Convolutional Layers: Space and Time

In our introductory examples

- ullet The spatial dimension of output $\mathbf{y}_{(l)}$
- ullet Is identical to the spatial dimension of input $\mathbf{y}_{(l-1)}$

There are different choices we can make when "sliding" the kernel over the input.

These choices impact

- The spatial dimension of the output
- And, in turn, the time requirements of subsequent layers (because of the size)

Let's do some quick calculations and then show choices for controlling the space consumed by $y_{\left(l\right)}.$

CNN Math: Time versus number of parameters

Consider input layer (l-1) with

- \bullet N spatial dimensions
- ullet $n_{(l-1)}$ feature maps/channels

$$||\mathbf{y}_{(l-1)}|| = (d_{(l-1),1} imes d_{(l-1),2} imes \ldots d_{(l-1),N} imes n_{(l-1)})$$

Layer l will apply a Convolution that preserves the spatial dimensions

$$||\mathbf{y}_{(l)}|| = (d_{(l-1),1} imes d_{(l-1),2} imes \ldots d_{(l-1),N} imes n_{(l)})$$

For simplicity of presentation: consider the case when N=2.

How many weights/parameters does layer l consume (i.e, what is size of $\mathbf{W}_{(l)}$) ?

- Each kernel $\mathbf{k}_{(l),j}$
 - lacktriangle Has spatial dimension $(f_{(l)} imes f_{(l)})$
 - $\ \ \, \ \ \,$ And "depth" $n_{(l-1)}$ (to match the number of input feature maps/channels)
- There are $n_{(l)}$ kernels in layer l

So the size of $W_{\left(l
ight)}$ (ignoring the optional bias term per output feature map)

$$||\mathbf{W}_{(l)}|| = n_{(l)} * (n_{(l-1)} * f_{(l)} * f_{(l)})$$

The part of the product that most concerns us is ($n_{(l)} st n_{(l-1)}$)

- $\bullet \;\; \mbox{Values for} \; n_{(l)}, n_{(l-1)} \; \mbox{in} \; \{32, 64, 256\} \; \mbox{are not uncommon} \; !$
- ullet Hence $||\mathbf{W}_{(l)}||$ is often easily several thousand
- State of the art image recognition models use several hundred million weights!

How many multiplications (in the dot product) are required for layer l?

- We will ignore additions (the part of the dot product that reduces pair-wise products to a scalar, and for the bias)
- Each kernel $\mathbf{k}_{(l),j}$ of dimension

$$(f_{(l)} imes f_{(l)} imes n_{(l-1)})$$

- ullet Applied over each location in the $(d_{(l-1),1} imes d_{(l-1),2})$ spatial dimension of the input layer (l-1)
- There are $n_{(l)}$ kernels in layer l

So the number of multiplications

$$n_{(l)}*(d_{(l-1),1}*d_{(l-1),2})*(n_{(l-1)}*f_{(l)}*f_{(l)})$$

Consider a grey-scale image of size $(d_{(l-1),1}*d_{(l-1),2})=(1024 imes1024)$

- Lower than your cell-phones camera!
- Easily several million multiplications

Expect the time to train a Neural Network with Convolutional layers to be long!

- That's why GPU's are important in training
- But GPU's have limited memory so space is important too
 - Can control with batch size

All of this ignores the final layer ${\cal L}$

- Often a Fully Connected layer implementing Regression or Classification
- With n_L output features
 - ullet e.g., For Classification over classes in set $C, \mathbf{y}_{(L)}$ is a One Hot Vector of length $n_L = ||C||$

Suppose layer
$$(L-1)$$
 has dimension $||\mathbf{y}_{(L-1)}||=(d_{(L-1),1} imes d_{(L-1),2} imes n_{(L-1)})$

Before we can use it as input to the Fully Connected Layer ${\cal L}$ we flatten it to a vector of length

