# EE-559 - Deep learning

# 4a. DAG networks, autograd, convolution layers

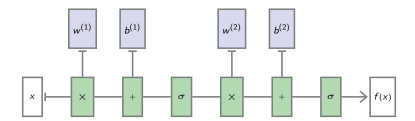
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https://fleuret.org/dlc/
[version of: March 10, 2018]



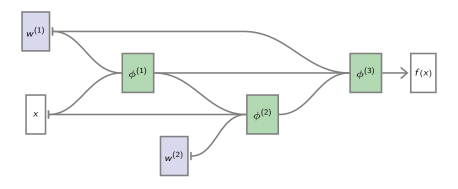


DAG networks

### Everything we have seen for an MLP



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



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Remember that we use tensorial notation.

If  $(a_1, ..., a_Q) = \phi(b_1, ..., b_R)$ , we have

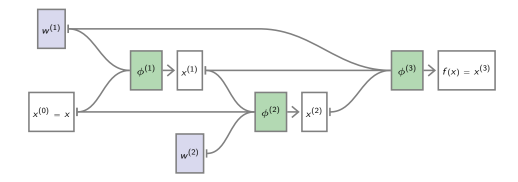
$$\left[rac{\partial {\sf a}}{\partial b}
ight] = J_\phi = \left(egin{array}{ccc} rac{\partial {\sf a}_1}{\partial b_1} & \cdots & rac{\partial {\sf a}_1}{\partial b_R} \ dots & \ddots & dots \ rac{\partial {\sf a}_Q}{\partial b_1} & \cdots & rac{\partial {\sf a}_Q}{\partial b_R} \end{array}
ight).$$

This notation does not specify at which point this is computed. 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

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

### Forward pass



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

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

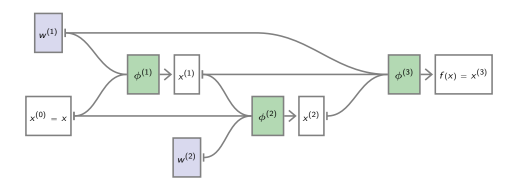
$$x^{(2)} = \phi^{(2)}(x^{(0)}, x^{(1)}; w^{(2)})$$

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

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### Backward pass, derivatives w.r.t activations

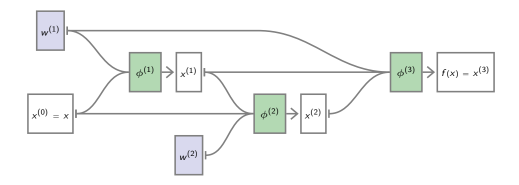


$$\begin{split} & \left[\frac{\partial \ell}{\partial x^{(2)}}\right] = \left[\frac{\partial x^{(3)}}{\partial x^{(2)}}\right] \left[\frac{\partial \ell}{\partial x^{(3)}}\right] = J_{\phi^{(3)}|x^{(2)}} \left[\frac{\partial \ell}{\partial x^{(3)}}\right] \\ & \left[\frac{\partial \ell}{\partial x^{(1)}}\right] = \left[\frac{\partial x^{(2)}}{\partial x^{(1)}}\right] \left[\frac{\partial \ell}{\partial x^{(2)}}\right] + \left[\frac{\partial x^{(3)}}{\partial x^{(1)}}\right] \left[\frac{\partial \ell}{\partial x^{(3)}}\right] = J_{\phi^{(2)}|x^{(1)}} \left[\frac{\partial \ell}{\partial x^{(2)}}\right] + J_{\phi^{(3)}|x^{(1)}} \left[\frac{\partial \ell}{\partial x^{(3)}}\right] \\ & \left[\frac{\partial \ell}{\partial x^{(0)}}\right] = \left[\frac{\partial x^{(1)}}{\partial x^{(0)}}\right] \left[\frac{\partial \ell}{\partial x^{(1)}}\right] + \left[\frac{\partial x^{(2)}}{\partial x^{(0)}}\right] \left[\frac{\partial \ell}{\partial x^{(2)}}\right] = J_{\phi^{(1)}|x^{(0)}} \left[\frac{\partial \ell}{\partial x^{(1)}}\right] + J_{\phi^{(2)}|x^{(0)}} \left[\frac{\partial \ell}{\partial x^{(2)}}\right] \end{split}$$

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### Backward pass, derivatives w.r.t parameters



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

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

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

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

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

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

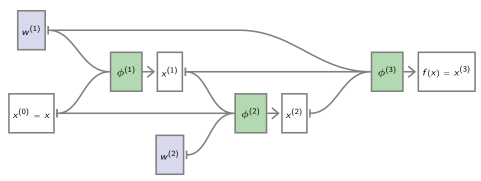
One approach is to define the nodes and edges of such a DAG statically (Torch, TensorFlow, Caffe, Theano, etc.)

