A Primer on Small-Area Estimation

A Case Study on Mortality in Massachusetts Census Tracts

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Small Area Estimation: What Is It and Why It Matters?

Small Area Estimation generally refers to the estimation of a quantity of interest for a small subpopulation. In demography, Small Area Estimation problems arise in many situations because most demographic events (birth, death, and migration) are relatively rare, especially when their occurrence is measured in specific age-ranges and periods.

For example, in 2021, the US mortality rates for age groups 1-4, 5-9, 10-14, and 15-19 are all below 1 every 1,000 residents (HMD, 2023). Which, given the US age-distribution, means we would need at least 15,000 residents to observe at least one death in each of these age-groups each year.

Given that about 1 every 3 counties in the US have fewer than 15,000 residents, you see why techniques for modelling demographic quantities in these areas are crucial.

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Why are Zero Death Counts a Problem?

The issue with zero death counts in small populations is that they provide little information about the underlying mortality rate. They give us a sense that mortality is low but make it very difficult for us to say where exactly the rate should be.

Clearly, a direct estimation approach would fail in this case because a mortality rate equal to zero is effectively impossible. Even if tiny, the probability of dying over a positive time interval should always be positive.

In other words, with a large enough population we should always observe a strictly positive number of deaths.

Reliable versus Unreliable Data

Our beloved demographic methods textbook (Preston et al., 2001) discusses few statistical models for mortality. The Gompertz model (Gompertz, 1825), the Brass relational model (Brass, 1971), and the Lee-Carter model for forecasting mortality (Lee and Carter, 1992).

With the exception of the Lee-Carter model, the other models are only introduced in the context of unreliable data (together with model life-tables). Indeed, the book gives the impression that direct estimation is always preferred when the data is deemed reliable.

Reliable versus Unreliable Data

However, models are useful even when the data is complete and reliable.

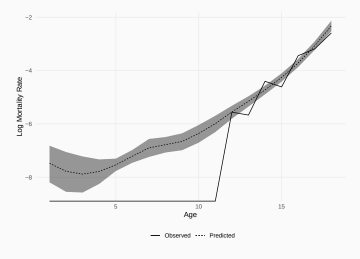


Figure 1: Direct vs. Model-Based Estimation

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Hierarchical Models: A General Purpose Tool for Harder Estimation Problems

Hierarchical models (also called multilevel models, mixed effects models, or random effects models - don't ask me why all these names) can often provide a solution to estimation issues arising with sparse data.

From the frequentist viewpoint, hierarchical models can be thought of as models in which the parameters themselves are random variables and have a distribution whose parameters we are trying to estimate.

If this sounds confusing it is likely because in the frequentist framework we make distributional statements about y but never about β (e.g. the rather absurd language about confidence intervals). If this does not sound right to you it's probably because you are thinking of $\hat{\beta}$, which is indeed a random variable but should not be confused with β .¹

This is probably confusing but it may become clearer with an example (after one more slide of theory).

¹This paragraph alone is a reason enough to switch over to a Bayesian approach.

Bayesian Data Analysis: Bayesian Model Specification in a Nutshell

Every Bayesian model has three components²:

- Likelihood p(data|unknowns). A function showing the probability of observing the data, for any particular value of the unknowns. A way of summarizing the information about unknowns contained in the data.
- Prior distribution p(unknowns). A probability distribution summarizing information about the unknowns, beyond what is contained in the data.
- Posterior distribution p(unknowns|data). A probability distribution summarizing information about the unknowns, after combining information from the likelihood and prior.

Let's go to an example.

²The excellent language describing each component is borrowed from Bryant and Zhang (2018)

A Demographer's Best Friend: The Poisson Model

A one-parameter Poisson model:

Likelihood:
$$p(y|\theta) = \prod_{i} \frac{\theta^{y_i} e^{-\theta}}{y_i!} = \frac{\theta^{\sum_{i} y_i} e^{-n\theta}}{\prod_{i} y_i!}$$
 (1)

Prior:
$$p(\theta|\alpha,\beta) = \frac{\beta^{\alpha}}{\Gamma(\alpha)} \theta^{\alpha-1} e^{-\beta\theta}$$
 (2)

