Neural Networks

Variational Autoencoders
Part 2
(based in part on slides from Dan Schwartz and Tom Manzini)
Recap: Neural nets as generative models

• We’ve seen how neural nets can perform classification
  – Or regression
  – MLPs, CNNs, RNNs..

• Next step: NNs as generic generative models
  – Model the distribution of any data
  – Such that we can draw samples from it
Recap: Generative models and Maximum Likelihood Estimation

What is a generative model

• In statistical estimation, a *generative model* is a functional or computational model for the probability distribution of a given data
  – Can be represented generically as $P(x; \theta)$, where $x$ represents a data instance and $\theta$ are the *parameters* of the model
  – But actually encodes a *generative story* for how the data were produced

• Utility of the model
  – Can compute the probability of observing a given value $x$
  – Can also be used to *generate* samples of (or statistically similar to) the data
Recap: Examples of Generative Models

• Generative models can be simple, one step models of generation
  – E.g. Gaussians, Multinomials

• Or a multi-step generating process
  – E.g. Gaussian Mixtures
  – E.g. Linear Gaussian Models
Recap: ML Estimation of Generative Models

• Must estimate the parameters of the model from observed data

• Maximum likelihood estimation: Choose parameters to maximize the (log) likelihood of observed data

\[
\theta^* = \arg\max_{\theta} \log(P(X; \theta)) \\
= \arg\max_{\theta} \sum_{x \in X} \log(P(x; \theta))
\]
Recap: ML estimation from incomplete data

• In many situations, our observed data are missing information
  – E.g. components of the data
  – E.g. “inside” information about how the data are drawn by the model

• In these cases, the ML estimate must only consider the observed data $O$

\[
\arg\max_\theta \sum_{o \in O} \log P(o; \theta)
\]
  – But the observed data are incomplete

• Observation probability $P(o)$ must be obtained from the complete data probability, by marginalizing out missing components
  – This can cause ML estimation to become challenging
Recap: ML estimation from incomplete data

- **ML estimate from observed data** $O$

  $$\arg\max_{\theta} \sum_{o \in O} \log P(o; \theta)$$

- $P(o)$ is obtained by marginalizing out missing components

  $$P(o; \theta) = \sum_{h} P(h, o; \theta)$$

- The ML estimate becomes

  $$\arg\max_{\theta} \sum_{o \in O} \log \left( \sum_{h} P(h, o; \theta) \right)$$

  - $h$ represent the hidden or missing components

- Minimizing the log of a sum of ugly functions usually doesn’t have nice solutions
Recap: The ELBO function and EM

• We can define an *Empirical Lower Bound* (or ELBO) for the log probability $\log(P(o))$ as:

$$
\log P(o; \theta) \geq \sum_h Q(h) \log \frac{P(h, o; \theta)}{Q(h)}
$$

  – The bound is tightest when $Q(h) = P(h|o; \theta^k)$

• We get a nice iterative ML estimator if we maximize the ELBO instead of $\log(P(o))$ directly
  – The *Expectation Maximization* algorithm
Recap: The Expectation Maximization Algorithm

• Define the auxiliary function:

\[ J(\theta, \theta^k) = \sum_{o \in O} \sum_{h} P(h|o; \theta^k) \log P(h, o; \theta) \]

• Which is the ELBO plus a term that doesn’t depend on \( \theta \)

• Iteratively compute

\[ \theta^{k+1} \leftarrow \arg \max_\theta J(\theta, \theta^k) \]

• Guaranteed to increase \( \log P(o) \) with every iteration
Recap: EM principle

• Problem: Train generative model using incomplete data
  – Gaussian from vectors with missing components
  – GMMs, where the Gaussian that generated each observation is unknown
  – Etc...

• E.M: “Complete” the data by “filling in” the missing components
  – Estimate models from “completed” data
  – Question: How to “complete” data
P(X,Y) and P(Y|X): Gaussian PDF

When you have to “guess” the missing data
P(X,Y) and P(Y|X): GMM

When you have to “guess” the missing data
Recap: EM principle

• Iteratively:
  • **Complete the data according to the posterior probabilities** $P(m|o)$ **computed by the current model**
    – By explicitly considering every possible value, with its posterior-based proportionality
    – Or by sampling the posterior probability distribution $P(m|o)$
      • Upon completion each incomplete observation implicitly or explicitly becomes many (potentially infinite) complete observations
  • **Reestimate the model from completed data**
Questions?
Principal Component Analysis

Given a (centered) set of data
find subspace such that
the projection of the data onto the subspace
results in the lowest total (squared) error

Minimize the sum of the squared lengths of these lines

- Find the principal subspace such that when all vectors are approximated as lying on that subspace, the approximation error is minimal
  - Assuming “centered” (zero-mean) data
A subspace is the set of all vectors that can be composed as a linear combination of a set of “basis” vectors.

- E.g., if $w_1$ and $w_2$ are two “basis” vectors, the set of all vectors that can be composed as $aw_1 + bw_2$ is a subspace.
  - In this case it will be a two-dimensional plane.
  - The 2-D plane in which both vectors lie.

Defining subspaces
A subspace is the set of all vectors that can be composed as a linear combination of a set of “basis” vectors.  

- E.g., if \( w_1 \) and \( w_2 \) are two “basis” vectors, the set of all vectors that can be composed as \( aw_1 + bw_2 \) is a subspace.  
  - In this case it will be a two-dimensional plane.  
  - The 2-D plane in which both vectors lie.

The same subspace can be composed by different sets of bases.  

- Any set of linearly independent vectors in the subspace.  
- The number of basis vectors is the dimensionality of the subspace, and will be the same in every set.
Given bases $W = [w_1 \ w_2]$, any vector $v$ on the subspace can be written in terms of a “position” vector $z$ such that $v = Wz$.
The “projection” of a point onto a subspace is finding the point on the subspace that is closest to it.

- Given bases $W$, the projection of any vector $v$ on the subspace is the vector $\hat{v} = Wz$ on it that is closest to $v$.

  \[
  \hat{v} = Wz \quad s.t. \quad z = \text{argmin}_{z'} ||v - Wz'||^2
  \]

  - $(v - Wz')$ is the projection error

- Trivial solution: $z = W^+ v$

The projection will always be perpendicularly below $v$ on the subspace.

- i.e. $\hat{v}^T (v - \hat{v}) = 0$
Finding the “best” subspace

• Given a data set $v_1, v_2, \ldots, v_N$, find the subspace such that the total squared projection error is minimum
  – If every data point were replaced by its projection, the total squared error incurred would be minimum

• Find $W$ and $z_i$ for every instance $v_i$ such that the total squared projection error is minimized

$$\text{argmin}_{W, \{z_i\}} \sum_i \|v_i - Wz_i\|^2$$

• If we arrange all the vectors in a matrix as $V = [v_1, v_2, \ldots, v_N]$ and their position vectors on the subspace as a matrix $Z = [z_1z_2, \ldots z_N]$, then

$$\text{argmin}_{W, Z} \sum_i \|V - WZ\|^2_F$$
Given a data set \( v_1, v_2, \ldots, v_N \), find the subspace such that the total squared projection error is minimum.

