EE-559 - Deep learning

4.1. DAG networks

François Fleuret

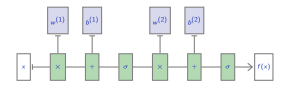
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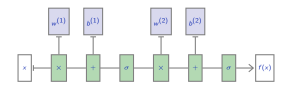




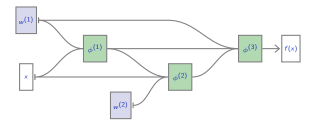
We can generalize an MLP



We can generalize an MLP



to an arbitrary "Directed Acyclic Graph" (DAG) of operators



If $(a_1, \ldots, a_Q) = \phi(b_1, \ldots, b_R)$, we use the notation

$$\begin{bmatrix} \frac{\partial \mathbf{a}}{\partial \mathbf{b}} \end{bmatrix} = J_{\phi} = \begin{pmatrix} \frac{\partial \mathbf{a}_{1}}{\partial b_{1}} & \cdots & \frac{\partial \mathbf{a}_{1}}{\partial b_{R}} \\ \vdots & \ddots & \vdots \\ \frac{\partial \mathbf{a}_{Q}}{\partial b_{1}} & \cdots & \frac{\partial \mathbf{a}_{Q}}{\partial b_{R}} \end{pmatrix}.$$

It does not specify at which point this is computed, but it will always be for the forward-pass activations.

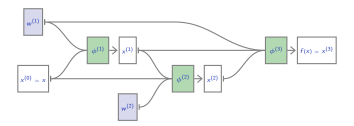
If $(a_1, \ldots, a_Q) = \phi(b_1, \ldots, b_R)$, we use the notation

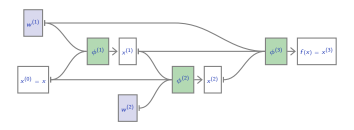
$$\begin{bmatrix} \frac{\partial \mathbf{a}}{\partial b} \end{bmatrix} = J_{\phi} = \begin{pmatrix} \frac{\partial a_1}{\partial b_1} & \dots & \frac{\partial a_1}{\partial b_R} \\ \vdots & \ddots & \vdots \\ \frac{\partial a_Q}{\partial b_1} & \dots & \frac{\partial a_Q}{\partial b_R} \end{pmatrix}.$$

It does not specify at which point this is computed, but it will always be for the forward-pass activations.

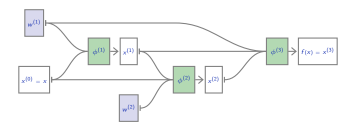
Also, if $(a_1, ..., a_Q) = \phi(b_1, ..., b_R, c_1, ..., c_S)$, we use

$$\left[\frac{\partial a}{\partial c}\right] = J_{\phi|c} = \begin{pmatrix} \frac{\partial a_1}{\partial c_1} & \cdots & \frac{\partial a_1}{\partial c_S} \\ \vdots & \ddots & \vdots \\ \frac{\partial a_Q}{\partial c_1} & \cdots & \frac{\partial a_Q}{\partial c_S} \end{pmatrix}.$$



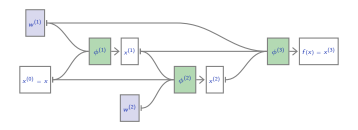


$$x^{(0)} = x$$

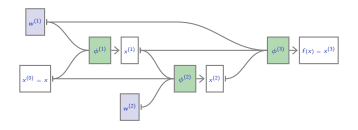


$$x^{(0)} = x$$

 $x^{(1)} = \phi^{(1)}(x^{(0)}; w^{(1)})$



$$\begin{aligned} x^{(0)} &= x \\ x^{(1)} &= \phi^{(1)}(x^{(0)}; w^{(1)}) \\ x^{(2)} &= \phi^{(2)}(x^{(0)}, x^{(1)}; w^{(2)}) \end{aligned}$$

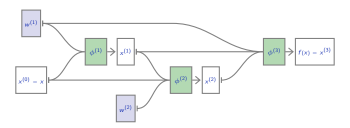


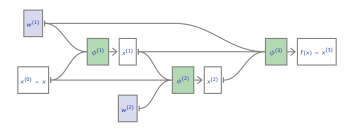
$$x^{(0)} = x$$

$$x^{(1)} = \phi^{(1)}(x^{(0)}; w^{(1)})$$

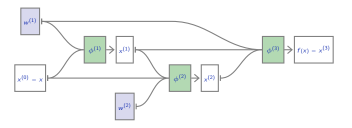
$$x^{(2)} = \phi^{(2)}(x^{(0)}, x^{(1)}; w^{(2)})$$

$$f(x) = x^{(3)} = \phi^{(3)}(x^{(1)}, x^{(2)}; w^{(1)})$$

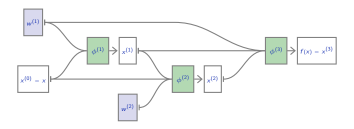




$$\left[\frac{\partial \ell}{\partial \mathbf{x}^{(2)}}\right] = \left[\frac{\partial \mathbf{x}^{(3)}}{\partial \mathbf{x}^{(2)}}\right] \left[\frac{\partial \ell}{\partial \mathbf{x}^{(3)}}\right] = J_{\phi^{(3)}|\mathbf{x}^{(2)}} \left[\frac{\partial \ell}{\partial \mathbf{x}^{(3)}}\right]$$



$$\begin{split} & \left[\frac{\partial \ell}{\partial x^{(2)}}\right] = \left[\frac{\partial x^{(3)}}{\partial x^{(2)}}\right] \left[\frac{\partial \ell}{\partial x^{(3)}}\right] = J_{\phi^{(3)}|x^{(2)}} \left[\frac{\partial \ell}{\partial x^{(3)}}\right] \\ & \left[\frac{\partial \ell}{\partial x^{(1)}}\right] = \left[\frac{\partial x^{(2)}}{\partial x^{(1)}}\right] \left[\frac{\partial \ell}{\partial x^{(2)}}\right] + \left[\frac{\partial x^{(3)}}{\partial x^{(1)}}\right] \left[\frac{\partial \ell}{\partial x^{(3)}}\right] = J_{\phi^{(2)}|x^{(1)}} \left[\frac{\partial \ell}{\partial x^{(2)}}\right] + J_{\phi^{(3)}|x^{(1)}} \left[\frac{\partial \ell}{\partial x^{(3)}}\right] \end{split}$$

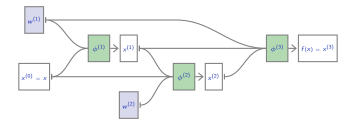


$$\begin{bmatrix} \frac{\partial \ell}{\partial x^{(2)}} \end{bmatrix} = \begin{bmatrix} \frac{\partial x^{(3)}}{\partial x^{(2)}} \end{bmatrix} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(3)}} \end{bmatrix} = J_{\phi^{(3)}|x^{(2)}} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(3)}} \end{bmatrix}$$

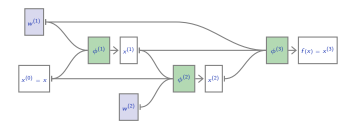
$$\begin{bmatrix} \frac{\partial \ell}{\partial x^{(1)}} \end{bmatrix} = \begin{bmatrix} \frac{\partial x^{(2)}}{\partial x^{(1)}} \end{bmatrix} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(2)}} \end{bmatrix} + \begin{bmatrix} \frac{\partial x^{(3)}}{\partial x^{(1)}} \end{bmatrix} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(3)}} \end{bmatrix} = J_{\phi^{(2)}|x^{(1)}} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(2)}} \end{bmatrix} + J_{\phi^{(3)}|x^{(1)}} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(3)}} \end{bmatrix}$$

$$\begin{bmatrix} \frac{\partial \ell}{\partial x^{(0)}} \end{bmatrix} = \begin{bmatrix} \frac{\partial x^{(1)}}{\partial x^{(0)}} \end{bmatrix} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(1)}} \end{bmatrix} + \begin{bmatrix} \frac{\partial x^{(2)}}{\partial x^{(0)}} \end{bmatrix} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(2)}} \end{bmatrix} = J_{\phi^{(1)}|x^{(0)}} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(1)}} \end{bmatrix} + J_{\phi^{(2)}|x^{(0)}} \begin{bmatrix} \frac{\partial \ell}{\partial x^{(2)}} \end{bmatrix}$$