Posterior:
$$p(\theta|y) \propto p(y|\theta)p(\theta|\alpha,\beta)$$
 (3)

$$\propto \frac{\theta^{\sum_{i} y_{i}} e^{-n\theta}}{\prod_{i} y_{i}!} \frac{\beta^{\alpha}}{\Gamma(\alpha)} \theta^{\alpha - 1} e^{-\beta \theta}$$
(4)

$$\propto \theta^{\sum_{i} y_{i}} e^{-n\theta} \frac{\beta^{\alpha}}{\Gamma(\alpha)} \theta^{\alpha - 1} e^{-\beta \theta} \tag{5}$$

$$\propto \theta^{\sum_i y_i + \alpha - 1} e^{-(n\theta + \beta)}$$
 (6)

$$\implies \theta | y \sim \text{Gamma}\left(\sum_{i} y_{i} + \alpha, n + \beta\right) \tag{7}$$

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A Demographer's Best Friend: The Poisson Model

You can run this example with the file BDAOverview.Rmd.

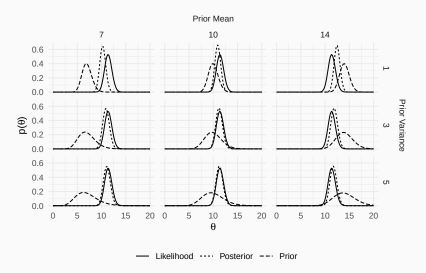


Figure 2: Bayesian Inference for the One-Parameter Poisson Model

Bayesian Data Analysis: Regression with Multiple Groups

Now we are going to work through a more complex example. Open the following files in RStudio:

- · hierarchicalRegression.Rmd
- · regressionModel.stan
- · regressionModelFE.stan
- · regressionModelH.stan

Estimating Life Expectancy for Massachusetts Census Tracts

We will now work through a real application of these models to the estimation of life expectancy for Massachusetts Census tracts. The codes for running this example are:

- cleanData.Rmd: Cleans county-subdivision data and creates age-mortality principal components. [This will not run but it's there to show how the PCs are computed].
- runAZBModelForTracts.Rmd: Prepares the data for the model, fits the model, and checks the model fit.
- · AZBModelForTracts.stan: Stan code for the model.
- createLifeTablesTracts.Rmd: Create life tables and estimates life expectancies for each tract.

Data

Now that we have gotten a sense of what hierarchical models do, we can get to a real application to estimate tract-level life-expectancies in Massachusetts.

- Deaths: Tract-Level death counts by five-year age groups for the period 2017-2021.
- Population: Population counts by five-year age groups from the 2020 Census.

I confidentialized the mortality data by applying random rounding to base three. The exposure data (population) is not ideal but no yearly population estimates are produced by the Census Bureau at the tract level.

Analytical Framework

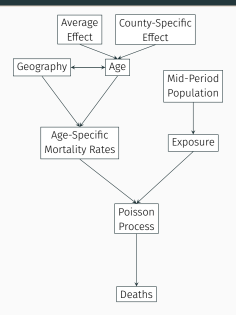


Figure 3: Analytical Framework

The Model: Overview

Denoting deaths for age group a, and tract s with $D_{a,s}$, the corresponding population with $P_{a,s}$, and the corresponding mortality rate with $m_{a,s}$, we have:

$$D_{a,s} \sim Poisson(m_{a,s} \cdot P_{a,s})$$
 (8)

$$log(m_{a,s}) = \beta_0 + \beta_1^A \cdot AC_{1,a} + (\beta_2^A + \beta_{2,s}^A) \cdot AC_{2,a} + (\beta_3^A + \beta_{3,s}^A) \cdot AC_{3,a} + \beta_s$$
 (9)

$$\beta_0, \beta_1^A, \beta_2^A, \beta_3^A \sim Unif(-\infty, +\infty)$$
 (10)

$$\beta_{2,s}^{A} \sim Normal(0, \sigma_2^{A})$$
 (11)

$$\beta_{3,s}^A \sim Normal(0, \sigma_3^A)$$
 (12)

$$\beta_{\rm S} \sim Normal(0, \sigma^{\rm S})$$
 (13)

This model is indebted to Alexander et al. (2017). More complex models are discussed in Bryant and Zhang (2018), which also provides a broad overview of applications of Bayesian statistics to demography.

The Model: Age

Age is modelled through tract-specific coefficients on three vectors (AC_1 , AC_2 and AC_3) representing common underlying features of the age-mortality curves. I obtained the three vectors as the first three principal components of county-level age-mortality curves.

