深度学习笔记

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1	Line	ear Algebra						
	1.1	Eigenvalues and Eigenvectors						
	1.2	Singular value decomposition						
		1.2.1 Dimensionality Reduction						
2	Statistics 9							
	2.1	Frequentist statistics VS Bayesian statistics						
	2.2	Mean						
	2.3	Variance						
	2.4	Best Unbiased Estimators						
	2.5	Divergence						
		2.5.1 <i>f</i> -divergence						
		2.5.2 Kullback–Leibler divergence / Relative entropy						
	2.6	Entropy						
		2.6.1 Informational value						
		2.6.2 Entropy						
		2.6.3 Cross Entropy						
		2.6.4 Conditional Entropy						
	2.7	Mutual Information						
	2.8	Maximum likelihood estimation						
	2.0	2.8.1 MLE & KLD						
	2.9	When P(x) is constant						
		Maximum A Posteriori						
	2.10	Wide Militarity A 1 Osteriori						
3	Basi	15 C						
	3.1	Mechine Learning						
	3.2	iid						
	3.3	Convolution						
	3.4	Transposed Convolution						
	3.5	Universal approximation theorem						
	3.6	Initialization scheme						
	0.0	3.6.1 constant initialization						
		3.6.2 random initialization						
		3.6.3 xavier initialization						
		3.6.4 orthogonal initalization						
	2.7	0						
	3.7	Regularization						
	3.8	Data proprocessing						
	•	3.8.1 whitening						
	3.9	Activation functions						
		3.9.1 sigmoid						
		3.9.2 $tanh(x)$						
		3.9.3 Rectified linear units						

		3.9.4 haha	19						
		3.9.5 Loss functions	19						
	3.10	Batch Normalization	19						
	3.11	How Does Batch Normalization Help Optimization?	20						
		* *							
4	Opt	Optimization Algorithms 23							
	4.1	Challenges	23						
		4.1.1 Local Minima	23						
		4.1.2 Saddle Points	23						
		4.1.3 Vanishing gradients	24						
		4.1.4 noise / noise-free	24						
	4.2	Gradient Descent	24						
		4.2.1 Adaptive Methods	24						
		4.2.2 Stochastic Gradient Descent	25						
		4.2.3 Minibatch Stochastic Gradient Descent	25						
	4.3	SGDM - SGD with Momentum	25						
	4.4	NAG - Nesterov Accelerated Gradient	26						
	4.5	Adagrad	26						
	4.6	RMSProp	26						
	4.7	Adadelta	27						
	4.8	Adam	27						
	4.9	AdaMax	27						
		Nadam	27						
	4.11	AMSGrad	27						
5	Mod	als	29						
J	14100		_,						
6	Skij	Connections	31						
	6.1	Residual Units	31						
	6.2	The Shattered Gradients Problem	32						
	6.3	DenseNet	00						
	6.4		33						
	0.4	FCN	<i>33</i>						
		FCN	33						
	6.5	UNet family	33 33						
		UNet family	33 33 33						
		UNet family	33 33 34						
		UNet family	33 33 34 34						
		UNet family	33 33 34						
7	6.5	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+	33 33 34 34						
7	6.5	UNet family	33 33 34 34 34 37						
7	6.5 Sem	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+	33 33 34 34 34						
7	6.5 Sem 7.1	UNet family	33 33 34 34 34 37						
	6.5 Sem 7.1	UNet family	33 33 34 34 34 37						
	6.5 Sem 7.1 Auto	UNet family	33 33 34 34 34 37 37						
	6.5 Sem 7.1 Auto 8.1	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network encoders Linear Autoencoders VS PCA Undercomplete Autoencoders	33 33 34 34 34 37 37 39						
	6.5 Sem 7.1 Auto 8.1 8.2	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network encoders Linear Autoencoders VS PCA Undercomplete Autoencoders Regularized Autoencoders	33 33 34 34 34 37 37 39 39						
	6.5 Sem 7.1 Auto 8.1 8.2	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network encoders Linear Autoencoders VS PCA Undercomplete Autoencoders Regularized Autoencoders 8.3.1 Sparse Autoencoders	33 33 34 34 34 37 37 39 39 39 39						
	6.5 Sem 7.1 Auto 8.1 8.2	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network sencoders Linear Autoencoders VS PCA Undercomplete Autoencoders Regularized Autoencoders 8.3.1 Sparse Autoencoders 8.3.2 DAE - Denoising Autoencoders	33 33 34 34 34 37 37 39 39 39 39 40						
	6.5 Sem 7.1 Auto 8.1 8.2	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network encoders Linear Autoencoders VS PCA Undercomplete Autoencoders Regularized Autoencoders 8.3.1 Sparse Autoencoders	33 33 34 34 34 37 37 39 39 39 39						
	6.5 Sem 7.1 Auto 8.1 8.2 8.3	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network sencoders Linear Autoencoders VS PCA Undercomplete Autoencoders Regularized Autoencoders 8.3.1 Sparse Autoencoders 8.3.2 DAE - Denoising Autoencoders	33 33 34 34 34 37 37 39 39 39 39 40						
8	6.5 Sem 7.1 Auto 8.1 8.2 8.3	UNet family 6.5.1 U-Net 6.5.2 VNet 6.5.3 UNet 2+ 6.5.4 UNet 3+ antic Segmentation FCN - Fully Convolutional Network encoders Linear Autoencoders VS PCA Undercomplete Autoencoders Regularized Autoencoders 8.3.1 Sparse Autoencoders 8.3.2 DAE - Denoising Autoencoders 8.3.3 CAE - Contractive Autoencoders	33 33 34 34 37 37 39 39 39 40 40						

10	Approximate Inference	4
	10.1 Monte Carlo Methods	4
	10.1.1 Markov Chains	4
	10.2 Variational Inference	4
	10.2.1 Bayesian mixture of Gaussians	4
11	Deep Generative Models	4
	11.1 Auto-Regressive Generative Models	4
	11.2 Variational Autoencoders	4
	11.2.1 Why do we need VAEs?	4
	11.2.2 VAĚ	
	11.2.3 The VAE Lower Bound	4
	11.2.4 Regularizer - Solution of $KL(Q(z) P(z))$, Gaussion case	4
	11.2.5 Reconstruction Error	
	11.3 Generative Adversarial Models	4
	11.3.1 Mode collapse	4
	11.3.2 Wasserstein GAN and the Kantorovich-Rubinstein Duality	
	11.3.3 vanilla GAN	
	11.3.4 Latent space	5
	11.3.5 Architecture	
	11.3.6 Object functions	

Linear Algebra

1.1 Eigenvalues and Eigenvectors

$$Au = \lambda u \tag{1.1}$$

特征值 λ 代表线性变化的伸缩倍数,特征向量u代表变换的方向

1.2 Singular value decomposition

定义 对于 $A \in C^{m \times n}$,rank(A) = r,矩阵 $A^H A$ 的特征值为 $\lambda_1 \geqslant \lambda_2 \geqslant ... \geqslant \lambda_r > 0$, $\lambda_{r+1} = \lambda_{r+2} = ... = \lambda_n = 0$,称正数 $\sigma_i = \sqrt{\lambda_i}(i = 1, 2, ..., r)$ 为矩阵A的**奇异值**

$$A = U\Sigma V^{H}$$

$$= (u_{1} \quad u_{2} \quad \cdots \quad u_{m}) \begin{pmatrix} \sigma_{1} & & & & \\ & \sigma_{2} & & & \\ & & \ddots & & \\ & & ---- & \bar{0} & - & \\ & & & \bar{0} \end{pmatrix} \begin{pmatrix} v_{1}^{H} \\ v_{2}^{H} \\ \vdots \\ v_{n}^{H} \end{pmatrix}$$

$$= \sigma_{1} u_{1} v_{1}^{H} + \sigma_{2} u_{2} v_{1}^{H} + \cdots + \sigma_{r} u_{r} v_{r}^{H}$$

$$(1.2)$$

1.2.1 Dimensionality Reduction

将图片作为矩阵进行奇异值分解,提取前n个奇异值,则可以达到图像压缩的目的.

Statistics

- sample space The set, S, of all possible outcomes of a particular experiment is called the sample space for the experiment.
- event An event is any collection of possible outcomes of an experiment, that is, any subset of S.
- **conditional probability** If A and B are events in S, and P(B) > 0, then the conditional probability of A given B, written P(A|B), is

$$P(A|B) = \frac{P(AB)}{P(B)} \tag{2.1}$$

• **Bayes' Rule** Let $A_1, A_2, ...$ be a partition of the sample space, and let B be any set. Then, for each i = 1, 2, 3, ...

$$P(A_i|B) = \frac{P(B|A_i)P(A_i)}{\sum P(B|A_i)P(A_i)}$$
(2.2)

• statistically independent Two events, A and B, are statistically independent if

$$P(AB) = P(A)P(B) \tag{2.3}$$

- random variable A random variable is a function from a sample space $\mathcal S$ into the real numbers
- **identically distributed** The random variables *X* and *Y* are identically distributed if, for every set $A \in \mathcal{B}^1$, $P(X \in A) = P(Y \in A)$.
- cdf cumulative distribution function.
- pmf probability mass function.
- pdf probability density function.
- **moment/central moment** For each integer *n*, the *n*th moment of *X*, μ'_n , is

$$\mu_n' = \mathbb{E}X^n \tag{2.4}$$

The *n*th central moment of X, μ_n , is

$$\mu_n = \mathbb{E}(X - \mu)^n \tag{2.5}$$

where $\mu = \mu'_1 = \mathbb{E}X$.

