# SK hynix i-TAP 반도체 Data Scientist를 위한 ML/DL 심화 커리큘럼

1강. Convolutional Neural Networks Ernest K. Ryu (류경석) 2020.11.06



#### 1강. Outline

- Basics of (non-adaptive) SGD
- PyTorch as a GPU-computing numerical library
- Backpropagation
- Multilayer perceptron
- Convolutional neural networks



## Optimization

We consider

$$\underset{\theta \in \mathbb{R}^p}{\text{minimize}} \quad \frac{1}{N} \sum_{i=1}^{N} f_i(\theta)$$

where  $f_1, ..., f_N$  are "differentiable" functions.

In DL, the ReLU activation function  $\sigma(x) = \max(0, x)$  is said to be "differentiable".



### Gradient Descent (GD)

Define 
$$F(\theta) = \frac{1}{N} \sum_{i=1}^{N} f_i(\theta)$$
. Then 
$$\min_{\theta \in \mathbb{R}^p} F(\theta)$$

GD:

$$\theta^{k+1} = \theta^k - \alpha_k \nabla F(\theta^k)$$

where  $\alpha_0, \alpha_1, ... \in \mathbb{R}$  are stepsizes.

Since  $\nabla F = \frac{1}{N} \sum_{i=1}^{N} \nabla f_i$ , can parallelize  $\nabla F(\theta^k)$  computation.



## Why does GD converge?

Taylor expansion of F about  $\theta^k$ :

$$F(\theta) = F(\theta^k) + \nabla F(\theta^k)^{\mathrm{T}} (\theta - \theta^k) + \mathcal{O}\left((\theta - \theta^k)^2\right)$$

Plug in  $\theta^{k+1}$ :

$$F(\theta^{k+1}) = F(\theta^k) - \alpha_k \|\nabla F(\theta^k)\|^2 + \mathcal{O}(\alpha_k^2)$$

 $-\nabla F(\theta^k)$  is steepest descent direction. For small (cautious)  $\alpha_k$ , GD step reduces function value.



### Stochastic Gradient Descent (SGD)

We consider

$$\underset{\theta \in \mathbb{R}^p}{\text{minimize}} \quad \frac{1}{N} \sum_{i=1}^{N} f_i(\theta) = \mathbb{E}_{I \sim \text{Uniform}\{1,\dots,N\}} \left[ f_I(\theta) \right]$$

SGD:

$$i(k) \sim \text{Uniform}\{1, ..., N\}$$
  
 $\theta^{k+1} = \theta^k - \alpha_k \nabla f_{i(k)}(\theta^k)$ 

where i(k) are uniform IID indices.



# Why does SGD converge?

 $\nabla f_{i(k)}(\theta^k)$  is an unbiased estimate of the gradient  $\nabla F(\theta^k)$ 

$$\mathbb{E}_{i(k)} \nabla f_{i(k)}(\theta^k) = \frac{1}{N} \sum_{i=1}^{N} \nabla f_i(\theta^k) = \nabla F(\theta^k)$$

(So  $\nabla f_{i(k)}(\theta^k)$  is a "stochastic gradient" of F at  $\theta^k$ )



# Why does SGD converge?

Plug  $\theta^{k+1}$  into Taylor expansion of F about  $\theta^k$ :

$$F(\theta^{k+1}) = F(\theta^k) - \alpha_k \nabla F(\theta^k)^{\mathrm{T}} \nabla f_{i(k)}(\theta^k) + \mathcal{O}(\alpha_k^2)$$

Expectation on both sides:

$$\mathbb{E}_{k}F(\theta^{k+1}) = F(\theta^{k}) - \alpha_{k} \|\nabla F(\theta^{k})\|^{2} + \mathcal{O}(\alpha_{k}^{2})$$

 $(\mathbb{E}_k \text{ is expectation conditioned on } \theta^k)$ 

 $-\nabla f_{i(k)}(\theta^k)$  is descent direction in expectation. For small (cautious)  $\alpha_k$ , SGD step reduces function value in expactation.



# SGD with general expectation

Consider general expectation

$$\underset{\theta \in \mathbb{R}^p}{\text{minimize}} \quad \mathbb{E}_{\omega} \left[ f_{\omega}(\theta) \right]$$

where  $\omega$  is a random variable. Expectation (not finite sum) appears when you have generative model of  $\omega$ . E.g. GAN.

SGD:

$$\theta^{k+1} = \theta^k - \alpha_k \nabla f_{\omega^k}(\theta^k)$$

where  $\omega^0, \omega^1, ...$  IID random samples of  $\omega$ .



