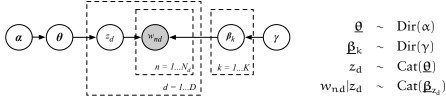
Latent Dirichlet Allocation for Topic Modeling

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Limitations of the mixture of categoricals model



A generative view of the mixture of categoricals model

- **1** Draw a distribution $\underline{\theta}$ over K topics from a Dirichlet(α).
- **2** For each topic k, draw a distribution $\underline{\beta}_k$ over words from a Dirichlet(γ).
- **3** For each document d, draw a topic z_d from a Categorical($\underline{\theta}$)
- **4** For each document d, draw N_d words w_{nd} from a Categorical($\underline{\beta_{z_d}}$)

Limitations:

- All words in each document are drawn from one specific topic distribution.
- This works if each document is exclusively about one topic, but if some documents span more than one topic, then "blurred" topics must be learnt.

NIPS dataset: LDA topics 1 to 7 out of 20.

network unit training weight input hidden output learning laver error set neural net number performance pattern problem trained generalization result

network node representation input unit learning activation nodes pattern level string structure grammar symbol recurrent system connectionist sequence order context

model data distribution probability parameter set gaussian error method likelihood prediction function mean density prior estimate estimation neural expert bayesian

problem constraint distance cluster point algorithm tangent energy clustering optimization cost graph method neural transformation matching code objective entropy set

neuron cell input model synaptic firing response activity potential current synapses membrane pattern output inhibitory effect system neural function network

network neural system model control output recurrent input signal controller forward error dvnamic problem training nonlinear prediction adaptive memory algorithm

cell model visual direction motion field eve unit cortex orientation map receptive neuron input head spatial velocity stimulus activity

cortical

NIPS dataset: LDA topics 8 to 14 out of 20.

circuit chip network neural analog output neuron current input system vlsi weight implementation voltage processor bit hardware data digital transistor

learning speech algorithm word recognition error gradient system weight training function network convergence hmm speaker vector context rate model parameter optimal set problem mlp method neural order acoustic descent phoneme equation output term input result letter noise performance solution segment

classifier classification pattern training character set vector class algorithm recognition data performance error number digit feature network neural nearest

network neuron dynamic system neural pattern phase point equation model function field attractor connection parameter oscillation fixed oscillator states activity

data memory performance genetic system set features model problem task patient human target similarity algorithm number population probability item

result

function linear vector input space matrix component dimensional point data basis output set approximation order method gaussian network algorithm

dimension

problem

NIPS dataset: LDA topics 15 to 20 out of 20.

function network bound neural threshold theorem result number size weight probability set proof net input class dimension case complexity distribution

learning action task function reinforcement algorithm control system path robot policy problem step environment optimal goal method states space sutton

model object movement motor point view position field arm trajectory learning control dvnamic hand joint surface subject data human inverse

image images system features feature recognition pixel network object visual map neural vision laver level information set segmentation task location

rules algorithm learning tree rule examples set neural prediction concept knowledge trees information query label structure model method data system

signal frequency noise spike information filter channel auditory temporal model sound rate train system processing analysis peak response correlation neuron

Latent Dirichlet Allocation (LDA)

Seeking Life's Bare (Genetic) Necessities

Haemophilus

genome 1703 genes

Genes

n common 233 genes

genome

COLD SPRING HARBOR, NEW YORK-How many genes does an organism need to survive? Last week at the genome meeting here,* two genome researchers with radically different approaches presented complementary views of the basic genes needed for life. One research team, using computer analyses to compare known genomes, concluded that today's organisms can be sustained with just 250 genes, and that the earliest life forms

required a mere 128 genes. The other researcher mapped genes in a simple parasite and estimated that for this organism. 800 genes are plenty to do the job-but that anything short of 100 wouldn't be enough.

