IV. Preliminary Knowledge: Python and PyTorch Implementations

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Introduction

- This lecture aims to leverage Python and PyTorch implementations to (i) deepen understanding of preliminary knowledge about Statistics and Statistical Learning, and (ii) empirically validate key theoretical results.
- This lecture omits some basics of Python. For additional resources on the basics of Python, see https://cs231n.github.io/python-numpy-tutorial/#scipy.
- Most of the code drafts in this lecture were generated by ChatGPT 4, a generative AI.

Outline

Law of Large Numbers and Central Limit Theorem

- 2 Logistic Regression
- 3 NEURAL NETWORK

Recapping Law of Large Numbers

- (Weak) Law of Large Numbers: Let \bar{X} be the mean of n i.i.d. samples from a distribution with a finite first moment μ . Then, $\bar{X} \stackrel{p}{\to} \mu$. That is, for any $\epsilon > 0$, $\mathbb{P}(|\bar{X} \mu| > \epsilon) \to 0$.
- Experiment: We will examine whether $\mathbb{P}(|\bar{X} \mu| > \epsilon)$ converges to zero:
 - Sample a dataset of size n = 1000, denoted as $(x_i)_{i=1}^n$, M = 50 times to draw a spaghetti plot having M trajectories of \bar{X} (x-axis: n; y-axis: \bar{X}).
 - ② Sample a dataset of size n=1000, totaling M=1000 times. Plot empirical estimates of $\mathbb{P}(|\bar{X}-\mu|>\epsilon=0.1)$ as a function of n.
- For the distribution of X, we consider the Student's t distribution with a degree of freedom ν : $p_{\theta:=\nu}(x) \propto (1+x^2/\nu)^{-(\nu+1)/2}$. Here, $\mu=0$ for $\nu>1$ and μ is undefined otherwise. Its variance is finite if and only if $\nu>2$.

Python Code 1: Law of Large Numbers I

```
import numpy as np
import matplotlib.pyplot as plt
from scipy.stats import t
# Part 1: Spaghetti plot
np.random.seed(0) # For reproducibility
# Parameters
n = 1000
M = 50
nu = 3  # degrees of freedom for t-distribution
# Plotting spaghetti plot
plt.figure(figsize=(10, 6))
for i in range(M):
    plt.plot(range(1, n+1), np.cumsum(t.rvs(df=nu, size=n)) / np.arange(1, n+1), alpha=0.5)
plt.title('Spaghetti plot of sample means for M=50 trials')
```

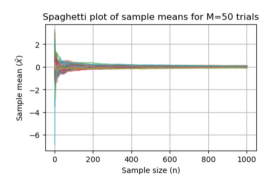
Python Code 1: Law of Large Numbers II

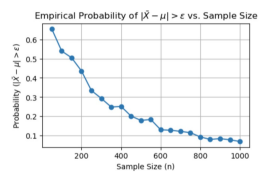
```
plt.xlabel('Sample size (n)')
plt.ylabel('Sample mean ($\\bar{X}$)')
plt.grid(True)
plt.show()
# Part 2: Empirical probability plot
N = 1000 # Max sample size
M = 1000 # Number of trials
epsilon = 0.1
# Computing probabilities
probabilities = []
sample_sizes = np.arange(50, N+1, 50) # Step through sample sizes from 50 to N
for n in sample sizes:
    count_within_epsilon = 0
    for _ in range(M):
```

Python Code 1: Law of Large Numbers III

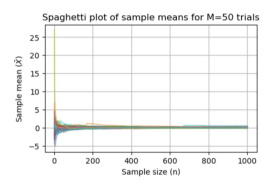
```
sample_mean = t.rvs(df=nu, size=n).mean()
        if abs(sample_mean) > epsilon:
            count within epsilon += 1
    probabilities.append(count_within_epsilon / M)
# Plotting the probabilities
plt.figure(figsize=(10, 6))
plt.plot(sample_sizes, probabilities, marker='o')
plt.title('Empirical Probability of $|\\bar{X} - \\mu| > \\epsilon$ vs. Sample Size')
plt.xlabel('Sample Size (n)')
plt.vlabel('Probability ($|\\bar{X} - \\mu| > \\epsilon$)')
plt.grid(True)
plt.show()
```

