SURVIVAL-SUPERVISED LATENT DIRICHLET ALLOCATION MODELS FOR GENOMIC ANALYSIS OF TIME-TO-EVENT OUTCOMES

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Two challenging problems in the clinical study of cancer are the characterization of cancer subtypes and the classification of individual patients according to those subtypes. Statistical approaches addressing these problems are hampered by population heterogeneity and challenges inherent in data integration across high-dimensional, diverse covariates. We have developed a survival-supervised latent Dirichlet allocation (survLDA) modeling framework to address these concerns. LDA models have proven extremely effective at identifying themes common across large collections of text, but applications to genomics have been limited. Our framework extends LDA to the genome by considering each patient as a “document” with “text” constructed from clinical and high-dimensional genomic measurements. We then further extend the framework to allow for supervision by a time-to-event response. The model enables the efficient identification of collections of clinical and genomic features that co-occur within patient subgroups, and then characterizes each patient by those features. An application of survLDA to The Cancer Genome Atlas (TCGA) ovarian project identifies informative patient subgroups that are characterized by different propensities for exhibiting abnormal mRNA expression and methylations, corresponding to differential rates of survival from primary therapy.

1. Introduction. Technological advances continue to increase both the ease and accuracy with which measurements of the genome and phenome can be obtained and, consequently, genomic-based studies of disease often involve highly diverse types of data collected on large groups of patients. The primary goals of such studies involve identifying genomic features useful for characterizing patient subgroups as well as predicting patient-specific disease course and/or likelihood of response to treatment. Doing so requires statistical methods that handle complex interactions, accommodate population heterogeneity, and allow for data integration across multiple sources.

A number of statistical methods are available for survival-related feature identification and prediction (for a review, see Witten and Tibshirani (2009); Li and Li (2004); Wei and Li (2007)). Most often, classical models for a survival response are coupled with some dimension-reduction method for individual (Li and Luan, 2003; Ghosh and Yuan, 2010; Pang, Datta and Zhao, 2010) or grouped predictors (Chen and Wang, 2009; Li and Li, 2004; Ma, Song and Huang, 2007; Chen, Wang and Ishwaran, 2010), providing a concise representation of the genomic features affecting patient outcome. Although useful, the majority of these methods identify a set of covariates common to all patients and as a result may “distort what is observed” in the presence of population heterogeneity (Aalen, 1988). Survival-supervised clustering approaches naturally accommodate heterogeneity, providing for efficient and effective identification of patient subgroups (Dettling and Buhlmann, 2002; Li and Gui, 2004). However, these approaches do not identify salient features associated with subgroups; and, as with the aforementioned methods, may sacrifice power and accuracy by focusing on one (or a few) data sets in isolation.

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Latent Dirichlet allocation (LDA, Blei, Ng and Jordan (2003)) models are particularly well-tailored for accommodating heterogeneity, selecting features, and characterizing complex interactions in a high-dimensional setting. By far the most common application concerns identifying groups of words that co-occur frequently (topics) across large collections of text (e.g., a collection of abstracts, emails, or manuscripts). The derived topics provide insight into the collections’ content overall as well as into the specific content within a document; and estimated document-specific distributions over topics are useful in classifying new documents (Blei, Ng and Jordan (2003); Porteous et al. (2008); Biro, Szabo and Beniczur (2008)).

A recent extension allows for topic estimation to be supervised by a response that is suitably described by a generalized linear model (Blei and McAuliffe, 2008). So-called supervised LDA (sLDA) debuted with a study of movie reviews (text) and estimated topics (collections of co-occurring words in a review) that determined the number of stars (supervising response) a movie received. Derived topics included ones having highest weight on words such as ‘power’, ‘perfect’, ‘fascinating’ and ‘complex’; another with highest weight on ‘routine’, ‘awful’, ‘featuring’, ‘dry’; a third on ‘unfortunately’, ‘least’, ‘flat’, ‘dull’; and so on. The movie-review-specific distribution over topics proved useful in classifying movies. Those with highest weight on the ‘power’ topic generally had a high number of stars while those with highest weight on the ‘unfortunately’ topic had a low number; those with weight on the ‘routine’ topic most often ended up in the middle. Differences between the distributions also provided insights into differences between movies that received a similar number of stars, as well as insights into the connotative nature of word choice (e.g., pictures described as ‘films’ rating higher than those referred to as ‘movies’).

Our interest here is not in evaluating movies. However, it is important to note that the questions addressed in Blei and McAuliffe (2008) are identical in structure to the most important questions we face in cancer genomics. In the former, questions include: ‘Given reviews and ratings for a group of movies, can we identify topics - collections of words that co-occur frequently in some reviews and less frequently in others ? Can each individual movie be described by a distribution over those topics ? Can distributions over topics provide insights into differences between similarly rated movies ? And can a movie-specific distribution over topics be used to predict what the rating of a new movie will be ?’ In cancer genomics, the questions include: ‘Given genomic, clinical, and survival information on a group of patients, can we identify topics that are collections of genomic and clinical features that co-occur frequently in some patients and less frequently in others ? Can a patient be well described by a distribution over those topics ? Can distributions over topics provide insights into the genomic differences between two patients with similar survival ? And can a patient-specific distribution over topics be used to predict survival of an individual patient ?’

To address these types of questions, we extend LDA for use in a clinical and genomic setting. Unlike in the textual domains of Blei, Ng and Jordan (2003), Porteous et al. (2008) and Biro, Szabo and Beniczur (2008), the definition of a document here is not obvious. Section 2.4 details the construction of documents, one for each patient, where words describe clinical events, treatment protocols, and genomic information from multiple sources. As we show in Section 3, application of traditional LDA to this collection of documents provides for the identification of topics useful in characterizing patient subpopulations as well as individual patients in a study of ovarian cancer conducted as part of The Cancer Genome Atlas (TCGA) project (National Cancer Institute and National Human Genome Research Institute, 2011). Survival supervised LDA (survLDA) is developed in Section 2.2 to facilitate topic supervision by a time-to-event response, which further improves patient-specific characterization and prediction.

