL, dim = -1L, norm = NULL)
```

## **Arguments**

self	(Tensor) the input tensor
n	(int) Signal length. If given, the input will either be zero-padded or trimmed to this length before computing the FFT.
dim	(int, optional) The dimension along which to take the one dimensional FFT.
norm	(str, optional) Normalization mode. For the forward transform, these correspond to:
	• "forward" - normalize by 1/n

torch\_fft\_fftfreq 405

- "backward" no normalization
- "ortho" normalize by 1/sqrt(n) (making the FFT orthonormal) Calling the backward transform (ifft()) with the same normalization mode will apply an overall normalization of 1/n between the two transforms. This is required to make IFFT the exact inverse. Default is "backward" (no normalization).

#### Note

The Fourier domain representation of any real signal satisfies the Hermitian property: X[i] = conj(X[-i]). This function always returns both the positive and negative frequency terms even though, for real inputs, the negative frequencies are redundant. rfft() returns the more compact one-sided representation where only the positive frequencies are returned.

#### **Examples**

```
if (torch_is_installed()) {
t <- torch_arange(start = 0, end = 3)
t
torch_fft_fft(t, norm = "backward")
}</pre>
```

torch\_fft\_fftfreq fftfreq

## **Description**

Computes the discrete Fourier Transform sample frequencies for a signal of size n.

## Usage

```
torch_fft_fftfreq(
   n,
   d = 1,
   dtype = NULL,
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

## **Arguments**

```
n (integer) – the FFT length

d (float, optional) – the sampling length scale. The spacing between individual samples of the FFT input. The default assumes unit spacing, dividing that result by the actual spacing gives the result in physical frequency units.

dtype (default: torch_get_default_dtype()) the desired data type of returned tensor.
```

406 torch\_fft\_ifft

layout (default: torch\_strided()) the desired layout of returned tensor.

device (default: NULL) the desired device of returned tensor. Default: If NULL, uses the

current device for the default tensor type.

requires\_grad (default: FALSE) If autograd should record operations on the returned tensor.

#### Note

By convention, torch\_fft\_fft() returns positive frequency terms first, followed by the negative frequencies in reverse order, so that f[-i] for all 0 < i <= n/2 gives the negative frequency terms. For an FFT of length n and with inputs spaced in length unit d, the frequencies are: f = [0, 1, ..., (n - 1) // 2, -(n // 2), ..., -1] / (d \* n)

For even lengths, the Nyquist frequency at f[n/2] can be thought of as either negative or positive. fftfreq() follows NumPy's convention of taking it to be negative.

## **Examples**

```
if (torch_is_installed()) {
  torch_fft_fftfreq(5) # Nyquist frequency at f[3] is positive
  torch_fft_fftfreq(4) # Nyquist frequency at f[3] is given as negative
}
```

torch\_fft\_ifft

*Ifft* 

#### Description

Computes the one dimensional inverse discrete Fourier transform of input.

#### Usage

```
torch_fft_ifft(self, n = NULL, dim = -1L, norm = NULL)
```

## **Arguments**

self (Tensor) the input tensor

n (int, optional) – Signal length. If given, the input will either be zero-padded or trimmed to this length before computing the IFFT.

dim (int, optional) – The dimension along which to take the one dimensional IFFT.

norm (str, optional) – Normalization mode. For the backward transform, these correspond to:

- "forward" no normalization
- "backward" normalize by 1/n
- "ortho" normalize by 1/sqrt(n) (making the IFFT orthonormal) Calling the forward transform with the same normalization mode will apply an overall normalization of 1/n between the two transforms. This is required to make ifft() the exact inverse. Default is "backward" (normalize by 1/n).

torch\_fft\_irfft 407

#### **Examples**

```
if (torch_is_installed()) {
t <- torch_arange(start = 0, end = 3)
t
x <- torch_fft_fft(t, norm = "backward")
torch_fft_ifft(x)
}</pre>
```

torch\_fft\_irfft

Irfft

#### **Description**

Computes the inverse of torch\_fft\_rfft(). Input is interpreted as a one-sided Hermitian signal in the Fourier domain, as produced by torch\_fft\_rfft(). By the Hermitian property, the output will be real-valued.

#### Usage

```
torch_fft_irfft(self, n = NULL, dim = -1L, norm = NULL)
```

# **Arguments**

self (Tensor) the input tensor representing a half-Hermitian signal

(int) Output signal length. This determines the length of the output signal. If given, the input will either be zero-padded or trimmed to this length before computing the real IFFT. Defaults to even output: n=2\*(input.size(dim) - 1).

dim (int, optional) – The dimension along which to take the one dimensional real IFFT.

norm (str, optional) – Normalization mode. For the backward transform, these correspond to:

- "forward" no normalization
- "backward" normalize by 1/n
- "ortho" normalize by 1/sqrt(n) (making the real IFFT orthonormal) Calling the forward transform (torch\_fft\_rfft()) with the same normalization mode will apply an overall normalization of 1/n between the two transforms. This is required to make irfft() the exact inverse. Default is "backward" (normalize by 1/n).

408 torch\_fft\_rfft

#### Note

Some input frequencies must be real-valued to satisfy the Hermitian property. In these cases the imaginary component will be ignored. For example, any imaginary component in the zero-frequency term cannot be represented in a real output and so will always be ignored.

The correct interpretation of the Hermitian input depends on the length of the original data, as given by n. This is because each input shape could correspond to either an odd or even length signal. By default, the signal is assumed to be even length and odd signals will not round-trip properly. So, it is recommended to always pass the signal length n.

## **Examples**

```
if (torch_is_installed()) {
  t <- torch_arange(start = 0, end = 4)
  x <- torch_fft_rfft(t)
  torch_fft_irfft(x)
  torch_fft_irfft(x, n = t$numel())
}</pre>
```

torch fft rfft

Rfft

#### **Description**

Computes the one dimensional Fourier transform of real-valued input.

#### Usage

```
torch_fft_rfft(self, n = NULL, dim = -1L, norm = NULL)
```

## **Arguments**

dim

norm

self (Tensor) the real input tensor

n (int) Signal length. If given, the input will either be zero-padded or trimmed to this length before computing the real FFT.

(int, optional) - The dimension along which to take the one dimensional real

FFT

norm (str, optional) – Normalization mode. For the forward transform, these correspond to:

- "forward" normalize by 1/n
- "backward" no normalization
- "ortho" normalize by 1/sqrt(n) (making the FFT orthonormal) Calling the backward transform (torch\_fft\_irfft()) with the same normalization mode will apply an overall normalization of 1/n between the two transforms. This is required to make irfft() the exact inverse. Default is "backward" (no normalization).

torch\_finfo 409

## **Details**

The FFT of a real signal is Hermitian-symmetric, X[i] = conj(X[-i]) so the output contains only the positive frequencies below the Nyquist frequency. To compute the full output, use  $torch_fft_fft()$ .

# **Examples**

```
if (torch_is_installed()) {
t <- torch_arange(start = 0, end = 3)
torch_fft_rfft(t)
}</pre>
```

torch\_finfo

Floating point type info

## Description

A list that represents the numerical properties of a floating point torch.dtype

# Usage

```
torch_finfo(dtype)
```

## **Arguments**

dtype

dtype to check information

torch\_fix

Fix

# Description

Fix

# Usage

```
torch_fix(self)
```

# Arguments

self

(Tensor) the input tensor.

# fix(input, \*, out=None) -> Tensor

```
Alias for torch_trunc()
```

410 torch\_flip

torch\_flatten

Flatten

# Description

Flatten

#### Usage

```
torch_flatten(self, dims, start_dim = 1L, end_dim = -1L, out_dim)
```

## **Arguments**

self (Tensor) the input tensor.

dims if tensor is named you can pass the name of the dimensions to flatten

start\_dim (int) the first dim to flatten end\_dim (int) the last dim to flatten

out\_dim the name of the resulting dimension if a named tensor.

# flatten(input, start\_dim=0, end\_dim=-1) -> Tensor

Flattens a contiguous range of dims in a tensor.

## **Examples**

```
if (torch_is_installed()) {
t = torch_tensor(matrix(c(1, 2), ncol = 2))
torch_flatten(t)
torch_flatten(t, start_dim=2)
}
```

torch\_flip

Flip

## **Description**

Flip

# Usage

```
torch_flip(self, dims)
```

torch\_fliplr 411

## **Arguments**

```
self (Tensor) the input tensor.
dims (a list or tuple) axis to flip on
```

# flip(input, dims) -> Tensor

Reverse the order of a n-D tensor along given axis in dims.

# **Examples**

```
if (torch_is_installed()) {
x <- torch_arange(1, 8)$view(c(2, 2, 2))
x
torch_flip(x, c(1, 2))
}</pre>
```

torch\_fliplr

Fliplr

# Description

Fliplr

# Usage

```
torch_fliplr(self)
```

## **Arguments**

self

(Tensor) Must be at least 2-dimensional.

# fliplr(input) -> Tensor

Flip array in the left/right direction, returning a new tensor.

Flip the entries in each row in the left/right direction. Columns are preserved, but appear in a different order than before.

## Note

Equivalent to input[,-1]. Requires the array to be at least 2-D.

412 torch\_flipud

#### **Examples**

```
if (torch_is_installed()) {

x <- torch_arange(start = 1, end = 4)$view(c(2, 2))
x
torch_fliplr(x)
}</pre>
```

torch\_flipud

Flipud

# Description

Flipud

# Usage

```
torch_flipud(self)
```

## **Arguments**

self

(Tensor) Must be at least 1-dimensional.

# flipud(input) -> Tensor

Flip array in the up/down direction, returning a new tensor.

Flip the entries in each column in the up/down direction. Rows are preserved, but appear in a different order than before.

## Note

Equivalent to input[-1,]. Requires the array to be at least 1-D.

```
if (torch_is_installed()) {
x <- torch_arange(start = 1, end = 4)$view(c(2, 2))
x
torch_flipud(x)
}</pre>
```

torch\_floor 413

torch\_floor

Floor

# Description

Floor

## Usage

```
torch_floor(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# floor(input, out=NULL) -> Tensor

Returns a new tensor with the floor of the elements of input, the largest integer less than or equal to each element.

$$out_i = \lfloor input_i \rfloor$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_floor(a)
}
```

torch\_floor\_divide

 $Floor\_divide$ 

# Description

Floor\_divide

## Usage

```
torch_floor_divide(self, other)
```

## Arguments

self (Tensor) the numerator tensor other (Tensor or Scalar) the denominator 414 torch\_fmod

#### floor\_divide(input, other, out=NULL) -> Tensor

Return the division of the inputs rounded down to the nearest integer. See torch\_div for type promotion and broadcasting rules.

$$\operatorname{out}_i = \left\lfloor \frac{\operatorname{input}_i}{\operatorname{other}_i} \right\rfloor$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_tensor(c(4.0, 3.0))
b = torch_tensor(c(2.0, 2.0))
torch_floor_divide(a, b)
torch_floor_divide(a, 1.4)
}
```

torch\_fmod

Fmod

#### **Description**

Fmod

#### Usage

```
torch_fmod(self, other)
```

#### **Arguments**

self (Tensor) the dividend

other (Tensor or float) the divisor, which may be either a number or a tensor of the same shape as the dividend

#### fmod(input, other, out=NULL) -> Tensor

Computes the element-wise remainder of division.

The dividend and divisor may contain both for integer and floating point numbers. The remainder has the same sign as the dividend input.

When other is a tensor, the shapes of input and other must be broadcastable.

```
if (torch_is_installed()) {
torch_fmod(torch_tensor(c(-3., -2, -1, 1, 2, 3)), 2)
torch_fmod(torch_tensor(c(1., 2, 3, 4, 5)), 1.5)
}
```

torch\_frac 415

torch\_frac

Frac

# Description

Frac

# Usage

```
torch_frac(self)
```

## **Arguments**

self

the input tensor.

# frac(input, out=NULL) -> Tensor

Computes the fractional portion of each element in input.

```
out_i = input_i - \lfloor |input_i| \rfloor * sgn(input_i)
```

# **Examples**

```
if (torch_is_installed()) {
  torch_frac(torch_tensor(c(1, 2.5, -3.2)))
}
```

torch\_full

Full

# Description

Full

# Usage

```
torch_full(
    size,
    fill_value,
    names = NULL,
    dtype = NULL,
    layout = NULL,
    device = NULL,
    requires_grad = FALSE
)
```

416 torch\_full\_like

#### Arguments

(int...) a list, tuple, or torch\_Size of integers defining the shape of the output size tensor. NA the number to fill the output tensor with. fill\_value names optional names of the dimensions (torch.dtype, optional) the desired data type of returned tensor. Default: if dtype NULL, uses a global default (see torch\_set\_default\_tensor\_type). layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided. device (torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types. requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

# full(size, fill\_value, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a tensor of size size filled with fill\_value.

fault: FALSE.

# Warning

In PyTorch 1.5 a bool or integral fill\_value will produce a warning if dtype or out are not set. In a future PyTorch release, when dtype and out are not set a bool fill\_value will return a tensor of torch.bool dtype, and an integral fill\_value will return a tensor of torch.long dtype.

## **Examples**

```
if (torch_is_installed()) {
torch_full(list(2, 3), 3.141592)
}
```

torch\_full\_like Full\_like

## **Description**

Full\_like

torch\_gather 417

#### Usage

```
torch_full_like(
  input,
  fill_value,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

#### **Arguments**

input (Tensor) the size of input will determine size of the output tensor.

fill\_value the number to fill the output tensor with.

dtype (torch.dtype, optional) the desired data type of returned Tensor. Default: if

NULL, defaults to the dtype of input.

layout (torch.layout, optional) the desired layout of returned tensor. Default: if

NULL, defaults to the layout of input.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, defaults to the device of input.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

memory\_format (torch.memory\_format, optional) the desired memory format of returned Ten-

sor. Default: torch\_preserve\_format.

## full\_like(input, fill\_value, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False,

memory\_format=torch.preserve\_format) -> Tensor

Returns a tensor with the same size as input filled with fill\_value. torch\_full\_like(input, fill\_value) is equivalent to torch\_full(input.size(), fill\_value, dtype=input.dtype, layout=input.layout, device=input.device).

## **Description**

Gather

## Usage

```
torch_gather(self, dim, index, sparse_grad = FALSE)
```

418 torch\_gcd

#### **Arguments**

self	(Tensor) the source tensor
dim	(int) the axis along which to index
index	(LongTensor) the indices of elements to gather
sparse_grad	(bool,optional) If TRUE, gradient w.r.t. input will be a sparse tensor.

#### gather(input, dim, index, sparse\_grad=FALSE) -> Tensor

Gathers values along an axis specified by dim.

For a 3-D tensor the output is specified by::

```
out[i][j][k] = input[index[i][j][k]][j][k]  # if dim == 0
out[i][j][k] = input[i][index[i][j][k]]  # if dim == 1
out[i][j][k] = input[i][j][index[i][j][k]]  # if dim == 2
```

If input is an n-dimensional tensor with size  $(x_0, x_1..., x_{i-1}, x_i, x_{i+1}, ..., x_{n-1})$  and dim = i, then index must be an n-dimensional tensor with size  $(x_0, x_1, ..., x_{i-1}, y, x_{i+1}, ..., x_{n-1})$  where  $y \ge 1$  and out will have the same size as index.

## **Examples**

```
if (torch_is_installed()) {

t = torch_tensor(matrix(c(1,2,3,4), ncol = 2, byrow = TRUE))

torch_gather(t, 2, torch_tensor(matrix(c(1,1,2,1), ncol = 2, byrow=TRUE), dtype = torch_int64()))
}
```

torch\_gcd

Gcd

## Description

Gcd

#### Usage

```
torch_gcd(self, other)
```

#### Arguments

```
self (Tensor) the input tensor.
other (Tensor) the second input tensor
```

# gcd(input, other, \*, out=None) -> Tensor

Computes the element-wise greatest common divisor (GCD) of input and other.

Both input and other must have integer types.

torch\_ge 419

## Note

This defines gcd(0,0) = 0.

## **Examples**

```
if (torch_is_installed()) {
   if (torch::cuda_is_available()) {
    a <- torch_tensor(c(5, 10, 15), dtype = torch_long(), device = "cuda")
   b <- torch_tensor(c(3, 4, 5), dtype = torch_long(), device = "cuda")
   torch_gcd(a, b)
   c <- torch_tensor(c(3L), device = "cuda")
   torch_gcd(a, c)
}
</pre>
```

torch\_ge

Ge

## **Description**

Ge

## Usage

```
torch_ge(self, other)
```

## **Arguments**

```
self (Tensor) the tensor to compare
other (Tensor or float) the tensor or value to compare
```

## ge(input, other, out=NULL) -> Tensor

Computes input  $\geq$  other element-wise.

The second argument can be a number or a tensor whose shape is broadcastable with the first argument.

420 torch\_geqrf

torch\_generator

Create a Generator object

# Description

A torch\_generator is an object which manages the state of the algorithm that produces pseudo random numbers. Used as a keyword argument in many In-place random sampling functions.

## Usage

```
torch_generator()
```

# **Examples**

```
if (torch_is_installed()) {
# Via string
generator <- torch_generator()
generator$current_seed()
generator$set_current_seed(1234567L)
generator$current_seed()
}</pre>
```

torch\_geqrf

Geqrf

# Description

Geqrf

# Usage

```
torch_geqrf(self)
```

# Arguments

self

(Tensor) the input matrix

torch\_ger 421

#### geqrf(input, out=NULL) -> (Tensor, Tensor)

This is a low-level function for calling LAPACK directly. This function returns a namedtuple (a, tau) as defined in LAPACK documentation for geqrf\_.

You'll generally want to use torch\_qr instead.

Computes a QR decomposition of input, but without constructing Q and R as explicit separate matrices.

Rather, this directly calls the underlying LAPACK function ?geqrf which produces a sequence of 'elementary reflectors'.

See LAPACK documentation for geqrf\_for further details.

torch\_ger

Ger

#### Description

Ger

## Usage

```
torch_ger(self, vec2)
```

#### **Arguments**

```
self (Tensor) 1-D input vector
vec2 (Tensor) 1-D input vector
```

# ger(input, vec2, out=NULL) -> Tensor

Outer product of input and vec2. If input is a vector of size n and vec2 is a vector of size m, then out must be a matrix of size  $(n \times m)$ .

## Note

This function does not broadcast.

```
if (torch_is_installed()) {
v1 = torch_arange(1., 5.)
v2 = torch_arange(1., 4.)
torch_ger(v1, v2)
```

422 torch\_greater

torch\_get\_rng\_state RNG state management

#### **Description**

Low level functionality to set and change the RNG state. It's recommended to use torch\_manual\_seed() for most cases.

## Usage

```
torch_get_rng_state()
torch_set_rng_state(state)
cuda_get_rng_state(device = NULL)
cuda_set_rng_state(state, device = NULL)
```

# Arguments

state A tensor with the current state or a list containing the state for each device - (for

CUDA).

device The cuda device index to get or set the state. If NULL gets the state for all avail-

able devices.

#### **Functions**

- torch\_set\_rng\_state(): Sets the RNG state for the CPU
- cuda\_get\_rng\_state(): Gets the RNG state for CUDA.
- cuda\_set\_rng\_state(): Sets the RNG state for CUDA.

torch\_greater

Greater

## Description

Greater

# Usage

```
torch_greater(self, other)
```

## **Arguments**

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

torch\_greater\_equal 423

# greater(input, other, \*, out=None) -> Tensor

Alias for torch\_gt().

torch\_greater\_equal Greater\_equal

# Description

Greater\_equal

## Usage

```
torch_greater_equal(self, other)
```

## **Arguments**

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

# greater\_equal(input, other, \*, out=None) -> Tensor

Alias for torch\_ge().

torch\_gt Gt

## **Description**

Gt

## Usage

```
torch_gt(self, other)
```

## **Arguments**

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

# gt(input, other, out=NULL) -> Tensor

Computes input > other element-wise.

The second argument can be a number or a tensor whose shape is broadcastable with the first argument.

## **Examples**

# Description

 $Hamming\_window$ 

# Usage

```
torch_hamming_window(
  window_length,
  periodic = TRUE,
  alpha = 0.54,
  beta = 0.46,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE
)
```

## **Arguments**

window_length	(int) the size of returned window
periodic	(bool, optional) If TRUE, returns a window to be used as periodic function. If False, return a symmetric window.
alpha	(float, optional) The coefficient $\alpha$ in the equation above
beta	(float, optional) The coefficient $\beta$ in the equation above
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch_set_default_tensor_type). Only floating point types are supported.
layout	(torch.layout, optional) the desired layout of returned window tensor. Only torch_strided (dense layout) is supported.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

torch\_hann\_window 425

hamming\_window(window\_length, periodic=TRUE, alpha=0.54, beta=0.46, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Hamming window function.

$$w[n] = \alpha - \beta \, \cos \left( \frac{2\pi n}{N-1} \right),$$

where N is the full window size.

The input window\_length is a positive integer controlling the returned window size. periodic flag determines whether the returned window trims off the last duplicate value from the symmetric window and is ready to be used as a periodic window with functions like torch\_stft. Therefore, if periodic is true, the N in above formula is in fact window\_length + 1. Also, we always have torch\_hamming\_window(L, periodic=TRUE) equal to torch\_hamming\_window(L + 1, periodic=False)[:-1]).

#### Note

```
If `window_length` \eqn{=1}, the returned window contains a single value 1.
```

This is a generalized version of `torch\_hann\_window`.

torch\_hann\_window

Hann\_window

#### **Description**

Hann\_window

#### Usage

```
torch_hann_window(
  window_length,
  periodic = TRUE,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE
)
```

#### **Arguments**

window\_length (int) the size of returned window

periodic (bool, optional) If TRUE, returns a window to be used as periodic function. If

False, return a symmetric window.

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type). Only

floating point types are supported.

426 torch\_heaviside

layout (torch.layout, optional) the desired layout of returned window tensor. Only

torch\_strided (dense layout) is supported.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

(bool, optional) If autograd should record operations on the returned tensor. Derequires\_grad

fault: FALSE.

## hann\_window(window\_length, periodic=TRUE, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Hann window function.

$$w[n] = \frac{1}{2} \left[ 1 - \cos\left(\frac{2\pi n}{N-1}\right) \right] = \sin^2\left(\frac{\pi n}{N-1}\right),$$

where N is the full window size.

The input window\_length is a positive integer controlling the returned window size. periodic flag determines whether the returned window trims off the last duplicate value from the symmetric window and is ready to be used as a periodic window with functions like torch\_stft. Therefore, if periodic is true, the N in above formula is in fact window\_length + 1. Also, we always have torch\_hann\_window(L, periodic=TRUE) equal to torch\_hann\_window(L + 1, periodic=False)[:-1]).

