## Extending the Hierarchical Model for Antibody Kinetics

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#### Overview

- Incorporates feedback from Dr. Morrison and Dr.Aiemjoy
- Focus exclusively on (Teunis and Eijkeren 2016) model
- Clarifies model dynamics: growth, clearance, decay
- $\bullet$  Uses updated parameter notation:  $\mu_{y}$ ,  $\mu_{b}$ ,  $\gamma$ ,  $\alpha$ ,  $\rho$
- Assumes block-diagonal covariance structure across biomarkers

## Observation Model (Data Level)

Observed (log-transformed) antibody levels:

$$\log(y_{\mathsf{obs},ij}) \sim \mathcal{N}(\mu_{\log y,ij}, \tau_j^{-1}) \tag{1}$$

#### Where:

- ullet  $y_{{
  m obs},ij}$ : Observed antibody level for subject i and biomarker j
- $\mu_{\log y,ij}$  is the **expected log antibody level**, computed from the two-phase model using subject-level parameters  $\theta_{ij}$ .
- $\theta_{ij}$ : Subject-level latent parameters (e.g.,  $y_0, \alpha, \rho$ ) used to define the predicted antibody curve

Measurement precision prior:

$$\tau_j \sim \mathsf{Gamma}(a_j, b_j) \tag{2}$$

## Parameter Summary

Table 1: Parameter summary for antibody kinetics model.

Symbol	Description
$\begin{array}{c} \mu_y \\ \mu_b \\ \gamma \end{array}$	Antibody production rate (growth phase) Pathogen replication rate Clearance rate (by antibodies)
$\alpha$	Antibody decay rate
$\rho$	Shape of antibody decay (power-law)
$t_1$	Time of peak response
$y_1$	Peak antibody concentration

**Note:** Only the first 6 are typically estimated.  $y_1$  is derived from the ODE solution at  $t_1$ .

## Within-Host ODE System (Teunis and Eijkeren 2016)

Two-phase within-host antibody kinetics:

$$\frac{dy}{dt} = \begin{cases} \mu_y y(t), & t \le t_1 \\ -\alpha y(t)^\rho, & t > t_1 \end{cases} \quad \text{with } \frac{db}{dt} = \mu_b b(t) - \gamma y(t) \tag{3}$$

Initial conditions:  $y(0) = y_0$ ,  $b(0) = b_0$ 

**Key transition:**  $t_1$  is the time when  $b(t_1)=0$ 

 $\mbox{ Derived quantity: } y_1 = y(t_1)$ 

#### Closed-Form Solutions

#### Antibody concentration y(t)

•  $t \le t_1$ :

$$y(t) = y_0 e^{\mu_y t}$$

•  $t > t_1$ :

$$y(t) = y_1 \left( 1 + (\rho - 1) \alpha y_1^{\rho - 1} (t - t_1) \right)^{-\frac{1}{\rho - 1}}$$

#### Pathogen load b(t)

•  $t \le t_1$ :

$$b(t) = b_0 e^{\mu_b t} - \frac{\gamma y_0}{\mu_y - \mu_b} \left( e^{\mu_y t} - e^{\mu_b t} \right)$$

•  $t > t_1$ :

## Time of Peak Response

Peak Time  $t_1$ 

$$t_1 = \frac{1}{\mu_y - \mu_b} \log \left( 1 + \frac{(\mu_y - \mu_b)b_0}{\gamma y_0} \right) \tag{4}$$

Peak Antibody Level  $y_1$ 

$$y_1 = y_0 e^{\mu_y t_1} \tag{5}$$

# Model Comparison: (Teunis and Eijkeren 2016) vs serodynamics

Table 2: Comparison of Teunis (2016) model and serodynamic's model assumptions.

Component	(Teunis and Eijkeren 2016)	serodynamics
Pathogen ODE	$\mu_0 b(t) - cy(t)$	$\mu_b b(t) - \gamma y(t)$
Antibody ODE	$\mu y(t)$	$\mu_{y}y(t)$
$(pre-t_1)$		J
Antibody ODE	$-\alpha y(t)^r$	$-\alpha y(t)^{ ho}$
$(post ext{-}t_1)$		
Antibody growth	Pathogen-driven	Self-driven
type		exponential
Antibody rate	$\mu$	$\mu_y$
name		Ü
$t_1$ formula $\;\;\;$	Uses $\mu_0$ , $\mu$ , $b_0$ , $c$ , $y_0$	Uses $\mu_b$ , $\mu_y$ , etc.

