

NPFL099 Statistical Dialogue Systems

4. Training Neural Nets

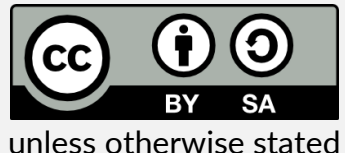
<http://ufal.cz/npfl099>

Ondřej Dušek, Zdeněk Kasner, Mateusz Lango, Ondřej Plátek

24. 10. 2024



Charles University
Faculty of Mathematics and Physics
Institute of Formal and Applied Linguistics

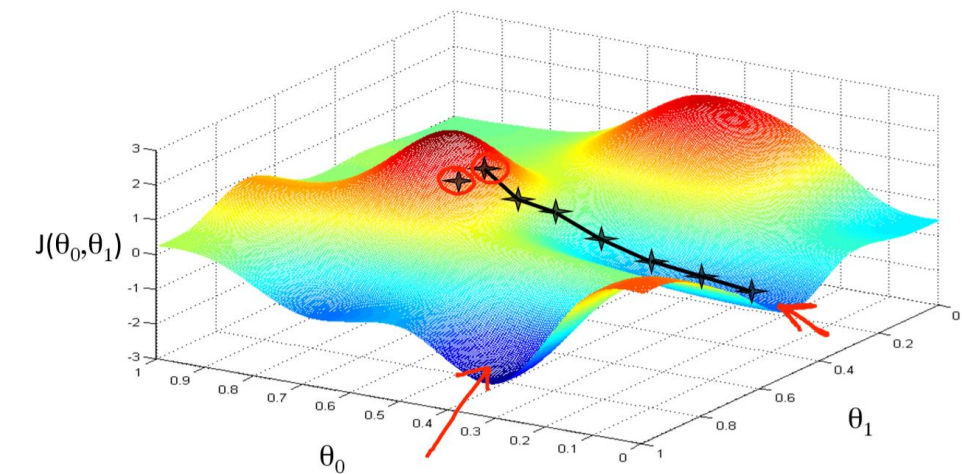


Recap: Neural Nets

- **complex functions, composed of simple functions** (=layers)
 - linear, ReLU, tanh, sigmoid, softmax
- **fully differentiable**
- different arrangements:
 - feed forward / multi-layer perceptron
 - CNNs
 - RNNs (LSTM/GRU)
 - attention
 - Transformer
- input: binary, float, embedding
- tasks/problems: classification, regression, structured (sequences/ranking)

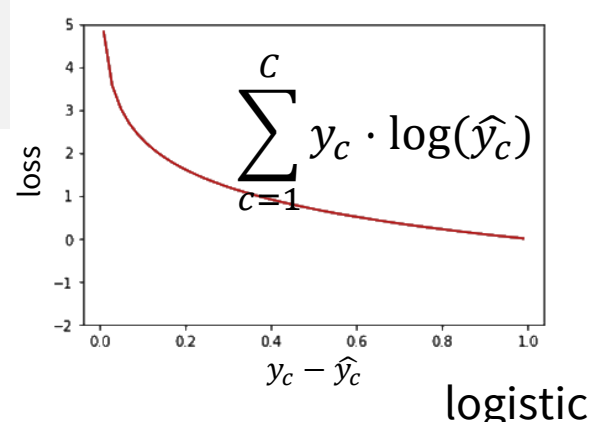
Supervised Training: Gradient Descent

- supervised training– **gradient descent** methods
 - minimizing a **cost/loss function**
(notion of error – given system output, how far off are we?)
 - calculus: derivative = steepness/slope
 - follow the slope to find the minimum – derivative gives the direction
 - **learning rate** = how fast we go (needs to be tuned)
- gradient typically computed (=averaged) over **mini-batches**
 - random bunches of a few training instances
 - not as erratic as using just 1 instance,
not as slow as computing over whole data
 - **stochastic gradient descent**
 - batches may be **accumulated** to fit into memory
 - e.g. your GPU only fits one instance
→ compute gradients multiple times, then do 1 update



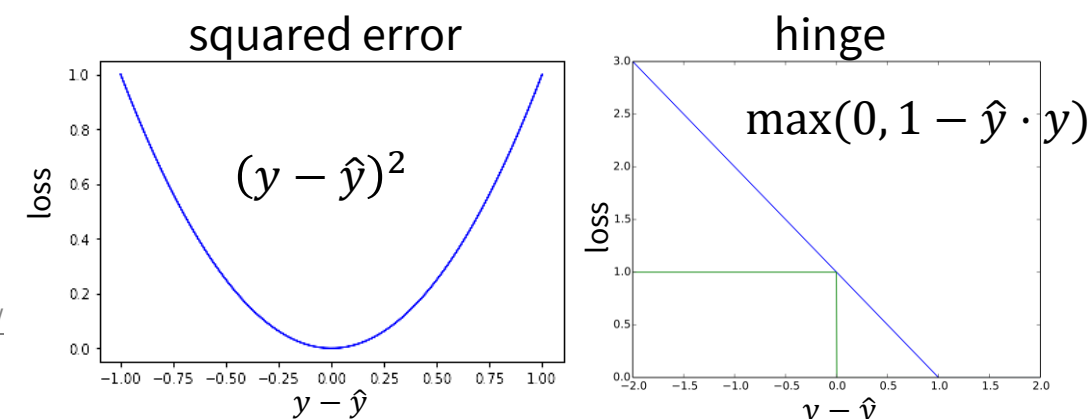
Cost/Loss Functions

- differ based on what we're trying to predict
- **logistic / log loss** (“cross entropy”)
 - for classification / softmax – including **word prediction**
 - classes from the whole dictionary
 - correct class <100% prob. → loss
 - pretty stupid for sequences, but works
 - sequence shifted by 1 ⇒ everything wrong
- **squared error loss** – for regression
 - forcing the predicted float value to be close to actual one
- **hinge loss** – binary classif. (SVMs), ranking
 - forcing the correct sign
- many others, variants



logistic

reference: *Blue Spice is expensive
expensive
cheap
pricey
in the expensive price range*



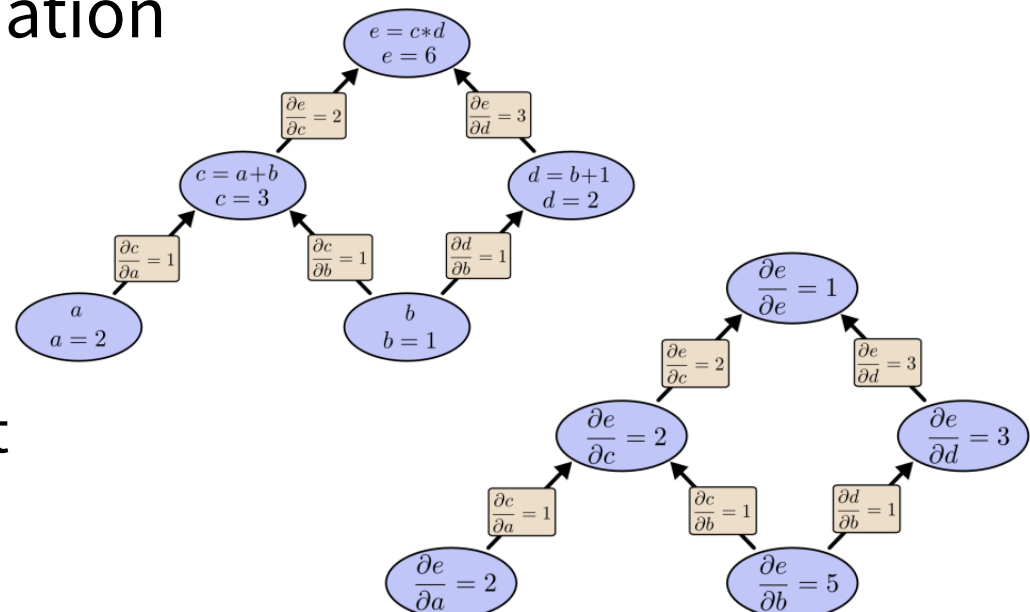
Backpropagation

<https://www.mathsisfun.com/calculus/derivatives-rules.html>

- network ~ computational graph
 - reflects function/layer composition
- composed function derivatives – simple rules
 - basically summing over different paths
 - factoring ~ merging paths at every node

