# Notes on INFO8010 - Deep Learning

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This document contains notes and additional readings for self-study.

### Lecture 4: Computer Vision

- Misc.
  - On cross-entropy
    - \* (Wiki page) Cross Entropy
    - \* A Gentle Introduction to Cross-Entropy for Machine Learning
- Classification
  - Image augmentation
  - Use pre-trained models for fine tuning and transfer learning
  - Large networks trained for classification are heavily re-used for object detection and semantic segmentation tasks.
- Object Detection
  - YOLO for object detection
    - \* EPFL EE-559, 8-3: Object Detection
  - R-CNN
    - \* Dive into Deep Learning 13.8. Region-based CNNs (R-CNNs)
  - Takeaways
    - \* One-stage detectors (YOLO, SSD, RetinaNet, etc) are fast for inference not as accurate.
    - \* Two-stage detectors (Fast R-CNN, Faster R-CNN, R-FCN, Light head R-CNN, etc) are usually slower but are more accurate.
    - \* Both depend on engineering decisions.
- Segmentation
  - Task: partitioning an image into regions of different semantic categories at pixel level.
  - Fully convolutional network(FCN) and transposed convolution
    - \* CS231n, Lecture 11, 2018.
  - Mask R-CNN
    - \* Object detection combined with mask prediction enables instance segmentation.
    - $\ast$  Dive into Deep Learning 13.8.4 Mask R-CNN

## Lecture 5: Training Neural Networks

- Optimizers
  - Gradient descent
    - \* GD, SGD, mini-batch SGD

- \* Rely on assumptions on 1) the magnitude of the local curvature to set the step size, and 2) isotropy in gradient so the step size makes sense in all directions
- Wolfe conditions ensures that both the loss function decreases sufficiently and the slope reduces sufficiently.
   However, line search will be too expensive for DL, and might lead to local minimum / overfitted solution.
- Momentem
  - \* Use momentum to add inertia in the choice of the step direction
  - \* Nesterov momentem
- Adaptive learning rate: without the assumption of istropic gradient
  - \* Per-parameter methods: AdaGrad, RMSProp, Adam
  - \* Scheduling
- Some additional reading on optimization: (Sebastian Ruder) An overview of gradient descent optimization algorithms
- Initialization
  - Principles
    - \* Break symmetry
    - \* Control variance of activation across layers during forward and backward pass
  - Xavier initialization
- Normalization
  - Batch normalization
  - Layer normalization

#### Lecture 6: Recurrent Neural Networks

- Types of tasks
  - Classification: sequence to classes
  - Synthesis: real values to sequence
  - Translation: sequence to sequence
- Temporal convolutions
- Recurent neural networks
  - Structure
    - \* maintain a recurrent state updated at each time step (a function of state the previous step, input of the current step, and weights),  $\mathbf{h}_t = \phi(\mathbf{x}_t, \mathbf{h}_{t-1}; \theta)$
    - \* Predictions can be computed at any step from the recurrent state  $y_t = \psi(\mathbf{h}_t; \theta)$
    - \* Elman netoworks apply non-linear activation functions as  $\phi$  and  $\psi$
  - Stacked RNN
    - \* Since RNNs can be viewed as layers producing sequences of activations, and can be stacked
  - Bidirectional RNNs. RNNs can be made bidirectional. run the same single direction RNN twice from both end and concatenate the states.
  - Gating
    - \* Similar to the skip connections in ResNet, RNN cells can inlude pass-throughs so recurrent state does not go repeatedly through a squashing non-linearity.
    - \* forget gate: current state update be a per-component weighted average of its previous value and a full update, with the weighting depending on input and the previous state.
  - LSTM is able to learn long-term dependencies, and the core idea is to use cell state and erase/update/output
    gates for cell state information.
    - \* See Understanding LSTM Networks by Colah
  - GRU (gated recurren unit) uses two (instead of three as in LSTM) gates (update/reset), and it performs similarly to LSTM but with fewer parameters (although LSTM is strictly stronger).
  - Graident
    - \* Note that gated units prevent gradients from vanishing, but not from exploding, which can be solved using **gradient norm clipping** (scaling of the norm).
    - \* Orthogonal initialization (of the weight matrix) will guarantee that activations will neigher vanish nore explode.
- Applications
  - Sentiment analysis
    - \* Document-level modeling for sentiment analysis (= text classification), with stacked, bidirectional and gated recurrent networks. (Duyu Tang et al, 2015)