#### Mini-batch SGD

For each k, let i(k, 1), ..., i(k, B) be IID indices.

$$\frac{1}{B} \sum_{j=1}^{B} \nabla f_{i(k,j)}(\theta^k)$$

is also an unbiased estimate of 
$$\nabla F(\theta^k)$$
 since  $\mathbb{E}\frac{1}{B}\sum_{j=1}^{B}\nabla f_{i(k,j)}(\theta^k) = \frac{1}{B}\sum_{j=1}^{B}\nabla F(\theta^k) = \nabla F(\theta^k)$ 



#### Mini-batch SGD

Mini-batch SGD:

$$g = 0$$
For  $j = 1, ..., B$ 

$$i(k, j) \sim \text{Uniform}\{1, ..., N\}$$

$$g = g + \frac{1}{B} \nabla f_{i(k, j)}$$

$$\theta^{k+1} = \theta^k - \alpha_k g$$

is also an instance of SGD.



#### Mini-batch Size

Mathematically (measuring performance per iteration)

- Use large batch is when noise/randomness is large.
- Use small batch is when noise/randomness is small.

Practically (measuring performance per unit time)

- Large batch allows more efficient communication and computation, up to the GPU memory limit.
- Often best to increase batch size up to the GPU memory limit.



### Cyclic (mini-batch) SGD

Cyclic SGD:

$$\theta^{k+1} = \theta^k - \alpha_k \nabla f_{i(k)}(\theta^k)$$

where i(k) is selected in a cyclic order.

Can also write as:

$$\theta^{k+1} = \theta^k - \alpha_k \nabla f_{\text{mod}(k,N)+1}(\theta^k)$$

Strictly speaking, is not an instance of SGD as unbiased estimation property lost.



# Epoch in Optimization and Training

Epoch: loosely defined as computation time of computing 1 full gradient. One iteration of GD is, by definition, an epoch. *N* iterations of SGD constitute an epoch.

Epoch is a convenient unit for counting iterations (rather than directly counting iteration numbers).



### Cyclic (mini-batch) SGD

#### Cyclic SGD advantage:

- Simpler than SGD
- Uses all datapoints within single epoch.

#### Cyclic SGD disadvantage:

- Worse than SGD in some cases, theoretically and empirically.
- In deep learning, neural nets learn to anticipate cyclic order.



# Shuffled Cyclic (mini-batch) SGD

Shuffled cyclic SGD:

$$\theta^{k+1} = \theta^k - \alpha_k \nabla f_{i(k)}(\theta^k)$$

where i(k) is selected in a cyclic order shuffled every epoch.

Can also write as:

$$\theta^{k+1} = \theta^k - \alpha_k \nabla f_{\sigma^{\left[\frac{k}{N}\right]}(\text{mod}(k,N)+1)}(\theta^k)$$

where  $\sigma^0, \sigma^1, ...$  is a sequence of random permutations.



# Shuffled Cyclic (mini-batch) SGD

Shuffled Cyclic SGD:

```
For e = 1, ..., E //for each epoch \sigma \sim \text{randomPermuation}(N)

For i = 1, ..., N
\theta^{k+1} = \theta^k - \alpha_k \nabla f_{\sigma(k)}(\theta^k)
k = k+1
```



## Shuffled Cyclic (mini-batch) SGD

#### Shuffled cyclic SGD advantage:

- Uses all datapoints within single epoch.
- Network cannot learn to anticipate data order.
- Generally best performance.

#### Shuffled cyclic SGD advantage:

- Not as simple. (But PyTorch makes it simple to use.)
- Theory not as strong as regular SGD.



# PyTorch: GPU Numerical Computing Library

PyTorch is a machine learning library of Python, but PyTorch is fundamentally a numerical computation library.

Features of PyTorch that make itsuitable for using neural networks and machine learning:

- PyTorch supports easy GPU computation.
- Automatic differentiation.
- Numerous ML libraries and sample code.



### GPU Computing Code With CUDA

#### GPU computing opeations:

- cudaMemcpy (CPU→GPU or GPU→CPU)
- CPU code
- GPU kernel calls (CPU instructs GPU to execute computation)

#### GPU computing workflow:

- (i) end data CPU→GPU
- (ii) Compute on GPU
- (iii) Receive result GPU→CPU



#### CPU vs. GPU variables

Variables either reside in CPU or GPU memory.