2. Methods.

2.1. The LDA model. We briefly review the LDA model as detailed in Blei et al. 2003. Assume there are $D$ documents indexed by $i = 1, \ldots, D$, each of which consists of $N_i$ words. The vocabulary
is the unique set of length \( V \) indexed by \( v = 1, \ldots, V \), from which the documents’ words arise, and is usually taken to be the union of all words over documents. Further assume that there are \( K \) latent ‘topics’ indexed by \( k = 1, \ldots, K \), that govern the assignment of words to documents. Each topic corresponds to a discrete distribution over the \( V \) words in the vocabulary, with parameters given by the \( V \)-vector \( \tau_k \). Likewise, each document is assumed to be associated with mixing coefficients \( \theta_i \) over the \( K \) topics, indicating its partiality with respect to word sources.

For a given document \( i \), \( N_i \) words arise from the following generative process given the system-wide hyperparameters \( \alpha \) (a \( K \)-vector Dirichlet parameter) and the \( \tau_{1:K} \) (the topic \( V \)-vectors):

1. Draw topic proportions \( \theta_i \sim \text{Dirichlet}(\alpha) \)
2. For each of the \( N_i \) words, indexed by \( j \):
   
   (a) Draw a topic assignment \( Z_{ij} | \theta_i \sim \text{Multinomial}(1, \theta_i) \) (Note \( Z_{ij} \in \{1, \ldots, K\} \))
   
   (b) Draw a word \( W_{ij} | Z_{ij}, \tau_{1:K} \sim \text{Multinomial}(1, \tau_{Z_{ij}}) \) (Note \( W_{ij} \in \{1, \ldots, V\} \))

With this model in place, a variational expectation-maximization (EM) algorithm may be used to estimate the joint posterior distribution of \( \theta \) and \( \tau \) given \( w_{i,1:N_i} \), \( \alpha \) and \( \tau_{1:K} \) for each document \( i \) (E-step) and then to estimate the system-wide hyperparameters \( \alpha \) and \( \tau_{1:K} \) (M-step). Upon convergence, the variational EM yields optimal values for the key quantities of interest, namely posterior estimates for \( \theta_{i,D} \) and \( \tau_{1:K} \).

We note that the topics engendered by unsupervised LDA are, well, unsupervised. If there is a relationship between certain words and an outcome of interest, the model may be hard pressed to find it as the algorithm does not have access to the outcome. While one may certainly be interested in the relationship between certain words and an outcome of interest, the model may be hard pressed to find them.

2.2. The survival supervised LDA model. Assume the same setup as in Section 2.1 with \( D \) documents indexed by \( i \), a vocabulary of size \( V \), and \( K \) topics with corresponding discrete distributions \( \tau_{1:K} \). Additionally, the introduction of supervision through a time-to-event outcome means that, just as a document’s partiality to certain topics through \( \theta_i \) impacts its constituent words, those topics affect the survival outcome \( T_i \). An indicator variable for death/censoring is also observed for each document and denoted by \( \delta_i \).

The system-wide model parameters for the survLDA model include a \( K \)-vector Dirichlet parameter \( \alpha \), the topic \( V \)-vectors \( \tau_{1:K} \), and survival response parameters \( \beta \) (a \( K \)-vector of regression coefficients) and \( h_0(\cdot) \) (baseline hazard). When they are available for a given document \( i \), \( N_i \) words and a survival response \( T_i \) arise from the following generative process:

1. Draw topic proportions \( \theta_i \sim \text{Dirichlet}(\alpha) \)
2. For each of the \( N_i \) words, indexed by \( j \)
   
   (a) Draw a topic assignment \( Z_{ij} | \theta_i \sim \text{Multinomial}(1, \theta_i) \) (Note \( Z_{ij} \in \{1, \ldots, K\} \))
   
   (b) Draw a word \( W_{ij} | Z_{ij}, \tau_{1:K} \sim \text{Multinomial}(1, \tau_{Z_{ij}}) \) (Note \( W_{ij} \in \{1, \ldots, V\} \))
3. Compute the \( K \)-vector \( \bar{Z}_i \) s.t. \( \bar{Z}_{ik} = \#\{Z_{ij} = k\}/N_i \)
4. Draw a survival response \( T_i | \bar{Z}_i, \beta, h_0 \) from the survival function corresponding to a Cox proportional hazards model with hazard function \( h(t|\bar{Z}_i) = h_0(t) \exp\{\beta'\bar{Z}_i\} \)

Note that we are using a Cox proportional hazards model (Cox, 1972), with each regression coefficient \( \beta_k \) exhibiting the beneficent (negative) or deleterious (positive) effect of topic \( k \) on survival. The form
of \( h_0 \) may be chosen by the user and may be parametric (such as using a Weibull survival model) or nonparametric.

As in LDA, a variational EM algorithm may be used to estimate the joint posterior distribution of \( \theta_i \) and \( Z_{i:1:N_i} \) given \( w_{i:1:N_i}, T_i, \delta_i, \alpha, \tau_{1:K}, \beta \) and \( h_0 \) for each document \( i \) (E-step) and then to estimate the system-wide hyperparameters \( \alpha, \tau_{1:K}, \beta \) and \( h_0 \) (M-step). The derivation of this variational EM is given in Appendix A. As detailed in Appendix B, we introduce into the variational EM an ‘uninteresting’ background topic to act as a benchmark with respect to the supervising outcome. Without loss of generality, say this is the last (\( K^{th} \)) topic. Then the distribution over the vocabulary for the background topic (\( \tau_K \)) may be identified by placing count weights on the known ‘background’ words, a tiny amount of weight on all other words (say, 0.001) and re-normalizing so that the sum over the entire \( V \)-vector is one. Non-background topics are then different deviations from this benchmark that express themselves through differential survival. In our application, the background topic would describe ‘featureless’ documents that contain only the ubiquitous adjuvant therapy information, nothing more. Upon convergence, the variational EM yields posterior estimates for the key quantities of interest: posterior estimates for the \( \theta_{1:D} \) as well as for the composition (\( \tau_{1:K} \)) and outcome effect (\( \beta \)) of the \( K \) topics.