- If every data point were replaced by its projection, the total squared error incurred would be minimum.

Find \( W \) and \( z_i \) for every instance \( v_i \) such that the total squared projection error is minimized:

\[
\text{argmin}_{W, \{z_i\}} \sum_i \left| |v_i - Wz_i| \right|^2
\]

Those of you who have worked on PCA knows that the principal subspace is obtained as the space composed by the Eigen vectors of the covariance matrix of \( \{v_i\} \).
PCA is also performed by a linear-activation autoencoder

\[ X \xrightarrow{W^+} Z \xrightarrow{W} \hat{X} \approx X \]

- Given \( X = [x_1 \ldots x_N] \) be the entire training set (arranged as a matrix)
- Initialize \( W \)
- Iterate until convergence:
  - Given \( W \), find the best position vectors \( Z \): \( Z \leftarrow W^+ X \)
  - Given position vectors \( Z \), find the best subspace: \( W \leftarrow XZ^+ \)
  - Guaranteed to find the principal subspace

- The decoder matrix will be the bases of the encoder and decoder weights are decoupled
The iterative solution

- The position vector $z$ is the “missing” component of the input
  - If given, the subspace bases can be identified
- Solution: Iterative solution
  - Initialize $W$
  - Iteratively “complete” the data by estimating $z$ from $W$ and $x$, and then reestimating $W$ from the complete data
- Looks like EM
The iterative solution is EM

- The position vector $z$ is the “missing” component of the input
  - If given, the subspace bases can be identified
- Solution: Iterative solution
  - Initialize $W$
  - Iteratively “complete” the data by estimating $z$ from $W$ and $x$, and then reestimating $W$ from the complete data
- But if this is EM, what is the generative model?
The generative story behind PCA

- The generative story derives from the fact that the projection error is orthogonal to the subspace

\[-(v - Wz)^T W = 0\] for the projection \( \hat{v} = Wz \)
The *generative* story behind PCA with standard-normal constraint on z

- Generative story for PCA:
  - $z$ is drawn from a $K$-dim isotropic Gaussian
    - $K$ is the dimensionality of the principal subspace
  - $A$ is “basis” matrix
    - Matrix of principal Eigen vectors scaled by Eigen values
  - $e$ is a 0-mean Gaussian noise that is orthogonal to the principal subspace
    - The covariance of the Gaussian is low-rank and orthogonal to the principal subspace!
The generative story behind PCA

1. Alternate view: Az stretches and rotates the $K$-dimensional planar space of $z$ into a $K$-dimensional planar subspace (manifold) of the data space.

2. The circular distribution of $z$ in the $K$-dimensional $z$ space transforms into an ellipsoidal distribution on a $K$-dimensional hyperplane the data space.

3. Samples are drawn from the ellipsoidal distribution on the hyperplane, and noise is added to them.
PCA models a Gaussian distribution:

- The probability density of $x$ is Gaussian lying mostly close to a hyperplane
- With correlated structure on the plane
- And uncorrelated components orthogonal to the plane

Equations:

$$\hat{x} = Az \Rightarrow P(\hat{x}) = N(0, AA^T)$$
$$x = \hat{x} + E \Rightarrow P(x) = N(0, AA^T + D)$$
Choose all that are true about PCA

- It actually performs Maximum Likelihood estimation of a generative model for the data
- The generative model for PCA is that in order to generate any point, the process first takes a Gaussian step on the principal hyperplane, followed by a Gaussian step perpendicular to the hyperplane
- It can also be iteratively estimated using Expectation Maximization
- It assumes the distribution of the data is a Gaussian that is centered on the principal hyperplane
Choose all that are true about PCA

- It actually performs Maximum Likelihood estimation of a generative model for the data
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- It can also be iteratively estimated using Expectation Maximization
- It assumes the distribution of the data is a Gaussian that is centered on the principal hyperplane
ML estimation of PCA parameters

$z \sim N(0, I) \rightarrow Az \rightarrow \hat{x} \rightarrow x$

$e \sim N(0, D)$ \hspace{1cm} $D \perp A$

$x = Az + e$

$P(x) = N(0, AA^T + D)$

- The parameters of the PCA generative model are $A$ and $D$
- The ML estimator is

$$\arg \max_{A,D} \sum_x \log \frac{1}{\sqrt{(2\pi)^d |AA^T + D|}} \exp(-0.5x^T (AA^T + D)^{-1}x)$$

  - Where $d$ is the dimensionality of the space

- Combined with the constraints on the number of columns in $A$ (dimensions of principal subspace), and that $A^T D = 0$, this will give us the principal subspace
  - But will require EM
Missing information for PCA

- There is missing information about the observation $X$
  - Information about intermediate values drawn in generating $X$
  - We don’t know $z$

- If we knew $z$ for each $X$, estimating $A$ (and $D$) would be simple
PCA with complete information

\[ x = Az + E \]
\[ P(x|z) = N(Az, D) \]

- Given complete information \((x_1, z_1), (x_2, z_2), \ldots\)
  - Representing \(X = [x_1, x_2, \ldots], \quad Z = [z_1, z_2, \ldots]\)

\[
\arg\max_{A,D} \sum_{(x,z)} \log P(x, z) = \arg\max_{A,D} \sum_{(x,z)} \log P(x|z)
\]

\[
= \arg\max_{A,D} \sum_{(x,z)} \log \frac{1}{\sqrt{(2\pi)^d |D|}} \exp\left(-0.5(x - Az)^T D^{-1} (x - Az)\right)
\]

- Differentiating w.r.t \(A\) and equating to 0, we get the easy solution

\[ A = XZ^+ \]

  - (Some sloppy math \((D\) is not invertible), but the solution is right)

But we don’t have \(z\). It is missing
EM for PCA

- Initialize the plane
  - Or rather, the bases for the plane
- “Complete” the data by computing the appropriate zs for the plane
  - \( P(z|X; A) \) is a delta, because \( E \) is orthogonal to \( A \)
- Reestimate the plane using the zs
- Iterate
Improving on PCA

- PCA assumes the noise is always orthogonal to the data
  - Not always true
  - Noise in images can look like images, random noise can sound like speech, etc.
- Let us generalize the model to permit non-orthogonal noise
The noise added to the output of the encoder can lie in any direction
  - Uncorrelated, but not just orthogonal to the principal subspace

Generative model: to generate any point
  - Take a Gaussian step on the hyperplane
  - Add full-rank Gaussian uncorrelated noise that is independent of the position on the hyperplane
    - Uncorrelated: diagonal covariance matrix
    - Direction of noise is unconstrained
      - Need not be orthogonal to the plane
The probability distribution modelled by the LGM

\[ x = Az + e \]