- CPU variable computation on CPU
- GPU variable computation on GPU

CPU and GPU variables cannot directly interact. Can interact only after CPU→GPU or GPU→CPU transfer.



# PyTorch Demo

Power iteration example on PyTorch

```
send A from host (CPU) to device (GPU)
send x=x0 from host (CPU) to device (GPU)
for _ in range(100):
  tell GPU to compute x=A*x
send x from device (GPU) to host (CPU)
```



# Back Propagation ⊆ Automatic Differentiation

Automates gradient computation! Only need to specify how to evaluate function.

Gradient costs roughly  $5 \times$  computation cost\* of evaluating function.

AutoDiff is not

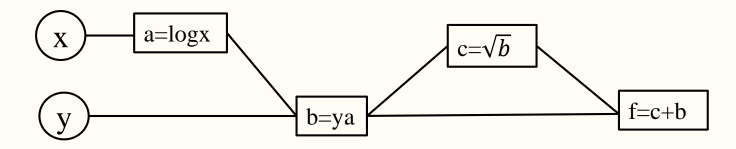
- Finite differencing
- Symbolic differentiation.

AutoDiff ≈ chain rule of vector calculus



#### Chain Rule

- · Consider  $f(x, y) = y \log x + \sqrt{y \log x}$
- $\cdot$  Evaluate f with the computation graph:



· Chain rule:

$$\frac{\partial f}{\partial x} = \frac{\partial f}{\partial c} \frac{\partial c}{\partial b} \left( \frac{\partial b}{\partial a} \frac{\partial a}{\partial x} \frac{\partial x}{\partial x} + \frac{\partial b}{\partial y} \frac{\partial y}{\partial x} \right) + \frac{\partial f}{\partial b} \left( \frac{\partial b}{\partial a} \frac{\partial a}{\partial x} \frac{\partial x}{\partial x} + \frac{\partial b}{\partial y} \frac{\partial y}{\partial x} \right)$$

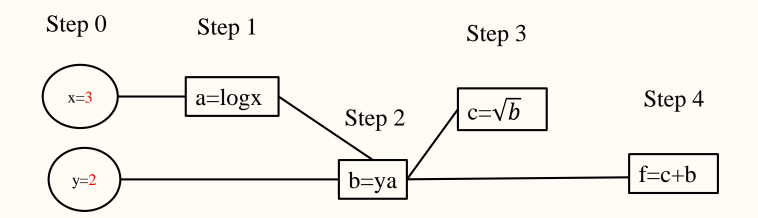
$$\frac{\partial f}{\partial y} = \frac{\partial f}{\partial c} \frac{\partial c}{\partial b} \left( \frac{\partial b}{\partial a} \frac{\partial a}{\partial x} \frac{\partial x}{\partial y} + \frac{\partial b}{\partial y} \frac{\partial y}{\partial x} \right) + \frac{\partial f}{\partial b} \left( \frac{\partial b}{\partial a} \frac{\partial a}{\partial x} \frac{\partial x}{\partial y} + \frac{\partial b}{\partial y} \frac{\partial y}{\partial y} \right)$$

$$\frac{\partial f}{\partial y} = \frac{\partial f}{\partial c} \frac{\partial c}{\partial b} \left( \frac{\partial b}{\partial \alpha} \frac{\partial \alpha}{\partial x} \frac{\partial x}{\partial y} + \frac{\partial b}{\partial y} \frac{\partial y}{\partial x} \right) + \frac{\partial f}{\partial b} \left( \frac{\partial b}{\partial \alpha} \frac{\partial \alpha}{\partial x} \frac{\partial x}{\partial y} + \frac{\partial b}{\partial y} \frac{\partial y}{\partial y} \right)$$

But in what order do you evaluate the chain rule expression?



#### Forward mode auto-diff



0. 
$$x = 3, y = 2, \frac{\partial x}{\partial x} = 1, \frac{\partial x}{\partial y} = 0, \frac{\partial y}{\partial x} = 0, \frac{\partial y}{\partial y} = 1$$

1. 
$$a = log x = log 3$$
,  $\frac{\partial a}{\partial x} = \frac{1}{x} \cdot \frac{\partial x}{\partial x}$ ,  $\frac{\partial a}{\partial y} = 0$ 

2. 
$$b = ya = 2log3$$
,  $\frac{\partial b}{\partial x} = \frac{\partial y}{\partial x}a + y\frac{\partial a}{\partial x} = \frac{2}{3}$ ,  $\frac{\partial b}{\partial y} = \frac{\partial y}{\partial y}a + y\frac{\partial a}{\partial y} = a = log3$  Computation does not involve 'x' or derivatives of 'x'

