# Appendix to "Integrated nested Laplace approximations for extended latent Gaussian models with application to the Naomi HIV model" Corresponding author: Adam Howes (ath19@ic.ac.uk)

# Contents

1	Modified Naomi model description	<b>2</b>
	1.1 Notation	2
	1.2 Process specification	2
	1.3 Likelihood specification	
	1.4 Identifiability constaints	6
2	MCMC convergence and suitability	7
3	C++ template	8
$\mathbf{R}$	eferences	20

# 1 Modified Naomi model description

#### 1.1 Notation

#### 1.1.1 Indexing

Consider a given country in sub-Saharan Africa. Let x refer to a district located within the Spectrum region  $R_x$ . We restrict our attention to the time period  $t=T_1$  of the most recent national household survey with HIV testing. Let  $s \in \{F, M\}$  be sex, and  $a \in \{0\text{-}5, 5\text{-}10, \dots, 75\text{-}80, 80+\}$  be five-year age bands. As short-hand, we write a=l to refer to the age band with lower bound l, e.g. a=20 for a=20-25. We index the following quantities by district, sex and age-band: population size  $N_{x,s,a}$ , HIV prevalence  $\rho_{x,s,a}$ , ART coverage  $\alpha_{x,s,a}$ , annual HIV incidence rate  $\lambda_{x,s,a}$ , and proportion of HIV positive persons recently infected  $\kappa_{x,s,a}$ . Sometimes data are observed at an aggregate level, rather than the modelled level of granularity. In this instance, we use  $\{\cdot\}$  to generically refer to a aggregate set over which an observation is made, e.g.  $\{a\} = \{15\text{-}19, \dots, 45\text{-}49\}$  for adults.

#### 1.1.2 Structured random effects

Consider a generic random effect u. We use use structured random effects to share information across units assessed as being similar, such as neighbouring districts or adjacent age-bands. We use  $u \sim \text{ICAR}(\sigma)$  to refer to the Besag intrinsic conditional auto-regressive model (ICAR) (Besag, York, and Mollié 1991) with full conditionals

$$u_i \mid u_{-i} \sim \mathcal{N}\left(\frac{\sum_{j:j \sim i} u_j}{n_{\delta i}}, \frac{\sigma^2}{n_{\delta i}}\right),$$

where  $u_{-i}$  is u with the ith unit removed,  $j \sim i$  if the units are defined as adjacent,  $n_{\delta i} = |\{j : j \sim i\}|$  is the total number of adjacent units, and  $\sigma > 0$  is the marginal standard deviation. We follow recommendations of Freni-Sterrantino, Ventrucci, and Rue (2018) on scaling of precision matrices, disconnected adjacency graph components, and islands regarding the ICAR model. For the reparameterised Besag-York-Mollie model (BYM2) (Simpson et al. 2017) we write  $u \sim \text{BYM2}(\sigma, \phi)$ , where u is comprised of a spatially structured ICAR component  $v^*$  with proportion  $\phi \in (0, 1)$  and spatially unstructured IID component  $w^*$  with proportion  $1 - \phi$ , both scaled to have generalised variance equal to one, and  $\sigma > 0$  is the marginal standard deviation such that

$$u = \sigma \left( \sqrt{\phi} \cdot v^* + \sqrt{1 - \phi} \cdot w^* \right).$$

We specify the first order auto-regressive model by  $u \sim AR1(\sigma, \phi)$  such that

$$u_1 \sim \left(0, \frac{1}{1 - \rho^2}\right),$$
  
 $u_i = \rho u_{i-1} + \epsilon_t, \quad i = 2, \dots$ 

where  $\epsilon_i \sim \mathcal{N}(0,1)$  is Gaussian white noise,  $|\rho| < 1$  is the lag-one correlation parameter.

### 1.2 Process specification

#### 1.2.1 HIV prevalence

We model HIV prevalence on the logit scale using

$$logit(\rho_{x,s,a}) = \beta_0^{\rho} + \beta_S^{\rho,s=M} + u_a^{\rho} + u_a^{\rho,s=M} + u_x^{\rho} + u_x^{\rho,s=M} + u_x^{\rho,s=M} + u_x^{\rho,a<15} + \eta_{R_x,s,a}^{\rho}$$