• **variance** The variance of a random variable X is its second central moment, $VarX = \mathbb{E}(X - \mathbb{E}X)^2$. The positive square root of VarX is the standard deviation of X.

- n-dimensional random vector An n-dimensional random vector is a function form a sample space S into \mathfrak{R}^n , n-dimensional Euclidean space.
- joint pdf / joint pmf
- · conditional pmf

2.1 Frequentist statistics VS Bayesian statistics

Throughout our subsequent discussions, we viewed θ as an unknown parameter of the world. This view of the θ as being constant-valued but unknown is taken in frequentist statistics. In the frequentist this view of the world, θ is not random—it just happens to be unknown—and it's our job to come up with statistical procedures (such as maximum likelihood) to try to estimate this parameter.

An alternative way to approach our parameter estimation problems is to take the Bayesian view of the world, and think of θ as being a random variable whose value is unknown. In this approach, we would specify a prior distribution $p(\theta)$ on θ that expresses our "prior beliefs" about the parameters. Given a training set $S = (x_i, y_i)$, make a prediction on a new value of x, we can then compute the posterior distribution on the parameters.(CS229)

2.2 Mean

- 若干个随机变量之和的期望等于各变量的期望之和, $\mathbb{E}(X_1+X_2+...+X_n)=\mathbb{E}(X_1)+E(X_2)+...+\mathbb{E}(X_n)$
- 若干个独立随机变量之和的期望等于各变量的期望之和, $\mathbb{E}(X_1X_2...X_n) = \mathbb{E}(X_1)\mathbb{E}(X_2)...\mathbb{E}(X_n)$

2.3 Variance

均匀分布 $X \sim R(a,b)$ 其期望为

$$\mathbb{E}(X) = \int_{a}^{b} x f(x) dx$$

$$= \frac{1}{b-a} \in t_{a}^{b} x dx$$

$$= \frac{1}{b-a} \frac{1}{2} (b^{2} - a^{2})$$

$$= \frac{1}{2} (b+a)$$
(2.6)

其方差为

$$Var(X) = \mathbb{E}(X - \mathbb{E}X)^{2}$$

$$= \mathbb{E}(X)^{2} - (\mathbb{E}X)^{2}$$

$$= \int_{a}^{b} x^{2} f(x) dx - (\mathbb{E}X)^{2}$$

$$= \frac{1}{3(b-a)} (b^{3} - a^{3}) - \frac{1}{4} (b+a)^{2}$$

$$= \frac{1}{12} (b-a)^{2}$$
(2.7)

对于正态分布 $\mathcal{N}(\mu, \sigma^2)$, 其期望 $\mathbb{E}(X) = \mu$, 方差为 $Var(X) = \sigma^2$.

2.4 Best Unbiased Estimators

2.5 Divergence

In statistics and information geometry, **divergence** or a **contrast function** is a function which establishes the "distance" of one probability distribution to the other on a statistical manifold. The divergence is a weaker notion than that of the distance, *in particular the divergence need not be symmetric*, and need not satisfy the triangle inequality. [Wikipedia: Divergence (statistics)]

Suppose $\mathcal S$ is a space of all probability distributions with common support. Then a divergence on $\mathcal S$ is a function $D(\cdot\|\cdot):\mathcal S\times\mathcal S\to R$ satisfying

$$D(p||q) \ge 0$$
 for all $p, q \in S$,
 $D(p||q) = 0$ if and only if $p = q$, (2.8)

The two most important divergences are the **KL divergence** (relative entropy), which is central to information theory and statistics, and the **squared Euclidean distance** (SED).

The two most important classes of divergences are the f-divergences and Bregman divergences; The only divergence that is both an f-divergence and a Bregman divergence is the KL divergence; the SED is a Bregman divergence, but not an f-divergence.

2.5.1 *f*-divergence

An f-divergence is a function $D_f(P||Q)$ that measures the difference between two probability distributions P and Q. It helps the intuition to think of the divergence as an average, weighted by the function f, of the odds ratio given by P and Q. [Wikipedia: f-divergence]

Definition

Let P and Q be two probability over a space Ω such that P is absolutely continuous with respect to Q. Then, for a convex function f such that f(1) = 0, the f-divergence of P from Q is defined as

$$D_f(P||Q) = \int_{\Omega} f\left(\frac{\mathrm{d}P}{\mathrm{d}Q}\right) \mathrm{d}Q \tag{2.9}$$

If P and Q are both absolutely continuous with respect to a reference distribution μ on Ω then their probability densities p and q satisfy $dP = pd\mu$ and $dQ = qd\mu$. In this case the f-divergence can be written as

$$D_f(P||Q) = \int_{\Omega} f\left(\frac{p(x)}{q(x)}\right) q(x) d\mu(x)$$
(2.10)

Instances of *f*-divergences

Divergence	$f(\cdot)$
KL-divergence	$t \log t$
reverse KL-Divergence	$-\log t$
Jensen-Shannon Divergence	$(t+1)\log(\frac{2}{t+1}) + t\log t$
squared Hellinger distance	$(\sqrt{t}-1)^2, 2(1-\sqrt{t})$
Total variation distance	$\frac{1}{2} t-1 $
Pearson \mathcal{X}^2 -divergence	$(t-1)^{\overline{2}}, t^2-1, t^2-t$
Neyman \mathcal{X}^2 -divergence(reverse Pearson)	$\frac{1}{t} - 1, \frac{1}{t} - t$

Properties

• **Non-negativity**: the *f*-divergence is always positive; it's zero if and only if the measures *P* and *Q* coincide. This follows immediately from Jensen's inequlity:

$$D_f(P||Q) = \int f\left(\frac{\mathrm{d}P}{\mathrm{d}Q}\right) \mathrm{d}Q \ge f\left(\frac{\mathrm{d}P}{\mathrm{d}Q}\mathrm{d}Q\right) = f(1) = 0 \tag{2.11}$$

- Monotonicity
- Joint Convexity

2.5.2 Kullback-Leibler divergence / Relative entropy

KLD is a measure of how one probability distribution is different from a second, reference probability distribution.

Consider two probability distribution P and Q. Usually, P reresents the data, the observations, or a probability distribution precisely measured. Q represents instead a theory, a model, a description or an approximation of P. The KL divergence is then interperted as the average difference of the number of bits required for encoding samples of P using a code optimized for Q rather than one optimized for P

Definition

$$D_{KL}(P||Q) = \mathbb{E}_{x \sim P(x)} \left[\log \frac{P(x)}{Q(x)}\right]$$

$$= \mathbb{E}_{x \sim P(x)} \left[\log P(x) - \log Q(x)\right]$$
(2.12)

2.6 Entropy

2.6.1 Informational value

The basic idea of information theory is that the "informational value" of a communicated message depends on the degree to which the content of the message is surprising. If an event is very probable, it is no surprise when that event happens as expected; hence transmission of such a message carries very little new information. However, if an event is unlikely to occur, it is much more informative to learn that the event happened or will happen.

The information content (also called the surprisal) of an event E is a function which decreases the probability p(E) of an event increases, defined by $I(E) = -\log p(E)$.

2.6.2 Entropy

In information theory, the entropy of a random variable is the average level of "information", "surprise", or "uncertainty" inherent in the variable's possible outcomes.

$$H(x) = -\mathbb{E}[\log P(x)] \tag{2.13}$$

2.6.3 Cross Entropy

The cross entropy between two probability distributions measures the average number of bits needed to identify an event from a set of possibilities, if a coding scheme is used based on a given probability distribution Q, rather than the "true" distribution P. The cross entropy for two distributions P and Q over the same probability space is thus defined as follows

$$H(P,Q) = \mathbb{E}_{x \sim P(x)}[-\log Q(x)] = H(P(x)) + KL(P(x)||Q(x))$$
(2.14)

When H(P) is constant, minimizing the H(P,Q) is equivalent to minimizing the KL divergence KL(P||Q).