- The noise added to the output of the encoder can lie in any direction
  \[ \hat{x} = Az \Rightarrow P(\hat{x}) = N(0, AA^T) \]
  \[ x = \hat{x} + E \Rightarrow P(x) = N(0, AA^T + D) \]
- The probability density of \( x \) is Gaussian lying mostly close to a hyperplane
  - With uncorrelated Gaussian
- Also
  \[ P(x|z) = N(Az, D) \]
The linear Gaussian model

- Is a generative model for Gaussians
- Data distribution are Gaussian lying largely on a hyperplane with some Gaussian “fuzz”
  - Only components on the plane are correlated with one another
    - No correlations off the plane
  - Which allows us to model *some* correlations between components
    - Halfway between a Gaussian with a diagonal covariance, and one with a full covariance
The parameters of the LGM generative model are $A$ and $D$

The ML estimator is

$$\text{argmax}_{A,D} \sum_{x} \log \frac{1}{\sqrt{(2\pi)^d |AA^T + D|}} \exp(-0.5x^T( AA^T + D)^{-1}x)$$

- Where $d$ is the dimensionality of the space

As it turns out, this does \textit{not} have a nice closed form solution
- Because $D$ is full rank

Will require EM
There is missing information about the observation $X$
- Information about intermediate values drawn in generating $X$
  - We don’t know $z$

If we knew the $z$ for each $X$, estimating $A$ (and $D$) would be very simple
LGM with complete information

\[ x = Az + e \]
\[ P(x|z) = N(Az, D) \]

- Given complete information \( X = [x_1, x_2, \ldots], \ Z = [z_1, z_2, \ldots] \)

\[
\arg\max_{A,D} \sum_{(x,z)} \log P(x, z) = \arg\max_{A,D} \sum_{(x,z)} \log P(x|z)
\]
\[
= \arg\max_{A,D} \sum_{(x,z)} \log \frac{1}{\sqrt{(2\pi)^d |D|}} \exp\left(-0.5(x - Az)^T D^{-1} (x - Az)\right)
\]
\[
= \arg\max_{A,D} \sum_{(x,z)} -\frac{1}{2} \log|D| -0.5(x - Az)^T D^{-1} (x - Az)
\]

- Differentiating w.r.t. \( A \) and \( D \) equating to 0, we get an easy solution
LGM with complete information

\[
\arg\max_{A,D} \sum_{(x,z)} -\frac{1}{2} \log|D| - 0.5(x - Az)^T D^{-1} (x - Az)
\]

- Differentiating w.r.t $A$ and $D$ and equating to 0, we get an easy solution
- Solution for $A$

\[
\nabla_A \sum_{(x,z)} 0.5(x - Az)^T D^{-1} (x - Az) = 0 \Rightarrow
\]

\[
\sum_{(x,z)} (x - Az)z^T = 0 \Rightarrow A = \left( \sum_{(x,z)} xz^T \right) \left( \sum_z zz^T \right)^{-1}
\]

- Solution for $D$

\[
\nabla_D \sum_{(x,z)} \frac{1}{2} \log|D| + 0.5(x - Az)^T D^{-1} (x - Az) = 0 \Rightarrow
\]

\[
D = \text{diag} \left( \frac{1}{N} \left( \sum_x xx^T - A \sum_{(x,z)} xz^T \right) \right)
\]
LGM with complete information

\[
\arg\max_{A,D} \sum_{(x,z)} -\frac{1}{2} \log|D| - 0.5 (x - Az)^T D^{-1} (x - Az)
\]

• Differentiating w.r.t \(A\) and \(D\) and equating to 0, we get an easy solution
• Solution for \(A\)

\[
\nabla_A \sum_{(x,z)} 0.5 (x - Az)^T D^{-1} (x - Az) = 0 \implies 
\]

\[
\sum_{(x,z)} (x - Az)z^T = 0 \implies A = \left( \sum_{(x,z)} xz^T \right) \left( \sum_z zz^T \right)^{-1}
\]

• Solution for \(D\)

\[
\nabla_D \sum_{(x,z)} \frac{1}{2} \log|D| + 0.5 (x - Az)^T D^{-1} (x - Az) = 0 \implies 
\]

\[
D = \text{diag} \left( \frac{1}{N} \left( \sum_x xx^T - A \sum_{(x,z)} xz^T \right) \right)
\]

Unfortunately we do not observe \(z\). It is missing; the observations are incomplete.
Expectation Maximization for LGM

Complete the data

Option 1:
- In *every possible way* proportional to $P(z|x)$
- Compute the solution from the completed data
The posterior $P(z|x)$

- $P(x)$ is Gaussian
  - We saw this
- The joint distribution of $x$ and $z$ is also Gaussian
  - Trust me
- The conditional distribution of $z$ given $x$ is also Gaussian
  
  $$P(z|x) = N(z; A^T(AA^T + D)^{-1}x, I - A^T(AA^T + D)^{-1}A)$$
  
  - Trust me
Expectation Maximization for LGM

- **Complete** the data
- **Option 1:**
  - In *every possible way* proportional to $P(z|x)$
  - Compute the solution from the completed data

$$P(z|x) = N(z; A^T(AA^T + D)^{-1}x, I - A^T(AA^T + D)^{-1}A)$$
Expectation Maximization for LGM

- **Complete** the data in *every possible way* proportional to $P(z|\mathbf{x})$
  - Compute the solution from the completed data
    
    $$\text{argmax}_{A,D} \sum_{(x,z)} -\frac{1}{2} \log|D| - 0.5(\mathbf{x} - \mathbf{Az})^T D^{-1} (\mathbf{x} - \mathbf{Az})$$

- The $z$ values for each $x$ are distributed according to $P(z|\mathbf{x})$. Segregating the summation by $x$

  $$\text{argmax}_{A,D} \sum_{x} \int_{-\infty}^{\infty} p(z|x) \left( -\frac{1}{2} \log|D| - 0.5(\mathbf{x} - \mathbf{Az})^T D^{-1} (\mathbf{x} - \mathbf{Az}) \right) dz$$
LGM with incomplete information

\[
\text{argmax}_{A,D} \sum_x \int_{-\infty}^{\infty} p(z|x) \left( -\frac{1}{2} \log |D| - 0.5 (x - Az)^T D^{-1} (x - Az) \right) dz
\]

- Differentiating w.r.t. \( A \) and \( D \) and equating to 0, we get an easy solution
- Solution for \( A \)

\[
\nabla_A \sum_x \int_{-\infty}^{\infty} p(z|x) (x - Az)^T D^{-1} (x - Az) dz = 0 \quad \Rightarrow \\
\sum_x \int_{-\infty}^{\infty} p(z|x) (x - Az) z^T dz = 0 \quad \Rightarrow \\
A = \left( \sum_x \int_{-\infty}^{\infty} p(z|x) xz^T dz \right) \left( \sum_x \int_{-\infty}^{\infty} p(z|x) z z^T dz \right)^{-1}
\]

- Solution for \( D \)

\[
\nabla_D \left( N \log |D| + \sum_x \int_{-\infty}^{\infty} p(z|x) (x - Az)^T D^{-1} (x - Az) dz \right) = 0 \quad \Rightarrow \\
D = \text{diag} \left( \frac{1}{N} \left( \sum_x xx^T - A \sum_x \int_{-\infty}^{\infty} p(z|x) xz^T dz \right) \right)
\]

These are closed form solutions, the details of which are not relevant to us.