where  $\beta_0^{\rho}$  is an intercept term,  $\beta_s^{\rho,s=\mathrm{M}}$  is the difference in logit prevalence for men compared to women,  $u_a^{\rho} \sim \mathrm{AR1}(\sigma_A^{\rho},\phi_A^{\rho})$  are age random effects for women,  $u_a^{\rho,s=\mathrm{M}} \sim \mathrm{UNKNOWN}$  are age random effects for the difference in logit prevalence for men compared to women age  $a, u_x^{\rho} \sim \mathrm{BYM2}(\sigma_X^{\rho},\phi_X^{\rho})$  are spatial random effects for women,  $u_x^{\rho,s=\mathrm{M}} \sim \mathrm{BYM2}(\sigma_{XS}^{\rho},\phi_{XS}^{\rho})$  are spatial random effects for the difference in logit prevalence for men compared to women in district  $x, u_x^{\rho,a<15} \sim \mathrm{ICAR}(0,\sigma_{XA}^{\rho})$  are spatial random effects for the ratio of paediatric prevalence to adult women prevalence, and  $\eta_{R_x,s,a}^{\rho}$  are fixed offsets specifying assumed odds ratios

for prevalence outside the age ranges for which data are available. We use the prior distributions  $\mathcal{N}(0,5)$  for the fixed effects  $\beta_0^{\rho}$  and  $\beta_s^{\rho,s=M}$ ,  $\mathcal{N}^+(0,2.5)$  for all standard deviation terms,  $\mathcal{U}(-1,1)$  for all AR1 correlation parameters, and Beta(0.5,0.5) for all BYM2 proportion parameters.

#### 1.2.2 ART coverage

We model ART coverage by

$$logit(\alpha_{x,s,a}) = \beta_0^{\alpha} + \beta_S^{\alpha,s=M} + u_a^{\alpha} + u_a^{\alpha,s=M} + u_x^{\alpha} + u_x^{\alpha,s=M} + u_x^{\alpha,a<15} + \eta_{R_r,s,a}^{\alpha}$$

with terms and priors analogous to the HIV prevalence model above.

#### 1.2.3 HIV incidence rate

We model HIV incidence rate by

$$\log(\lambda_{x,s,a}) = \beta_0^{\lambda} + \beta_S^{\lambda,s=\mathrm{M}} + \log(\rho_x^{15\text{-}49}) + \log(1 - \omega \cdot \alpha_x^{15\text{-}49}) + u_x^{\lambda} + \eta_{R_x,s,a}^{\lambda}$$

where  $\beta_0^{\lambda}$  is an intercept term proportional to the average HIV transmission rate for untreated HIV positive adults,  $\beta_S^{\lambda,s=\mathrm{M}}$  is the log incidence rate ratio for men compared to women,  $\rho_x^{15\text{-}49}$  is the HIV prevalence among adults 15-49 calculated by

$$\rho_x^{15\text{-}49} = \frac{\sum_{s \in \{F,M\}} \sum_{a=15}^{45} N_{x,s,a} \cdot \rho_{x,s,a}}{\sum_{s \in \{F,M\}} \sum_{a=15}^{45} N_{x,s,a}},$$

 $\alpha_x^{15-49}$  is the ART coverage among adults 15-49 calculated by

$$\alpha_x^{15\text{-}49} = \frac{\sum_{s \in \{F,M\}} \sum_{a=15}^{45} N_{x,s,a} \cdot \rho_{x,s,a} \cdot \alpha_{x,s,a}}{\sum_{s \in \{F,M\}} \sum_{a=15}^{45} N_{x,s,a} \cdot \rho_{x,s,a}},$$

 $\omega$  is the average reduction in HIV transmission rate per 1% increase in population ART coverage and is fixed at  $\omega = 0.7$  via the EPP model,  $u_x^{\lambda} \sim \mathcal{N}(0, \sigma^{\lambda})$  with  $\sigma^{\lambda} \sim \mathcal{N}^+(0, 1)$  are IID spatial random effects, and  $\eta_{R_x, s, a}^{\lambda}$  specify log incidence rate ratios by sex and age group calculated from Spectrum model output

#### 1.2.4 ANC testing

HIV prevalence  $\rho_{x,a}^{\mathrm{ANC}}$  and ART coverage  $\alpha_{x,a}^{\mathrm{ANC}}$  among pregnant women modelled with logit-linear models

$$\operatorname{logit}(\rho_{x,a}^{\mathrm{ANC}}) = \operatorname{logit}(\rho_{x,F,a}) + \beta^{\rho^{\mathrm{ANC}}} + u_x^{\rho^{\mathrm{ANC}}} + \eta_{R_x,a}^{\rho^{\mathrm{ANC}}}$$
 (1)

$$\operatorname{logit}(\alpha_{x,a}^{\mathrm{ANC}}) = \operatorname{logit}(\alpha_{x,F,a}) + \beta^{\alpha^{\mathrm{ANC}}} + u_x^{\alpha^{\mathrm{ANC}}} + \eta_{R_x,a}^{\alpha^{\mathrm{ANC}}}, \tag{2}$$