2.6.4 Conditional Entropy

In information theory, the conditional entropy quantifies the amount of information needed to describe the outcome of a random variable Y given that the value of another random variable X is known.

$$H(Y|X) = -\sum P(x,y) \log \frac{P(x,y)}{P(x)}$$
(2.15)

2.7 Mutual Information

In probability theory and information theory, the mutual information (MI) of two random variables is a measure of the **mutual dependence between the two variables**. More specifically, it quantifies the "amount of information" obtained about one random variable through observing the other random variable. The concept of mutual information is intimately linked to that of entropy of a random variable, a fundamental notion in information theory that quantifies the expected "amount of information" held in a random variable.

$$I(X;Y) = KL(P(X,Y)||P(X)P(Y))$$

$$= \mathbb{E}_{x}[KL(P(Y|X)||P(Y))]$$

$$= \mathbb{E}_{y}[KL(P(X|Y)||P(X))]$$

$$= H(X) - H(X|Y)$$

$$= H(Y) - H(Y|X)$$

$$= H(X) + H(Y) - H(X,Y)$$

$$= H(X,Y) - H(X|Y) - H(Y|X)$$

$$= \sum_{y} \sum_{x} P(x,y) \log \frac{P(x,y)}{P(x)P(y)}$$
(2.16)

2.8 Maximum likelihood estimation

Frequentist statistics

深度学习来就是用模型 $Q(x;\theta)$ 来估计数据的真实分布P(x),对于一组确定的数据集X,在样本已被观察到的情况下,需要找到使得 $Q(X;\theta)$ 出现可能性最大的一组参数 θ ,也就是最大似然估计:

$$\begin{split} \hat{\theta}_{MLE} &= \arg\max_{\theta} Q(X; \theta) \\ &= \arg\max_{\theta} \prod_{i=1}^{m} Q(x^{i}; \theta) \\ &\Rightarrow \arg\max_{\theta} \sum_{i=0}^{m} \log Q(x^{i}; \theta) \\ &\Rightarrow \arg\min_{\theta} - \sum_{i=0}^{m} \log Q(x^{i}; \theta) \\ &\Rightarrow \arg\min_{\theta} - \mathbb{E}_{x \sim P(x)} [\log Q(x; \theta)] \end{split}$$
 (2.17)

2.8.1 MLE & KLD

如果使用KLD来衡量模型 $Q(x;\theta)$ 和真实分布P(x)之间的差距,其中 $\mathbb{E}_{x\sim P(x)}\log P(x)$ 为常量,则有

$$\arg\min KL(P(x)||Q(x;\theta)) = \arg\min_{\theta} \mathbb{E}_{x \sim P(x)}[\log P(x) - \log Q(x;\theta)]$$

$$\Rightarrow \arg\min_{\theta} -\mathbb{E}_{x \sim P(x)}[\log Q(x;\theta)]$$
(2.18)

在估计真实分布的情况下,计算KLD和使用MLE是等价的。

2.9 When P(x) is constant

When P(x) is contant, these equtions are equivalent.

•
$$KLD = KL(P(x)||Q(x;\theta)) = \mathbb{E}_{x \sim P(x)}[\log P(x) - \log Q(x;\theta)] = -\mathbb{E}_{x \sim P(x)}[\log Q(x;\theta)]$$

•
$$CE = H(P,Q) = \mathbb{E}_{x \sim P(x)}[-\log Q(x)] = KL(P(x)||Q(x)) = -\mathbb{E}_{x \sim P(x)}[\log Q(x;\theta)]$$

•
$$MLE = \arg\min -\mathbb{E}_{x \sim P(x)}[\log Q(x; \theta)]$$

2.10 Maximum A Posteriori

 $P(\theta)$ 先验概率分布, $P(X|\theta)$ 是似然函数,根据贝叶斯定理,可用以下公式计算后验概率

$$P(\theta|X) = \frac{P(X|\theta)P(\theta)}{P(X)}$$
 (2.19)

模型估计时,估计整个后验概率分布 $P(\theta|X)$,如需给出一个模型,通常取后验概率最大的模型。

$$\hat{\theta}_{MAP} = \arg \max P(\theta|X)$$

$$= \arg \min - \log P(\theta|X)$$

$$= \arg \min - \log P(X|\theta) - \log P(\theta) + \log P(X)$$

$$= \arg \min - \log P(X|\theta) - \log P(\theta)$$
(2.20)

 $\log P(X)$ 与 θ 无关,可以丢掉。 $-\log P(X|\theta)$ 其实就是NLL,所以MLE和MAP不同在 $-\log P(\theta)$ 。假定先验是一个高斯分布

$$P(\theta) = C \times e^{-\frac{\theta^2}{2\sigma^2}} \tag{2.21}$$

那么

$$\log P(\theta) = C + \frac{\theta^2}{2\sigma^2} \tag{2.22}$$

在MAP中,使用一个高斯分布的先验等价于在MLE中采用 L^2 的正则化。MAP贝叶斯推断提供了一个直观的方法来设计复杂但可解释的正则化,更复杂的惩罚项可以通过混合高斯分布作为先验得到,而不是一个单独的高斯分布。

预测新的观察数据x时,计算数据对后验概率分布的期望值:

$$p(x|X) = \int p(x|\theta, X)p(\theta|X)d\theta$$
 (2.23)

Basic

3.1 Mechine Learning

- supervised learning
- semi-supervised learning
- unsupervised learning
- reinforcement learning
- · active learning
- transfer learning

3.2 iid

3.3 Convolution

3.4 Transposed Convolution

```
H_{out} = (H_{in} - 1) \times \text{stride}[0] - 2 \times \text{padding}[0] + \text{dilation}[0] \times (\text{kernel\_size}[0] - 1) + \text{output\_padding}[0] + 1
W_{out} = (W_{in} - 1) \times \text{stride}[1] - 2 \times \text{padding}[1] + \text{dilation}[1] \times (\text{kernel\_size}[1] - 1) + \text{output\_padding}[1] + 1
(3.1)
```

3.5 Universal approximation theorem

Any continuous function can be uniformly approximated by a continuous neural network having only one internal, hidden layer and with an arbitrary continuous sigmoidal nonlinearity.[2]

万能近似定理最初以sigmoidal激活函数来描述,后被证明对于更广泛的激活函数也适用[7],包括ReLU。

3.6 Initialization scheme

3.6.1 constant initialization

3.6.2 random initialization

按照某一分布随机初始化

normal initialization

$$W \sim \mathcal{N}(\mu, \sigma^2) \tag{3.2}$$

uniform initalization

$$W \sim U[-\frac{1}{\sqrt{n}}, \frac{1}{\sqrt{n}}] \tag{3.3}$$

3.6.3 xavier initalization

针对使用**对称激活函数**tanh(x)的网络进行参数初始化,对于ReLU激活函数并不适用[3].

Forward

对于一个卷积层来说

$$y_{l} = W_{l}X_{l} + b_{l}$$

$$X_{l} = f(y_{l-1})$$
(3.4)

在如下前提和假设下

- 初始化W₁元素为独立同分布
- 假设X₁元素也为独立同分布
- W₁, X₁互相独立

则有

$$Var(y_{l}) = Var(\sum W_{l}X_{l} + b_{l})$$

$$= Var(\sum W_{l}X_{l})$$

$$= n_{l}Var(W_{l}X_{l})$$
(3.5)

令 W_l 期望为0, $E(W_l) = 0$, $Var(W_l) = E(W_l - E(W_l))^2 = EW_l^2$,则

$$Var(y_{l}) = n_{l}E(W_{l}^{2}X_{l}^{2}) - n_{l}E^{2}W_{l}E^{2}X_{l}$$

$$= n_{l}E(W_{l}^{2}X_{l}^{2})$$

$$= n_{l}Var(W_{l})E(X_{l}^{2})$$
(3.6)

若 $EX_l=0$,则

$$Var(y_l) = n_l Var(W_l) Var(X_l)$$
(3.7)

若要实现 $Var(y_l) = Var(X_l)$, 则需要满足 $n_l Var(W_l) = 1$, 即

$$Var(W_l) = \frac{1}{n_l} \tag{3.8}$$

- 若 W_l 服从正态分布,则 $W_l \sim N(0, \frac{1}{n_l})$
- 若 W_l 服从均匀分布,则 $W_l \sim U(-\sqrt{\frac{3}{n_l}},\sqrt{\frac{3}{n_l}})$

Backword

反向传播过程中,需要保证梯度的方差不变,每一层的梯度为:

$$\Delta X_l = \hat{W}_l \Delta y_l \tag{3.9}$$

假设

- W_l和Δy_l互相独立
- $EW_l = 0, E\Delta X_l = 0$

同前向传播,可得

$$Var(\Delta X_l) = \hat{n}_l Var(W_l) Var(\Delta y_l)$$
(3.10)

- 若 W_l 服从正态分布,则 $W_l \sim N(0, \frac{1}{n})$
- 若 W_l 服从均匀分布,则 $W_l \sim U(-\sqrt{\frac{3}{\hat{n}_l}},\sqrt{\frac{3}{\hat{n}_l}})$

除非 $n=\hat{n}_l$,否则同时保证信号在向前向后传播时的Var不变,取调和平均数, $Var(W_l)=\frac{2}{n_l+\hat{n}_l}$,可得

- Normal distribution: $W_l \sim \mathcal{N}(0, \frac{2}{n_l + \hat{n}_l})$
- Uniform distribution: $W_l \sim \mathcal{U}(-\sqrt{\frac{6}{n_l+\hat{n}_l}},\sqrt{\frac{6}{n_l+\hat{n}_l}})$

3.6.4 orthogonal initalization

3.6.5 kaiming initalization

Xavier 针对对称激活函数的层权重初始化进行设计,但对于使用ReLu激活函数的层并不适用[4]。 -> 期望为0,方差为1

对ReLU层来说, $E(X_i^2) = \frac{1}{2}Var(y_i)$,因此

$$Var(y_l) = \frac{1}{2} n_l Var(W_l) Var(X_l)$$

$$Var(\Delta X_l) = \frac{1}{2} \hat{n}_l Var(W_l) Var(\Delta y_l)$$
(3.11)