- Language models
  - \* Language as Markov Chain  $p(\mathbf{w}_t|\mathbf{w}_{1:t-1})$
  - \* An instance of sequence synthesis where predictions are computed at all time steps
  - \* Text generation (Max Woolf 2018)
- Sequence synthesis
- Neural machine translation (Google's Neural Machine Translation System: Bridging the Gap between Human and Machine Translation)
- Text-to-speech synthesis
- Beyond sequences
  - Neural computers
  - Programs as neural nets
  - Graph neural network
- Reference
  - Kyunghyun Cho, "Natural Language Understanding with Distributed Representation", 2015

### Lecture 7: Auto-encoders and generative models

- Auto-encoders (AE)
  - An **auto-encoder** is a composite function made of
    - \* encoder f from the original space  $\mathcal{X}$  to a latent space  $\mathcal{Z}$
    - \* decoder g to map back to  $\mathcal{X}$
    - \* such that  $g \circ f$  is close to the identity on the data, i.e.  $\mathbb{E}_{\mathbf{x} \sim p(\mathbf{x})}[||\mathbf{x} g \circ f(\mathbf{x})||^2] \approx 0$
    - \* Training an auto-encoder consists of minimizing this loss function to find the best parameterization of f and g.
  - Interpolation on latent space can be made to get an intuition of the learned latent representation.
  - Denoising auto-encoders
    - \* The goal is to optimize  $h = g \circ f : \mathcal{X} \to \mathcal{X}$  such that a perturbation  $\tilde{\mathbf{x}}$  is restored to  $\mathbf{x}$ .
    - \* A weakness of denoising auto-encoder is that the posterior  $p(\mathbf{x}|\tilde{\mathbf{x}})$  may be multi-modal.
- Generative models
  - a probabilistic model that can be used to simulate the data,  $\mathbf{x} \sim p(\mathbf{x}; \theta)$ .
  - Applications
    - \* Supper-resolution, Compression, text-to-speech
    - \* Proteomics, drug discovery, astronomy
    - \* Planning, exploration, model-based RL
  - The decoder g can be assessed by introducing a density model q over the latent space  $\mathcal{Z}$  for sampling and mapping back into the data space  $\mathcal{X}$ . (e.g., Gaussian  $q(\mathbf{z}) = \mathcal{N}(\hat{\mu}, \hat{\Sigma})$ )
  - Sampled and generated results are not satisfactory because the density model p on the latent space is too simple and inadequate.
- Variational inference (VI)
  - A prescribed latent variable model that defines a joint probability  $p(\mathbf{x}, \mathbf{z}) = p(\mathbf{x}|\mathbf{z})p(\mathbf{z})$
  - Bayes rule gives  $p(\mathbf{z}|\mathbf{x}) = \frac{p(\mathbf{x}|\mathbf{z})p(\mathbf{z})}{p(\mathbf{x})}$ , which is intractable for integrating over  $\mathbf{x}$ .
  - VI turns the posterior inference into an optimization problem that minimize the KL divergence between  $p(\mathbf{z}|\mathbf{x})$  and the approximation  $q(\mathbf{z}|\mathbf{x};\nu)$ 
    - \* See slides pp. 44 pp.47 for details of the KL divergence, evidence lower bound objective (ELBO), and the optimization setups.
    - \* ELBO encourages distributions to place their mass on configurations of latent variables that explain the oberved data, and close to the prior.
- Variational auto-encoders
  - Variational auto-encoder is a deep latent model where
    - \*  $p(\mathbf{x}|\mathbf{z};\theta)$  is parameterized with a **generative network**  $NN_{\theta}$  (decoder) that takes input  $\mathbf{z} \in \mathcal{Z}$  and outputs parameters  $\phi = NN_{\theta}(\mathbf{z})$  to the data distribution, i.e.

$$\mu, \sigma = NN_{\theta}(\mathbf{z}), \quad p(\mathbf{x}|\mathbf{z}; \theta) = \mathcal{N}(\mathbf{x}; \mu, \sigma^2 \mathbf{I})$$

\* The approximate posterior  $q(\mathbf{z}|\mathbf{x};\varphi)$  is parameterized with an **inference network**  $NN_{\varphi}$  (encoder)

that takes as input x and outputs parameters  $\nu = NN_{\varphi}(x)$  to the approximate posterior. E.g.

$$\mu, \sigma = \text{NN}_{\varphi}(\mathbf{x}), \quad q(\mathbf{z}|\mathbf{x}; \varphi) = \mathcal{N}(\mathbf{z}; \mu, \sigma^2 \mathbf{I})$$