2.3. Prediction. Given a new patient with document \( w_{1:N} \) and a fitted model \( \{ \alpha, \tau_{1:K} \} \), the posterior mean \( \bar{Z}_{\text{new}} = \bar{Z}|w_{1:N}, \alpha, \tau_{1:K} \) can be obtained in order to estimate from what topics this new patient draws words, and in what proportions. As was the case during model fitting, this posterior must be approximated via variational inference. We do so by following the same procedure as outlined in the first subsection of Appendix A, except that all survival-related terms in the evidence lower bound are dropped; see the third subsection of Appendix A for details. We note that this approach is analogous to that in Blei and McAuliffe (2008), where they point out that the prediction protocol does not depend on the particular response type.

Given \( \bar{Z}_{\text{new}} \), measures related to topic membership can be predicted for the new patient. This may be done qualitatively (e.g., “This patient is predicted to belong strongly to the second topic and survival for that topic is poor, hence her prognosis is bad.”) or quantitatively (e.g., predicting median survival time using the parametric survival model; see Appendix A).

2.4. Document construction in the TCGA cohort. In most applications of LDA, the definition of ‘document’ is obvious. That is not the case here. In studies of cancer genomics, data are available in the form of text (e.g., treatments received, a clinician’s evaluation of response to treatment, etc.); there are also often disparate non-textual measurements (e.g., binary, count, continuous, factor). By translating these measurements into words, and thus patients into textual documents, the LDA and survLDA models may be used for inference.

Our population of interest is the ovarian cancer cohort from The Cancer Genome Atlas (TCGA) (National Cancer Institute and National Human Genome Research Institute, 2011). For these women, clinical information such as time of surgery, adjuvant therapies, time of recurrence, treatment at recurrence, overall survival, and dozens of other variables are available. Also available are high-throughput measurements of gene expression, methylation, SNP/CNVs, and microRNAs. For document construction, we used words associated with adjuvant therapy, expression as assayed by the Affymetrix HT Human Genome U133A chip, and methylation as assayed using the Illumina Infinium HumanMethylation27 BeadChip. Out of the cohort as accessed in April of 2011, 448 of the women were not missing more than half of their gene expression or methylation measurements and received at least the standard therapy of a platinum and taxane. A document was constructed for each of the 448 patients.

For a given patient, a drug-related word was added to her document for every drug given to her during adjuvant treatment. For example, a patient receiving two platinums and a taxane would have the words ‘platinum’, ‘platinum’ and ‘taxane’ added to her document. Related drugs were collected together (e.g., the many varied platinums) and adjuvant drug therapies not given to at least 10% of the women were
To help ensure that documents contained words corresponding to meaningful genes, we considered those genes for which expression or methylation is multi-modal, as assessed by MCLUST (Fraley and Raftery, 2002, 2006). Specifically, a gene’s expression distribution was deemed multi-modal if MCLUST preferred a two-component over a one-component model when given only those choices. Filtering by known genes, non-missingness, and multi-modality reduced the number of gene expression measurements considered from 14,500 to 7,727. Given the multi-modal genes, a patient’s document received a gene word, given by the gene name, if that patient showed extreme expression for that gene. The same word was added again if the patient showed extreme methylation. As most genes have multiple sites at which methylation was measured, we considered each methylation site in each gene separately. So a patient’s document receives copies of the word “geneX” if that patient had extreme expression for geneX (one copy added) as well as multiple hyper (or hypo) methylated sites (one copy added per hyper- or hypo-methylated site). For both types of data, extreme was defined as being in the minor mode of the genes’ multi-modal distribution, if the probability of being so, as estimated by MCLUST, exceeded 95%. As extreme may manifest as high or low expression (or methylation), one cannot infer direction of the gene expression or methylation measurement from word frequency; i.e., high frequency does not necessarily correspond to high expression. This is a consequence of the way in which documents were constructed and could be changed.

In an effort to improve power to detect survival-related topics, a second filter was applied. Specifically, we considered the 234 uncensored patients and partitioned them into three groups of 78 based on survival times. A gene’s word was removed if word frequency did not vary across groups. For gene expression (methylation), we required a difference of at least 10 (15) words between at least two of the survival groupings, leaving 201 (1,063) vocabulary words. Note that the censored women still received words based on gene expression and methylations; they merely did not contribute to the decisions made during this pre-filtering. A typical document contained approximately 350 words (mean 347; median = 132), on the order of a PubMed abstract.

3. Application to TCGA data. Given documents constructed as described above for each of the 448 women considered, we applied LDA as well as survLDA. The outcome of interest is all-cause mortality and for the survLDA application, we used a Weibull model for the baseline hazard. The background topic (the seventh topic) was assigned non-trivial weight only on the adjuvant therapy words platinum and taxane, as this setup would constitute an ‘uninteresting’ document since all patients received these two treatments (they were inclusion criteria in the TCGA ovarian project). In all analyses, we use \( K = 7 \) topics, the last being the background topic.

3.1. Results. Application of (unsupervised) LDA provides two quantities of primary interest. The first are the topics \( \tau_{1,K} \), or estimated distributions over words; and the second are the document-specific distributions over topics \( \theta_{1,D} \). The left panel of Figure 1 presents the topic-specific distributions over words for each topic. Red (blue) indicates an overabundance (dearth) of a word’s weight in the corpus belonging to a particular topic; white indicates an amount equivalent to an even spread over all non-background topics. The overabundance of some words and paucity of others characterize the topics, allowing one to differentiate among them. For example, consider topics 5 and 6. The right panel of Figure 1 shows twenty words having high weight in topic 5 (upper), and twenty with high weight in topic 6 (lower). It is clear that patients described primarily by these topics differ with respect to aberrations for this collection of genes. Of interest is determining whether this difference translates to a difference in overall survival.

The top left panel of Figure 2 shows a heat map of estimated topic membership for the six non-background topics (rows), clustered over patients (columns). As expected, none of the patients exhibited more than minimal weight in the background topic (not shown). Patient membership within a topic ranges from 0 (nil weight in the topic, deep blue) to 1 (wholly belonging to the topic, red). As shown, most patients have high weight in a single topic, while a few are best described by mixtures over topics.
Topics 5 and 6 are the largest topics, in the sense that they contain the most weight over the cohort. The top right panel of Figure 2 shows Kaplan-Meier (KM) curves for each topic shown in the left panel. The curves are generated by weighting the TCGA patients’ survival information by their topic membership; hence a patient whose $\theta_i$ had 50% weight in topic one and 50% weight in topic two would count as ‘half a person’ in the KM estimation of those curves, but would not contribute to the survival curves associated with other topics. As shown, there is some, albeit limited, separation with respect to survival over topics. Topics 5 and 6, for example, have rather similar survival (80% vs 70% at two years), in spite of the genomic differences highlighted in Figure 1. This type of finding is consistent with recent studies showing remarkable genetic heterogeneity among cancer patients that appear to be clinically similar (Jones et al., 2008).