Key: All terms integrate over all possible completion of incomplete observations, where the proportionality attached to any completion of \( x \) is \( P(z|x) \)
LGM with incomplete information

• It is actually an iterative algorithm (EM):

• Solution for $A$

$$A^{k+1} = \left( \sum_x \int_{-\infty}^{\infty} p(z|x; A^k, D^k) x z^T \, dz \right) \left( \sum_x \int_{-\infty}^{\infty} p(z|x; A^k, D^k) z z^T \, dz \right)^{-1}$$

• Solution for $D$

$$D = \text{diag} \left( \frac{1}{N} \left( \sum_x xx^T - A \sum_x \int_{-\infty}^{\infty} p(z|x; A^k, D^k) x z^T \, dz \right) \right)$$

These are closed form solutions, the details of which are not relevant to us.

Key: All terms integrate over all possible completion of incomplete observations, where the proportionality attached to any completion of $x$ is $P(z|x)$.
Expectation Maximization for LGM

- Complete the data

- Option 2:
  - **By drawing samples from** $P(z|x)$
  - Compute the solution from the completed data

\[
P(z|x) = N(z; A^T( AA^T + D)^{-1}x, I-A^T(AA^T + D)^{-1}A)
\]
LGM from drawn samples

- Since we now have a collection of complete vectors, we can use the usual complete-data formulae.
- Solution for $A$

$$A^{k+1} = \left( \sum_{(x,z)} xz^T \right) \left( \sum_z zz^T \right)^{-1}$$

- Solution for $D$

$$D^{k+1} = \text{diag} \left( \frac{1}{N} \left( \sum_x xx^T - A^k \sum_{(x,z)} xz^T \right) \right)$$

These are closed form solutions.

Draw missing components from $P(z|x; A^k, D^k)$ to complete the data.

Estimate parameters from completed data.
LGMs: The intuition

- The linear transform stretches and rotates the K-dimensional input space onto a K-dimensional hyperplane in the data space.
- The isotropic Gaussian in the input space becomes a stretched and rotated Gaussian on the hyperplane.

\[ z \sim N(0,I) \]

\[ E \sim N(0,D) \]

\[ x = Az + E \]
LGMs: The intuition

- Drawing samples: The first step places the $z$ somewhere on the plane described by $A$
  - The distribution of points on the plane is also Gaussian
LGMs: The intuition

- **LGM model**: The first step places the $z$ somewhere on the plane described by $A$
  - The distribution of points on the plane is also Gaussian
- **Second step**: Add Gaussian noise to produce points that aren’t necessarily on the plane
  - Noise added is not revealed

\[ z \sim N(0, I) \]
\[ E \sim N(0, D) \]
\[ X = Az + E \]
EM for LGMs: The intuition

- In an LGM the way to produce any data instance is not unique
- Conversely, given only the data point, the “shadow” on the principal plane cannot be uniquely known

**Mathematical Expressions:***

\[ z \sim N(0, I) \]
\[ E \sim N(0, D) \]
\[ X = Az + E \]

**Diagram:**
- Red arrows are different possibilities for \( E \)
- Blue arrows are different possibilities for \( \hat{X} \)

**Equation:**
\[ X = \hat{X} + E \]
$z \sim N(0, I)$

$E \sim N(0, D)$

$x = Az + E$

- The posterior probability $P(z|x)$ gives you the location of all the points on the plane that could have generated $x$ and their probabilities.
• Attach the point to *every* location on the plane, according to $P(z|x)$
  – Or to a sample of points on the plane drawn from $P(z|x)$
• There will be more attachments where $P(z|x)$ is higher, and fewer where it is lower

\[
\begin{align*}
z &\sim N(0, I) \\
E &\sim N(0, D) \\
X &= Az + E
\end{align*}
\]
• Attach *every* training point in this manner
• Let the plane rotate and stretch until the total tension (sum squared length) of all the attachments is minimize
• Repeat attachment and rotation until convergence...
Summarizing LGMs

- LGMs are models for *Gaussian* distributions
- Specifically, they model the distribution of data as Gaussian, where most of the variation is along a *linear* manifold
  - They do this by transforming a Gaussian RV $z$ through a linear transform $f(z) = Az$ that transforms the $K$-dim input space of $z$ into a $K$-dimensional hyperplane (linear manifold) in the data space

- They are excellent models for data that actually fit these assumptions
  - Often, we can simply assume that data lie near linear manifolds and model them with LGMs
  - PCA, an instance of LGMs, is very popular
Choose all that are true about Factor Analysis (Linear Gaussian Models)

- It models the distribution of the data as a Gaussian centered on a principal (hyper)plane
- The generative model is that in order to generate any point, the process first takes a Gaussian step on the principal hyperplane, followed by addition of Gaussian noise.
- The parameters of the distribution are the bases of the hyperplane and the covariance of the noise.
- The parameters can be easily estimated if the location of the first step on the principal hyperplane is known for every data point.
- The actual estimation is performed using EM, which iteratively “completes” each data instance with the location of this first step, and then estimates the parameters.
Choose all that are true about Factor Analysis (Linear Gaussian Models)

- It models the distribution of the data as a Gaussian centered on a principal (hyper)plane.
- The generative model is that in order to generate any point, the process first takes a Gaussian step on the principal hyperplane, followed by addition of Gaussian noise.
- The parameters of the distribution are the bases of the hyperplane and the covariance of the noise.
- The parameters can be easily estimated if the location of the first step on the principal hyperplane is known for every data point.
- The actual estimation is performed using EM, which iteratively “completes” each data instance with the location of this first step, and then estimates the parameters.
Where LGMs fail

• What about data that are not Gaussian distributed close to a plane
  – The distributions lie close to a curved or otherwise non-linear manifold?
• You can model these as Gaussian data centered on a plane that has been warped into the observed shape
The non-linear Gaussian model

\[ x = f(z) + e \]

- The non-linear function that produces a curved manifold
  - Like the decoder of a non-linear AE
- The samples of \( z \) are placed on this curved manifold
- The actual data are produced by adding noise to samples on the manifold
NLGMs

• **Step1**: Find a function that warps a lower-dimensional input plane to the target manifold in the data space
  – The non-linear version of the linear transform in the LGM
NLGMs

• **Step 1:** Find a function that warps a lower-dimensional input plane to the target manifold in the data space
  – The non-linear version of the linear transform in the LGM

• **Step 2:** Transform a Gaussian distribution on the input plane to a distribution on the curved manifold
**NLGMs**