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## For instance, in TensorFlow, to run a forward/backward pass on



with

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

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

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

#### we can do

```
w1 = tf.Variable(tf.random_normal([5, 5]))
w2 = tf.Variable(tf.random_normal([5, 5]))
x = tf.Variable(tf.random_normal([5, 5]))
x0 = x
x1 = tf.matmul(w1, x0)
x2 = x0 + tf.matmul(w2, x1)
x3 = tf.matmul(w1, x1 + x2)
q = tf.norm(x3)
gw1, gw2 = tf.gradients(q, [w1, w2])
with tf.Session() as sess:
    sess.run(tf.global_variables_initializer())
    _grads = sess.run(grads)
```

## Autograd

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The forward pass is "just" a computation as usual. The graph structure is needed for the backward pass only.

The specification of the graph looks a lot like the forward pass, and the operations of the forward pass fully define those of the backward.

PyTorch provides Variable s, which can be used as Tensor s, with the advantage that during any computation, the graph of operations to computes the gradient wrt any quantity is automatically constructed.

This "autograd" mechanism has two main benefits:

- Simpler syntax: one just need to write the forward pass as a standard computation,
- greater flexibility: Since the graph is not static, the forward pass can be dynamically modulated.

To use autograd, use torch.autograd.Variable instead of torch.Tensor. Most of the Tensor operations [have corresponding operations that] accept Variable.

A Variable is first a wrapper around a Tensor. It has the following fields

- data is the Tensor containing the data itself,
- grad is a Variable of same dimension to sum the gradient,
- requires\_grad is a Boolean stating if we need the gradient w.r.t this Variable (default is False).

A Parameter is a Variable with requires\_grad to True by default, and known to be a parameter by various utility functions.



A Variable can only embed a Tensor, so functions returning a scalar (e.g. a loss) now return a 1d Variable with a single value.

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torch.autograd.grad(outputs, inputs) computes and returns the sum of gradients of outputs wrt the specified inputs. This is always a tuple of Variable.

An alternative is to use torch.autograd.backward(variables) or Variable.backward(), which accumulates the gradients in the grad fields of the leaf Variable s.

Consider a simple example  $(x_1, x_2, x_3) = (1, 2, 2)$ , and

$$\ell = \|x\| = \sqrt{x_1^2 + x_2^2 + x_3^2}.$$

We have  $\ell = 3$  and

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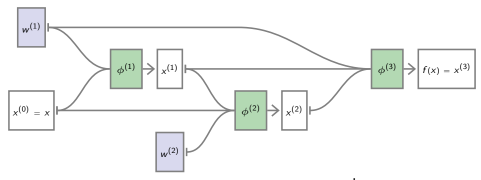
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Alternatively, Variable.backward() accumulates the gradient in the variable's grad fields.

```
>>> from torch import Tensor
>>> from torch.autograd import Variable
>>> x = Variable(Tensor([1, 2, 2]), requires_grad = True)
>>> 1 = x.norm()
>>> 1
Variable containing:
3
[torch.FloatTensor of size 1]
>>> 1.backward()
>>> x.grad
Variable containing:
0.3333
0.6667
0.6667
[torch.FloatTensor of size 3]
```

## For instance, in PyTorch, to run a forward/backward pass on



with

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

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

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

we can do

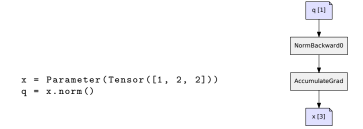
```
w1 = Parameter(Tensor(5, 5).normal_())
w2 = Parameter(Tensor(5, 5).normal_())
x = Variable(Tensor(5).normal_())
x0 = x
x1 = w1.mv(x0)
x2 = x0 + w2.mv(x1)
x3 = w1.mv(x1 + x2)
q = x3.norm()
q.backward()
```