## Full Parameter Model (7 Parameters)

#### Subject-level parameters:

$$\theta_{ij} \sim \mathcal{N}(\mu_j, \, \Sigma_j), \quad \theta_{ij} = \begin{bmatrix} y_{0,ij} \\ b_{0,ij} \\ \mu_{b,ij} \\ \mu_{y,ij} \\ \gamma_{ij} \\ \alpha_{ij} \\ \rho_{ij} \end{bmatrix}$$

#### **Hyperparameters** – Means:

- ullet  $\mu_j$ : population-level mean vector for biomarker j
- Prior on  $\mu_j$ :

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

#### From Full 7 Parameters to 5 Latent Parameters

• Although the model estimates 7 parameters, for modeling antibody kinetics y(t), we focus on **5-parameter subset**:

$$y_0, \ t_1(\text{derived}), \ y_1(\text{derived}), \ \alpha, \ \rho$$

• These 5 parameters are **log-transformed** into the latent parameters  $\theta_{-}ij$  used for modeling.

## 5 Core Parameters Used for Curve Drawing

In this presentation, we focus on **5 key parameters** required to draw antibody curves:

- $y_0$ : initial antibody level
- $t_1$ : time of peak antibody response
- $y_1$ : peak antibody level
- $\alpha$ : decay rate
- $\rho$ : shape of decay

Note:  $t_1$  and  $y_1$  are  $\mbox{derived}\mbox{ from the full model}$  - These 5 are sufficient for prediction and plotting

# Classifying Model Parameters ((Teunis and Eijkeren 2016) Structure)

#### **Estimated Parameters (7 total):**

- Core model parameters (5):  $\mu_b$ ,  $\mu_y$ ,  $\gamma$ ,  $\alpha$ ,  $\rho$
- Initial conditions (2):  $y_0$ ,  $b_0$

#### **Derived Quantity (not estimated):**

ullet  $y_1$ : peak antibody level computed as  $y(t_1)$ 

# Subject-Level Parameters (Latent Version = serodynamics)

Each subject i and biomarker j has latent parameters:

$$\theta_{ij} = \begin{bmatrix} \log(y_{0,ij}) \\ \log(y_{1,ij} - y_{0,ij}) \\ \log(t_{1,ij}) \\ \log(\alpha_{ij}) \\ \log(\rho_{ij} - 1) \end{bmatrix}$$
 (6)

Distribution:

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

## Why $y_1$ Is Not Fit Directly

•  $y_1$  is the antibody level at the time the pathogen is cleared:

$$y_1 = y(t_1)$$
 where  $b(t_1) = 0$ 

- $y_1$  is not an "input" it is **computed** from:
  - $\mu_{y}$ ,  $y_{0}$ ,  $b_{0}$ ,  $\mu_{b}$ ,  $\gamma$
  - ullet via solution of ODEs to find  $t_1$  and compute  $y(t_1)$

In other words:  $y_1$  is a **derived output**, not a fit parameter.

## How $y_1$ Is Computed

•  $y_1$  is computed by solving the ODE system:

$$\frac{dy}{dt} = \mu_y y(t), \quad \frac{db}{dt} = \mu_b b(t) - \gamma y(t)$$

• Evaluate y(t) at  $t = t_1$  using ODE solution:

$$y_1=y(t_1;\mu_y,y_0,b_0,\mu_b,\gamma)$$

## Recap: What We Estimate

#### Seven model parameters (7-parameter model for full dynamics):

- $\mu_b$ ,  $\mu_u$ ,  $\gamma$ ,  $\alpha$ ,  $\rho$  (biological process)
- $y_0$ ,  $b_0$  (initial state)

#### **Derived quantity:**

ullet  $y_1=y(t_1)$  — not directly estimated, computed

#### 5-parameter subset for curve visualization:

•  $y_0$ ,  $y_1$ ,  $t_1$ ,  $\alpha$ ,  $\rho$ 

## Hierarchical Bayesian Structure (serodynamics)

#### Individual parameters:

$$\theta_{ij} = \begin{bmatrix} \log(y_{0,ij}) \\ \log(y_{1,ij} - y_{0,ij}) \\ \log(t_{1,ij}) \\ \log(\alpha_{ij}) \\ \log(\rho_{ij} - 1) \end{bmatrix} \sim \mathcal{N}(\mu_j, \, \Sigma_j)$$