#### Note

If `window\_length` \eqn{=1}, the returned window contains a single value 1.

torch\_heaviside Heaviside

## **Description**

Heaviside

#### **Usage**

torch\_heaviside(self, values)

#### **Arguments**

self (Tensor) the input tensor.

values (Tensor) The values to use where input is zero. torch\_histc 427

## heaviside(input, values, \*, out=None) -> Tensor

Computes the Heaviside step function for each element in input. The Heaviside step function is defined as:

```
\label{eq:heaviside} \begin{aligned} \text{heaviside}(input, values) &= \begin{array}{c} 0, & \text{if input} < 0 \\ values, & \text{if input} == 0 \\ 1, & \text{if input} > 0 \\ \end{aligned}
```

## **Examples**

```
if (torch_is_installed()) {
input <- torch_tensor(c(-1.5, 0, 2.0))
values <- torch_tensor(c(0.5))
torch_heaviside(input, values)
values <- torch_tensor(c(1.2, -2.0, 3.5))
torch_heaviside(input, values)
}</pre>
```

torch\_histc

Histc

## **Description**

Histc

## Usage

```
torch_histc(self, bins = 100L, min = 0L, max = 0L)
```

## Arguments

self	(Tensor) the input tensor.
bins	(int) number of histogram bins
min	(int) lower end of the range (inclusive)
max	(int) upper end of the range (inclusive)

#### histc(input, bins=100, min=0, max=0, out=NULL) -> Tensor

Computes the histogram of a tensor.

The elements are sorted into equal width bins between min and max. If min and max are both zero, the minimum and maximum values of the data are used.

```
if (torch_is_installed()) {
  torch_histc(torch_tensor(c(1., 2, 1)), bins=4, min=0, max=3)
}
```

428 torch\_hypot

torch\_hstack

Hstack

# Description

Hstack

## Usage

```
torch_hstack(tensors)
```

# Arguments

tensors

(sequence of Tensors) sequence of tensors to concatenate

# hstack(tensors, \*, out=None) -> Tensor

Stack tensors in sequence horizontally (column wise).

This is equivalent to concatenation along the first axis for 1-D tensors, and along the second axis for all other tensors.

# **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(c(1, 2, 3))
b <- torch_tensor(c(4, 5, 6))
torch_hstack(list(a,b))
a <- torch_tensor(rbind(1,2,3))
b <- torch_tensor(rbind(4,5,6))
torch_hstack(list(a,b))
}</pre>
```

torch\_hypot

Hypot

# Description

Hypot

# Usage

```
torch_hypot(self, other)
```

torch\_i0 429

## **Arguments**

self (Tensor) the first input tensor other (Tensor) the second input tensor

## hypot(input, other, \*, out=None) -> Tensor

Given the legs of a right triangle, return its hypotenuse.

$$\operatorname{out}_i = \sqrt{\operatorname{input}_i^2 + \operatorname{other}_i^2}$$

The shapes of input and other must be broadcastable.

## **Examples**

```
if (torch_is_installed()) {
  torch_hypot(torch_tensor(c(4.0)), torch_tensor(c(3.0, 4.0, 5.0)))
}
```

torch\_i0

*I0* 

## **Description**

I0

# Usage

```
torch_i0(self)
```

## **Arguments**

self

(Tensor) the input tensor

#### i0(input, \*, out=None) -> Tensor

Computes the zeroth order modified Bessel function of the first kind for each element of input.

$$\mathrm{out}_i = I_0(\mathrm{input}_i) = \sum_{k=0}^{\infty} \frac{(\mathrm{input}_i^2/4)^k}{(k!)^2}$$

```
if (torch_is_installed()) {
torch_i0(torch_arange(start = 0, end = 5, dtype=torch_float32()))
}
```

430 torch\_imag

torch\_iinfo

Integer type info

# Description

A list that represents the numerical properties of a integer type.

# Usage

```
torch_iinfo(dtype)
```

# Arguments

dtype

dtype to get information from.

torch\_imag

Imag

# Description

Imag

## Usage

```
torch_imag(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# imag(input) -> Tensor

Returns the imaginary part of the input tensor.

# Warning

Not yet implemented.

```
out_i = imag(input_i)
```

```
if (torch_is_installed()) {
## Not run:
torch_imag(torch_tensor(c(-1 + 1i, -2 + 2i, 3 - 3i)))
## End(Not run)
}
```

torch\_index 431

# Description

Helper functions to index tensors.

# Usage

```
torch_index(self, indices)
```

# Arguments

self (Tensor) Tensor that will be indexed.

indices (List[Tensor]) List of indices. Indices are torch tensors with torch\_long()

dtype.

torch_index_put	Modify values selected by indices.	
-----------------	------------------------------------	--

# Description

Modify values selected by indices.

# Usage

```
torch_index_put(self, indices, values, accumulate = FALSE)
```

# Arguments

self	(Tensor) Tensor that will be indexed.
indices	(List[Tensor]) List of indices. Indices are torch tensors with torch_long() dtype.
values	$(Tensor)\ values\ that\ will\ be\ replaced\ the\ indexed\ location.\ Used\ for\ torch\_index\_put\ and\ torch\_index\_put\$
accumulate	(bool) Wether instead of replacing the current values with values, you want to add them.

432 torch\_index\_select

torch_index_put_	<pre>In-place version of torch_index_put.</pre>
------------------	---

#### **Description**

In-place version of torch\_index\_put.

## Usage

```
torch_index_put_(self, indices, values, accumulate = FALSE)
```

## **Arguments**

self (Tensor) Tensor that will be indexed.

indices (List[Tensor]) List of indices. Indices are torch tensors with torch\_long()

dtype.

values (Tensor) values that will be replaced the indexed location. Used for torch\_index\_put

and torch\_index\_put\_.

accumulate (bool) Wether instead of replacing the current values with values, you want to

add them.

## Description

Index\_select

#### Usage

```
torch_index_select(self, dim, index)
```

#### **Arguments**

self (Tensor) the input tensor.

dim (int) the dimension in which we index

index (LongTensor) the 1-D tensor containing the indices to index

# index\_select(input, dim, index, out=NULL) -> Tensor

Returns a new tensor which indexes the input tensor along dimension dim using the entries in index which is a LongTensor.

The returned tensor has the same number of dimensions as the original tensor (input). The dim\ th dimension has the same size as the length of index; other dimensions have the same size as in the original tensor.

torch\_install\_path 433

## Note

The returned tensor does **not** use the same storage as the original tensor. If out has a different shape than expected, we silently change it to the correct shape, reallocating the underlying storage if necessary.

# **Examples**

```
if (torch_is_installed()) {
x = torch_randn(c(3, 4))
x
indices = torch_tensor(c(1, 3), dtype = torch_int64())
torch_index_select(x, 1, indices)
torch_index_select(x, 2, indices)
}
```

torch\_install\_path

A simple exported version of install\_path Returns the torch installation path.

## Description

A simple exported version of install\_path Returns the torch installation path.

## Usage

```
torch_install_path()
```

torch\_inverse

Inverse

## Description

Inverse

# Usage

```
torch_inverse(self)
```

## Arguments

self

(Tensor) the input tensor of size  $(\ast,n,n)$  where  $\star$  is zero or more batch dimensions

torch\_isclose

#### inverse(input, out=NULL) -> Tensor

Takes the inverse of the square matrix input. input can be batches of 2D square tensors, in which case this function would return a tensor composed of individual inverses.

## Note

```
Irrespective of the original strides, the returned tensors will be transposed, i.e. with strides like `input.contiguous().transpose(-2, -1).stride()`
```

## **Examples**

```
if (torch_is_installed()) {
## Not run:
x = torch_rand(c(4, 4))
y = torch_inverse(x)
z = torch_mm(x, y)
z
torch_max(torch_abs(z - torch_eye(4))) # Max non-zero
# Batched inverse example
x = torch_randn(c(2, 3, 4, 4))
y = torch_inverse(x)
z = torch_matmul(x, y)
torch_max(torch_abs(z - torch_eye(4)$expand_as(x))) # Max non-zero
## End(Not run)
}
```

torch\_isclose

Isclose

## **Description**

Isclose

## Usage

```
torch_isclose(self, other, rtol = 1e-05, atol = 1e-08, equal_nan = FALSE)
```

#### **Arguments**

self	(Tensor) first tensor to compare
other	(Tensor) second tensor to compare
rtol	(float, optional) relative tolerance. Default: 1e-05
atol	(float, optional) absolute tolerance. Default: 1e-08
egual_nan	(bool, optional) if TRUE, then two NaN s will be considered equal. Default: FALSE

torch\_isfinite 435

#### isclose(input, other, rtol=1e-05, atol=1e-08, equal\_nan=FALSE) -> Tensor

Returns a new tensor with boolean elements representing if each element of input is "close" to the corresponding element of other. Closeness is defined as:

```
|input - other| \le atol + rtol \times |other|
```

where input and other are finite. Where input and/or other are nonfinite they are close if and only if they are equal, with NaNs being considered equal to each other when equal\_nan is TRUE.

## Examples

```
if (torch_is_installed()) {

torch_isclose(torch_tensor(c(1., 2, 3)), torch_tensor(c(1 + 1e-10, 3, 4)))
torch_isclose(torch_tensor(c(Inf, 4)), torch_tensor(c(Inf, 6)), rtol=.5)
}
```

torch\_isfinite

**Isfinite** 

## Description

**Isfinite** 

## Usage

```
torch_isfinite(self)
```

#### **Arguments**

self

(Tensor) A tensor to check

#### **TEST**

Returns a new tensor with boolean elements representing if each element is Finite or not.

```
if (torch_is_installed()) {
torch_isfinite(torch_tensor(c(1, Inf, 2, -Inf, NaN)))
}
```

torch\_isnan

torch\_isinf

Isinf

# Description

Isinf

## Usage

```
torch_isinf(self)
```

## **Arguments**

self

(Tensor) A tensor to check

## **TEST**

Returns a new tensor with boolean elements representing if each element is +/-INF or not.

## **Examples**

```
if (torch_is_installed()) {
  torch_isinf(torch_tensor(c(1, Inf, 2, -Inf, NaN)))
}
```

torch\_isnan

Isnan

# Description

Isnan

## Usage

```
torch_isnan(self)
```

# Arguments

self

(Tensor) A tensor to check

## **TEST**

Returns a new tensor with boolean elements representing if each element is NaN or not.

torch\_isneginf 437

## **Examples**

```
if (torch_is_installed()) {
  torch_isnan(torch_tensor(c(1, NaN, 2)))
}
```

torch\_isneginf

Isneginf

# Description

Isneginf

## Usage

```
torch_isneginf(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# isneginf(input, \*, out=None) -> Tensor

Tests if each element of input is negative infinity or not.

## **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(c(-Inf, Inf, 1.2))
torch_isneginf(a)
}</pre>
```

 $torch\_isposinf$ 

 ${\it Is posinf}$ 

## Description

Isposinf

# Usage

```
torch_isposinf(self)
```

## Arguments

self

(Tensor) the input tensor.

438 torch\_isreal

## isposinf(input, \*, out=None) -> Tensor

Tests if each element of input is positive infinity or not.

# Examples

```
if (torch_is_installed()) {
a <- torch_tensor(c(-Inf, Inf, 1.2))
torch_isposinf(a)
}</pre>
```

torch\_isreal

Is real

# Description

Isreal

## Usage

```
torch_isreal(self)
```

## **Arguments**

self

(Tensor) the input tensor.

#### isreal(input) -> Tensor

Returns a new tensor with boolean elements representing if each element of input is real-valued or not. All real-valued types are considered real. Complex values are considered real when their imaginary part is 0.

```
if (torch_is_installed()) {
  if (FALSE) {
  torch_isreal(torch_tensor(c(1, 1+1i, 2+0i)))
  }
}
```

torch\_istft 439

# Description

Inverse short time Fourier Transform. This is expected to be the inverse of torch\_stft().

## Usage

```
torch_istft(
   self,
   n_fft,
   hop_length = NULL,
   win_length = NULL,
   window = list(),
   center = TRUE,
   normalized = FALSE,
   onesided = NULL,
   length = NULL,
   return_complex = FALSE
)
```

## **Arguments**

self	(Tensor) The input tensor. Expected to be output of torch_stft(), can either be complex (channel, fft_size, n_frame), or real (channel, fft_size, n_frame, 2) where the channel dimension is optional.
n_fft	(int) Size of Fourier transform
hop_length	(Optional[int]) The distance between neighboring sliding window frames. (Default: n_fft %% 4)
win_length	(Optional[int]) The size of window frame and STFT filter. (Default: $n_{fft}$ )
window	$(Optional (torch. Tensor)) \ The \ optional \ window \ function. \ (Default: \ torch\_ones(win\_length))$
center	(bool) Whether input was padded on both sides so that the $t$ -th frame is centered at time $t \times \text{hop\_length}$ . (Default: TRUE)
normalized	(bool) Whether the STFT was normalized. (Default: FALSE)
onesided	(Optional(bool)) Whether the STFT was onesided. (Default: TRUE if n_fft != fft_size in the input size)
length	(Optional(int)]) The amount to trim the signal by (i.e. the original signal length). (Default: whole signal)
return_complex	(Optional(bool)) Whether the output should be complex, or if the input should be assumed to derive from a real signal and window. Note that this is incompatible with onesided=TRUE. (Default: FALSE)

440 torch\_is\_complex

#### **Details**

It has the same parameters (+ additional optional parameter of length) and it should return the least squares estimation of the original signal. The algorithm will check using the NOLA condition (nonzero overlap).

Important consideration in the parameters window and center so that the envelop created by the summation of all the windows is never zero at certain point in time. Specifically,  $\sum_{t=-\infty}^{\infty} |w|^2 (n-t \times hop_length) \neq 0$ .

Since torch\_stft() discards elements at the end of the signal if they do not fit in a frame, istft may return a shorter signal than the original signal (can occur if center is FALSE since the signal isn't padded).

If center is TRUE, then there will be padding e.g. 'constant', 'reflect', etc. Left padding can be trimmed off exactly because they can be calculated but right padding cannot be calculated without additional information.

Example: Suppose the last window is: [c(17, 18, 0, 0, 0) vs c(18, 0, 0, 0, 0)]

The n\_fft, hop\_length, win\_length are all the same which prevents the calculation of right padding. These additional values could be zeros or a reflection of the signal so providing length could be useful. If length is None then padding will be aggressively removed (some loss of signal).

D. W. Griffin and J. S. Lim, "Signal estimation from modified short-time Fourier transform," IEEE Trans. ASSP, vol.32, no.2, pp.236-243, Apr. 1984.

torch\_is\_complex

Is\_complex

#### **Description**

Is complex

#### Usage

torch\_is\_complex(self)

## **Arguments**

self

(Tensor) the PyTorch tensor to test

#### is\_complex(input) -> (bool)

Returns TRUE if the data type of input is a complex data type i.e., one of torch\_complex64, and torch.complex128.

torch\_is\_floating\_point

```
torch\_is\_floating\_point \\ \textit{Is\_floating\_point}
```

# Description

Is\_floating\_point

## Usage

```
torch_is_floating_point(self)
```

## Arguments

self

(Tensor) the PyTorch tensor to test

# $is\_floating\_point(input) \rightarrow (bool)$

Returns TRUE if the data type of input is a floating point data type i.e., one of torch\_float64, torch.float32 and torch.float16.

torch\_is\_installed

Verifies if torch is installed

# Description

Verifies if torch is installed

```
torch_is_installed()
```

442 torch\_kaiser\_window

torch\_is\_nonzero

Is\_nonzero

## **Description**

Is\_nonzero

## Usage

```
torch_is_nonzero(self)
```

#### **Arguments**

self

(Tensor) the input tensor.

## is\_nonzero(input) -> (bool)

Returns TRUE if the input is a single element tensor which is not equal to zero after type conversions. i.e. not equal to  $torch_tensor(c(0))$  or  $torch_tensor(c(0))$  or  $torch_tensor(c(FALSE))$ . Throws a RuntimeError if  $torch_numel()$ != 1 (even in case of sparse tensors).

## **Examples**

```
if (torch_is_installed()) {
  torch_is_nonzero(torch_tensor(c(0.)))
  torch_is_nonzero(torch_tensor(c(1.5)))
  torch_is_nonzero(torch_tensor(c(FALSE)))
  torch_is_nonzero(torch_tensor(c(3)))
  if (FALSE) {
    torch_is_nonzero(torch_tensor(c(1, 3, 5)))
    torch_is_nonzero(torch_tensor(c()))
  }
}
```

torch\_kaiser\_window

Kaiser\_window

## **Description**

Kaiser\_window

torch\_kaiser\_window 443

#### Usage

```
torch_kaiser_window(
  window_length,
  periodic,
  beta,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = NULL)
```

#### **Arguments**

window\_length (int) length of the window.

periodic (bool, optional) If TRUE, returns a periodic window suitable for use in spectral

analysis. If FALSE, returns a symmetric window suitable for use in filter design.

beta (float, optional) shape parameter for the window.

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type). If dtype is not given, infer the data type from the other input arguments. If any of start, end, or stop are floating-point, the dtype is inferred to be the default dtype, see ~torch.get\_default\_dtype. Otherwise, the dtype is inferred to be

torch.int64.

layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided.

device (torch.device, optional) the desired device of returned tensor. Default: if

 ${\tt NULL, uses the current device for the default tensor type (see {\tt torch\_set\_default\_tensor\_type)}.}$ 

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

kaiser\_window(window\_length, periodic=TRUE, beta=12.0, \*, dtype=None, layout=torch.strided, device=None, requires\_grad=FALSE) -> Tensor

Computes the Kaiser window with window length window\_length and shape parameter beta.

Let  $I_0$  be the zeroth order modified Bessel function of the first kind (see torch\_i0()) and N = L - 1 if periodic is FALSE and L if periodic is TRUE, where L is the window\_length. This function computes:

$$out_i = I_0 \left(\beta \sqrt{1 - \left(\frac{i - N/2}{N/2}\right)^2}\right) / I_0(\beta)$$

Calling torch\_kaiser\_window(L, B, periodic=TRUE) is equivalent to calling torch\_kaiser\_window(L + 1, B, periodic The periodic argument is intended as a helpful shorthand to produce a periodic window as input to functions like torch\_stft().

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#### Note

If window\_length is one, then the returned window is a single element tensor containing a one.

torch\_kron Kronecker product

## **Description**

Computes the Kronecker product of self and other.

## Usage

```
torch_kron(self, other)
```

## Arguments

self (Tensor) input Tensor other (Tensor) other tensor.

torch\_kthvalue K

Kthvalue

## **Description**

Kthvalue

#### Usage

```
torch_kthvalue(self, k, dim = -1L, keepdim = FALSE)
```

#### **Arguments**

self (Tensor) the input tensor.

k (int) k for the k-th smallest element

dim (int, optional) the dimension to find the kth value along keepdim (bool) whether the output tensor has dim retained or not.

## kthvalue(input, k, dim=NULL, keepdim=False, out=NULL) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the k th smallest element of each row of the input tensor in the given dimension dim. And indices is the index location of each element found.

If dim is not given, the last dimension of the input is chosen.

If keepdim is TRUE, both the values and indices tensors are the same size as input, except in the dimension dim where they are of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in both the values and indices tensors having 1 fewer dimension than the input tensor.

torch\_layout 445

## **Examples**

```
if (torch_is_installed()) {

x <- torch_arange(1, 6)
x
torch_kthvalue(x, 4)
x <- torch_arange(1,6)$resize_(c(2,3))
x
torch_kthvalue(x, 2, 1, TRUE)
}</pre>
```

torch\_layout

Creates the corresponding layout

## **Description**

Creates the corresponding layout

## Usage

```
torch_strided()
torch_sparse_coo()
```

 $torch_lcm$ 

Lcm

## Description

Lcm

## Usage

```
torch_lcm(self, other)
```

## **Arguments**

self (Tensor) the input tensor.

other (Tensor) the second input tensor

# lcm(input, other, \*, out=None) -> Tensor

Computes the element-wise least common multiple (LCM) of input and other.

Both input and other must have integer types.

torch\_le

## Note

This defines lcm(0,0) = 0 and lcm(0,a) = 0.

#### **Examples**

```
if (torch_is_installed()) {
   if (torch::cuda_is_available()) {
    a <- torch_tensor(c(5, 10, 15), dtype = torch_long(), device = "cuda")
   b <- torch_tensor(c(3, 4, 5), dtype = torch_long(), device = "cuda")
   torch_lcm(a, b)
   c <- torch_tensor(c(3L), device = "cuda")
   torch_lcm(a, c)
}
</pre>
```

torch\_le

Le

## Description

Le

## Usage

```
torch_le(self, other)
```

## Arguments

```
self (Tensor) the tensor to compare
other (Tensor or float) the tensor or value to compare
```

#### le(input, other, out=NULL) -> Tensor

Computes input  $\leq$  other element-wise.

The second argument can be a number or a tensor whose shape is broadcastable with the first argument.

torch\_lerp 447

torch\_lerp

Lerp

## Description

Lerp

#### Usage

```
torch_lerp(self, end, weight)
```

## **Arguments**

```
self (Tensor) the tensor with the starting points
end (Tensor) the tensor with the ending points
weight (float or tensor) the weight for the interpolation formula
```

## lerp(input, end, weight, out=NULL)

Does a linear interpolation of two tensors start (given by input) and end based on a scalar or tensor weight and returns the resulting out tensor.

```
\operatorname{out}_i = \operatorname{start}_i + \operatorname{weight}_i \times (\operatorname{end}_i - \operatorname{start}_i)
```

The shapes of start and end must be broadcastable . If weight is a tensor, then the shapes of weight, start, and end must be broadcastable .

```
if (torch_is_installed()) {

start = torch_arange(1, 4)
end = torch_empty(4)$fill_(10)
start
end
torch_lerp(start, end, 0.5)
torch_lerp(start, end, torch_full_like(start, 0.5))
}
```

448 torch\_less\_equal

torch\_less

Less

## Description

Less

## Usage

```
torch_less(self, other)
```

## Arguments

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

## less(input, other, \*, out=None) -> Tensor

Alias for torch\_lt().

torch\_less\_equal

Less\_equal

## Description

```
Less_equal
```

## Usage

```
torch_less_equal(self, other)
```

## Arguments

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

## less\_equal(input, other, \*, out=None) -> Tensor

```
Alias for torch_le().
```

torch\_lgamma 449

torch\_lgamma

Lgamma

## Description

Lgamma

## Usage

```
torch_lgamma(self)
```

## Arguments

self

(Tensor) the input tensor.