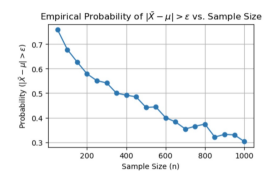
Python Code 1: Law of Large Numbers IV





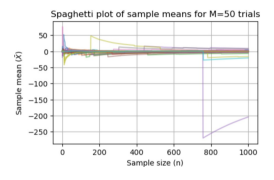
Advanced Discussion

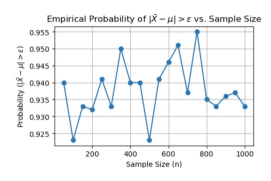




- Results when $\nu=2$. In this case, $\mathbb{E}_X X=0$ and $\operatorname{Var}_X X=\infty$.
- Given n=1000, the empirical estimate of $\mathbb{P}(|\bar{X}-\mu|>\epsilon)$ increased a lot. We need larger sample sizes to empirically confirm the convergence of \bar{X} .

Advanced Discussion





• Results when $\nu=1$. In this case, $\mathbb{E}_X X$ is undefined. This distribution is referred to as the *Cauchy distribution*. In the right figure, $\mu=0$ was chosen for consistent visualization. **Q:** Conditions for the Law of Large Numbers do not hold; However, this does NOT guarantee that \bar{X} diverges. Based on plots, should we conclude that \bar{X} does not converge to zero (or any other point), or could this result be due to a small sample size issue?

Recapping Central Limit Theorem

• <u>Central Limit Theorem:</u> Let \bar{X} be the mean of n i.i.d. samples from a distribution with a finite population mean μ and standard deviation σ . Then,

$$\sqrt{n}(\bar{X} - \mu)/\sigma \xrightarrow{d} Z \sim N(0, 1).$$
 (1)

That is, $\mathbb{P}(\sqrt{n}(\bar{X} - \mu)/\sigma \le z) \to \mathbb{P}(Z \le z)$ for any $z \in \mathbb{R}$.

- Experiment: We will examine whether $\mathbb{P}(\sqrt{n}(\bar{X}-\mu)/\sigma \leq z)$ converges to $\mathbb{P}(Z \leq z)$:
 - **●** Sample a dataset of size $n \in \{25, 50, 100, 200\}$, denoted as $(x_i)_{i=1}^n$, M = 1000 times to compute M estimates of $\sqrt{n}(\bar{X} \mu)/\sigma$ for each n.
 - Examine whether the empirical distribution functions of $\sqrt{n}(\bar{X} \mu)/\sigma$ converge to the distribution function of a standard normal N(0,1).
- For the distribution of X, we consider a Gaussian mixture model: $p_{\theta}(x) = \pi_1 p(x; \mu_1, \sigma_1^2) + \pi_2 p(x; \mu_2, \sigma_2^2)$ where $\theta := (\pi_1, \pi_2, \mu_1, \sigma_1, \mu_2, \sigma_2)^T$. Here, $\mu = \pi_1 \mu_1 + \pi_2 \mu_2$ and $\sigma = \sqrt{(\pi_1 \sigma_1^2 + \pi_2 \sigma_2^2) + (\pi_1 (\mu_1 \mu)^2 + \pi_2 (\mu_2 \mu)^2)}$.

Python Code 2: Central Limit Theorem I

```
import numpy as np
import matplotlib.pyplot as plt
from scipy.stats import norm
# Parameters for Gaussian Mixture
mul, sigmal = 0, 1 # Mean and std dev of first normal distribution
mu2, sigma2 = 3, 1.5 # Mean and std dev of second normal distribution
pi1, pi2 = 0.5, 0.5 # Proportion of the first and second distribution
# Combined mean and variance
mu = pi1 * mu1 + pi2 * mu2
sigma = np.sqrt(pi1 * (sigma1**2) + pi2 * (sigma2**2)
                + pi1 * ((mu - mu1)**2) + pi2 * ((mu - mu2)**2))
# Sample sizes to test
sample sizes = [25, 50, 100, 200]
M = 1000 # Number of experiments
```

Python Code 2: Central Limit Theorem II

```
# Function to generate samples from a Gaussian Mixture
def generate_gaussian_mixture(n):
   n1 = np.random.binomial(n, pi1)
   n2 = n - n1
    samples1 = np.random.normal(mu1, sigma1, n1)
    samples2 = np.random.normal(mu2, sigma2, n2)
    return np.concatenate((samples1, samples2))
# Function to calculate empirical distribution function
def ecdf(data):
   n = len(data)
   x = np.sort(data)
    y = np.arange(1, n+1) / n
   return x, y
# Create figure for ECDF and CDF comparison
```

