# A model of MRSA transmission incorporating epidemiological and genetic data

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# Observed data (Y)

For each patient i = 1, ..., N admitted to one of the wards in the study period, we denote

- $w_i$  the ward where the patient is admitted (1 for adult ICU, 2 for paediatric ICU)
- $k_i$  the number of times the patient is admitted (1 if no readmission)
- $A_i$  and  $D_i$  vectors containing the times of admission and discharge from the ward
- $P_i$  and  $N_i$  vectors containing the times of positive and negative swabs (positive defined as any of the samples taken is positive; negative defined as all samples taken are negative).
- $p_i$  and  $n_i$  the size of those vectors, ie the number of positive and negative swabs.
- $S_i = \{s_i^1, \dots, s_i^{m_i}\}$  a set of  $m_i$  genetic sequences of MRSA isolated in patient i at times  $T_i = \{t_i^1, \dots, t_i^{m_i}\}$ ; collection dates  $T_i$  are ordered so that  $t_i^k \leq t_i^{k+1}$ .
- $d_{s_i^k, s_j^q}$  the number of transitions between sequence k isolated in patient i and sequence q isolated in patient j.
- $g_{s_i^k, s_j^q}$  the number of transversions between sequence k isolated in patient i and sequence q isolated in patient j.
- $l_{s_i^k, s_i^q}$  the number of typed nucleotides common to sequences  $s_i^k$  and  $s_j^q$ .

# Augmented (unobserved) data (Z)

For each patient i admitted to one of the wards in the study period, we denote

- $C_i$  the colonisation time (we assume no supercolonisations)
- $E_i$  the time of end of colonisation.

We denote  $I_w(t) = \sum_{i=1}^N \mathbf{1}_{\{w_i = w\}} \mathbf{1}_{\{C_i \le t < E_i\}} \sum_{l=1}^{k_i} \mathbf{1}_{\{A_i[l] \le t < D_i[l]\}}$  the number of patients in ward w who are colonized at time t.

## Parameters $(\theta)$

Parameters of the model are:

- $\beta$  a 2 by 2 matrix containing  $\beta_{i\leftarrow j}$ , the person to person transmission rate from ward j to ward i
- $\beta_{\text{ward}\leftarrow\text{out}}$  the force of infection from outside the 2 wards applied to patients in the wards
- $\beta_{\text{out}\leftarrow\text{out}}$  the force of infection from outside the 2 wards applied to patients when they are not in the wards (eg inbetween two admissions)
- Sp the specificity of the testing, ie the probability of getting a negative test given uncolonized (assumed 100%)
- Se the sensitivity of the testing, ie the probability of getting a positive test given colonized
- $\pi$  the probability of being already colonized at first admission
- $\mu$  and  $\sigma$  the mean and standard deviation of the duration of colonization.
- $\nu_1$  and  $\nu_2$  the rate of transitions  $(A \leftrightarrow G \text{ and } C \leftrightarrow T)$  and transversions (other changes) of the DNA sequences. In practice, we will use  $\nu_2 = \kappa \nu_1$  with  $\kappa \in \mathbb{R}_+$ .
- $\alpha$  is the 'within-host pathogenic diversity', defined as the number of pathogenic lineages infecting a given patient; all lineages are supposed to be as likely to be have been sequenced. See description of the genetic likelihood for the definition of a lineage.

### Statistical Model

In the following,  $\mathbf{1}_{\{.\}}$  denotes the indicator function, defined by  $\mathbf{1}_{\{X\}} = 1$  if X is true, and 0 otherwise. The joint density of the observed data, the augmented data, and the model parameters is:

$$P(Y, Z, \theta) = P(Y|Z) P(Z|\theta) P(\theta)$$

where P(Y|Z),  $P(Z|\theta)$  and  $P(\theta)$  refer to the observation level, the transmission level and the prior level respectively.