The bottom left and right panels of Figure 2 similarly show estimated topic membership and KM survival curves for the survLDA analysis. While the analysis uses a parametric Weibull survival model, empirical (weighted) KM curves are presented to facilitate comparison to the (unsupervised) LDA approach. In survLDA, the largest topics are 1 and 4; and better separation among all topics is observed. Two topics have poorer than baseline survival (topics 1 and 2), and two have better (topics 5 and 6).

Figure 3 is similar to Figure 1, but shows results from survLDA. For comparison, the word order shown in Figure 1 is preserved in Figure 3. Topics have also been reordered in Figure 3 to stress similarity between topics 5 and 6 derived from the LDA analysis and topics 4 and 1 derived from survLDA. As shown in Figure 3, their topic-specific distributions over words are very similar. This is not an artifact. Indeed, many of the topics’ high-weight words are consistent between LDA and survLDA. The consistency as we shift from free-formed topics to survival supervision suggests that there are subpopulations within the cohort whose constituent topics have strong enough effects on survival that they are evident even without survival supervision. When survival information is available and used to supervise topic creation, the subsequent alterations to topics result in a much sharper disparity in survival rates. Recall that two year survival is about 80% vs. 70% for topics 5 and 6 in the LDA analysis; that spread increases to 92% vs. 65% under survival supervision for topics 4 and 1 (which are most similar in structure to topics 5 and 6). We note that this magnitude of differential survival is on par with results of other ovarian cancer studies (Tothill et al., 2008; The Cancer Genome Atlas Research Network, 2011).

With respect to words that distinguish between topics 4 and 1, we note that the following words consistently show high frequency in the poor survival group: NDC80 (a spindle checkpoint regulator associated with breast cancer (Bieche et al., 2011)) RXRA (a transcriptional regulator associated with breast cancer (Lawrence et al., 1998; Ditsch et al., 2012)), MANF (a gene coding a highly conserved protein with unknown function with mutations often observed in lung, breast, prostate (Shridhar et al., 1996) and pancreatic cancers (Shridhar et al., 1997)), and INTS6 (a tumor suppressor known to be involved in prostate cancer (Filleur et al., 2009)). Further note that NDC80, RXRA and INTS6 would not likely have been identified in this cohort by another approach, as the marginal p-values from a Cox proportional hazards test are far from overwhelming (NDC80 p=0.6, 0.98, 0.07; RXRA p=0.22, 0.37, 0.27; INTS6 p=0.38, 0.8, 0.59; all p-values are for gene expression and then for two methylations measured in the gene). MANF (p=0.02, 0.46, 0.08) is the only one of these that shows even a nominally significant marginal effect on survival. Further investigation of these and other genes that display markedly different abundance patterns between these patient subtypes might elucidate the mechanisms that underlie differences between the groups. To this end, word cloud representations such as those shown in Figure 4 may prove useful.

Additionally, the patient-specific distributions over topics are useful in that they characterize the genomic aberrations underlying individual patients. Figure 5 shows 20 high-weight words from topic 4 and 20 from topic 5 (rows). Fifteen patients are shown (columns): five identified as strongly belonging to topic four ($\theta_{i,4} > 0.99$, left column), five identified as strongly belonging to topic five ($\theta_{i,5} > 0.99$, right column).
right column), and five whom the model classifies as a mixture of both ($\theta_{i,5} > 0.4$ and $\theta_{i,5} > 0.4$, middle column). (Note that topics 4 and 5 were chosen since most patients are described almost exclusively by a single topic as shown in Figure 2; however, there are some best described by a mixture. In the survLDA analysis, most of these latter patients are a mixture between topics 4 and 5.)

The differences in genomic aberrations shown among the groups in Figure 5 are clear. Patients defined primarily by topic 4 have many of the high-weight topic 4 words in their document. The same is true for topic 5, while those who are a mixture have some realizations from both sets of words. The right side of the plot shows p-values from Cox proportional hazards tests conducted on the entire cohort. For each gene, a test was conducted for expression as well as methylation measurements associated with that gene. The p-value reported is the minimum among those tests. As shown, few of these genes would be identified as significant using a standard Cox based test since the gene expression measurements are not significantly associated with survival across the entire cohort, even though they show clear differences for some subsets of patients.

To evaluate the utility of survLDA for patient-specific prediction, we split the TCGA cohort into a training and test set (75% and 25% of the cohort, respectively). The full barcoding, document creation and survLDA model fitting procedures were applied to the training set. Documents for the test set women were derived using the abnormality indications for the genes and methylations surviving filtering that arose from the training set document creation. These documents in hand, the survLDA output was used to predict topic membership for the test set, using the prediction approach given in Section 2.3.

There were 23 women in the test set with more than 80% weight in topic 1. The patient-specific distributions over topics for the other women in the test set were not concentrated on a different topic, but had weight rather evenly spread out across topics 2-6. The pink (red) line in Figure 6 shows the Kaplan-Meier curve for the test (training) set women predicted as having more than 80% weight in topic 1, while the light blue line is the survival curve for test set women predicted as having less than 20% weight in topic 1. As shown in Figure 6, survival of women largely described by topic 1 in the training and test sets is similar, while survival for those test cases predicted as not being well described by topic 1 is considerably different, indicating some predictive power for patients strongly described by a single topic. Simulations (not shown) indicate that topic specificity as well as prediction improves with either more or larger documents; and work toward these ends is ongoing.