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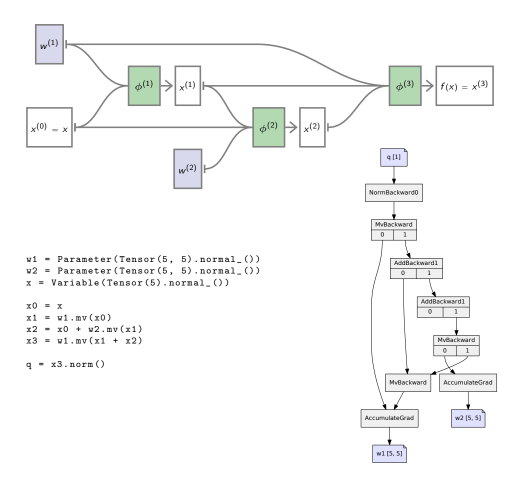
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We can look precisely at the graph built during a computation.



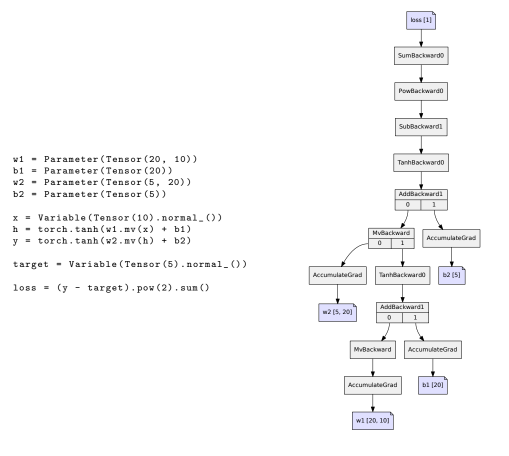
This graph was generated with

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

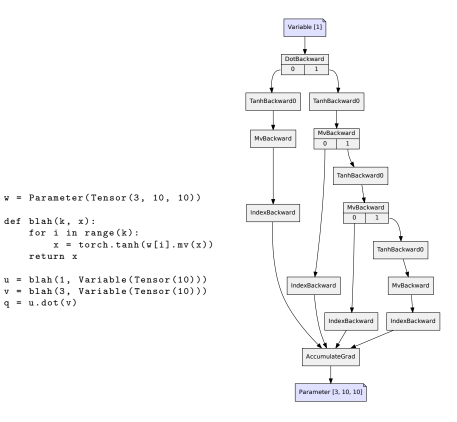


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Variable.backward() accumulates the gradients in the different Variable s, so one may have to zero them before.

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.

```
>>> x = Variable(Tensor([3, 4]), requires_grad = True)
>>> a = x.norm()
>>> a.backward()
>>> b = x.dot(Variable(Tensor([-10, 10])))
>>> b.backward()
>>> x.grad
Variable containing:
 -9.4000
10.8000
[torch.FloatTensor of size 2]
>>> x = Variable(Tensor([3, 4]), requires_grad = True)
>>> q = x.norm() + x.dot(Variable(Tensor([-10, 10])))
>>> q.backward()
>>> x.grad
Variable containing:
-9.4000
 10.8000
[torch.FloatTensor of size 2]
```

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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.

Finally, since the gradient itself is a Variable, autograd can generate the computational graph for computing higher-order derivatives.

This is done by passing create\_graph=True to torch.autograd.grad(...)

```
>>> x = Variable(Tensor([ 1, 2, 3 ]), requires_grad = True)
>>> s1 = x.pow(2).sum()
>>> g1, = torch.autograd.grad(s1, x, create_graph = True)
>>> g1
Variable containing:
2
4
6
[torch.FloatTensor of size 3]

>>> s2 = g1[0].exp() - g1[2].exp()
>>> g2

Variable containing:
14.7781
0.0000
-806.8576
[torch.FloatTensor of size 3]
```

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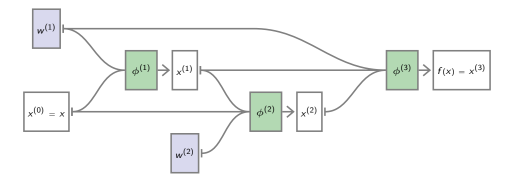
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## Weight sharing

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  $\phi^{(3)}$ .