且在权重初始化时,使用上述任意一个即可。

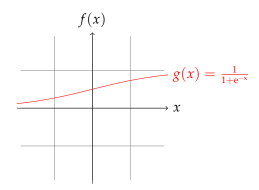
3.7 Regularization

3.8 Data proprocessing

3.8.1 whitening

3.9 Activation functions

3.9.1 sigmoid



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

$$g(x)' = g(x)(1 - g(x))$$
(3.12)

$3.9.2 \quad tanh(x)$

$$f(x)$$

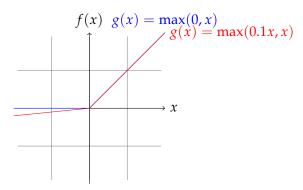
$$g(x) = \tanh(x)$$

$$g(x) = tanh(x)$$

$$g(x)' = 1 - g(x)^{2}$$

$$= 1 - tanh(x)^{2}$$
(3.13)

3.9.3 Rectified linear units



ReLU

$$g(x) = \max(0, x) \tag{3.14}$$

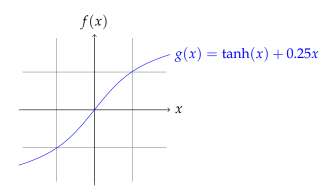
Leaky ReLU

$$g(x) = \max(0.01x, x) \tag{3.15}$$

PReLU

$$g(x) = \max(\alpha x, x) \tag{3.16}$$

3.9.4 haha



$$g(x) = \tanh(x) + 0.25x$$

$$g(x)' = 0.75 - g(x)^{2}$$

$$= 0.75 - \tanh(x)^{2}$$
(3.17)

3.9.5 Loss functions

3.10 Batch Normalization

机器学习中假设:源空间和目标空间的数据分布是一致的,如果不一致,就是新的机器学习问题,例如transfer learning/domain adaptation 等。而covariate shift就是分布不一致假设下的一个分支问题,它指的是源空间和目标空间的条件概率是一致的,但是其边缘概率不同.对于所有 $x \in \chi$

$$P_s(Y|X=x) = P_t(Y|X=x)$$

$$P_s(X) \neq P_t(X)$$
(3.18)

Internal Covariate Shift Training Deep Neural Networks is complicated by the fact that the distribution of each layer's inputs changes during training, as the parameters of the previous layers change. And this slows down the training by requring lower lr ant careful parameter initialization, and makes it notoriously hard to train models with saturating nonlinearities.

解决方案: 白化操作可以使得数据变为独立同分布

By whitening the inputs to each layer, we would take a step towards achieving the fixed distribution of inputs that would remove the ill effects of the ICS.

但是每层的白化会影响梯度下降优化过程。每层进行白化计算量过大。保证模型的表达能力不因为规范化而下降。

Simply normalizaing each input of a layer may change what the layer can represent. So, we make sure that the transformation inserted in the network can represent the identity transform.

Input: Values of
$$x$$
 over a mini-batch: $\mathcal{B} = \{x_{1...m}\}$;

Parameters to be learned: γ , β

Output: $\{y_i = \mathrm{BN}_{\gamma,\beta}(x_i)\}$

$$\mu_{\mathcal{B}} \leftarrow \frac{1}{m} \sum_{i=1}^m x_i \qquad \text{// mini-batch mean}$$

$$\sigma_{\mathcal{B}}^2 \leftarrow \frac{1}{m} \sum_{i=1}^m (x_i - \mu_{\mathcal{B}})^2 \qquad \text{// mini-batch variance}$$

$$\widehat{x}_i \leftarrow \frac{x_i - \mu_{\mathcal{B}}}{\sqrt{\sigma_{\mathcal{B}}^2 + \epsilon}} \qquad \text{// normalize}$$

$$y_i \leftarrow \gamma \widehat{x}_i + \beta \equiv \mathrm{BN}_{\gamma,\beta}(x_i) \qquad \text{// scale and shift}$$

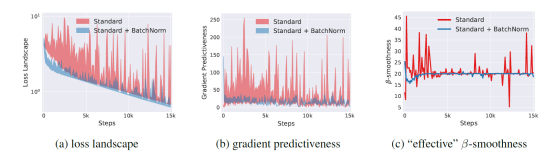
Algorithm 1: Batch Normalizing Transform, applied to activation x over a mini-batch.

图 3.1: Batch Normalization

3.11 How Does Batch Normalization Help Optimization?

Batch Normalization 对减少ICS起到了很少的作用,it makes the optimization landscape significantly smoother.

- Is the effectiveness of BatchNorm indeed related to internal covariate shift? The 'noisy' BatchNorm network has qualitatively less stable distributions than even the standard, non-BatchNoram network, yet it still better performs better in terms of training.
- **Is BatchNorm reducing internal covariate shift?** We Observe that networks with BatchNorm often exhibit an increase in ICS.



Why does BatchNorm work?

BatchNorm reparametrizes the underlying optimization problem to make its landscape significatly more smooth. The key implication of BatchNorm's reparametrization is that it makes the gradients more reliable and predictive. After all, improved Lipschitzness of the gradients gives us confidence that when we take a larger step in a direction of a computed gradient, this gradient direction remains a fairly accurate estimate of the actual gradient direction after taking that step.

Is BatchNorm the best way to smoothen the landscape?

In fact, for deep linear networks, l_1 -normalization performs even better than BatchNorm. Note that, qualitatively, the l_p -normalization techniques lead to larger distributional shift than the vanilla, i.e., unnormalized, networks, yet they still yield improved optimization performance.

Optimization Algorithms

4.1 Challenges

4.1.1 Local Minima

When the numerical solution of an optimization problem is near the local optimum, the numerical solution obtained by the final iteration may only minimize the objective function locally, rather than globally, as the gradient of the objective function's solutions approaches or becomes zero. Only some degree of noise might knock the parameter out of the local minimium. In fact, this is the one of the beneficial properties of stochastic gradient descent where the natural variation of gradients over minibatches is able to dislodge the parameters from loacl minima.[10]

4.1.2 Saddle Points

We assume that the input of a function is *k*-dimensional vector and its ouput is a scalar, so its *Hession matrix* will have *k eigenvalues*. The solution of the function could be a local minimum, a local maximum or a saddle point at a position where the function gradient is zero:

- When the eigenvalues of the function's Hession matrix at the zero-gradient position are all positive, we have a local minimum for the function.
- When the eigenvalues of the function's Hession matrix at the zero-gradient position are all negative, we have a local maximum for the function.
- When the eigenvalues of the function's Hession matrix at the zero-gradient position are negative and positive, we have a saddle point for the function.

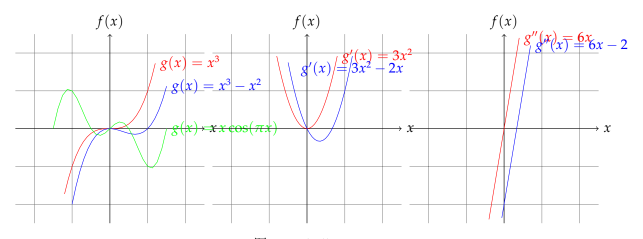


图 4.1: Challenges

• 同号目至少有一个为0,不确定

For high-dimensional problem the likehood that at least some of the eigenvalues are negative is quite high. This makes sadlle points are more likely then local minima.[10]

4.1.3 Vanishing gradients

Vanishing gradients can cause optimization to stall. Often a reparameterization of the problem helps. Good initialization of the parameters can be beneficial, too.[10]

4.1.4 noise / noise-free

noise对优化过程中saddle point的影响? what is nosie? and how it effect the optimization. 无偏估计、一致性

4.2 Gradient Descent

Using a Taylor expansion we obtain that:

$$f(x+\epsilon) = f(x) + \epsilon f'(x) + \mathcal{O}(\epsilon^2) \tag{4.1}$$

For small ϵ moving in the direction of negative gradient will decrease f. Choose $\epsilon = -\eta f'(x)$, $\eta > 0$, then we get:

$$f(x - \eta f'(x)) = f(x) - \eta f'^{2}(x) + \mathcal{O}((\eta f'(x))^{2})$$
(4.2)

Choose η samll enough for the higher order terms to become irrelevant. then

$$f(x - \eta f'(x)) \lesssim f(x) \tag{4.3}$$

that means, if we use

$$x \leftarrow x - \eta f'(x) \tag{4.4}$$

to iterate x, the value of f(x) decline.

4.2.1 Adaptive Methods

Getting the learning rate 'just right' is tricky. What if we could determine η automatically or get rid of having to select a setp size at all? Second order methods that look not only at the value and gradient of the objective but alse at its *curvature* can help in this case. These methods cannot be applied to deep learning directly due to the computational cost.

Newton's Method

当Taylor expansion展开到二阶导数时:

$$f(\mathbf{x} + \epsilon) = f(\mathbf{x}) + \epsilon^{\top} \nabla f(\mathbf{x}) + \frac{1}{2} \epsilon^{\top} H_f \epsilon + \mathcal{O}(\|\epsilon\|^3)$$
(4.5)

we define $H_f := \nabla \nabla^\top f(\mathbf{x})$ to be the *Hession* of f. H is a $d \times d$ matrix and may be prohibitively large, due to the cost of storing $\mathcal{O}(d^2)$ entries.