where  $\eta_{R_x,a}^{\theta^{\text{ANC}}}$  for  $\theta \in \{\rho,\alpha\}$  are offsets for the log fertility rate ratios for HIV positive women compared to HIV negative women and for women on ART to HIV positive women not on ART, calculated from Spectrum model outputs for region  $R_x$ ,  $\beta^{\theta^{\text{ANC}}}$  for  $\theta \in \{\rho,\alpha\}$  are the average differences between population and ANC outcomes after removing the offset, and  $u_x^{\theta^{\text{ANC}}} \sim \mathcal{N}(0,\sigma_X^{\theta^{\text{ANC}}})$  for  $\theta \in \{\rho,\alpha\}$  are district random effects.

#### 1.2.5 ART attendance

Let  $\gamma_{x,x'} \in [0,1]$  be the probability that a person on ART residing in district x recieves ART in district x'. We assume that  $\gamma_{x,x'} = 0$  for  $x \notin \{x, \text{ne}(x)\}$  such that individuals seek treatment only in their residing district and its neighbours  $\text{ne}(x) = \{x' : x' \sim x\}$ , where  $\sim$  is an adjacency relation and  $\sum_{x' \in \{x, \text{ne}(x)\}} \gamma_{x,x'} = 1$ .

To model  $\gamma_{x,x'}$  for  $x \sim x'$  we use a multinomial logistic regression model where  $\tilde{\gamma}_{x,x'}$  is the log odds ratio of seeking ART

$$\tilde{\gamma}_{x,x'} = \tilde{\gamma}_0 + u_x^{\tilde{\gamma}} \tag{3}$$

$$\tilde{\gamma}_0 = -4 \tag{4}$$

$$u_x^{\tilde{\gamma}} \sim \mathcal{N}(0, \sigma_X^{\tilde{\gamma}})$$
 (5)

where  $\tilde{\gamma}_0$  is an intercept specifying the prior mean for the log odds of seeking ART in each neighbouring district compared to the home district, and  $u_x^{\tilde{\gamma}}$  are district random effects. We fix  $\tilde{\gamma}_{x,x} = 0$  and recover the multinomial probabilities using the softmax

$$\gamma_{x,x'} = \frac{\exp(\tilde{\gamma}_{x,x'})}{\sum_{x^* \in \{x, \text{ne}(x)\}} \exp(\tilde{\gamma}_{x,x^*})}$$

Then, given the number of PLHIV on ART  $A_{x,s,a} = N_{x,s,a} \cdot \rho_{x,s,a} \cdot \alpha_{x,s,a}$ , the number of ART clients who reside in district x and obtain ART in district x' are

$$A_{x,x',s,a} = A_{x,s,a} \cdot \gamma_{x,x'},$$

and the total attending ART facilities in district x' are

$$\tilde{A}_{x',s,a} = \sum_{x \in \{x', \text{ne}(x')\}} A_{x,x',s,a}.$$

## 1.3 Likelihood specification

## 1.3.1 Household survey data

Let  $\nu$  be a household survey occurring at  $T_1$ , furnishing weighted observations  $\hat{\theta}_{\{x\},\{s\},\{a\},\nu}$  for  $\theta \in \{\rho,\alpha,\kappa\}$  with respective Kish effective sample sizes  $M^{\hat{\theta}}_{\{x\},\{s\},\{a\},\nu}$ , and observed number of cases

$$Y_{\{x\},\{s\},\{a\},\nu}^{\hat{\theta}} = M_{\{x\},\{s\},\{a\},\nu}^{\hat{\theta}} \cdot \hat{\theta}_{\{x\},\{s\},\{a\},\nu}.$$

For HIV prevalence and ART coverage we use binomial working likelihoods as follows

$$Y_{\{x\},\{s\},\{a\},\nu}^{\hat{\rho}} \sim x \text{Bin}(M_{\{x\},\{s\},\{a\},\nu}^{\hat{\rho}}, \rho_{\{x\},\{s\},\{a\},T_1})$$
(6)

$$\rho_{\{x\},\{s\},\{a\}} = \frac{\sum_{x \in \{x\}} \sum_{s \in \{s\}} \sum_{a \in \{a\}} N_{x,s,a} \cdot \rho_{x,s,a}}{\sum_{x \in \{x\}} \sum_{s \in \{s\}} \sum_{a \in \{a\}} N_{x,s,a}},$$
(7)

and

$$Y_{\{x\},\{s\},\{a\},\nu}^{\hat{\alpha}} \sim x \text{Bin}(M_{\{x\},\{s\},\{a\},\nu}^{\hat{\alpha}}, \alpha_{\{x\},\{s\},\{a\},T_1})$$
(8)

$$\alpha_{\{x\},\{s\},\{a\}} = \frac{\sum_{x \in \{x\}} \sum_{s \in \{s\}} \sum_{a \in \{a\}} N_{x,s,a} \cdot \rho_{x,s,a} \cdot \alpha_{x,s,a}}{\sum_{x \in \{x\}} \sum_{s \in \{s\}} \sum_{a \in \{a\}} N_{x,s,a} \cdot \rho_{x,s,a}}.$$
(9)