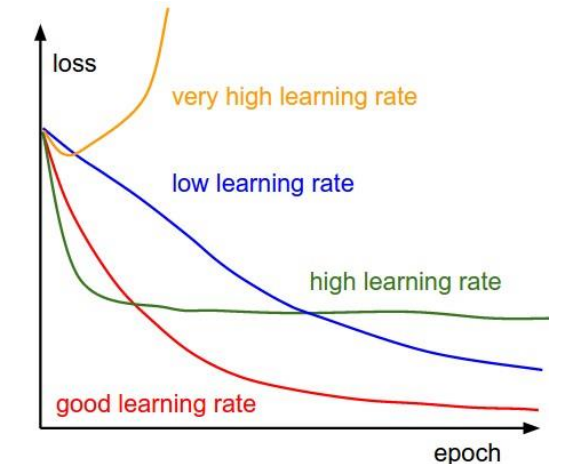
- **backpropagation** = reverse-mode differentiation
 - going back from output to input
 - ~ how every node affects the output
 - your graph **output = cost function**
 - → derivatives of all parameters w. r. t. cost
 - one pass through the network only → easy & fast
 - NN frameworks do this automatically

Rules	Function	Derivative
Multiplication by constant	cf	cf'
<u>Power Rule</u>	x^n	nx^{n-1}
Sum Rule	$f + g$	$f' + g'$
Difference Rule	$f - g$	$f' - g'$
<u>Product Rule</u>	fg	$f'g + f'g$
Quotient Rule	f/g	$\frac{f'g - g'f}{g^2}$
Reciprocal Rule	$1/f$	$-f'/f^2$
Chain Rule (as " Composition of Functions ")	$f \circ g$	$(f' \circ g) \times g'$

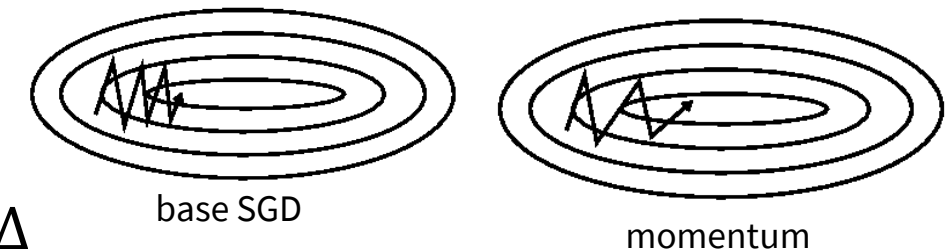


Learning Rate (α) & Momentum

- α : most important parameter in (stochastic) gradient descent
- tricky to tune:
 - too high α = may not find optimum
 - too low α = may take forever
- Learning rate decay: start high, lower α gradually
 - make bigger steps (to speed learning)
 - slow down when you're almost there (to avoid overshooting)
 - linear, stepwise, exponential
 - **reduce-on-plateau** – check every now and then if we're still improving, reduce LR if not
- Momentum: moving average of gradients
 - make learning less erratic
 - $m = \beta \cdot m + (1 - \beta) \cdot \Delta$, update by m instead of Δ



<http://cs231n.github.io/neural-networks-3/>



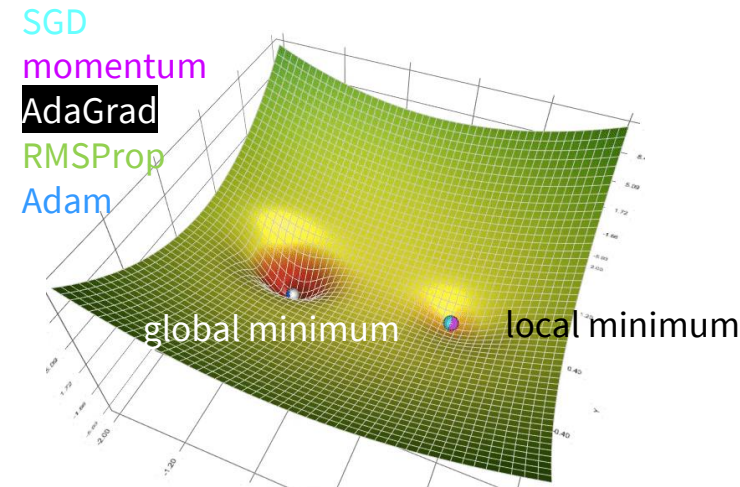
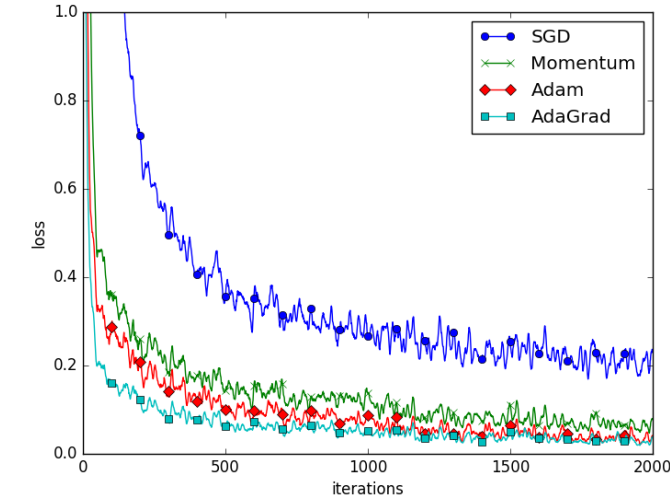
Optimizers

<http://kaeken.hatenablog.com/entry/2016/11/10/203151>

- Better LR management
 - change LR based on gradients, less sensitive to settings
- **AdaGrad** – all history
 - remember sum of total gradients squared: $\sum_t \Delta_t^2$
 - divide LR by $\sqrt{\sum \Delta_t^2}$
 - variants: **Adadelta**, **RMSProp** – slower LR drop
- **Adam** – per-parameter momentum
 - moving averages for Δ & Δ^2 :
 $m = \beta_1 \cdot m + (1 - \beta_1)\Delta, v = \beta_2 \cdot v + (1 - \beta_2)\Delta^2$
 - use m instead of Δ , divide LR by \sqrt{v}
 - often used as default nowadays
 - variant: **AdamW** – better regularization
 - not much difference though

(Kingma & Ba, 2015)
<https://arxiv.org/abs/1412.6980>

(Loshchilov & Hutter, 2019)
<https://arxiv.org/abs/1711.05101>



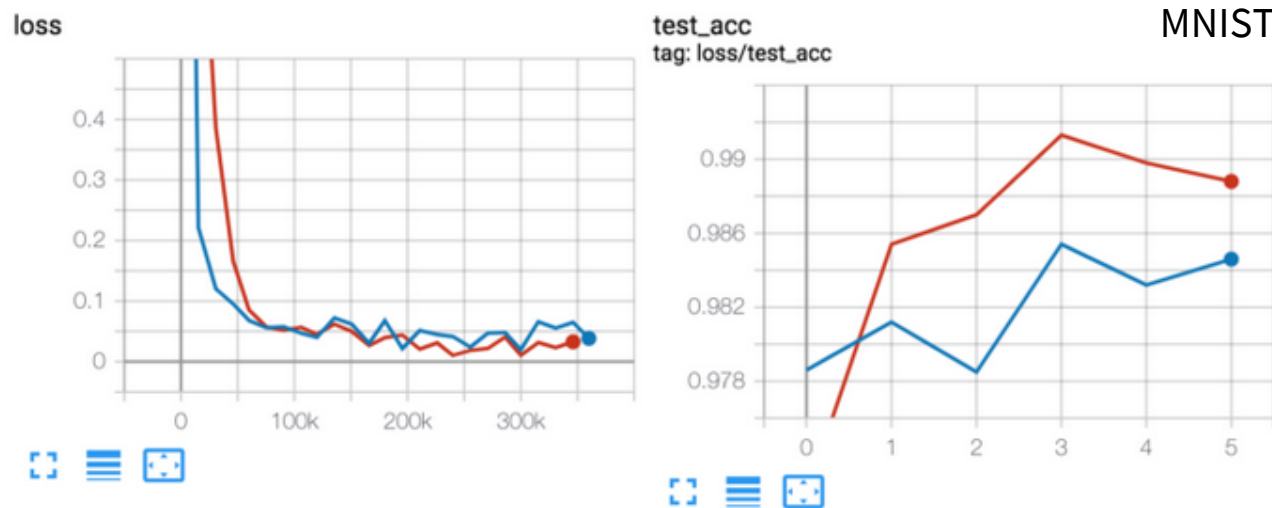
<https://ruder.io/optimizing-gradient-descent/>

<https://towardsdatascience.com/a-visual-explanation-of-gradient-descent-methods-momentum-adagrad-rmsprop-adam-f898b102325c>