- **Step 1:** Find a function that warps a lower-dimensional input plane to the target manifold in the data space
  - The non-linear version of the linear transform in the LGM

- **Step 2:** Transform a Gaussian distribution on the input plane to a distribution on the curved manifold

- **Step 3:** Add some uncorrelated Gaussian “fuzz” to account for off-manifold variations
The non-linear function warps the input space into a curved manifold in the data space.
NLGMs

The non-linear function warps the input space into a curved manifold in the data space

- Samples drawn from $z$ are placed on this manifold
- The distribution of $\hat{x}$ on the manifold will follow the distribution of $z$
  - High-density regions of $\hat{x}$ correspond to high-density regions of $z$
The non-linear function warps the input space into a curved manifold in the data space:
- Samples drawn from $z$ are placed on this manifold.
- The distribution of $\hat{x}$ on the manifold will follow the distribution of $z$.
  - High-density regions of $\hat{x}$ correspond to high-density regions of $z$.
- The final observations are obtained by adding uncorrelated full-dimensional Gaussian noise to the samples.
NLGM Generating Process

\[ x = f(z) + e \]

- Generating process:
  - Draw a sample \( z \) from a Uniform Gaussian
  - Transform \( z \) by \( f(z) \)
    - This places \( z \) on the curved manifold

\[ z \sim N(0, I) \]
\[ f(z) \]
\[ \hat{x} \]
\[ x \]
\[ e \sim N(0, D) \]

Drawn \( z \)
NLGMs Generating Process

- Generating process: Draw a sample $z$ from a Uniform Gaussian
  - Draw a sample $z$ from a Uniform Gaussian
  - Transform $z$ by $f(z)$
    - This places $z$ on the curved manifold
  - Add uncorrelated Gaussian noise to get the final observation
• The NLGM can model very complicated distributions
  – Distributions that may be viewed as lying close to a curved $K$-dimensional surface in the data space
    • Or even a linear surface: $f(z) = Az$ is a special case
    • $K$ is the dimensionality of $z$
The NLGM can model very complicated distributions

- Distributions that may be viewed as lying close to a curved $K$-dimensional surface in the data space
  - Or even a linear surface: $f(z) = Az$ is a special case
  - $K$ is the dimensionality of $z$

Key requirement:

- Identifying the dimensionality $K$ of the curved manifold
- Having a function $f(z)$ that can transform the (linear) $K$-dimensional input space (space of $z$) to the desired $K$-dimensional manifold in the data space
Designing NLGMs

- Key design issues:
  - Select (or guess) the dimensionality of the manifold
    - This is the dimensionality of $z$
  - Choosing the right function $f(z)$ that is capable of learning the shape of the manifold
    - We will choose a Neural Network $f(z; \theta)$
Learning the NLGM

- Given a collection of training data $X = \{x\}$
  - Estimate the parameters $\theta$ of $f(z; \theta)$
  - Estimate $D$
- The NLGM is a generative model that actually models a distribution
  - The distribution obtained when $N(0, I)$ is transformed by $f(z)$
- We will use ML estimation to learn its parameters to best match the training data
Probabilities modelled by the NLGM

\[ x = f(z) + e \]

- The conditional probability of \( x \) given \( z \)
  \[ P(x|z) = N(x; f(z; \theta), D) \]

- The marginal probability of \( x \)
  \[ P(x) = \int_{-\infty}^{\infty} P(x|z)P(z)dz = \int_{-\infty}^{\infty} N(x; f(z; \theta), D) N(z; 0, D) dz \]
  - For most nonlinear functions \( f(z; \theta) \) this math is not tractable, and we cannot get a closed form for \( P(x) \)
  - \textit{That won’t prevent us from being able to estimate \( \theta \) and \( D \)}
Choose all that are true of Nonlinear Gaussian Models

- They are generative models
- They model the distribution of the data as Gaussian distributed about a curved manifold
- The generative model is that in order to generate any point, the process first takes a Gaussian step on along the curved manifold, followed by the addition of Gaussian noise
- The parameters of the distribution are the parameters of the function that transform a K-dimensional plane into a K-dimensional manifold, and the covariance of the noise
- The NLGM is the decoder component of a variational autoencoder
Choose all that are true of Nonlinear Gaussian Models

- They are generative models
- They model the distribution of the data as Gaussian distributed about a curved manifold
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- The NLGM is the decoder component of a variational autoencoder
Learning the NLGM with *complete data*

- Drawing a sample from the NLGM is a two-step process
  - First a $z$ is drawn
    - And transformed
  - Then an $e$ is drawn
    - And added

- The complete data to describe any draw are the outcomes of every stage of the drawing process, i.e. $(x, z)$
  - Actually $(x, e, z)$, but let’s work without $e$
NLGM with complete data

- Let us first consider a *glass-box* process that gives us *complete* data
  - The output and the intermediate steps of the generation process
  - I.e. both the $x$ and the $z$ for every draw

- We will derive estimation rules for the model parameters using the complete data
ML estimation with complete information

Given complete information $X = [x_1, x_2, ...], \ Z = [z_1, z_2, ...]$

\[
\theta^*, D^* = \arg\max_{\theta, D} \sum_{(x,z)} \log P(x, z) = \arg\max_{\theta, D} \sum_{(x,z)} \log P(x|z)
\]

\[
= \arg\max_{\theta, D} \sum_{(x,z)} \log \frac{1}{\sqrt{(2\pi)^d |D|}} \exp\left(-0.5(x - f(z; \theta))^T D^{-1} (x - f(z; \theta))\right)
\]

\[
= \arg\max_{\theta, D} \sum_{(x,z)} -\frac{1}{2} \log|D| - 0.5(x - f(z; \theta))^T D^{-1} (x - f(z; \theta))
\]
There isn’t a nice closed form solution, but we could learn the parameters using backpropagation, which minimizes the following loss:

$$L(\theta, D) = \sum_{(x,z)} \frac{1}{2} \log |D| + 0.5 (x - f(z; \theta))^T D^{-1} (x - f(z; \theta))$$

$$\theta^*, D^* = \arg\min_{\theta, D} L(\theta, D)$$

Unfortunately we do not observe $z$. It is missing; the observations are incomplete.
NLGM with incomplete data

• We could estimate the model parameters if $z$ were known for every data observation
  – I.e. if the data were complete
• Unfortunately we don’t know $z$
  – The data are incomplete
• Solution: EM!
  – Complete the data
**Expectation Maximization for NLGM**

- **Complete** the data
- **Option 1:**
  - In every possible way proportional to $P(z|x)$
  - Compute the solution from the completed data
**Expectation Maximization for LGM**

- **Complete** the data

- **Option 2:**
  - *By drawing samples from* $P(z|x)$
  - Compute the solution from the completed data
**Expectation Maximization for LGM**

- **Complete** the data
- **Option 2:**
  - By drawing samples from $P(z|x)$
  - Compute the solution from the completed data