## lgamma(input, out=NULL) -> Tensor

Computes the logarithm of the gamma function on input.

$$\mathsf{out}_i = \log \Gamma(\mathsf{input}_i)$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_arange(0.5, 2, 0.5)
torch_lgamma(a)
}
```

torch\_linspace

Linspace

## Description

Linspace

```
torch_linspace(
   start,
   end,
   steps = 100,
   dtype = NULL,
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

450 torch\_load

## **Arguments**

start	(float) the starting value for the set of points
end	(float) the ending value for the set of points
steps	(int) number of points to sample between start and end. Default: 100.
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch_set_default_tensor_type).
layout	(torch.layout, optional) the desired layout of returned Tensor. Default: torch_strided.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

# $linspace (start, end, steps=100, out=NULL, dtype=NULL, layout=torch. strided, device=NULL, requires\_grad=False) -> Tensor$

Returns a one-dimensional tensor of steps equally spaced points between start and end.

The output tensor is 1-D of size steps.

## **Examples**

```
if (torch_is_installed()) {
  torch_linspace(3, 10, steps=5)
  torch_linspace(-10, 10, steps=5)
  torch_linspace(start=-10, end=10, steps=5)
  torch_linspace(start=-10, end=10, steps=1)
}
```

torch\_load

Loads a saved object

## Description

Loads a saved object

#### Usage

```
torch_load(path, device = "cpu")
```

#### **Arguments**

path a path to the saved object

device a device to load tensors to. By default we load to the cpu but you can also load

them to any cuda device. If NULL then the device where the tensor has been

saved will be reused.

torch\_log 451

## See Also

Other torch\_save: torch\_save(), torch\_serialize()

torch\_log

Log

## Description

Log

## Usage

```
torch_log(self)
```

## Arguments

self

(Tensor) the input tensor.

# log(input, out=NULL) -> Tensor

Returns a new tensor with the natural logarithm of the elements of input.

$$y_i = \log_e(x_i)$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(5))
a
torch_log(a)
}
```

torch\_log10

Log10

# Description

Log10

```
torch_log10(self)
```

452 torch\_log1p

## **Arguments**

self

(Tensor) the input tensor.

## log10(input, out=NULL) -> Tensor

Returns a new tensor with the logarithm to the base 10 of the elements of input.

$$y_i = \log_{10}(x_i)$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_rand(5)
a
torch_log10(a)
}
```

torch\_log1p

Log1p

## Description

Log1p

## Usage

```
torch_log1p(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# log1p(input, out=NULL) -> Tensor

Returns a new tensor with the natural logarithm of (1 + input).

$$y_i = \log_e(x_i + 1)$$

#### Note

This function is more accurate than torch\_log for small values of input

torch\_log2 453

# Examples

```
if (torch_is_installed()) {
a = torch_randn(c(5))
a
torch_log1p(a)
}
```

torch\_log2

Log2

# Description

Log2

# Usage

```
torch_log2(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# log2(input, out=NULL) -> Tensor

Returns a new tensor with the logarithm to the base 2 of the elements of input.

$$y_i = \log_2(x_i)$$

```
if (torch_is_installed()) {
a = torch_rand(5)
a
torch_log2(a)
}
```

454 torch\_logaddexp2

torch\_logaddexp

Logaddexp

#### **Description**

Logaddexp

#### Usage

```
torch_logaddexp(self, other)
```

#### **Arguments**

```
self (Tensor) the input tensor.
other (Tensor) the second input tensor
```

## logaddexp(input, other, \*, out=None) -> Tensor

Logarithm of the sum of exponentiations of the inputs.

Calculates pointwise  $\log{(e^x + e^y)}$ . This function is useful in statistics where the calculated probabilities of events may be so small as to exceed the range of normal floating point numbers. In such cases the logarithm of the calculated probability is stored. This function allows adding probabilities stored in such a fashion.

This op should be disambiguated with torch\_logsumexp() which performs a reduction on a single tensor.

## **Examples**

```
if (torch_is_installed()) { torch_logaddexp(torch_tensor(c(-1.0)), torch_tensor(c(-1.0, -2, -3))) \\ torch_logaddexp(torch_tensor(c(-100.0, -200, -300)), torch_tensor(c(-1.0, -2, -3))) \\ torch_logaddexp(torch_tensor(c(1.0, 2000, 30000)), torch_tensor(c(-1.0, -2, -3))) \\ \}
```

torch\_logaddexp2

Logaddexp2

#### **Description**

Logaddexp2

```
torch_logaddexp2(self, other)
```

torch\_logcumsumexp 455

#### **Arguments**

self (Tensor) the input tensor.

other (Tensor) the second input tensor

## logaddexp2(input, other, \*, out=None) -> Tensor

Logarithm of the sum of exponentiations of the inputs in base-2.

Calculates pointwise  $\log_2(2^x + 2^y)$ . See torch\_logaddexp() for more details.

torch\_logcumsumexp

Logcumsumexp

#### **Description**

Logcumsumexp

#### Usage

```
torch_logcumsumexp(self, dim)
```

## **Arguments**

self (Tensor) the input tensor.

dim (int) the dimension to do the operation over

# logcumsumexp(input, dim, \*, out=None) -> Tensor

Returns the logarithm of the cumulative summation of the exponentiation of elements of input in the dimension dim.

For summation index j given by dim and other indices i, the result is

$$\operatorname{logcumsumexp}(x)_{ij} = \log \sum_{j=0}^{i} \exp(x_{ij})$$

```
if (torch_is_installed()) {
a <- torch_randn(c(10))
torch_logcumsumexp(a, dim=1)
}</pre>
```

456 torch\_logdet

torch\_logdet

Logdet

## **Description**

Logdet

#### Usage

```
torch_logdet(self)
```

## Arguments

self

(Tensor) the input tensor of size (\*, n, n) where \* is zero or more batch dimensions.

#### logdet(input) -> Tensor

Calculates log determinant of a square matrix or batches of square matrices.

## Note

```
Result is '-inf' if 'input' has zero log determinant, and is 'NaN' if 'input' has negative determinant.
```

Backward through `logdet` internally uses SVD results when `input` is not invertible. In this case, double backward through `logdet` will be unstable in when `input` doesn't have distinct singular values. See `~torch.svd` for details.

```
if (torch_is_installed()) {
A = torch_randn(c(3, 3))
torch_det(A)
torch_logdet(A)
A
A$det()
A$det()$log()
}
```

torch\_logical\_and 457

torch\_logical\_and

Logical\_and

#### **Description**

Logical\_and

#### Usage

```
torch_logical_and(self, other)
```

## Arguments

```
self (Tensor) the input tensor.
```

other (Tensor) the tensor to compute AND with

## logical\_and(input, other, out=NULL) -> Tensor

Computes the element-wise logical AND of the given input tensors. Zeros are treated as FALSE and nonzeros are treated as TRUE.

## **Examples**

```
if (torch_is_installed()) {

torch_logical_and(torch_tensor(c(TRUE, FALSE, TRUE)), torch_tensor(c(TRUE, FALSE, FALSE)))
a = torch_tensor(c(0, 1, 10, 0), dtype=torch_int8())
b = torch_tensor(c(4, 0, 1, 0), dtype=torch_int8())
torch_logical_and(a, b)
## Not run:
torch_logical_and(a, b, out=torch_empty(4, dtype=torch_bool()))
## End(Not run)
}
```

torch\_logical\_not

Logical\_not

## **Description**

Logical\_not

## **Arguments**

self

(Tensor) the input tensor.

458 torch\_logical\_or

#### logical\_not(input, out=NULL) -> Tensor

Computes the element-wise logical NOT of the given input tensor. If not specified, the output tensor will have the bool dtype. If the input tensor is not a bool tensor, zeros are treated as FALSE and non-zeros are treated as TRUE.

#### **Examples**

```
if (torch_is_installed()) {

torch_logical_not(torch_tensor(c(TRUE, FALSE)))

torch_logical_not(torch_tensor(c(0, 1, -10), dtype=torch_int8()))

torch_logical_not(torch_tensor(c(0., 1.5, -10.), dtype=torch_double()))
}
```

torch\_logical\_or

 $Logical\_or$ 

#### **Description**

```
Logical_or
```

## Usage

```
torch_logical_or(self, other)
```

# **Arguments**

```
self (Tensor) the input tensor.
other (Tensor) the tensor to compute OR with
```

#### logical\_or(input, other, out=NULL) -> Tensor

Computes the element-wise logical OR of the given input tensors. Zeros are treated as FALSE and nonzeros are treated as TRUE.

```
if (torch_is_installed()) {

torch_logical_or(torch_tensor(c(TRUE, FALSE, TRUE)), torch_tensor(c(TRUE, FALSE, FALSE)))
a = torch_tensor(c(0, 1, 10, 0), dtype=torch_int8())
b = torch_tensor(c(4, 0, 1, 0), dtype=torch_int8())
torch_logical_or(a, b)
## Not run:
torch_logical_or(a$double(), b$double())
torch_logical_or(a$double(), b)
torch_logical_or(a, b, out=torch_empty(4, dtype=torch_bool()))
## End(Not run)
}
```

torch\_logical\_xor 459

torch\_logical\_xor

Logical\_xor

## **Description**

```
Logical_xor
```

## Usage

```
torch_logical_xor(self, other)
```

## Arguments

self (Tensor) the input tensor.

other (Tensor) the tensor to compute XOR with

## logical\_xor(input, other, out=NULL) -> Tensor

Computes the element-wise logical XOR of the given input tensors. Zeros are treated as FALSE and nonzeros are treated as TRUE.

## **Examples**

```
if (torch_is_installed()) {

torch_logical_xor(torch_tensor(c(TRUE, FALSE, TRUE)), torch_tensor(c(TRUE, FALSE, FALSE)))
a = torch_tensor(c(0, 1, 10, 0), dtype=torch_int8())
b = torch_tensor(c(4, 0, 1, 0), dtype=torch_int8())
torch_logical_xor(a, b)
torch_logical_xor(a$to(dtype=torch_double()), b$to(dtype=torch_double()))
torch_logical_xor(a$to(dtype=torch_double()), b)
}
```

torch\_logit

Logit

## **Description**

Logit

```
torch_logit(self, eps = NULL)
```

460 torch\_logspace

#### **Arguments**

```
self (Tensor) the input tensor.
eps (float, optional) the epsilon for input clamp bound. Default: None
```

## logit(input, eps=None, \*, out=None) -> Tensor

Returns a new tensor with the logit of the elements of input. input is clamped to [eps, 1 - eps] when eps is not None. When eps is None and input < 0 or input > 1, the function will yields NaN.

$$y_i = \ln(\frac{z_i}{1-z_i})z_i = \begin{cases} x_i & \text{if eps is None} \\ \text{eps} & \text{if } x_i < \text{eps} \\ x_i & \text{if eps} \le x_i \le 1 - \text{eps} \\ 1 - \text{eps} & \text{if } x_i > 1 - \text{eps} \end{cases}$$

## **Examples**

```
if (torch_is_installed()) {
a <- torch_rand(5)
a
torch_logit(a, eps=1e-6)
}</pre>
```

 $torch\_logspace$ 

Logspace

## **Description**

Logspace

```
torch_logspace(
   start,
   end,
   steps = 100,
   base = 10,
   dtype = NULL,
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

torch\_logsumexp 461

## **Arguments**

start	(float) the starting value for the set of points
end	(float) the ending value for the set of points
steps	(int) number of points to sample between start and end. Default: 100.
base	(float) base of the logarithm function. Default: 10.0.
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch_set_default_tensor_type).
layout	(torch.layout, optional) the desired layout of returned Tensor. Default: torch_strided.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

# $log space (start,\ end,\ steps=100,\ base=10.0,\ out=NULL,\ dtype=NULL,\ layout=torch.strided,\ device=NULL,\ requires\_grad=False) -> Tensor$

Returns a one-dimensional tensor of steps points logarithmically spaced with base base between base start and base end.

The output tensor is 1-D of size steps.

## **Examples**

```
if (torch_is_installed()) {
torch_logspace(start=-10, end=10, steps=5)
torch_logspace(start=0.1, end=1.0, steps=5)
torch_logspace(start=0.1, end=1.0, steps=1)
torch_logspace(start=2, end=2, steps=1, base=2)
}
```

## **Description**

Logsumexp

```
torch_logsumexp(self, dim, keepdim = FALSE)
```

462 torch\_lstsq

#### **Arguments**

self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

#### logsumexp(input, dim, keepdim=False, out=NULL)

Returns the log of summed exponentials of each row of the input tensor in the given dimension dim. The computation is numerically stabilized.

For summation index j given by dim and other indices i, the result is

$$\operatorname{logsumexp}(x)_i = \log \sum_j \exp(x_{ij})$$

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3, 3))
torch_logsumexp(a, 1)
}
```

torch\_lstsq

Lstsq

## **Description**

Lstsq

## **Arguments**

self (Tensor) the matrix B

A (Tensor) the m by n matrix A

## lstsq(input, A, out=NULL) -> Tensor

Computes the solution to the least squares and least norm problems for a full rank matrix A of size  $(m \times n)$  and a matrix B of size  $(m \times k)$ .

If  $m \ge n$ , torch\_lstsq() solves the least-squares problem:

$$\min_X \|AX - B\|_2$$
.

torch\_lt 463

If m < n, torch\_lstsq() solves the least-norm problem:

```
\min_X \|X\|_2 subject to AX = B.
```

Returned tensor X has shape  $(\max(m,n) \times k)$ . The first n rows of X contains the solution. If  $m \ge n$ , the residual sum of squares for the solution in each column is given by the sum of squares of elements in the remaining m-n rows of that column.

#### Note

The case when  $eqn\{m < n\}$  is not supported on the GPU.

torch\_lt

Lt

## **Description**

Lt

#### Usage

```
torch_lt(self, other)
```

## **Arguments**

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

## lt(input, other, out=NULL) -> Tensor

Computes input < other element-wise.

The second argument can be a number or a tensor whose shape is broadcastable with the first argument.

464 torch\_lu\_solve

torch\_lu

LU

## Description

Computes the LU factorization of a matrix or batches of matrices A. Returns a tuple containing the LU factorization and pivots of A. Pivoting is done if pivot is set to True.

#### Usage

```
torch_lu(A, pivot = TRUE, get_infos = FALSE, out = NULL)
```

## **Arguments**

A (Tensor) the tensor to factor of size (, m, n)(,m,n)

pivot (bool, optional) – controls whether pivoting is done. Default: TRUE

get\_infos (bool, optional) – if set to True, returns an info IntTensor. Default: FALSE

out (tuple, optional) – optional output tuple. If get\_infos is True, then the elements

in the tuple are Tensor, IntTensor, and IntTensor. If get\_infos is False, then the

elements in the tuple are Tensor, IntTensor. Default: NULL

## **Examples**

```
if (torch_is_installed()) {
A <- torch_randn(c(2, 3, 3))
torch_lu(A)
}</pre>
```

torch\_lu\_solve

Lu\_solve

## **Description**

Lu\_solve

```
torch_lu_solve(self, LU_data, LU_pivots)
```

torch\_lu\_unpack 465

## **Arguments**

self	(Tensor) the RHS tensor of size $(*,m,k)$ , where $*$ is zero or more batch dimensions.
LU_data	(Tensor) the pivoted LU factorization of A from torch_lu of size $(*,m,m)$ , where $*$ is zero or more batch dimensions.
LU_pivots	(IntTensor) the pivots of the LU factorization from torch_lu of size $(*, m)$ , where $*$ is zero or more batch dimensions. The batch dimensions of LU_pivots must be equal to the batch dimensions of LU data.

# lu\_solve(input, LU\_data, LU\_pivots, out=NULL) -> Tensor

Returns the LU solve of the linear system Ax = b using the partially pivoted LU factorization of A from torch\_lu.

## **Examples**

```
if (torch_is_installed()) {
A = torch_randn(c(2, 3, 3))
b = torch_randn(c(2, 3, 1))
out = torch_lu(A)
x = torch_lu_solve(b, out[[1]], out[[2]])
torch_norm(torch_bmm(A, x) - b)
}
```

# Description

Lu\_unpack

# Usage

```
torch_lu_unpack(LU_data, LU_pivots, unpack_data = TRUE, unpack_pivots = TRUE)
```

# Arguments

LU_data	(Tensor) – the packed LU factorization data
LU_pivots	(Tensor) - the packed LU factorization pivots
unpack_data	$(logical)-flag\ indicating\ if\ the\ data\ should\ be\ unpacked.\ If\ FALSE,\ then\ the\ returned\ L\ and\ U\ are\ NULL\ Default:\ TRUE$
unpack_pivots	(logical) – flag indicating if the pivots should be unpacked into a permutation matrix P. If FALSE, then the returned P is None. Default: TRUE

466 torch\_manual\_seed

#### lu\_unpack(LU\_data, LU\_pivots, unpack\_data = TRUE, unpack\_pivots=TRUE) -> Tensor

Unpacks the data and pivots from a LU factorization of a tensor into tensors L and U and a permutation tensor P such that LU\_data\_and\_pivots <- torch\_lu(P\$matmul(L)\$matmul(U)). Returns a list of tensors as list(the P tensor (permutation matrix), the L tensor, the U tensor)

torch\_manual\_seed

Sets the seed for generating random numbers.

## Description

Sets the seed for generating random numbers.

## Usage

```
torch_manual_seed(seed)
local_torch_manual_seed(seed, .env = parent.frame())
with_torch_manual_seed(code, ..., seed)
```

#### **Arguments**

```
seed integer seed.

.env environment that will take the modifications from manual_seed.

code expression to run in the context of the seed

... unused currently.
```

#### **Functions**

- local\_torch\_manual\_seed(): Modifies the torch seed in the environment scope.
- with\_torch\_manual\_seed(): A with context to change the seed during the function execution.

#### Note

Currently the local\_torch\_manual\_seed and with\_torch\_manual\_seed won't work with Tensors in the MPS device. You can sample the tensors on CPU and move them to MPS if reproducibility is required.

torch\_masked\_select 467

## Description

Masked\_select

#### Usage

```
torch_masked_select(self, mask)
```

## **Arguments**

```
self (Tensor) the input tensor.
```

mask (BoolTensor) the tensor containing the binary mask to index with

## masked\_select(input, mask, out=NULL) -> Tensor

Returns a new 1-D tensor which indexes the input tensor according to the boolean mask mask which is a BoolTensor.

The shapes of the mask tensor and the input tensor don't need to match, but they must be broadcastable .

#### Note

The returned tensor does not use the same storage as the original tensor

```
if (torch_is_installed()) {
x = torch_randn(c(3, 4))
x
mask = x$ge(0.5)
mask
torch_masked_select(x, mask)
}
```

468 torch\_matmul

torch\_matmul

Matmul

#### Description

Matmul

#### Usage

```
torch_matmul(self, other)
```

#### **Arguments**

self (Tensor) the first tensor to be multiplied other (Tensor) the second tensor to be multiplied

## matmul(input, other, out=NULL) -> Tensor

Matrix product of two tensors.

The behavior depends on the dimensionality of the tensors as follows:

- If both tensors are 1-dimensional, the dot product (scalar) is returned.
- If both arguments are 2-dimensional, the matrix-matrix product is returned.
- If the first argument is 1-dimensional and the second argument is 2-dimensional, a 1 is prepended to its dimension for the purpose of the matrix multiply. After the matrix multiply, the prepended dimension is removed.
- If the first argument is 2-dimensional and the second argument is 1-dimensional, the matrix-vector product is returned.
- If both arguments are at least 1-dimensional and at least one argument is N-dimensional (where N > 2), then a batched matrix multiply is returned. If the first argument is 1-dimensional, a 1 is prepended to its dimension for the purpose of the batched matrix multiply and removed after. If the second argument is 1-dimensional, a 1 is appended to its dimension for the purpose of the batched matrix multiple and removed after. The non-matrix (i.e. batch) dimensions are broadcasted (and thus must be broadcastable). For example, if input is a  $(j \times 1 \times n \times m)$  tensor and other is a  $(k \times m \times p)$  tensor, out will be an  $(j \times k \times n \times p)$  tensor.

#### Note

The 1-dimensional dot product version of this function does not support an `out` parameter.

torch\_matrix\_exp 469

### **Examples**

```
if (torch_is_installed()) {
# vector x vector
tensor1 = torch_randn(c(3))
tensor2 = torch_randn(c(3))
torch_matmul(tensor1, tensor2)
# matrix x vector
tensor1 = torch_randn(c(3, 4))
tensor2 = torch_randn(c(4))
torch_matmul(tensor1, tensor2)
# batched matrix x broadcasted vector
tensor1 = torch_randn(c(10, 3, 4))
tensor2 = torch_randn(c(4))
torch_matmul(tensor1, tensor2)
# batched matrix x batched matrix
tensor1 = torch_randn(c(10, 3, 4))
tensor2 = torch_randn(c(10, 4, 5))
torch_matmul(tensor1, tensor2)
# batched matrix x broadcasted matrix
tensor1 = torch_randn(c(10, 3, 4))
tensor2 = torch_randn(c(4, 5))
torch_matmul(tensor1, tensor2)
}
```

torch\_matrix\_exp

Matrix\_exp

# **Description**

Matrix exp

# Usage

```
torch_matrix_exp(self)
```

### **Arguments**

self

(Tensor) the input tensor.

## matrix\_power(input) -> Tensor

Returns the matrix exponential. Supports batched input. For a matrix A, the matrix exponential is defined as

$$\exp^A = \sum_{k=0}^{\infty} A^k / k!.$$

The implementation is based on: Bader, P.; Blanes, S.; Casas, F. Computing the Matrix Exponential with an Optimized Taylor Polynomial Approximation. Mathematics 2019, 7, 1174.

470 torch\_matrix\_power

### **Examples**

```
if (torch_is_installed()) {
    a <- torch_randn(c(2, 2, 2))
    a[1, , ] <- torch_eye(2, 2)
    a[2, , ] <- 2 * torch_eye(2, 2)
    a
    torch_matrix_exp(a)

x <- torch_tensor(rbind(c(0, pi/3), c(-pi/3, 0)))
x$matrix_exp() # should be [[cos(pi/3), sin(pi/3)], [-sin(pi/3), cos(pi/3)]]
}</pre>
```

torch\_matrix\_power

Matrix\_power

# **Description**

Matrix\_power

## Usage

```
torch_matrix_power(self, n)
```

## **Arguments**

```
self (Tensor) the input tensor.

n (int) the power to raise the matrix to
```

### matrix\_power(input, n) -> Tensor

Returns the matrix raised to the power n for square matrices. For batch of matrices, each individual matrix is raised to the power n.

If n is negative, then the inverse of the matrix (if invertible) is raised to the power n. For a batch of matrices, the batched inverse (if invertible) is raised to the power n. If n is 0, then an identity matrix is returned.

```
if (torch_is_installed()) {
a = torch_randn(c(2, 2, 2))
a
torch_matrix_power(a, 3)
}
```

torch\_matrix\_rank 471

# **Description**

Matrix\_rank

## **Arguments**

self (Tensor) the input 2-D tensor

tol (float, optional) the tolerance value. Default: NULL

symmetric (bool, optional) indicates whether input is symmetric. Default: FALSE

# matrix\_rank(input, tol=NULL, symmetric=False) -> Tensor

Returns the numerical rank of a 2-D tensor. The method to compute the matrix rank is done using SVD by default. If symmetric is TRUE, then input is assumed to be symmetric, and the computation of the rank is done by obtaining the eigenvalues.

tol is the threshold below which the singular values (or the eigenvalues when symmetric is TRUE) are considered to be 0. If tol is not specified, tol is set to S.max()\*max(S.size())\*eps where S is the singular values (or the eigenvalues when symmetric is TRUE), and eps is the epsilon value for the datatype of input.