Optimizers

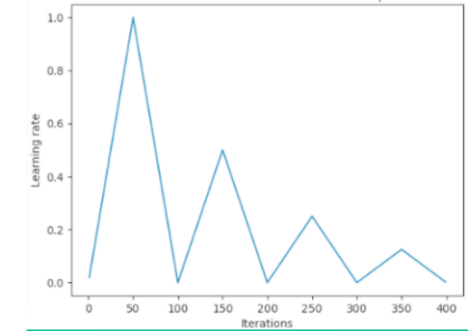
(You et al., 2020)
[http://arxiv.org/abs/1904.00962](https://arxiv.org/abs/1904.00962)

- **LAMB** – Layer-wise Adaptive Moments for Batches
 - for larger batches & allowing to use larger LR (~unstable otherwise)
- **LARS** layer-wise adaptive rate scaling
 - layer-wise LRs, always multiplied by a **trust ratio**:
$$\alpha^l = \alpha \cdot \frac{\|w^l\|}{\|\Delta^l\|}$$
 – norm of weights/ norm of gradients
 - higher trust ratio = faster updates
 - start of training:
low w , high $\Delta \rightarrow$ slow **warm up**
 - towards convergence:
higher w , low $\Delta \rightarrow$ faster training
- $\text{LAMB} \approx \text{LARS} + \text{AdamW}$

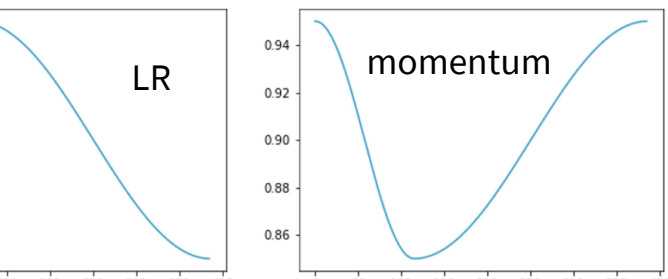
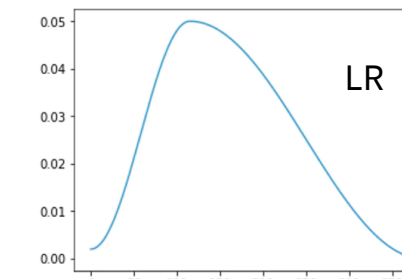


Schedulers

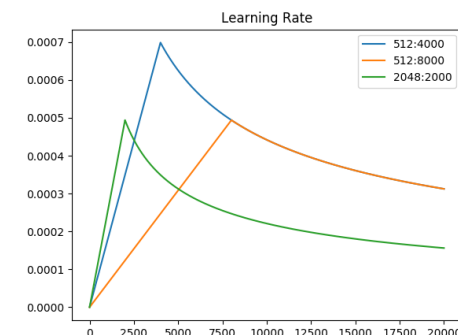
- more fiddling with LR – **warm-ups**
 - start learning slowly, then increase LR, then reduce again
 - may be repeated (**warm restarts**), with lowered maximum LR
 - allow to diverge slightly – work around local minima
- multiple options:
 - cyclical (=warm restarts) – linear, cosine annealing
 - **one cycle** – same, just don't restart
 - **Noam scheduler** – linear warm-up, decay by $\sqrt{\text{steps}}$
- combine with base SGD or Adam/Adadelta etc.
 - momentum updated inversely to LR
 - may have less effect with optimizers
 - trade-off: speed vs. sensitivity to parameter settings



cyclical scheduler (warm restarts)



one cycle with cosine annealing

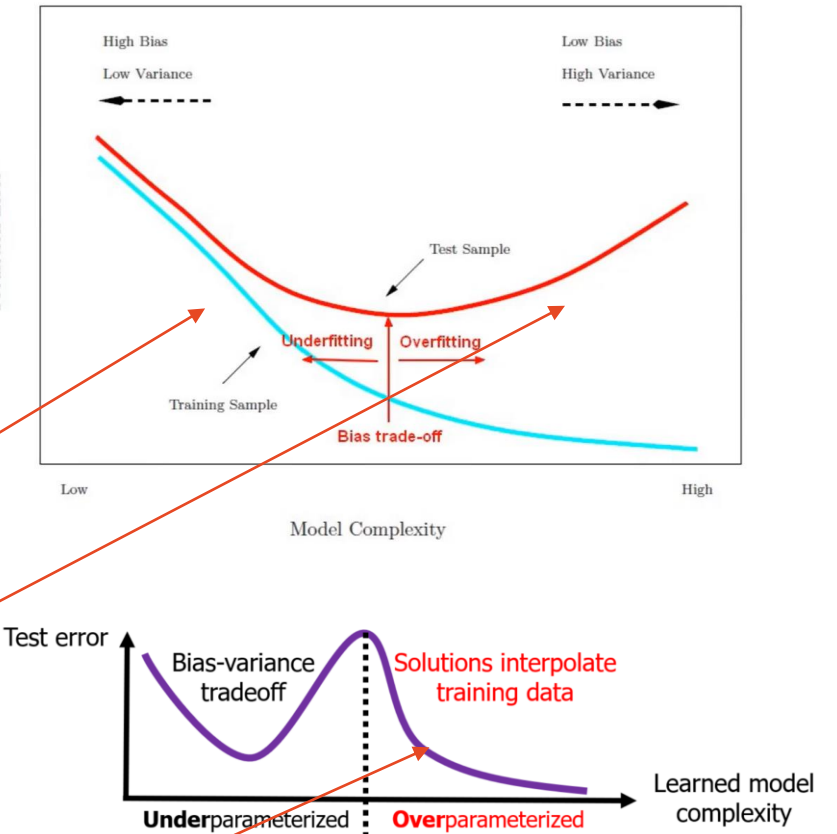


Noam scheduler with different parameters

When to stop training

<https://www.andreaperlato.com/theorypost/bias-variance-trade-off/>

- generally, when cost stops going down
 - despite all the LR fiddling
- problem: **overfitting**
 - cost low on training set, high on validation set
 - network essentially memorized the training set
 - → **check on validation set** after each epoch (pass through data)
 - stop when cost goes up on validation set
 - regularization (see →) helps delay overfitting
- **bias-variance** trade-off:
 - smaller models may underfit (high bias, low variance = not flexible enough)
 - larger models likely to overfit (too flexible, memorize data)
 - XXL models: “**grokking**” – overfit soo much they interpolate data → good (��?)

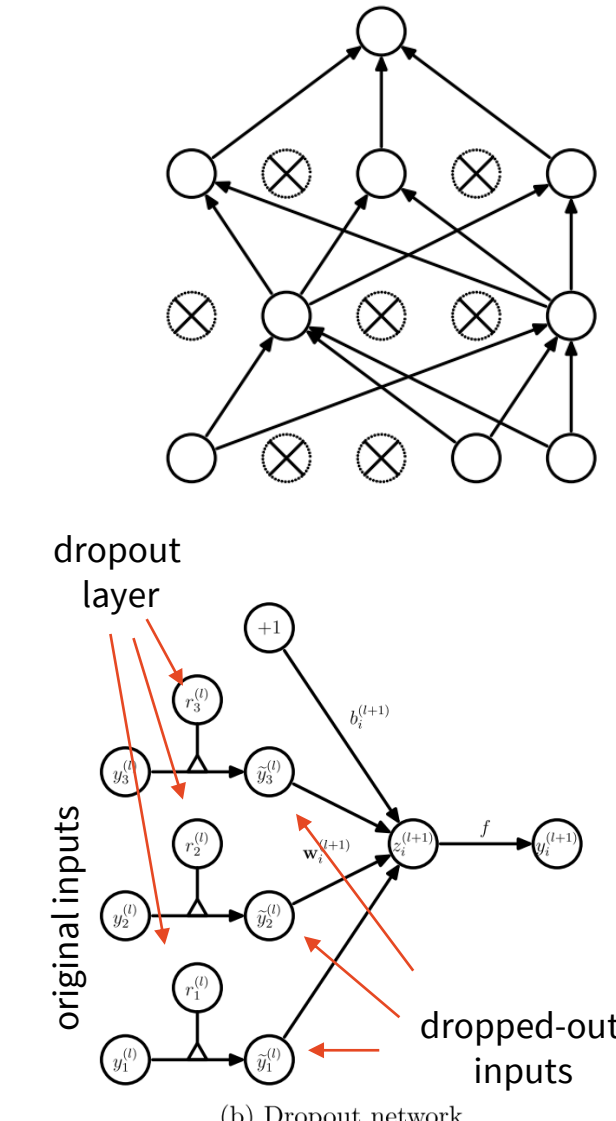


(Dar et al., 2021) <https://arxiv.org/abs/2109.02355>
(Power et al., 2022) <http://arxiv.org/abs/2201.02177>