Using every possible value (option 1) is to be preferred over sampling (option 2) if the former produces tractable closed form solutions. Otherwise we must use option 2.
Problem with completing the data

• The posterior probability is given by

\[
P(z|x) = \frac{P(x|z)P(z)}{P(x)}
\]

• The denominator

\[
P(x) = \int_{-\infty}^{\infty} N(x; f(z; \theta), D) N(z; 0, D) \, dz
\]

  – This is intractable to compute in closed form for most \( f(z; \theta) \)

• \( P(z|x) \) is intractable as a closed form solution
  
  – Makes it challenging to integrate over it or draw samples from it
  – But we could try to *approximate it*
Approximating $P(z|x)$

- We will approximate $P(z|x)$ as

$$P(z|x) \approx Q(z, x) = \text{Gaussian } N(z; \mu(x), \Sigma(x))$$

  - where $\mu(x; \varphi)$ and $\Sigma(x; \varphi)$ are estimated such that $Q(z, x)$ approximates $P(z|x)$ as closely as possible
  - For convenience, we will assume $\Sigma(x; \varphi)$ is a diagonal matrix, represented entirely by its diagonal elements

- We will use $Q(z, x)$ as our proxy for $P(z|x)$
Overall Solution

- Initialize $f(z; \theta)$
- Iterate:
  - Estimate $\mu(x; \varphi)$ and $\Sigma(x; \varphi)$ to give you the best $Q(x, z)$
  - “Complete” the data using $Q(z, x)$
  - Reestimate $f(z; \theta)$
The complete pipeline

• Initialize $\theta$ and $\varphi$

• Iterate:
  – Sample $z$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
    • “Completing” the data
  – Reestimate $\theta$ from the entire “complete” data
  – Estimate $\varphi$ using the entire “complete” data
The complete pipeline

• Initialize $\theta$ and $\varphi$

• Iterate:
  – Sample $z$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
    • “Completing” the data
  – Reestimate $\theta$ from the entire “complete” data
  – Estimate $\varphi$ using the entire “complete” data
Sampling $z$

- Sample $z$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
  - “Completing” the data

- We use a standard “reparametrization” step to sample $z$
  - Sample $z$ from a standard Gaussian, and scale and shift it such that it appears as a sample from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$

- For each training instance $x$
  - Compute $\mu(x; \varphi)$ and $\Sigma(x; \varphi)$
  - Draw one or more samples from the Gaussian $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$
    - Draw $K$-dimensional vector $\epsilon$ from $N(0, I)$
    - Compute $z = \mu(x; \varphi) + \Sigma(x; \varphi)^{0.5} \epsilon$
Sampling $z$

- **Sample $z$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance**
  - “Completing” the data

- We use a standard “reparametrization” step to sample $z$
  - Sample $z$ from a standard Gaussian, and scale and shift it such that it appears as a sample from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$

- For each training instance $x$
  - Compute $\mu(x; \varphi)$ and $\Sigma(x; \varphi)$
  - Draw one or more samples from the Gaussian $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$
    - Draw $K$-dimensional vector $\varepsilon$ from $N(0, I)$
    - Compute $z = \mu(x; \varphi) + \Sigma(x; \varphi)^{0.5} \varepsilon$

Remember this one  
\[ \nabla_\varphi z = \nabla_\varphi \mu(x; \varphi) + \text{diag}(\varepsilon) \nabla_\varphi \Sigma(x; \varphi)^{0.5} \]

This will be specific to $x$ and to the specific sample of $z$ for that $x$ (via $\varepsilon$)
The complete pipeline

- Initialize $\theta$ and $\varphi$

- Iterate:
  - Sample $z$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
    - "Completing" the data
  - Reestimate $\theta$ from the entire "complete" data
  - Estimate $\varphi$ using the entire "complete" data

We know how to do this

\[ e \sim N(0, D) \]
\[ f(z; \theta) \]
\[ N(z; \mu(x), \Sigma(x)) \]
\[ \mu(x; \varphi) \]
\[ \Sigma(x; \varphi) \]
### NLGM with complete information

\[\theta^*, D^* = \arg \max_{\theta, D} \sum_{(x,z)} -\frac{1}{2} \log |D| - 0.5 (x - f(z; \theta))^T D^{-1} (x - f(z; \theta))\]

- We can learn the parameters using backpropagation, which minimizes the following loss

\[L(\theta, D) = \sum_{(x,z)} \log |D| + (x - f(z; \theta))^T D^{-1} (x - f(z; \theta))\]

\[\theta^*, D^* = \arg \min_{\theta, D} L(\theta, D)\]

- It is common to assume that all the (diagonal) entries of \(D\) are identical, with value \(\sigma^2\)

\[L(\theta, \sigma^2) = d \log \sigma^2 + \sum_{(x,z)} \frac{1}{\sigma^2} ||x - f(z; \theta)||^2\]

- The derivative of this w.r.t \(\theta\) and \(\sigma^2\) is trivially computed for backprop
The complete pipeline

• Initialize $\theta$ and $\varphi$

• Iterate:
  – Sample $z$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
    • “Completing” the data
    – Reestimate $\theta$ from the entire “complete” data
    – Estimate $\varphi$ using the entire “complete” data

We know how to do this
Approximating $P(z|x)$ by $Q(z, x)$

- Recall $Q(z, x) = N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ must approximate $P(z|x)$ as closely as possible.

- Estimate $\varphi$ to minimize the error between $Q(z, x)$ and $P(z|x)$
  - Define a divergence between $Q(z, x)$ and $P(z|x)$ and minimize it w.r.t. $\varphi$
  - Following the literature, we will use the KL divergence
    - Then I will give you a simpler explanation
Approximating $P(z|x)$ by $Q(z, x)$

$$KL(Q(z, x)P(z|x)) = E_{z \sim Q} \log \frac{Q(z, x)}{P(z|x)}$$

$$= E_{z \sim Q} \log Q(z, x) - E_{z \sim Q} \log P(z|x)$$

$$= E_{z \sim Q} \log Q(z, x) - E_{z \sim Q} \log \frac{P(z)P(x|z)}{P(x)}$$

$$= E_{z \sim Q} \log Q(z, x) - E_{z \sim Q} \log P(z) - E_{z \sim Q} \log P(x|z) + E_{z \sim Q} \log P(x)$$

$$= KL(Q(z, x), P(z)) - E_{z \sim Q} \log P(x|z) + E_{z \sim Q} \log P(x)$$

- $Q(z, x)$ is a function of $\varphi$. Minimizing the loss w.r.t. $\varphi$ we get

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

$$= \arg\min_{\varphi} KL(Q(z, x), P(z)) - E_{z \sim Q} \log P(x|z)$$

Find $\varphi$ to minimize the (empirical estimate of the) KL divergence between $Q(z,x)$ and $P(z)$ while simultaneously maximizing the (empirical estimate of) the expectation of $\log P(x|z)$
Let’s try that again...
NLGM with complete data