4. Discussion. A problem pervasive in genomic based studies of disease concerns taking large, diverse data sets collected on a cohort of patients and using the information contained therein to characterize patient subtypes as well as individuals. Computational scientists often address this problem by performing analysis within a single data type and comparing results subsequently in an effort to identify a signal supported by the disparate analyses (e.g., a gene’s SNPs, expression, and methylation all associate with a phenotype). Comparing results manually has its obvious disadvantages. At the same time, meta-analysis approaches such as Fisher’s combined probability test can be limited by low power (Zaykin et al., 2002); and efforts to combine data directly are challenged by measurements on different scales with differential dependencies. The LDA based framework proposed here addresses these challenges by transforming the information contained in high-throughput genomic screens into text. Doing so has both advantages and disadvantages.

One advantage is that data integration is seamless. In the implementation presented, a word for a gene is assigned to a patient’s document if the gene shows extreme expression; the same word is assigned if the gene shows extreme methylation. In this way, a document may contain copies of words associated with extreme genomic features, measured from expression and/or methylation. The number of copies is proportional to the number of measurements for which the gene is extreme: a gene with extreme expression and methylation will have more copies of that word than a gene showing extreme expression alone. Although we have demonstrated the approach using expression and methylation measurements (and treatment information), applications that use additional types of data are easily incorporated
into the framework.

A second advantage is that the threshold required for a gene to be included in the analysis is much lower than would be required with other methods. As detailed in Section 2.4, some pre-selection of genes is done, but the selection does not require even nominally significant association with a survival end-point, as is often required in survival studies with high-dimensional covariates (Li and Luan, 2003; Chen and Wang, 2009; Liu et al., 2009). As shown in Figure 5, this allows for the identification of many important genes, some previously known to be involved in cancer, that would not otherwise have been considered.

Application of LDA to patient documents reveals groups of genomic aberrations that co-occur together (topics) and then characterizes individual patients by those groups. The topics themselves are useful in that they define collections of genes, methylations or other covariates among which undiscovered interactions might occur, while the patient-specific distributions over topics give insights into the similarities and differences among patients that go beyond the information that can be gained from grouping by like outcome. In our application to the TCGA ovarian project, there was some consistency between the topics derived under unsupervised and supervised analyses, suggesting that the approach may produce topics whose constituents have similar survival profiles even in settings where survival information is not available. Of course, when survival is available, survLDA may be used to improve inference.

Our analysis of the TCGA ovarian cohort identified several genes whose products and methylations bear further interrogation, some already known to be involved in cancer. Our investigations of the predictive ability of the approach are mixed. While there is some ability for prediction, improvements are expected with increases in the number of patients as well as improvements in document creation strategies. For instance, in the testing set we could evaluate predictive inference in one group, the only one for which the test women had strong indications of topic membership or exclusion. Simulations (not shown) suggest that this limitation is largely due to the relatively small sample size in the test set and the minimal amount of word replication in our documents. More work is required to ascertain the power associated with sample size, document size, word frequencies, and replication, which in turn requires further experimentation with the method of document construction.

As this framework relies heavily on the words contained in a patient’s document, much work is required to develop and evaluate methods of document construction. Our approach to assign a word for any gene showing extreme expression or methylation was motivated by Ziliox and Irizarry (2007), where the authors identify bi-modal genes and, for each individual and each gene, assign a binary variable indicative of mode membership. The resulting gene expression ‘barcode’ for each patient proved extremely useful in classifying patients into biologically meaningful groups in Ziliox and Irizarry (2007); and, as demonstrated, the extrapolation of their approach proved to be an effective strategy here. At the same time, one could imagine assigning a word associated with a pathway if any gene in that pathway was extreme as assessed by expression, methylation, SNP profile, etc. This would increase the frequency with which words appeared; and our preliminary evaluation suggests that this can be useful, but can also result in topics that are not clearly distinct with respect to high-weight words. Another possibility is to assign an increasing number of words in direct proportion with signal. For example, consider breaking a gene’s expression into deciles, say, and assign one to ten words for each document (e.g., a value between the sixth and seventh deciles gets seven words). We did not favor this approach for two main reasons. First, the approach assumes linearity of expression and methylation which is often not the case. Second, the approach results in documents having few unique words, which reduces specificity of topics as well as document specific distributions over topics. Document construction continues to be explored, and improvements are expected to prove useful in a number of settings.

In addition to the means by which covariates are translated into words, there are many aspects of the proposed methods that require further development. In particular, survLDA assumes the simplest of
Dirichlet priors on the distributions of topics over patients and therefore the documents are considered conditionally independent given $\alpha$. While this is a reasonable assumption for the TCGA data set we considered, there are other realms where correlation among the documents could arise. For example, one could have multiple documents arising from the same subject, one for each time point or tissue; or, when integrating multiple cancer types, subjects with the same type of cancer would be expected to be more alike than subjects with differing cancer types. Adding such hierarchy has already been explored for traditional LDA (Teh et al., 2006), presenting a starting point for methodological extension.

Similarly, the composition of the topics themselves is essentially free. Were it not for our imposition of a background topic, the topics would be completely unstructured a priori. As it is, $K - 1$ topics are still governed solely by the data. This need not be the case, as methods similar to those proposed for construction of a background topic (see Appendix B) could be extended. In particular, the Dirichlet prior could be modified directly, or a set of restrictions could be imposed for each topic and groups of words so that certain words cannot appear together, or may only appear together in certain topics. Some of these modifications were considered in Andrzejewski and Zhu (2009), providing another starting point for future extension.

In summary, it is becoming increasingly clear that studies aimed at solving the most challenging problems in cancer genomics involve highly diverse types of data collected on large groups of patients. Many methods will prove useful. We suspect that advantage will be gained from methods that are able to integrate data and account for cohort heterogeneity, allow supervision by outcomes of interest such as survival, provide for patient specific inference, and facilitate prediction of unobserved outcomes. The proposed approach provides tools for these purposes in an effort to help ensure that maximal information is obtained from powerful genomic based studies of disease.