Regularization: Dropout

- regularization: preventing overfitting
 - making it harder for the network to learn, adding noise
- **Dropout** – simple regularization technique
 - more effective than e.g. weight decay (L2)
 - **zero out some neurons/connections** in the network at random
 - technically: multiply by dropout layer
 - 0/1 with some probability (typically 0.5–0.8)
 - at training time only – full network for prediction
 - weights scaled down after training
 - they end up larger than normal because there's fewer nodes
 - done by libraries automatically
 - may need larger networks to compensate

(Srivastava et al., 2014)
<http://jmlr.org/papers/v15/srivastava14a.html>

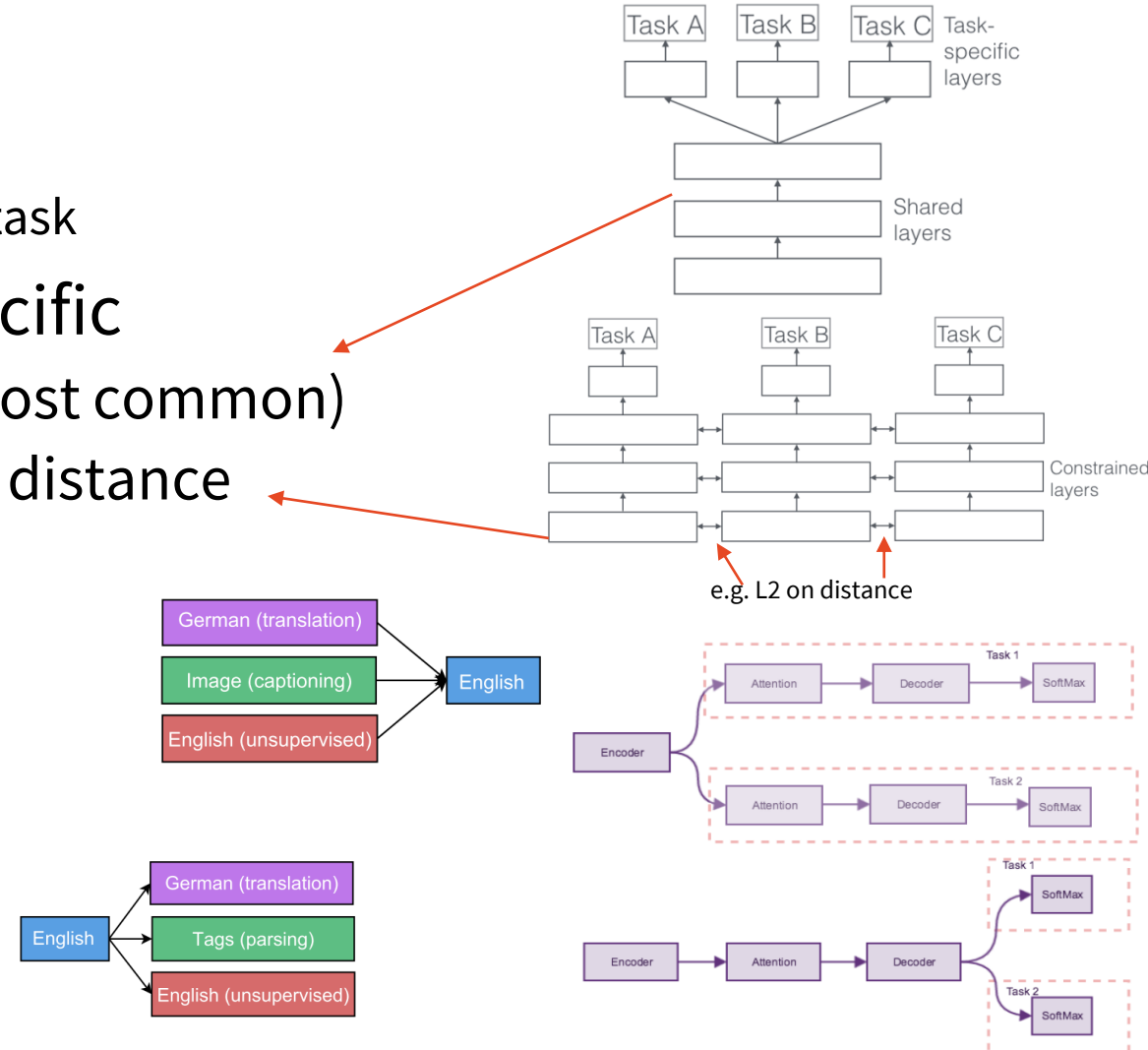


(b) Dropout network

Regularization: Multi-task Learning

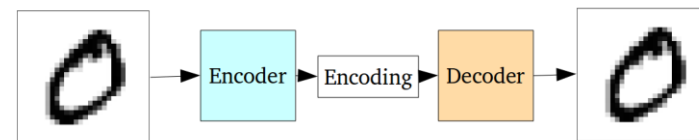
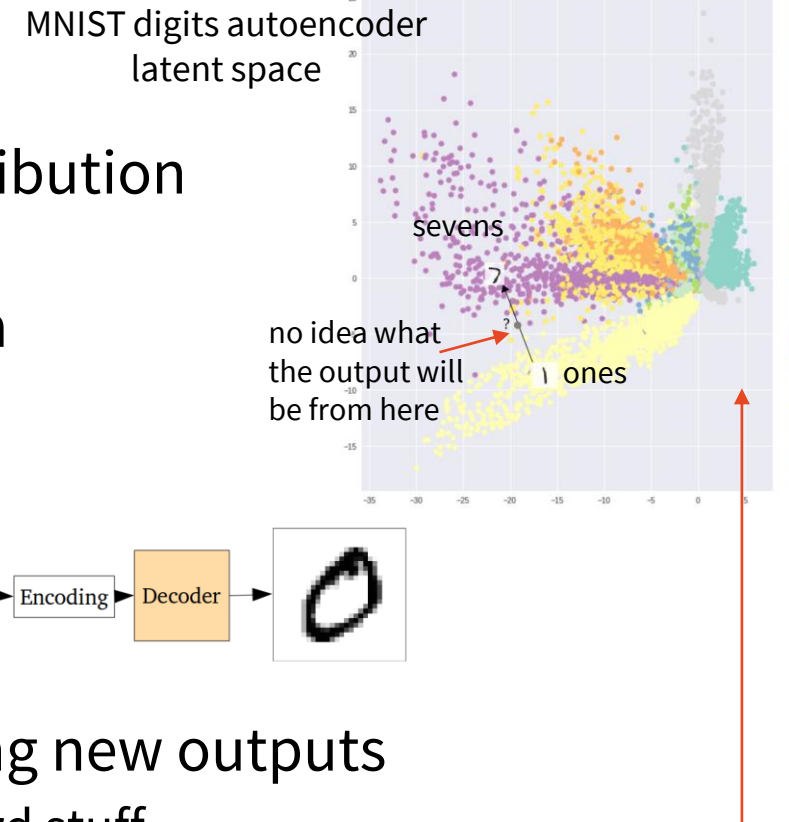
(Ruder, 2017)
<http://arxiv.org/abs/1706.05098>
(Fan et al., 2017)
<http://arxiv.org/abs/1706.04326>
(Luong et al., 2016)
<http://arxiv.org/abs/1511.06114>

- achieve better generalization by **learning more things at once**
 - a form of regularization
 - implicit data augmentation
 - biasing/focusing the model
 - e.g. by explicitly training for an important subtask
- parts of network shared, parts task-specific
 - hard sharing = parameters truly shared (most common)
 - soft sharing = regularization by parameter distance
 - different approaches w. r. t. what to share
- training – **alternating** between tasks
 - **catastrophic forgetting:**
if you don't alternate,
the network forgets previous tasks



Autoencoders

- Using NNs as **generative models**
 - more than just classification – modelling the whole distribution
 - (of e.g. possible texts, images)
 - generate new instances that look similar to training data
- **Autoencoder:** input → encoding → input
 - encoding ~ “embedding” in latent space (i.e. some vector)
 - trained by reconstruction loss
 - problem: can’t easily get valid embeddings for generating new outputs
 - parts of embedding space might be unused – will generate weird stuff
 - no easy interpretation of embeddings – no idea what the model will generate
- extension – **denoising autoencoder:**
 - add noise to inputs, train to generate clean outputs
 - use in multi-task learning, representations for use in downstream tasks