Although the numbers don't match precisely, those predictions Arcady Mushegian, a computational molecular biologist at the National Center for Biotechnology Information (NCBI) in Bethesda, Maryland. Comparing an

Redundant and

Related and

Genes parasite-specific genes removed hahaan for biochemical -122 genes pathways +22 genes Minimal Mycoplasma genes 250 genes Ancestral

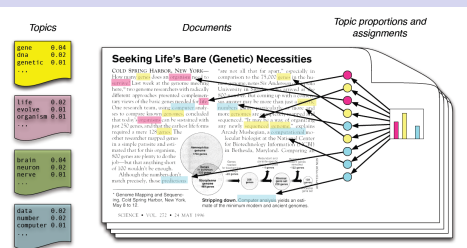
Stripping down. Computer analysis yields an estimate of the minimum modern and ancient genomes.

SCIENCE • VOL. 272 • 24 MAY 1996

[&]quot;are not all that far apart," especially in comparison to the 75,000 genes in the human genome, notes Siv Andersson of Uppsala University in Sweden, who arrived at the 800 number. But coming up with a consensus answer may be more than just a genetic numbers game, particularly as more and more genomes are completely mapped and sequenced. "It may be a way of organizing any newly sequenced genome," explains

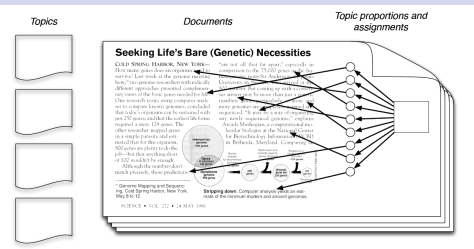
^{*} Genome Mapping and Sequencing, Cold Spring Harbor, New York, May 8 to 12.

Generative model for LDA



- Each *topic* is a distribution over words.
- Each *document* is a mixture of corpus-wide topics.
- Each *word* is drawn from one of those topics.

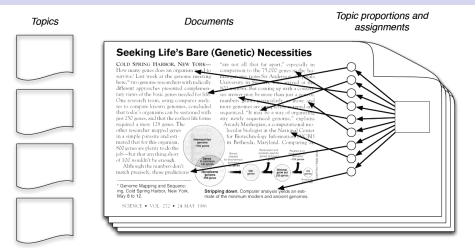
The posterior distribution



- In reality, we only observe the documents.
- The other structure are *hidden* variables.
 i.e. the "topics" (left) & the "topic proportions and assignments" (right).

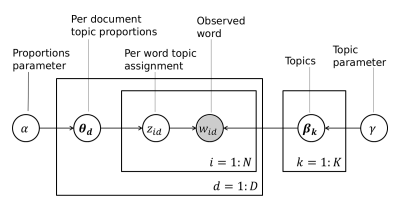
8/18

The posterior distribution



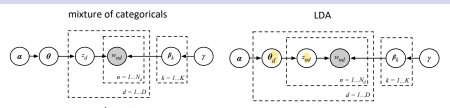
- Our goal is to *infer* the hidden variables.
- This means computing their distribution conditioned on the documents p(topics, proportions, assignments|documents)

The LDA graphical model



- Nodes are random variables; edges indicate dependence.
- Shaded nodes indicate observed variables.

The difference between mixture of categoricals and LDA



A generative view of LDA

- **1** For each document d draw a distribution θ_d over topics from a Dirichlet(α).
 - **2** For each topic k draw a distribution β_k over words from a Dirichlet(γ).
 - **3** Draw a topic z_{nd} for the n-th word in document d from a Categorical(θ_d)
- **4** Draw word w_{nd} from a Categorical($\beta_{z_{nd}}$)

Differences with the mixture of categoricals model:

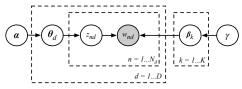
• In LDA, every word in a document can be drawn from a different topic,



• and every document has its own distribution over topics θ_d .