APPENDIX A: THE SURVLDA VARIATIONAL EM

Posterior inference. For a given document $i$ with survival response dyad $(T_i, \delta_i)$, the key quantity of interest is

$$p(\theta_i, Z_{i,1:N_i}|w_{i,1:N_i}, T_i, \delta_i, \alpha, \tau_{1:K}, \beta, h_0) =$$

$$\frac{p(\theta_i|\alpha) \left( \prod_{j=1}^{N_j} p(Z_{ij}|\theta_i)p(W_{ij}|Z_{ij}, \tau_{1:K}) \right) p(T_i, \delta_i|Z_{i,1:N_i}, \beta, h_0)}{\int p(\theta_i|\alpha) \sum_{Z_{i,1:N_i}} \left( \prod_{j=1}^{N_i} p(Z_{ij}|\theta_i)p(W_{ij}|Z_{ij}, \tau_{1:K}) \right) p(T_i, \delta_i|Z_{i,1:N_i}, \beta, h_0) d\theta}$$

(A.1)

where the normalizing value is known as the evidence. As in LDA (Blei, Ng and Jordan, 2003) and sLDA (Blei and McAuliffe, 2008), the evidence cannot be exactly computed efficiently, so we will use mean-field variational inference using Jensen’s inequality to approximate it. For reviews of this and other variational methods, see Wainwright and Jordan (2008) and Jordan et al. (1999).

Let $\pi = \{\alpha, \tau_{1:K}, \beta, h_0\}$ and $q_i(\theta_i, Z_{i,1:N_i})$ denote a variational distribution of the latent variables. For computational tractability, we choose a fully factorized variational distribution:

$$q_i(\theta_i, Z_{i,1:N_i}|\gamma_i, \phi_{1,1:N_i}) = q_i(\theta_i|\gamma_i) \prod_{j=1}^{N_i} q_i(Z_{ij}|\phi_{ij})$$

(A.2)

where

$$\theta_i|\gamma_i \sim \text{Dir}(\gamma_i) \quad \text{and} \quad Z_{ij}|\phi_{ij} \sim \text{Discrete}(\phi_{ij}).$$
With this quantity defined, the lower bound for the evidence given by Jensen’s inequality is

\[
\log p(W_{i,1:N_i}, T_i, \delta_i | \pi) = \log \int_{\theta_i Z_{i,1:N_i}} p(\theta_i, Z_{i,1:N_i}, W_{i,1:N_i}, T_i, \delta_i | \pi) \, d\theta
\]

\[
= \log \int_{\theta_i Z_{i,1:N_i}} p(\theta_i, Z_{i,1:N_i}, W_{i,1:N_i}, T_i, \delta_i | \pi) \frac{q(\theta_i, Z_{i,1:N_i})}{q(\theta_i, Z_{i,1:N_i})} \, d\theta
\]

\[
= \log E_{q_i} \left[ p(\theta_i, Z_{i,1:N_i}, W_{i,1:N_i}, T_i, \delta_i | \pi) \frac{1}{q(\theta_i, Z_{i,1:N_i})} \right]
\]

\[
\geq E_{q_i} \left[ \log p(\theta_i, Z_{i,1:N_i}, W_{i,1:N_i}, T_i, \delta_i | \pi) \right]
- E_{q_i} [\log q(\theta_i, Z_{i,1:N_i})]
\]

(A.3)

where the second term in the lower bound is the entropy \(H(q_i)\) of the variational distribution. We will use \(\mathcal{L}(\cdot)\) to refer to the so-called evidence lower bound (ELBO) given in (A.3). We can expand the ELBO:

\[
\mathcal{L}(W_{i,1:N_i}, T_i, \delta_i | \pi) = E_{q_i} [\log p(\theta_i | \alpha)] + \sum_{j=1}^{N_i} E_{q_i} [\log p(Z_{ij} | \theta_i)] + \sum_{j=1}^{N_i} E_{q_i} [\log p(W_{ij} | Z_{ij}, \tau_{1:K})]
\]

(A.4)

\[
+ E_{q_i} [\log p(T_i, \delta_i | Z_{i,1:N_i}, \beta, h_0)] + H(q_i)
\]

Thus, an approximation of the posterior given in (A.1) is obtained by maximizing \(\mathcal{L}\) with respect to \(\gamma_i\) and \(\phi_{i,1:N_i}\). The first, second and third terms in (A.4), as well as the entropy \(H(q_i)\), are identical to the corresponding terms in the ELBO for LDA (Blei, Ng and Jordan, 2003) and sLDA (Blei and McAuliffe, 2008):

\[
E_{q_i} [\log p(\theta_i | \alpha)] = \log \Gamma \left( \sum_{k=1}^{K} \alpha_k \right) - \sum_{k=1}^{K} \log \Gamma (\alpha_k)
\]

(A.5)

\[
+ \sum_{k=1}^{K} (\alpha_k - 1) \left[ \Psi(\alpha_k) - \Psi \left( \sum_{g=1}^{K} \alpha_g \right) \right]
\]

(A.6)

\[
\sum_{j=1}^{N_i} E_{q_i} [\log p(Z_{ij} | \theta_i)] = \sum_{j=1}^{N_i} \sum_{k=1}^{K} \phi_{ijk} \left[ \Psi(\alpha_k) - \Psi \left( \sum_{g=1}^{K} \alpha_g \right) \right]
\]

(A.7)

\[
\sum_{j=1}^{N_i} E_{q_i} [\log p(W_{ij} | Z_{ij}, \tau_{1:K})] = \sum_{j=1}^{N_i} \sum_{k=1}^{K} \sum_{v=1}^{V} \phi_{ijk} W_{ijv} \log \tau_{kv}
\]

\[
H(q_i) = \left\{ \log \Gamma \left( \sum_{k=1}^{K} \gamma_{ik} \right) - \sum_{k=1}^{K} \log \Gamma (\gamma_{ik}) \right\}
\]

(A.8)

\[
+ \sum_{k=1}^{K} (\gamma_{ik} - 1) \left[ \Psi(\gamma_{ik}) - \Psi \left( \sum_{g=1}^{K} \gamma_{ig} \right) \right]
\]

\[
+ \sum_{j=1}^{N_i} \sum_{k=1}^{K} \phi_{ijk} \log \phi_{ijk}
\]

where \(\Psi\) denotes the digamma function. All that remains is to derive the fourth term of (A.4):
\[ E_q[p(T_i, \delta_i|Z_{i,1:N}, \beta, h_0)] = E_q[\log \left\{ h_0(T_i) \exp (\beta'Z_i) \right\}^\delta_i \times \exp \left\{ -H_0(T_i) \exp (\beta'Z_i) \right\}] \]
\[ = E_q [\delta_i \log h_0(T_i) + \delta_i \beta'Z_i - H_0(T_i) \exp (\beta'Z_i)] \]
\[ = \delta_i \log h_0(T_i) + \delta_i \beta'\bar{\phi}_i - H_0(T_i) \prod_{j=1}^{N_i} \left( \exp (\frac{\beta}{N_i} \phi_{ij}) \right) \]
\[ (A.9) \]

where the \( K \)-vector
\[ \bar{\phi}_i = (1/N_i) \sum_{j=1}^{N_i} \phi_{ij} \]