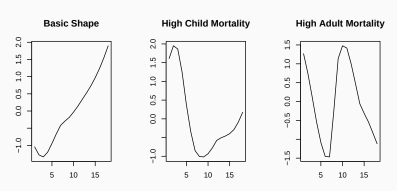


Figure 4: First Three Principal Components of County-Level Age-Mortality Curves

The Model: Geography

Geography is modelled with a set of county-specific intercepts assumed to come from the same Normal distribution. This hierarchical structure induces global smoothing, especially for less populated counties.

The Model: Summary

The model combines all the components we have seen and is estimated (with several implementation challenges) in Stan (Stan Development Team, 2023).

$$\begin{split} &D_{a,s} \sim Poisson(m_{a,s} \cdot P_{a,s}) \\ &log(m_{a,s}) = \beta_0 + \beta_1^A \cdot AC_{1,a} + (\beta_2^A + \beta_{2,s}^A) \cdot AC_{2,a} + (\beta_3^A + \beta_{3,s}^A) \cdot AC_{3,a} + \beta_s \end{split}$$

The result are 4,000 samples from the posterior distribution of each parameter that are used to construct posterior samples for the log mortality rates of all 5,187 data points.

Results: Does the Model Fit the Data? Age Patterns

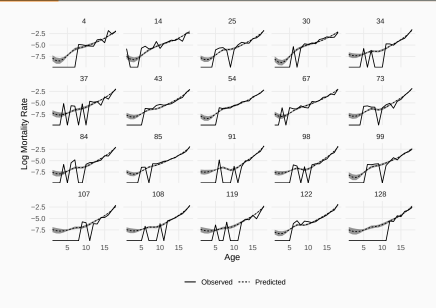


Figure 5: Comparing Estimated and Observed Mortality Rates by Age

Results: Does the Model Fit the Data? Age Patterns

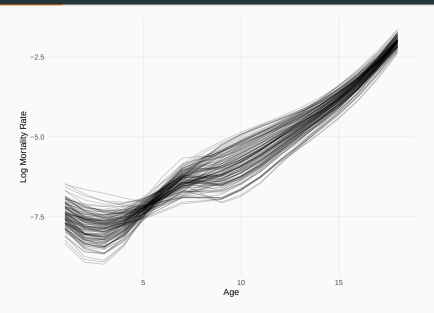


Figure 6: Age-Mortality Curves

Results: Does the Model Fit the Data? Geography

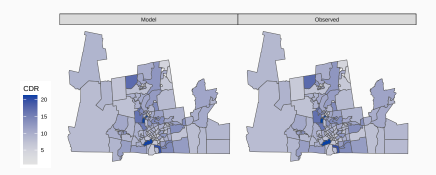


Figure 7: Crude Death Rates by Tract

Results: Tract-Level Life Expectancies

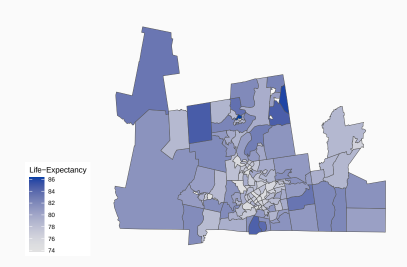


Figure 8: Life Expectancies by Tract

Summary and Key Takeaways: The Good Stuff

The good stuff:

- · Models for mortality rates are a very useful tool in a demographer's toolkit.
- They are useful even when the data is complete and reliable and make it much easier to deal with uncertainty in a systematic way (sorry Chiang (1984), it was good until it lasted).
- (personal opinion) The Bayesian approach (especially with pure Stan) forces you
 to think hard about the model and demands a very good understanding of what
 you are doing (which is not always the case with frequentist procedures).

Summary and Key Takeaways: The Caveats

The caveats:

- Care should be used when selecting an appropriate model. Are we capturing all important features of the data? (Not always easy to answer).
- The Achilles's heel of Bayesian inference is that it is computationally intensive and hard to scale when the number of data points increases (Stan is the fastest of the MCMC approaches but approximation approaches like INLA or Template Model Builder are generally faster). See Dwyer-Lindgren et al. (2016) for an example of county-level modeling using Template Model Builder.
- Bayesian approaches are not the only option. The National Center for Health Statistics undertook a massive (and surprisingly ignored) effort to produce tract-level life-expectancy estimates using a frequentist (but poorly documented) approach (Arias et al., 2018).

References i

References

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