Backward pass, derivatives w.r.t parameters

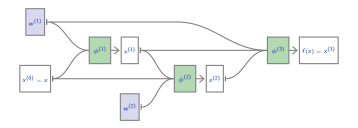


Backward pass, derivatives w.r.t parameters



$$\left[\frac{\partial \ell}{\partial w^{(1)}}\right] = \left[\frac{\partial x^{(1)}}{\partial w^{(1)}}\right] \left[\frac{\partial \ell}{\partial x^{(1)}}\right] + \left[\frac{\partial x^{(3)}}{\partial w^{(1)}}\right] \left[\frac{\partial \ell}{\partial x^{(3)}}\right] = J_{\phi^{(1)}|w^{(1)}} \left[\frac{\partial \ell}{\partial x^{(1)}}\right] + J_{\phi^{(3)}|w^{(1)}} \left[\frac{\partial \ell}{\partial x^{(3)}}\right]$$

Backward pass, derivatives w.r.t parameters



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So if we have a library of "tensor operators", and implementations of

$$(x_1, \dots, x_d, w) \mapsto \phi(x_1, \dots, x_d; w)$$

$$\forall c, (x_1, \dots, x_d, w) \mapsto J_{\phi|x_c}(x_1, \dots, x_d; w)$$

$$(x_1, \dots, x_d, w) \mapsto J_{\phi|w}(x_1, \dots, x_d; w),$$

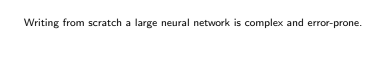
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$$(x_1, \ldots, x_d, w) \mapsto J_{\phi|w}(x_1, \ldots, x_d; w),$$

we can build an arbitrary directed acyclic graph with these operators at the nodes, compute the response of the resulting mapping, and compute its gradient with back-prop.



Writing from scratch a large neural network is complex and error-prone.

Multiple frameworks provide libraries of tensor operators and mechanisms to combine them into DAGs and automatically differentiate them.

	Language(s)	License	Main backer
PyTorch	Python	BSD	Facebook
Caffe2	C++, Python	Apache	Facebook
TensorFlow	Python, $C++$	Apache	Google
MXNet	Python, C++, R, Scala	Apache	Amazon
CNTK	Python, $C++$	MIT	Microsoft
Torch	Lua	BSD	Facebook
Theano	Python	BSD	U. of Montreal
Caffe	C++	BSD 2 clauses	U. of CA, Berkeley

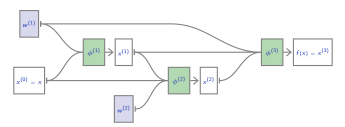
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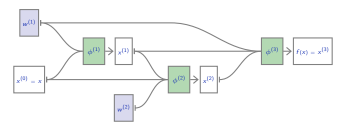
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One approach is to define the nodes and edges of such a DAG statically (Torch, TensorFlow, Caffe, Theano, etc.)

In TensorFlow, to run a forward/backward pass on



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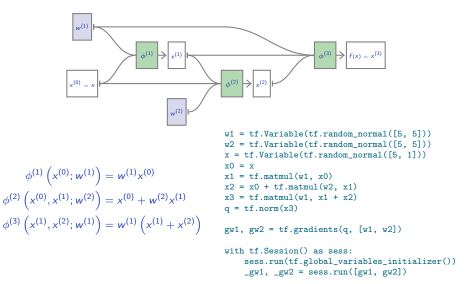


$$\phi^{(1)}\left(x^{(0)}; w^{(1)}\right) = w^{(1)}x^{(0)}$$

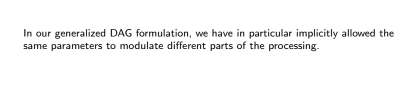
$$\phi^{(2)}\left(x^{(0)}, x^{(1)}; w^{(2)}\right) = x^{(0)} + w^{(2)}x^{(1)}$$

$$\phi^{(3)}\left(x^{(1)}, x^{(2)}; w^{(1)}\right) = w^{(1)}\left(x^{(1)} + x^{(2)}\right)$$

In TensorFlow, to run a forward/backward pass on

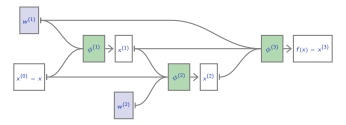


Weight sharing

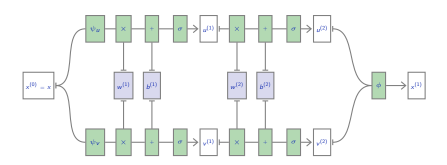


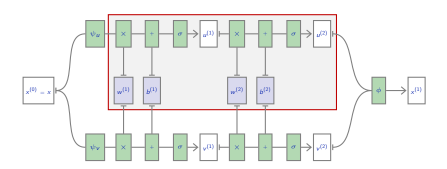
In our generalized DAG formulation, we have in particular implicitly allowed the same parameters to modulate different parts of the processing.

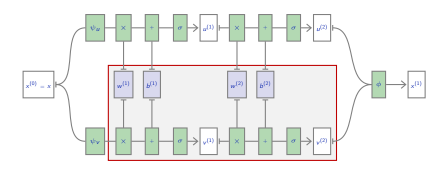
For instance $w^{(1)}$ in our example parametrizes both $\phi^{(1)}$ and $\phi^{(3)}$.

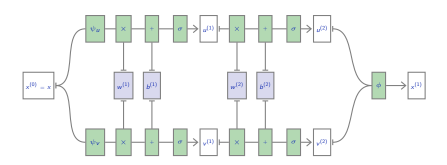


This is called weight sharing.











EE-559 – Deep learning

4.2. Autograd

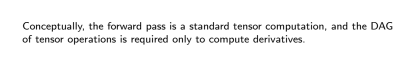
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https://fleuret.org/ee559/

Mon Feb 18 13:34:25 UTC 2019







Conceptually, the forward pass is a standard tensor computation, and the DAG of tensor operations is required only to compute derivatives.

When executing tensor operations, PyTorch can automatically construct on-the-fly the graph of operations to compute the gradient of any quantity with respect to any tensor involved.

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This "autograd" mechanism (Paszke et al., 2017) has two main benefits:

- Simpler syntax: one just need to write the forward pass as a standard sequence of Python operations,
- greater flexibility: since the graph is not static, the forward pass can be dynamically modulated.

A Tensor has a Boolean field requires_grad, set to False by default, which states if PyTorch should build the graph of operations so that gradients with respect to it can be computed.

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The result of a tensorial operation has this flag to True if any of its operand has it to True.

```
>>> x = torch.tensor([ 1., 2. ])
>>> y = torch.tensor([ 4., 5. ])
>>> z = torch.tensor([ 7., 3. ])
>>> x.requires_grad
False
>>> (x + y).requires_grad
False
>>> z.requires_grad = True
>>> (x + z).requires_grad
True
```



Only floating point type tensors can have their gradient computed.

```
>>> x = torch.tensor([1., 10.])
>>> x.requires_grad = True
>>> x = torch.tensor([1, 10])
>>> x.requires_grad = True
Traceback (most recent call last):
   File "<stdin>", line 1, in <module>
RuntimeError: only Tensors of floating point dtype can require gradients
```



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RuntimeError: only Tensors of floating point dtype can require gradients
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The method requires_grad_(value = True) set requires_grad to value, which is True by default.

torch.autograd.grad(outputs, inputs) computes and returns the gradient of outputs with respect to inputs.

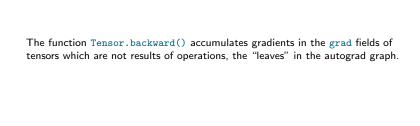
```
>>> t = torch.tensor([1., 2., 4.]).requires_grad_()
>>> u = torch.tensor([10., 20.]).requires_grad_()
>>> a = t.pow(2).sum() + u.log().sum()
>>> torch.autograd.grad(a, (t, u))
(tensor([2., 4., 8.]), tensor([0.1000, 0.0500]))
```

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(tensor([2., 4., 8.]), tensor([0.1000, 0.0500]))
```

inputs can be a single tensor, but the result is still a [one element] tuple.

If outputs is a tuple, the result is the sum of the gradients of its elements.



The function Tensor.backward() accumulates gradients in the grad fields of tensors which are not results of operations, the "leaves" in the autograd graph.