We use block coordinate-ascent variational inference, maximizing (A.4) with respect to \( \gamma_i \) and then each \( \phi_{ij} \) in turn. As in sLDA (Blei and McAuliffe, 2008), the terms of (A.4) involving \( \gamma_i \) are unchanged from LDA and hence the update for \( \gamma_i \) is
\[ \gamma_i^{new} = \alpha + \sum_{j=1}^{N_i} \phi_{ij} \]
\[ (A.10) \]

The update for a given \( \phi_{ij} \), however, must be derived anew. We first define the following quantities:
\[ \psi_i = \left[ \Psi(\gamma_{i1}) - \Psi \left( \sum_{g=1}^{K} \gamma_{ig} \right), \ldots, \Psi(\gamma_{iK}) - \Psi \left( \sum_{g=1}^{K} \gamma_{ig} \right) \right] \]
\[ \xi_{ij} = \left[ \sum_{v=1}^{V} I(W_{ij} = v) \log \tau_{iv}, \ldots, \sum_{v=1}^{V} I(W_{ij} = v) \log \tau_{Kv} \right] \]

and then take the partial derivative of (A.4) with respect to \( \phi_{ijk} \):
\[ \frac{\partial \mathcal{L}}{\partial \phi_{ijk}} = 0 + \psi_{ik} + \xi_{ijk} + [-\log \phi_{ijk} - 1] \]
\[ + \delta_i \beta_k \bar{N}_i - H_0(T_i) \left( \prod_{m \neq j} \exp \left( \frac{\beta}{N_i} \phi_{im} \right) \right) \exp \left( \frac{\beta_k}{N_i} \right) \]
\[ (A.11) \]

Setting this equal to zero and plugging in \( \phi_{ijk}^{new} \) yields:
\[ \phi_{ijk}^{new} \propto \exp \left[ \psi_{ik} + \xi_{ijk} + \delta_i \beta_k \bar{N}_i - H_0(T_i) \left( \prod_{m \neq j} \exp \left( \frac{\beta}{N_i} \phi_{im} \right) \right) \exp \left( \frac{\beta_k}{N_i} \right) \right] \]
\[ (A.12) \]

where proportionality means that the components of \( \phi_{ijk}^{new} \) are evaluated according to (A.12) and then normalized so that their sum is one. Variational inference proceeds by iteratively updating the variational parameters \( \{\gamma_i, \phi_{i,1:N_i}\} \) according to (A.10) and (A.12) in order to find a local optimum for the ELBO, which in turn best approximates the evidence given in (A.1).
Parameter estimation. We use maximum likelihood estimation based on variational expectation-maximization. Our data are $D = \{W_{i,1:N_i}, T_i, \delta_i\}$.

\begin{equation}
L(\alpha, \tau_{1:K}, \beta, h_0; D) = \sum_{i=1}^{D} \{E_q[\log p(\theta_i, Z_{i,1:N_i}, W_{i,1:N_i}, T_i, \delta_i)] + H(q_i)\}
\end{equation}

In the expectation step (E-step), we use the variational inference algorithm outlined in the first subsection of Appendix A to estimate the approximate posterior distribution for each document-response pair. In the maximization step (M-step), we maximize the corpus-level ELBO with respect to $\pi$, subject to some constraints. First, we take $\alpha$ to be $(\alpha_0/K, \ldots, \alpha_0/K)$ where $\alpha_0$ is specified a priori. This is not necessary; further structure could be placed on $\alpha$, ranging from a simple Dirichlet prior as in Blei, Ng and Jordan (2003) to more complicated structures allowing dependence among the documents more complex than simple conditional independence as in Teh et al. (2006). However we, like Blei and McAuliffe 2008, prefer letting $\alpha$ be user-defined, which is simple and straightforward, yet allows some flexibility in the model specification.

The $\tau_{1:K}$ updates are unchanged from unsupervised LDA (Blei, Ng and Jordan, 2003; Blei and McAuliffe, 2008) and are thus calculated in this manner:

\begin{equation}
\tau_{k}^{new} \propto \sum_{i=1}^{D} \sum_{j=1}^{N_i} I(W_{ij} = v) \phi_{ijk}
\end{equation}

where proportionality means that each $\tau_{k}^{new}$ is normalized to sum to one.

The regression coefficients that comprise $\beta$ and the baseline hazard $h_0$ must be numerically optimized w.r.t. maximizing the portion of the joint ELBO that depends on them. Numerical optimization is required as no closed form can be derived in general for the maximizing choice of $\beta$. The specific computations this process entails depend on the choice for $h_0$. For example, when an exponential survival model is chosen, so that $h_0 = \lambda$, $\beta$ and $\lambda$ are numerically optimized by finding the solutions:

\begin{equation}
(\hat{\beta}^{new}, \lambda^{new}) = \underset{\beta, \lambda}{\text{argmax}} \ L(\beta, \lambda) = \underset{\beta}{\text{argmax}} \ \sum_{i=1}^{D} \left[ \delta_i \log \lambda + \delta_i \beta' \phi_i - \lambda T_i \times \prod_{j=1}^{N_i} \exp \left( \frac{\beta'}{N_i} \phi_{ij} \right) \right]
\end{equation}

Numerical optimization for a Weibull survival model is similar. In contrast, if we use a non-parametric Breslow estimate (Breslow, 1974) for $h_0$, we first update $\beta$ given the current value for $h_0$:

\begin{equation}
\hat{\beta}^{new} = \underset{\beta}{\text{argmax}} \ L(\beta) = \underset{\beta}{\text{argmax}} \ \sum_{i=1}^{D} \left[ \delta_i \log h_0 + \delta_i \beta' \phi_i - H_0(T_i) \prod_{j=1}^{N_i} \exp \left( \frac{\beta'}{N_i} \phi_{ij} \right) \right]
\end{equation}