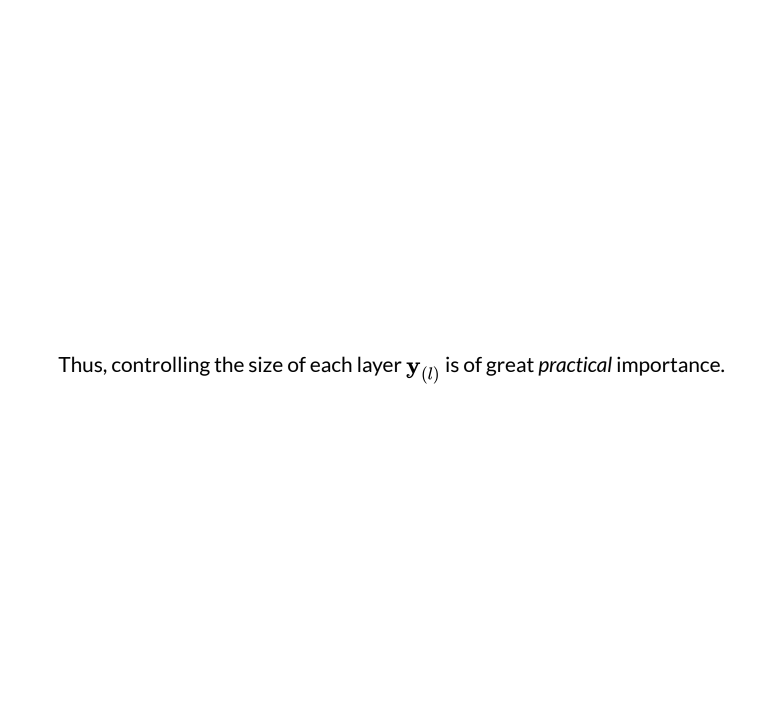
$$(d_{(L-1),1}*d_{(L-1),2}*n_{(L-1)})$$

The number of weights (ignoring biases) and multiplications is

$$||W_L|| = n_{(L)} * (d_{(L-1),1} * d_{(L-1),2} * n_{(L-1)})$$

- ullet $n_{(L)}*n_{(L-1)}$ on the order of several thousand
- $(d_{(L-1),1} * d_{(L-1),2})$ on the order of several million, for images

This may not even be feasible!



Controlling the output spatial dimensions

Padding

In our examples thus far

- When a location in the spatial dimensions of the input
- Is such that, when the kernel is placed there, it extends beyond the input
- We have added "padding"

This is not strictly necessary

- But has advantage that the spatial dimension of output $\mathbf{y}_{(l)}$ is the same as the input $\mathbf{y}_{(l-1)}$
- One can simply *not* produce an output for such locations
- ullet It just means the output spatial dimension shrinks in each dimension by $f_{(l)}-1$
 - Assuming $f_{(l)}$ is odd
 - The number of locations in which the kernel extends over the border
 - lacksquare Is Half of the filter size $(f_{(l)}-1)/2$ times two (for each edge)

Stride

Thus far, we have placed the kernel over *each* location in the spatial dimension of the input layer.

This, along with padding, ensures that the spatial dimension of the input and output layers are identical.

Consider two adjacent locations in the spatial dimension of the input layer

• The values of the input layer that appear in each dot product overlap

 By placing the kernel over <i>every other</i> location of the spatial dimension of the input layer We may still be able to recognize features And reduce the spatial dimension of the output layer by a factor of 2 for each dimension.

In general, we can choose to choose to pass over (S-1) locations in the spatial dimension of the input layer

- ullet S is called the *stride*
- ullet Up until now: S=1
- But you are free to choose

Size of output

We can combine choices of Padding and Stride to control the spatial dimension of the output layer l:

Let

- ullet $d_{(l-1),j}$ denote the number of elements in spatial dimension j of layer (l-1)
- ullet denote the number of elements added as padding on each border
- \bullet S denote the stride
- ullet $f_{(l)}$ be the size of the filter (for each spatial dimension)

Then the number of elements in spatial dimension
$$j$$
 of output layer (l) is $d_{(l),j}=rac{d_{(l-1),j}+2P-f_{(l)}}{S}+1$

You can see that increasing the stride has the biggest impact on reducing the spatial dimension of the output.

Pooling layer

There is a layer type with the specific purpose of changing the spatial dimension of the output.

This is called a Pooling Layer.