For recent infections we also use a binomial working likelihood

$$Y_{\{x\},\{s\},\{a\},\nu}^{\hat{\kappa}} \sim x \text{Bin}(M_{\{x\},\{s\},\{a\},\nu}^{\hat{\kappa}}, \kappa_{\{x\},\{s\},\{a\},T_1})$$
(10)

$$\kappa_{x,s,a} = 1 - \exp(-\lambda_{x,s,a} \cdot \frac{1 - \rho_{x,s,a}}{\rho_{x,s,a}} \cdot (\Omega_T - \beta_T) - \beta_T)$$
(11)

$$\Omega_T \sim \mathcal{N}(\Omega_{T_0}, \sigma^{\Omega_T}) \tag{12}$$

$$\beta_T \sim \mathcal{N}(\beta_{T_0}, \sigma^{\beta_T}) \tag{13}$$

where  $\kappa_{x,s,a}$  are the predicted proportion recently infected among HIV positive persons,  $\Omega_T$  is the mean duration of recent infection (MDRI), and  $\beta_T$  is the false recent ratio (FRR). We use an informative prior on  $\Omega_T$  based on the characteristics of the recent infection testing algorithm (RITA). For PHIA surveys this is  $\Omega_{T_0} = 130$  days and  $\sigma^{\Omega_T} = 6.12$  days. For PHIA surveys we assume there is no false recency, such that  $\beta_{T_0} = 0.0$  and  $\sigma^{\beta_T} = 0.0$ .

#### 1.3.2 ANC testing data

For women 15-49 the predicted number of ANC clients  $\Psi_{x,a}$  is a log-linear model

$$\log(\Psi_{x,a}) = \log(N_{x,F,a}) + \psi_{R_x,a} + \beta^{\psi} + u_x^{\psi}$$

where  $N_{x,\mathrm{F},a}$  are the female population sizes,  $\psi_{R_x,a}$  are ASFR in Spectrum region  $R_x$  at time t,  $\beta^{\psi}$  are the log rate ratio for the number of ANC clients relative to the predicted fertility,  $u_x^{\psi} \sim \mathcal{N}(0, \sigma^{\psi})$  are district random effects.

We include ANC testing data for the year of the most recent survey  $Y[T_1]$ . Let  $W_{\{x\},Y[t]}^{\text{ANC}}$  be the number of ANC clients,  $X_{\{x\},Y[t]}^{\text{ANC}}$  the number of those with ascertained status,  $Y_{\{x\},Y[t]}^{\text{ANC}}$  the number of those with positive status (either known or tested) and  $Z_{\{x\},Y[t]}^{\text{ANC}}$  the number of ANC clients already on ART prior to first ANC, such that  $W_{x,Y[t]}^{\text{ANC}} \geq X_{x,Y[t]}^{\text{ANC}} \geq Y_{x,Y[t]}^{\text{ANC}} \geq Z_{x,Y[t]}^{\text{ANC}}$  When ANC testing data are only available for part of a given year, we denote  $M_{Y[t]}^{\text{ANC}} \in \{1, \dots, 12\}$  the number of months of reported data reflected in counts for year Y[t].

The observed number of HIV positive and already on ART among ANC clients at  $Y[T_1]$  is modelled by

$$\begin{split} &Y_{\{x\}Y[T_1]}^{\text{ANC}} \sim \text{Bin}\left(X_{\{x\}Y[T_1]}^{\text{ANC}}, \rho_{\{x\},\{15,\dots45\}}^{\text{ANC}}\right) \\ &Z_{\{x\}Y[T_1]}^{\text{ANC}} \sim \text{Bin}\left(Y_{\{x\}Y[T_1]}^{\text{ANC}}, \alpha_{\{x\},\{15,\dots45\}}^{\text{ANC}}\right) \end{split}$$

where predicted prevalence and ART coverage are aggregated weighted by the predicted number of pregnant women by age  $\Psi_{x,a}$ 