#### **Hyperparameters:**

- $\mu_j$ : population-level means (per biomarker j)
- $\Sigma_{j}$ :  $5 \times 5$  covariance matrix over parameters

# Subject-Level Parameters: $\theta_{ij}$

$$\theta_{ij} \sim \mathcal{N}(\mu_j,\, \Sigma_j), \quad \theta_{ij} \in \mathbb{R}^5$$

Where:

$$\theta_{ij} = \begin{bmatrix} \log(y_{0,ij}) \\ \log(y_{1,ij} - y_{0,ij}) \\ \log(t_{1,ij}) \\ \log(\alpha_{ij}) \\ \log(\rho_{ij} - 1) \end{bmatrix}, \quad \Sigma_j \in \mathbb{R}^{5 \times 5}$$

Each subject i has a unique 5-parameter vector per biomarker j, capturing individual-level variation in antibody dynamics.

## Hyperparameters: Priors on Population Means

#### Population-level means:

$$\boldsymbol{\mu}_j \sim \mathcal{N}(\boldsymbol{\mu}_{\mathrm{hyp},j}, \boldsymbol{\Omega}_{\mathrm{hyp},j})$$

#### Interpretation:

- ullet  $\mu_j$ : average parameter vector for biomarker j
- $\mu_{\text{hyp.},i}$ : prior guess (e.g., vector of zeros)
- $\bullet$   $\Omega_{\mathrm{hyp},j}^{\mathrm{TTS}}$ : covariance matrix encoding uncertainty

#### Example:

$$\mu_{\mathrm{hyp},j} = 0, \quad \Omega_{\mathrm{hyp},j} = 100 \cdot I_5$$

## Hyperparameters: Priors on Covariance

#### Covariance across parameters:

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

- $\Sigma_i$ : variability/covariance in subject-level parameters
- $\Omega_i$ : prior scale matrix
- $\nu_i$ : degrees of freedom

#### **Example:**

$$\Omega_j = 0.1 \cdot I_5, \quad \nu_j = 6$$

#### Measurement Error and Precision Priors

#### Observed antibody levels:

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

#### **Precision prior:**

$$\tau_j \sim \mathsf{Gamma}(a_j,b_j)$$

- $\tau_i$ : shared measurement precision for biomarker j
- Gamma prior allows flexible noise modeling

## Matrix Algebra Computation

Let P=5 (parameters), J biomarkers. Then:

$$\Theta_i = \begin{bmatrix} \theta_{i1} & \theta_{i2} & \cdots & \theta_{iB} \end{bmatrix} \in \mathbb{R}^{P \times B}$$

Assume:

$$\mathrm{vec}(\Theta_i) \sim \mathcal{N}(\mathrm{vec}(M), \Sigma_P \otimes I_B)$$

## Matrix Algebra - Simplified Structure

Setup: 
$$\Theta_i \in \mathbb{R}^{P \times B}$$

Model:

$$\mathrm{vec}(\Theta_i) \sim \mathcal{N}(\mathrm{vec}(M), \Sigma_P \otimes I_B)$$

- $\Sigma_P$ : 5×5 covariance (same across biomarkers)
- $\bullet$   $I_B$ : biomarkers assumed uncorrelated
- Block-diagonal covariance

# Understanding $vec(\Theta_i)$

Each  $\theta_{ij} \in \mathbb{R}^7$ :

$$\theta_{ij} = \begin{bmatrix} y_0 \\ b_0 \\ \mu_0 \\ \mu_1 \\ c \\ \alpha \\ r \end{bmatrix}$$

Flattening:

$$\operatorname{vec}(\Theta_i) \in \mathbb{R}^{5B \times 1}$$

## Understanding vec(M)

Let 
$$M = [\mu_1 \, \mu_2 \, \cdots \, \mu_B] \in \mathbb{R}^{5 \times B}$$

Example for B=3:

$$M = \begin{bmatrix} \mu_{1,1} & \mu_{1,2} & \mu_{1,3} \\ \mu_{2,1} & \mu_{2,2} & \mu_{2,3} \\ \mu_{3,1} & \mu_{3,2} & \mu_{3,3} \\ \mu_{4,1} & \mu_{4,2} & \mu_{4,3} \\ \mu_{5,1} & \mu_{5,2} & \mu_{5,3} \end{bmatrix}$$