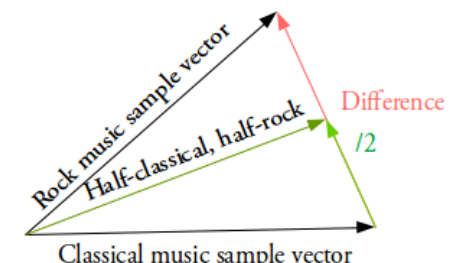
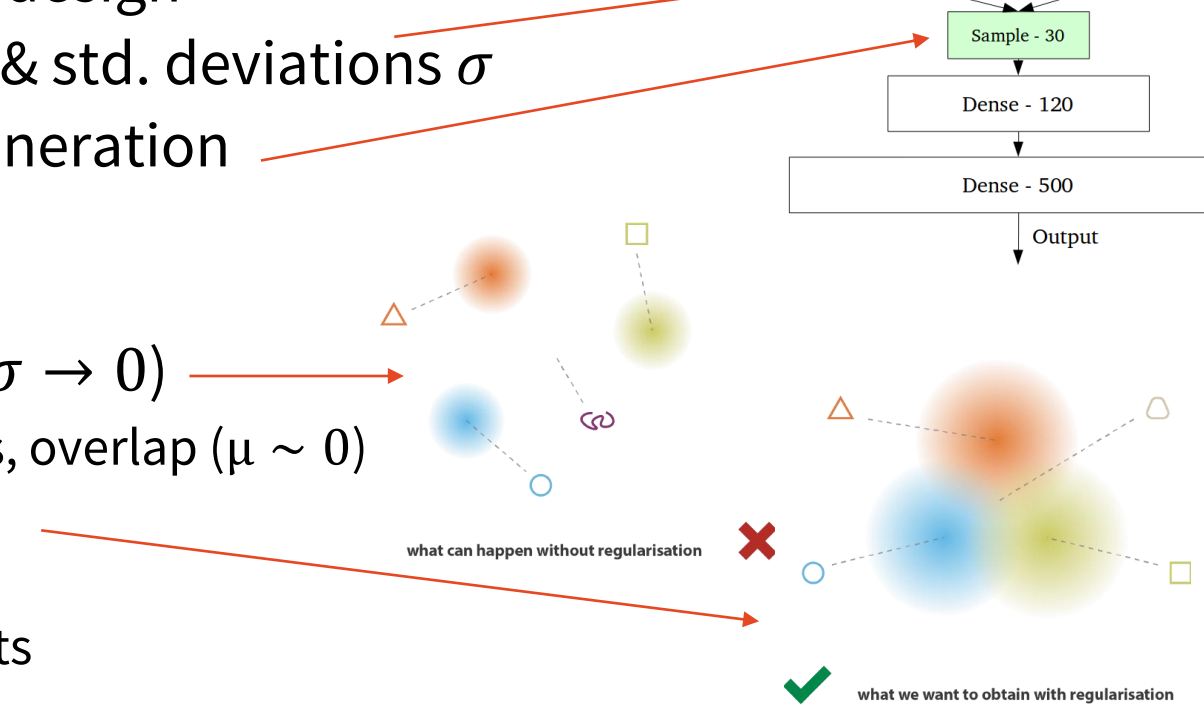
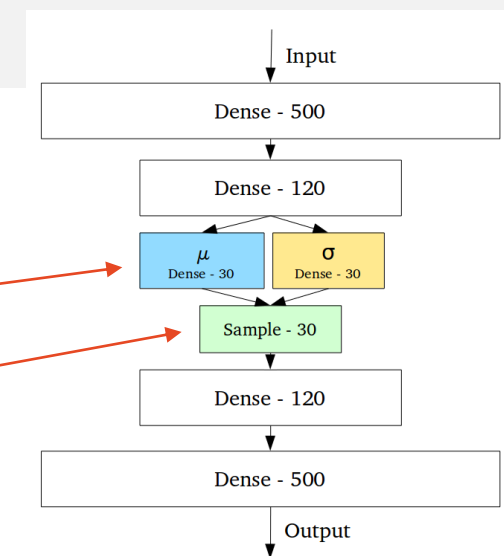


Variational Autoencoders

- Making the encoding latent space more useful

- using **Gaussians** – continuous space by design
- encoding input into vectors of means μ & std. deviations σ
- sampling encodings from $N(\mu, \sigma)$ for generation
 - samples vary a bit even for the same input
 - decoder learns to be more robust
- model can degenerate into normal AE ($\sigma \rightarrow 0$)
 - we need to encourage some σ , smoothness, overlap ($\mu \sim 0$)
 - add **2nd loss: KL divergence** from $N(0,1)$
 - VAE learns a trade-off between using unit Gaussians & reconstructing inputs

- Problem: still not too much control of the embeddings
 - we can only guess what kind of output the model will generate



<https://towardsdatascience.com/intuitively-understanding-variational-autoencoders-1bfe67eb5daf>
<https://towardsdatascience.com/understanding-variational-autoencoders-vae-f70510919f73>
<http://kvfrans.com/variational-autoencoders-explained/>

VAE details

- VAE objective:

- “AE” {
 - **reconstruction loss** (maximizing $p(x|z)$ in the decoder), MLE as per usual
- “V” {
 - **latent loss** (KL-divergence from ideal $p(z) \sim \mathcal{N}(0,1)$ in the encoder)

$$\mathcal{L} = -\mathbb{E}_q[\log p(x|z)] + KL[q(z|x)||p(z)]$$

- This is equivalent to maximizing true $\log p(x)$ with some error
 - i.e. maximizing **evidence lower bound** (ELBO) / variational lower bound:

$$\mathbb{E}_q[\log p(x|z)] - KL[q(z|x)||p(z)] = \log p(x) - KL[q(z|x)||p(z|x)]$$

↑
“evidence”
(i.e. data) ELBO

Normal noise

error incurred
by using q
instead of true
distribution p

- Sidestepping sampling – **reparameterization trick**

- $z \sim \mu + \sigma \cdot \mathcal{N}(0,1)$, then differentiate w. r. t. μ and σ
 - differentiating w. r. t. μ & σ still works, no hard sampling on that path

Discrete VAE: Gumbel-Softmax

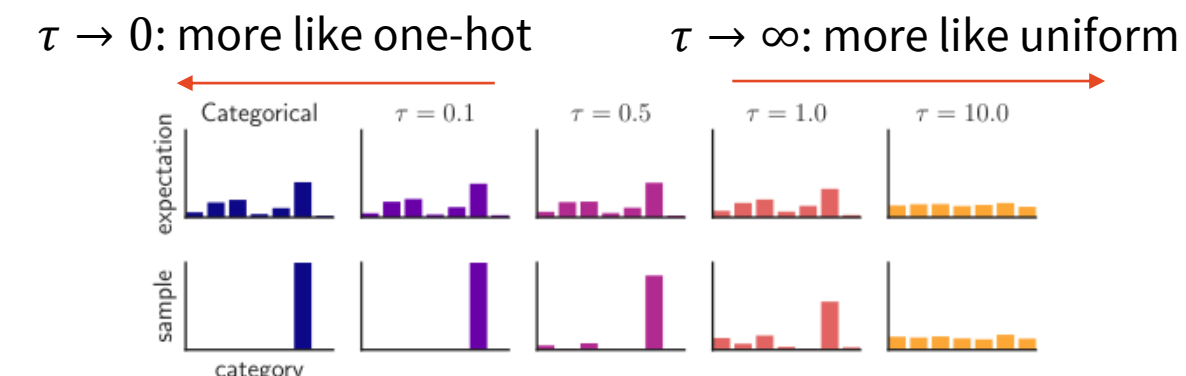
(Jang et al., 2017)
<https://arxiv.org/abs/1611.01144>

- “reparameterization trick for discrete distributions”
 - same idea, just with a **discrete/categorial distribution**
 - this makes the latent space better interpretable
- **Gumbel-max trick:**
 - categorial distribution π with probabilities π_i
 - sampling from π : $z = \text{onehot}(\arg \max_i (\log \pi_i + g_i))$
- Swap argmax for softmax with temperature τ
 - differs from π if $\tau > 0$, but may be close to it
 - approx. sample of the true distribution
 - fully differentiable
 - g_i bypassed in differentiation, same as $\mathcal{N}(0,1)$ in Gaussian sampling

Gumbel noise:

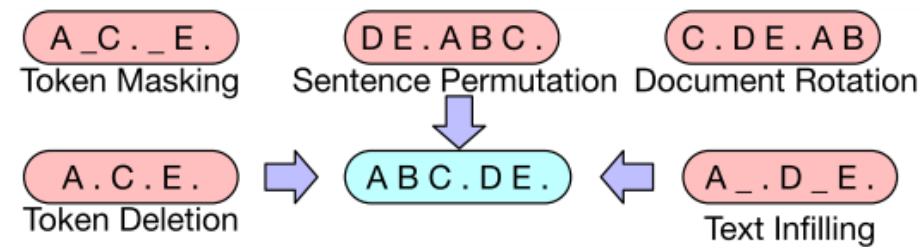
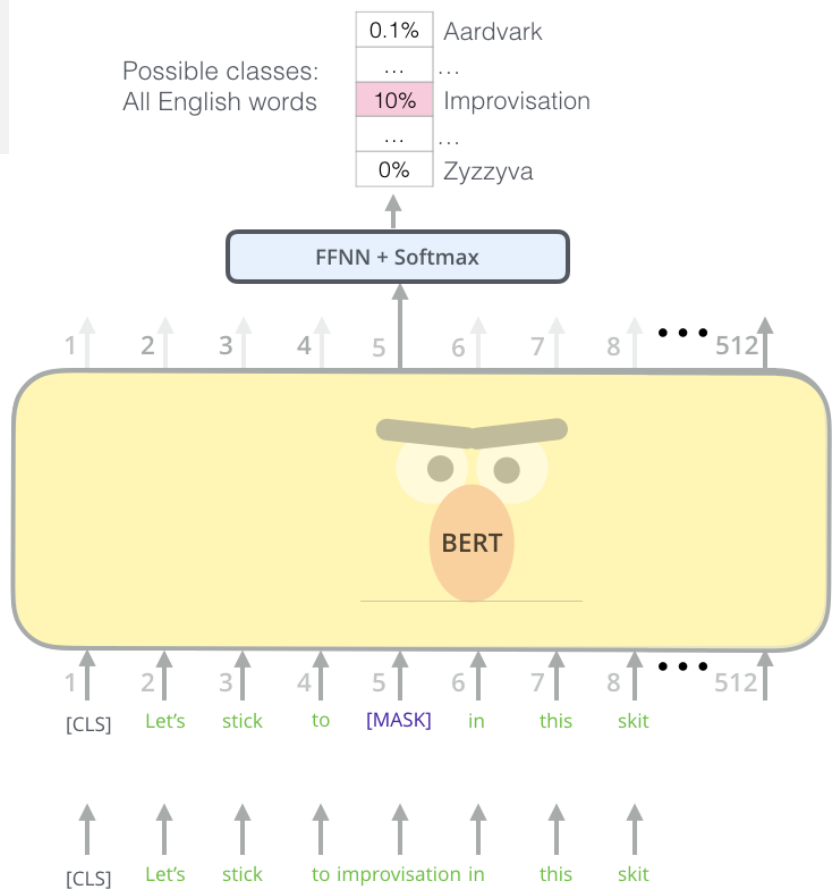
$$g_i = -\log(-\log(\text{Uniform}(0,1)))$$

$$y_i = \frac{\exp\left(\frac{\log(\pi_i) + g_i}{\tau}\right)}{\sum_{j=1}^N \exp\left(\frac{\log(\pi_j) + g_j}{\tau}\right)}$$



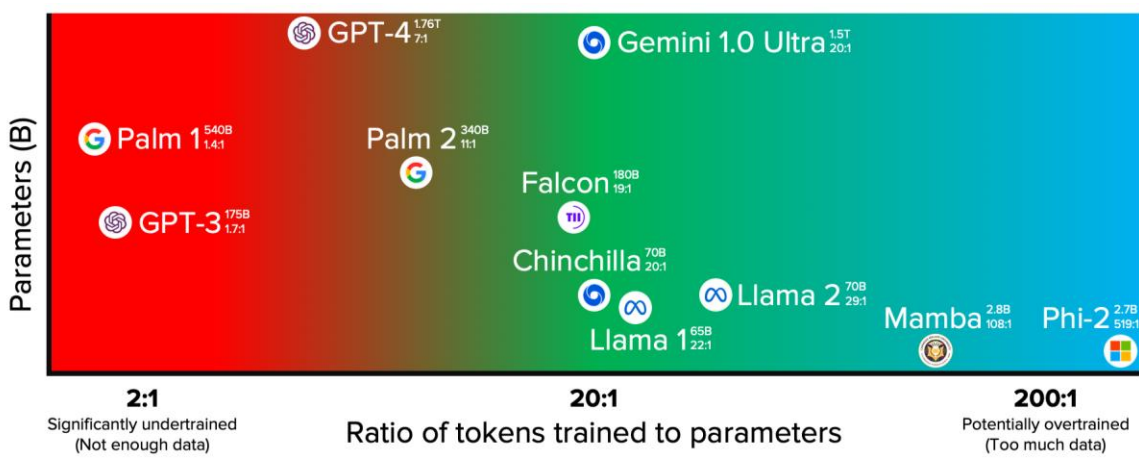
Self-supervised training

- train supervised, but **don't provide labels**
 - use naturally occurring labels
 - create labels automatically somehow
 - use specific tasks that don't require manual labels
- good to train on huge amounts of data
 - language modelling
 - **next-word prediction**
 - **MLM** – masked word prediction (~like word2vec)
 - masked span prediction: MLM for multiple words
 - **autoencoding**: predict your own input
 - denoising: corrupt data & learn to fix them
 - learn from rule-based annotation (not ideal)
- unsupervised, but with supervised approaches



Pretraining & Finetuning: Pretrained LMs

- 2-step training:
 1. **Pretrain** a model on a huge dataset (**self-supervised**, language-based tasks)
 2. **Fine-tune** for your own task on your smaller data (**supervised**)
- ~ pretrained “contextual embeddings” (“better word2vec”, typically Transformer)
- Model capability is all about the data
 - the larger model, the more you need (“Chinchilla scaling laws”)
 - anyway the more, the better



<https://lifearchitect.ai/chinchilla/>

<https://www.harmdevries.com/post/model-size-vs-compute-overhead/>

https://twitter.com/Thom_Wolf/status/1766783830839406596

Thomas Wolf 
@Thom_Wolf

this contrarian thing I keep repeating in my "LLMs in 2024" lectures – surprisingly hard to get this message across

2 Pretraining

Our approach to pretraining is to train a standard dense transformer architecture on a heavily engineered large pretraining corpora, where our underlying assumption is that when trained on extensive data of high-enough quality, a standard architecture can exhibit advanced capability. This is to say, we may not need much architectural modification, although we have indeed conducted extensive preliminary architectural experiments. In the following subsections, we first detail our data engineering pipeline, then briefly discuss the model architecture.

Thomas Wolf 
@Thom_Wolf · Mar 10

i guess we all want to believe that models are magic

Pretrained (Large) Language Models (PLMs/LLMs)

(Zhao et al., 2023)
<http://arxiv.org/abs/2303.18223>

- **BERT/RoBERTa**: Transformer encoder
 - trained by masked language modeling, good for classification
- **GPT-2, most LLMs (GPT-3, Llama, Mistral, Gemma, Phi, Qwen...)**: decoder
 - trained by next-word prediction (=language modeling), good for generation / prompting
- **BART, T5** – encoder-decoder (many training tasks, good for generation)
- multilingual: **XLM-RoBERTa, mBART, mT5, Aya**
- many models released plug-and-play
 - **!!** others (GPT-3/ChatGPT/GPT-4, Claude... closed & API-only)
- PLM vs. LLM distinction a bit vague
 - generally >1B, but more on behavior
 - PLMs: ready to finetune
 - LLMs: ready to prompt (→ →)

<https://huggingface.co/>
<https://ollama.com/>



(((J0(J0 'yoav)))) 🌈 ✅ ...

@yoavgo

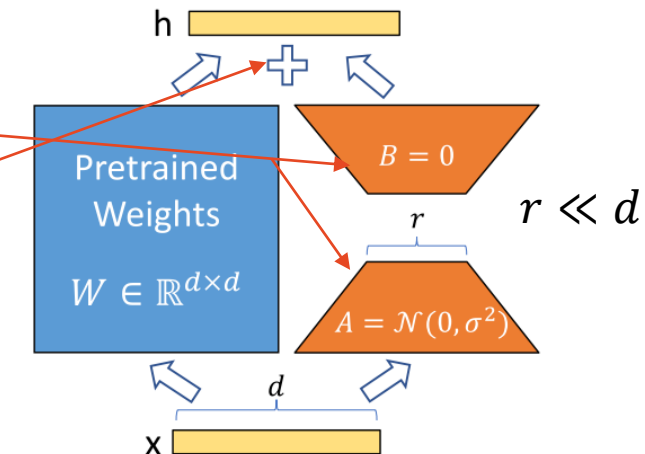
"How large should a model be to qualify as an LLM" is a vacuous question. LLMs are NOT about size, they are about having a set of behaviors that happen to correlate with those exhibited by GPT-3/ChatGPT, which were large (and not exhibited by GPT2, BERT, T5, which were smaller).