Python Code 2: Central Limit Theorem III

```
fig, ax = plt.subplots(figsize=(12, 8))
# Define line styles for different sample sizes
line styles = ['-', '--', '-.', ':']
mise values = []
for i, n in enumerate(sample_sizes):
    sample_means = []
    for _ in range(M):
        samples = generate_gaussian_mixture(n)
        sample_mean = np.mean(samples)
        sample means.append(sample mean)
    # Normalize sample means
    sample_means = np.array(sample_means)
    normalized_means = np.sqrt(n) * ((sample_means - mu) / sigma)
```

Python Code 2: Central Limit Theorem IV

```
# Calculate and plot ECDF
    x, y = ecdf(normalized means)
    ax.step(x, y, where='post', label=f'ECDF for n={n}', linestyle=line styles[i])
    # Calculate MISE
    x_grid = np.linspace(min(x), max(x), 500)
    ecdf_values = np.interp(x_grid, x, y)
    cdf_values = norm.cdf(x_grid)
   mise = np.mean((ecdf_values - cdf_values)**2)
   mise values.append(mise)
# Plotting CDF of standard normal distribution
x_grid = np.linspace(-3, 3, 100)
ax.plot(x grid, norm.cdf(x grid), 'k--', label='CDF N(0,1)', color='gray')
ax.set title('ECDF and CDF Comparison for Gaussian Mixture')
```

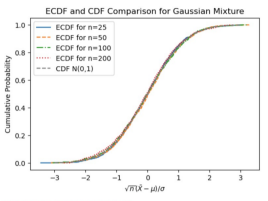
Python Code 2: Central Limit Theorem V

```
ax.set_xlabel(r'$\sqrt{n}(\bar{X}-\mu)/\sigma$')
ax.set_ylabel('Cumulative Probability')
ax.legend()

plt.show()

# Print MISE values for each sample size
for size, mise in zip(sample_sizes, mise_values):
    print(f'MISE for n={size}: {mise}')
```

Python Code 2: Central Limit Theorem VI



MISE for n=25: 2.926529079375994e-05 MISE for n=50: 4.158184313277996e-05 MISE for n=100: 2.709211076327643e-05 MISE for n=200: 0.00012699633237750485

Q: Why mean integrated squared errors (MISEs) do not monotonically decrease?

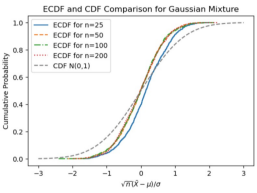
Python Code 2': Central Limit Theorem? I

• The following is the draft version of the function *generate_gaussian_mixture* that ChatGPT gave:

```
# Function to generate samples from a Gaussian Mixture

def generate_gaussian_mixture(n):
    n1 = int(n*pi1) # previous example: n1 = np.random.binomial(n, pi1)
    n2 = n - n1
    samples1 = np.random.normal(mu1, sigma1, n1)
    samples2 = np.random.normal(mu2, sigma2, n2)
    return np.concatenate((samples1, samples2))
```

Python Code 2': Central Limit Theorem? II



MISE for n=25: 0.007840964793214642 MISE for n=50: 0.005802817753966564 MISE for n=100: 0.0049223933000111094 MISE for n=200: 0.0052024150821751965

Q: What happened?

Outline

Law of Large Numbers and Central Limit Theorem

2 Logistic Regression

3 NEURAL NETWORK

Optimization

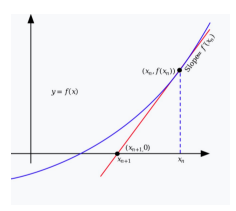
• The estimator \hat{f}_n usually does not have a (computable) closed-from expression:

$$\hat{f}_n \in \underset{f \in \mathcal{F}}{\arg\min} n^{-1} \sum_{i=1}^n I(f(\vec{x}_i), y_i). \tag{2}$$

- Typical remedies are to apply iterative algorithms to solve the above optimization problem.
- In this subsection, we focus on simple logistic regression where we have an univariate X, $\mathcal{F} = \{f_{\theta}(x) = \sigma(\beta_0 + \beta_1 x) \mid \theta = (\beta_0, \beta_1)^T \in \mathbb{R}^2\}$, and we use the cross-entropy loss:

$$I(f_{\theta}(x), y) = -I(y = 1)\log \sigma(f_{\theta}(x)) - I(y = -1)\log(1 - \sigma(f_{\theta}(x)))$$

where $\sigma(x) = 1/(1 + \exp(-x))$ denotes the standard logistic function (or sigmoid function), I is the indicator function, and y takes values in $\{1, -1\}$ indicating the class labels.