# Covariance Structure: $\Sigma_P \otimes I_B$

$$\mathsf{Cov}(\mathsf{vec}(\Theta_i)) = \Sigma_P \otimes I_B$$

- $\Sigma_P$ : parameter covariance matrix
- ullet  $I_B$ : biomarker-wise independence
- Kronecker product yields block-diagonal matrix

### Example: Kronecker Product with P=2, B=3

Let:

$$\Sigma_P = \begin{bmatrix} \sigma_{11} & \sigma_{12} \\ \sigma_{21} & \sigma_{22} \end{bmatrix}, \quad I_B = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

Then:

$$\Sigma_P \otimes I_B \in \mathbb{R}^{6 \times 6}$$

# Expanded Matrix: $\Sigma_P \otimes I_B$

$$\Sigma_P \otimes I_B = \begin{bmatrix} \sigma_{11} & 0 & 0 & \sigma_{12} & 0 & 0 \\ 0 & \sigma_{11} & 0 & 0 & \sigma_{12} & 0 \\ 0 & 0 & \sigma_{11} & 0 & 0 & \sigma_{12} \\ \sigma_{21} & 0 & 0 & \sigma_{22} & 0 & 0 \\ 0 & \sigma_{21} & 0 & 0 & \sigma_{22} & 0 \\ 0 & 0 & \sigma_{21} & 0 & 0 & \sigma_{22} \end{bmatrix}$$

## Next Steps: Modeling Correlation Across Biomarkers

#### Current Limitation:

ullet Biomarkers assumed independent:  $I_B$ 

#### Planned Extension:

• Use full covariance  $\Sigma_B$ :

$$\mathsf{Cov}(\mathsf{vec}(\Theta_i)) = \Sigma_P \otimes \Sigma_B$$

## Extending to Correlated Biomarkers

Assume 
$$P=3$$
,  $B=3$ 

Define:

$$\Sigma_K = \begin{bmatrix} \sigma_{11} & \sigma_{12} & \sigma_{13} \\ \sigma_{21} & \sigma_{22} & \sigma_{23} \\ \sigma_{31} & \sigma_{32} & \sigma_{33} \end{bmatrix}, \quad \Sigma_J = \begin{bmatrix} \tau_{11} & \tau_{12} & \tau_{13} \\ \tau_{21} & \tau_{22} & \tau_{23} \\ \tau_{31} & \tau_{32} & \tau_{33} \end{bmatrix}$$

## Kronecker Product Structure: $\Sigma_K \otimes \Sigma_J$

$$\Sigma_K \otimes \Sigma_J = \begin{bmatrix} \sigma_{11} \Sigma_J & \sigma_{12} \Sigma_J & \sigma_{13} \Sigma_J \\ \sigma_{21} \Sigma_J & \sigma_{22} \Sigma_J & \sigma_{23} \Sigma_J \\ \sigma_{31} \Sigma_J & \sigma_{32} \Sigma_J & \sigma_{33} \Sigma_J \end{bmatrix}$$

Now biomarkers and parameters can be correlated.

# Expanded Form: $\Sigma_K \otimes \Sigma_J$ (3x3)

The  $9\times 9$  matrix contains all combinations  $\sigma_{ab}\tau_{cd}$ 

Not block-diagonal — includes cross-biomarker correlation

## Practical To-Do List (for Chapter 2)

#### **Model Implementation:**

- $\bullet$  Define full  $\Sigma_J$  and prior:  $\Sigma_J^{-1} \sim \mathcal{W}(\Psi, \nu)$
- $\bullet$  Implement  $\Sigma_K \otimes \Sigma_J$  in JAGS

#### Simulation + Validation:

- Simulate individuals with correlated biomarkers
- Fit both block-diagonal and full-covariance models
- Compare fit: DIC, WAIC, predictive checks
- Teunis, Peter F. M., and J. C. H. van Eijkeren. 2016. "Linking the Seroresponse to Infection to Within-Host Heterogeneity in Antibody Production." *Epidemics* 16: 33–39. https://doi.org/10.1016/j.epidem.2016.04.001.