Then, given the updated $\beta = \hat{\beta}^{new}$, the maximum likelihood estimate of the baseline hazard $h_0$ at the $r^{th}$ ordered survival time $t_r$ is given by Breslow (1974):

\begin{equation}
\hat{h}_0^{new}(t_r | \beta) = \frac{m_r}{(t_r - t_{r-1}) \sum_{j \in R_r} \exp \left( \beta' \phi_j \right)}
\end{equation}

where $m_r$ is the number of failures at time $t_r$ and $R_r$ is the set of patients that have not failed or been censored by time $t_r$. Regardless, once $h_0$ has been updated an estimate of the cumulative baseline hazard $H_0$ follows immediately.
**Prediction for a new document.** Given a new patient with document $w_1:N$ and a fitted model \{\alpha, \tau_{1:K}\}, the posterior mean $\bar{Z}_{new} = \bar{Z}|w_1:N, \alpha, \tau_{1:K}$ can be obtained in order to estimate from what topics this new patient draws words, and in what proportions. This is similar to the procedure outlined in the *Posterior Inference* subsection, except that all survival-related terms in the ELBO are dropped. Thus, under the same variational distribution as given in (A.2), the coordinate ascent updates are

(A.18) \[
\gamma_{new} = \alpha + \sum_{j=1}^{N} \phi_j
\]

(A.19) \[
\phi_{jk}^{new} \propto \exp (\psi_{ik} + \xi_{jk})
\]

where again $j$ indexes words, $k$ indexes topics and proportionality means that the components of $\phi_{jk}^{new}$ are evaluated according to the above update and then normalized so that their sum is one. Note that this variational sequence is identical to that in Blei and McAuliffe (2008), as they point out that it does not depend on the particular response type.

Given $\bar{Z}_{new}$, measures related to topic membership can be predicted for a new document. This may be done qualitatively or quantitatively using the chosen survival model. For example, the predicted median lifetime can be obtained by solving the following equation for $\hat{t}_{med}$:

(A.20) \[
\exp \left[ -H_0 \left( \hat{t}_{med} \right) \exp \left( \beta' \bar{Z}_{new} \right) \right] = \frac{1}{2}
\]

where $H_0$ and $\beta$ are taken from the fitted survLDA model.

**APPENDIX B: IMPOSING A BACKGROUND TOPIC**

As mentioned in the text, the documents we are working on are of our own creation and hence we know their nature. In particular, we know that each patient in the TCGA ovarian cohort received platinum and taxane as treatment with each of these following surgery was an inclusion criteria. Consequently an uninteresting background document would include platinum and taxane and nothing else. The distribution on the background topic may thus be found by placing those weights on those words and re-normalizing so that their sum is one. In order to avoid negative infinities on the log scale, a small weight such as a thousandth of a word may be given to each of the words that do not appear in this background document.

With respect to the variational EM algorithm, imposition of a background topic into LDA may be achieved by specifying one (say the $K^{th}$) topic as the background. In the variational EM, inference in the E-step proceeds as usual but the M-step is modified so that $\tau_K$ is not updated like the other $\tau_{1:(K-1)}$ but is instead fixed according to the distribution designated by the user. Modification of survLDA so that it accepts a background topic is similar except that, in addition to the fixation just outlined, $\beta_K$ is set to 0. In this way the background topic also becomes the topic corresponding to the baseline hazard, and the $\beta_k$ values of the other topics will reflect increased or decreased force of mortality relative to that baseline.

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Fig 1: A heat map of the word distributions for the topics derived from LDA, with two insets marked by black boxes. The background (seventh) topic is not shown. Words are clustered along the rows, topics are clustered across the columns. The colors range from blue (word under-represented in the topic) to red (word over-represented in the topic), with white in the middle (average representation). The insets are close-up views of the boxed regions, which contain the twenty highest-weight words for topics five (upper right heatmap) and six (lower right heatmap).
Fig 2: Top left: A heat map of estimates of $\theta$ (patient-specific distributions over topics) for each patient derived from LDA (upper left) and survLDA (lower left). Topics are given across the rows and patients are clustered across the columns. Colors range from deep blue (0) to red (1). The right panels show Kaplan-Meier curves for the LDA (upper) and survLDA (lower) topics. The background (seventh) topic is not shown. Topic $k$’s curve is generated using all 448 documents, weighted by the $\theta_{ik}$. 
Fig 3: A heat map of the word distributions for the topics derived from survLDA, with two insets marked by black boxes. The background (seventh) topic is not shown. The rows are in the same order as in Figure 2, topics are clustered across the columns. The colors range from blue (word under-represented in the topic) to red (word over-represented in the topic), with white in the middle (average representation). The insets are close-up views of the boxed regions, which contain the twenty highest-weight words for topics 5 and 6. Note the similarities between topics 4 and 1 derived from survLDA and topics 5 and 6 derived from LDA.
Fig 4: Word clouds generated using the twenty highest-weight words for topics 1 (left) and 4 (right) derived from survLDA.
Fig 5: The rows show the twenty highest-weight words from the fourth and fifth topics derived from survLDA. The columns show words (presence in black and absence in white) for fifteen patients: 5 identified as strongly belonging to topic 4 ($\theta_{i,4} > 0.99$, left column), 5 identified as strongly belonging to topic 5 ($\theta_{i,5} > 0.99$, right column), and 5 for whom the model indicates a mixture of the two ($\theta_{i,5} > 0.4$ and $\theta_{i,5} > 0.4$, middle column). The bars on the right indicate the marginal p-values for each gene, which is taken to be the minimum of all p-values from Cox proportional hazard-based score tests among all expression and methylation measurements associated with that gene. Note that only three of the genes shown would have survived a pre-filtering step using a nominal 0.01 level.
Fig 6: Kaplan Meier curves for selected topic structures within the training and testing data. The red (pink) line represents the 99 training set (23 testing set) women whose documents are identified as having more than 80% weight in topic 1; the light blue line is the survival curve for the 26 test set women predicted as having less than 20% weight in topic 1.