A Pooling Layer combines the information from adjacent locations in the spatial dimension of the input layer.

- The "combining" operation may be average or maximum
- Sacrificing the exact location in the spatial dimension
- Often in exchange for reduced space

A Pooling Layer is similar in some respects to a Convolution.

Recall that the One Dimensional Convolutional Layer (Conv1d) with a single input feature computes the following for output feature/channel j:

$$\mathbf{y}_{(l),j} = egin{pmatrix} a_{(l)} \left(\ N(\mathbf{y}_{(l-1)}, \mathbf{W}_{(l),j}, 1) \cdot \mathbf{W}_{(l)} \ a_{(l)} \left(\ N(\mathbf{y}_{(l-1)}, \mathbf{W}_{(l),j}, 2) \cdot \mathbf{W}_{(l)} \ \end{pmatrix} \ & \vdots \ a_{(l)} \left(\ N(\mathbf{y}_{(l-1)}, \mathbf{W}_{(l),j}, n_{(l-1)} \cdot \mathbf{W}_{(l)} \ \end{pmatrix} \end{pmatrix}$$

The analogous One Dimensional Pooling Layer (Pooling1D) computes

Dimensional Pooling Layer (Pooling1D) computes
$$\mathbf{y}_{(l),j} = egin{pmatrix} p_{(l)} \left(\ N'(\mathbf{y}_{(l-1)},f_{(l)},1) \ p_{(l)} \left(\ N'(\mathbf{y}_{(l-1)},f_{(l)},2) \ dots \ p_{(l)} \left(\ N'(\mathbf{y}_{(l-1)},f_{(l)},n_{(l-1)} \ \end{pmatrix} \end{pmatrix}$$

where
$$N'(|\mathbf{y}_{(l-1)},f_{(l)},j|)$$

- selects a subsequence of $\mathbf{y}_{(l-1)}$ centered at $\mathbf{y}_{(l-1),j}$
- of length $f_{(l)}$

and $p_{\left(l\right)}$ is a pooling operation

That is, similar to a Convolutional Layer, the Pooling Layer

- ullet Selects a region of length $f_{(l)}$
- ullet Centered at each location in the spatial dimension of the input layer (l-1)

and produces a value in the corresponding spatial location of output layer l

• That summarizes the selected region

Observe that

- There are *no* weights
- No dot product
- Just a pooling operation

Similar to Convolution, we can extend pooling to higher spatial dimension (N>1) and higher number of input channels $n_{(l-1)}>1$.

Suppose the input $\mathbf{y}_{(l-1)}$ is (N+1) dimensional of shape

$$||\mathbf{y}_{(l-1)}|| = (d_{(l-1),1} imes d_{(l-1),2} imes \ldots d_{(l-1),N} imes n_{(l-1)})$$

Pooling:

- ullet Selects an N-dimensional region, where each dimension is of length $f_{(l)}$
- Centered at each location in the spatial dimension
 - lacksquare Of a single feature map $m{j}$ of the input layer (l-1): $\mathbf{y}_{(l-1),\ldots,j}$

and produces a value in the corresponding spatial location of output layer \boldsymbol{l}

ullet That $\mathit{summarizes}$ the selected region by applying $p_{(l)}$ to the selected region

Pooling with a stride S>1

- "Down samples" the spatial dimension
- Sacrificing some information about locality

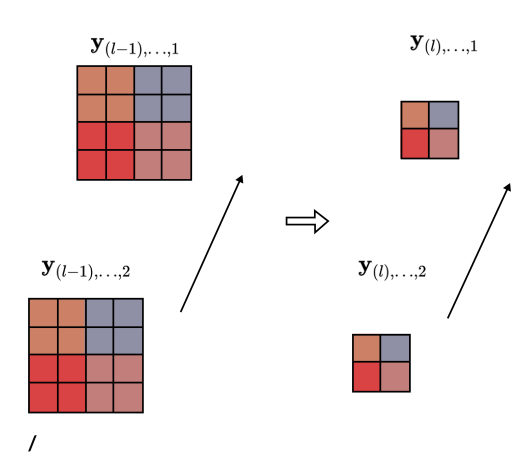
It effectively asks the question

• Does the feature exist in a broader neighborhood of the spatial dimension

Here is a two dimensional example with a filter size and stride of 2:

- \bullet N=2
- $ullet f_{(l)}=2 \ ullet S=2$

Conv 2D: Pooling (Max/Average)



The key difference between Pooling and Convolution (other than the absence of the dot product and kernel weights)

- The pooling operation is applied to each input feature map *separately*
- Versus all the input feature maps at a given location in the spatial dimension of the input

Pooling operations

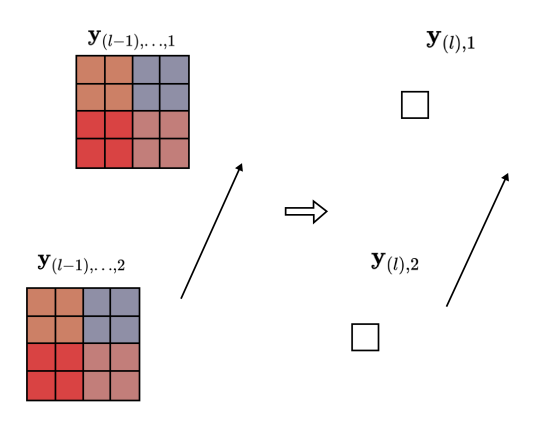
- Max pooling
 - Maximum over the selected region
 - Good for answering the question: "Does the feature exist" in the neighborhood
- Average pooling
 - average over the selected region
 - "blurs" the location in the spatial dimension when it is unimportant or highly variable

Global Pooling

Each feature map $m{j}$ of the input layer $(\mathbf{y}_{(l-1),\ldots,j})$

• Is summarized by a single value produced by Max Pooling operation $p'_{(l)}$ $\mathbf{y}_{(l),j}=p'_{(l)}(\mathbf{y}_{(l-1),\ldots,j})$

Conv 2D: Global Pooling (Max/Average)



Notice that each input feature map has been reduced to a single value in the output.

• No spatial dimension in $\mathbf{y}_{(l)}$ (hence no "...")

The Global Pooling operation effectively asks the question

- Does the feature occur anywhere in the feature map?
- Losing information about the exact location in the spatial dimensions

Global pooling operations

- Global average pooling
 - Maximum over the feature map
- K-Max pooling
 - lacktriangledown replace one dimension of the volume with the $m{K}$ largest elements of the dimension

Review

Let's summarize our knowledge of controlling the size of $y_{(l-1)}$:

- Controlling spatial dimensions
 - Increase stride
 - Pooling
 - Global average pooling often used in final Convolutional Layer
- Control number of feature maps per layer
 - Choice of $n_{(l),1}$
 - Kernel size $f_{(l)}=1$
 - preserve spatial dimension
 - $\circ~$ change number of feature maps from $oldsymbol{n}_{(l-1),1}$ to $oldsymbol{n}_{(l),1}$

Striding and Pooling

- increase receptive field
- typically small values (e.g., S=2)
 - limited reduction

Kernel size
$$\emph{f}_{(l)}=1$$

- ullet reduction depends on the ratio of $oldsymbol{n}_{(l),1}$ to $oldsymbol{n}_{(l-1),1}$
 - unlimited reduction possible

Kernel size 1

A less obvious way to control the size of $\mathbf{y}_{(l)}$ is to use a kernel with $f_{(l)}=1$

Why might that be?

Recall that a Convolutional Layer

- Preserves the spatial dimension
- Replaces the channel/feature dimension (number of feature maps)

That is\

$$||\mathbf{y}_{(l-1)}|| = (n_{(l-1),1} \times n_{(l-1),2} \times \dots n_{(l-1),N}, \mathbf{n}_{(l-1)})$$

 $||\mathbf{y}_{(l)}|| = (n_{(l-1),1} \times n_{(l-1),2} \times \dots n_{(l-1),N}, \mathbf{n}_{(l)})$

So a kernel of size $oldsymbol{f}_{(l)}=\mathbf{1}$ in all N spatial dimensions

- With "depth" $oldsymbol{n}_{(l-1)}$
- Is just a way to resize $\mathbf{y}_{(l-1)}$ from $n_{(l-1)}$ feature maps to a *single* feature map
 - That sums, across feature maps, the elements in each feature map at the same spatial location

In other words:

• Yet another way to reduce the size of $y_{(l)}$.