After all, the minimum of f statifies $\nabla f(\mathbf{x}) = 0$. Taking derivatives of above equtaion with regard to η and ignoring higher order terms we arrive at

$$\nabla f(\mathbf{x}) + H_f \epsilon = 0$$

$$\epsilon = -H_f^{-1} \nabla f(\mathbf{x})$$
(4.6)

then, $\epsilon = -\eta \nabla f(\mathbf{x})$, we get

$$\eta = -H_f^{-1} \tag{4.7}$$

For
$$f(x) = (x-2)(x-4) = x^2 - 6x + 8$$
, $f'(x) = 2x - 6$, $f''(x) = 2$, then
$$\epsilon = -f''(0)^{-1}f'(0) = -\frac{1}{2} \times -6 = 3 \tag{4.8}$$

Hessian

Hession 矩阵是是对称的,可以表示为一组特征值和一组特征向量的正交基底,在特定方向上g上的二阶导数为g^THg,当g为特征向量时,这个二阶导数就是对应的特征值。最大特征值确定最大二阶导数,最小特征值确定最小导数。

在g方向上的learning rate为

$$\eta_g = \frac{1}{g^\top H_f g} \tag{4.9}$$

最差的情况下,g与H最大的特征值 λ_{max} 对应的特征向量对齐,此时的最优步长为 $\frac{1}{\lambda_{max}}$,当要最小化的目标函数能用二次函数很好近似的情况下,Hession的特征值决定了学习率的量级。

4.2.2 Stochastic Gradient Descent

Dynamic Learning rate

$$\eta(t) = \eta_i \text{ if } t_i \leq t \leq t_{i+1} \quad \text{piecewise constant}$$

$$\eta(t) = \eta_0 e^{-\lambda t} \quad \text{exponential}$$

$$\eta(t) = \eta_0 (\beta t + 1)^{-\alpha} \quad \text{polynomial}$$
(4.10)

In the case of convex optimization there are a number of proofs which show that this rate is well behaved.

4.2.3 Minibatch Stochastic Gradient Descent

At each iteration of stochastic gradient descent, we uniformly sample an index $i \in i, ..., n$ for data examples at random, and compute the gradient $\nabla f_i(\mathbf{x})$ to update \mathbf{x} :

$$\mathbf{x} \leftarrow \mathbf{x} - \eta \nabla f_i(\mathbf{x}) \tag{4.11}$$

The stochastic gradient $\nabla f_i(\mathbf{x})$ is the *unbiased estimate* of gradient $\nabla f(\mathbf{x})$. That means the stochastic gradient is a good estimate of the gradient.

$$\mathbb{E}_{i}\nabla f_{i}(\mathbf{x}) = \frac{1}{n}\sum_{i=1}^{n}\nabla f_{i}(\mathbf{x}) = \nabla f(\mathbf{x})$$
(4.12)

4.3 SGDM - SGD with Momentum

$$\mathbf{v}_{t} \leftarrow \beta \mathbf{v}_{t-1} + \nabla f(\mathbf{x}) \mathbf{x}_{t} \leftarrow \mathbf{x}_{t-1} - \eta_{t} \mathbf{v}_{t}$$

$$(4.13)$$

v is called *momentum*. Momentum replaces gradients with a leaky average over past gradients. This accelerates convergence significantly. And it prevents stalling of the optimization process that is much more likely to occur for stochastic gradient descent.

An Ill-conditioned Problem

$$f(\mathbf{x}) = 0.1x_1^2 + 2x_2^2 \tag{4.14}$$

and we know that f has minimum at (0,0). Then we get

$$\nabla f(\mathbf{x}) = \begin{bmatrix} 0.2x_1 \\ 4x_2 \end{bmatrix} \tag{4.15}$$

let's start at (2,2), $\eta = 0.4$, $\beta = 0.6$.

4.4 NAG - Nesterov Accelerated Gradient

$$\mathbf{v}_{t} = \alpha \mathbf{v}_{t-1} + \eta \nabla_{\theta} J(\theta - \alpha \mathbf{v}_{t-1})$$

$$\theta = \theta - \mathbf{v}_{t}$$
(4.16)

4.5 Adagrad

Consider a model training on spare features, i.e., features that occur only infrequently. Parameters associated with infrequent features only receive meaningful updates whenever these features occur. And to get good accuracy we want to decrease the lr as well keep on training, usually at a lr of $\mathcal{O}(t^{-\frac{1}{2}})$. we might end up in situation where the parameters from common features converge rather quickly to their optimal values.

$$\mathbf{g}_{t} = \partial_{\mathbf{w}} l(y_{t}, f(\mathbf{x}_{t}, \mathbf{w}))$$

$$\mathbf{s}_{t} = \mathbf{s}_{t-1} + \mathbf{g}_{t}^{2}$$

$$\mathbf{w}_{t} = \mathbf{w}_{t-1} - \frac{\eta}{\sqrt{\mathbf{s}_{t} + \epsilon}} \cdot \mathbf{g}_{t}$$

$$(4.17)$$

learning rate不再简单的是时间的函数,而是和梯度相关,lr的减小速度和梯度正相关,每个维度都有自己的learning rate.

- It uses the magnitude of the gradient as a means of adjusting how quickly progress is achieved coordinates with large gradients are compensated with a smaller learning rate.
- On deep learning problems Adagrad can sometimes be too aggressive in reducing learning rates

4.6 RMSProp

A variant of AdaGrad.

As a result \mathbf{s}_t keeps on growing without bound due to the lack of normalization, essentially linearly as the algorithm converges.

$$\mathbf{s}_{t} \leftarrow \gamma \mathbf{s}_{t-1} + (1 - \gamma) \mathbf{g}_{t}^{2}$$

$$\mathbf{x}_{t} \leftarrow \mathbf{x}_{t-1} - \frac{\eta}{\sqrt{\mathbf{s}_{t} + \epsilon}} \odot \mathbf{g}_{t}$$

$$(4.18)$$

$$\mathbf{s}_{t} = (1 - \gamma)\mathbf{g}_{t}^{2} + \gamma\mathbf{s}_{t-1}$$

$$= (1 - \gamma)(\mathbf{g}_{t}^{2} + \gamma\mathbf{g}_{t-1}^{2} + \gamma^{2}\mathbf{g}_{t-2}^{2} + ...)$$
(4.19)

4.7 Adadelta

A variant of AdaGrad.

$$\mathbf{s}_{t} = \rho \mathbf{s}_{t-1} + (1 - \rho) \mathbf{g}_{t}^{2}$$

$$\mathbf{g}_{t}' = \frac{\sqrt{\Delta \mathbf{x}_{t-1} + \epsilon}}{\sqrt{\mathbf{s}_{t} + \epsilon}} \odot \mathbf{g}_{t}$$

$$\Delta \mathbf{x}_{t} = \rho \Delta \mathbf{x}_{t-1} + (1 - \rho) \mathbf{g}_{t}'^{2}$$

$$\mathbf{x}_{t} = \mathbf{x}_{t-1} - \mathbf{g}_{t}'$$

$$(4.20)$$

4.8 Adam

$$\mathbf{v}_{t} \leftarrow \beta_{1} \mathbf{v}_{t-1} + (1 - \beta_{1}) \mathbf{g}_{t}$$

$$\mathbf{s}_{t} \leftarrow \beta_{2} \mathbf{s}_{t-1} + (1 - \beta_{2}) \mathbf{g}_{t}^{2}$$

$$(4.21)$$

Common choices for them are $\beta_1=0.9$ and $\beta_2=0.9999$. That is, the variance estimate moves *much more slowly* than the momentum term. Note that if we initialize $\mathbf{v}_0=\mathbf{s}_0=0$ we have a significant amount of bias initally towards smaller values. This can be addressed by using the fact that $\sum_{i=0}^t = \frac{1-\beta^t}{1-\beta}$ to re-normalize terms.

$$\hat{\mathbf{v}}_{t} = \frac{\mathbf{v}_{t}}{1 - \beta_{1}^{t}}
\hat{\mathbf{s}}_{t} = \frac{\mathbf{s}_{t}}{1 - \beta_{2}^{t}}
\mathbf{x}_{t} \leftarrow \mathbf{x}_{t-1} - \frac{\eta \hat{\mathbf{v}}_{t}}{\sqrt{\hat{\mathbf{s}}_{t}} + \epsilon}$$
(4.22)

- 4.9 AdaMax
- 4.10 Nadam
- 4.11 AMSGrad

Models

Skip Connections

6.1 Residual Units

为了训练deeper model,并解决梯度消失/爆炸以及网络degradation problem,在ResNets[5]中提出,如果在shallower model 中添加identity mapping,那么deeper model理应不会有更高的错误率。

Residual unit表示如下:

$$\mathbf{y}_{l} = h(\mathbf{x}_{l}) + \mathcal{F}(\mathbf{x}_{l}, \mathcal{W}_{l})$$

$$\mathbf{x}_{l+1} = f(\mathbf{y}_{l})$$
(6.1)

In ResNet, $h(\mathbf{x}) = \mathbf{x}_l$ is an identity mapping and f is a ReLU function.