• The Newton-Raphson method: $x^{(t+1)} = x^{(t)} - f(x^{(t)})/f'(x^{(t)})$ is used to find a solution to f(x) = 0.

Image source: https://en.wikipedia.org/wiki/Newton%27s_method.

• Let $R_n(f_\theta)$ be the empirical risk. In logistic regression, the minimizer $(\hat{f}_n =) f_{\hat{\theta}_n}$ uniquely exists and is the solution to:

$$\frac{\partial R_n(\theta)}{\partial \theta} = \begin{pmatrix} n^{-1} \sum_{i=1}^n \left(\sigma(\beta_0 + \beta_1 x_i) - I(y_i = 1) \right) \\ n^{-1} \sum_{i=1}^n x_i \left(\sigma(\beta_0 + \beta_1 x_i) - I(y_i = 1) \right) \end{pmatrix} = \begin{pmatrix} 0 \\ 0 \end{pmatrix}. \tag{3}$$

The $R_n(\theta)$ is the same as the negative log-likelihood, so $\hat{\theta}_n$ is a maximum likelihood estimator.¹

• The Newton-Raphson method for the logistic regression can be expressed as:

$$\hat{\theta}_n^{(t+1)} = \hat{\theta}_n^{(t)} - \left(\partial^2 R_n(\theta) / \partial \theta \partial \theta^T \right)^{-1} (\partial R_n(\theta) / \partial \theta) \big|_{\theta = \hat{\theta}_n}. \tag{4}$$

¹In this context, Equation (3) is a score equation.

• The Hessian can be expressed as:

$$\frac{\partial^2 R_n(\theta)}{\partial \theta \partial \theta^T} = \begin{pmatrix} n^{-1} \sum_{i=1}^n w_i(\theta) & n^{-1} \sum_{i=1}^n w_i(\theta) x_i \\ n^{-1} \sum_{i=1}^n w_i(\theta) x_i & n^{-1} \sum_{i=1}^n w_i(\theta) x_i^2 \end{pmatrix}$$
(5)

where
$$w_i(\theta) := \sigma(\beta_0 + \beta_1 x_i) (1 - \sigma(\beta_0 + \beta_1 x_i)).$$

- Experiment: We will examine whether $\hat{\theta}_n^{(t)}$ converges to the true θ^* :
 - **①** Sample a dataset of size $n \in \{100, 200, 400, 800\}$, denoted as $((x_i, y_i))_{i=1}^n$.
 - ② Apply the Newton-Raphson method to find $\hat{\theta}_n^{(T=25)}$. The initial point $\hat{\theta}_n^{(0)}$ is randomly sampled from $U([-0.1,0.1]^2)$.
 - **3** Repeat the previous steps M = 500 times to compute M estimates of $\hat{\theta}_n^{(T)}$ for each n.
 - Plot $\|\hat{\theta}_n^{(T)} \theta^*\|$ as a function of n and plot a scattar plot to visualize $\sqrt{n}(\hat{\theta}_n^{(T)} \theta^*)$.
- For the distribution of (X,Y), we consider the following setting: $Y \sim 2 \operatorname{Ber}(p) 1$ and $X|Y = y \sim N(x; \mu_y, \sigma_\epsilon^2)$. Then, $\theta^* = ((\mu_{-1}^2 \mu_1^2)/2\sigma_\epsilon^2, (\mu_1 \mu_{-1})/\sigma_\epsilon^2)^T$.

Python Code 3: Newton-Raphson Method I

```
import numpy as np
import matplotlib.pyplot as plt
from scipy.stats import bernoulli, norm
def sigmoid(x):
    """Stable sigmoid function to prevent overflow."""
   return 1 / (1 + np.exp(-x))
def logistic regression(x, y, theta_init, max_iter=25):
    """Performs logistic regression using the Newton-Raphson method."""
   n = len(y)
    theta = theta init
    indicator = (y+1)/2 # Convert class labels from {1, -1} to {1, 0}
    for in range(max iter):
        p = sigmoid(x @ theta) # Current estimate of probability
        gradient = x.T @ (p - indicator) / n
```