$$\rho_{\{x\}\{a\}}^{\text{ANC}} = \frac{\sum_{x \in \{x\}} \sum_{a \in \{a\}} \Psi_{x,a} \cdot \rho_{x,a}^{\text{ANC}}}{\sum_{x \in \{x\}} \sum_{a \in \{a\}} \Psi_{x,a}}$$

$$\alpha_{\{x\}\{a\}}^{\text{ANC}} = \frac{\sum_{x \in \{x\}} \sum_{a \in \{a\}} \Psi_{x,a} \cdot \rho_{x,a}^{\text{ANC}} \cdot \alpha_{x,a}^{\text{ANC}}}{\sum_{x \in \{x\}} \sum_{a \in \{a\}} \Psi_{x,a} \cdot \rho_{x,a}^{\text{ANC}}}$$

#### 1.3.3 Number receiving ART

Let  $\dot{A}_{\{x\},\{s\},\{a\}}$  be data for the number receiving ART

$$\dot{A}_{\{x\},\{s\},\{a\}} = \sum_{s \in \{s\}} \sum_{a \in \{a\}} \sum_{x \in \{x\}} \sum_{x \sim x', x = x'} \dot{A}_{x',x,s,a},$$

We model the unobserved numbers of ART clients travelling from x' to x as

$$\dot{A}_{x',x,s,a} \sim \operatorname{Bin}(N_{x',s,a}, \pi_{x',x,s,a})$$

where  $\pi_{x',x,s,a} = \rho_{x',s,a} \cdot \alpha_{x',s,a} \cdot \gamma_{x',x,s,a}$ . This likelihood is approximated using a normal for the sum of binomials by

$$\dot{A}_{\{x\},\{s\},\{a\}} \sim \mathcal{N}(\tilde{A}_{\{x\},\{s\},\{a\}},\sigma_{\{x\},\{s\},\{a\}}^{\tilde{A}})$$

where

$$\tilde{A}_{\{x\},\{s\}\{a\}} = \sum_{s \in \{s\}} \sum_{a \in \{a\}} \sum_{x \in \{x\}} \sum_{x \sim x', x = x'} N_{x',s,a} \cdot \pi_{x',x,s,a},$$

and

$$\sigma_{\{x\},\{s\}\{a\}}^{\tilde{A}} = \sqrt{\sum_{s \in \{s\}} \sum_{a \in \{a\}} \sum_{x \in \{x\}} \sum_{x \sim x', x = x'} N_{x',s,a} \cdot \pi_{x',x,s,a} \cdot (1 - \pi_{x',x,s,a})}.$$

## 1.4 Identifiability constaints

If data are missing, some parameters are fixed to default values to help with identifiability. In particular:

- If survey data on ART coverage by age and sex are not available then we set  $u_a^{\alpha} = 0$  and  $u_{a,s=M}^{\alpha} = 0$  and use the average age/sex pattern of ART coverage from the Spectrum offset  $\eta_{R_x,s,a}^{\alpha}$ .
- If no ART data (survey or ART programme) are available at  $T_1$  but data on ART coverage among ANC clients are available, the level of ART coverage is not identifiable, but spatial variation is identifiable. In this instance, overall ART coverage is determined by the Spectrum offset, and only area random effects are estimated

$$logit (\alpha_{x,s,a}) = u_x^{\alpha} + \eta_{R_x,s,a}^{\alpha}$$

• If survey data on recent HIV infection are not included in the model, then  $\beta_0^{\lambda} = \beta_S^{\lambda,s=M} = u_x^{\lambda} = 0$ . The sex ratio for HIV incidence is determined by the sex incidence rate ratio from Spectrum in the same years and the incidence rate in all districts is modelled assuming the same average HIV transmission rate for untreated adults, but varies according to district estimates of HIV prevalence and ART coverage.

# 2 MCMC convergence and suitability

We assessed MCMC convergence and suitability using a range of graphical and numerical tests. These included the potential scale reduction factor  $\hat{R}$ , bulk and tail effective sample size (ESS), autocorrelation decay plots, univariate traceplots, pairs density plots, and NUTS specific divergent transition and energy assessments.

For the time being, this analysis is available from athows.github.io/elgm-inf/mcmc-convergence. Once the MCMC results are finalised, the analysis will be moved to this appendix and expanded upon. The following draft text and references may be useful in that expanded write-up.