在ResNet中,因为f不是identity mapping,所以残差只能ResNet units中学习且信息不能直接 传达到后面的层中, In [6],希望*propagating information*可以*through theentire network*.

Our derivations reveal that *if both* $h(\mathbf{x}_l)$ *and* $f(\mathbf{y}_l)$ *are identity mappings*, the signal could be *directly* propagated from one unit to any other units, in both forward and backward passes.[6]

if f is also an identity mapping:

$$\mathbf{x}_{l+1} = \mathbf{x}_l + \mathcal{F}(\mathbf{x}_l, \mathbf{W}_l) \tag{6.2}$$

then we will have:

$$\mathbf{x}_{l+n} = \mathbf{x}_{l+n-1} + \mathcal{F}(\mathbf{x}_{l+n-1}, \mathcal{W}_{l+n-1})$$

$$= \mathbf{x}_{l+n-2} + \mathcal{F}(\mathbf{x}_{l+n-2}, \mathcal{W}_{l+n-2}) + \mathcal{F}(\mathbf{x}_{l+n-1}, \mathcal{W}_{l+n-1})$$

$$= \mathbf{x}_{l} + \sum_{i=1}^{l+n-1} \mathcal{F}(\mathbf{x}_{i}, \mathcal{W}_{i})$$
(6.3)

相较于plain network(ignoring BN and ReLU is an identity mapping):

$$\mathbf{x}_{l+n} = \prod_{i=l}^{l+n-1} \mathbf{W}_i \mathbf{x}_i \tag{6.4}$$

反向传播过程:

$$\frac{\partial \mathbb{E}}{\partial \mathbf{x}_{l}} = \frac{\partial \mathbb{E}}{\partial \mathbf{x}_{l+n}} \frac{\partial \mathbf{x}_{l+n}}{\partial \mathbf{x}_{l}}$$

$$= \frac{\partial \mathbb{E}}{\partial \mathbf{x}_{l+n}} \left(1 + \frac{\partial}{\partial \mathbf{x}_{l}} \sum_{i=l}^{l+n-1} \mathcal{F}(\mathbf{x}_{i}, \mathcal{W}_{i}) \right)$$
(6.5)

可以看出,上式可以分为两部分, $\frac{\partial \mathbb{E}}{\partial x_{1+n}}$ 不经过任何权重信息,可以直接传播到浅层,只要括号中的第二部分不总是为-1,那么**即使权重任意小也不会发生梯度消失**。

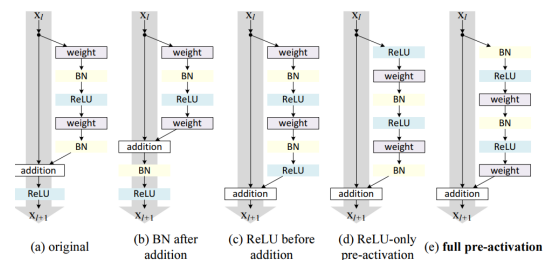


图 6.1: Residual Units

以上的分析基于f是一个恒等变换,但实际上f会影响之前分析的两条信息传播的路径:

$$\mathbf{x}_{l+1} = f(\mathbf{x}_l) + \mathcal{F}(f(\mathbf{x}_l), \mathcal{W}_l)$$
(6.6)

因此,在[6]提出了新的Residual unit结构pre-activation,等式变为

$$\mathbf{x}_{l+1} = \mathbf{x}_l + \mathcal{F}(\hat{f}(\mathbf{x}_l), \mathcal{W}_l) \tag{6.7}$$

6.2 The Shattered Gradients Problem

A previously unnoticed difficulty with gradients in deep rectifier networks that orthogonal to vanishing and exploding gradients. The shattering gradients problem is that, as depth increases, gradients in standard feedforward networks increasingly resemble white noise[1].

- Gradients of shallow networks resemble brown noise(布朗噪声).
- Gradients of deep networks resemble white noise(白噪声).
- Training is difficult when gradients behave like white noise.
- Gradients of deep resnets lie in between brown and white noise.

在标准前馈神经网络中,神经元相关性按指数级减少($\frac{1}{2^L}$),同时,梯度的空间结构也随着深度增加被逐渐消除。使用BatchNorm的ResNets中梯度相关系减少的速度从指数级减少到亚线性级($\frac{1}{\sqrt{(L)}}$),极大的保留梯度的空间结构。

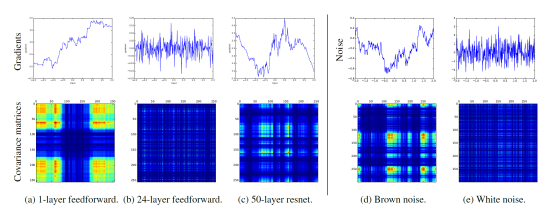


图 6.2: The shattered Gradients Problem

6.3 DenseNet



图 6.3: DenseNet

6.4 FCN

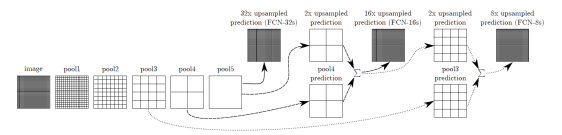


图 6.4: FCN

6.5 UNet family

6.5.1 U-Net

Architecture

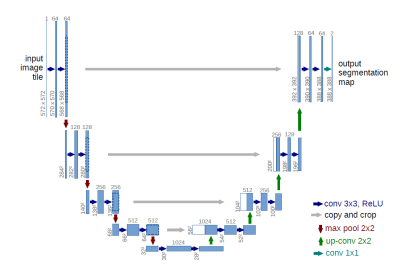


图 6.5: UNet

UNet[9]: The architecture consists of a **contracting path** to capture context and a symmetric **expanding path** that enables precise localization.

 $[offical\ implementation] \verb|http://lmb.informatik.uni-freiburg.de/people/ronneber/u-net|\\$

Metrics

6.5.2 VNet

Dice coeffcient

$$dice = \frac{2|X \cap Y|}{|X| + |Y|}$$

6.5.3 UNet 2+

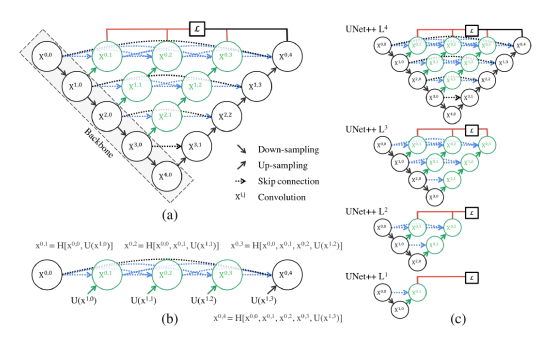


图 6.6: UNet 2+

多深合适?降采样对分割网络到底是不是必须的?不一定要降到第四次才上采样,使用浅层和深层的特征。

6.5.4 UNet 3+

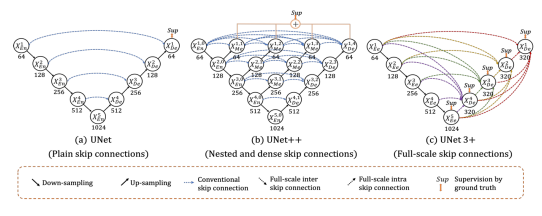


图 6.7: UNet 3+

都缺乏从全尺度探索足够信息的能力,未能明确了解器官的位置和边界。UNet 3+中的每一个解码器层都融合了来自编码器中的小尺度和同尺度的特征图,以及来自解码器的大尺度的特征图,

这些特征图捕获了全尺度下的细粒度语义和粗粒度语义。

Semantic Segmentation

7.1 FCN - Fully Convolutional Network

Autoencoders

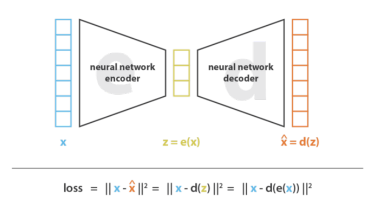


图 8.1: Autoencoder

8.1 Linear Autoencoders VS PCA

8.2 Undercomplete Autoencoders

编码维度小于输入维度。学习欠完备的表示将强制自编码器捕捉训练数据中**最显著的特征**,可用于**降维**。

8.3 Regularized Autoencoders

如果隐藏层的编码的维度允许与输入相等,或隐藏编码维数大于输入的过完备的情况下,会学习将输入复制到输出,而学不到任何有关数据分布的的有用信息。正则自编码器使用的损失函数可以鼓励模型学习其他特性(除了将输入复制到输出),而不必限制使用浅层的编码器和解码器以及小的编码维度来限制模型的容量。这些特性包括稀疏表示、表示的小导数以及对噪声或输入缺失的鲁棒性,即使模型容量大到足以学习一个无意义的恒等函数,非线性且过完备的正则自编码器仍然能够从数据中学到一些关于数据分布的有用信息。

8.3.1 Sparse Autoencoders

$$L(\mathbf{x}, D(E(\mathbf{x}))) + \Omega(h) \tag{8.1}$$

8.3.2 DAE - Denoising Autoencoders

$$L(\mathbf{x}, D(E(\tilde{\mathbf{x}}))) \tag{8.2}$$

 $\hat{\mathbf{x}}$ is a copy of \mathbf{x} that has been corrupted by some form of noise.