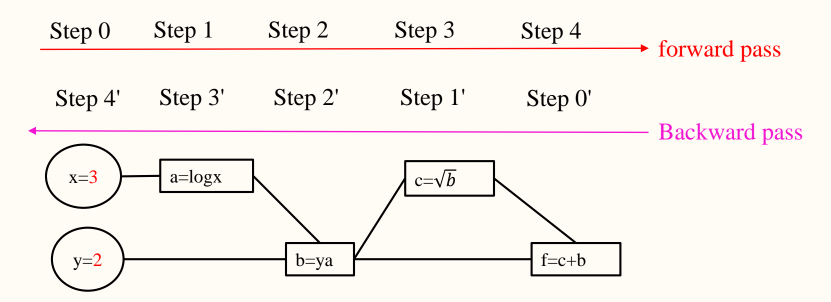
3. 
$$c = \sqrt{b} = \sqrt{2\log 3}$$
,  $\frac{\partial c}{\partial x} = \frac{1}{\sqrt{b}} \frac{\partial b}{\partial x} = \frac{2}{3\sqrt{2\log 3}}$ ,  $\frac{\partial c}{\partial y} = \frac{1}{\sqrt{b}} \frac{\partial b}{\partial y} = \sqrt{\frac{\log 3}{2}}$  Computation only depends on node 'b'

$$4. f = c + b = \sqrt{2\log 3} + 2\log 3, \frac{\partial f}{\partial x} = \frac{\partial c}{\partial x} + \frac{\partial b}{\partial x} = \frac{2}{3} \left( 1 + \frac{1}{\sqrt{2\log 3}} \right), \frac{\partial f}{\partial y} = \frac{\partial c}{\partial y} + \frac{\partial b}{\partial y} = \sqrt{\frac{\log 3}{2}} + \log 3$$



Computation only depends on node 'b' and 'c'

### Reverse mode auto-diff (Backpropagation)



$$0. x = 3, y = 2$$

1. 
$$a = log 3$$

$$2. b = 2log3$$

3. 
$$c = \sqrt{2log3}$$

$$4. f = \sqrt{2\log 3} + 2\log 3$$

$$0' \cdot \frac{\partial f}{\partial f} = 1$$

1'. 
$$\frac{\partial f}{\partial c} = \frac{\partial f}{\partial f} \frac{\partial f}{\partial c} = 1$$

$$1 \cdot \frac{\partial}{\partial c} - \frac{\partial}{\partial f} \frac{\partial}{\partial c} - 1$$

$$2' \cdot \frac{\partial f}{\partial b} = \frac{\partial}{\partial c} \frac{\partial}{\partial b} + \frac{\partial}{\partial f} \frac{\partial}{\partial c} = \frac{1}{\sqrt{b}} + 1 = \frac{1}{\sqrt{2log3}} + 1$$

3'. 
$$\frac{\partial f}{\partial a} = \frac{\partial f}{\partial b} \frac{\partial b}{\partial a} = 2(\frac{1}{\sqrt{2log^3}} + 1)$$

4'. 
$$\frac{\partial f}{\partial x} = \frac{\partial f}{\partial a} \frac{\partial a}{\partial x} = \frac{2}{3} \left( \frac{1}{\sqrt{2 \log 3}} + 1 \right)$$

$$\frac{\partial f}{\partial y} = \frac{\partial f}{\partial b} \frac{\partial b}{\partial y} = \left(\frac{1}{\sqrt{2\log 3}} + 1\right) a = \sqrt{\frac{\log 3}{2}} + \log 3$$



Backward pass depends on node values

computed in forward pass.

### Autodiff by Jacobian multiplication

Consider  $g = f_1 \circ f_2 \circ \cdots \circ f_N$  where  $f_i: \mathbb{R}^{n_i} \to \mathbb{R}^{n_{i-1}}$  for  $i = 1, \dots, N$ .

Chain rule: 
$$\nabla_{x}g(x) = Df_1 Df_2 \cdots Df_N D$$
 denotes Jacobian.