Python Code 3: Newton-Raphson Method II

```
W = np.diag(p * (1 - p))
        Hessian = x.T @ W @ x / n
        theta -= np.linalg.inv(Hessian) @ gradient
    return theta
def generate_data(n, mu_y, sigma_epsilon):
    """Generates binary classification data with logistic features."""
    y = 2 * bernoulli.rvs(0.5, size=n) - 1
    x = norm.rvs(loc=[mu_y[yy] for yy in y], scale=sigma_epsilon, size=n)
    return np.vstack([np.ones(n), x]).T, y # Add intercept
# Setup parameters
sample sizes = [100, 200, 400, 800]
M = 500
mu_v = \{1: 1, -1: -2\}
sigma_epsilon = 1.0
theta star = np.arrav(\lceil (\text{mu v} \lceil -1] **2 - \text{mu v} \lceil 1] **2) / (2 * sigma epsilon**2).
```

Python Code 3: Newton-Raphson Method III

```
(mu_y[1] - mu_y[-1]) / (sigma_epsilon**2)])
norm diffs = []
scaled_diffs = {n: [] for n in sample_sizes}
# Perform simulations
for n in sample_sizes:
   theta_estimates = []
    for _ in range(M):
        x, y = generate_data(n, mu_y, sigma_epsilon)
        theta init = np.random.uniform(-0.1, 0.1, 2)
        theta_hat = logistic regression(x, y, theta_init)
        norm_diff = np.linalg.norm(theta_hat - theta_star)
        theta estimates.append(norm diff)
        scaled diff = np.sqrt(n) * (theta hat - theta star)
        scaled_diffs[n].append(scaled_diff)
    norm diffs.append(np.mean(theta estimates))
```

Python Code 3: Newton-Raphson Method IV

```
# Plot for norm differences
plt.figure(figsize=(8, 5))
plt.plot(sample_sizes, norm_diffs, marker='o')
plt.xlabel('Sample Size (n)')
plt.ylabel(r'\frac{r'}{\hat{t}}_{n}^{(T)} - \frac{*}{|\dot{t}}
plt.title('Convergence of Newton-Raphson Estimates')
plt.grid(True)
plt.show()
# Plot scaled differences as scatter plots
fig, axes = plt.subplots(1, len(sample sizes), figsize=(20, 5), sharex=True, sharey=True)
for i, n in enumerate(sample_sizes):
    diffs = np.array(scaled_diffs[n])
    sigma_x = np.std(diffs[:, 0])
    sigma_y = np.std(diffs[:, 1])
    limit x = 3 * sigma x
```

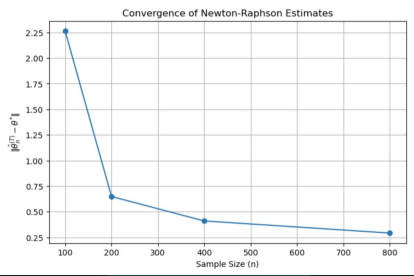
Python Code 3: Newton-Raphson Method V

```
limit_y = 3 * sigma_y

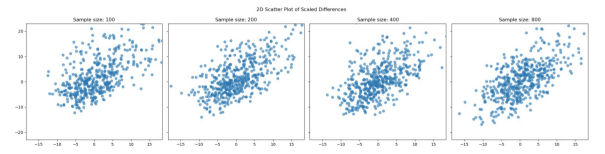
axes[i].scatter(diffs[:, 0], diffs[:, 1], alpha=0.6)
axes[i].set_title(f'Sample size: {n}')
axes[i].set_xlim([-limit_x, limit_x])
axes[i].set_ylim([-limit_y, limit_y])

plt.suptitle('2D Scatter Plot of Scaled Differences')
plt.tight_layout()
plt.show()
```

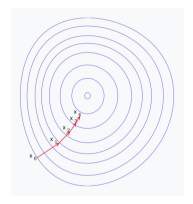
Results: Newton-Raphson Method



Results: Newton-Raphson Method



Optimization: Gradient Descent Method



• The gradient descent method: $x^{(t+1)} = x^{(t)} - \gamma_t f'(x^{(t)})$ is used to find a solution to f(x) = 0 where γ_t represents learning rates.

Image source: https://en.wikipedia.org/wiki/Gradient_descent.

Optimization: Gradient Descent Method

• The gradient descent method for the logistic regression can be expressed as:

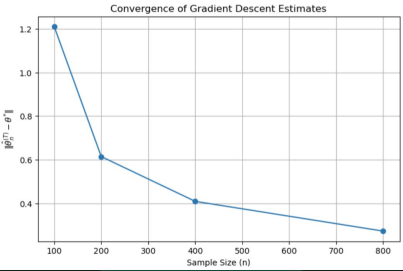
$$\hat{\theta}_n^{(t+1)} = \hat{\theta}_n^{(t)} - \gamma_t \left(\partial R_n(\theta) / \partial \theta \right) |_{\theta = \hat{\theta}_n}. \tag{6}$$

• Experiment: We use the experimental setting from the previous example for the Newton-Raphson method while updating the optimization part to the gradient descent method with T=1000 and $\gamma_t=50/(t+50)$.