### **Description**

Max

## **Arguments**

self	(Tensor) the input tensor.
dim	(int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not. Default: FALSE.

out (tuple, optional) the result tuple of two output tensors (max, max\_indices)

other (Tensor) the second input tensor

# max(input) -> Tensor

Returns the maximum value of all elements in the input tensor.

472 torch\_max

## max(input, dim, keepdim=False, out=NULL) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the maximum value of each row of the input tensor in the given dimension dim. And indices is the index location of each maximum value found (argmax).

### Warning

indices does not necessarily contain the first occurrence of each maximal value found, unless it is unique. The exact implementation details are device-specific. Do not expect the same result when run on CPU and GPU in general.

If keepdim is TRUE, the output tensors are of the same size as input except in the dimension dim where they are of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensors having 1 fewer dimension than input.

### max(input, other, out=NULL) -> Tensor

Each element of the tensor input is compared with the corresponding element of the tensor other and an element-wise maximum is taken.

The shapes of input and other don't need to match, but they must be broadcastable.

```
out_i = max(tensor_i, other_i)
```

#### Note

When the shapes do not match, the shape of the returned output tensor follows the broadcasting rules .

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_max(a)

a = torch_randn(c(4, 4))
a
torch_max(a, dim = 1)

a = torch_randn(c(4))
a
b = torch_randn(c(4))
b
torch_max(a, other = b)
}
```

torch\_maximum 473

torch\_maximum

Maximum

# Description

Maximum

# Usage

```
torch_maximum(self, other)
```

# **Arguments**

```
self (Tensor) the input tensor.
other (Tensor) the second input tensor
```

# maximum(input, other, \*, out=None) -> Tensor

Computes the element-wise maximum of input and other.

### Note

If one of the elements being compared is a NaN, then that element is returned. torch\_maximum() is not supported for tensors with complex dtypes.

# **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(c(1, 2, -1))
b <- torch_tensor(c(3, 0, 4))
torch_maximum(a, b)
}</pre>
```

torch\_mean

Mean

# Description

Mean

```
torch_mean(self, dim, keepdim = FALSE, dtype = NULL)
```

474 torch\_median

## **Arguments**

self	(Tensor) the input tensor.
dim	(int or tuple of ints) the dimension or dimensions to reduce.
keepdim	(bool) whether the output tensor has dim retained or not.
dtype	the resulting data type.

# mean(input) -> Tensor

Returns the mean value of all elements in the input tensor.

# mean(input, dim, keepdim=False, out=NULL) -> Tensor

Returns the mean value of each row of the input tensor in the given dimension dim. If dim is a list of dimensions, reduce over all of them.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_mean(a)

a = torch_randn(c(4, 4))
a
torch_mean(a, 1)
torch_mean(a, 1, TRUE)
}
```

torch\_median Median

# **Description**

Median

```
torch_median(self, dim, keepdim = FALSE)
```

torch\_memory\_format 475

## **Arguments**

```
self (Tensor) the input tensor.

dim (int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not.
```

# median(input) -> Tensor

Returns the median value of all elements in the input tensor.

# median(input, dim=-1, keepdim=False, out=NULL) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the median value of each row of the input tensor in the given dimension dim. And indices is the index location of each median value found.

By default, dim is the last dimension of the input tensor.

If keepdim is TRUE, the output tensors are of the same size as input except in the dimension dim where they are of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the outputs tensor having 1 fewer dimension than input.

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_median(a)

a = torch_randn(c(4, 5))
a
torch_median(a, 1)
}
```

torch\_memory\_format

Memory format

# **Description**

Returns the correspondent memory format.

```
torch_contiguous_format()
torch_preserve_format()
torch_channels_last_format()
```

476 torch\_min

torch\_meshgrid

Meshgrid

# Description

Take N tensors, each of which can be either scalar or 1-dimensional vector, and create N N-dimensional grids, where the i th grid is defined by expanding the i th input over dimensions defined by other inputs.

# Usage

```
torch_meshgrid(tensors, indexing)
```

## Arguments

tensors

(list of Tensor) list of scalars or 1 dimensional tensors. Scalars will be treated

(1,).

indexing

(str, optional): the indexing mode, either "xy" or "ij", defaults to "ij". See warning for future changes. If "xy" is selected, the first dimension corresponds to the cardinality of the second input and the second dimension corresponds to the cardinality of the first input. If "ij" is selected, the dimensions are in the same order as the cardinality of the inputs.

# Warning

In the future torch\_meshgrid will transition to indexing='xy' as the default. This issue tracks this issue with the goal of migrating to NumPy's behavior.

# **Examples**

```
if (torch_is_installed()) {
x = torch_tensor(c(1, 2, 3))
y = torch_tensor(c(4, 5, 6))
out = torch_meshgrid(list(x, y))
out
}
```

torch\_min

Min

# Description

Min

torch\_min 477

## Arguments

self	(Tensor) the input tensor.
dim	(int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

out (tuple, optional) the tuple of two output tensors (min, min\_indices)

other (Tensor) the second input tensor

## min(input) -> Tensor

Returns the minimum value of all elements in the input tensor.

## min(input, dim, keepdim=False, out=NULL) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the minimum value of each row of the input tensor in the given dimension dim. And indices is the index location of each minimum value found (argmin).

### Warning

indices does not necessarily contain the first occurrence of each minimal value found, unless it is unique. The exact implementation details are device-specific. Do not expect the same result when run on CPU and GPU in general.

If keepdim is TRUE, the output tensors are of the same size as input except in the dimension dim where they are of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensors having 1 fewer dimension than input.

## min(input, other, out=NULL) -> Tensor

Each element of the tensor input is compared with the corresponding element of the tensor other and an element-wise minimum is taken. The resulting tensor is returned.

The shapes of input and other don't need to match, but they must be broadcastable.

```
out_i = min(tensor_i, other_i)
```

#### Note

When the shapes do not match, the shape of the returned output tensor follows the broadcasting rules.

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_min(a)
```

478 torch\_minimum

```
a = torch_randn(c(4, 4))
a
torch_min(a, dim = 1)

a = torch_randn(c(4))
a
b = torch_randn(c(4))
b
torch_min(a, other = b)
}
```

torch\_minimum

Minimum

# Description

Minimum

# Usage

```
torch_minimum(self, other)
```

# **Arguments**

```
self (Tensor) the input tensor.
other (Tensor) the second input tensor
```

# minimum(input, other, \*, out=None) -> Tensor

Computes the element-wise minimum of input and other.

## Note

If one of the elements being compared is a NaN, then that element is returned. torch\_minimum() is not supported for tensors with complex dtypes.

```
if (torch_is_installed()) {
a <- torch_tensor(c(1, 2, -1))
b <- torch_tensor(c(3, 0, 4))
torch_minimum(a, b)
}</pre>
```

torch\_mm 479

torch\_mm

Mm

# Description

Mm

### Usage

```
torch_mm(self, mat2)
```

## **Arguments**

```
self (Tensor) the first matrix to be multiplied
mat2 (Tensor) the second matrix to be multiplied
```

# mm(input, mat2, out=NULL) -> Tensor

Performs a matrix multiplication of the matrices input and mat2.

```
If input is a (n \times m) tensor, mat 2 is a (m \times p) tensor, out will be a (n \times p) tensor.
```

### Note

This function does not broadcast . For broadcasting matrix products, see torch\_matmul.

# **Examples**

```
if (torch_is_installed()) {
mat1 = torch_randn(c(2, 3))
mat2 = torch_randn(c(3, 3))
torch_mm(mat1, mat2)
}
```

torch\_mode

Mode

# Description

Mode

```
torch_mode(self, dim = -1L, keepdim = FALSE)
```

480 torch\_movedim

#### **Arguments**

self (Tensor) the input tensor.
dim (int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

### mode(input, dim=-1, keepdim=False, out=NULL) -> (Tensor, LongTensor)

Returns a namedtuple (values, indices) where values is the mode value of each row of the input tensor in the given dimension dim, i.e. a value which appears most often in that row, and indices is the index location of each mode value found.

By default, dim is the last dimension of the input tensor.

If keepdim is TRUE, the output tensors are of the same size as input except in the dimension dim where they are of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensors having 1 fewer dimension than input.

#### Note

This function is not defined for torch\_cuda. Tensor yet.

# **Examples**

```
if (torch_is_installed()) {
a = torch_randint(0, 50, size = list(5))
a
torch_mode(a, 1)
}
```

torch\_movedim

Movedim

# Description

Movedim

### Usage

```
torch_movedim(self, source, destination)
```

## **Arguments**

self (Tensor) the input tensor.

source (int or tuple of ints) Original positions of the dims to move. These must be

inique

destination (int or tuple of ints) Destination positions for each of the original dims. These

must also be unique.

torch\_mul 481

### movedim(input, source, destination) -> Tensor

Moves the dimension(s) of input at the position(s) in source to the position(s) in destination.

Other dimensions of input that are not explicitly moved remain in their original order and appear at the positions not specified in destination.

### **Examples**

```
if (torch_is_installed()) {
t <- torch_randn(c(3,2,1))
t
torch_movedim(t, 2, 1)$shape
torch_movedim(t, 2, 1)
torch_movedim(t, c(2, 3), c(1, 2))$shape
torch_movedim(t, c(2, 3), c(1, 2))
}</pre>
```

torch\_mul

Mul

# **Description**

Mul

## Usage

```
torch_mul(self, other)
```

# Arguments

```
self (Tensor) the first multiplicand tensor
other (Tensor) the second multiplicand tensor
```

# mul(input, other, out=NULL)

Multiplies each element of the input input with the scalar other and returns a new resulting tensor.

```
out_i = other \times input_i
```

If input is of type FloatTensor or DoubleTensor, other should be a real number, otherwise it should be an integer

Each element of the tensor input is multiplied by the corresponding element of the Tensor other. The resulting tensor is returned.

The shapes of input and other must be broadcastable.

```
out_i = input_i \times other_i
```

482 torch\_multinomial

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3))
a
torch_mul(a, 100)

a = torch_randn(c(4, 1))
a
b = torch_randn(c(1, 4))
b
torch_mul(a, b)
}
```

torch\_multinomial

Multinomial

# **Description**

Multinomial

# Usage

```
torch_multinomial(self, num_samples, replacement = FALSE, generator = NULL)
```

## **Arguments**

```
self (Tensor) the input tensor containing probabilities
```

num\_samples (int) number of samples to draw

replacement (bool, optional) whether to draw with replacement or not

generator (torch.Generator, optional) a pseudorandom number generator for sampling

# $multinomial (input, num\_samples, replacement = False, *, generator = NULL, out = NULL) -> Long Tensor$

Returns a tensor where each row contains num\_samples indices sampled from the multinomial probability distribution located in the corresponding row of tensor input.

## Note

```
The rows of `input` do not need to sum to one (in which case we use
the values as weights), but must be non-negative, finite and have
a non-zero sum.
```

Indices are ordered from left to right according to when each was sampled (first samples are placed in first column).

If input is a vector, out is a vector of size num\_samples.

torch\_multiply 483

If input is a matrix with m rows, out is an matrix of shape  $(m \times \text{num\_samples})$ .

If replacement is TRUE, samples are drawn with replacement.

If not, they are drawn without replacement, which means that when a sample index is drawn for a row, it cannot be drawn again for that row.

```
When drawn without replacement, `num_samples` must be lower than number of non-zero elements in `input` (or the min number of non-zero elements in each row of `input` if it is a matrix).
```

# **Examples**

```
if (torch_is_installed()) {
weights = torch_tensor(c(0, 10, 3, 0), dtype=torch_float()) # create a tensor of weights
torch_multinomial(weights, 2)
torch_multinomial(weights, 4, replacement=TRUE)
}
```

torch\_multiply

Multiply

# **Description**

Multiply

# Usage

```
torch_multiply(self, other)
```

# Arguments

```
self (Tensor) the first multiplicand tensor
other (Tensor) the second multiplicand tensor
```

# multiply(input, other, \*, out=None)

```
Alias for torch_mul().
```

484 torch\_mvlgamma

torch\_mv

Mv

# Description

Mv

# Usage

```
torch_mv(self, vec)
```

# **Arguments**

```
self (Tensor) matrix to be multiplied vec (Tensor) vector to be multiplied
```

# mv(input, vec, out=NULL) -> Tensor

Performs a matrix-vector product of the matrix input and the vector vec.

If input is a  $(n \times m)$  tensor, vec is a 1-D tensor of size m, out will be 1-D of size n.

# Note

This function does not broadcast.

# **Examples**

```
if (torch_is_installed()) {
mat = torch_randn(c(2, 3))
vec = torch_randn(c(3))
torch_mv(mat, vec)
}
```

torch\_mvlgamma

Mvlgamma

# Description

Mvlgamma

```
torch_mvlgamma(self, p)
```

torch\_nanquantile 485

## Arguments

self (Tensor) the tensor to compute the multivariate log-gamma function p (int) the number of dimensions

### mvlgamma(input, p) -> Tensor

Computes the multivariate log-gamma function <a href="https://en.wikipedia.org/wiki/Multivariate\_gamma\_functio">https://en.wikipedia.org/wiki/Multivariate\_gamma\_functio</a> with dimension p element-wise, given by

$$\log(\Gamma_p(a)) = C + \sum_{i=1}^p \log\left(\Gamma\left(a - \frac{i-1}{2}\right)\right)$$

where  $C = \log(\pi) \times \frac{p(p-1)}{4}$  and  $\Gamma(\cdot)$  is the Gamma function.

All elements must be greater than  $\frac{p-1}{2}$ , otherwise an error would be thrown.

# **Examples**

```
if (torch_is_installed()) {
a = torch_empty(c(2, 3))$uniform_(1, 2)
a
torch_mvlgamma(a, 2)
}
```

torch\_nanguantile

Nanquantile

# Description

Nanquantile

### Usage

```
torch_nanquantile(
  self,
  q,
  dim = NULL,
  keepdim = FALSE,
  interpolation = "linear")
```

# **Arguments**

self (Tensor) the input tensor.

q (float or Tensor) a scalar or 1D tensor of quantile values in the range [0, 1]

dim (int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

interpolation The interpolation method.

486 torch\_nansum

## nanquantile(input, q, dim=None, keepdim=FALSE, \*, out=None) -> Tensor

This is a variant of torch\_quantile() that "ignores" NaN values, computing the quantiles q as if NaN values in input did not exist. If all values in a reduced row are NaN then the quantiles for that reduction will be NaN. See the documentation for torch\_quantile().

# **Examples**

```
if (torch_is_installed()) {

t <- torch_tensor(c(NaN, 1, 2))

t$quantile(0.5)

t <- torch_tensor(rbind(c(NaN, NaN), c(1, 2)))

t

t$nanquantile(0.5, dim=1)

t$nanquantile(0.5, dim=2)

torch_nanquantile(t, 0.5, dim = 1)

torch_nanquantile(t, 0.5, dim = 2)
}</pre>
```

torch\_nansum

Nansum

#### **Description**

Nansum

# Usage

```
torch_nansum(self, dim = NULL, keepdim = FALSE, dtype = NULL)
```

### **Arguments**

self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

dtype the desired data type of returned tensor. If specified, the input tensor is casted to

dtype before the operation is performed. This is useful for preventing data type

overflows. Default: NULL.

# nansum(input, \*, dtype=None) -> Tensor

Returns the sum of all elements, treating Not a Numbers (NaNs) as zero.

torch\_narrow 487

## nansum(input, dim, keepdim=FALSE, \*, dtype=None) -> Tensor

Returns the sum of each row of the input tensor in the given dimension dim, treating Not a Numbers (NaNs) as zero. If dim is a list of dimensions, reduce over all of them.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

# **Examples**

```
if (torch_is_installed()) {
    a <- torch_tensor(c(1., 2., NaN, 4.))
    torch_nansum(a)

torch_nansum(torch_tensor(c(1., NaN)))
    a <- torch_tensor(rbind(c(1, 2), c(3., NaN)))
    torch_nansum(a)
    torch_nansum(a, dim=1)
    torch_nansum(a, dim=2)
}</pre>
```

torch\_narrow

Narrow

## Description

Narrow

# Usage

```
torch_narrow(self, dim, start, length)
```

## **Arguments**

```
self (Tensor) the tensor to narrow

dim (int) the dimension along which to narrow

start (int) the starting dimension

length (int) the distance to the ending dimension
```

## narrow(input, dim, start, length) -> Tensor

Returns a new tensor that is a narrowed version of input tensor. The dimension dim is input from start to start + length. The returned tensor and input tensor share the same underlying storage.

488 torch\_ne

# **Examples**

```
if (torch_is_installed()) {
x = torch_tensor(matrix(c(1:9), ncol = 3, byrow= TRUE))
torch_narrow(x, 1, 1, 2)
torch_narrow(x, 2, 2, 2)
}
```

torch\_ne

Ne

# Description

Ne

# Usage

```
torch_ne(self, other)
```

# Arguments

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

# ne(input, other, out=NULL) -> Tensor

Computes  $input \neq other$  element-wise.

The second argument can be a number or a tensor whose shape is broadcastable with the first argument.

torch\_neg 489

torch\_neg

Neg

# Description

Neg

# Usage

```
torch_neg(self)
```

# Arguments

self

(Tensor) the input tensor.

# neg(input, out=NULL) -> Tensor

Returns a new tensor with the negative of the elements of input.

$$out = -1 \times input$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(5))
a
torch_neg(a)
}
```

torch\_negative

Negative

# Description

Negative

# Usage

```
torch_negative(self)
```

# **Arguments**

self

(Tensor) the input tensor.

# negative(input, \*, out=None) -> Tensor

Alias for torch\_neg()

490 torch\_nonzero

torch\_nextafter

Nextafter

# Description

Nextafter

# Usage

```
torch_nextafter(self, other)
```

# **Arguments**

```
self (Tensor) the first input tensor
other (Tensor) the second input tensor
```

# nextafter(input, other, \*, out=None) -> Tensor

Return the next floating-point value after input towards other, elementwise.

The shapes of input and other must be broadcastable .

# **Examples**

```
if (torch_is_installed()) {
eps <- torch_finfo(torch_float32())$eps
torch_nextafter(torch_tensor(c(1, 2)), torch_tensor(c(2, 1))) == torch_tensor(c(eps + 1, 2 - eps))
}</pre>
```

torch\_nonzero

Nonzero

# **Description**

Nonzero elements of tensors.

```
torch_nonzero(self, as_list = FALSE)
```

torch\_norm 491

### **Arguments**

self
as\_list

(Tensor) the input tensor.

If FALSE, the output tensor containing indices. If TRUE, one 1-D tensor for each dimension, containing the indices of each nonzero element along that dimension.

### When as\_list is FALSE (default):

Returns a tensor containing the indices of all non-zero elements of input. Each row in the result contains the indices of a non-zero element in input. The result is sorted lexicographically, with the last index changing the fastest (C-style).

If input has n dimensions, then the resulting indices tensor out is of size  $(z \times n)$ , where z is the total number of non-zero elements in the input tensor.

#### When as\_list is TRUE:

Returns a tuple of 1-D tensors, one for each dimension in input, each containing the indices (in that dimension) of all non-zero elements of input.

If input has n dimensions, then the resulting tuple contains n tensors of size z, where z is the total number of non-zero elements in the input tensor.

As a special case, when input has zero dimensions and a nonzero scalar value, it is treated as a one-dimensional tensor with one element.

### **Examples**

```
if (torch_is_installed()) {
torch_nonzero(torch_tensor(c(1, 1, 1, 0, 1)))
}
```

torch\_norm

Norm

### **Description**

Norm

## Usage

```
torch_norm(self, p = 2L, dim, keepdim = FALSE, dtype)
```

# **Arguments**

 492 torch\_normal

dim (int, 2-tuple of ints, 2-list of ints, optional) If it is an int, vector norm will be

calculated, if it is 2-tuple of ints, matrix norm will be calculated. If the value is NULL, matrix norm will be calculated when the input tensor only has two dimensions, vector norm will be calculated when the input tensor only has one dimension. If the input tensor has more than two dimensions, the vector norm

will be applied to last dimension.

keepdim (bool, optional) whether the output tensors have dim retained or not. Ignored if

dim = NULL and out = NULL. Default: FALSE Ignored if dim = NULL and out =

NULL.

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to 'dtype' while performing the operation. Default:

NULL.

#### **TEST**

Returns the matrix norm or vector norm of a given tensor.

## **Examples**

```
if (torch_is_installed()) {
a <- torch_arange(1, 9, dtype = torch_float())
b <- a$reshape(list(3, 3))
torch_norm(a)
torch_norm(b)
torch_norm(a, Inf)
torch_norm(b, Inf)
}</pre>
```

torch\_normal

Normal

### **Description**

Normal

Normal distributed

### Usage

```
torch_normal(mean, std, size = NULL, generator = NULL, ...)
```

### **Arguments**

mean

(tensor or scalar double) Mean of the normal distribution. If this is a torch\_tensor() then the output has the same dim as mean and it represents the per-element mean. If it's a scalar value, it's reused for all elements.

torch\_normal 493

std	(tensor or scalar double) The standard deviation of the normal distribution. If this is a torch_tensor() then the output has the same size as std and it represents the per-element standard deviation. If it's a scalar value, it's reused for all elements.
size	(integers, optional) only used if both mean and std are scalars.
generator	a random number generator created with torch_generator(). If NULL a default generator is used.
	Tensor option parameters like dtype, layout, and device. Can only be used when mean and std are both scalar numerics.

# normal(mean, std, \*) -> Tensor

Returns a tensor of random numbers drawn from separate normal distributions whose mean and standard deviation are given.

The mean is a tensor with the mean of each output element's normal distribution

The std is a tensor with the standard deviation of each output element's normal distribution

The shapes of mean and std don't need to match, but the total number of elements in each tensor need to be the same.

# normal(mean=0.0, std) -> Tensor

Similar to the function above, but the means are shared among all drawn elements.

## normal(mean, std=1.0) -> Tensor

Similar to the function above, but the standard-deviations are shared among all drawn elements.

# normal(mean, std, size, \*) -> Tensor

Similar to the function above, but the means and standard deviations are shared among all drawn elements. The resulting tensor has size given by size.

# Note

When the shapes do not match, the shape of mean is used as the shape for the returned output tensor

```
if (torch_is_installed()) {

torch_normal(mean=0, std=torch_arange(1, 0, -0.1) + 1e-6)
torch_normal(mean=0.5, std=torch_arange(1., 6.))
torch_normal(mean=torch_arange(1., 6.))
torch_normal(2, 3, size=c(1, 4))
}
```

494 torch\_ones

torch\_not\_equal

Not\_equal

# **Description**

Not\_equal

# Usage

```
torch_not_equal(self, other)
```

# Arguments

self (Tensor) the tensor to compare

other (Tensor or float) the tensor or value to compare

# not\_equal(input, other, \*, out=None) -> Tensor

Alias for torch\_ne().

torch\_ones

Ones

# Description

Ones

# Usage

```
torch_ones(
    ...,
    names = NULL,
    dtype = NULL,
    layout = NULL,
    device = NULL,
    requires_grad = FALSE
)
```

# **Arguments**

... (int...) a sequence of integers defining the shape of the output tensor. Can be a

variable number of arguments or a collection like a list or tuple.

names optional names for the dimensions

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type).

torch\_ones\_like 495

layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

# ones(\*size, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a tensor filled with the scalar value 1, with the shape defined by the variable argument size.

#### **Examples**

```
if (torch_is_installed()) {
torch_ones(c(2, 3))
torch_ones(c(5))
}
```

torch\_ones\_like

Ones\_like

## Description

Ones\_like

#### Usage

```
torch_ones_like(
  input,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

## **Arguments**

input (Tensor) the size of input will determine size of the output tensor.

dtype (torch.dtype, optional) the desired data type of returned Tensor. Default: if

NULL, defaults to the dtype of input.

layout (torch.layout, optional) the desired layout of returned tensor. Default: if

NULL, defaults to the layout of input.