## Observation level

The observation level ensures that the observed data are consistent with the augmented data:

$$\begin{split} P\left(\boldsymbol{Y}|\boldsymbol{Z}\right) = & & \prod_{i=1}^{N} & \mathbf{1}_{\{C_{i} < E_{i}\}} \\ & & \prod_{j=1}^{p_{i}} \left( \left(\mathbf{1}_{\{P_{i}[j] < C_{i}\}} + \mathbf{1}_{\{P_{i}[j] \ge E_{i}\}}\right) \times (1 - Sp) + \mathbf{1}_{\{C_{i} \le P_{i}[j] < E_{i}\}} \times Se \right) \\ & & \prod_{k=1}^{n_{i}} \left( \left(\mathbf{1}_{\{N_{i}[k] < C_{i}\}} + \mathbf{1}_{\{N_{i}[k] \ge E_{i}\}}\right) \times Sp + \mathbf{1}_{\{C_{i} \le N_{i}[k] < E_{i}\}} \times (1 - Se) \right) \end{split}$$

The first line describes the positive tests, which can be either false positives (first term) or true positives (second term). The second line describes the negative tests, which can be either true negatives (first term) or false negatives (second term).

## Transmission level (discrete time version; time step = half day or day?)

In the discrete version  $A_i[k]$  is the first time step where individual i is in hospital (for his/her  $k_{th}$  stay), and  $D_i[k]$  is the first time step where he/she is out of hospital (after his/her  $k_{th}$  stay). Individual i can transmit staph aureus from time step  $C_i$  to time step  $E_i - 1$ .

$$P\left(\mathbf{Z}|\boldsymbol{\theta}\right) = \prod_{i=1}^{N} \left(\Omega_{i}^{(1)} + \Omega_{i}^{(2)}\right) \times \left(\Phi_{\mu,\sigma}\left(E_{i} - C_{i} + 0.5\right) - \Phi_{\mu,\sigma}\left(E_{i} - C_{i} - 0.5\right)\right)$$

where  $\Phi_{\mu,\sigma}$  is the cumulative density function of a Gamma distribution with mean  $\mu$  and standard deviation  $\sigma$  (we assume that the duration of colonisation is Gamma distributed), and:

$$\Omega_{i}^{(1)} = \pi \times \mathbf{1}_{\{C_{i} < A_{i}[1]\}} 
\Omega_{i}^{(2)} = (1 - \pi) \times \mathbf{1}_{\{C_{i} \ge A_{i}[1]\}} \times e^{-\sum_{t=A_{i}[1]}^{C_{i}-1} \lambda_{i}(t)} \left(1 - e^{-\lambda_{i}(C_{i})}\right) \eta_{i}(C_{i})$$

 $\Omega_i^{(1)}$  is the probability that individual i is colonized before his/her first admission in the wards;  $\Omega_i^{(2)}$  is the probability that individual i is colonized after his/her first admission in the wards.  $\lambda_i(t)$  is the force of transmission applied to individual i at time t. It is equal to:

$$\lambda_{i}^{(t)} = \sum_{w=1}^{2} \beta_{w_{i} \leftarrow w} I_{w}(t) + \beta_{\text{ward} \leftarrow \text{out}} \text{ if individual } i \text{ is in a ward at time } t$$
$$= \beta_{\text{out} \leftarrow \text{out}} \text{ otherwise}$$

 $\eta_i(C_i)$  is the probability of observing sequences  $S_i$  given the possible contamination sources at time  $C_i$ .

It is equal to:

$$\eta_{i}\left(C_{i}\right) = \frac{\sum_{j \text{ colonized and in a ward at time } C_{i}} \beta_{w_{i} \leftarrow w_{j}} f_{i \leftarrow j} + \beta_{\text{ward} \leftarrow \text{out}} f_{i, \text{ward} \leftarrow \text{out}}}{\sum_{j \text{ colonized and in a ward at time } C_{i}} \beta_{w_{i} \leftarrow w_{j}} + \beta_{\text{ward} \leftarrow \text{out}}} \text{ if individual } i \text{ is in a ward at time } C_{i}$$

$$= f_{i, \text{out} \leftarrow \text{out}} \text{ otherwise}}$$

 $f_{i,j}$  is the probability of observing sequences  $S_i$  and  $S_j$  at respective times  $T_i$  and  $T_j$  given that individual j infected individual i (see next section on the genetic likelihood). Similarly,  $f_{i,\text{ward}\leftarrow\text{out}}$  and  $f_{i,\text{out}\leftarrow\text{out}}$  are the probability of observing sequence  $s_i$  at time  $t_i$  given that individual i was infected from outside the wards while he was in or outside hospital respectively.