The LDA inference problem



"Always write down the probability of everything." (Steve Gull)

$$\begin{split} p(\underline{\beta}_{1:K}, \underline{\theta}_{1:D}, \{z_{nd}\}, \{w_{nd}\}|_{\pmb{\gamma}, \pmb{\alpha}}) & & \text{given the topic proportion parameter `alpha` \&} \\ & & \text{vocab "topic" parameter `gamma`.} \\ & = \prod_{k=1}^K p(\beta_k|\gamma) & \prod_{d=1}^D \left[p(\theta_d|\alpha) \prod_{n=1}^{N_d} \left[p(z_{nd}|\theta_d) p(w_{nd}|\beta_{1:K}, z_{nd}) \right] \right] \end{split}$$

Learning involves computing the posterior over the parameters, $\beta_{1:K}$ and $\theta_{1:D}$ given the words $\{w_{nd}\}$, but this requires the we marginalize out the latent $\{z_{nd}\}$.

How many configurations are there?

This computation is *intractable*.

The intractability of LDA

The evidence (normalising constant of the posterior):

$$p(\{w_{id}\}) = \iint \sum_{z_{id}} \prod_{d=1}^{D} \prod_{k=1}^{K} \prod_{n=1}^{N_d} p(z_{nd}|\theta_d) p(\theta_d|\alpha) p(w_{nd}|\beta_{1:K}, z_{nd}) p(\beta_k|\gamma) d\beta_k d\theta_d$$

We need to average over all possible set of values of all z_{nd} . If every document had N words, this means K^N configurations per document.

Gibbs to the Rescue

The posterior is *intractable* because there are too many possible latent $\{z_{nd}\}$.

Sigh, ... if only we knew the $\{z_{nd}\}$...?

Which might remind us of Gibbs sampling ... could we sample each latent variable given the values of the other ones?

Refresher on Beta and Dirichlet

If we had a $p(\pi) = \text{Beta}(\alpha, \beta)$ prior on a binomial probability π , and observed k successes and n - k failures, then the posterior probability

$$p(\pi|n,k) = Beta(\alpha + k, \beta + n - k),$$

and the predictive probability of success at the next experiment

$$p(success|n,k) = \int_{\text{This term is just pi.}} p(success|\pi) p(\pi|n,k) d\pi = \mathbb{E}[\pi|n,k] \ = \ \frac{\alpha+k}{\alpha+\beta+n}.$$

Analogously, if we had a prior $p(\pi) = Dir(\alpha_1, ..., \alpha_k)$ on the parameter π of a multinomial, and $c_1, ..., c_k$ observed counts of each value, the posterior is

$$p(\boldsymbol{\pi}|c_1,\ldots,c_k) = \mathrm{Dir}(\alpha_1+c_1,\ldots,\alpha_k+c_k),$$

and the predictive probability that the next item takes value j is:

$$p(j|c_1,\ldots,c_k) = \int_{\substack{p|j|j}} p(j|\pi)p(\pi|c_1,\ldots,c_k) d\pi = \mathbb{E}[\pi|c_1,\ldots,c_k] \ = \ \frac{\alpha_j+c_j}{\sum_i \alpha_i+c_i}.$$

Collapsed Gibbs sampler for LDA

In the LDA model, we can integrate out the parameters of the multinomial distributions, θ_d and β , and just keep the latent counts z_{nd} . Sampling these z_{nd} in turn is called a *collapsed* Gibbs sampler.

Recall, that the predictive distribution for a symmetric Dirichlet is given by

$$p_i = \frac{\alpha + c_i}{\sum_j \alpha + c_j}.$$

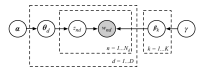
Now, for Gibbs sampling, we need the predictive distribution for a single z_{nd} given all other z_{nd} , ie, given all the counts except for the word n in document d. The Gibbs update contains two parts, one from the topic distribution and one from the word distribution: (check out comments for underlines.)