```
>>> x = torch.tensor([ -3., 2., 5. ]).requires_grad_()
>>> u = x.pow(3).sum()
>>> x.grad
>>> u.backward()
>>> x.grad
tensor([27., 12., 75.])
```

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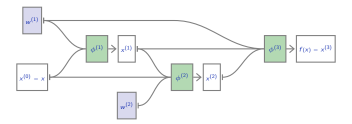
This function is an alternative to torch.autograd.grad(...) and standard for training models.



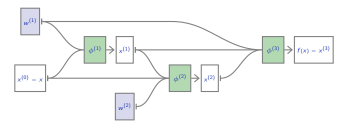
Tensor.backward() accumulates the gradients in the different Tensors, so one may have to set them to zero before calling it.

This accumulating behavior is desirable in particular to compute the gradient of a loss summed over several "mini-batches," or the gradient of a sum of losses.

So we can run a forward/backward pass on

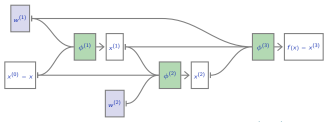


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$$\begin{split} \phi^{(1)}\left(x^{(0)};w^{(1)}\right) &= w^{(1)}x^{(0)} \\ \phi^{(2)}\left(x^{(0)},x^{(1)};w^{(2)}\right) &= x^{(0)} + w^{(2)}x^{(1)} \\ \phi^{(3)}\left(x^{(1)},x^{(2)};w^{(1)}\right) &= w^{(1)}\left(x^{(1)} + x^{(2)}\right) \end{split}$$

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$$\phi^{(1)}\left(x^{(0)}; w^{(1)}\right) = w^{(1)}x^{(0)}$$

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$$\phi^{(3)}\left(x^{(1)}, x^{(2)}; w^{(1)}\right) = w^{(1)}\left(x^{(1)} + x^{(2)}\right)$$

q.backward()

The autograd machinery

The autograd graph is encoded through the fields grad_fn of Tensors, and the fields next_functions of Functions.

```
>>> x = torch.tensor([ 1.0, -2.0, 3.0, -4.0 ]).requires_grad_()
>>> a = x.abs()
>>> s = a.sum()
>>> s
tensor(10., grad_fn=<SumBackward0>)
>>> s.grad_fn.next_functions
((<AbsBackward object at 0x7ffb2b1462b0>, 0),)
>>> s.grad_fn.next_functions[0][0].next_functions
((<AbcGumulateGrad object at 0x7ffb2b146278>, 0),)
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((<AccumulateGrad object at 0x7ffb2b146278>, 0),)
```

We will come back to this later to write our own Functions.

We can visualize the full graph built during a computation.

```
x = torch.tensor([1., 2., 2.]).requires_grad_()
q = x.norm()
```

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```
x = torch.tensor([1., 2., 2.]).requires_grad_()
q = x.norm()

AccumulateGrad
x[3]
```

This graph was generated with

 ${\tt https://fleuret.org/git/agtree2dot}$ and ${\tt Graphviz}.$

```
w1 = torch.rand(20, 10).requires_grad_()
b1 = torch.rand(20).requires_grad_()
w2 = torch.rand(5, 20).requires_grad_()
b2 = torch.rand(5).requires_grad_()

x = torch.rand(10)
h = torch.tanh(w1 @ x + b1)
y = torch.tanh(w2 @ h + b2)

target = torch.rand(5)
loss = (y - target).pow(2).mean()
```

```
MeanBackward1
                                                                          PowBackward0
                                                                          ThSubBackward
w1 = torch.rand(20, 10).requires_grad_()
b1 = torch.rand(20).requires_grad_()
                                                                          TanhBackward
w2 = torch.rand(5, 20).requires_grad_()
b2 = torch.rand(5).requires_grad_()
                                                                          ThAddBackward
x = torch.rand(10)
h = torch.tanh(w1 @ x + b1)
                                                                     MvBackward
                                                                               AccumulateGrad
y = torch.tanh(w2 @ h + b2)
                                                           AccumulateGrad
                                                                       TanhBackward
                                                                                 b2 [5]
target = torch.rand(5)
loss = (y - target).pow(2).mean()
                                                                      ThAddBackward
                                                             w2 [5, 20]
                                                                  MvBackward
                                                                           AccumulateGrad
                                                                 AccumulateGrad
                                                                              b1 [20]
                                                                  w1 [20, 10]
```

loss []

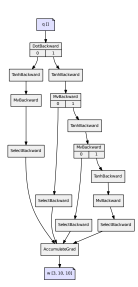
```
w = torch.rand(3, 10, 10).requires_grad_()
def blah(k, x):
    for i in range(k):
        x = torch.tanh(w[i] @ x)
    return x

u = blah(1, torch.rand(10))
v = blah(3, torch.rand(10))
```

q = u.dot(v)

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u = blah(1, torch.rand(10))
v = blah(3, torch.rand(10))
q = u.dot(v)
```





Although they are related, the autograd graph is not the network's structure, but the graph of operations to compute the gradient. It can be data-dependent and miss or replicate sub-parts of the network.

The torch.no_grad() context switches off the autograd machinery, and can be used for operations such as parameter updates.

```
w = torch.empty(10, 784).normal_(0, 1e-3).requires_grad_()
b = torch.empty(10).normal_(0, 1e-3).requires_grad_()

for k in range(10001):
    y_hat = x @ w.t() + b
    loss = (y_hat - y).pow(2).mean()

    w.grad, b.grad = None, None
    loss.backward()

    with torch.no_grad():
        w -= eta * w.grad
        b -= eta * b.grad
```

The detach() method creates a tensor which shares the data, but does not require gradient computation, and is not connected to the current graph.

This method should be used when the gradient should not be propagated beyond a variable, or to update leaf tensors.

```
a = torch.tensor( 0.5).requires_grad_()
b = torch.tensor(-0.5).requires_grad_()

for k in range(100):
    l = (a - 1)**2 + (b + 1)**2 + (a - b)**2
    ga, gb = torch.autograd.grad(l, (a, b))
    with torch.no_grad():
        a -= eta * ga
        b -= eta * gb

print('%.06f' % a.item(), '%.06f' % b.item())

prints
```

0.333333 -0.333333

```
a = torch.tensor( 0.5).requires_grad_()
b = torch.tensor(-0.5).requires_grad_()

for k in range(100):
    1 = (a - 1)**2 + (b + 1)**2 + (a.detach() - b)**2
    ga, gb = torch.autograd.grad(1, (a, b))
    with torch.no_grad():
        a -= eta * ga
        b -= eta * gb

print('%.06f' % a.item(), '%.06f' % b.item())
```

François Fleuret

1,000000 -0,000000

Autograd can also track the computation of the gradient itself, to allows higher-order derivatives. This is specified with create_graph = True:

Autograd can also track the computation of the gradient itself, to allows higher-order derivatives. This is specified with create_graph = True:



In-place operations may corrupt values required to compute the gradient, and this is tracked down by autograd.