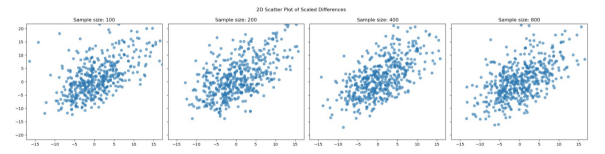
Python Code 4: Gradient Descent Method

```
def gd logistic regression(x, y, theta init, init learning rate=20.0, max iter=1000):
    """Performs logistic regression using Gradient Descent."""
   n = len(y)
    theta = theta_init
    indicator = (y+1)/2 # Convert class labels from {1, -1} to {1. 0}
    for t in range(max_iter):
        learning_rate = init_learning_rate*(50/(t+50))
        p = sigmoid(x @ theta)
        gradient = x.T @ (p - indicator)/n
        theta -= learning rate * gradient
    return theta
```

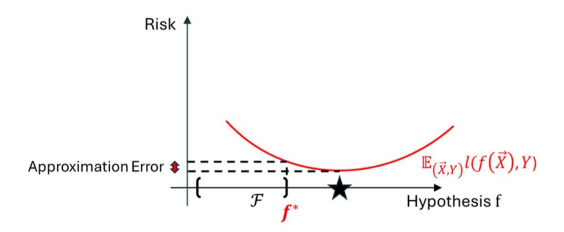
Results: Gradient Descent Method



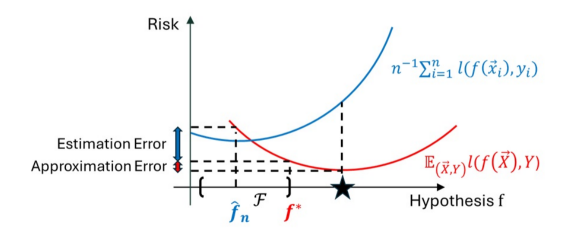
Results: Gradient Descent Method



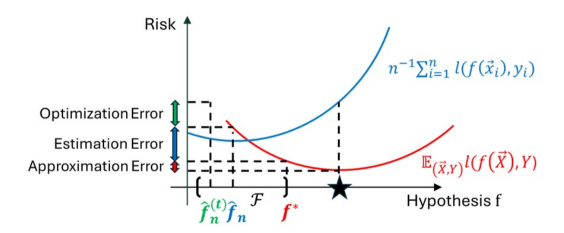
Recapping Approximation Error



Recapping Estimation Error



Optimization Error



Outline

Law of Large Numbers and Central Limit Theorem

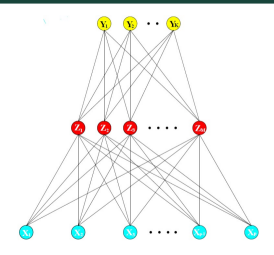
2 Logistic Regression

3 NEURAL NETWORK

Introduction to PyTorch

- PyTorch is a popular open-source machine learning library for Python, known for its flexibility and ease of use in research.
- It provides two high-level features:
 - Tensor computation (like NumPy) with strong GPU acceleration
 - Automatic differentiation for building and training neural networks
- Further details about PyTorch will be provided in a later main lecture, along with implementations of several deep generative models. In this lecture, we will review two basic neural network-based classifiers on the MNIST dataset.

Recapping Neural Network



The figure is from Hastie (2009).

Recapping Neural Network

• Outputs from neural networks with one hidden layer can be expressed as:

$$f_{\theta}(\vec{x}) = \beta_0 + \sum_{h=1}^{H} \beta_h \sigma(\alpha_h + \vec{w}_h^T \vec{x})$$
 (7)

where $\theta := (\beta_0, \dots, \beta_H, \alpha_1, \dots, \alpha_H, \vec{w}_1, \dots, \vec{w}_H)^T$ and $g_h(\cdot) = \beta_h \sigma(\cdot)$ where σ is the sigmoid function (can be replaced with other activation functions). Here, $\alpha_h + \vec{w}_h^T \vec{x}$ represents the h-th hidden node value, and $(\beta_1, \dots, \beta_H)^T$ and β_0 are weight vector and bias term, respectively, for the output layer.

