

# Hierarchical Bayesian Model

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# Big Picture: What Are We Modeling?

We are modeling **how antibody levels change over time** in response to infection, using data from multiple individuals and multiple **biomarkers** (10 antigen-isotype combinations, so (  $j = 1, 2, \dots, 10$  )).

We want to:

- Understand the average pattern for each biomarker
- Allow each person's response to vary
- Share information across individuals to improve estimates

This is a perfect use case for a **hierarchical Bayesian model**.

## Step 1: Individual-Level Parameters (Subject-Level)

Each person ( $i$ ), for biomarker ( $j$ ), has their own unique set of parameters:

$$\theta_{ij} = \begin{bmatrix} y_{0,ij} \\ b_{0,ij} \\ \mu_{b,ij} \\ \mu_{y,ij} \\ \gamma_{ij} \\ \alpha_{ij} \\ \rho_{ij} \end{bmatrix}$$

These describe the antibody curve for person ( $i$ ) and biomarker ( $j$ ): the starting level, how fast it rises, peaks, and decays.

## Step 2: Population-Level Parameters (Per Biomarker $j$ )

Now we summarize how people typically behave for each **biomarker**:

$\mu_j$  = population mean vector for biomarker  $j$

This means:

- For biomarker (  $j$  ), we believe the true average antibody trajectory is governed by parameters (  $\mu_j$  ).
- But we don't know (  $\mu_j$  ) — so we estimate it using data across all individuals.

## Step 3: Hierarchical Modeling Structure

We assume each individual's parameter vector  $\theta_{ij}$  is drawn from a multivariate normal distribution:

$$\theta_{ij} \sim \mathcal{N}(\mu_j, \Sigma_j)$$

- $\mu_j$  : the population-level mean for biomarker (  $j$  )
- $\Sigma_j$  :  $j \times j$  covariance matrix describing how the parameters co-vary

This is where the “**borrowing strength**” happens. Even if someone has sparse data, we can still make good inferences by **using the group-level pattern**.

## Step 4: Priors on Population Means — “Hyperpriors”

But wait — since we are Bayesian, so we also need a prior belief about  $\mu_j$  :

$$\mu_j \sim \mathcal{N}(\mu_{\text{hyp},j}, \Omega_{\text{hyp},j})$$

Where:

- $\mu_{\text{hyp},j}$  : prior guess for the population mean (e.g., a vector of zeros )
- $\Omega_{\text{hyp},j}$  : uncertainty about that guess (e.g.,  $100 \cdot I_7$  for weakly informative prior)

This is a **hyperprior**, because it's a prior on a prior-level parameter.

## Step 5: Priors on Covariance — “Priors on Variability”

We also don't know how much individual parameters vary. So we assign a **Wishart prior** to the **inverse** covariance matrix:

$$\Sigma_j^{-1} \sim \mathcal{W}(\Omega_j, \nu_j)$$

- $\Omega_j$  : prior scale matrix (small variance across parameters, often  $0.1 \cdot I_7$ )
- $\nu_j$  : degrees of freedom

This tells the model how much we expect individuals to vary from the average for biomarker  $j$ .

## Step 6: Measurement Error Model

Our observations are noisy! So we model the observed log-antibody levels  $\log(y_{obs,ij})$  like this:

$$\log(y_{obs,ij}) \sim \mathcal{N}(\log(y_{pred,ij}), \tau_j^{-1})$$

Where  $\tau_j \sim \text{Gamma}(a_j, b_j)$  is a prior on measurement precision for biomarker  $j$ .



## Step 7: Putting It All Together

The model is built hierarchically across five conceptual levels:

- 1 **Observed data:** log antibody concentrations from serum samples
- 2 **Individual-level parameters:** specific antibody dynamics for each subject-biomarker pair
- 3 **Population-level means:** average antibody parameters for each biomarker
- 4 **Hyperpriors on means:** our belief about the likely range of population means
- 5 **Priors on variability:** our belief about individual variation around those means

This structure lets us account for uncertainty at every level, while borrowing strength across subjects and biomarkers.

# Summary of the Hierarchy

Let's stack it up top-down:

## ① Top Level:

- For each biomarker  $j$ , the true mean antibody trajectory parameters  $\mu_j$  come from a prior  $\mathcal{N}(\mu_{\text{hyp},j}, \Omega_{\text{hyp},j})$

## ② Middle Level:

- For each person  $i$ , their parameters  $\theta_{ij} \sim \mathcal{N}(\mu_j, \Sigma_j)$

## ③ Bottom Level:

- Their actual observed antibody levels are noisy measurements of predictions from  $\theta_{ij}$

## Step 3: Modeling Individuals

We say:

$$\theta_{ij} \sim \mathcal{N}(\mu_j, \Sigma_j)$$

This means:

- For biomarker  $j$ , each subject  $i$  has their own parameter vector  $\theta_{ij}$
- These vectors come from a **Normal distribution** centered at  $\mu_j$  (the population mean for that biomarker)
- $\Sigma_j$  is the covariance matrix capturing variation across individuals for that biomarker

But here's the catch: **we don't know**  $\mu_j$  or  $\Sigma_j$  yet.

# So How Do We Handle the Unknowns?

In **Bayesian modeling**, we treat unknowns as **random variables** too. So instead of fixing  $\mu_j$  and  $\Sigma_j$ , we say:

*“Let’s estimate them, but we’ll put a prior belief on them to guide the learning.”*

This brings us to:

## Step 4: Priors on $\mu_j$

$$\mu_j \sim \mathcal{N}(\mu_{\text{hyp},j}, \Omega_{\text{hyp},j})$$

### Explanation:

- $\mu_j$ : unknown population-level mean of the parameters for biomarker  $j$
- We say:
  - “We believe  $\mu_j$  comes from another normal distribution”*
    - *Centered at  $\mu_{\text{hyp},j}$  — a guess for what the mean might be*
    - *With spread  $\Omega_{\text{hyp},j}$  — how confident we are in that guess*

If we want to be very flexible, we make this prior **weakly informative**:

- Set  $\mu_{\text{hyp},j} = 0$
- Set  $\Omega_{\text{hyp},j} = 100 \cdot I_7$ , where  $I_7$  is the identity matrix (saying we are uncertain)

This is a **prior on a population-level parameter** — a “hyperprior”.

## Step 5: Priors on $\Sigma_j$

We also don't know how much individual parameters vary, so we say:

$$\Sigma_j^{-1} \sim \mathcal{W}(\Omega_j, \nu_j)$$

This is a **Wishart prior** on the **precision matrix** (inverse of covariance).  
Why?

- In multivariate stats, it's common to use the Wishart distribution as a prior for covariance matrices
- $\Omega_j$ : the scale (like the average covariance we expect)
- $\nu_j$ : degrees of freedom (how confident we are)

If we want to be uninformative, we might say:

- $\Omega_j = 0.1 \cdot I_7$
- $\nu_j = 8$

That allows a wide range of possible covariance matrices.

# Summary of Why Priors Show Up

Priors appear at step 4 and 5 because we are now **modeling the parameters themselves**.

In Bayesian statistics:

- Every unknown quantity is treated as a random variable
- Every random variable must have a probability distribution
- That's what **priors** are

They let us encode our beliefs, and importantly, they let us **regularize the model** so it doesn't overfit sparse or noisy data.