```
>>> x = torch.tensor([1., 2., 3.]).requires_grad_()
>>> v = x.sin()
>>> 1 = v.sum()
>>> 1.backward()
>>> y = x.sin()
>>> v += 1
>>> 1 = v.sum()
>>> 1.backward()
>>> y = x.sin()
>>> y *= y
>>> 1 = v.sum()
>>> 1.backward()
Traceback (most recent call last):
1.../
RuntimeError: one of the variables needed for gradient computation has
been modified by an inplace operation
```

They are also prohibited on so-called "leaf" tensors, which are not the results of operations but the initial inputs to the whole computation.



References

A. Paszke, S. Gross, S. Chintala, G. Chanan, E. Yang, Z. DeVito, Z. Lin, A. Desmaison, L. Antiga, and A. Lerer. Automatic differentiation in pytorch. In *Proceedings of the NIPS Autodiff workshop*, 2017.

EE-559 - Deep learning

4.3. PyTorch modules and batch processing

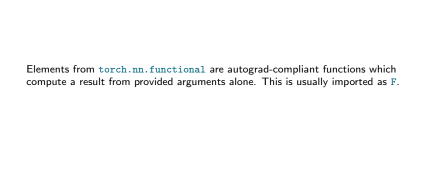
François Fleuret

https://fleuret.org/ee559/

Mon Feb 18 13:34:30 UTC 2019







Elements from ${\tt torch.nn.functional}$ are autograd-compliant functions which compute a result from provided arguments alone. This is usually imported as F.

Subclasses of torch.nn.Module are losses and network components. The latter embed parameters to be optimized during training.

Parameters are of the type torch.nn.Parameter which is a Tensor with requires_grad to True, and known to be a model parameter by various utility functions, in particular torch.nn.Module.parameters().



Functions and modules from torch.nn process batches of inputs stored in a tensor whose first dimension indexes them, and produce a corresponding tensor with the same additional dimension.

E.g. a fully connected layer $\mathbb{R}^C \to \mathbb{R}^D$ expects as input a tensor of size $N \times C$ and computes a tensor of size $N \times D$, where N is the number of samples and can vary from a call to another.

torch.nn.functional.relu(input, inplace=False)

takes a tensor of any size as input, applies ReLU on each value to produce a result tensor of same size.

```
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```

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```
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```

takes a tensor of any size as input, applies ReLU on each value to produce a result tensor of same size.

inplace indicates if the operation should modify the argument itself. This may be desirable to reduce the memory footprint of the processing.

torch.nn.Linear(in_features, out_features, bias=True)

implements a $\mathbb{R}^C \to \mathbb{R}^D$ fully-connected layer. It takes as input a tensor of size $N \times C$ and produce a tensor of size $N \times D$.

```
implements a \mathbb{R}^C \to \mathbb{R}^D fully-connected layer. It takes as input a tensor of size N \times C and produce a tensor of size N \times D.

>>> f = nn.Linear(in_features = 10, out_features = 4)
>>> for n, p in f.named_parameters(): print(n, p.size())
...

weight torch.Size([4, 10])
bias torch.Size([4])
>>> x = torch.empty(523, 10).normal_()
>>> y = f(x)
>>> y.size()
torch.Size([523, 4])
```

```
torch.nn.Linear(in_features, out_features, bias=True) implements a \mathbb{R}^C \to \mathbb{R}^D fully-connected layer. It takes as input a tensor of size N \times C and produce a tensor of size N \times D.

>>> f = nn.Linear(in_features = 10, out_features = 4)
>>> for n, p in f.named_parameters(): print(n, p.size())
...

weight torch.Size([4, 10])
bias torch.Size([4])
>>> x = torch.empty(523, 10).normal_()
>>> y = f(x)
>>> y.size()
torch.Size([523, 4])
```



The weights and biases are automatically randomized at creation. We will come back to that later.

torch.nn.MSELoss()

implements the Mean Square Error loss: the sum of the component-wise squared difference, divided by the total number of components in the tensors.

```
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```

implements the Mean Square Error loss: the sum of the component-wise squared difference, divided by the total number of components in the tensors.

```
>>> f = torch.nn.MSELoss()
>>> x = torch.tensor([[ 3. ]])
>>> y = torch.tensor([[ 0. ]])
>>> f(x, y)
tensor(9.)
>>> x = torch.tensor([[ 3., 0., 0., 0. ]])
>>> y = torch.tensor([[ 0., 0., 0., 0. ]])
>>> f(x, y)
tensor(2.2500)
```

```
torch.nn.MSELoss()
```

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>>> f(x, y)
tensor(2.2500)
```

The first parameter of a loss is traditionally called the **input** and the second the **target**. These two quantities may be of different dimensions or even types for some losses (e.g. for classification).



Criteria do not accept a tensor with requires_grad to True for target.

```
>>> import torch
>>> f = torch.nn.MSELoss()
>>> x = torch.tensor([ 3., 2. ]).requires_grad_()
>>> y = torch.tensor([ 0., -2. ]).requires_grad_()
>>> f(x, y)
Traceback (most recent call last):
/.../
AssertionError: nn criterions don't compute the gradient w.r.t.
targets - please mark these tensors as not requiring gradients
```

Batch processing

Functions and modules from torch.nn process samples by batches. This is motivated by the computational speed-up it induces.

To evaluate a module on a sample, both the module's parameters and the sample have to be first copied into **cache memory**, which is fast but small.

For any model of reasonable size, only a fraction of its parameters can be kept in cache, so a module's parameters have to be copied there every time it is used. Functions and modules from torch.nn process samples by batches. This is motivated by the computational speed-up it induces.

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These memory transfers are slower than the computation itself.

This is the main reason for batch processing: it cuts down to one per module per batch the number of copies of parameters to the cache. Functions and modules from torch.nn process samples by batches. This is motivated by the computational speed-up it induces.

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These memory transfers are slower than the computation itself.

This is the main reason for batch processing: it cuts down to one per module per batch the number of copies of parameters to the cache.

It also cuts down the use of Python loops, which are awfully slow.

Consider a model composed of three modules

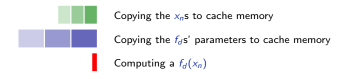
$$\textit{f} = \textit{f}_{3} \circ \textit{f}_{2} \circ \textit{f}_{1},$$

and we want to compute $f(x_1), f(x_2), f(x_3)$.

Consider a model composed of three modules

$$f=f_3\circ f_2\circ f_1,$$

and we want to compute $f(x_1), f(x_2), f(x_3)$.



Processing samples one by one:

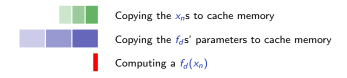


Time

Consider a model composed of three modules

$$f = f_3 \circ f_2 \circ f_1$$
,

and we want to compute $f(x_1), f(x_2), f(x_3)$.



Processing samples one by one:



Time

Batch processing:



With

```
def timing(x, w, batch = False, nb = 101):
    t = torch.zeros(nb)

for u in range(0, t.size(0)):
    t0 = time.perf_counter()
    if batch:
        y = x.mm(w.t())
    else:
        y = torch.empty(x.size(0), w.size(0))
        for k in range(y.size(0)): y[k] = w.mv(x[k])
    y.is_cuda and torch.cuda.synchronize()
    t[u] = time.perf_counter() - t0

return t.median().item()
```

prints

Batch-processing speed-up on CPU 4.6 Batch-processing speed-up on GPU 144.4

Formally, we have to revisit a bit some expressions we saw previously for fully connected layers. We had

$$\forall l, n, \ w^{(l)} \in \mathbb{R}^{d_l \times d_{l-1}}, \ x_n^{(l-1)} \in \mathbb{R}^{d_{l-1}}, \ s_n^{(l)} = w^{(l)} x_n^{(l-1)}.$$

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From now on, we will use row vectors, so that we can represent a series of samples as a 2d array with the first index being the sample's index.

$$x = \begin{pmatrix} x_{1,1} & \dots & x_{1,D} \\ \vdots & \ddots & \vdots \\ x_{N,1} & \dots & x_{N,D} \end{pmatrix} = \begin{pmatrix} (x_1)^T \\ \vdots \\ (x_N)^T \end{pmatrix},$$

which is an element of $\mathbb{R}^{N \times D}$.

To make all sample row vectors and apply a linear operator, we want

$$\forall n, \ s_n^{(l)} = \left(w^{(l)} \left(x_n^{(l-1)} \right)^T \right)^T = x_n^{(l-1)} \left(w^{(l)} \right)^T$$

which gives a tensorial expression for the full batch

$$s^{(l)} = x^{(l-1)} \left(w^{(l)} \right)^T.$$

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which gives a tensorial expression for the full batch

$$s^{(l)} = x^{(l-1)} \left(w^{(l)} \right)^T.$$

And in torch/nn/functional.py

```
def linear(input, weight, bias=None):
   if input.dim() == 2 and bias is not None:
      # fused op is marginally faster
      return torch.addmm(bias, input, weight.t())
   output = input.matmul(weight.t())
   if bias is not None:
      output += bias
   return output
```

Similarly for the backward pass of a linear layer we get

$$\left[\!\!\left[\frac{\partial \mathcal{L}}{\partial w^{(l)}}\right]\!\!\right] = \left[\!\!\left[\frac{\partial \mathcal{L}}{\partial x^{(l)}}\right]\!\!\right]^T x^{(l-1)},$$

and

$$\left[\left[\frac{\partial \mathcal{L}}{\partial x^{(l)}}\right]\right] = \left[\left[\frac{\partial \ell}{\partial x^{(l+1)}}\right]\right] w^{(l+1)}.$$



EE-559 - Deep learning

4.4. Convolutions

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https://fleuret.org/ee559/

Mon Feb 18 13:34:32 UTC 2019





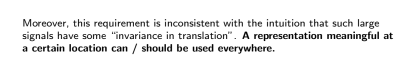
If they were handled as normal "unstructured" vectors, large-dimension signals such as sound samples or images would require models of intractable size.