8.3.3 CAE - Contractive Autoencoders

$$L(\mathbf{x}, D(E(\mathbf{x}))) + \Omega(\mathbf{h}, \mathbf{x})$$

$$\Omega(\mathbf{h}, \mathbf{x}) = \lambda \sum_{i} \|\nabla_{x} h_{i}\|^{2}$$
(8.3)

Structured Probabilistic Model

- 9.1 Directed Graphical Model / Bayesian Network
- 9.2 Undirected Graphical Model / MRF Markov Random Field

Approximate Inference

Statistical inference is the process of using data analysis to infer properties of an underlying distribution of probability. **Bayesian inference** is a method of statistical inference in which *Bayes' theorem* is used to update the probability for a hypothesis as more evidence or information becomes available.

Bayesian inference derives the **posterior probability** as a consequence of two antecedents: a **prior probability** and a "**likelihood function**" derived from a statistical model for the observed data. Bayesian inference computes the posterior probability according to Bayes' theorem:

$$P(z|x) = \frac{P(x|z)P(z)}{P(x)}$$
(10.1)

- P(x) **evidence**: probability distribution of the observed data independenty from any parameter value or latent variables.
- P(z) **prior**: probability distribution of the parameter or latent variables independently from any observation.
- $P(x \mid z)$ **likelihood**: probability distribution of the observed data given a parameter value or latent variables.
- $P(z \mid x)$ **posterior**: probability distribution of the parameter or latent variables given the observed data.

The computation of the posterior requires three terms: *a prior, a likelihood and an evidence*. The first two can be expressed easily as they are part of the assumed model. But the evdience requires to be computed such that

$$P(x) = \int P(x|z)P(z)dz$$
 (10.2)

It can become intractable in higher dimensions. In this last case, the exact computation of the posterior distribution is practically infeasible and some *approximation techniques* have to be used to get solutions to problems that require to know this posterior, such as **Markov Chain Monte Carlo** and **Variational Inference** methods.

10.1 Monte Carlo Methods

Monte Carlo method, 也称为 statistical simulation method, 是通过从概率模型的随机抽样进行近似数值计算的方法。Markov Chain Monte Carlo 则是以Markov chain为概率模型的Monte carlo Method.

Metropolis-Hastings 算法是最基本的MCMC, Gibbs sampling 是更简单、使用更广泛的MCMC. MC要解决的问题是,假设概率分布已知,通过抽样获得概率分布的随机样本,并通过得到的随机样本对概率分布的特征进行分析. 所以MC的核心是random sampling.

一般的蒙特卡罗法有直接抽样法、接受-拒绝抽样法、重要性抽样法. [8]

10.1.1 Markov Chains

10.2 Variational Inference

The main idea behind variational inference is to first posit a family of densities and then to find the member of that family which is close to the target. Closeness is measured by Kullback-Leibler divergence. Variational inference thus turns the inference problem into an optimization problem.

First, we posit a family of approximate densities Q. This is a set of densities over the latent variables. Then, we try to find the member of that family that minimizes the KL divergence to the exact posterior,

$$Q^*(z) = \arg\min_{Q(z) \in \mathcal{Q}} KL(Q(z) || P(z|x))$$

$$\tag{10.3}$$

Finally, we approximate the posterior with the optimized member of the family $Q^*(z)$. The reach of the family $\mathcal Q$ manages the complexity of this optimization. One of the key ideas behind variational inference is to choos $\mathcal Q$ to be flexible enough to capture a density close to p(z|x), but simple enough for efficient optimization.

However, this objective is not computable because it requires computing the evidence $\log p(x)$

$$KL(Q(z)||P(z|x)) = \mathbb{E}_{z \sim Q}[\log Q(z)] - \mathbb{E}_{z \sim Q}[\log P(z|x)]$$

$$= \mathbb{E}_{z \sim Q}[\log Q(z)] - \mathbb{E}_{z \sim Q}[\log P(z,x)] + \mathbb{E}_{z \sim Q}[\log P(x)]$$

$$= \mathbb{E}_{z \sim Q}[\log Q(z) - \log P(z,x)] + \log P(x)$$
(10.4)

$$\log P(x) = KL(Q(z)||P(z|x)) + \mathbb{E}_{z \sim Q}[\log P(z, x) - \log Q(z)]$$

 $\log P(x)$ is constant with respect to Q(z), $KL(\cdot) \geq 0$, then

$$ELBO(Q) = \mathbb{E}_{z \sim Q}[\log P(z, x) - \log Q(z)] \le \log P(x)$$
(10.5)

The function is call **evdience low bound(ELBO)**. Maximizing the ELBO is equivalent to minimizing the KL divergence.

$$ELBO(Q) = \mathbb{E}_{z \sim Q}[\log P(z, x)] - \mathbb{E}_{z \sim Q}[\log Q(z)]$$

$$= \mathbb{E}_{z \sim Q}[\log P(z)] + \mathbb{E}_{z \sim Q}[\log P(x|z)] - \mathbb{E}_{z \sim Q}[\log Q(z)]$$

$$= \mathbb{E}_{z \sim Q}[\log P(x|z)] - KL(Q(z)||P(z))$$
(10.6)

The first termis an expected likelihood; it encourages densities that place their mass on configurations of the latent variables that explain the observed data. The second term is the negative divergence between the variational density and the prior; it encourages densities close to the prior.

10.2.1 Bayesian mixture of Gaussians

Consider to Bayesian mixture of unit-variance univariate Gaussians. There are K mixture components, corresponding to K Gaussian distributions with means $\mu = \{\mu_1, ..., \mu_k\}$. The mean parameters are drawn independently from a common prior $p(\mu_k)$, which we assume to be as Gaussian $\mathcal{N}(0,\sigma^2)$; the prior variance σ^2 is a hyperparameter. To generate an observation x_i comes from the model, we first choose a cluster assignment c_i . It indicates which latent cluster x_i comes from and is drawn from a categorical distribution over $\{1,...K\}$. (We encode c_i as an indicator K-vector, all zeros except for a one in the position corresponding to x_i 's cluster.) We then draw x_i from the corresponding Gaussian $\mathcal{N}(c_i^T\mu,1)$.

The full hierarchical model is:

$$\mu_k \sim \mathcal{N}(0, \sigma^2),$$
 $c_i \sim Categorical(\frac{1}{K}, ..., \frac{1}{K}),$
 $x_i | c_i, \mu \sim \mathcal{N}(c_i^T \mu, 1)$ (10.7)

Deep Generative Models

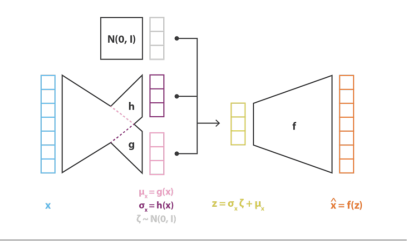
- **Generative** models can capture the joint probability P(X, Y), or just P(x) if there are no labels
- **Discriminative** models capture the conditional probability P(Y|X)

11.1 Auto-Regressive Generative Models

PixelRNNs and PixelCNNs model the joint distribution of pixels over an image *x* as the following product of conditional distributions.

$$P(x) = \prod_{i=1}^{n^2} P(x_i|x_1, ..., x_{i-1})$$
 (11.1)

11.2 Variational Autoencoders



loss = $C ||x - \hat{x}||^2 + KL[N(\mu_x, \sigma_x), N(0, I)] = C ||x - f(z)||^2 + KL[N(g(x), h(x)), N(0, I)]$

图 11.1: vae

11.2.1 Why do we need VAEs?

Autoencoder的encoder生成的latent space不够regurlarity(不连续),我们无法从latent space的中随机采样来生成或修改图片

The lack of interpretable and exploitable structures in the latent space (lack of regularity). the regularity of the latent space for autoencoders is a difficult point that depends on the distribution of the data in the initial space, the dimension of the latent space and the architecture of the encoder.

So, it is pretty difficult (if not impossible) to ensure, a priori, that the encoder will organize the latent space in a smart way compatible with the generative process we just described.

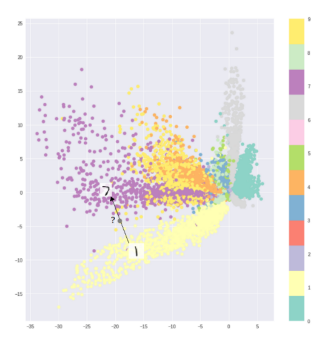


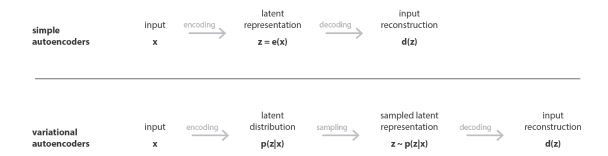
图 11.2: the encodings from a 2D latent space / MNIST

11.2.2 VAE

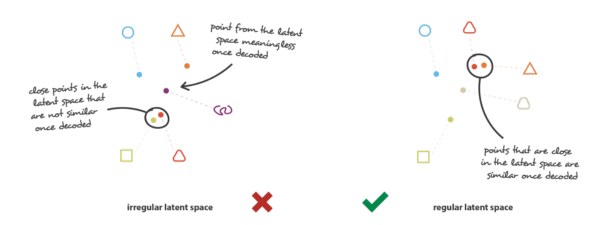
A variational autoencoder can be defined as being an autoencoder whose training is regularised to avoid overfitting and ensure that the latent space has good properties that enable generative process.