(controversial! see discussion 🎬) <https://x.com/yoavgo/status/1828383882317549765>

Parameter-efficient Finetuning

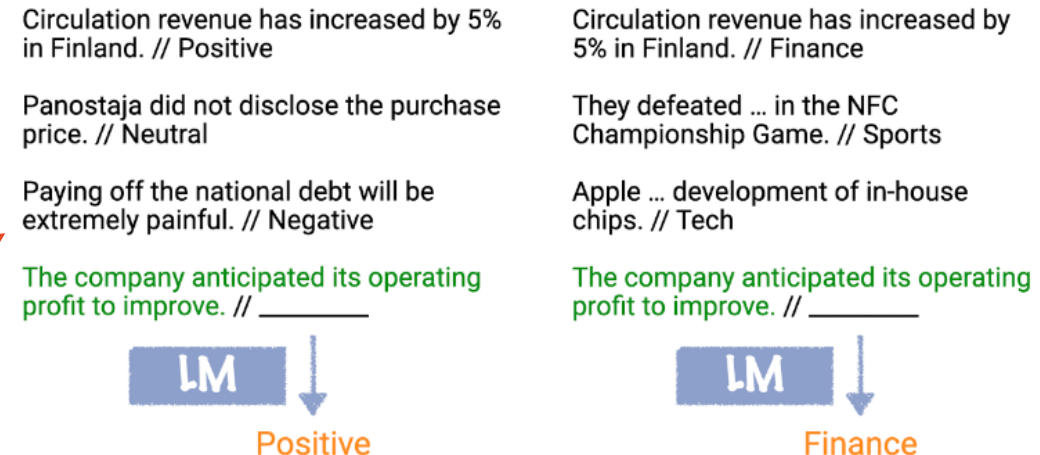
(Lialin et al., 2023)
<http://arxiv.org/abs/2303.15647>
(Sabry & Belz, 2023)
<http://arxiv.org/abs/2304.12410>

- Finetuning large models: don't update all parameters
 - faster, less memory-hungry (fewer gradients/momentums etc.)
 - trains faster
 - less prone to overfitting (~ regularization)
- Add few parameters & only update these
 - **Adapters** – small feed-forward networks after/on top of each layer
 - **Soft prompts** – tune a few special embeddings & use them in a prompt
 - **LoRA** (low-rank adaptation):
 - updates = 2 decomposition matrixes A, B (parallel to each layer)
 - update = multiplication AB
 - $2 \times r \times d$ is much smaller than full weights (d^2)
 - update is added to original weights on the fly
 - **QLoRA** – LoRA + quantized 4/8-bit computation
 - to fit large models onto a small GPU



LLMs: Prompting = In-context Learning

- No model finetuning, just show a few examples in the input (=prompt)
- pretrained LMs can do various tasks, given the right prompt
 - they've seen many tasks in training data
 - only works with the larger LMs (>1B)
- adjusting prompts often helps
 - **“prompt engineering”**
 - zero-shot (no examples) vs. **few-shot**
 - **chain-of-thought**
prompting: “let's think step by step”
 - adding / rephrasing **instructions** (see → →)



<http://ai.stanford.edu/blog/understanding-incontext/>

The diagram shows two examples of how a chain-of-thought prompt can guide an LLM to provide a correct answer.

Example 1: Q: A juggler can juggle 16 balls. Half of the balls are golf balls, and half of the golf balls are blue. How many blue golf balls are there?
A: The answer (arabic numerals) is
(Output) 8 X

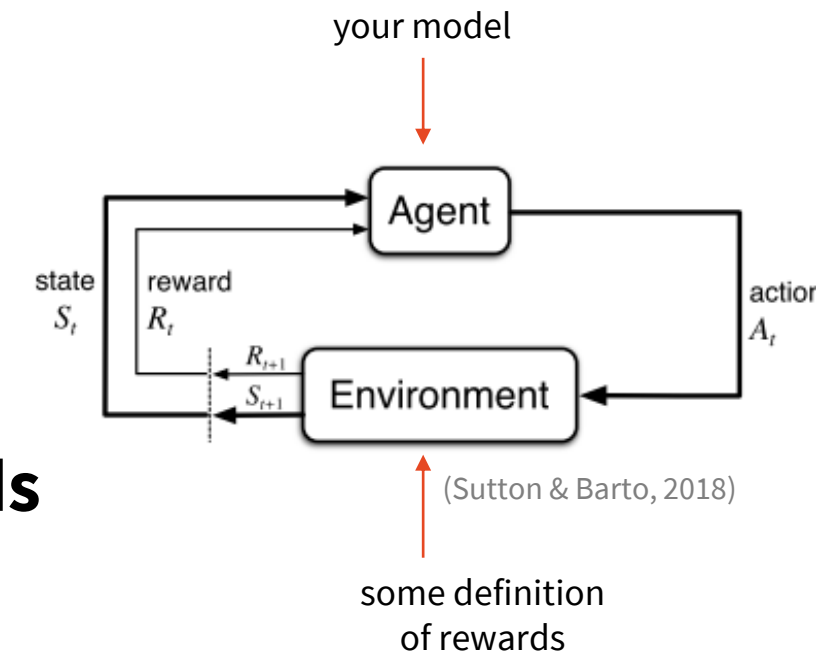
Example 2: Q: A juggler can juggle 16 balls. Half of the balls are golf balls, and half of the golf balls are blue. How many blue golf balls are there?
A: **Let's think step by step.**
(Output) There are 16 balls in total. Half of the balls are golf balls. That means that there are 8 golf balls. Half of the golf balls are blue. That means that there are 4 blue golf balls. ✓

<https://lilianweng.github.io/posts/2023-03-15-prompt-engineering/>

(Liu et al., 2023) <https://arxiv.org/abs/2107.13586>

Reinforcement Learning

- Learning from **weaker supervision**
 - only get feedback once in a while, not for every output
 - good for globally optimizing sequence generation
 - you know if the whole sequence is good
 - you don't know if step X is good
 - sequence = e.g. sentence, dialogue
- Framing the problem as **states & actions & rewards**
 - "robot moving in space", but works for dialogue too
 - state = generation so far (sentence, dialogue state)
 - action = one generation output (word, system dialogue act)
 - defining rewards might be an issue
- Training: **maximizing long-term reward**
 - via state/action values (Q function)
 - directly – optimizing policy



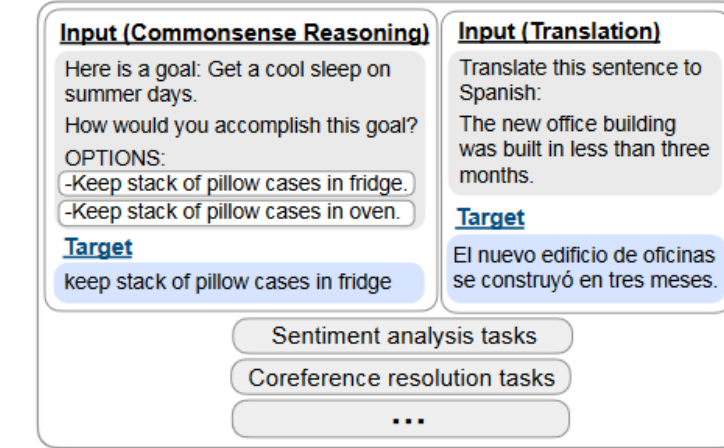
Instruction Tuning / RL from Human Feedback

(Wei et al., 2022) <https://arxiv.org/abs/2109.01652>

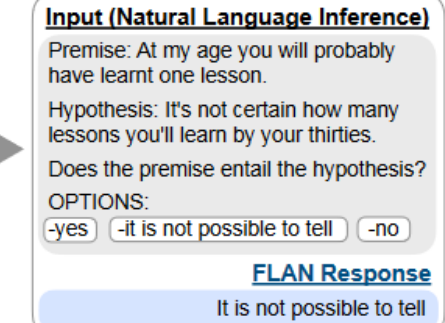
- Finetune for use with prompting
 - “in-domain” for what it’s used later
 - Use **instructions** (task description) + **solution** in prompts
 - Many different tasks
 - Specific datasets available
 - Some LLMs released as base (“foundation”) & instruction-tuned versions