• Assume we have completed the data using $Q(z, x)$
  – We have a collection of $(x, z)$ pairs
    • More precisely denoted as $(x, z_{x,j})$ since the value $z_{x,j}$ used to complete the observation is specific to $x$ (subscript $x$)
      – Also, a single $x$ may be completed in multiple ways (subscript $j$)
    • We will use $(x, z)$ as our shorthand notation, though
  – We can work with complete data!
Representing the completed data as \([X, Z] = \{(x, z)\}\), the actual posterior probability for \(Z\) given \(X\) as computed by the model is

\[
P(Z|X; \theta) = \prod_{(x,z)\in[X,Z]} P(z|x; \theta),
\]

- Because the observations are independent

The approximation using \(Q(Z, X)\) is

\[
Q(Z, X; \varphi) = \prod_{(x,z)\in[X,Z]} Q(z, x; \varphi),
\]

\[
\log Q(Z, X; \varphi) = \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi)
\]
NLGM with complete data

Representing the completed data as \([X, Z] = \{(x, z)\}\), the actual posterior probability for \(Z\) given \(X\) as computed by the model is

\[
\log P(Z|X; \theta) = \sum_{(x,z) \in [X,Z]} \log P(z|x; \theta)
\]

\[
\log Q(Z,X; \phi) = \sum_{(x,z) \in [X,Z]} \log Q(z,x; \phi)
\]

We will estimate \(\phi\) to minimize the discrepancy between these two probabilities.
Estimating $\varphi$

- We will minimize the following error

$$\log Q(Z, X; \varphi) - \log P(Z|X; \theta)$$

$$= \sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z|x; \theta)$$
Estimating $\varphi$

- We will minimize the following error
  \[ \log Q(Z, X; \varphi) - \log P(Z|X; \theta) \]
  \[ = \sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z|x; \theta) \]

- By Bayes rule $P(z|x; \theta) = P(z)P(x|z)/P(x)$
- The error becomes
  \[ \sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta) + \log P(x; \theta) \]
Estimating $\varphi$

- We will minimize the following error
  \[ \log Q(Z, X; \varphi) - \log P(Z|X; \theta) \]
  \[ = \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi) - \log P(z|x; \theta) \]

- By Bayes rule $P(z|x; \theta) = P(z)P(x|z)/P(x)$
- The error becomes
  \[ \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta) + \log P(x; \theta) \]

- $\varphi$ influences $Q(z, x; \varphi)$ directly and $z$, because it is sampled from $Q(z, x; \varphi)$. $P(x; \theta)$ is not related to either $\varphi$ or $z$ and can be ignored.
Estimating $\varphi$

- We will minimize the following error
  $$\log Q(Z, X; \varphi) - \log P(Z|X; \theta)$$
  $$= \sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z|x; \theta)$$

- By Bayes rule $P(z|x; \theta) = P(z)P(x|z)/P(x)$
- The error becomes
  $$\sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta) + \log P(x; \theta)$$

- $\varphi$ influences $Q(z, x; \varphi)$ directly and $z$, because it is sampled from $Q(z, x; \varphi)$. $P(x; \theta)$ is not related to either $\varphi$ or $z$ and can be ignored.
- This gives us the loss function
  $$L_Q(\varphi) = \sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta)$$

- This must be minimized w.r.t. $\varphi$
Estimating $\varphi$

- We will minimize the following error
  \[
  \log Q(Z, X; \varphi) - \log P(Z|X; \theta)
  \]
  \[
  = \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi) - \log P(z|x; \theta)
  \]
- By Bayes rule $P(z|x; \theta) = P(z)P(x|z)/P(x)$
- The error becomes
  \[
  \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta) + \log P(x; \theta)
  \]
- $\varphi$ influences $Q(z, x; \varphi)$ directly and $z$, because it is sampled from $Q(z, x; \varphi)$.
- $P(x; \theta)$ is not related to either $\varphi$ or $z$ and can be ignored.
- This gives us the loss function
  \[
  L_Q(\varphi) = \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta)
  \]
- This must be minimized w.r.t. $\varphi$
Estimating $\varphi$

• The loss function

$$L_Q(\varphi) = \sum_{(x,z)\in[X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta)$$

• Multiple choices for data completion

• Simple option: Simply use samples drawn from $Q(z, x; \varphi)$
  – You can skip the next couple of slides if you do

$$\nabla_{\varphi} L_Q(\varphi) = \sum_{(x,z)\in[X,Z]} \nabla_{\varphi} \log Q(z, x; \varphi) + \nabla_z \log Q(z, x; \varphi) \nabla_{\varphi} z$$

$$- \nabla_z \log P(z) \nabla_{\varphi} z - \nabla_z \log P(x|z; \theta) \nabla_{\varphi} z$$
Estimating $\phi$

- The loss function

$$L_Q(\phi) = \sum_{(x,z) \in [X,Z]} \log Q(z, x; \phi) - \log P(z) - \log P(x|z; \theta)$$

- Multiple choices for data completion
- Simple option: Simply use samples drawn from $Q(z, x; \phi)$
  - You can skip the next couple of slides if you do

- Orrrrr try to be more precise....
Expectation Maximization for LGM

Using every possible value (option 1) is to be preferred over sampling (option 2) if the former produces tractable closed form solutions. Otherwise we must use option 2.

- **Complete** the data
  - Option 1: Consider *every possible value* for $z$
  - Option 2: *By drawing samples from* $P(z|x)$
- Compute the solution from the completed data
Estimating $\varphi$

- The loss function

$$L_Q(\varphi) = \sum_{(x, z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta)$$

- It turns out that the portion underlined in blue can be computed in closed form if you consider every possible value of $z$

- The portion underlined in red cannot

- So sum the first portion over all possible values of $z$ from and the second one over only the drawn samples

$$L_Q(\varphi) = \sum_{(x) \in [X]} \int_{-\infty}^{\infty} Q(z, x; \varphi)(\log Q(z, x; \varphi) - \log P(z)) \, dz - \sum_{(x, z) \in [X,Z]} \log P(x|z; \theta)$$
Estimating $\varphi$

- The loss function

$$L_Q(\varphi) = \sum_{(x,z) \in [X,Z]} \log Q(z, x; \varphi) - \log P(z) - \log P(x|z; \theta)$$

- It turns out that the portion underlined in blue can be computed in closed form if you consider every possible value of $z$

- The portion underlined in red cannot

- So sum the first portion over all possible values of $z$ from and the second one over only the drawn samples

$$L_Q(\varphi) = \sum_{(x) \in [X]} \int_{-\infty}^{\infty} Q(z, x; \varphi) (\log Q(z, x; \varphi) - \log P(z)) \, dz - \sum_{(x,z) \in [X,Z]} \log P(x|z; \theta)$$

$$KL(Q(z, x; \varphi), P(z))$$
Estimating $\varphi$

- The loss function

$$L_Q(\varphi) = \sum_{x \in X} KL(Q(z, x; \varphi), P(z)) - \sum_{(x, z) \in [X, Z]} \log P(x|z; \theta)$$