$$p(z_{\mathrm{nd}} = \mathrm{k} | \{z_{-\mathrm{nd}}\}, \{w\}, \gamma, \alpha) \propto \frac{\alpha + c_{-\mathrm{nd}}^{\mathrm{k}}}{\displaystyle\sum_{\mathrm{j=1}}^{\mathrm{K}} (\alpha + c_{-\mathrm{nd}}^{\mathrm{j}})} \frac{\gamma + \tilde{c}_{-w_{\mathrm{nd}}}^{\mathrm{k}}}{\displaystyle\sum_{\mathrm{m=1}}^{\mathrm{M}} (\gamma + \tilde{c}_{-\mathrm{m}}^{\mathrm{k}})}$$

k here is not a power.

where $c_{-n,d}^{k}$ is the count of words from document d, excluding n, assigned to topic k, and \tilde{c}_{-m}^k is the number of times word m was generated from topic k (again, excluding the observation nd).

Derivation of the collapsed Gibbs sampler



The probability of everything: from pg 12/18.

$$p(\boldsymbol{\beta}_{1:K},\boldsymbol{\theta}_{1:D},\!\{\boldsymbol{z}_{nd}\},\!\{\boldsymbol{w}_{nd}\}\!|\boldsymbol{\gamma},\boldsymbol{\alpha}) = \prod_{k=1}^{K} p(\boldsymbol{\beta}_{k}|\boldsymbol{\gamma}) \ \prod_{d=1}^{D} \left[p(\boldsymbol{\theta}_{d}|\boldsymbol{\alpha}) \prod_{n=1}^{N_{d}} \left[p(\boldsymbol{z}_{nd}|\boldsymbol{\theta}_{d}) p(\boldsymbol{w}_{nd}|\boldsymbol{\beta}_{1:K},\boldsymbol{z}_{nd}) \right] \right] \left[p(\boldsymbol{\beta}_{1:K},\boldsymbol{\theta}_{1:D},\!\{\boldsymbol{z}_{nd}\},\!\{\boldsymbol{\omega}_{nd}\},\!\{$$

What we want for Gibbs sampling is:

$$\begin{split} p(z_{nd} = k | \{z_{-nd}\}, & \{w\}, \gamma, \alpha \} \quad \text{prior} \quad \text{likelihood of z_nd} \\ & \propto p(z_{nd} = k | \{z_{-nd}\}, \alpha) \; p(w_{nd} | z_{nd} = k, \{w_{-nd}\}, \{z_{-nd}\}, \gamma) \\ & = \frac{\alpha + c_{-nd}^k}{\sum\limits_{j=1}^K (\alpha + c_{-nd}^j)} \; \frac{\gamma + \tilde{c}_{-w_{nd}}^k}{\sum\limits_{m=1}^M (\gamma + \tilde{c}_{-m}^k)} \end{split}$$

where $c_{-nd}^j \stackrel{\text{def}}{=} \sum_{n' \neq n} \mathbb{I}(z_{n'd} = j)$ and $\tilde{c}_{-m}^k \stackrel{\text{def}}{=} \sum_{(n',d') \neq (n,d)} \mathbb{I}(w_{n'd'} = m) \mathbb{I}(z_{n'd'} = k)$.

Per word Perplexity

In text modeling, performance is often given in terms of per word *perplexity*. The perplexity for a document is given by

$$\exp(-\ell/n)$$
,

where ℓ is the log joint probability over the words in the document, and n is the number of words. Note, that the average is done in the log space.

A perplexity of g corresponds to the uncertainty associated with a die with g sides, which generates each new word. Example:

$$p(w_1, w_2, w_3, w_4) = \frac{1}{6} \frac{1}{6} \frac{1}{6} \frac{1}{6}$$
 (1)

$$\frac{1}{n}\log p(w_1, \dots, w_4) = \frac{1}{4}\log(\frac{1}{6})^4 = -\log 6$$
 (2)

$$perplexity = \exp(-\frac{1}{n}\log p(w_1, \dots, w_4)) = 6$$
 (3)