- RL improvements on top of this (~InstructGPT/ChatGPT):
 - 1) generate lots of outputs for instructions
 - 2) have humans rate them
 - 3) learn a rating model (some kind of other LM: instruction + solution → score)
 - 4) use rating model score as reward in RL
 - main point: **reward is global** (not token-by-token) – RL-free alternatives exist

Finetune on many tasks (“instruction-tuning”)



Inference on unseen task type

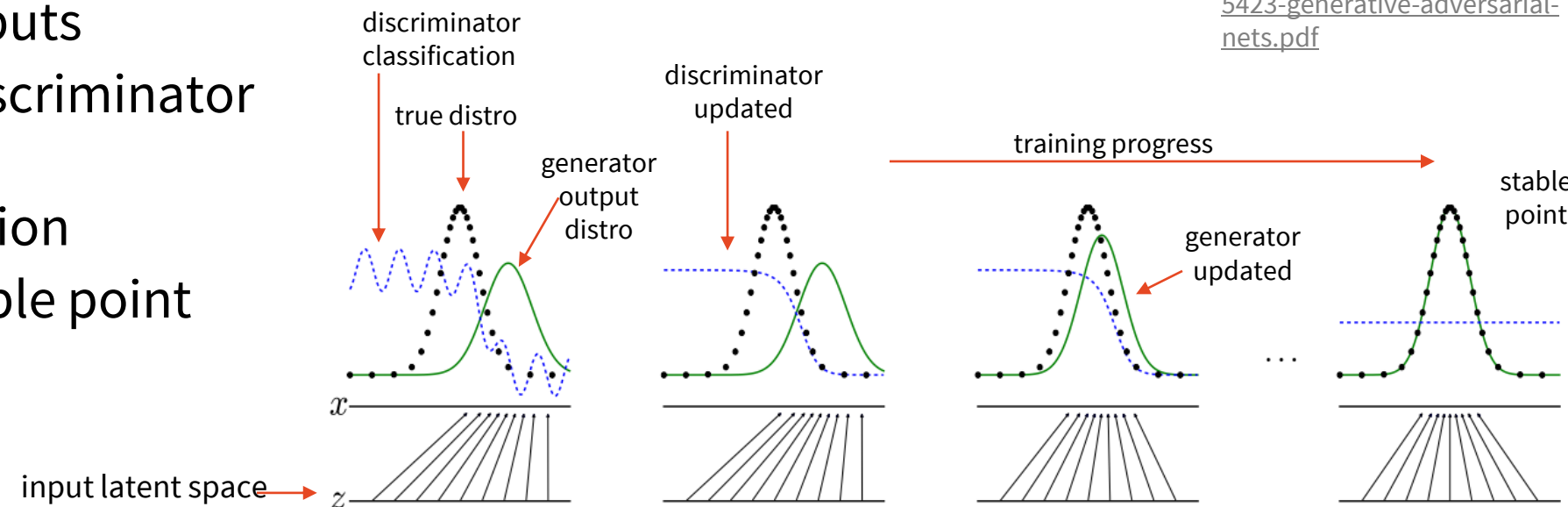


(Ouyang et al., 2022)
<http://arxiv.org/abs/2203.02155>
<https://openai.com/blog/chatgpt>

Adversarial Learning / Generative Adversarial Nets

- Training generative models to generate **believable** outputs
 - to do so, they necessarily get a better grasp on the distribution
- Getting loss from a 2nd model:
 - **discriminator D** – “adversary” classifying real vs. generated samples
 - **generator G** – trained to fool the discriminator
 - the best chance to fool the discriminator is to generate likely outputs
- Training iteratively (EM style)
 - generate some outputs
 - classify + update discriminator
 - update generator based on classification
 - this will reach a stable point

(Goodfellow et al, 2014)
<http://papers.nips.cc/paper/5423-generative-adversarial-nets.pdf>

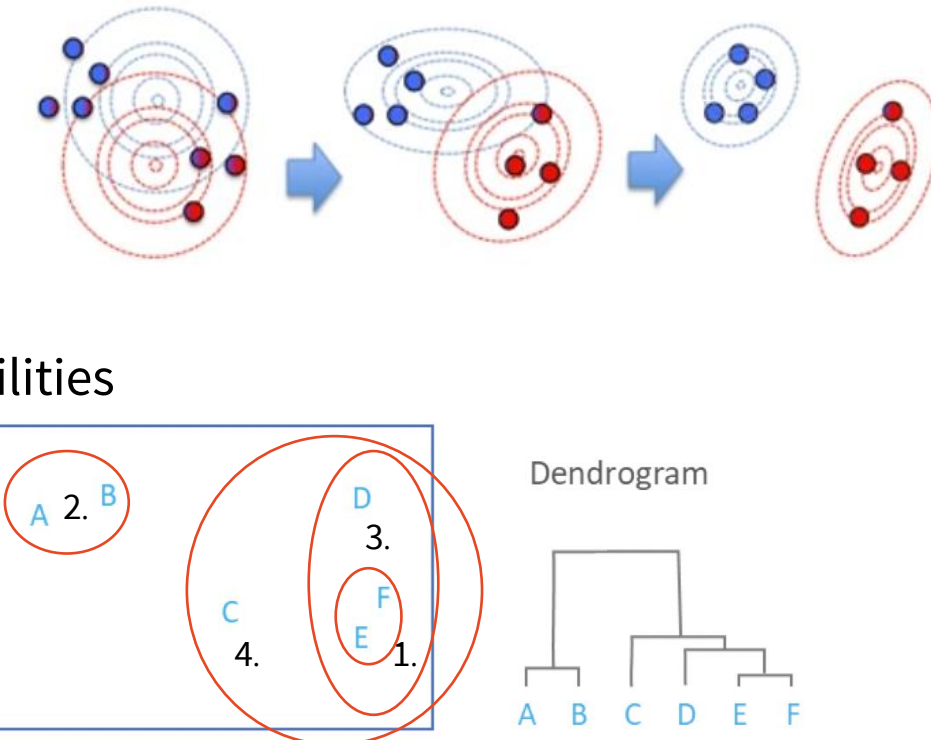


Clustering

https://en.wikipedia.org/wiki/K-means_clustering
<https://www.displayr.com/what-is-hierarchical-clustering/>
<https://towardsdatascience.com/gaussian-mixture-models-d13a5e915c8e>

- Unsupervised, finding similarities in data
- basic algorithms
 - **k-means**: assign into k clusters randomly, iterate:
 - compute means (centroids)
 - reassign to nearest centroid
 - **Gaussian mixture**: similar, but soft & variance
 - clusters = multivariate Gaussian distributions
 - estimating probabilities of belonging to each cluster
 - cluster mean/variance based on data weighted by probabilities
 - **hierarchical** (bottom up):
start with one cluster per instance, iterate:
 - merge 2 closest clusters
 - end when you have k clusters / distance is too big
 - hierarchical top-down (reversed \rightarrow)
 - distance metrics & features decide what ends up together

<https://www.youtube.com/watch?v=9YA2t78Ha68>



Summary

- Supervised training
 - cost function
 - stochastic **gradient descent** – minibatches
 - backpropagation
 - **learning rate** tricks – optimizers (Adam), schedulers
 - regularization: dropout, multi-task training
- Self-supervised learning (~kinda unsupervised)
 - autoencoders, denoising, variational autoencoders
 - (masked) language models
- PLMs/LLMs: **pretraining & finetuning, prompting, instruction tuning**
- Reinforcement learning (more to come later)
- Unsupervised: GANs, clustering

Thanks

Contact us:

<https://ufaldsg.slack.com/>

{odusek,hudecek,kasner}@ufal.mff.cuni.cz

Zoom/Skype/Troja

Labs in 10 mins

Next Thu 10:40

Get the slides here:

<http://ufal.cz/npfl099>

References/Further:

Goodfellow et al. (2016): Deep Learning, MIT Press (<http://www.deeplearningbook.org>)

Kim et al. (2018): Tutorial on Deep Latent Variable Models of Natural Language (<http://arxiv.org/abs/1812.06834>)

Milan Straka's Deep Learning slides: <http://ufal.mff.cuni.cz/courses/npfl114/1819-summer>

Neural nets tutorials:

- <https://codelabs.developers.google.com/codelabs/cloud-tensorflow-mnist/#0>
- <https://minitorch.github.io/index.html>
- <https://objax.readthedocs.io/en/latest/>

GPT in Excel: <https://www.youtube.com/watch?v=FyeN5tXMnJ8>