- We have:

$$Q(z, x) = N(z; \mu(x; \varphi), \Sigma(x; \varphi)), \quad P(z) = N(0, I)$$

- The KL between the two Gaussians works out to

$$KL(Q(z, x; \varphi), P(z)) = \frac{1}{2} \left( tr(\Sigma(x; \varphi)) + \mu(x; \varphi)^T (\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right)$$

- We have

$$\log P(x|z; \theta) = \sum_{(x, z)} -\frac{1}{2} \log|D| -0.5(x - f(z; \theta))^T D^{-1}(x - f(z; \theta))$$

- Plugging it all in:

$$L_Q(\varphi) = \sum_{x \in X} \frac{1}{2} \left( tr(\Sigma(x; \varphi)) + \mu(x; \varphi)^T (\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right)$$

$$+ \sum_{(x, z) \in [X, Z]} \frac{1}{2} \log|D| +0.5(x - f(z; \theta))^T D^{-1}(x - f(z; \theta))$$
Estimating $\varphi$

- So we finally have the loss function (ignoring unnecessary terms and factors)

$$
L_Q(\varphi) = \sum_{x \in X} \left( tr(\Sigma(x; \varphi)) + \mu(x; \varphi)^T(\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right) + 
\sum_{(x, z) \in [X, Z]} (x - f(z; \theta))^T D^{-1}(x - f(z; \theta))
$$

- Assuming that $D$ is diagonal with identical values $\sigma^2$ for the diagonal elements gives us the simplification

$$
L_Q(\varphi) = \sum_{x \in X} \left( tr(\Sigma(x; \varphi)) + \mu(x; \varphi)^T(\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right) + \frac{1}{\sigma^2} \sum_{(x, z) \in [X, Z]} \|x - f(z; \theta)\|^2
$$

- To estimate $\varphi$ we will compute

$$
\varphi^* = \arg\min_{\varphi} L_Q(\varphi)
$$

- To perform the minimization we will use gradient descent
Estimating $\varphi$

$$\varphi^* = \arg\min_{\varphi} L_Q(\varphi)$$

- To perform the minimization we will use gradient descent
  $$\nabla_{\varphi} L_Q(\varphi)$$
  $$= \sum_{x \in X} \nabla_{\varphi} \left( \text{tr}(\Sigma(x; \varphi)) + \mu(x; \varphi)^T (\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right)$$
  $$+ \frac{1}{\sigma^2} \sum_{(x,z) \in [X,Z]} \nabla_{z}\| (x - f(z; \theta)) \|^2 \nabla_{\varphi} z$$
The complete training pipeline

- Initialize $\theta$ and $\phi$

- Iterate:
  - Sample $z_{x,e}$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
    - “Completing” the data
  - Reestimate $\theta$ from the entire “complete” data
    \[
    L(\theta, \sigma^2) = d \log \sigma^2 + \frac{1}{\sigma^2} \sum_{(x,z)} \|x - f(z; \theta)\|^2
    \]
  - Estimate $\varphi$ using the entire “complete” data
    \[
    L_Q(\varphi) = \sum_{x \in X} \left( tr(\Sigma(x; \varphi)) + \mu(x; \varphi)^T (\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right) + \frac{1}{\sigma^2} \sum_{(x,z) \in [X,Z]} \|(x - f(z; \theta))\|^2
    \]
The complete training pipeline:
Single step update

• Initialize $\theta$ and $\varphi$

• Iterate:
  – Sample $z_{x,\epsilon}$ from $N(z; \mu(x; \varphi), \Sigma(x; \varphi))$ for each training instance
    • “Completing” the data
  – Reestimate $\theta$ and $\varphi$ from the entire “complete” data

\[
L(\theta, \sigma^2, \varphi) = \sum_{x \in X} \left( tr(\Sigma(x; \varphi)) + \mu(x; \varphi)^T (\mu(x; \varphi) - d - \log|\Sigma(x; \varphi)|) \right)
+ \frac{1}{\sigma^2} \sum_{(x,z) \in [X,Z]} \|x - f(z; \theta)\|^2 + d \log \sigma^2
\]

• (Merged the updates of $\theta$ and $\varphi$ into a single step)
  – Gradient computation doesn’t change
The complete training pipeline

• Once trained the approximation function $Q(z, x)$ can be discarded.

• The rest of the function gives us a generative model for $x$.

• Generating data using this part of the model should (ideally) give us data similar to the training data.
But where are the neural nets?

• $f(z; \theta)$ is generally modelled by a neural network

• $\mu(x; \varphi)$ and $\Sigma(x; \varphi)$ are generally modelled by a common network with two outputs
  – The combined parameters of the network are $\varphi$
The Variational AutoEncoder

The decoder is the actual generative model.

The encoder is primarily needed for training. It can also be used to generate the (approximate) distribution of latent space representations conditioned on specific inputs (much like a regular autoencoder).

\( z \) is a latent-space representation of the data. \( \mu(x) \) can also be used as a expected latent representation of \( x \).
Mark all that are true

- The decoder in a Variational Auto Encoder is a non-linear Gaussian model
- The NLGM in the VAE is estimated using EM
- The encoder in a VAE is a module that generates the samples of $z$ needed to complete the data, in order to estimate the parameters of the NLGM (decoder)
- The encoder approximates $P(z|x)$ to enable sampling of $z$, to complete the data
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VAEs

• VAEs are, unfortunately, strictly generative models

• They can be used to generate samples of the data

• But they cannot be used to compute the likelihood of data
  – At least not directly
  – Because $P(x; \theta)$ is generally intractable

• Nevertheless, they are highly effective as generators
  – They can learn highly complex distributions
VAE examples

• Top: VAE trained on MNIST and used to generate new data

• Below: VAE trained on faces, and used to generate new data
**VAE and latent spaces**

- The latent space $z$ often captures underlying structure in the data $x$ in a smooth manner.

- Varying $z$ continuously in different directions can result in plausible variations in the drawn output.
  - Typically, manipulations are performed by wiggling $z$ around its expected value $\mu(x)$.

- Typically, in these draws, you do not add the noise $e$.
  - The output is the *expected* generation for a given latent value $z$. 

![Diagram of VAE and latent spaces](image-url)
VAE conclusions

• Simple non-linear extensions of linear Gaussian models

• Excellent generative models for the distribution of data $P(x)$
  – Various extensions such as Conditional VAEs, which model conditional distributions, such as $P(x|y)$
    • Straight-forward extension where the conditioning variable $y$ is an additional input to the encoder and decoder

• Have also been successfully embedded into dynamical system models
  – $P(z)$ now becomes a mixture, or a Markov model instead of $N(0, I)$

• In all cases, the arithmetic for learning is similar to that presented here
• Read the literature on the topic, it is vast