496 torch\_orgqr

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, defaults to the device of input.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

memory\_format (torch.memory\_format, optional) the desired memory format of returned Ten-

sor. Default: torch\_preserve\_format.

# ones\_like(input, dtype=NULL, layout=NULL, device=NULL, requires\_grad=False, memory\_format=torch.preserve\_form -> Tensor

```
Returns a tensor filled with the scalar value 1, with the same size as input. torch_ones_like(input) is equivalent to torch_ones(input.size(), dtype=input.dtype, layout=input.layout, device=input.device).
```

### Warning

As of 0.4, this function does not support an out keyword. As an alternative, the old torch\_ones\_like(input, out=output) is equivalent to torch\_ones(input.size(), out=output).

## **Examples**

```
if (torch_is_installed()) {
input = torch_empty(c(2, 3))
torch_ones_like(input)
}
```

torch\_orgqr

Orgqr

# Description

Orgqr

### Usage

```
torch_orgqr(self, input2)
```

# **Arguments**

```
self (Tensor) the a from torch_geqrf. input2 (Tensor) the tau from torch_geqrf.
```

# orgqr(input, input2) -> Tensor

Computes the orthogonal matrix Q of a QR factorization, from the (input, input2) tuple returned by torch\_geqrf.

This directly calls the underlying LAPACK function ?orgqr. See LAPACK documentation for orgqr\_for further details.

torch\_ormqr 497

torch_ormqr	Ormqr
-------------	-------

# **Description**

Ormqr

# Usage

```
torch_ormqr(self, input2, input3, left = TRUE, transpose = FALSE)
```

# **Arguments**

self (Tensor) the a from torch\_geqrf.
input2 (Tensor) the tau from torch\_geqrf.
input3 (Tensor) the matrix to be multiplied.
left see LAPACK documentation
transpose see LAPACK documentation

### ormqr(input, input2, input3, left=TRUE, transpose=False) -> Tensor

Multiplies mat (given by input3) by the orthogonal Q matrix of the QR factorization formed by torch\_geqrf() that is represented by (a, tau) (given by (input, input2)).

This directly calls the underlying LAPACK function ?ormqr.

### **Description**

Outer

## Usage

```
torch_outer(self, vec2)
```

## **Arguments**

```
self (Tensor) 1-D input vector
vec2 (Tensor) 1-D input vector
```

### outer(input, vec2, \*, out=None) -> Tensor

Outer product of input and vec2. If input is a vector of size n and vec2 is a vector of size m, then out must be a matrix of size  $(n \times m)$ .

498 torch\_pdist

### Note

This function does not broadcast.

# **Examples**

```
if (torch_is_installed()) {
v1 <- torch_arange(1., 5.)
v2 <- torch_arange(1., 4.)
torch_outer(v1, v2)
}</pre>
```

torch\_pdist

Pdist

## Description

**Pdist** 

## Usage

```
torch_pdist(self, p = 2L)
```

# **Arguments**

```
self NA input tensor of shape N \times M.

p NA p value for the p-norm distance to calculate between each vector pair \in [0,\infty].
```

## pdist(input, p=2) -> Tensor

Computes the p-norm distance between every pair of row vectors in the input. This is identical to the upper triangular portion, excluding the diagonal, of torch\_norm(input[:, NULL] - input, dim=2, p=p). This function will be faster if the rows are contiguous.

```
If input has shape N \times M then the output will have shape \frac{1}{2}N(N-1).
```

```
This function is equivalent to scipy.spatial.distance.pdist(input, 'minkowski', p=p) if p \in (0,\infty). When p=0 it is equivalent to scipy.spatial.distance.pdist(input, 'hamming') * M. When p=\infty, the closest scipy function is scipy.spatial.distance.pdist(xn, lambda x, y: np.abs(x - y).max(
```

torch\_pinverse 499

torch_pinverse	Pinverse
----------------	----------

# **Description**

Pinverse

#### Usage

```
torch_pinverse(self, rcond = 1e-15)
```

## **Arguments**

self (Tensor) The input tensor of size (\*, m, n) where \* is zero or more batch dimen-

sions

rcond (float) A floating point value to determine the cutoff for small singular values.

Default: 1e-15

# pinverse(input, rcond=1e-15) -> Tensor

Calculates the pseudo-inverse (also known as the Moore-Penrose inverse) of a 2D tensor. Please look at Moore-Penrose inverse\_ for more details

## Note

This method is implemented using the Singular Value Decomposition.

The pseudo-inverse is not necessarily a continuous function in the elements of the matrix `[1]`\_. Therefore, derivatives are not always existent, and exist for a constant rank only `[2]`\_. However, this method is backprop-able due to the implementation by using SVD results, and could be unstable. Double-backward will also be unstable due to the usage of SVD internally. See `~torch.svd` for more details.

```
if (torch_is_installed()) {
input = torch_randn(c(3, 5))
input
torch_pinverse(input)
# Batched pinverse example
a = torch_randn(c(2,6,3))
b = torch_pinverse(a)
torch_matmul(b, a)
}
```

500 torch\_poisson

# Description

Pixel\_shuffle

# Usage

```
torch_pixel_shuffle(self, upscale_factor)
```

# **Arguments**

```
self (Tensor) the input tensor
upscale_factor (int) factor to increase spatial resolution by
```

# Rearranges elements in a tensor of shape

```
math: (*, C \times r^2, H, W) to a : Rearranges elements in a tensor of shape (*, C \times r^2, H, W) to a tensor of shape (*, C, H \times r, W \times r). See ~torch.nn.PixelShuffle for details.
```

# **Examples**

```
if (torch_is_installed()) {
input = torch_randn(c(1, 9, 4, 4))
output = nnf_pixel_shuffle(input, 3)
print(output$size())
}
```

torch\_poisson

Poisson

# Description

Poisson

# Usage

```
torch_poisson(self, generator = NULL)
```

# Arguments

```
self (Tensor) the input tensor containing the rates of the Poisson distribution generator (torch.Generator, optional) a pseudorandom number generator for sampling
```

torch\_polar 501

### poisson(input \*, generator=NULL) -> Tensor

Returns a tensor of the same size as input with each element sampled from a Poisson distribution with rate parameter given by the corresponding element in input i.e.,

```
out_i \sim Poisson(input_i)
```

## **Examples**

```
if (torch_is_installed()) {

rates = torch_rand(c(4, 4)) * 5  # rate parameter between 0 and 5
torch_poisson(rates)
}
```

torch\_polar

Polar

# **Description**

Polar

# Usage

```
torch_polar(abs, angle)
```

# Arguments

```
abs (Tensor) The absolute value the complex tensor. Must be float or double.

angle (Tensor) The angle of the complex tensor. Must be same dtype as abs.
```

### polar(abs, angle, \*, out=None) -> Tensor

Constructs a complex tensor whose elements are Cartesian coordinates corresponding to the polar coordinates with absolute value abs and angle angle.

```
out = abs \cdot cos(angle) + abs \cdot sin(angle) \cdot j
```

```
if (torch_is_installed()) {
  abs <- torch_tensor(c(1, 2), dtype=torch_float64())
  angle <- torch_tensor(c(pi / 2, 5 * pi / 4), dtype=torch_float64())
  z <- torch_polar(abs, angle)
  z
}</pre>
```

502 torch\_polygamma

torch\_polygamma

Polygamma

# Description

Polygamma

# Usage

```
torch_polygamma(n, input)
```

## **Arguments**

n (int) the order of the polygamma function

input (Tensor) the input tensor.

# polygamma(n, input, out=NULL) -> Tensor

Computes the  $n^{th}$  derivative of the digamma function on input.  $n \geq 0$  is called the order of the polygamma function.

$$\psi^{(n)}(x) = \frac{d^{(n)}}{dx^{(n)}}\psi(x)$$

# Note

This function is not implemented for  $\leq 2$ .

```
if (torch_is_installed()) {
## Not run:
a = torch_tensor(c(1, 0.5))
torch_polygamma(1, a)
## End(Not run)
}
```

torch\_pow 503

torch\_pow

Pow

# **Description**

Pow

# Usage

```
torch_pow(self, exponent)
```

# **Arguments**

self (float) the scalar base value for the power operation

exponent (float or tensor) the exponent value

# pow(input, exponent, out=NULL) -> Tensor

Takes the power of each element in input with exponent and returns a tensor with the result.

exponent can be either a single float number or a Tensor with the same number of elements as input.

When exponent is a scalar value, the operation applied is:

$$\operatorname{out}_i = x_i^{\operatorname{exponent}}$$

When exponent is a tensor, the operation applied is:

$$\operatorname{out}_i = x_i^{\operatorname{exponent}_i}$$

When exponent is a tensor, the shapes of input and exponent must be broadcastable .

# pow(self, exponent, out=NULL) -> Tensor

self is a scalar float value, and exponent is a tensor. The returned tensor out is of the same shape as exponent

The operation applied is:

$$out_i = self^{exponent_i}$$

504 torch\_prod

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_pow(a, 2)
exp <- torch_arange(1, 5)
a <- torch_arange(1, 5)
a
exp
torch_pow(a, exp)

exp <- torch_arange(1, 5)
base <- 2
torch_pow(base, exp)
}</pre>
```

torch\_prod

Prod

## Description

Prod

#### **Usage**

```
torch_prod(self, dim, keepdim = FALSE, dtype = NULL)
```

## **Arguments**

self (Tensor) the input tensor.
dim (int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

# prod(input, dtype=NULL) -> Tensor

Returns the product of all elements in the input tensor.

# prod(input, dim, keepdim=False, dtype=NULL) -> Tensor

Returns the product of each row of the input tensor in the given dimension dim.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 fewer dimension than input.

torch\_promote\_types 505

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_prod(a)

a = torch_randn(c(4, 2))
a
torch_prod(a, 1)
}
```

torch\_promote\_types

Promote\_types

## **Description**

Promote\_types

### Usage

```
torch_promote_types(type1, type2)
```

# Arguments

```
type1 (torch.dtype)
type2 (torch.dtype)
```

## promote\_types(type1, type2) -> dtype

Returns the torch\_dtype with the smallest size and scalar kind that is not smaller nor of lower kind than either type1 or type2. See type promotion documentation for more information on the type promotion logic.

```
if (torch_is_installed()) {
  torch_promote_types(torch_int32(), torch_float32())
  torch_promote_types(torch_uint8(), torch_long())
}
```

506 torch\_qr

torch\_qr

Qr

## **Description**

Qr

#### Usage

```
torch_qr(self, some = TRUE)
```

#### **Arguments**

self (Tensor) the input tensor of size (\*, m, n) where \* is zero or more batch dimen-

sions consisting of matrices of dimension  $m \times n$ .

some (bool, optional) Set to TRUE for reduced QR decomposition and FALSE for com-

plete QR decomposition.

#### qr(input, some=TRUE, out=NULL) -> (Tensor, Tensor)

Computes the QR decomposition of a matrix or a batch of matrices input, and returns a namedtuple (Q, R) of tensors such that input = QR with Q being an orthogonal matrix or batch of orthogonal matrices and R being an upper triangular matrix or batch of upper triangular matrices.

If some is TRUE, then this function returns the thin (reduced) QR factorization. Otherwise, if some is FALSE, this function returns the complete QR factorization.

#### Note

precision may be lost if the magnitudes of the elements of input are large

While it should always give you a valid decomposition, it may not give you the same one across platforms - it will depend on your LAPACK implementation.

```
if (torch_is_installed()) {
a = torch_tensor(matrix(c(12., -51, 4, 6, 167, -68, -4, 24, -41), ncol = 3, byrow = TRUE))
out = torch_qr(a)
q = out[[1]]
r = out[[2]]
torch_mm(q, r)$round()
torch_mm(q$t(), q)$round()
}
```

torch\_qscheme 507

torch	gscheme
COI CII_	_q3CHCIIIC

Creates the corresponding Scheme object

# Description

Creates the corresponding Scheme object

# Usage

```
torch_per_channel_affine()
torch_per_tensor_affine()
torch_per_channel_symmetric()
torch_per_tensor_symmetric()
```

torch\_quantile

Quantile

# Description

Quantile

## Usage

```
torch_quantile(self, q, dim = NULL, keepdim = FALSE, interpolation = "linear")
```

## **Arguments**

self (Tensor) the input tensor.

q (float or Tensor) a scalar or 1D tensor of quantile values in the range [0, 1]

dim (int) the dimension to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

interpolation The interpolation method.

## quantile(input, q) -> Tensor

Returns the q-th quantiles of all elements in the input tensor, doing a linear interpolation when the q-th quantile lies between two data points.

### quantile(input, q, dim=None, keepdim=FALSE, \*, out=None) -> Tensor

Returns the q-th quantiles of each row of the input tensor along the dimension dim, doing a linear interpolation when the q-th quantile lies between two data points. By default, dim is None resulting in the input tensor being flattened before computation.

If keepdim is TRUE, the output dimensions are of the same size as input except in the dimensions being reduced (dim or all if dim is NULL) where they have size 1. Otherwise, the dimensions being reduced are squeezed (see torch\_squeeze). If q is a 1D tensor, an extra dimension is prepended to the output tensor with the same size as q which represents the quantiles.

#### **Examples**

```
if (torch_is_installed()) {
    a <- torch_randn(c(1, 3))
    a
    q <- torch_tensor(c(0, 0.5, 1))
    torch_quantile(a, q)

a <- torch_randn(c(2, 3))
    a
    q <- torch_tensor(c(0.25, 0.5, 0.75))
    torch_quantile(a, q, dim=1, keepdim=TRUE)
    torch_quantile(a, q, dim=1, keepdim=TRUE)$shape
}</pre>
```

torch\_quantize\_per\_channel

Quantize\_per\_channel

## **Description**

Quantize\_per\_channel

#### Usage

```
torch_quantize_per_channel(self, scales, zero_points, axis, dtype)
```

## Arguments

self	(Tensor) float tensor to quantize	
scales	(Tensor) float 1D tensor of scales to use, size should match input.size(axis)	
zero_points	(int) integer 1D tensor of offset to use, size should match input.size(axis)	
axis	(int) dimension on which apply per-channel quantization	
dtype	(torch.dtype) the desired data type of returned tensor. Has to be one of the quantized dtypes: torch_quint8, torch.qint8, torch.qint32	

### quantize\_per\_channel(input, scales, zero\_points, axis, dtype) -> Tensor

Converts a float tensor to per-channel quantized tensor with given scales and zero points.

#### **Examples**

```
torch_quantize_per_tensor
```

Quantize\_per\_tensor

## **Description**

```
Quantize_per_tensor
```

#### Usage

```
torch_quantize_per_tensor(self, scale, zero_point, dtype)
```

#### **Arguments**

self (Tensor) float tensor to quantize scale (float) scale to apply in quantiza

scale (float) scale to apply in quantization formula zero\_point (int) offset in integer value that maps to float zero

dtype (torch.dtype) the desired data type of returned tensor. Has to be one of the

quantized dtypes: torch\_quint8, torch.qint8, torch.qint32

#### quantize\_per\_tensor(input, scale, zero\_point, dtype) -> Tensor

Converts a float tensor to quantized tensor with given scale and zero point.

```
if (torch_is_installed()) {
  torch_quantize_per_tensor(torch_tensor(c(-1.0, 0.0, 1.0, 2.0)), 0.1, 10, torch_quint8())
  torch_quantize_per_tensor(torch_tensor(c(-1.0, 0.0, 1.0, 2.0)), 0.1, 10, torch_quint8())$int_repr()
}
```

510 torch\_rand

torch\_rad2deg

Rad2deg

## **Description**

Rad2deg

# Usage

```
torch_rad2deg(self)
```

# Arguments

self

(Tensor) the input tensor.

# rad2deg(input, \*, out=None) -> Tensor

Returns a new tensor with each of the elements of input converted from angles in radians to degrees.

## **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(rbind(c(3.142, -3.142), c(6.283, -6.283), c(1.570, -1.570)))
torch_rad2deg(a)
}</pre>
```

torch\_rand

Rand

# Description

Rand

# Usage

torch\_randint 511

## **Arguments**

•••	(int) a sequence of integers defining the shape of the output tensor. Can be a variable number of arguments or a collection like a list or tuple.
names	optional dimension names
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch_set_default_tensor_type).
layout	(torch.layout, optional) the desired layout of returned Tensor. Default: torch_strided.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

# rand(\*size, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a tensor filled with random numbers from a uniform distribution on the interval [0,1). The shape of the tensor is defined by the variable argument size.

# **Examples**

```
if (torch_is_installed()) {
torch_rand(4)
torch_rand(c(2, 3))
}
```

torch\_randint

Randint

# Description

Randint

# Usage

```
torch_randint(
  low,
  high,
  size,
  generator = NULL,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

512 torch\_randint\_like

### **Arguments**

low (int, optional) Lowest integer to be drawn from the distribution. Default: 0. high (int) One above the highest integer to be drawn from the distribution. (tuple) a tuple defining the shape of the output tensor. size generator (torch. Generator, optional) a pseudorandom number generator for sampling dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch\_set\_default\_tensor\_type). layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided. device (torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types. (bool, optional) If autograd should record operations on the returned tensor. Derequires\_grad fault: FALSE.

## randint(low=0, high, size, \*, generator=NULL, out=NULL, \

dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

memory format for the resulting tensor.

Returns a tensor filled with random integers generated uniformly between low (inclusive) and high (exclusive).

The shape of the tensor is defined by the variable argument size.

.. note: With the global dtype default (torch\_float32), this function returns a tensor with dtype torch\_int64.

## **Examples**

memory\_format

```
if (torch_is_installed()) {
torch_randint(3, 5, list(3))
torch_randint(0, 10, size = list(2, 2))
torch_randint(3, 10, list(2, 2))
}
```

#### Description

Randint\_like

torch\_randn 513

## Usage

```
torch_randint_like(
  input,
  low,
  high,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE
)
```

# Arguments

input	(Tensor) the size of input will determine size of the output tensor.	
low	(int, optional) Lowest integer to be drawn from the distribution. Default: 0.	
high	(int) One above the highest integer to be drawn from the distribution.	
dtype	(torch.dtype, optional) the desired data type of returned Tensor. Default: if NULL, defaults to the dtype of input.	
layout	(torch.layout, optional) the desired layout of returned tensor. Default: if NULL, defaults to the layout of input.	
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, defaults to the device of input.	
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.	

# randint\_like(input, low=0, high, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False,

memory\_format=torch.preserve\_format) -> Tensor

Returns a tensor with the same shape as Tensor input filled with random integers generated uniformly between low (inclusive) and high (exclusive).

 $\dots$  note: With the global dtype default (torch\_float32), this function returns a tensor with dtype torch\_int64.

# **Description**

Randn

514 torch\_randn

## Usage

```
torch_randn(
    ...,
    names = NULL,
    dtype = NULL,
    layout = NULL,
    device = NULL,
    requires_grad = FALSE
)
```

# Arguments

	(int) a sequence of integers defining the shape of the output tensor. Can be a variable number of arguments or a collection like a list or tuple.
names	optional names for the dimensions
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, uses a global default (see torch_set_default_tensor_type).
layout	(torch.layout, optional) the desired layout of returned Tensor. Default: torch_strided.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

# randn(\*size, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a tensor filled with random numbers from a normal distribution with mean 0 and variance 1 (also called the standard normal distribution).

$$\operatorname{out}_i \sim \mathcal{N}(0,1)$$

The shape of the tensor is defined by the variable argument size.

```
if (torch_is_installed()) {
  torch_randn(c(4))
  torch_randn(c(2, 3))
}
```

torch\_randn\_like 515

torch\_randn\_like Randn\_like

## **Description**

Randn\_like

## Usage

```
torch_randn_like(
  input,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

## **Arguments**

input	(Tensor) the size of input will determine size of the output tensor.	
dtype	(torch.dtype, optional) the desired data type of returned Tensor. Default: if NULL, defaults to the dtype of input.	
layout	(torch.layout, optional) the desired layout of returned tensor. Default: if NULL, defaults to the layout of input.	
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, defaults to the device of input.	
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.	
memory_format	(torch.memory_format, optional) the desired memory format of returned Tensor. Default: torch_preserve_format.	

# randn\_like(input, dtype=NULL, layout=NULL, device=NULL, requires\_grad=False, memory\_format=torch.preserve\_for -> Tensor

Returns a tensor with the same size as input that is filled with random numbers from a normal distribution with mean 0 and variance 1. torch\_randn\_like(input) is equivalent to torch\_randn(input.size(), dtype=input.dtype, layout=input.layout, device=input.device).

516 torch\_randperm

torch\_randperm Randperm

## Description

Randperm

## Usage

```
torch_randperm(
   n,
   dtype = torch_int64(),
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

## Arguments

```
n (int) the upper bound (exclusive)

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: torch_int64.

layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch_strided.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires_grad (bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.
```

# randperm(n, out=NULL, dtype=torch.int64, layout=torch.strided, device=NULL, requires\_grad=False) -> LongTensor

Returns a random permutation of integers from 0 to n-1.

```
if (torch_is_installed()) {
torch_randperm(4)
}
```

torch\_rand\_like 517

torch\_rand\_like Rand\_like

# Description

Rand\_like

## Usage

```
torch_rand_like(
  input,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

## **Arguments**

input	(Tensor) the size of input will determine size of the output tensor.	
dtype	(torch.dtype, optional) the desired data type of returned Tensor. Default: if NULL, defaults to the dtype of input.	
layout	(torch.layout, optional) the desired layout of returned tensor. Default: if NULL, defaults to the layout of input.	
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, defaults to the device of input.	
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.	
memory_format	(torch.memory_format, optional) the desired memory format of returned Tensor. Default: torch_preserve_format.	

# rand\_like(input, dtype=NULL, layout=NULL, device=NULL, requires\_grad=False, memory\_format=torch.preserve\_form -> Tensor

Returns a tensor with the same size as input that is filled with random numbers from a uniform distribution on the interval [0,1). torch\_rand\_like(input) is equivalent to torch\_rand(input.size(), dtype=input.dtype, layout=input.layout, device=input.device).

518 torch\_range

torch\_range Range

## **Description**

Range

# Usage

```
torch_range(
   start,
   end,
   step = 1,
   dtype = NULL,
   layout = NULL,
   device = NULL,
   requires_grad = FALSE
)
```

## **Arguments**

start

device

end (float) the ending value for the set of points

step (float) the gap between each pair of adjacent points. Default: 1.

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type). If dtype

is not given, infer the data type from the other input arguments. If any of

start, end, or stop are floating-point, the dtype is inferred to be the default

dtype, see ~torch.get\_default\_dtype. Otherwise, the dtype is inferred to be

torch.int64.

layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided.

(float) the starting value for the set of points. Default: 0.

(torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

# range(start=0, end, step=1, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a 1-D tensor of size  $\left\lfloor \frac{\text{end-start}}{\text{step}} \right\rfloor + 1$  with values from start to end with step step. Step is the gap between two values in the tensor.