Forward mode:  $Df_1(Df_2(\cdots(Df_{N-1}Df_N)\cdots))$ 

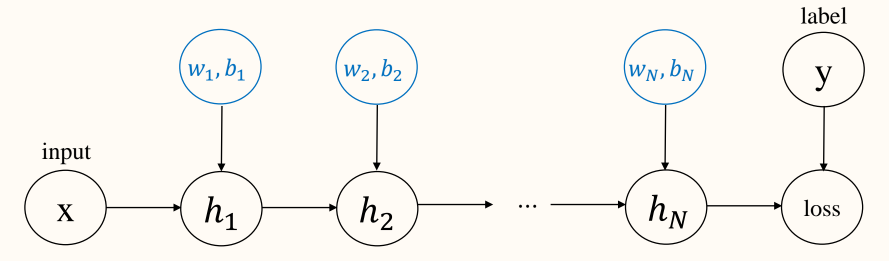
Reverse mode:  $(((Df_1 \ Df_2) \ Df_3) \cdots) \ Df_N$ 



Optimal if  $n_0 \le n_1 \le \cdots \le n_N$ . Proof by dynamic programming.



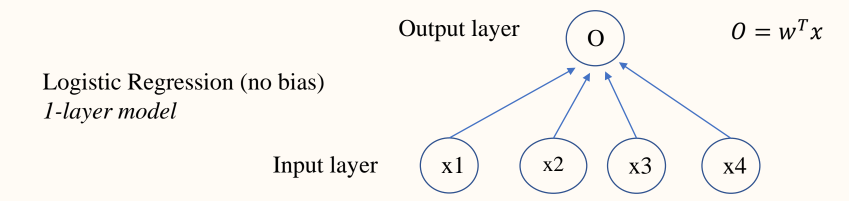
#### Backprop on multi-layer perceptron

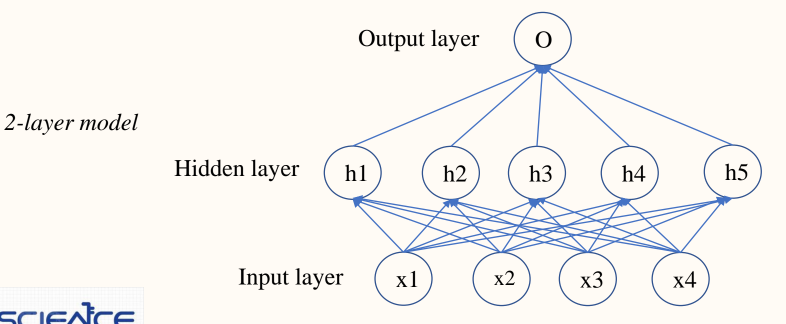


- In NN training, parameters and fixed inputs are distinguished.
- In Pytorch, 1. evaluate the loss function
  - 2. call ·backward() to perform backward pass and compute gradients.
- When performing the forward pass, intermediate node values are stored so that they can later be used in backward pass.
- In testing loop, we don't compute gradients so this is unnecessary.
- The torch.no\_grad() context marger allows intermediate node values to discarded or not be stored. This saves memory and can reduce the time to compute the test loop.



#### Multilayer Perceptron (fully connected deep neural network)





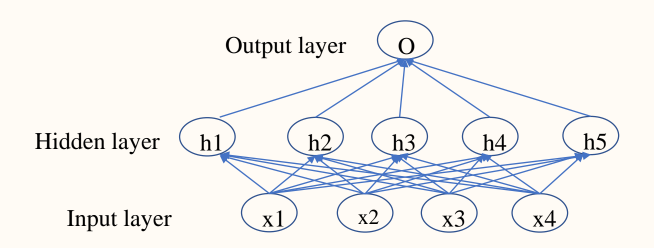
$$0 = W_2 h$$

$$h = W_1 x$$
  $h_i = (W_1)_i x$ ,  
 $(W_1 \text{ is } 5x4)$   $(i = 1,2,3,4,5.)$   
 $W_1 \text{ is } 1x4$ 

$$O=W_2(W_1x) = (W_2W_1)x$$
  
(This equation is linear in x !!!)



#### Deep Neural Network

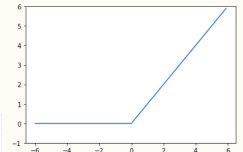


$$O=W_2h$$
  $W_2$  is  $1 \times 5$ 

$$h = \sigma(W_1 x)$$
  $W_1$  is  $5 \times 4$   
 $\sigma$  is a non-linear function  
Applied elementwise

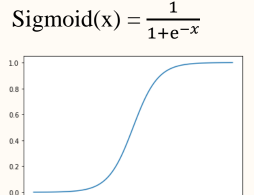
#### Use non-linear activation functions