- We use variational inference to jointly optimize the generative and inference networks.
  - \* Doing so involves Monte Carlo integration (for computing gradients of the ELBO w.r.t.  $\theta$ ) and reparameterization trick + Monte Carlo (for computing gradients of ELBO w.r.t.  $\varphi$ )

#### Lecture 8: Generative Adversarial Networks

- Generative adversarial networks (GANs)
  - Two-player game
    - \* Generator network  $g(\cdot; \theta) : \mathcal{Z} \to \mathcal{X}$  with prior  $p(\mathbf{z})$  on latent space, thereby inducing a generative distribution

$$\mathbf{x} \sim q(\mathbf{x}; \theta) \iff \mathbf{z} \sim p(\mathbf{z}), \mathbf{x} = g(\mathbf{z}; \theta)$$

- \* Discriminative network  $d(\cdot; \phi) : \mathcal{X} \to [0, 1]$  as a classifier to distinguish between true samples  $\mathbf{x} \sim p(\mathbf{x})$  and generated samples  $\mathbf{x} \sim q(\mathbf{x}; \theta)$
- Objective of GANs
  - \* Use cross-entropy loss for d and we have the value function (log-likelihood)

$$V(\phi, \theta) = \mathbb{E}_{\mathbf{x} \sim p(\mathbf{x})}[\log d(\mathbf{x}; \theta)] + \mathbb{E}_{\mathbf{z} \sim p(\mathbf{z})}[1 - d(g(\mathbf{z}; \theta); \phi))]$$

- \* The ultimate goal is  $\theta^* = \arg\min_{\theta} \max_{\phi} V(\phi, \theta)$
- \* Mathematically  $\theta^*$  is minimum  $\iff p(\mathbf{x}) = q(\mathbf{x}; \theta)$ , that is, the corresponding generative model can perfectly reproduce the true data distribution.
- Learning process
  - \* Alternating SGD

$$\theta \leftarrow \theta - \gamma \nabla_{\theta} V(\phi, \theta); \quad \phi \leftarrow \phi - \gamma \nabla_{\phi} V(\theta, \phi)$$

- \* For each step of  $\theta$  we can take k steps on  $\phi$  to make the classifier near optimal
- \* Computing  $\nabla_{\theta}$  requires backprop through d before computing the partial derivatives w.r.t. g's internals.
- Open problems
  - \* Training standard GAN often results in pathological behaviors due to
    - · Oscillations without convergence
    - · Vanishing gradients
    - · Mode collapse: g models well on a small sub-population concentrating on a few modes of the data distribution.
    - · Performance diffucult to assess in practice.
- Wasserstein GANs
  - Original GAN as defined above suffers from vanishing gradients, especially when initialy  $\mathbf{x} \sim q(\mathbf{x}; \theta)$  can be so bad that the response of d saturates, meaning d is nearly perfect there fore  $V(\phi, \theta) = ...(definedabove)$  has near-zero gradients which halts the optimization.
    - \* One of the reason for the setback is that Jensen-SHannon divergence poorly accounts for the metric structure of the space.
  - Wasserstein GAN uses Wasserstein-1 distance

$$\theta^* = \arg\min_{\theta} W_1(p(\mathbf{x}||q(\mathbf{x};\theta))) = \arg\min_{\theta} \max_{\phi:||d(\cdot;\phi)||_L \le 1} \mathbb{E}_{\mathbf{x} \sim p(\mathbf{x})}[d(\cdot;\phi)] - \mathbb{E}_{\mathbf{x} \sim q(\mathbf{x};\theta)[d(\mathbf{x};\theta)]}$$

- \* In this case,  $d: \mathcal{X} \to \mathbb{R}$  is a critic function that satisfies 1-Lipschitzness.
- \* See Arjovsky et al (2017) and Gulrajani et al (2017) for details.
- As a result, Wasserstein GANS benefit from
  - \* A meaningful loss metric
  - \* Improved stability (no mode collapse is observed)
- Convergence of GANs
- State of the art
- Applications

# Resrouces

- EPFL EE-559 Deep Learning EE-559 "Deep Learning", taught by François Fleuret in the School of Engineering of the École Polytechnique Fédérale de Lausanne, Switzerland.
- Dive into Deep Learning: An interactive deep learning book with code, math, and discussions, based on the NumPy interface.
- Notes of deep learning specialization, good for reviewing the fundamentals of DL.