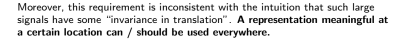
For instance a linear layer taking a 256×256 RGB image as input, and producing an image of same size would require

$$(256 \times 256 \times 3)^2 \simeq 3.87e + 10$$

parameters, with the corresponding memory footprint ($\simeq\!150\mbox{Gb}$!), and excess of capacity.

1 / 23



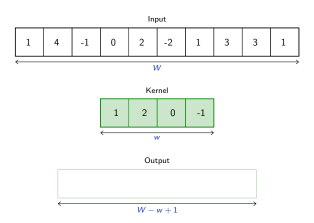


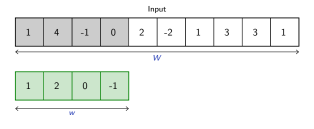
A convolution layer embodies this idea. It applies the same linear transformation locally, everywhere, and preserves the signal structure.

Input

1	4	-1	0	2	-2	1	3	3	1
←									

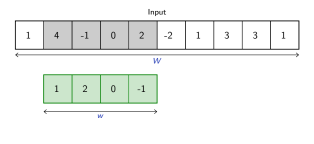
W





Output



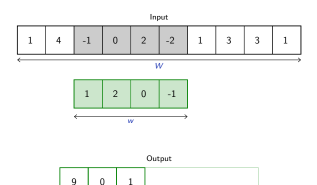


Output

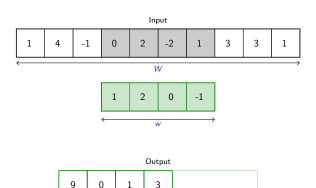
W - w + 1

9

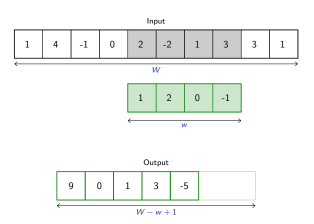
0

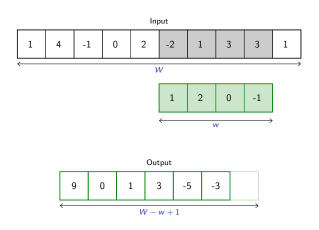


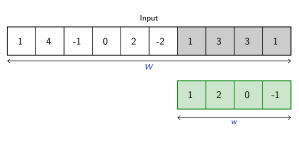
W - w + 1

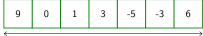


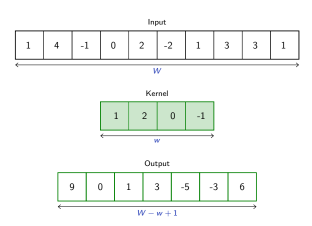
W - w + 1











Formally, in 1d, given

$$x = (x_1, \ldots, x_W)$$

and a "convolution kernel" (or "filter") of width w

$$u=(u_1,\ldots,u_w)$$

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$$u = (u_1, \ldots, u_w)$$

the convolution $x \circledast u$ is a vector of size W - w + 1, with

$$(x \circledast u)_i = \sum_{j=1}^w x_{i-1+j} u_j$$

= $(x_i, \dots, x_{i+w-1}) \cdot u$

for instance

$$(1,2,3,4) \circledast (3,2) = (3+4,6+6,9+8) = (7,12,17).$$

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= $(x_i, \dots, x_{i+w-1}) \cdot u$

for instance

$$(1,2,3,4) \otimes (3,2) = (3+4,6+6,9+8) = (7,12,17).$$

 \triangle

This differs from the usual convolution since the kernel and the signal are both visited in increasing index order.

$$(0,0,0,0,1,2,3,4,4,4,4) \otimes (-1,1) = (0,0,0,1,1,1,1,0,0,0).$$

$$(0,0,0,0,1,2,3,4,4,4,4) \circledast (-1,1) = (0,0,0,1,1,1,1,0,0,0).$$



$$(0,0,0,0,1,2,3,4,4,4,4) \otimes (-1,1) = (0,0,0,1,1,1,1,0,0,0).$$



or crude "template matcher", e.g.



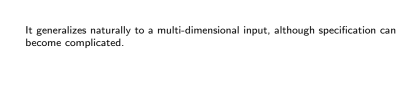
$$(0,0,0,0,1,2,3,4,4,4,4) \circledast (-1,1) = (0,0,0,1,1,1,1,0,0,0).$$



or crude "template matcher", e.g.



Both of these computation examples are indeed "invariant by translation".



It generalizes naturally to a multi-dimensional input, although specification can become complicated.

Its most usual form for "convolutional networks" processes a 3d tensor as input (i.e. a multi-channel 2d signal) to output a 2d tensor. The kernel is not swiped across channels, just across rows and columns.

In this case, if the input tensor is of size $C \times H \times W$, and the kernel is $C \times h \times w$, the output is $(H - h + 1) \times (W - w + 1)$.

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We say "2d signal" even though it has $\mathcal C$ channels, since it is a feature vector indexed by a 2d location without structure on the feature indexes.

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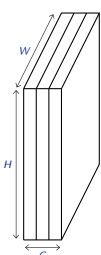
In this case, if the input tensor is of size $C \times H \times W$, and the kernel is $C \times h \times w$, the output is $(H - h + 1) \times (W - w + 1)$.



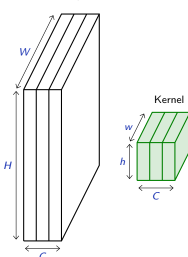
We say "2d signal" even though it has C channels, since it is a feature vector indexed by a 2d location without structure on the feature indexes.

In a standard convolution layer, D such convolutions are combined to generate a $D \times (H-h+1) \times (W-w+1)$ output.

Input



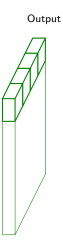
Input

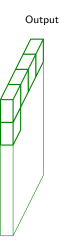










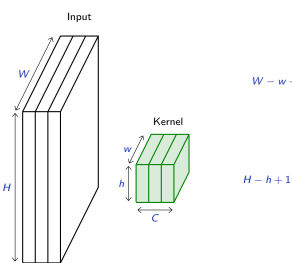


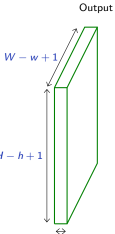


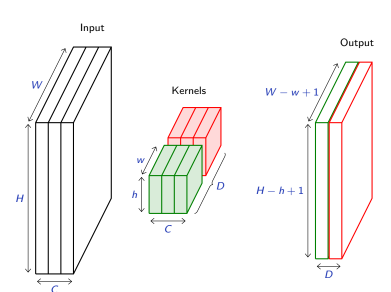


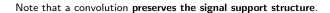










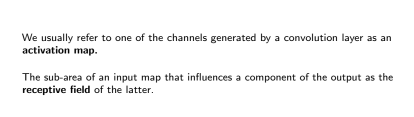


A 1d signal is converted into a 1d signal, a 2d signal into a 2d, and neighboring parts of the input signal influence neighboring parts of the output signal.

Note that a convolution preserves the signal support structure.

A 1d signal is converted into a 1d signal, a 2d signal into a 2d, and neighboring parts of the input signal influence neighboring parts of the output signal.

A 3d convolution can be used if the channel index has some metric meaning, such as time for a series of grayscale video frames. Otherwise swiping across channels makes no sense.



We usually refer to one of the channels generated by a convolution layer as an **activation map.**

The sub-area of an input map that influences a component of the output as the **receptive field** of the latter.