```
out_{i+1} = out_i + step.
```

torch\_real 519

## Warning

This function is deprecated in favor of torch\_arange.

# **Examples**

```
if (torch_is_installed()) {
torch_range(1, 4)
torch_range(1, 4, 0.5)
}
```

torch\_real

Real

# Description

Real

## Usage

```
torch_real(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# real(input) -> Tensor

Returns the real part of the input tensor. If input is a real (non-complex) tensor, this function just returns it.

# Warning

Not yet implemented for complex tensors.

```
out_i = real(input_i)
```

```
if (torch_is_installed()) {
## Not run:
torch_real(torch_tensor(c(-1 + 1i, -2 + 2i, 3 - 3i)))
## End(Not run)
}
```

520 torch\_reduction

torch\_reciprocal

Reciprocal

# Description

Reciprocal

# Usage

```
torch_reciprocal(self)
```

# Arguments

self

(Tensor) the input tensor.

# reciprocal(input, out=NULL) -> Tensor

Returns a new tensor with the reciprocal of the elements of input

$$\mathsf{out}_i = \frac{1}{\mathsf{input}_i}$$

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_reciprocal(a)
}
```

torch\_reduction

Creates the reduction objet

# Description

Creates the reduction objet

## Usage

```
torch_reduction_sum()
torch_reduction_mean()
torch_reduction_none()
```

torch\_relu 521

torch\_relu

Relu

# Description

Relu

# Usage

```
torch_relu(self)
```

# Arguments

self

the input tensor

# relu(input) -> Tensor

Computes the relu tranformation.

torch\_relu\_

Relu\_

# Description

Relu\_

# Usage

```
torch_relu_(self)
```

# Arguments

self

the input tensor

# relu\_(input) -> Tensor

In-place version of torch\_relu().

522 torch\_renorm

torch\_remainder

Remainder

# Description

Remainder

# Usage

```
torch_remainder(self, other)
```

# Arguments

self (Tensor) the dividend

other (Tensor or float) the divisor that may be either a number or a Tensor of the same

shape as the dividend

# remainder(input, other, out=NULL) -> Tensor

Computes the element-wise remainder of division.

The divisor and dividend may contain both for integer and floating point numbers. The remainder has the same sign as the divisor.

When other is a tensor, the shapes of input and other must be broadcastable.

## **Examples**

```
if (torch_is_installed()) {

torch_remainder(torch_tensor(c(-3., -2, -1, 1, 2, 3)), 2)
torch_remainder(torch_tensor(c(1., 2, 3, 4, 5)), 1.5)
}
```

torch\_renorm

Renorm

# Description

Renorm

## Usage

```
torch_renorm(self, p, dim, maxnorm)
```

## Arguments

self	(Tensor) the input tensor.
------	----------------------------

p (float) the power for the norm computation

dim (int) the dimension to slice over to get the sub-tensors maxnorm (float) the maximum norm to keep each sub-tensor under

#### renorm(input, p, dim, maxnorm, out=NULL) -> Tensor

Returns a tensor where each sub-tensor of input along dimension dim is normalized such that the p-norm of the sub-tensor is lower than the value maxnorm

#### Note

If the norm of a row is lower than maxnorm, the row is unchanged

## **Examples**

```
if (torch_is_installed()) {
x = torch_ones(c(3, 3))
x[2,]$fill_(2)
x[3,]$fill_(3)
x
torch_renorm(x, 1, 1, 5)
}
```

torch\_repeat\_interleave

Repeat\_interleave

## **Description**

Repeat\_interleave

#### Usage

```
torch_repeat_interleave(self, repeats, dim = NULL, output_size = NULL)
```

### **Arguments**

self	(Tensor) the input tensor.
------	----------------------------

repeats (Tensor or int) The number of repetitions for each element. repeats is broad-

casted to fit the shape of the given axis.

dim (int, optional) The dimension along which to repeat values. By default, use the

flattened input array, and return a flat output array.

output\_size (int, optional) - Total output size for the given axis (e.g. sum of repeats). If

given, it will avoid stream syncronization needed to calculate output shape of

the tensor.

524 torch\_reshape

### repeat\_interleave(input, repeats, dim=NULL) -> Tensor

Repeat elements of a tensor.

# Warning

```
This is different from `torch_Tensor.repeat` but similar to `numpy.repeat`.
```

## repeat\_interleave(repeats) -> Tensor

```
If the repeats is tensor([n1, n2, n3, ...]), then the output will be tensor([0, 0, ..., 1, 1, ..., 2, 2, ..., ...]) where 0 appears n1 times, 1 appears n2 times, 2 appears n3 times, etc.
```

# **Examples**

```
if (torch_is_installed()) {
## Not run:
x = torch_tensor(c(1, 2, 3))
x$repeat_interleave(2)
y = torch_tensor(matrix(c(1, 2, 3, 4), ncol = 2, byrow=TRUE))
torch_repeat_interleave(y, 2)
torch_repeat_interleave(y, 3, dim=1)
torch_repeat_interleave(y, torch_tensor(c(1, 2)), dim=1)
## End(Not run)
}
```

torch\_reshape

Reshape

## **Description**

Reshape

## Usage

```
torch_reshape(self, shape)
```

# Arguments

```
self (Tensor) the tensor to be reshaped shape (tuple of ints) the new shape
```

torch\_result\_type 525

### reshape(input, shape) -> Tensor

Returns a tensor with the same data and number of elements as input, but with the specified shape. When possible, the returned tensor will be a view of input. Otherwise, it will be a copy. Contiguous inputs and inputs with compatible strides can be reshaped without copying, but you should not depend on the copying vs. viewing behavior.

See torch\_Tensor.view on when it is possible to return a view.

A single dimension may be -1, in which case it's inferred from the remaining dimensions and the number of elements in input.

## **Examples**

```
if (torch_is_installed()) {
a <- torch_arange(0, 3)
torch_reshape(a, list(2, 2))
b <- torch_tensor(matrix(c(0, 1, 2, 3), ncol = 2, byrow=TRUE))
torch_reshape(b, list(-1))
}</pre>
```

torch\_result\_type

Result\_type

#### **Description**

Result\_type

## Usage

```
torch_result_type(tensor1, tensor2)
```

#### **Arguments**

```
tensor1 (Tensor or Number) an input tensor or number
tensor2 (Tensor or Number) an input tensor or number
```

#### result\_type(tensor1, tensor2) -> dtype

Returns the torch\_dtype that would result from performing an arithmetic operation on the provided input tensors. See type promotion documentation for more information on the type promotion logic.

```
if (torch_is_installed()) {
torch_result_type(tensor1 = torch_tensor(c(1, 2), dtype=torch_int()), tensor2 = 1)
}
```

526 torch\_rot90

torch\_roll

Roll

#### **Description**

Roll

#### **Usage**

```
torch_roll(self, shifts, dims = list())
```

## **Arguments**

self (Tensor) the input tensor.

shifts (int or tuple of ints) The number of places by which the elements of the tensor

are shifted. If shifts is a tuple, dims must be a tuple of the same size, and each

dimension will be rolled by the corresponding value

dims (int or tuple of ints) Axis along which to roll

# roll(input, shifts, dims=NULL) -> Tensor

Roll the tensor along the given dimension(s). Elements that are shifted beyond the last position are re-introduced at the first position. If a dimension is not specified, the tensor will be flattened before rolling and then restored to the original shape.

## **Examples**

```
if (torch_is_installed()) {
x = torch_tensor(c(1, 2, 3, 4, 5, 6, 7, 8))$view(c(4, 2))
x
torch_roll(x, 1, 1)
torch_roll(x, -1, 1)
torch_roll(x, shifts=list(2, 1), dims=list(1, 2))
}
```

torch\_rot90

Rot90

## **Description**

Rot90

#### Usage

```
torch_rot90(self, k = 1L, dims = c(0, 1))
```

torch\_round 527

## Arguments

```
self (Tensor) the input tensor.

k (int) number of times to rotate

dims (a list or tuple) axis to rotate
```

# rot90(input, k, dims) -> Tensor

Rotate a n-D tensor by 90 degrees in the plane specified by dims axis. Rotation direction is from the first towards the second axis if k > 0, and from the second towards the first for k < 0.

## **Examples**

```
if (torch_is_installed()) {

x <- torch_arange(1, 4)$view(c(2, 2))
x
torch_rot90(x, 1, c(1, 2))
x <- torch_arange(1, 8)$view(c(2, 2, 2))
x
torch_rot90(x, 1, c(1, 2))
}</pre>
```

torch\_round

Round

## Description

Round

# Usage

```
torch_round(self, decimals)
```

## **Arguments**

self (Tensor) the input tensor.

decimals Number of decimal places to round to (default: 0). If decimals is negative, it

specifies the number of positions to the left of the decimal point.

## round(input, out=NULL) -> Tensor

Returns a new tensor with each of the elements of input rounded to the closest integer.

528 torch\_rrelu\_

## **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_round(a)
}
```

torch\_rrelu\_

Rrelu\_

# Description

Rrelu\_

# Usage

# Arguments

self the input tensor

lower lower bound of the uniform distribution. Default: 1/8 upper upper bound of the uniform distribution. Default: 1/3 training bool wether it's a training pass. DEfault: FALSE

generator random number generator

# rrelu\_(input, lower=1./8, upper=1./3, training=False) -> Tensor

In-place version of torch\_rrelu.

torch\_rsqrt 529

torch\_rsqrt

Rsqrt

# Description

Rsqrt

# Usage

```
torch_rsqrt(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# rsqrt(input, out=NULL) -> Tensor

Returns a new tensor with the reciprocal of the square-root of each of the elements of input.

$$\mathsf{out}_i = \frac{1}{\sqrt{\mathsf{input}_i}}$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_rsqrt(a)
}
```

torch\_save

Saves an object to a disk file.

# Description

This function is experimental, don't use for long term storage.

# Usage

```
torch_save(obj, path, ..., compress = TRUE)
```

530 torch\_scalar\_tensor

#### **Arguments**

obj the saved object

path a connection or the name of the file to save.

... not currently used.

compress a logical specifying whether saving to a named file is to use "gzip" compression,

or one of "gzip", "bzip2" or "xz" to indicate the type of compression to be used.

Ignored if file is a connection.

#### See Also

Other torch\_save: torch\_load(), torch\_serialize()

torch\_scalar\_tensor Scalar tensor

# Description

Creates a singleton dimension tensor.

## Usage

torch\_scalar\_tensor(value, dtype = NULL, device = NULL, requires\_grad = FALSE)

#### **Arguments**

value the value you want to use

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type).

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

torch\_searchsorted 531

torch\_searchsorted

Searchsorted

#### **Description**

Searchsorted

#### Usage

```
torch_searchsorted(
  sorted_sequence,
  self,
  out_int32 = FALSE,
  right = FALSE,
  side = NULL,
  sorter = list()
)
```

#### **Arguments**

side

sorted\_sequence

(Tensor) N-D or 1-D tensor, containing monotonically increasing sequence on

the innermost dimension.

self (Tensor or Scalar) N-D tensor or a Scalar containing the search value(s).

out\_int32 (bool, optional) – indicate the output data type. torch\_int32() if True, torch\_int64()

otherwise. Default value is FALSE, i.e. default output data type is torch\_int64().

right (bool, optional) – if False, return the first suitable location that is found. If

True, return the last such index. If no suitable index found, return 0 for non-numerical value (eg. nan, inf) or the size of boundaries (one pass the last index). In other words, if False, gets the lower bound index for each value in input from houndaries. If True gets the upper hound index instead. Default value in False.

boundaries. If True, gets the upper bound index instead. Default value is False.

the same as right but preferred. "left" corresponds to FALSE for right and "right" corresponds to TRUE for right. It will error if this is set to "left" while right is

TRUE.

sorter if provided, a tensor matching the shape of the unsorted sorted\_sequence con-

taining a sequence of indices that sort it in the ascending order on the innermost

dimension.

#### searchsorted(sorted\_sequence, values, \*, out\_int32=FALSE, right=FALSE, out=None) -> Tensor

Find the indices from the *innermost* dimension of sorted\_sequence such that, if the corresponding values in values were inserted before the indices, the order of the corresponding *innermost* dimension within sorted\_sequence would be preserved. Return a new tensor with the same size as values. If right is FALSE (default), then the left boundary of sorted\_sequence is closed.

532 torch\_selu\_

## **Examples**

```
if (torch_is_installed()) {
    sorted_sequence <- torch_tensor(rbind(c(1, 3, 5, 7, 9), c(2, 4, 6, 8, 10)))
    sorted_sequence
    values <- torch_tensor(rbind(c(3, 6, 9), c(3, 6, 9)))
    values
    torch_searchsorted(sorted_sequence, values)
    torch_searchsorted(sorted_sequence, values, right=TRUE)
    sorted_sequence_1d <- torch_tensor(c(1, 3, 5, 7, 9))
    sorted_sequence_1d
    torch_searchsorted(sorted_sequence_1d, values)
}</pre>
```

torch\_selu

Selu

# Description

Selu

# Usage

torch\_selu(self)

#### **Arguments**

self

the input tensor

## selu(input) -> Tensor

Computes the selu transformation.

torch\_selu\_

Selu\_

## **Description**

Selu\_

# Usage

```
torch_selu_(self)
```

## **Arguments**

self

the input tensor

torch\_serialize 533

#### selu\_(input) -> Tensor

In-place version of torch\_selu().

torch\_serialize

Serialize a torch object returning a raw object

# Description

It's just a wraper around torch\_save().

## Usage

```
torch_serialize(obj, ...)
```

## **Arguments**

obj the saved object

... Additional arguments passed to torch\_save(). obj and path are not accepted

as they are set by torch\_serialize().

## Value

A raw vector containing the serialized object. Can be reloaded using torch\_load().

## See Also

```
Other torch_save: torch_load(), torch_save()
```

```
torch_set_default_dtype
```

Gets and sets the default floating point dtype.

# Description

Gets and sets the default floating point dtype.

## Usage

```
torch_set_default_dtype(d)
torch_get_default_dtype()
```

## Arguments

d The default floating point dtype to set. Initially set to torch\_float().

534 torch\_sigmoid

torch\_sgn

Sgn

## **Description**

Sgn

## Usage

```
torch_sgn(self)
```

## **Arguments**

self

(Tensor) the input tensor.

## sgn(input, \*, out=None) -> Tensor

For complex tensors, this function returns a new tensor whose elemants have the same angle as that of the elements of input and absolute value 1. For a non-complex tensor, this function returns the signs of the elements of input (see torch\_sign).

```
\operatorname{out}_i = 0, if |\operatorname{input}_i| == 0 \operatorname{out}_i = \frac{\operatorname{input}_i}{|\operatorname{input}_i|}, otherwise
```

## **Examples**

```
if (torch_is_installed()) {
  if (FALSE) {
  x <- torch_tensor(c(3+4i, 7-24i, 0, 1+2i))
  x$sgn()
  torch_sgn(x)
  }
}</pre>
```

torch\_sigmoid

Sigmoid

## **Description**

Sigmoid

## Usage

```
torch_sigmoid(self)
```

# Arguments

self

(Tensor) the input tensor.

torch\_sign 535

# sigmoid(input, out=NULL) -> Tensor

Returns a new tensor with the sigmoid of the elements of input.

$$\mathsf{out}_i = \frac{1}{1 + e^{-\mathsf{input}_i}}$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_sigmoid(a)
}
```

torch\_sign

Sign

# Description

Sign

# Usage

```
torch_sign(self)
```

## **Arguments**

self

(Tensor) the input tensor.

# sign(input, out=NULL) -> Tensor

Returns a new tensor with the signs of the elements of input.

$$out_i = sgn(input_i)$$

```
if (torch_is_installed()) {
a = torch_tensor(c(0.7, -1.2, 0., 2.3))
a
torch_sign(a)
}
```

536 torch\_sin

torch\_signbit

Signbit

# Description

Signbit

## Usage

```
torch_signbit(self)
```

# Arguments

self

(Tensor) the input tensor.

# signbit(input, \*, out=None) -> Tensor

Tests if each element of input has its sign bit set (is less than zero) or not.

## **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(c(0.7, -1.2, 0., 2.3))
torch_signbit(a)
}</pre>
```

torch\_sin

Sin

# Description

Sin

# Usage

```
torch_sin(self)
```

#### **Arguments**

self

(Tensor) the input tensor.

# sin(input, out=NULL) -> Tensor

Returns a new tensor with the sine of the elements of input.

$$out_i = sin(input_i)$$

torch\_sinh 537

# Examples

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_sin(a)
}
```

torch\_sinh

Sinh

# Description

Sinh

# Usage

```
torch_sinh(self)
```

# Arguments

self

(Tensor) the input tensor.

# sinh(input, out=NULL) -> Tensor

Returns a new tensor with the hyperbolic sine of the elements of input.

```
\operatorname{out}_i = \sinh(\operatorname{input}_i)
```

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_sinh(a)
}
```

538 torch\_slogdet

torch\_slogdet

Slogdet

## **Description**

Slogdet

#### Usage

```
torch_slogdet(self)
```

#### **Arguments**

self

(Tensor) the input tensor of size (\*, n, n) where \* is zero or more batch dimensions.

## slogdet(input) -> (Tensor, Tensor)

Calculates the sign and log absolute value of the determinant(s) of a square matrix or batches of square matrices.

## Note

```
If 'input' has zero determinant, this returns '(0, -inf)'.
```

Backward through `slogdet` internally uses SVD results when `input` is not invertible. In this case, double backward through `slogdet` will be unstable in when `input` doesn't have distinct singular values. See `~torch.svd` for details.

```
if (torch_is_installed()) {
A = torch_randn(c(3, 3))
A
torch_det(A)
torch_logdet(A)
torch_slogdet(A)
}
```

torch\_sort 539

## Description

Sort

## **Arguments**

self (Tensor) the input tensor.

dim (int, optional) the dimension to sort along

descending (bool, optional) controls the sorting order (ascending or descending)

stable (bool, optional) – makes the sorting routine stable, which guarantees that the

order of equivalent elements is preserved.

## sort(input, dim=-1, descending=FALSE) -> (Tensor, LongTensor)

Sorts the elements of the input tensor along a given dimension in ascending order by value.

If dim is not given, the last dimension of the input is chosen.

If descending is TRUE then the elements are sorted in descending order by value.

A namedtuple of (values, indices) is returned, where the values are the sorted values and indices are the indices of the elements in the original input tensor.

#### **Examples**

```
if (torch_is_installed()) {
x = torch_randn(c(3, 4))
out = torch_sort(x)
out
out = torch_sort(x, 1)
out
}
```

```
torch_sparse_coo_tensor
```

Sparse\_coo\_tensor

## Description

Sparse\_coo\_tensor

### Usage

```
torch_sparse_coo_tensor(
  indices,
  values,
  size = NULL,
  dtype = NULL,
  device = NULL,
  requires_grad = FALSE
)
```

#### **Arguments**

indices	(array_like) Initial data for the tensor. Can be a list, tuple, NumPy ndarray, scalar, and other types. Will be cast to a torch_LongTensor internally. The indices are the coordinates of the non-zero values in the matrix, and thus should be two-dimensional where the first dimension is the number of tensor dimensions and the second dimension is the number of non-zero values.
values	(array_like) Initial values for the tensor. Can be a list, tuple, NumPy ndarray, scalar, and other types.
size	(list, tuple, or torch. Size, optional) Size of the sparse tensor. If not provided the size will be inferred as the minimum size big enough to hold all non-zero elements.
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, infers data type from values.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.

# sparse\_coo\_tensor(indices, values, size=NULL, dtype=NULL, device=NULL, requires\_grad=False) -> Tensor

Constructs a sparse tensors in COO(rdinate) format with non-zero elements at the given indices with the given values. A sparse tensor can be uncoalesced, in that case, there are duplicate coordinates in the indices, and the value at that index is the sum of all duplicate value entries: torch\_sparse\_.

```
if (torch_is_installed()) {
i = torch_tensor(matrix(c(1, 2, 2, 3, 1, 3), ncol = 3, byrow = TRUE), dtype=torch_int64())
v = torch_tensor(c(3, 4, 5), dtype=torch_float32())
torch_sparse_coo_tensor(i, v)
torch_sparse_coo_tensor(i, v, c(2, 4))
```

torch\_split 541

```
# create empty sparse tensors
S = torch_sparse_coo_tensor(
  torch_empty(c(1, 0), dtype = torch_int64()),
  torch_tensor(numeric(), dtype = torch_float32()),
  c(1)
)
S = torch_sparse_coo_tensor(
  torch_empty(c(1, 0), dtype = torch_int64()),
  torch_empty(c(0, 2)),
  c(1, 2)
)
}
```

torch\_split

Split

# **Description**

Splits the tensor into chunks. Each chunk is a view of the original tensor.

# Usage

```
torch_split(self, split_size, dim = 1L)
```

# **Arguments**

self (Tensor) tensor to split.

split\_size (int) size of a single chunk or list of sizes for each chunk

dim (int) dimension along which to split the tensor.

### **Details**

If split\_size is an integer type, then tensor will be split into equally sized chunks (if possible). Last chunk will be smaller if the tensor size along the given dimension dim is not divisible by split\_size.

If split\_size is a list, then tensor will be split into length(split\_size) chunks with sizes in dim according to split\_size\_or\_sections.