- Improved  $\hat{R}$  statistic from Vehtari et al. (2021). Recommended only to use the sample if the value is less than 1.05.
- ESS and autocorrelation related to efficiency
- Traceplots helpful for diagnosis of problems
- Pairs density plots helpful for understanding relationships between parameters, including possible indentifiability issues
- Energy plot from Betancourt (2017)

# 3 C++ template

```
// #define TMB_LIB_INIT R_init_naomi_simple
#include <TMB.hpp>
/** Log posterior density of BYM2 with INLA conditional parameterisation
 * Calculate the joint LPDF of parameter vector (x, u) where
 *x = sigma * (sqrt(phi) * u + sqrt(1-phi) * v) with u a ICAR structured
 * component u \sim N(0, Q^{-1}) and v is an IID effect v \sim N(0, 1). Calculation
 * proceeds by conditioning P(\langle x, u \rangle) = P(x \mid u) * P(u). See Reibler et al.
 * Section 3.4.
 * Oparam x vector of random effects.
 * Oparam u vector of spatial component of random effect.
 * Oparam sigma marginal standard deviation (>0).
 * Oparam phi proportion of marginal variance explained by spatial structured
              component u (phi \in [0, 1]).
 * Oparam Q scaled structure matrix for spatial component.
 * Creturn Log probability density of x and u.
 * Onote
 * The \ \sqrt(2\pi)^{-2*n}$ and $|Q|^{1/2}$ terms are dropped.
 * Returns the _positive_ log PDF (different from builtin TMB
 * functions. Thus should typically be implemented as `nll -= bym2_conditional_lpdf(...)`.
template < class Type >
Type bym2_conditional_lpdf(const vector<Type> x,
                           const vector<Type> u,
                           const Type sigma,
                           const Type phi,
                           const Eigen::SparseMatrix<Type> Q) {
  Type val(0.0);
  // constant terms omitted: -0.5 * (n + rank(Q)) * log(2*pi) + 0.5 * log(Q)
  val += -0.5 * x.size() * (2 * log(sigma) + log(1 - phi)); // normalising constant
  val += -0.5 / (sigma * sigma * (1 - phi)) * (x * x).sum();
  val += sqrt(phi) / (sigma * (1 - phi)) * (x * u).sum();
  val += -0.5 * (u * (Q * u)).sum();
  val += -0.5 * phi / (1 - phi) * (u * u).sum();
  return(val);
template<class Type>
Type objective_function<Type>::operator() ()
  // indexing:
```

```
// * rho: HIV prevalence model
// * alpha: ART coverage model
// * lambda: HIV incidence model
// * _x: area
// * _a: age
// * _s: sex
// * t: time
using namespace density;
// ** Data **
// Population
DATA_VECTOR(population_t1);
// Design matrices
DATA_MATRIX(X_rho);
DATA_MATRIX(X_alpha);
DATA_MATRIX(X_lambda);
DATA_MATRIX(X_ancrho);
DATA_MATRIX(X_ancalpha);
DATA SPARSE MATRIX(Z rho x);
DATA_SPARSE_MATRIX(Z_rho_xs);
DATA_SPARSE_MATRIX(Z_rho_a);
DATA_SPARSE_MATRIX(Z_rho_as);
DATA_SPARSE_MATRIX(Z_rho_xa);
DATA_SPARSE_MATRIX(Z_alpha_x);
DATA_SPARSE_MATRIX(Z_alpha_xs);
DATA_SPARSE_MATRIX(Z_alpha_a);
DATA_SPARSE_MATRIX(Z_alpha_as);
DATA_SPARSE_MATRIX(Z_alpha_xa);
DATA_SPARSE_MATRIX(Z_x);
DATA_SPARSE_MATRIX(Z_lambda_x);
DATA_VECTOR(logit_rho_offset);
DATA_VECTOR(logit_alpha_offset);
DATA_VECTOR(log_asfr_t1_offset);
DATA_VECTOR(logit_anc_rho_t1_offset);
DATA_VECTOR(logit_anc_alpha_t1_offset);
DATA_SPARSE_MATRIX(Z_ancrho_x);