In the context of convolutional networks, a standard linear layer is called a **fully connected layer** since every input influences every output.

Implements a 2d convolution, where weight contains the kernels, and is $D \times C \times h \times w$, bias is of dimension D, input is of dimension

$$N \times C \times H \times W$$

and the result is of dimension

$$N \times D \times (H-h+1) \times (W-w+1)$$
.

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$$N \times C \times H \times W$$

and the result is of dimension

$$N \times D \times (H-h+1) \times (W-w+1)$$
.

```
>>> weight = torch.empty(5, 4, 2, 3).normal_()
>>> bias = torch.empty(5).normal_()
>>> input = torch.empty(117, 4, 10, 3).normal_()
>>> output = torch.nn.functional.conv2d(input, weight, bias)
>>> output.size()
torch.Size([117, 5, 9, 1])
```

Implements a 2d convolution, where weight contains the kernels, and is $D \times C \times h \times w$, bias is of dimension D, input is of dimension

$$N \times C \times H \times W$$

and the result is of dimension

$$N \times D \times (H-h+1) \times (W-w+1)$$
.

```
>>> weight = torch.empty(5, 4, 2, 3).normal_()
>>> bias = torch.empty(5).normal_()
>>> input = torch.empty(117, 4, 10, 3).normal_()
>>> output = torch.nn.functional.conv2d(input, weight, bias)
>>> output.size()
torch.Size([117, 5, 9, 1])
```

Similar functions implement 1d and 3d convolutions.

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```
x = mnist train.train data[12].float().view(1, 1, 28, 28)
weight = torch.emptv(5, 1, 3, 3)
weight[0, 0] = torch.tensor([ [ 0., 0., 0. ],
                              [ 0., 1., 0.],
[ 0., 0., 0.]])
weight[1, 0] = torch.tensor([ [ 1., 1., 1.],
                              [ 1., 1., 1.],
[ 1., 1., 1.])
weight[2, 0] = torch.tensor([ [ -1., 0., 1. ],
                              [ -1.. 0.. 1. ].
                              [-1., 0., 1. ] ])
weight[3, 0] = torch.tensor([ [ -1., -1., -1. ],
                              [ 0., 0., 0.],
[ 1., 1., 1.])
weight[4, 0] = torch.tensor([ [ 0., -1., 0. ],
                              Г-1.. 4.. -1. ].
                              [0, -1, 0, 1]
v = torch.nn.functional.conv2d(x, weight)
```

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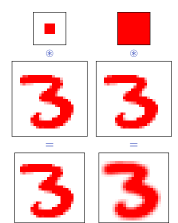


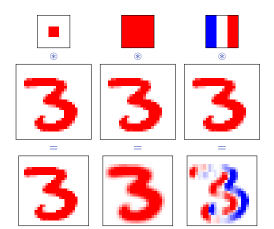
*

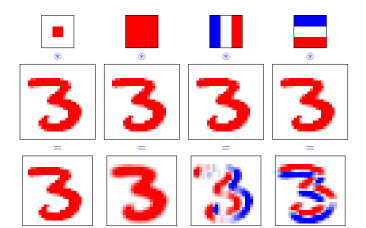


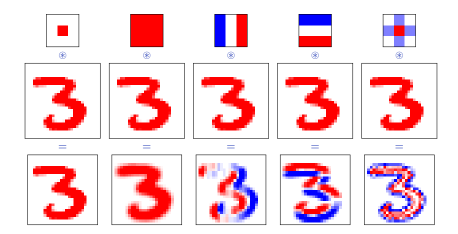
_











Wraps the convolution into a Module, with the kernels and biases as Parameter properly randomized at creation.

The kernel size is either a pair (h, w) or a single value k interpreted as (k, k).

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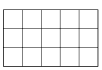
```
>>> f = nn.Conv2d(in_channels = 4, out_channels = 5, kernel_size = (2, 3))
>>> for n, p in f.named_parameters(): print(n, p.size())
...
weight torch.Size([5, 4, 2, 3])
bias torch.Size([5])
>>> x = torch.empty(117, 4, 10, 3).normal_()
>>> y = f(x)
>>> y.size()
torch.Size([117, 5, 9, 1])
```

Padding and stride

Convolutions have two additional standard parameters:

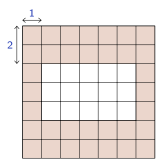
- The padding specifies the size of a zeroed frame added around the input,
- the **stride** specifies a step size when moving the kernel across the signal.

Here with $C \times 3 \times 5$ as input



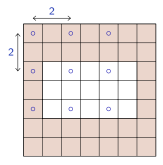
Input

Here with $C \times 3 \times 5$ as input, a padding of (2,1)

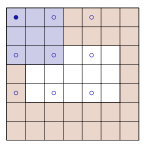


Input

Here with $C \times 3 \times 5$ as input, a padding of (2,1), a stride of (2,2)

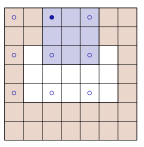


Input



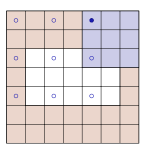
Output

Input



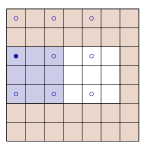
Output

Input



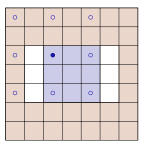


Input



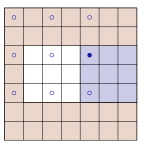


Input



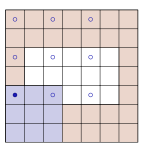


Input



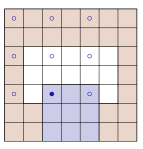


Input



Output

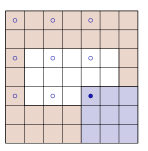
Input



Output

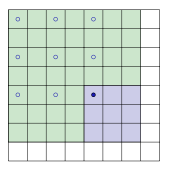
Input

Here with $C \times 3 \times 5$ as input, a padding of (2,1), a stride of (2,2), and a kernel of size $C \times 3 \times 3$, the output is $1 \times 3 \times 3$.



Output

Input





A convolution with a stride greater than ${\bf 1}$ may not cover the input map completely, hence may ignore some of the input values.

Dilated convolution

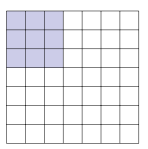
Convolution operations admit one more standard parameter that we have not discussed yet: The dilation, which modulates the expansion of the filter support (Yu and Koltun, 2015).

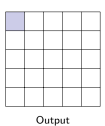
It is 1 for standard convolutions, but can be greater, in which case the resulting operation can be envisioned as a convolution with a regularly sparsified filter.

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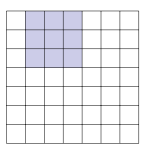
It is 1 for standard convolutions, but can be greater, in which case the resulting operation can be envisioned as a convolution with a regularly sparsified filter.

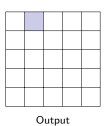
This notion comes from signal processing, where it is referred to as *algorithme à trous*. hence the term sometime used of "convolution à trous".



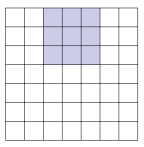


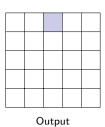
Input



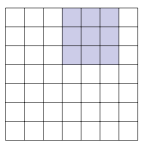


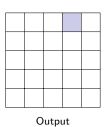
Input



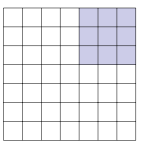


Input



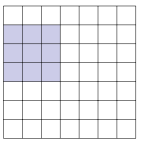


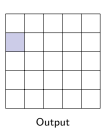
Input

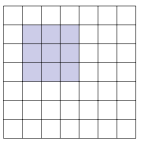


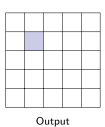
Output

Input

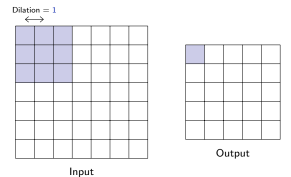


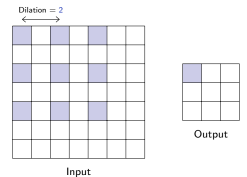


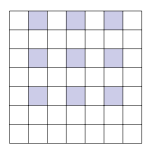




Input

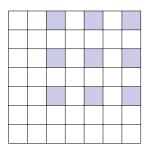






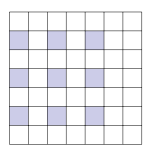


Input



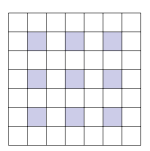


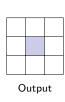
Input



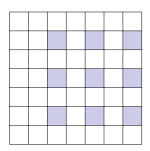


Input



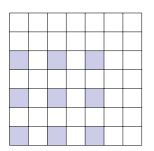


Input



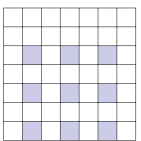


Input

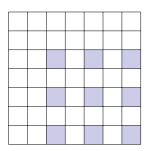




Input









Input

A convolution with a 1d kernel of size k and dilation d can be interpreted as a convolution with a filter of size 1 + (k-1)d with only k non-zero coefficients.

For with k=3 and d=4, the difference between the input map size and the output map size is 1+(3-1)4-1=8.