#### Genetic likelihood

#### Ancestors and lineages

We say that B is an ancestor of A if and only if there is a path leading from B to A in the directed acyclic graph (DAG) representing the genealogy of A. Put simply, B is an ancestor of A if A derives from B.

We will say that B is the *most recent ancestor* (MRA) of A if there is no ancestor of A observed later in the considered set. A *lineage* is defined as a set of (temporally ordered) individuals  $\{x_1, ..., x_n\}$  so that  $x_i$  is the MRA of  $x_{i+1}$  for i=1, ..., n-1. For instance, in the lineage  $(D \to C \to B \to A)$ , B, C, D are all ancestors of A and B is the MRA of A. The genetic likelihood of A given B, C, D will be defined as the probability of the observed mutations between B and A, and is not conditional on previous ancestries. The following statements (and thus the associated probabilities) are equivalent:

- $\bullet$  A and B are from the same lineage and B is older than A
- $\bullet$  B is an ancestor of A
- $\bullet$  A is a descendent of B

#### Genetic likelihood of an infection

The genetic likelihood of the infection of i by j (noted  $i \leftarrow j$ ) relies on how likely it is to observe the genetic differences between sequences in  $S_i$  and their most recent ancestors (MRA) in  $S_j$ . We first focus on the probability of observing a given sequence  $s_i^k$  in i given that j infected i. We note  $\phi(s_i^k, s_j^q)$  the probability of observing  $s_i^k$  given an ancestor  $s_j^k$  (which is also the probability that  $s_j^q$  is an ancestor of  $s_i^k$ ), defined as:

$$\phi(s_i^k, s_j^q) = \mathbf{1}_{\{t_i^k \geq t_j^q\}} \times \underbrace{\mathcal{P}\left(d_{s_i^k, s_j^q} | \nu_1(t_i^k - t_j^q) l_{s_i^k, s_j^q}\right)}_{\text{transitions}} \times \underbrace{\mathcal{P}\left(g_{s_i^k, s_j^q} | \nu_2(t_i^k - t_j^q) l_{s_i^k, s_j^q}\right)}_{\text{transversions}}$$

with:

- $\mathbf{1}_{\{\text{statement}\}}$ : indicator function, 1 if 'statement' is true, 0 otherwise
- $\mathcal{P}(.|\lambda)$ : the probability mass function of a Poisson distribution with parameter  $\lambda$

. The three terms respectively correspond to the indicator function ensuring that  $s_j^q$  is older than  $s_i^k$ , the probability of the observed transitions  $(d_{s_i^k, s_j^q})$ , and the probability of the observed transversions  $(g_{s_i^k, s_j^q})$ .

We are now interested in  $\xi(s_i^k, s_j^q)$ , the probability that the sequence  $s_j^q$  is the MRA of  $s_i^k$ . This requires two elements: i) that  $s_j^q$  is an ancestor of  $s_i^k$ , and ii) that no ancestor of  $s_i^k$  has been collected after  $s_j^q$ . This is given by:

$$\xi(s_i^k, s_j^q) = \underbrace{\phi(s_i^k, s_j^q)}_{s_j^q \text{ ances. of } s_i^k} \times \prod_{r=q+1}^{m_j} \underbrace{(1 - \phi(s_i^k, s_j^r))}_{s_j^r \text{ not ances. of } s_i^k}$$

The genetic likelihood also needs to account for the possibility that, due to the sampling process, no ancestor of  $s_i^k$  may have been isolated and sequenced in  $S_j$ . Assuming that all lineages are as likely to have been sequenced, the probability  $\gamma(s_i^k, S_j)$  that  $S_j$  contains at least one ancestor of  $s_i^k$  is:

$$\gamma(s_i^k, S_j) = 1 - \mathcal{B}\left(0 | \sum_{j=1}^{m_j} \mathbf{1}_{\{t_i^k \ge t_j^q\}}, 1/\alpha\right)$$

with:

- $\mathcal{B}(.|n,p)$ : probability mass function of the Binomial distribution with n draws and probability p
- $\sum_{j=1}^{m_j} \mathbf{1}_{\{t_i^k \geq t_j^q\}}$ : number of isolates sequenced in patient j and collected before the sequence  $s_i^k$

•  $\alpha$ : number of lineages in patient j

The probability  $p(s_i^k|S_j, i \leftarrow j)$  of observing the sequence  $s_i^k$  given that patient j infected patient i can now be computed as:

$$p(s_i^k|S_j, i \leftarrow j) = (\underbrace{\gamma(s_i^k, S_j)}_{\text{ances. in } S_j} \times \underbrace{\sum_{q=1}^{m_j} \xi(s_i^k, s_j^q)}_{\text{prob. MRA for each } S_j}) + \underbrace{1 - \gamma(s_i^k, S_j)}_{\text{ances. not sampled}}$$

The probability of observing the set of sequences  $S_i$  given that j infected i is simply computed as the product over all sequences in  $S_i$ :

$$p(S_i|S_j, i \leftarrow j) = \prod_{k=1}^{m_i} p(s_i^k|S_j, i \leftarrow j)$$

For the sake of simplicity, we shall refer to this quantity as  $f_{i,j}$ .

#### Assumptions of the genetic model

The genetic model makes a few key assumptions:

- different types of mutations happen independently
- all patients share the same number of pathogen lineages  $\alpha$ ; this could be relaxed by using a distribution for  $\alpha$ , e.g by taking  $\alpha \sim 1 + \mathcal{P}(\mu_a)$ .
- all lineages within a host are as likely to have been sampled and sequenced; when lineages have different within-host population sizes, this may still be ensured by extensive swabbing and retaining the sequences of new haplotypes only.

#### Prior level

For all model parameters, independent prior distributions were chosen:

- uniform on [0,1] for Sp, Se, and  $\pi$ ,
- flat exponential (mean 1000) for all other parameters.

#### Parameter Estimation

A Markov chain Monte Carlo (MCMC) method was used to sample the joint posterior distribution  $P(\mathbf{Y}, \mathbf{Z}, \boldsymbol{\theta})$ . Sp, Se and  $\pi$  were updated using the Gibbs sampler, and all other parameters using a Metropolis algorithm.

# Appendix: Full formulation of the transmission level

In the discrete version  $A_i[k]$  is the first time step where individual i is in hospital (for his/her  $k_{th}$  stay), and  $D_i[k]$  is the first time step where he/she is out of hospital (after his/her  $k_{th}$  stay). Individual i can transmit staph aureus from time step  $C_i$  to time step  $E_i - 1$ .

$$P(\mathbf{Z}|\boldsymbol{\theta}) = \prod_{i=1}^{N} \left( \Omega_{i}^{(1)} + \Omega_{i}^{(2)} + \Omega_{i}^{(3)} \right) \times \left( \Phi_{\mu,\sigma} \left( E_{i} - C_{i} + 0.5 \right) - \Phi_{\mu,\sigma} \left( E_{i} - C_{i} - 0.5 \right) \right)$$

where  $\Phi_{\mu,\sigma}$  is the cumulative density function of a Gamma distribution with mean  $\mu$  and standard deviation  $\sigma$  (we assume that the duration of colonisation is Gamma distributed), and:

$$\begin{split} &\Omega_{i}^{(1)} = \pi \times \mathbf{1}_{\{C_{i} < A_{i}[1]\}} \\ &\Omega_{i}^{(2)} = (1 - \pi) \sum_{l=1}^{k} \mathbf{1}_{\{A_{i}[l] \leq C_{i} < D_{i}[l]\}} \\ &\times \exp\left(-\mathbf{1}_{\{l \geq 2\}} \sum_{s=1}^{l-1} \left[\sum_{t=A_{i}[s]}^{D_{i}[s]-1} \left(\sum_{w=1}^{2} \beta_{w_{i} \leftarrow w} I_{w}\left(t\right)\right) + \beta_{\text{ward} \leftarrow \text{out}}\left(D_{i}[s] - A_{i}[s]\right)\right]\right) \\ &\times \exp\left(-\mathbf{1}_{\{l \geq 2\}} \sum_{s=1}^{l-1} \left[\beta_{\text{out} \leftarrow \text{out}}\left(A_{i}[s+1] - D_{i}[s]\right)\right]\right) \\ &\times \exp\left(-\sum_{t=A_{i}[l]}^{C_{i}-1} \left(\sum_{w=1}^{2} \beta_{w_{i} \leftarrow w} I_{w}\left(t\right)\right) - \beta_{\text{ward} \leftarrow \text{out}}\left(C_{i} - A_{i}[l]\right)\right) \\ &\times \left(1 - \exp\left(-\sum_{w=1}^{2} \beta_{w_{i} \leftarrow w} I_{w}\left(C_{i}\right) - \beta_{\text{ward} \leftarrow \text{out}}\right)\right) \\ &\times \frac{\sum_{j=1}^{N} \mathbf{1}_{\{C_{j} \leq C_{i} < E_{j}\}} \sum_{r=1}^{k_{j}} \mathbf{1}_{\{A_{j}[r] \leq C_{i} < D_{j}[r]\}} \beta_{w_{i} \leftarrow w_{j}} + \beta_{\text{ward} \leftarrow \text{out}} f_{i,\text{ward} \leftarrow \text{out}}}{\sum_{j=1}^{N} \mathbf{1}_{\{C_{j} \leq C_{i} < E_{j}\}} \sum_{r=1}^{k_{j}} \mathbf{1}_{\{A_{j}[r] \leq C_{i} < D_{j}[r]\}} \beta_{w_{i} \leftarrow w_{j}} + \beta_{\text{ward} \leftarrow \text{out}}}{\Omega_{i}^{(3)} = \mathbf{1}_{\{k_{i} > 1\}} \left(1 - \pi\right) \sum_{l=1}^{k_{i}-1} \mathbf{1}_{\{D_{i}[l] \leq C_{i} < A_{i}[l+1]\}} \\ &\times \exp\left(-\sum_{s=1}^{l} \left[\sum_{t=A_{i}[s]}^{D_{i}[s]-1} \left(\sum_{w=1}^{2} \beta_{w_{i} \leftarrow w} I_{w}\left(t\right)\right) + \beta_{\text{ward} \leftarrow \text{out}}\left(D_{i}[s] - A_{i}[s]\right)\right]\right) \\ &\times \exp\left(-\mathbf{1}_{\{l \geq 2\}} \sum_{s=1}^{l-1} \left[\beta_{\text{out} \leftarrow \text{out}}\left(A_{i}[s+1] - D_{i}[s]\right)\right]\right) \\ &\times \exp\left(-\beta_{\text{out} \leftarrow \text{out}}\left(C_{i} - D_{i}[l]\right)\right) \\ &\times \left(1 - \exp\left(-\beta_{\text{out} \leftarrow \text{out}}\right)\right) \\ &\times$$

 $\Omega_i^{(1)}$  is the probability that individual i is colonized before his/her first admission in the wards;  $\Omega_i^{(2)}$  is the probability that individual i is colonized during one of his/her stays in the wards;  $\Omega_i^{(3)}$  is the probability, that individual i, if admitted several times, is colonized between successive stays in the wards.

 $f_{i,j}$  is the probability of observing sequences  $s_i$  and  $s_j$  at respective time steps  $t_i$  and  $t_j$  given that individual j infected individual i (see next section on the genetic likelihood). Similarly,  $f_{i,\text{ward}\leftarrow\text{out}}$  and  $f_{i,\text{out}\leftarrow\text{out}}$  are the probability of observing sequence  $s_i$  at time  $t_i$  given that individual i was infected from outside the wards while he was in or outside hospital respectively.