Receptive field

The filter size $m{f}_{(l)}$ also plays a role in the space and time requirements of a Convolutional Layer.

It turns out that

- We can achieve the effect of a large $m{f}_{(l)}$
- With a smaller $oldsymbol{f}_{(l)}$ in conjunction with more Convolutional Layers

The receptive field of a layer $m{l}$ feature at a single spatial location in feature map $m{k}$

• are the spatial locations of Layer 0 (input) features that affect this single feature

For ease of notation:

- ullet we assume N=2 as the dimension of the kernel
- ullet we assume that all N dimensions of the kernel are the same ($f_{(l)}$)

We can determine spatial locations of the layer $oldsymbol{0}$ features influencing a single layer $oldsymbol{l}$ location by working backwards from layer $m{l}$ • As we will demonstrate shortly

So we will assume without loss of generality that

- the "height" and "width" of a single kernel kernel is $(f \times f)$ the full dimensionality of a single layer l kernel is $(f \times f \times n_{(l-1)})$

Increasing the Receptive Field

There are several ways to "widen" the receptive field

- Increasing $f_{(l)}$, the size of the kernel
- Stacking Convolutional Layers
- Stride
- Pooling

Striding and Pooling also have the effect of reducing the size of the output feature map.

Increase the size of the kernel

Although this is the most *obvious* way of increasing the receptive field, we tend to avoid it!

We will see that adding layers is a more efficient way of achieving a bigger receptive field.

Stacking Convolutional Layers

Let's introduce the idea of stacking multiple Convolutional Layers using N=1.

Consider

- N = 1
- $f_{(l)} = f_{(l-1)} = 3$
- (L-1)=2: Two Convolutional Layers

Conv 1D Receptive field: 2 layers

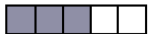
 $\mathbf{y}_{(l-1)}$

 $\mathbf{y}_{(l)}$

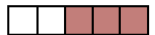
 $\mathbf{y}_{(l+1)}$

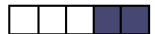












The elements in $\mathbf{y}_{(l)}$

- Are colored
- ullet The same color as the elements of $\mathbf{y}_{(l-1)}$ that they depend on

Each element of layer l depends on $f_{(l)}=3$ elements of layer (l-1).

Consider the element in the center of the second layer: $m{y}_{(l+1)}$, i.e., $m{y}_{(l+1),3}$

- ullet It depends on the Red, Light Blue and Pink elements of $\mathbf{y}_{(l)}$
- ullet Which in turn depend on the Red, Light Blue and Pink elements of $\mathbf{y}_{(l-1)}$
- ullet This includes ullet elements of $\mathbf{y}_{(l-1)}$

So the Two layer network with $oldsymbol{f}_{(l)} = oldsymbol{f}_{(l-1)} = oldsymbol{3}$

- ullet Is exposed to the same layer (l-1) elements
- ullet As a Single Convolutional Layer with $f_{(l)}=f_{(l-1)}=5$

One can trace an element in layer l+1

- Backwards through layers
- To input layer 0

in order to determine the receptive field of layer l+1/2

Bottom line: The size of the receptive field increases with the depth of the layer.

One can apply similar logic to N=2 spatial dimensions.

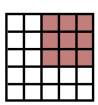
As you go one layer deeper in the NN, the receptive field width and height increase by (2 stride*)

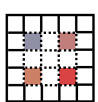
Conv 2D Receptive field: 2 layers

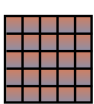


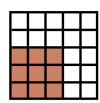
 $\mathbf{y}_{(l)}$

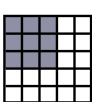
$$\mathbf{y}_{(l+1)}$$













Receptive field

- ullet The spatial locations in layer $oldsymbol{l}$
- ullet Are color coded to match the spatial locations in layer (l-1)
- That affect it