#### -Rectified Linear Unit (ReLU) ReLU(z) = max(z,0)



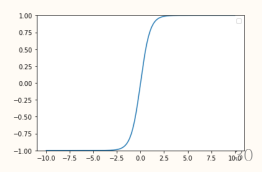
#### Common activation functions

-Sigmoid



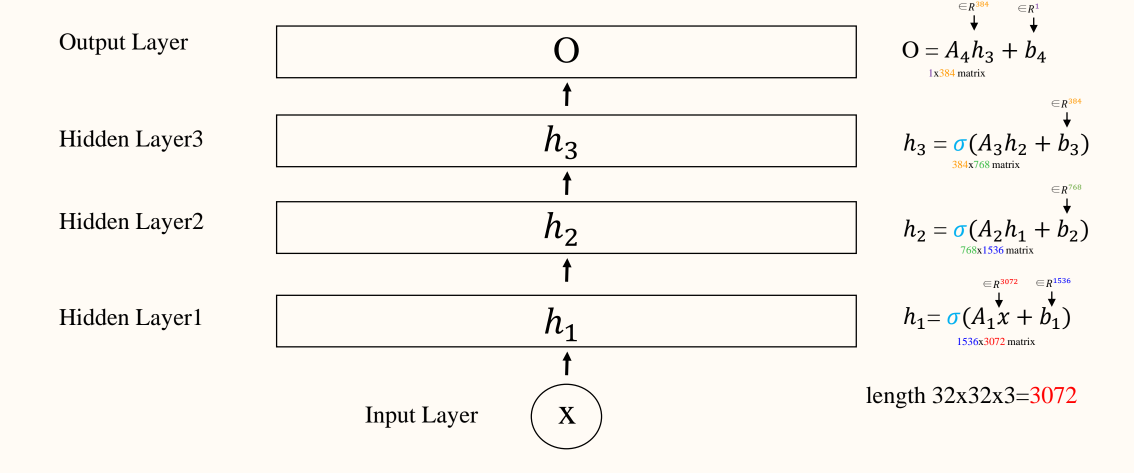
-Hyperbolic Tangent  

$$tanh(x) = \frac{1 - e^{-2x}}{1 + e^{-2x}}$$



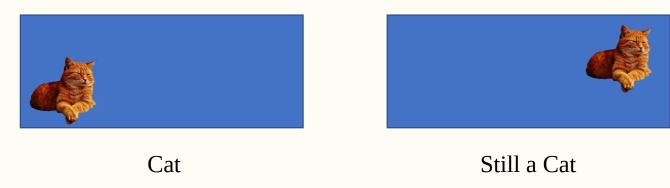


### Architecture for CIFAR10 Binary classification

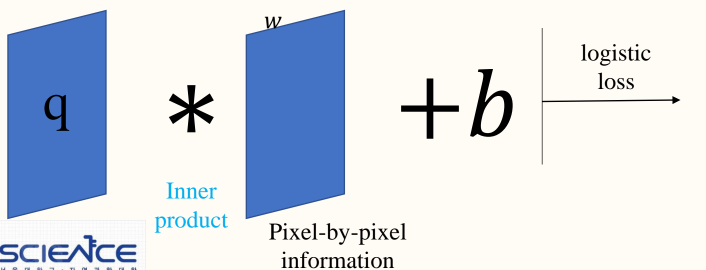


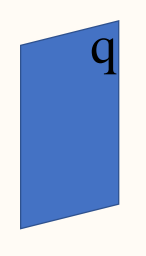


### Shift Invariance in Vision to Convolution



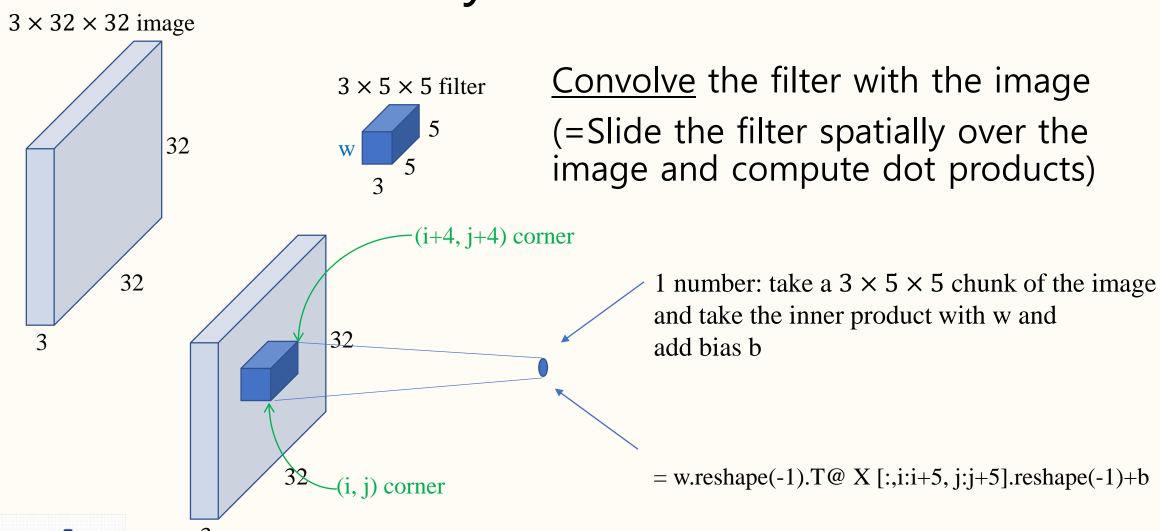
Logistic regression (with a single fully connected layer) does not encode shift invariance