DATA_SPARSE_MATRIX(Z_ancalpha_x);
// Precision matrix for ICAR area model
```

```
DATA_SPARSE_MATRIX(Q_x);
DATA_SCALAR(Q_x_rankdef);
DATA VECTOR(n prev);
DATA_VECTOR(x_prev);
DATA_SPARSE_MATRIX(A_prev);
DATA VECTOR(n artcov);
DATA VECTOR(x artcov);
DATA SPARSE MATRIX(A artcov);
DATA_VECTOR(n_recent);
DATA_VECTOR(x_recent);
DATA_SPARSE_MATRIX(A_recent);
DATA_VECTOR(n_anc_prev_t1);
DATA_VECTOR(x_anc_prev_t1);
DATA_SPARSE_MATRIX(A_anc_prev_t1);
DATA_VECTOR(n_anc_artcov_t1);
DATA VECTOR(x anc artcov t1);
DATA_SPARSE_MATRIX(A_anc_artcov_t1);
DATA_SPARSE_MATRIX(A_artattend_t1);
DATA VECTOR(x artnum t1);
DATA SPARSE MATRIX(A artattend mf);
DATA_SPARSE_MATRIX(A_art_reside_attend);
DATA_IVECTOR(n_nb);
DATA_IVECTOR(adj_i);
DATA_IVECTOR(adj_j);
DATA_SPARSE_MATRIX(Xgamma);
DATA_VECTOR(log_gamma_offset);
DATA SPARSE MATRIX(Xart idx);
DATA_SPARSE_MATRIX(Xart_gamma);
// Incidence model
DATA_SCALAR(omega);
DATA SCALAR (OmegaTO);
DATA SCALAR(sigma OmegaT);
DATA_SCALAR(betaT0);
DATA_SCALAR(sigma_betaT);
DATA_SCALAR(ritaT);
DATA_SPARSE_MATRIX(X_15to49);
DATA_VECTOR(log_lambda_t1_offset);
// Paediatric prevalence and incidence ratio model
```

```
DATA_SPARSE_MATRIX(X_15to49f);
DATA_SPARSE_MATRIX(X_paed_rho_ratio);
DATA_VECTOR(paed_rho_ratio_offset);
DATA_SPARSE_MATRIX(X_paed_lambda_ratio_t1);
// ** Initialize nll **
Type val(0);
// ** Parameters **
// fixed effects
// diffuse N(0.0, 5.0) prior distribution
PARAMETER_VECTOR(beta_rho);
val -= dnorm(beta_rho, 0.0, 5.0, true).sum();
PARAMETER_VECTOR(beta_alpha);
val -= dnorm(beta_alpha, 0.0, 5.0, true).sum();
PARAMETER VECTOR(beta lambda);
val -= dnorm(beta_lambda, 0.0, 5.0, true).sum();
PARAMETER_VECTOR(beta_anc_rho);
val -= dnorm(beta anc rho, 0.0, 5.0, true).sum();
PARAMETER VECTOR (beta anc alpha);
val -= dnorm(beta_anc_alpha, 0.0, 5.0, true).sum();
// * HIV prevalence model *
// hyper parameters
PARAMETER(logit_phi_rho_x);
Type phi_rho_x(invlogit(logit_phi_rho_x));
val -= log(phi_rho_x) + log(1 - phi_rho_x); // change of variables: logit_phi_x ->v phi_x
val -= dbeta(phi_rho_x, Type(0.5), Type(0.5), true);
PARAMETER(log_sigma_rho_x);
Type sigma_rho_x(exp(log_sigma_rho_x));
val -= dnorm(sigma_rho_x, Type(0.0), Type(2.5), true) + log_sigma_rho_x;
PARAMETER(logit_phi_rho_xs);
Type phi_rho_xs(invlogit(logit_phi_rho_xs));
val -= log(phi_rho_xs) + log(1 - phi_rho_xs); // change of variables: logit_phi_xs -> phi_xs
val -= dbeta(phi_rho_xs, Type(0.5), Type(0.5), true);
PARAMETER(log_sigma_rho_xs);
Type sigma_rho_xs(exp(log_sigma_rho_xs));
val -= dnorm(sigma_rho_xs, Type(0.0), Type(2.5), true) + log_sigma_rho_xs;
PARAMETER(logit_phi_rho_a);
val -= dnorm(logit_phi_rho_a, Type(0.0), Type(2.582), true); // INLA default
```

```
Type phi_rho_a(2.0 * invlogit(logit_phi_rho_a) - 1.0);
PARAMETER(log_sigma_rho_a);
Type sigma rho a(exp(log sigma rho a));
val -= dnorm(sigma_rho_a, Type(0.0), Type(2.5), true) + log_sigma_rho_a;
PARAMETER(logit_phi_rho_as);
val -= dnorm(logit_phi_rho_as, Type(0.0), Type(2.582), true); // INLA default
Type phi_rho_as(2.0 * invlogit(logit_phi_rho_as) - 1.0);
PARAMETER(log_sigma_rho_as);
Type sigma_rho_as(exp(log_sigma_rho_as));
val -= dnorm(sigma_rho_as, Type(0.0), Type(2.5), true) + log_sigma_rho_as;