PyTorch Code 1: Dense Network with One Hidden Layer I

```
import torch
import torch.nn as nn
import torch.optim as optim
import torchvision.transforms as transforms
import torchvision.datasets as datasets
import matplotlib.pyplot as plt
from torch.utils.data import DataLoader
# Define the neural network with one hidden layer
class FCNet(nn.Module):
    def __init__(self):
        super(FCNet, self). init ()
        self.fc1 = nn.Linear(28*28, 512) # Flatten 28x28 images to a 784 vector for each image
        self.relu = nn.ReLU()
        self.fc2 = nn.Linear(512, 10) # 10 output classes
    def forward(self, x):
```

PyTorch Code 1: Dense Network with One Hidden Layer II

```
x = x.view(-1, 28*28) # Flatten the images
        x = self.relu(self.fc1(x))
        x = self.fc2(x)
        return x
# Load the MNTST dataset
transform = transforms.Compose([
    transforms.ToTensor(),
    transforms.Normalize((0.5,), (0.5,))
1)
train dataset = datasets.MNIST(root='./data', train=True, download=True, transform=transform)
test dataset = datasets.MNIST(root='./data', train=False, download=True, transform=transform)
train loader = DataLoader(dataset=train dataset, batch size=64, shuffle=True)
test loader = DataLoader(dataset=test dataset, batch size=1000, shuffle=False)
```

PyTorch Code 1: Dense Network with One Hidden Layer III

```
# Initialize the network and optimizer
model = FCNet()
criterion = nn.CrossEntropyLoss()
optimizer = optim.SGD(model.parameters(), lr=0.01, momentum=0.5)
# Training the network
def train(num_epochs, model, loaders):
    train loader, test loader = loaders
    for epoch in range(num epochs):
        model train()
        for data, target in train_loader:
            optimizer.zero_grad()
            output = model(data)
            loss = criterion(output, target)
            loss.backward()
            optimizer.step()
```

PyTorch Code 1: Dense Network with One Hidden Layer IV

```
model eval()
        test_loss = 0
        correct = 0
        with torch.no grad():
            for data, target in test_loader:
                output = model(data)
                test loss += criterion(output, target).item()
                pred = output.data.max(1, keepdim=True)[1]
                correct += pred.eq(target.data.view_as(pred)).sum()
        test_loss /= len(test_loader.dataset)
        print(f'Epoch: {epoch+1}, Test Loss: {test_loss:.4f},
        Accuracy: {correct}/{len(test loader.dataset)} ({100. * correct / len(test loader.data
# Visualization of results
def visualize(model, loader):
   model.eval()
   data, _ = next(iter(loader))
```

PyTorch Code 1: Dense Network with One Hidden Layer V

```
with torch.no_grad():
        output = model(data)
    plt.figure(figsize=(10, 8))
    for i in range(6):
        plt.subplot(2, 3, i + 1)
       plt.tight_layout()
        plt.imshow(data[i][0], cmap='gray', interpolation='none')
        plt.title("Prediction: {}".format(output.data.max(1, keepdim=True)[1][i].item()))
        plt.xticks([])
        plt.yticks([])
   plt.show()
# Run training and visualization
train(5, model, (train_loader, test_loader))
visualize(model. test loader)
```

Result: Dense Neural Network

```
Epoch: 1, Test Loss: 0.0003, Accuracy: 9108/10000 (91%)
Epoch: 2, Test Loss: 0.0003, Accuracy: 9231/10000 (92%)
Epoch: 3, Test Loss: 0.0002, Accuracy: 9339/10000 (93%)
Epoch: 4, Test Loss: 0.0002, Accuracy: 9449/10000 (94%)
Epoch: 5, Test Loss: 0.0002, Accuracy: 9449/10000 (94%)
```

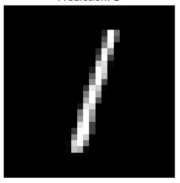
Prediction: 7



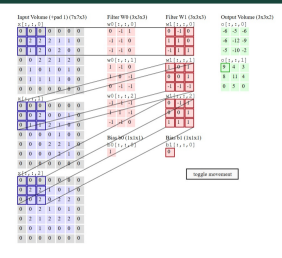
Prediction: 2



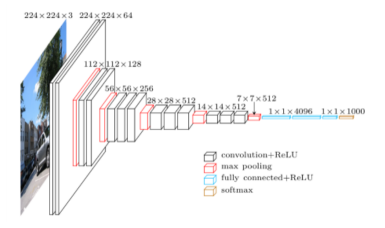
Prediction: 1



Recapping Convolutional Neural Network



Recapping Convolutional Neural Network



The figure is from Simonyan (2014).