```
>>> x = torch.empty(1, 1, 20, 30).normal_()
>>> 1 = nn.Conv2d(1, 1, kernel_size = 3, dilation = 4)
>>> 1(x).size()
torch.Size([1, 1, 12, 22])
```

Having a dilation greater than one increases the units' receptive field size without increasing the number of parameters.

Convolutions with stride or dilation strictly greater than one reduce the activation map size, for instance to make a final classification decision.

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Convolutions with stride or dilation strictly greater than one reduce the activation map size, for instance to make a final classification decision.

Such networks have the advantage of simplicity:

- non-linear operations are only in the activation function,
- joint operations that combine multiple activations to produce one are only in linear layers.

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References

abs/1511.07122v3, 2015.

F. Yu and V. Koltun. Multi-scale context aggregation by dilated convolutions. CoRR,

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4.5. Pooling

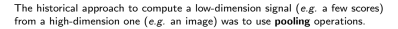
François Fleuret

https://fleuret.org/ee559/

Mon Feb 18 13:34:33 UTC 2019





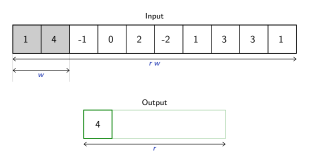


Such an operation aims at grouping several activations into a single "more meaningful" one.

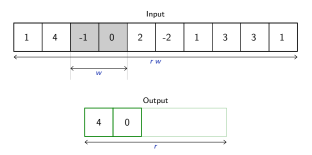
For instance in 1d with a kernel of size 2:

Input									
1	4	-1	0	2	-2	1	3	3	1
<i>rw</i> →									

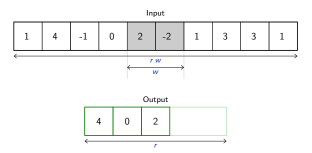
For instance in 1d with a kernel of size 2:



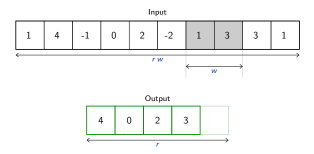
For instance in 1d with a kernel of size 2:



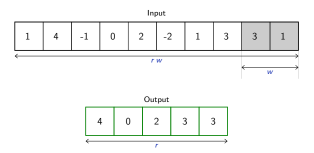
For instance in 1d with a kernel of size 2:



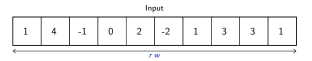
For instance in 1d with a kernel of size 2:



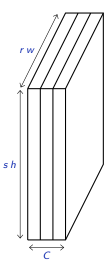
For instance in 1d with a kernel of size 2:

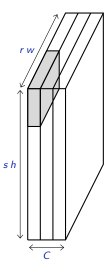


For instance in 1d with a kernel of size 2:

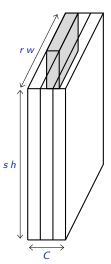


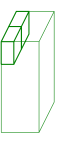


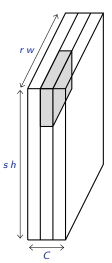


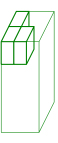


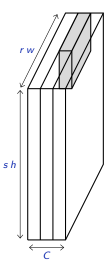


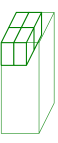


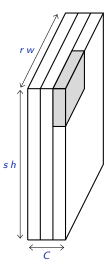


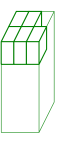


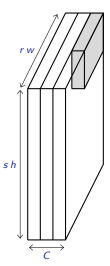


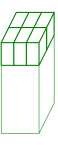


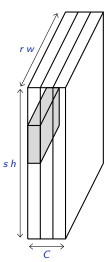


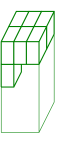


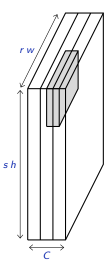




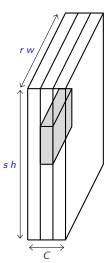


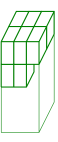


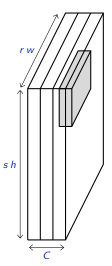


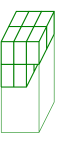


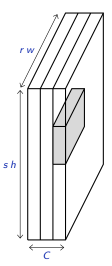


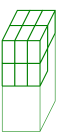


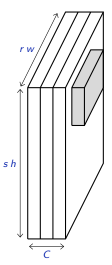


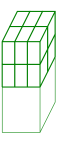


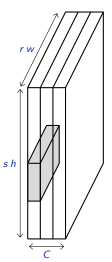






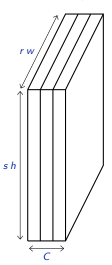




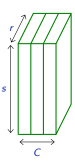




Input

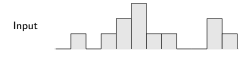


Output

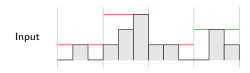


More practically, it provides a pseudo-invariance to deformations that result into local translations.

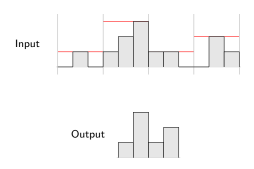
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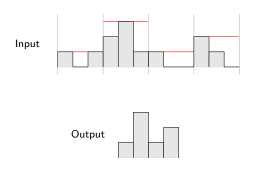


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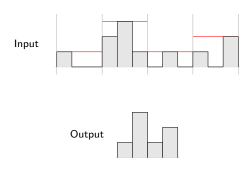
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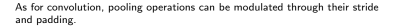
```
torch.nn.functional.max_pool2d(input, kernel_size,
                               stride=None, padding=0, dilation=1,
                               ceil_mode=False, return_indices=False)
```

takes as input a $N \times C \times H \times W$ tensor, and a kernel size (h, w) or kinterpreted as (k, k), applies the max-pooling on each channel of each sample separately, and produce if the padding is 0 a $N \times C \times |H/h| \times |W/w|$ output.

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Similar functions implements 1d and 3d max-pooling, and average pooling.



While for convolution the default stride is 1, for pooling it is equal to the kernel size, but this not obligatory.

Default padding is zero.

Wraps the max-pooling operation into a Module.

As for convolutions, the kernel size is either a pair (h, w) or a single value k interpreted as (k, k).

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EE-559 - Deep learning

4.6. Writing a PyTorch module

François Fleuret

https://fleuret.org/ee559/

Mon Feb 18 13:34:37 UTC 2019





We now have all the bricks needed to build our first convolutional network from scratch. The last technical points is the tensor shape between layers.

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Both the convolutional and pooling layers take as input batches of samples, each one being itself a 3d tensor $C \times H \times W$.

The output has the same structure, and tensors have to be explicitly reshaped before being forwarded to a fully connected layer.

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The output has the same structure, and tensors have to be explicitly reshaped before being forwarded to a fully connected layer.