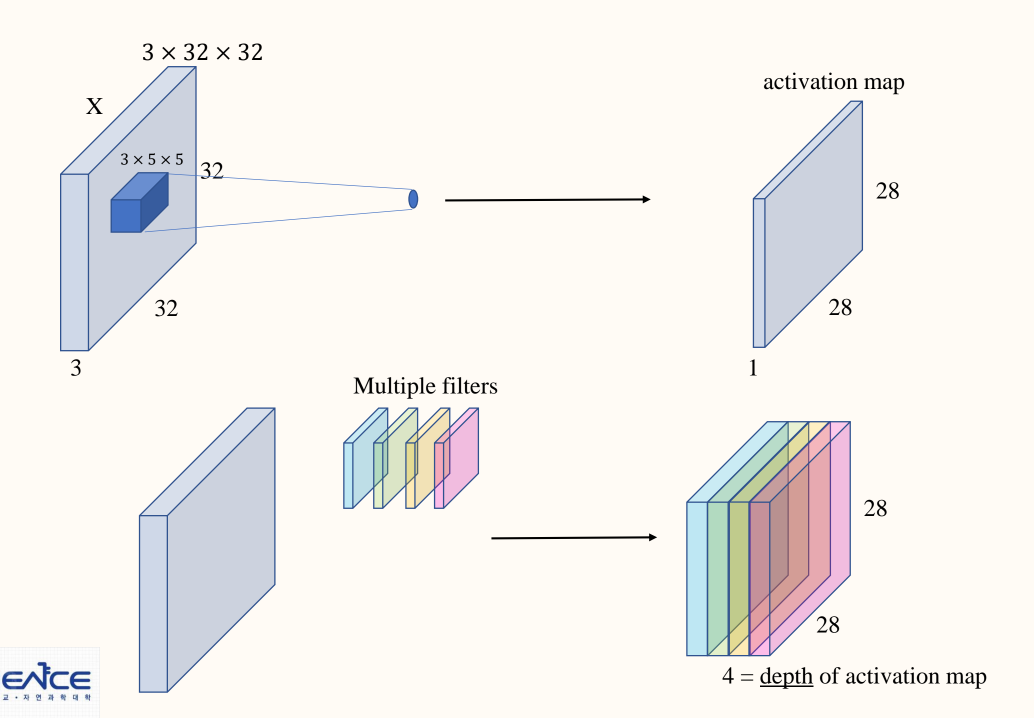




Translating digit changes output significantly

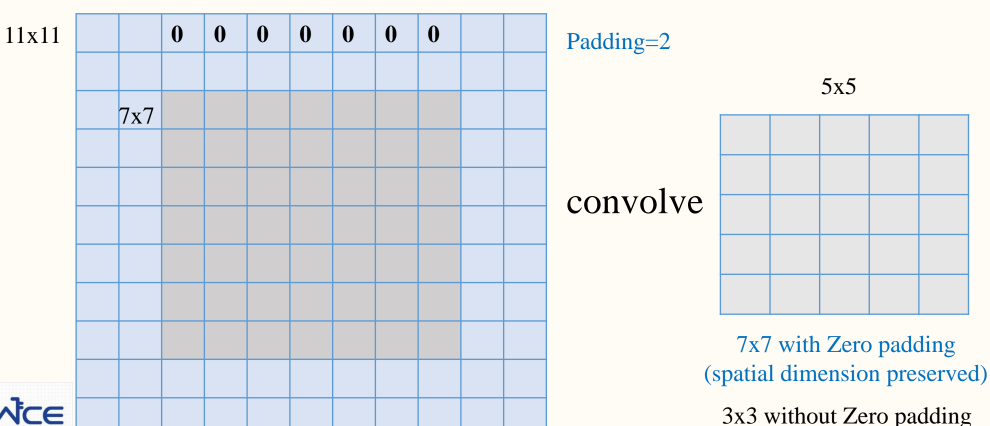
# Convolutional Layer





# Convolution options: Zero Padding

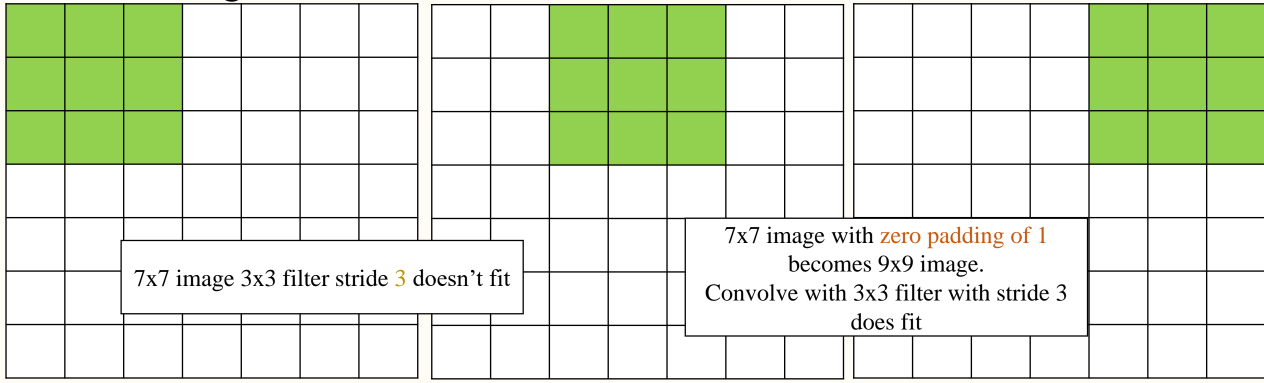
 $3 \times 32 \times 32$  convolved with  $3 \times 5 \times 5$  filter  $\Rightarrow 1 \times 28 \times 28$  activation map Spatial dimension 32 reduced to 28





## Convolution options : Stride

7x7 image convolved with 3x3 filter with stride 2





## Summary

Input  $W_1 \times H_1 \times D_1$ 

Conv layer parameters

⊩ K filters

 $\models$  F spatial extent ( $F \times F \times D_1$  filters)

**►** S stride

P padding

Output  $W_2 \times H_2 \times D_2$ 

$$F^2D_1K + K$$

$$\downarrow$$
filters biases

$$W_2 = \frac{W_1 - F + 2P}{S} + 1$$

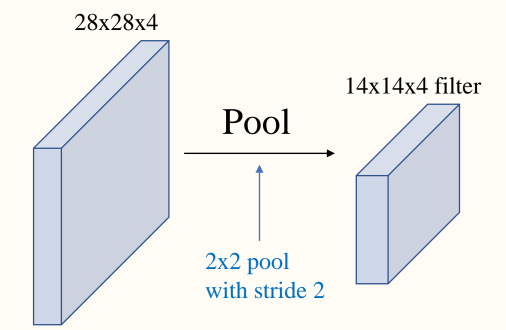
$$H_2 = \frac{H_1 - F + 2P}{S} + 1$$



$$D_2 = K$$

# Pooling

- Similar to convolutions
- Used to reduce the size of the output
- Operates over each activation map independently





#### Single depth size

1	1	2	4
5	6	7	8
3	2	1	0
1	2	3	4

#### Max Pool

2x2 filters and stride 2

6	8
3	4

Not an instance of convolution

Precise definitions in torch.nn.MaxPool2D torch.nn.AvgPool2D

### Average Pool 1 1 2 4

6 7 8 2x2 fil

3 2 1 0

1 2 3 4

2x2 filters and stride 2

Effect is subsampling (lowering image resolution)

3.25	5.25
2	2

Instance of convolution with fixed (untrainable) weights, including the independent operation over each activation map.

(Why?)



#### LeNet5

(LeCun, Bottou, Bengio, Haffner 1998)

Modern instances of LeNet5 use

- $\sigma = \text{ReLu}$
- MaxPool instead of avg. pool
- No  $\sigma$  after S2, S4 (Why?)
- No Gaussian connections
- Complete C4 connections

