# 1 Fast latent variable estimates for new patient data

Ideally, physicians like to give patients fast, in-visit risk estimates whenever new lab results are acquired. A standard implementation of our approach would entail refitting the entire model to get updated posteriors for the subject's latent variables, which can take hours to complete. Instead, we use importance sampling (CITE TEXTBOOK) to get fast latent variable posterior estimates. This can be combined with periodic refitting of the entire model (e.g. every two weeks) as more patients are acquired, to update the population-level parameter posteriors. It may also be possible to attain a fast, "online" update of the population-level parameter posteriors, but there are known obstacles to type of updating which push its solution beyond the scope of our current work (see supplemental materials).

In order to generate proposal values for importance sampling, we start with draws from the posterior of the population-level parameters, obtained by fitting model refXX on the previously observed data. For each draw, we use the conditional distributions in Equation refXX to generate proposed latent variable values for the subject with new data. The importance weights for these proposed latent variable values are then proportional to the likelihood of the newly acquired data, given the proposed parameters and latent variables. Note that proposals can be pre-generated before patients enter the clinic, so that only the weights need to be calculated in real time.

Using this approach, latent variable posterior mean estimates can be computed in approximately 2 seconds. These fast estimates have a correlation of XX% with the estimates from refitting the model on the entire dataset. For reference, estimates from two different runs of the JAGS model have a correlation of XX, due to the stochastic nature of the posterior sampling. We give further details of the importance weighting calculations, and their performance, in the supplementary materials.

## 2 Supplement

#### 2.1 Importance Sampling Procedure

For the purposes of this section, we introduce the following abbreviated form of the model in XX. Let the posterior for our model be

$$p(\theta, b_{1:n}|y_{1:n}) \propto \prod_{i=1}^{n} [f(y_i|b_i, \theta)g(b_i|\theta)]\pi(\theta)$$
(1)

Where  $y_i = ...$  is the vector of measurements for subject i,  $y_{1:n}$  is the list of measurements for the first n subjects,  $b_i = ...$  is a vector of latent variables for

<sup>&</sup>lt;sup>1</sup>This approach is conceptually very similar to the approach of Lee and Chia (2002), who combine a particle filter with periodic MCMC on all dynamic parameters. The dynamic parameters in their work are analogous to the subject-specific parameters in ours.

subject i,  $b_{1:n}$  is a list of latent variables for the first n subjects,  $\theta$  contains the population-level parameters,  $\pi$  is the prior for  $\theta$ , and f and g are multivariate distributions coming from the likelihood in XX. The goal of this section is to use importance weighting to estimate latent for a new subject (indexed by n+1) entering the study.

Our goal is to calculate expectations with respect to the posterior distribution based on all n+1 subjects (i.e.  $p(\theta,b_{1:(n+1)}|y_{1:(n+1)})$ ). Unfortunately, we cannot immediately draw from this distribution, but we can evaluate a function that is proportional to its density (Equation 1). To carry out importance sampling, we need choose a proposal distribution q from which to generate candidate values of  $(\theta,b_{1:(n+1)})$ . We propose the posterior distribution based on the first n subjects.

$$q(\theta, b_{1:(n+1)}) := g(b_{n+1}|\theta)p(\theta, b_{1:n}|y_{1:n})$$

Practically, this consists of taking J draws of  $\theta$  and  $b_{1:n}$  from the previously fitted posterior in Eq 1. Then, conditional on  $\theta$ , we draw  $b_{n+1}$  from the distribution g. We index each of the resulting draws  $(\theta^{(j)}, b_{1:(n+1)}^{(j)})$  by  $j = 1, \ldots, J$ . The importance weights  $w_j$  are then proportional to

$$\begin{array}{ll} w_{j} & \propto & \frac{p(\theta^{(j)},b_{1:(n+1)}^{(j)}|y_{1:(n+1)})}{q(\theta^{(j)},b_{1:(n+1)}^{(j)})} \\ \\ & \propto & \frac{\prod_{i=1}^{n+1}[f(y_{i}|b_{i}^{(j)},\theta^{(j)})g(b_{i}^{(j)}|\theta^{(j)})]\pi(\theta^{(j)})}{g(b_{n+1}^{(j)}|\theta^{(j)})\prod_{i=1}^{n}[f(y_{i}|b_{i}^{(j)},\theta^{(j)})g(b_{i}|\theta^{(j)})]\pi(\theta^{(j)})} \\ \\ & = & f(y_{i}|b_{i}^{(j)},\theta^{(j)}) \end{array}$$

The final weights are then standardized to sum to 1.

#### 2.2 Full model online updates

We generally propose that importance sampling be used for fast, in-visit estimates of patient's latent risk. This can be combin with periodic full model re-fitting to update the population-level parameter posteriors. The issue with this approach is that computational costs increase as more and more patients are required, making the total computation take no less than quadratic time. The task of updating a hierarchical model in constant time is an open problem.

Some initial work on online updates has been proposed in the field of text analysis. Hoffman et al. (2010) applied a variational Bayesian approach, but this has some problems [need to explore/talk to Beka]? Canini et al. (2009) consider online sampling methods for text analysis, and recommend a particle filter approach (also known as Sequential Importance Resampling) (Need canonical citation). However, all of the online methods considered by Canini et al. do not perform as well as refitting on the entire dataset, in a non-online fashion.

Our model also differs from Canini et al. (2009) in a way that further complicate the use of particle filters. Like Canini et al., we assume that our population

distribution is *static* over time. In other words, we believe that the populationlevel parameters do not change as we acquire new data. The presence of such static parameters is known to cause particle filters to break down (see Andrieu et al. (2005), section II, for an intuitive illustration). Canini et al. mitigate this issue by analytically integrating out the population-level parameters, but this approach is not feasible in our case.

### References

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