542 torch\_square

torch\_sqrt

Sqrt

# Description

Sqrt

# Usage

```
torch_sqrt(self)
```

# Arguments

self

(Tensor) the input tensor.

# sqrt(input, out=NULL) -> Tensor

Returns a new tensor with the square-root of the elements of input.

$$\operatorname{out}_i = \sqrt{\operatorname{input}_i}$$

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_sqrt(a)
}
```

torch\_square

Square

# Description

Square

# Usage

```
torch_square(self)
```

# Arguments

self

(Tensor) the input tensor.

torch\_squeeze 543

### square(input, out=NULL) -> Tensor

Returns a new tensor with the square of the elements of input.

### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_square(a)
}
```

torch\_squeeze

Squeeze

# **Description**

Squeeze

### Usage

```
torch_squeeze(self, dim)
```

# **Arguments**

```
self (Tensor) the input tensor.
```

dim (int, optional) if given, the input will be squeezed only in this dimension

# squeeze(input, dim=NULL, out=NULL) -> Tensor

Returns a tensor with all the dimensions of input of size 1 removed.

For example, if input is of shape:  $(A \times 1 \times B \times C \times 1 \times D)$  then the out tensor will be of shape:  $(A \times B \times C \times D)$ .

When dim is given, a squeeze operation is done only in the given dimension. If input is of shape:  $(A \times 1 \times B)$ , squeeze(input, 0) leaves the tensor unchanged, but squeeze(input, 1) will squeeze the tensor to the shape  $(A \times B)$ .

### Note

The returned tensor shares the storage with the input tensor, so changing the contents of one will change the contents of the other.

544 torch\_std

### **Examples**

```
if (torch_is_installed()) {
x = torch_zeros(c(2, 1, 2, 1, 2))
x
y = torch_squeeze(x)
y
y = torch_squeeze(x, 1)
y
y = torch_squeeze(x, 2)
y
}
```

torch\_stack

Stack

# Description

Stack

# Usage

```
torch_stack(tensors, dim = 1L)
```

# **Arguments**

tensors (sequence of Tensors) sequence of tensors to concatenate

dim (int) dimension to insert. Has to be between 0 and the number of dimensions of

concatenated tensors (inclusive)

# stack(tensors, dim=0, out=NULL) -> Tensor

Concatenates sequence of tensors along a new dimension.

All tensors need to be of the same size.

torch\_std

Std

# **Description**

Std

```
torch_std(self, dim, unbiased = TRUE, keepdim = FALSE)
```

torch\_std\_mean 545

# **Arguments**

self	(Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

unbiased (bool) whether to use the unbiased estimation or not keepdim (bool) whether the output tensor has dim retained or not.

# std(input, unbiased=TRUE) -> Tensor

Returns the standard-deviation of all elements in the input tensor.

If unbiased is FALSE, then the standard-deviation will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

# std(input, dim, unbiased=TRUE, keepdim=False, out=NULL) -> Tensor

Returns the standard-deviation of each row of the input tensor in the dimension dim. If dim is a list of dimensions, reduce over all of them.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

If unbiased is FALSE, then the standard-deviation will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_std(a)

a = torch_randn(c(4, 4))
a
torch_std(a, dim=1)
}
```

torch\_std\_mean

Std\_mean

### **Description**

```
Std_mean
```

```
torch_std_mean(self, dim, unbiased = TRUE, keepdim = FALSE)
```

546 torch\_stft

# **Arguments**

self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

unbiased (bool) whether to use the unbiased estimation or not

keepdim (bool) whether the output tensor has dim retained or not.

### std\_mean(input, unbiased=TRUE) -> (Tensor, Tensor)

Returns the standard-deviation and mean of all elements in the input tensor.

If unbiased is FALSE, then the standard-deviation will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

#### std\_mean(input, dim, unbiased=TRUE, keepdim=False) -> (Tensor, Tensor)

Returns the standard-deviation and mean of each row of the input tensor in the dimension dim. If dim is a list of dimensions, reduce over all of them.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

If unbiased is FALSE, then the standard-deviation will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_std_mean(a)

a = torch_randn(c(4, 4))
a
torch_std_mean(a, 1)
}
```

torch\_stft

Stft

### Description

Stft

torch\_stft 547

### Usage

```
torch_stft(
  input,
  n_fft,
  hop_length = NULL,
  win_length = NULL,
  window = NULL,
  center = TRUE,
  pad_mode = "reflect",
  normalized = FALSE,
  onesided = NULL,
  return_complex = NULL
)
```

# **Arguments**

input	(Tensor) the input tensor
n_fft	(int) size of Fourier transform
hop_length	(int, optional) the distance between neighboring sliding window frames. Default: NULL (treated as equal to floor( $n_fft/4$ ))
win_length	(int, optional) the size of window frame and STFT filter. Default: NULL (treated as equal to $n\_fft$ )
window	(Tensor, optional) the optional window function. Default: NULL (treated as window of all $1\ \mbox{s})$
center	(bool, optional) whether to pad input on both sides so that the $t\text{-th}$ frame is centered at time $t\times hop\_length.$ Default: TRUE
pad_mode	(string, optional) controls the padding method used when center is TRUE. Default: "reflect" $$
normalized	(bool, optional) controls whether to return the normalized STFT results Default: $\ensuremath{FALSE}$
onesided	(bool, optional) controls whether to return half of results to avoid redundancy Default: $\ensuremath{TRUE}$
return_complex	(bool, optional) controls whether to return complex tensors or not.

### **Short-time Fourier transform (STFT).**

Short-time Fourier transform (STFT).

Ignoring the optional batch dimension, this method computes the following expression:

$$X[m,\omega] = \sum_{k=0}^{\text{win\_length-1}} \text{window}[k] \ \text{input}[m \times \text{hop\_length} + k] \ \exp\left(-j\frac{2\pi \cdot \omega k}{\text{win\_length}}\right),$$

548 torch\_stft

where m is the index of the sliding window, and  $\omega$  is the frequency that  $0 \le \omega < n$ \_fft. When onesided is the default value TRUE,

- \* `input` must be either a 1-D time sequence or a 2-D batch of time sequences.
- \* If `hop\_length` is `NULL` (default), it is treated as equal to `floor(n\_fft / 4)`.
- \* If `win\_length` is `NULL` (default), it is treated as equal to `n\_fft`.
- \* `window` can be a 1-D tensor of size `win\_length`, e.g., from
   `torch\_hann\_window`. If `window` is `NULL` (default), it is
   treated as if having \eqn{1} everywhere in the window. If
   \eqn{\mbox{win\\_length} < \mbox{n\\_fft}}, `window` will be padded on
   both sides to length `n\_fft` before being applied.</pre>
- \* If `center` is `TRUE` (default), `input` will be padded on both sides so that the \eqn{t}-th frame is centered at time \eqn{t \times \mbox{hop\\_length}}. Otherwise, the \eqn{t}-th frame begins at time \eqn{t \times \mbox{hop\\_length}}.
- \* `pad\_mode` determines the padding method used on `input` when `center` is `TRUE`. See `torch\_nn.functional.pad` for all available options. Default is `"reflect"`.
- \* If `onesided` is `TRUE` (default), only values for \eqn{\omega}
  in \eqn{\left[0, 1, 2, \dots, \left\lfloor \frac{\mbox{n\\_fft}}{2} \right\rfloor + 1\right]}
  are returned because the real-to-complex Fourier transform satisfies the
  conjugate symmetry, i.e., \eqn{X[m, \omega] = X[m, \mbox{n\\_fft} \omega]^\*.
- \* If `normalized` is `TRUE` (default is `FALSE`), the function returns the normalized STFT results, i.e., multiplied by \eqn{(\mbox{frame\\_length})^{-0.5}}.

Returns the real and the imaginary parts together as one tensor of size  $\eqn{(* \times N \times R \times N)}, where \eqn{*} is the optional batch size of `input`, \eqn{N} is the number of frequencies where STFT is applied, \eqn{T} is the total number of frames used, and each pair in the last dimension represents a complex number as the real part and the imaginary part.$ 

# Warning

This function changed signature at version 0.4.1. Calling with the previous signature may cause error or return incorrect result.

torch\_sub 549

torch\_sub

Sub

# Description

Sub

# Usage

```
torch_sub(self, other, alpha = 1L)
```

### **Arguments**

self (Tensor) the input tensor.

other (Tensor or Scalar) the tensor or scalar to subtract from input

alpha the scalar multiplier for other

# sub(input, other, \*, alpha=1, out=None) -> Tensor

Subtracts other, scaled by alpha, from input.

```
\operatorname{out}_i = \operatorname{input}_i - \operatorname{alpha} \times \operatorname{other}_i
```

Supports broadcasting to a common shape, type promotion, and integer, float, and complex inputs.

# **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(c(1, 2))
b <- torch_tensor(c(0, 1))
torch_sub(a, b, alpha=2)
}</pre>
```

torch\_subtract

Subtract

# Description

Subtract

```
torch_subtract(self, other, alpha = 1L)
```

550 torch\_sum

### **Arguments**

self (Tensor) the input tensor.

other (Tensor or Scalar) the tensor or scalar to subtract from input

alpha the scalar multiplier for other

# subtract(input, other, \*, alpha=1, out=None) -> Tensor

Alias for torch\_sub().

# **Description**

Sum

#### Usage

```
torch_sum(self, dim, keepdim = FALSE, dtype = NULL)
```

# **Arguments**

self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

keepdim (bool) whether the output tensor has dim retained or not.

dtype (torch.dtype, optional) the desired data type of returned tensor. If specified,

the input tensor is casted to dtype before the operation is performed. This is

useful for preventing data type overflows. Default: NULL.

#### sum(input, dtype=NULL) -> Tensor

Returns the sum of all elements in the input tensor.

### sum(input, dim, keepdim=False, dtype=NULL) -> Tensor

Returns the sum of each row of the input tensor in the given dimension dim. If dim is a list of dimensions, reduce over all of them.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

torch\_svd 551

### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_sum(a)

a <- torch_randn(c(4, 4))
a
torch_sum(a, 1)
b <- torch_arange(1, 4 * 5 * 6)$view(c(4, 5, 6))
torch_sum(b, list(2, 1))
}</pre>
```

torch\_svd

Svd

# Description

Svd

### Usage

```
torch_svd(self, some = TRUE, compute_uv = TRUE)
```

# **Arguments**

self (Tensor) the input tensor of size (\*, m, n) where \* is zero or more batch dimensions consisting of  $m \times n$  matrices.

some (bool, optional) controls the shape of returned U and V compute\_uv (bool, optional) option whether to compute U and V or not

### svd(input, some=TRUE, compute\_uv=TRUE) -> (Tensor, Tensor, Tensor)

This function returns a namedtuple (U, S, V) which is the singular value decomposition of a input real matrix or batches of real matrices input such that  $input = U \times diag(S) \times V^T$ .

If some is TRUE (default), the method returns the reduced singular value decomposition i.e., if the last two dimensions of input are m and n, then the returned U and V matrices will contain only min(n,m) orthonormal columns.

If compute\_uv is FALSE, the returned U and V matrices will be zero matrices of shape  $(m \times m)$  and  $(n \times n)$  respectively. some will be ignored here.

552 torch\_t

#### Note

The singular values are returned in descending order. If input is a batch of matrices, then the singular values of each matrix in the batch is returned in descending order.

The implementation of SVD on CPU uses the LAPACK routine ?gesdd (a divide-and-conquer algorithm) instead of ?gesvd for speed. Analogously, the SVD on GPU uses the MAGMA routine gesdd as well.

Irrespective of the original strides, the returned matrix U will be transposed, i.e. with strides U.contiguous().transpose(-2, -1).stride()

Extra care needs to be taken when backward through U and V outputs. Such operation is really only stable when input is full rank with all distinct singular values. Otherwise, NaN can appear as the gradients are not properly defined. Also, notice that double backward will usually do an additional backward through U and V even if the original backward is only on S.

When some = FALSE, the gradients on U[..., :, min(m, n):] and V[..., :, min(m, n):] will be ignored in backward as those vectors can be arbitrary bases of the subspaces.

When compute\_uv = FALSE, backward cannot be performed since U and V from the forward pass is required for the backward operation.

# **Examples**

```
if (torch_is_installed()) {
    a = torch_randn(c(5, 3))
    a
    out = torch_svd(a)
    u = out[[1]]
    s = out[[2]]
    v = out[[3]]
    torch_dist(a, torch_mm(torch_mm(u, torch_diag(s)), v$t()))
    a_big = torch_randn(c(7, 5, 3))
    out = torch_svd(a_big)
    u = out[[1]]
    s = out[[2]]
    v = out[[3]]
    torch_dist(a_big, torch_matmul(torch_matmul(u, torch_diag_embed(s)), v$transpose(-2, -1)))
}
```

torch\_t

T

### **Description**

T

```
torch_t(self)
```

torch\_take 553

### **Arguments**

self

(Tensor) the input tensor.

# t(input) -> Tensor

Expects input to be  $\leq$  2-D tensor and transposes dimensions 0 and 1.

0-D and 1-D tensors are returned as is. When input is a 2-D tensor this is equivalent to transpose(input, 0, 1).

# **Examples**

```
if (torch_is_installed()) {
x = torch_randn(c(2,3))
x
torch_t(x)
x = torch_randn(c(3))
x
torch_t(x)
x = torch_randn(c(2, 3))
x
torch_t(x)
}
```

torch\_take

Take

# Description

Take

# Usage

```
torch_take(self, index)
```

# **Arguments**

self (Tensor) the input tensor.

index (LongTensor) the indices into tensor

# take(input, index) -> Tensor

Returns a new tensor with the elements of input at the given indices. The input tensor is treated as if it were viewed as a 1-D tensor. The result takes the same shape as the indices.

554 torch\_tan

# **Examples**

```
if (torch_is_installed()) {
src = torch_tensor(matrix(c(4,3,5,6,7,8), ncol = 3, byrow = TRUE))
torch_take(src, torch_tensor(c(1, 2, 5), dtype = torch_int64()))
}
```

 $torch\_tan$ 

Tan

# Description

Tan

# Usage

```
torch_tan(self)
```

# Arguments

self

(Tensor) the input tensor.

# tan(input, out=NULL) -> Tensor

Returns a new tensor with the tangent of the elements of input.

```
out_i = tan(input_i)
```

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_tan(a)
}
```

torch\_tanh 555

torch\_tanh

Tanh

# Description

Tanh

# Usage

```
torch_tanh(self)
```

# Arguments

self

(Tensor) the input tensor.

# tanh(input, out=NULL) -> Tensor

Returns a new tensor with the hyperbolic tangent of the elements of input.

```
out_i = tanh(input_i)
```

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_tanh(a)
}
```

 $torch\_tensor$ 

Converts R objects to a torch tensor

# **Description**

Converts R objects to a torch tensor

```
torch_tensor(
  data,
  dtype = NULL,
  device = NULL,
  requires_grad = FALSE,
  pin_memory = FALSE
)
```

556 torch\_tensordot

# **Arguments**

```
data an R atomic vector, matrix or array

dtype a torch_dtype instance

device a device creted with torch_device()

requires_grad if autograd should record operations on the returned tensor.

pin_memory If set, returned tensor would be allocated in the pinned memory.
```

### **Examples**

```
if (torch_is_installed()) {
torch_tensor(c(1, 2, 3, 4))
torch_tensor(c(1, 2, 3, 4), dtype = torch_int())
}
```

torch\_tensordot

Tensordot

# Description

Returns a contraction of a and b over multiple dimensions. tensordot implements a generalized matrix product.

### Usage

```
torch_tensordot(a, b, dims = 2)
```

#### **Arguments**

```
a (Tensor) Left tensor to contract
b (Tensor) Right tensor to contract
dims (int or tuple of two lists of integers) number of dimensions to contract or explicit lists of dimensions for a and b respectively
```

```
if (torch_is_installed()) {

a <- torch_arange(start = 1, end = 60)$reshape(c(3, 4, 5))
b <- torch_arange(start = 1, end = 24)$reshape(c(4, 3, 2))
torch_tensordot(a, b, dims = list(c(2, 1), c(1, 2)))
## Not run:
a = torch_randn(3, 4, 5, device='cuda')
b = torch_randn(4, 5, 6, device='cuda')
c = torch_tensordot(a, b, dims=2)$cpu()

## End(Not run)
}</pre>
```

```
torch_tensor_from_buffer
```

Creates a tensor from a buffer of memory

# **Description**

It creates a tensor without taking ownership of the memory it points to. You must call clone if you want to copy the memory over a new tensor.

### Usage

```
torch_tensor_from_buffer(buffer, shape, dtype = "float")
buffer_from_torch_tensor(tensor)
```

### **Arguments**

buffer An R atomic object containing the data in a contiguous array.

shape The shape of the resulting tensor.

dtype A torch data type for the tresulting tensor.

tensor Tensor object that will be converted into a buffer.

#### **Functions**

• buffer\_from\_torch\_tensor(): Creates a raw vector containing the tensor data. Causes a data copy.

torch\_threshold\_

Threshold\_

# Description

Threshold\_

# Usage

```
torch_threshold_(self, threshold, value)
```

### **Arguments**

self input tensor

threshold The value to threshold at value The value to replace with

# threshold\_(input, threshold, value) -> Tensor

In-place version of torch\_threshold.

558 torch\_topk

torch_topk	Topk
cor cri_copic	торк

# **Description**

Topk

### Usage

```
torch_topk(self, k, dim = -1L, largest = TRUE, sorted = TRUE)
```

# **Arguments**

```
self (Tensor) the input tensor.

k (int) the k in "top-k"

dim (int, optional) the dimension to sort along

largest (bool, optional) controls whether to return largest or smallest elements

sorted (bool, optional) controls whether to return the elements in sorted order
```

# topk(input, k, dim=NULL, largest=TRUE, sorted=TRUE) -> (Tensor, LongTensor)

Returns the k largest elements of the given input tensor along a given dimension.

If dim is not given, the last dimension of the input is chosen.

If largest is FALSE then the k smallest elements are returned.

A namedtuple of (values, indices) is returned, where the indices are the indices of the elements in the original input tensor.

The boolean option sorted if TRUE, will make sure that the returned k elements are themselves sorted

```
if (torch_is_installed()) {
x = torch_arange(1., 6.)
x
torch_topk(x, 3)
}
```

torch\_trace 559

torch\_trace

Trace

# Description

Trace

# Usage

```
torch_trace(self)
```

# **Arguments**

self

the input tensor

# trace(input) -> Tensor

Returns the sum of the elements of the diagonal of the input 2-D matrix.

# Examples

```
if (torch_is_installed()) {
x <- torch_arange(1, 9)$view(c(3, 3))
x
torch_trace(x)
}</pre>
```

torch\_transpose

Transpose

# Description

Transpose

# Usage

```
torch_transpose(self, dim0, dim1)
```

# **Arguments**

self (Tensor) the input tensor.

dim0 (int) the first dimension to be transposed dim1 (int) the second dimension to be transposed 560 torch\_trapz

# transpose(input, dim0, dim1) -> Tensor

Returns a tensor that is a transposed version of input. The given dimensions dim0 and dim1 are swapped.

The resulting out tensor shares it's underlying storage with the input tensor, so changing the content of one would change the content of the other.

# **Examples**

```
if (torch_is_installed()) {
x = torch_randn(c(2, 3))
x
torch_transpose(x, 1, 2)
}
```

torch\_trapz

**Trapz** 

# **Description**

Trapz

# Usage

```
torch_trapz(y, dx = 1L, x, dim = -1L)
```

# Arguments

У	(Tensor) The values of the function to integrate
dx	(float) The distance between points at which y is sampled.
X	(Tensor) The points at which the function y is sampled. If x is not in ascending order, intervals on which it is decreasing contribute negatively to the estimated integral (i.e., the convention $\int_a^b f = -\int_b^a f$ is followed).
dim	(int) The dimension along which to integrate. By default, use the last dimension.

# $trapz(y, x, *, dim=-1) \rightarrow Tensor$

Estimate  $\int y \, dx$  along dim, using the trapezoid rule.

# trapz(y, \*, dx=1, dim=-1) -> Tensor

As above, but the sample points are spaced uniformly at a distance of dx.

torch\_triangular\_solve 561

# Examples

```
if (torch_is_installed()) {

y = torch_randn(list(2, 3))
y
x = torch_tensor(matrix(c(1, 3, 4, 1, 2, 3), ncol = 3, byrow=TRUE))
torch_trapz(y, x = x)
}
```

torch\_triangular\_solve

Triangular\_solve

# Description

Triangular\_solve

# Usage

```
torch_triangular_solve(
   self,
   A,
   upper = TRUE,
   transpose = FALSE,
   unitriangular = FALSE)
```

# Arguments

self	(Tensor) multiple right-hand sides of size $(\ast,m,k)$ where $\ast$ is zero of more batch dimensions $(b)$
A	(Tensor) the input triangular coefficient matrix of size $(\ast,m,m)$ where $\ast$ is zero or more batch dimensions
upper	(bool, optional) whether to solve the upper-triangular system of equations (default) or the lower-triangular system of equations. Default: TRUE.
transpose	(bool, optional) whether ${\cal A}$ should be transposed before being sent into the solver. Default: FALSE.
unitriangular	(bool, optional) whether $A$ is unit triangular. If TRUE, the diagonal elements of $A$ are assumed to be 1 and not referenced from $A$ . Default: FALSE.

562 torch\_tril

# triangular\_solve(input, A, upper=TRUE, transpose=False, unitriangular=False) -> (Tensor, Tensor)

Solves a system of equations with a triangular coefficient matrix A and multiple right-hand sides b. In particular, solves AX = b and assumes A is upper-triangular with the default keyword arguments.

torch\_triangular\_solve(b, A) can take in 2D inputs b, A or inputs that are batches of 2D matrices. If the inputs are batches, then returns batched outputs X

# **Examples**

```
if (torch_is_installed()) {
A = torch_randn(c(2, 2))$triu()
A
b = torch_randn(c(2, 3))
b
torch_triangular_solve(b, A)
}
```

torch\_tril

Tril

### Description

Tril

### Usage

```
torch_tril(self, diagonal = 0L)
```

#### Arguments

self (Tensor) the input tensor.

diagonal (int, optional) the diagonal to consider

### tril(input, diagonal=0, out=NULL) -> Tensor

Returns the lower triangular part of the matrix (2-D tensor) or batch of matrices input, the other elements of the result tensor out are set to 0.

The lower triangular part of the matrix is defined as the elements on and below the diagonal.