So the yellow location in layer $m{l}$ is a function of the yellow locations in layer (l-1)

The central location in layer (l+1)

- Is a function of the spatial locations in layer $m{l}$ that are encircled by the dashed square
- ullet The layer l locations are a function of all the layer (l-1) locations

So the receptive field for the central location in layer (l+1)

• Includes all the locations of layer (l-1)

In other words: the size of the receptive field grows with layer depth.,

Layer Receptive field

 $1|(3 \times 3)2|(5 \times 5)3|(7 \times 7)$:

Let's compare

- ullet The math of 2 layers with $f_{(l)}=f_{(l-1)}=3$
- ullet The math of 1 layer with $f_{(l)}=5$

In terms of number of weights:

• The one layer network uses

$$egin{array}{lll} ||\mathbf{W}|| &=& n_{(l)}*(n_{(l-1)}*f'(l)*f'(l)) \ &=& 25*n_{(l)}*n_{(l-1)} & ext{when } f'(l)=5 \end{array}$$

• The two layer network uses \$\$ \begin{array}[III]\ || \W\|IIp || & = & n\|Ip (n_{(\II-1)}) f\|Ip *f\|Ip) \ || \W{(\II+1)}|| & = & n{(\II+1)} (n_\|Ip f{(\II+1)} *f{(\II+1)}) \ || \W || & = & || \W\|Ip || + || \W{(\II+1)}|| \

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& = & (9 * n_\llp * n_{(\ll-1)}) + 9 *( n_\llp * n_{(\ll+1)}) ) & \text{ when } f_\llp = f_{(\ll+1)} = 3 \\
```

\end{array}\$\$

The two layer network uses fewer weights when

$$9*(n_{(l)}*n_{(l+1)})) < (25-9)*n_{(l)}*n_{(l-1)}$$

This will be the case when the number of feature maps in all layers is roughly the same.

- The advantage of the smaller network increases as $m{f}_{(l)}' m{f}_l$ increases
 - For example: $f'_{(l)} = 7$
 - Versus 3 Convolutional Layers
 - $lacksquare ext{With } f_{(l)} = f_{(l-1)} = f_{(l+1)} = 3$

CNN advantages/disadvantages

Advantages

- Translational invariance
 - feature can be anywhere
- Locality
 - feature depends on nearby features, not the entire set of features
 - reduced number of parameters compared to a Fully Connected layer

Disadvantages

- Output feature map is roughly same size as input
 - lots of computation to compute a single output feature
 - one per feature of input map
 - higher computation cost
 - training and inference
- Translational invariance not always a positive

How many feature maps to use (What value to choose for $oldsymbol{n_{(l)}}$)

Bag of Tricks for Image Classification with CNNs (https://arxiv.org/abs/1812.01187)

Remember that a larger value for $oldsymbol{n}_{(l)}$ will increase space and time requirements.

One rule of thumb:

- For N=2
- With filter size $f_{(l)}$
- $\bullet\,$ The number of elements in the spatial dimension of input $\mathbf{y}_{(l-1)}$ involved in the dot product is

$$e = (n_{(l-1)} * f_{(l)} * f_{(l)})$$

- ullet It may not make sense to create *more* than e output features $n_{(l)}>e$
 - We would generate more features than input elements

Inverting convolution

The typical flow for multiple layers of Convolutions

- Is for the spatial dimension of successive layers to get smaller
- ullet By using stride S>1
- By using Pooling Layers

This brings up the question: Can we invert the process?

 \bullet That is, go from a smaller spatial dimension back to the spatial dimension of input layer 0

The answer is yes.

This process is sometimes called *Deconvolution* or *Transposed Convolution*.

- In a Deeper Dive, we relate Convolution to Matrix Multiplication
- So the inverting matrix's *dimensions* are the transpose of the matrix implementing the convolution

We will revisit this in the lecture addressing "What is a CNN looking for?"

Technical points

Convolution versus Cross Correlation

- math definition of convolution
 - dot product of input and reversed filter
 - we are doing <u>cross correlation</u>
 (<u>https://en.wikipedia.org/wiki/Convolution</u>)

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In [4]: print("Done")
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Done