VAE模型需要符合以下条件

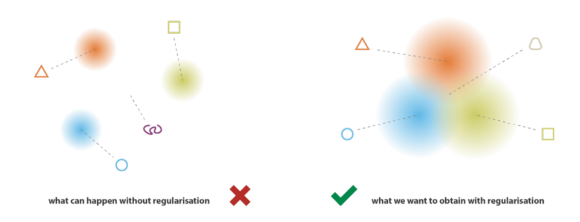
- the input is **encoded as distribution** over the latent space, instead of encoding an input as a single point
- a point from the latent space is sampled from that distribution
- the sampled point is decoded and the reconstruction error can be computed
- the reconstruction error is backpropagated through the network



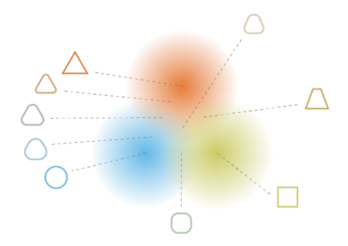
- Continuity two close points in the latent space should not give two completely different contents once decoded
- **Completeness** for a chosen distribution, a point sampled from the latent space should give "meaningful" content once decoded



The only fact that VAEs encode inputs as distributions instead of simple points is not sufficient to ensure continuity and completeness. Without a well defined regularisation term, the model can learn, in order to minimise its reconstruction error, to "ignore" the fact that distributions are returned and behave almost like classic autoencoders (leading to overfitting). To do so, the encoder can either return distributions with tiny variances (that would tend to be punctual distributions (点 分布)) or return distributions with very different means (that would then be really far apart from each other in the latent space).



we have to regularise both the covariance matrix and the mean of the distributions returned by the encoder. In practice, this regularisation is done by **enforcing distributions to be close to a standard normal distribution**



11.2.3 The VAE Lower Bound

We approximate the posterior with the optimized member of the family $Q^*(z|x)$.

$$Q_{\phi}^*(z|x) = \arg\min_{Q_{\phi}(z|x) \in \mathcal{Q}} KL(Q_{\phi}(z|x)||P_{\theta}(z|x))$$

$$\tag{11.2}$$

That is equivalent to ELBO:

$$\mathcal{L} = -ELBO(\theta, \phi, x) = KL(Q_{\phi}(z|x)||P(z)) - \mathbb{E}_{z \sim Q_{\phi}(z|x)}[\log P_{\theta}(x|z)]$$
(11.3)

11.2.4 Regularizer - Solution of KL(Q(z)||P(z)), Gaussion case

VAEs take an unusual approach to dealing with this problem: they assume that there is no simple interpretation of the dimensions of z, and instead assert that samples of z can be drawn from a simple distribution, i.e., $\mathcal{N}(0,I)$, where I is the identity matrix. The key is to notice that any distribution in d dimensions can be generated by taking a set of d variables that are normally distributed and mapping them through a sufficiently complicated function.

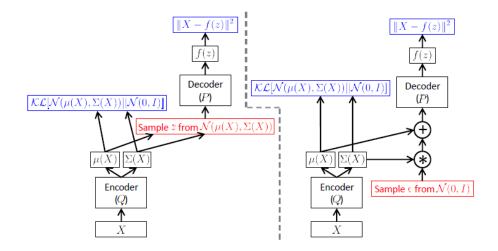
Let the prior over the latent variables be the **centered isotropic multivariate Gaussian** $P(z) = \mathcal{N}(z; 0, I)$. In this case, we can let the variational approximate posterior be a **multivariate Gaussian** with a **diagonal covariance structure** - $Q_{\phi}(z|x) = \mathcal{N}(z; \mu(x; \phi), \Sigma(x; \phi))$.

$$KL(Q_{\phi}(z|x)||P(z)) = KL(\mathcal{N}(\mu(x), \Sigma(x))||\mathcal{N}(0, I))$$

$$= \frac{1}{2} ((\mu(x))^{T}(\mu(x)) + tr(\Sigma(x)) - k - \log \det(\Sigma(x)))$$
(11.4)

11.2.5 Reconstruction Error

We want to **differentiate** and optimize the lower bound $ELBO(\theta, \phi, x)$ w.r.t both the variational parameters ϕ and generativet parameters θ .由于隐变量z从Q(z|x)的采样过程不可微,所以需要改写成一个可微的形式. we can sample from $\mathcal{N}(\mu(x), \Sigma(x))$ by first sampling $\epsilon \sim \mathcal{N}(0, I)$, then computing $z = \mu(x) + \Sigma^{\frac{1}{2}}(x) * \epsilon$ - **reparameterization trick**.



11.3 Generative Adversarial Models

11.3.1 Mode collapse

https://aiden.nibali.org/blog/2017-01-18-mode-collapse-gans/

11.3.2 Wasserstein GAN and the Kantorovich-Rubinstein Duality

https://vincentherrmann.github.io/blog/wasserstein/

11.3.3 vanilla GAN

D and G play the following two-player minimax game with value function V(G, D):

$$\min_{G} \max_{D} V(D, G) = \mathbb{E}_{x \sim P(x)}[\log D(x)] + \mathbb{E}_{z \sim P(z)}[\log(1 - D(G(z)))]$$
(11.5)

The training criterion for the discriminator D, given any generator G, is to maximize the quantity V(G,D)

$$V(G, D) = \mathbb{E}_{x \sim P(x)}[\log D(x)] + \mathbb{E}_{z \sim P(z)}[\log(1 - D(G(z)))]$$

= $\mathbb{E}_{x \sim P_d(x)}[\log D(x)] + \mathbb{E}_{x \sim P_g(x)}[\log(1 - D(x))]$ (11.6)

For any $(a,b) \in \mathbb{R}^2 \setminus \{0,0\}$, the function $y \to a \log(y) + b \log(1-y)$ achieves its maximum in [0,1] at $\frac{a}{a+b}$. So, for G fixed, the optimal discriminator D is

$$D_G^*(x) = \frac{P_d(x)}{P_d(x) + P_g(x)}$$
(11.7)

Note that the training objective for D can be interpreted as maximizing the log-likelihood for estimating the conditional probability P(Y = y|x), $x \in P_d or \in P_g$, then

$$C(G) = \max_{D} V(G, D)$$

$$= \mathbb{E}_{x \sim P_{d}(x)} [\log D_{G}^{*}(x)] + \mathbb{E}_{z \sim P(z)} [\log (1 - D_{G}^{*}(G(z)))]$$

$$= \mathbb{E}_{x \sim P_{d}(x)} [\log D_{G}^{*}(x)] + \mathbb{E}_{x \sim P_{g}(x)} [\log D_{G}^{*}(x)]$$

$$= \mathbb{E}_{x \sim P_{d}(x)} [\log \frac{P_{d}(x)}{P_{d}(x) + P_{g}(x)}] + \mathbb{E}_{x \sim P_{g}(x)} [\log \frac{P_{g}(x)}{P_{d}(x) + P_{g}(x)}]$$
(11.8)

The global minimum of C(G) is achieved if and onl if $P_g = P_d$. And

$$C(G) = -\log 4 \tag{11.9}$$

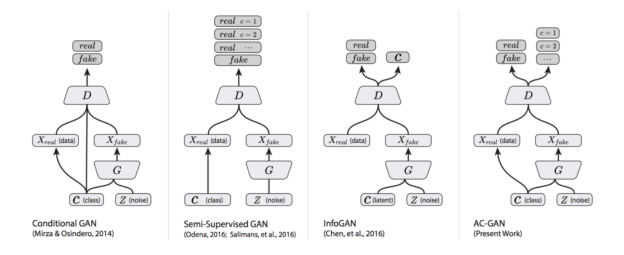
and that by substracting this expression from $C(G) = V(D_G^*, G)$, we abtain:

$$C(G) = -\log(4) + KL(P_d \| \frac{P_d + P_g}{2}) + KL(P_g \| \frac{P_d + P_g}{2})$$

= $-2\log(2) + 2JSD(P_d \| P_g)$ (11.10)

在D最优的情况下,等价于优化ISD

11.3.4 Latent space



Conditional GAN

使用辅助信息——label,将label和image绑定,分别输入到D和G中,

$$\min_{G} \max_{D} V(D,G) = \mathbb{E}_{x \sim P_d(x)}[\log D(x|y)] + \mathbb{E}_{z \sim P_z(z)}[\log(1 - D(G(z|y)))]$$
(11.11)

可以使用标签控制生成指定类型的图片

Semi-Supervised GAN

InfoGAN

$$\min_{G} \max_{D} V_{I}(D,G) = V(D,G) - \lambda I(c;G(z,c))$$
(11.12)

ACGAN

11.3.5 Architecture

11.3.6 Object functions

Reinforcement Learning

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