```
>>> from torchvision.datasets import MNIST
>>> mnist = MNIST('./data/mnist/', train = True, download = True)
>>> d = mnist.train_data
>>> d.size()
torch.Size([60000, 28, 28])
>>> x = d.view(d.size(0), 1, d.size(1), d.size(2))
>>> x.size()
torch.Size([60000, 1, 28, 28])
>>> x = x.view(x.size(0), -1)
>>> x.size()
torch.Size([60000, 784])
```

Input sizes / operations	Nb. parameters	Nb. products
1 × 28 × 28		

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1 × 28 × 28		
nn.Conv2d(1, 32, kernel_size=5)	$32 \times (5^2 + 1) = 832$	$32 \times 24^2 \times 5^2 = 460,800$
$32 \times 24 \times 24$		

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F.max_pool2d(., kernel_size=3)	0	0
32 × 8 × 8		

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$32 \times 24 \times 24$		
F.max_pool2d(., kernel_size=3)	0	0
$32 \times 8 \times 8$		
F.relu(.)	0	0
32×8×8		
nn.Conv2d(32, 64, kernel_size=5)	$64 \times (32 \times 5^2 + 1) = 51,264$	$32 \times 64 \times 4^2 \times 5^2 = 819,200$
$64 \times 4 \times 4$		

Input sizes / operations	Nb. parameters	Nb. products
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$32 \times 24 \times 24$		
<pre>F.max_pool2d(., kernel_size=3)</pre>	0	0
$32 \times 8 \times 8$		
F.relu(.)	0	0
$32 \times 8 \times 8$		
nn.Conv2d(32, 64, kernel_size=5)	$64 \times (32 \times 5^2 + 1) = 51,264$	$32 \times 64 \times 4^2 \times 5^2 = 819,200$
$64 \times 4 \times 4$		
<pre>F.max_pool2d(., kernel_size=2)</pre>	0	0
$64 \times 2 \times 2$		

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nn.Conv2d(1, 32, kernel_size=5)	$32 \times (5^2 + 1) = 832$	$32 \times 24^2 \times 5^2 = 460,800$
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F.max_pool2d(., kernel_size=3)	0	0
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$64 \times 4 \times 4$		
F.max_pool2d(., kernel_size=2)	0	0
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F.relu(.)	0	0
$64 \times 2 \times 2$		
x.view(-1, 256)	0	0
256		

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$64 \times 4 \times 4$		
F.max_pool2d(., kernel_size=2)	0	0
$64 \times 2 \times 2$		
F.relu(.)	0	0
$64 \times 2 \times 2$		
x.view(-1, 256)	0	0
256		
nn.Linear(256, 200)	$200 \times (256 + 1) = 51,400$	$200 \times 256 = 51,200$
200		

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x.view(-1, 256)	0	0
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$64 \times 4 \times 4$		
F.max_pool2d(., kernel_size=2)	0	0
$64 \times 2 \times 2$		
F.relu(.)	0	0
$64 \times 2 \times 2$		
x.view(-1, 256)	0	0
256		
nn.Linear(256, 200)	$200 \times (256 + 1) = 51,400$	$200 \times 256 = 51,200$
200		
F.relu(.)	0	0
200		
nn.Linear(200, 10)	$10 \times (200 + 1) = 2,010$	$10 \times 200 = 2,000$
10		

Input sizes / operations	Nb. parameters	Nb. products
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nn.Conv2d(1, 32, kernel_size=5)	$32 \times (5^2 + 1) = 832$	$32 \times 24^2 \times 5^2 = 460,800$
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x.view(-1, 256)	0	0
256		
nn.Linear(256, 200)	$200 \times (256 + 1) = 51,400$	$200 \times 256 = 51,200$
200		
F.relu(.)	0	0
200		
nn.Linear(200, 10)	$10 \times (200 + 1) = 2,010$	$10 \times 200 = 2,000$
10		

Total 105,506 parameters and 1,333,200 products for the forward pass.

Creating a module

PyTorch offers a sequential container module torch.nn.Sequential to build simple architectures.

For instance a MLP with a 10 dimension input, 2 dimension output, ReLU activation function and two hidden layers of dimensions 100 and 50 can be written as:

```
model = nn.Sequential(
    nn.Linear(10, 100), nn.ReLU(),
    nn.Linear(100, 50), nn.ReLU(),
    nn.Linear(50, 2)
);
```

However for any model of practical complexity, the best is to write a sub-class of torch.nn.Module.

To create a Module, one has to inherit from the base class and implement the constructor $_$ init $_$ (self, ...) and the forward pass forward(self, x).

To create a Module, one has to inherit from the base class and implement the constructor __init__(self, ...) and the forward pass forward(self, x).

```
class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
        self.conv1 = nn.Conv2d(1, 32, kernel_size=5)
        self.conv2 = nn.Conv2d(32, 64, kernel_size=5)
        self.fc1 = nn.Linear(256, 200)
        self.fc2 = nn.Linear(200, 10)

    def forward(self, x):
        x = F.relu(F.max_pool2d(self.conv1(x), kernel_size=3, stride=3))
        x = F.relu(F.max_pool2d(self.conv2(x), kernel_size=2, stride=2))
        x = x.view(-1, 256)
        x = F.relu(self.fc1(x))
        x = self.fc2(x)
        return x
```

Inheriting from torch.nn.Module provides many mechanisms implemented in the superclass.

First, the (...) operator is redefined to call the forward(...) method and run additional operations. The forward pass should be executed through this operator and not by calling forward explicitly.

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First, the (...) operator is redefined to call the forward(...) method and run additional operations. The forward pass should be executed through this operator and not by calling forward explicitly.

Using the class Net we just defined

```
model = Net()
input = torch.empty(12, 1, 28, 28).normal_()
output = model(input)
print(output.size())
...
```

prints

```
torch.Size([12, 10])
```

Also, all Parameters added as class attributes are seen by Module.parameters().

```
class Net(nn.Module):
    def __init__(self):
        super(Net, self).__init__()
        self.conv1 = nn.Conv2d(1, 32, kernel_size=5)
        self.conv2 = nn.Conv2d(32, 64, kernel_size=5)
        self.fc1 = nn.Linear(256, 200)
        self.fc2 = nn.Linear(200, 10)
/.../
model = Net()

for k in model.parameters():
    print(k.size())
```

Also, all Parameters added as class attributes are seen by Module.parameters().

```
class Net(nn.Module):
    def init (self):
        super(Net, self).__init__()
        self.conv1 = nn.Conv2d(1, 32, kernel_size=5)
        self.conv2 = nn.Conv2d(32, 64, kernel size=5)
        self.fc1 = nn.Linear(256, 200)
        self.fc2 = nn.Linear(200, 10)
1.../
model = Net()
for k in model.parameters():
    print(k.size())
prints
torch.Size([32, 1, 5, 5])
torch.Size([32])
torch.Size([64, 32, 5, 5])
torch.Size([64])
torch.Size([200, 256])
torch.Size([200])
torch.Size([10, 200])
torch.Size([10])
```



Parameters added in dictionaries or arrays are not seen.

```
class Buggy(nn.Module):
    def __init__(self):
        super(Buggy, self).__init__()
        self.conv = nn.Conv2d(1, 32, kernel_size=5)
        self.param = Parameter(torch.zeros(123, 456))
        self.other_stuff = [ nn.Linear(543, 21) ]

model = Buggy()

for k in model.parameters():
    print(k.size())
```



Parameters added in dictionaries or arrays are not seen.

```
class Buggy(nn.Module):
    def init (self):
        super(Buggy, self).__init__()
        self.conv = nn.Conv2d(1, 32, kernel_size=5)
        self.param = Parameter(torch.zeros(123, 456))
        self.other_stuff = [ nn.Linear(543, 21) ]
model = Buggv()
for k in model.parameters():
    print(k.size())
prints
torch.Size([123, 456])
torch.Size([32, 1, 5, 5])
torch.Size([32])
```

A simple option is to add modules in a torch.nn.ModuleList, which is a list of modules properly dealt with by PyTorch's machinery.

```
class AnotherNotBuggy(nn.Module):
    def __init__(self):
        super(AnotherNotBuggy, self).__init__()
        self.conv = nn.Conv2d(1, 32, kernel_size=5)
        self.param = Parameter(torch.zeros(123, 456))
        self.other_stuff = nn.ModuleList()
        self.other_stuff.append(nn.Linear(543, 21))
model = AnotherNotBuggy()
for k in model.parameters():
    print(k.size())
prints
torch.Size([123, 456])
torch.Size([32, 1, 5, 5])
torch.Size([32])
torch.Size([21, 543])
torch.Size([21])
```

As long as you use autograd-compliant operations, the backward pass is implemented automatically.

This is crucial to allow the optimization of the Parameters with gradient descent.