The argument diagonal controls which diagonal to consider. If diagonal = 0, all elements on and below the main diagonal are retained. A positive value includes just as many diagonals above the main diagonal, and similarly a negative value excludes just as many diagonals below the main diagonal. The main diagonal are the set of indices  $\{(i,i)\}$  for  $i \in [0, \min\{d_1, d_2\} - 1]$  where  $d_1, d_2$  are the dimensions of the matrix.

torch\_tril\_indices 563

### **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3, 3))
a
torch_tril(a)
b = torch_randn(c(4, 6))
b
torch_tril(b, diagonal=1)
torch_tril(b, diagonal=-1)
}
```

torch\_tril\_indices

Tril\_indices

# **Description**

Tril\_indices

# Usage

```
torch_tril_indices(
  row,
  col,
  offset = 0,
  dtype = NULL,
  device = NULL,
  layout = NULL
)
```

# Arguments

(int) number of rows in the 2-D matrix. row (int) number of columns in the 2-D matrix. col offset (int) diagonal offset from the main diagonal. Default: if not provided, 0. (torch.dtype, optional) the desired data type of returned tensor. Default: if dtype NULL, torch\_long. (torch.device, optional) the desired device of returned tensor. Default: if device NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types. layout (torch.layout, optional) currently only support torch\_strided.

564 torch\_triu

### tril\_indices(row, col, offset=0, dtype=torch.long, device='cpu', layout=torch.strided) -> Tensor

Returns the indices of the lower triangular part of a row-by- col matrix in a 2-by-N Tensor, where the first row contains row coordinates of all indices and the second row contains column coordinates. Indices are ordered based on rows and then columns.

The lower triangular part of the matrix is defined as the elements on and below the diagonal.

The argument offset controls which diagonal to consider. If offset = 0, all elements on and below the main diagonal are retained. A positive value includes just as many diagonals above the main diagonal, and similarly a negative value excludes just as many diagonals below the main diagonal. The main diagonal are the set of indices  $\{(i,i)\}$  for  $i \in [0, \min\{d_1, d_2\} - 1]$  where  $d_1, d_2$  are the dimensions of the matrix.

#### Note

When running on CUDA, `row  $\star$  col` must be less than \eqn{2^{59}} to prevent overflow during calculation.

# **Examples**

```
if (torch_is_installed()) {
## Not run:
a = torch_tril_indices(3, 3)
a
a = torch_tril_indices(4, 3, -1)
a
a = torch_tril_indices(4, 3, 1)
a
## End(Not run)
}
```

torch\_triu

Triu

# Description

Triu

# Usage

```
torch_triu(self, diagonal = 0L)
```

#### **Arguments**

```
self (Tensor) the input tensor.
diagonal (int, optional) the diagonal to consider
```

torch\_triu\_indices 565

### triu(input, diagonal=0, out=NULL) -> Tensor

Returns the upper triangular part of a matrix (2-D tensor) or batch of matrices input, the other elements of the result tensor out are set to 0.

The upper triangular part of the matrix is defined as the elements on and above the diagonal.

The argument diagonal controls which diagonal to consider. If diagonal = 0, all elements on and above the main diagonal are retained. A positive value excludes just as many diagonals above the main diagonal, and similarly a negative value includes just as many diagonals below the main diagonal. The main diagonal are the set of indices  $\{(i,i)\}$  for  $i \in [0, \min\{d_1, d_2\} - 1]$  where  $d_1, d_2$  are the dimensions of the matrix.

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(3, 3))
a
torch_triu(a)
torch_triu(a, diagonal=1)
torch_triu(a, diagonal=-1)
b = torch_randn(c(4, 6))
b
torch_triu(b, diagonal=1)
torch_triu(b, diagonal=-1)
}
```

torch\_triu\_indices

Triu\_indices

# Description

Triu\_indices

```
torch_triu_indices(
  row,
  col,
  offset = 0,
  dtype = NULL,
  device = NULL,
  layout = NULL
```

566 torch\_triu\_indices

# Arguments

row	(int) number of rows in the 2-D matrix.
col	(int) number of columns in the 2-D matrix.
offset	(int) diagonal offset from the main diagonal. Default: if not provided, 0.
dtype	(torch.dtype, optional) the desired data type of returned tensor. Default: if NULL, torch_long.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, uses the current device for the default tensor type (see torch_set_default_tensor_type). device will be the CPU for CPU tensor types and the current CUDA device for CUDA tensor types.
layout	(torch.layout, optional) currently only support torch_strided.

### triu\_indices(row, col, offset=0, dtype=torch.long, device='cpu', layout=torch.strided) -> Tensor

Returns the indices of the upper triangular part of a row by col matrix in a 2-by-N Tensor, where the first row contains row coordinates of all indices and the second row contains column coordinates. Indices are ordered based on rows and then columns.

The upper triangular part of the matrix is defined as the elements on and above the diagonal.

The argument offset controls which diagonal to consider. If offset = 0, all elements on and above the main diagonal are retained. A positive value excludes just as many diagonals above the main diagonal, and similarly a negative value includes just as many diagonals below the main diagonal. The main diagonal are the set of indices  $\{(i,i)\}$  for  $i \in [0, \min\{d_1, d_2\} - 1]$  where  $d_1, d_2$  are the dimensions of the matrix.

# Note

When running on CUDA, `row \* col` must be less than  $\eqn{2^{59}}$  to prevent overflow during calculation.

```
if (torch_is_installed()) {
## Not run:
a = torch_triu_indices(3, 3)
a
a = torch_triu_indices(4, 3, -1)
a
a = torch_triu_indices(4, 3, 1)
a
## End(Not run)
}
```

torch\_true\_divide 567

torch\_true\_divide

TRUE\_divide

# **Description**

TRUE\_divide

# Usage

```
torch_true_divide(self, other)
```

# Arguments

self (Tensor) the dividend other (Tensor or Scalar) the divisor

# true\_divide(dividend, divisor) -> Tensor

Performs "true division" that always computes the division in floating point. Analogous to division in Python 3 and equivalent to torch\_div except when both inputs have bool or integer scalar types, in which case they are cast to the default (floating) scalar type before the division.

$$out_i = \frac{dividend_i}{divisor}$$

# **Examples**

```
if (torch_is_installed()) {
    dividend = torch_tensor(c(5, 3), dtype=torch_int())
    divisor = torch_tensor(c(3, 2), dtype=torch_int())
    torch_true_divide(dividend, divisor)
    torch_true_divide(dividend, 2)
}
```

torch\_trunc

Trunc

# Description

Trunc

```
torch_trunc(self)
```

568 torch\_unbind

# **Arguments**

self

(Tensor) the input tensor.

# trunc(input, out=NULL) -> Tensor

Returns a new tensor with the truncated integer values of the elements of input.

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(4))
a
torch_trunc(a)
}
```

torch\_unbind

Unbind

# Description

Unbind

# Usage

```
torch_unbind(self, dim = 1L)
```

# Arguments

self (Tensor) the tensor to unbind dim (int) dimension to remove

# unbind(input, dim=0) -> seq

Removes a tensor dimension.

Returns a tuple of all slices along a given dimension, already without it.

```
if (torch_is_installed()) {
  torch_unbind(torch_tensor(matrix(1:9, ncol = 3, byrow=TRUE)))
}
```

torch\_unique\_consecutive

Unique\_consecutive

### **Description**

Unique\_consecutive

### Usage

```
torch_unique_consecutive(
  self,
  return_inverse = FALSE,
  return_counts = FALSE,
  dim = NULL
)
```

# **Arguments**

```
self (Tensor) the input tensor

return_inverse (bool) Whether to also return the indices for where elements in the original input ended up in the returned unique list.

return_counts (bool) Whether to also return the counts for each unique element.

dim (int) the dimension to apply unique. If NULL, the unique of the flattened input is returned, default: NULL
```

### TEST

Eliminates all but the first element from every consecutive group of equivalent elements.

.. note:: This function is different from [`torch\_unique`] in the sense that this function
 only eliminates consecutive duplicate values. This semantics is similar to `std::unique`
 in C++.

```
if (torch_is_installed()) {
x = torch_tensor(c(1, 1, 2, 2, 3, 1, 1, 2))
output = torch_unique_consecutive(x)
output
torch_unique_consecutive(x, return_inverse=TRUE)
torch_unique_consecutive(x, return_counts=TRUE)
}
```

570 torch\_unsafe\_split

torch\_unsafe\_chunk

Unsafe\_chunk

### **Description**

Unsafe\_chunk

# Usage

```
torch_unsafe_chunk(self, chunks, dim = 1L)
```

# **Arguments**

self (Tensor) the tensor to split

chunks (int) number of chunks to return

dim (int) dimension along which to split the tensor

# unsafe\_chunk(input, chunks, dim=0) -> List of Tensors

Works like torch\_chunk() but without enforcing the autograd restrictions on inplace modification of the outputs.

# Warning

This function is safe to use as long as only the input, or only the outputs are modified inplace after calling this function. It is user's responsibility to ensure that is the case. If both the input and one or more of the outputs are modified inplace, gradients computed by autograd will be silently incorrect.

# **Description**

Unsafe\_split

# Usage

```
torch_unsafe_split(self, split_size, dim = 1L)
```

# **Arguments**

self (Tensor) tensor to split.

split\_size (int) size of a single chunk or list of sizes for each chunk

dim (int) dimension along which to split the tensor.

torch\_unsqueeze 571

### unsafe\_split(tensor, split\_size\_or\_sections, dim=0) -> List of Tensors

Works like torch\_split() but without enforcing the autograd restrictions on inplace modification of the outputs.

# Warning

This function is safe to use as long as only the input, or only the outputs are modified inplace after calling this function. It is user's responsibility to ensure that is the case. If both the input and one or more of the outputs are modified inplace, gradients computed by autograd will be silently incorrect.

torch\_unsqueeze

Unsqueeze

### **Description**

Unsqueeze

# Usage

```
torch_unsqueeze(self, dim)
```

# Arguments

```
self (Tensor) the input tensor.

dim (int) the index at which to insert the singleton dimension
```

### unsqueeze(input, dim) -> Tensor

Returns a new tensor with a dimension of size one inserted at the specified position.

The returned tensor shares the same underlying data with this tensor.

A dim value within the range [-input.dim() - 1, input.dim() + 1) can be used. Negative dim will correspond to unsqueeze applied at dim = dim + input.dim() + 1.

```
if (torch_is_installed()) {
x = torch_tensor(c(1, 2, 3, 4))
torch_unsqueeze(x, 1)
torch_unsqueeze(x, 2)
}
```

572 torch\_var

torch\_vander

Vander

# **Description**

Vander

# Usage

```
torch_vander(x, N = NULL, increasing = FALSE)
```

### **Arguments**

x (Tensor) 1-D input tensor.

N (int, optional) Number of columns in the output. If N is not specified, a square

array is returned (N = len(x)).

increasing (bool, optional) Order of the powers of the columns. If TRUE, the powers in-

crease from left to right, if FALSE (the default) they are reversed.

# vander(x, N=None, increasing=FALSE) -> Tensor

Generates a Vandermonde matrix.

The columns of the output matrix are elementwise powers of the input vector  $x^{(N-1)}, x^{(N-2)}, ..., x^0$ . If increasing is TRUE, the order of the columns is reversed  $x^0, x^1, ..., x^{(N-1)}$ . Such a matrix with a geometric progression in each row is named for Alexandre-Theophile Vandermonde.

# Examples

```
if (torch_is_installed()) {
x <- torch_tensor(c(1, 2, 3, 5))
torch_vander(x)
torch_vander(x, N=3)
torch_vander(x, N=3, increasing=TRUE)
}</pre>
```

torch\_var

Var

# **Description**

Var

```
torch_var(self, dim, unbiased = TRUE, keepdim = FALSE)
```

torch\_var\_mean 573

### **Arguments**

self	(Tensor) the input tensor.
------	----------------------------

dim (int or tuple of ints) the dimension or dimensions to reduce.

unbiased (bool) whether to use the unbiased estimation or not keepdim (bool) whether the output tensor has dim retained or not.

# var(input, unbiased=TRUE) -> Tensor

Returns the variance of all elements in the input tensor.

If unbiased is FALSE, then the variance will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

# var(input, dim, keepdim=False, unbiased=TRUE, out=NULL) -> Tensor

Returns the variance of each row of the input tensor in the given dimension dim.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

If unbiased is FALSE, then the variance will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_var(a)

a = torch_randn(c(4, 4))
a
torch_var(a, 1)
}
```

torch\_var\_mean

Var\_mean

# **Description**

Var\_mean

```
torch_var_mean(self, dim, unbiased = TRUE, keepdim = FALSE)
```

574 torch\_vdot

# **Arguments**

self (Tensor) the input tensor.

dim (int or tuple of ints) the dimension or dimensions to reduce.

unbiased (bool) whether to use the unbiased estimation or not keepdim (bool) whether the output tensor has dim retained or not.

### var\_mean(input, unbiased=TRUE) -> (Tensor, Tensor)

Returns the variance and mean of all elements in the input tensor.

If unbiased is FALSE, then the variance will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

# var\_mean(input, dim, keepdim=False, unbiased=TRUE) -> (Tensor, Tensor)

Returns the variance and mean of each row of the input tensor in the given dimension dim.

If keepdim is TRUE, the output tensor is of the same size as input except in the dimension(s) dim where it is of size 1. Otherwise, dim is squeezed (see torch\_squeeze), resulting in the output tensor having 1 (or len(dim)) fewer dimension(s).

If unbiased is FALSE, then the variance will be calculated via the biased estimator. Otherwise, Bessel's correction will be used.

# **Examples**

```
if (torch_is_installed()) {
a = torch_randn(c(1, 3))
a
torch_var_mean(a)

a = torch_randn(c(4, 4))
a
torch_var_mean(a, 1)
}
```

torch\_vdot

Vdot

# **Description**

Vdot

```
torch_vdot(self, other)
```

575

# **Arguments**

```
self (Tensor) first tensor in the dot product. Its conjugate is used if it's complex. other (Tensor) second tensor in the dot product.
```

# vdot(input, other, \*, out=None) -> Tensor

Computes the dot product (inner product) of two tensors. The vdot(a, b) function handles complex numbers differently than dot(a, b). If the first argument is complex, the complex conjugate of the first argument is used for the calculation of the dot product.

#### Note

This function does not broadcast.

### **Examples**

```
if (torch_is_installed()) {

torch_vdot(torch_tensor(c(2, 3)), torch_tensor(c(2, 1)))
if (FALSE) {
    a <- torch_tensor(list(1 +2i, 3 - 1i))
    b <- torch_tensor(list(2 +1i, 4 - 0i))
torch_vdot(a, b)
torch_vdot(b, a)
}
}</pre>
```

### torch\_view\_as\_complex View\_as\_complex

# Description

```
View_as_complex
```

#### **Usage**

```
torch_view_as_complex(self)
```

### **Arguments**

self (Tensor) the input tensor.

# view\_as\_complex(input) -> Tensor

Returns a view of input as a complex tensor. For an input complex tensor of size  $m1, m2, \ldots, mi, 2$ , this function returns a new complex tensor of size  $m1, m2, \ldots, mi$  where the last dimension of the input tensor is expected to represent the real and imaginary components of complex numbers.

576 torch\_view\_as\_real

### Warning

torch\_view\_as\_complex is only supported for tensors with torch\_dtype torch\_float64() and torch\_float32(). The input is expected to have the last dimension of size 2. In addition, the tensor must have a stride of 1 for its last dimension. The strides of all other dimensions must be even numbers.

### **Examples**

```
if (torch_is_installed()) {
  if (FALSE) {
    x=torch_randn(c(4, 2))
    x
    torch_view_as_complex(x)
  }
}
```

torch\_view\_as\_real

View\_as\_real

# **Description**

```
View_as_real
```

### Usage

```
torch_view_as_real(self)
```

### **Arguments**

self

(Tensor) the input tensor.

# view\_as\_real(input) -> Tensor

Returns a view of input as a real tensor. For an input complex tensor of size  $m1, m2, \ldots, mi$ , this function returns a new real tensor of size  $m1, m2, \ldots, mi$ , 2, where the last dimension of size 2 represents the real and imaginary components of complex numbers.

### Warning

torch\_view\_as\_real() is only supported for tensors with complex dtypes.

```
if (torch_is_installed()) {

if (FALSE) {
    x <- torch_randn(4, dtype=torch_cfloat())
    x

torch_view_as_real(x)</pre>
```

torch\_vstack 577

} }

torch\_vstack

Vstack

# Description

Vstack

#### Usage

```
torch_vstack(tensors)
```

## Arguments

tensors

(sequence of Tensors) sequence of tensors to concatenate

# vstack(tensors, \*, out=None) -> Tensor

Stack tensors in sequence vertically (row wise).

This is equivalent to concatenation along the first axis after all 1-D tensors have been reshaped by torch\_atleast\_2d().

#### **Examples**

```
if (torch_is_installed()) {
a <- torch_tensor(c(1, 2, 3))
b <- torch_tensor(c(4, 5, 6))
torch_vstack(list(a,b))
a <- torch_tensor(rbind(1,2,3))
b <- torch_tensor(rbind(4,5,6))
torch_vstack(list(a,b))</pre>
```

torch\_where

Where

#### **Description**

Where

# Usage

```
torch_where(condition, self = NULL, other = NULL)
```

578 torch\_zeros

## **Arguments**

```
condition (BoolTensor) When TRUE (nonzero), yield x, otherwise yield y self (Tensor) values selected at indices where condition is TRUE other (Tensor) values selected at indices where condition is FALSE
```

#### where(condition, x, y) -> Tensor

Return a tensor of elements selected from either x or y, depending on condition.

The operation is defined as:

$$out_i = \left\{ \begin{array}{ll} \mathbf{x}_i & \text{if condition}_i \\ \mathbf{y}_i & \text{otherwise} \end{array} \right.$$

#### where(condition) -> tuple of LongTensor

torch\_where(condition) is identical to torch\_nonzero(condition, as\_tuple=TRUE).

#### Note

```
The tensors `condition`, `x`, `y` must be broadcastable . See also torch_nonzero().
```

# **Examples**

```
if (torch_is_installed()) {
## Not run:
x = torch_randn(c(3, 2))
y = torch_ones(c(3, 2))
x
torch_where(x > 0, x, y)
## End(Not run)
}
```

torch\_zeros

Zeros

# Description

Zeros

torch\_zeros\_like 579

#### Usage

```
torch_zeros(
    ...,
    names = NULL,
    dtype = NULL,
    layout = NULL,
    device = NULL,
    requires_grad = FALSE
)
```

#### **Arguments**

... a sequence of integers defining the shape of the output tensor. Can be a variable

number of arguments or a collection like a list or tuple.

names optional dimension names

dtype (torch.dtype, optional) the desired data type of returned tensor. Default: if

NULL, uses a global default (see torch\_set\_default\_tensor\_type).

layout (torch.layout, optional) the desired layout of returned Tensor. Default: torch\_strided.

device (torch.device, optional) the desired device of returned tensor. Default: if

NULL, uses the current device for the default tensor type (see torch\_set\_default\_tensor\_type).

device will be the CPU for CPU tensor types and the current CUDA device for

CUDA tensor types.

Zeros\_like

requires\_grad (bool, optional) If autograd should record operations on the returned tensor. De-

fault: FALSE.

# zeros(\*size, out=NULL, dtype=NULL, layout=torch.strided, device=NULL, requires\_grad=False) -> Tensor

Returns a tensor filled with the scalar value 0, with the shape defined by the variable argument size.

#### **Examples**

```
if (torch_is_installed()) {
torch_zeros(c(2, 3))
torch_zeros(c(5))
}
```

torch\_zeros\_like

#### **Description**

Zeros\_like

580 torch\_zeros\_like

#### Usage

```
torch_zeros_like(
  input,
  dtype = NULL,
  layout = NULL,
  device = NULL,
  requires_grad = FALSE,
  memory_format = torch_preserve_format()
)
```

# Arguments

input	(Tensor) the size of input will determine size of the output tensor.
dtype	(torch.dtype, optional) the desired data type of returned Tensor. Default: if NULL, defaults to the dtype of input.
layout	(torch.layout, optional) the desired layout of returned tensor. Default: if NULL, defaults to the layout of input.
device	(torch.device, optional) the desired device of returned tensor. Default: if NULL, defaults to the device of input.
requires_grad	(bool, optional) If autograd should record operations on the returned tensor. Default: FALSE.
memory_format	(torch.memory_format, optional) the desired memory format of returned Tensor. Default: torch_preserve_format.

# zeros\_like(input, dtype=NULL, layout=NULL, device=NULL, requires\_grad=False, memory\_format=torch.preserve\_form

Returns a tensor filled with the scalar value 0, with the same size as input. torch\_zeros\_like(input) is equivalent to torch\_zeros(input.size(), dtype=input.dtype, layout=input.layout, device=input.device).

#### Warning

As of 0.4, this function does not support an out keyword. As an alternative, the old torch\_zeros\_like(input, out=output) is equivalent to torch\_zeros(input.size(), out=output).

```
if (torch_is_installed()) {
input = torch_empty(c(2, 3))
torch_zeros_like(input)
}
```

with\_detect\_anomaly 581

with\_detect\_anomaly

Context-manager that enable anomaly detection for the autograd engine.

# Description

This does two things:

#### Usage

```
with_detect_anomaly(code)
```

#### **Arguments**

code

Code that will be executed in the detect anomaly context.

#### **Details**

- Running the forward pass with detection enabled will allow the backward pass to print the traceback of the forward operation that created the failing backward function.
- Any backward computation that generate "nan" value will raise an error.

## Warning

This mode should be enabled only for debugging as the different tests will slow down your program execution.

```
if (torch_is_installed()) {
x <- torch_randn(2, requires_grad = TRUE)
y <- torch_randn(1)
b <- (x^y)$sum()
y$add_(1)

try({
   b$backward()

   with_detect_anomaly({
      b$backward()
   })
})
})</pre>
```

582 with\_enable\_grad

with\_enable\_grad

Enable grad

# Description

Context-manager that enables gradient calculation. Enables gradient calculation, if it has been disabled via with\_no\_grad.

#### Usage

```
with_enable_grad(code)
local_enable_grad(.env = parent.frame())
```

#### **Arguments**

code code to be executed with gradient recording.

. env The environment to use for scoping.

#### **Details**

This context manager is thread local; it will not affect computation in other threads.

#### **Functions**

• local\_enable\_grad(): Locally enable gradient computations.

```
if (torch_is_installed()) {

x <- torch_tensor(1, requires_grad = TRUE)
with_no_grad({
    with_enable_grad({
        y <- x * 2
    })
})
y$backward()
x$grad
}</pre>
```

with\_no\_grad 583

with\_no\_grad

Temporarily modify gradient recording.

# Description

Temporarily modify gradient recording.

# Usage

```
with_no_grad(code)
local_no_grad(.env = parent.frame())
```

# Arguments

code code to be executed with no gradient recording.

. env The environment to use for scoping.

# **Functions**

• local\_no\_grad(): Disable autograd until it goes out of scope

```
if (torch_is_installed()) {
x <- torch_tensor(runif(5), requires_grad = TRUE)
with_no_grad({
    x$sub_(torch_tensor(as.numeric(1:5)))
})
x
x$grad
}</pre>
```

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