PyTorch Code 2: Convolutional Neural Network I

```
import torch
import torch.nn as nn
import torch.optim as optim
import torchvision.transforms as transforms
import torchvision.datasets as datasets
import matplotlib.pyplot as plt
from torch.utils.data import DataLoader
# Define the neural network architecture
class SimpleCNN(nn.Module):
    def __init__(self):
        super(SimpleCNN, self). init ()
        self.conv1 = nn.Conv2d(1, 10, kernel size=5)
        self.conv2 = nn.Conv2d(10, 20, kernel size=5)
        self.pool = nn.MaxPool2d(2)
        self.fc1 = nn.Linear(320, 50)
        self.fc2 = nn.Linear(50.10)
```

PyTorch Code 2: Convolutional Neural Network II

```
def forward(self. x):
        x = self.pool(torch.relu(self.conv1(x)))
        x = self.pool(torch.relu(self.conv2(x)))
        x = x.view(-1.320)
        x = torch.relu(self.fc1(x))
        x = self.fc2(x)
        return x
# Load the MNIST dataset
transform = transforms.Compose([
    transforms.ToTensor().
    transforms.Normalize((0.5,),(0.5,))
1)
train dataset = datasets.MNIST(root='./data', train=True, download=True, transform=transform)
test dataset = datasets.MNIST(root='./data', train=False, download=True, transform=transform)
```

PyTorch Code 2: Convolutional Neural Network III

```
train_loader = DataLoader(dataset=train_dataset, batch_size=64, shuffle=True)
test loader = DataLoader(dataset=test dataset, batch size=1000, shuffle=False)
# Initialize the network and optimizer
model = SimpleCNN()
criterion = nn.CrossEntropyLoss()
optimizer = optim.SGD(model.parameters(), lr=0.01, momentum=0.5)
# Train the network
def train(num epochs, model, loaders):
    train_loader, test_loader = loaders
    for epoch in range(num_epochs):
        model.train()
        for data, target in train loader:
            optimizer.zero_grad()
            output = model(data)
```

PyTorch Code 2: Convolutional Neural Network IV

```
loss = criterion(output, target)
    loss backward()
    optimizer.step()
model.eval()
test loss = 0
correct = 0
with torch.no grad():
    for data, target in test loader:
        output = model(data)
        test_loss += criterion(output, target).item()
        pred = output.data.max(1, keepdim=True)[1]
        correct += pred.eq(target.data.view as(pred)).sum()
test loss /= len(test loader.dataset)
print(f'Epoch: {epoch+1}, Test Loss: {test loss: .4f},
Accuracy: {correct}/{len(test loader.dataset)} ({100. * correct / len(test loader.data
```

PyTorch Code 2: Convolutional Neural Network V

```
# Visualizing some results
def visualize(model, loader):
   model eval()
    data, = next(iter(loader))
    with torch.no grad():
        output = model(data)
    plt.figure(figsize=(10, 8))
    for i in range(6):
        plt.subplot(2, 3, i + 1)
        plt.tight_layout()
        plt.imshow(data[i][0], cmap='gray', interpolation='none')
        plt.title("Prediction: {}".format(output.data.max(1, keepdim=True)[1][i].item()))
        plt.xticks([])
        plt.yticks([])
   plt.show()
```

Run training and visualization

PyTorch Code 2: Convolutional Neural Network VI

```
train(5, model, (train_loader, test_loader))
visualize(model, test_loader)
```

Result: Convolutional Neural Network

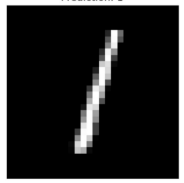
```
Epoch: 1, Test Loss: 0.0001, Accuracy: 9614/10000 (96%)
Epoch: 2, Test Loss: 0.0001, Accuracy: 9741/10000 (97%)
Epoch: 3, Test Loss: 0.0001, Accuracy: 9762/10000 (98%)
Epoch: 4, Test Loss: 0.0001, Accuracy: 9819/10000 (98%)
Epoch: 5, Test Loss: 0.0000, Accuracy: 9832/10000 (98%)
```

Prediction: 7

Prediction: 2



Prediction: 1



References I

Hastie, T. (2009). The elements of statistical learning: data mining, inference, and prediction. Simonyan, K. (2014). Very deep convolutional networks for large-scale image recognition. arXiv preprint arXiv:1409.1556.