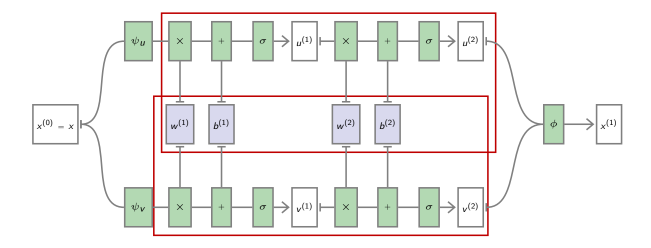
This is called weight sharing.

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Weight sharing allows in particular to build **siamese networks** where a full sub-network is replicated several times.



## Convolutional layers

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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  $256 \times 256$  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 ( $\simeq 150 \text{Gb}$  !), and excess of capacity.

Moreover, this requirement is inconsistent with the intuition that such large signals have some "invariance in translation". A representation meaningful at a certain location can / should be used everywhere.

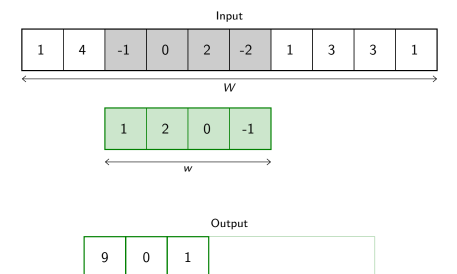
A convolutional layer embodies this idea. It applies the same linear transformation locally, everywhere, and preserves the signal structure.

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W-w+1

Formally, in 1d, given

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

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

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

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

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

 $\Lambda$ 

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

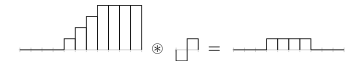
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Convolution can implement a differential operator

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



or a crude "template matcher"



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



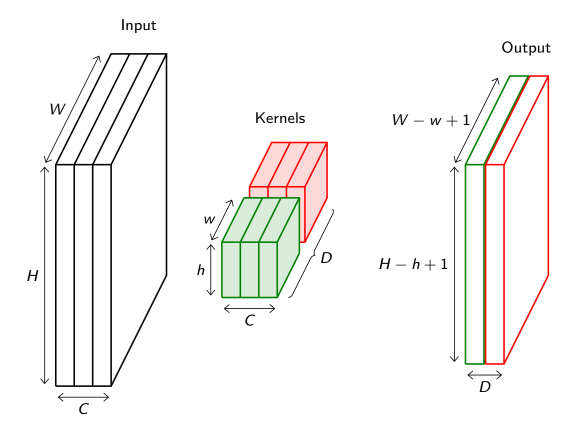
We say "2d signal" even though it has C channels, since it is a feature vector indexed by a 2d location without structure on the feature indexes.

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

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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.

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We usually refer to one of the channels generated by a convolutional 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.

## **Pooling**

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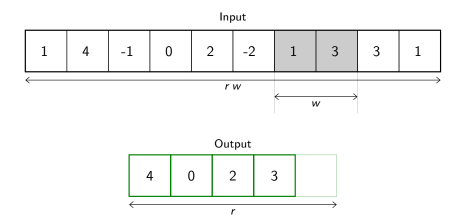
In many cases, a feed-forward network computes a low-dimension signal (e.g. a few scores) from a very high-dimension signal (e.g. an image).

As for convolution, it makes sense to reduce the signal's size in a way that preserves its structure, just "down-scaling it".

This operation is called **pooling**, and aims at grouping several activations into a single "more meaningful" one.

The most standard type of pooling is the **max-pooling**, which computes max values over non-overlapping blocks.

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

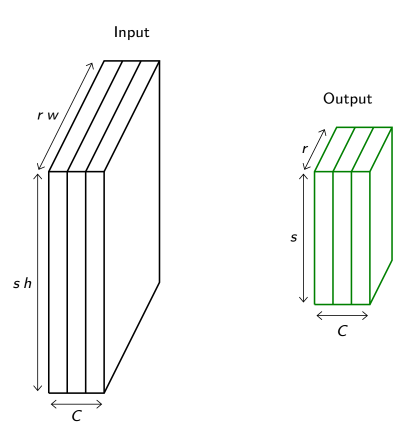


The average pooling computes average values per block instead of max values.

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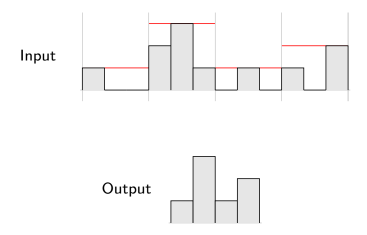
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Pooling provides invariance to any permutation inside one of the cell.

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



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