PARAMETER(log_sigma_rho_xa);
Type sigma_rho_xa(exp(log_sigma_rho_xa));
val -= dnorm(sigma_rho_xa, Type(0.0), Type(0.5), true) + log_sigma_rho_xa;
// latent effects
PARAMETER VECTOR(u rho x);
PARAMETER_VECTOR(us_rho_x);
val -= dnorm(sum(us_rho_x), Type(0.0), Type(0.001) * us_rho_x.size(), true); // soft sum-to-zero cons
val -= bym2_conditional_lpdf(u_rho_x, us_rho_x, sigma_rho_x, phi_rho_x, Q_x);
PARAMETER VECTOR(u rho xs);
PARAMETER_VECTOR(us_rho_xs);
if (u_rho_xs.size()) {
  val -= dnorm(sum(us_rho_xs), Type(0.0), Type(0.001) * us_rho_xs.size(), true); // soft sum-to-zero
  val -= bym2_conditional_lpdf(u_rho_xs, us_rho_xs, sigma_rho_xs, phi_rho_xs, Q_x);
PARAMETER_VECTOR(u_rho_a);
if(u_rho_a.size() > 0)
  val += SCALE(AR1(phi_rho_a), sigma_rho_a)(u_rho_a);
PARAMETER VECTOR(u rho as);
if(u_rho_a.size() > 0)
  val += SCALE(AR1(phi_rho_as), sigma_rho_as)(u_rho_as);
PARAMETER_VECTOR(u_rho_xa);
if (u \text{ rho xa.size}() > 0) {
  val -= dnorm(sum(u_rho_xa), Type(0.0), sigma_rho_xa * Type(0.001) * u_rho_xa.size(), true); // soft
 val -= -(Q_x.rows() - Q_x_rankdef) * log_sigma_rho_xa -
    0.5 / (sigma_rho_xa * sigma_rho_xa) * (u_rho_xa * (Q_x * u_rho_xa)).sum();
// * ART coverage model *
PARAMETER(logit_phi_alpha_x);
```

```
Type phi_alpha_x(invlogit(logit_phi_alpha_x));
val -= log(phi_alpha_x) + log(1 - phi_alpha_x); // change of variables: logit_phi_x -> phi_x
val -= dbeta(phi_alpha_x, Type(0.5), Type(0.5), true);
PARAMETER(log_sigma_alpha_x);
Type sigma_alpha_x(exp(log_sigma_alpha_x));
val -= dnorm(sigma_alpha_x, Type(0.0), Type(2.5), true) + log_sigma_alpha_x;
PARAMETER(logit_phi_alpha_xs);
Type phi_alpha_xs(invlogit(logit_phi_alpha_xs));
val -= log(phi_alpha_xs) + log(1 - phi_alpha_xs); // change of variables: logit_phi_xs -> phi_xs
val -= dbeta(phi_alpha_xs, Type(0.5), Type(0.5), true);
PARAMETER(log_sigma_alpha_xs);
Type sigma_alpha_xs(exp(log_sigma_alpha_xs));
val -= dnorm(sigma_alpha_xs, Type(0.0), Type(2.5), true) + log_sigma_alpha_xs;
PARAMETER(logit_phi_alpha_a);
val -= dnorm(logit_phi_alpha_a, Type(0.0), Type(2.582), true); // INLA default
Type phi_alpha_a(2.0 * invlogit(logit_phi_alpha_a) - 1.0);
PARAMETER(log_sigma_alpha_a);
Type sigma_alpha_a(exp(log_sigma_alpha_a));
val -= dnorm(sigma_alpha_a, Type(0.0), Type(2.5), true) + log_sigma_alpha_a;
PARAMETER(logit_phi_alpha_as);
val -= dnorm(logit_phi_alpha_as, Type(0.0), Type(2.582), true); // INLA default
Type phi_alpha_as(2.0 * invlogit(logit_phi_alpha_as) - 1.0);
PARAMETER(log_sigma_alpha_as);
Type sigma_alpha_as(exp(log_sigma_alpha_as));
val -= dnorm(sigma_alpha_as, Type(0.0), Type(2.5), true) + log_sigma_alpha_as;
PARAMETER(log_sigma_alpha_xa);
Type sigma_alpha_xa(exp(log_sigma_alpha_xa));
val -= dnorm(sigma_alpha_xa, Type(0.0), Type(2.5), true) + log_sigma_alpha_xa;
PARAMETER VECTOR(u alpha x);
PARAMETER_VECTOR(us_alpha_x);
val -= dnorm(sum(us_alpha_x), Type(0.0), Type(0.001) * us_alpha_x.size(), true); // soft sum-to-zero
val -= bym2_conditional_lpdf(u_alpha_x, us_alpha_x, sigma_alpha_x, phi_alpha_x, Q_x);
PARAMETER VECTOR(u alpha xs);
PARAMETER VECTOR(us alpha xs);
if (u_alpha_xs.size()) {
  val -= dnorm(sum(us_alpha_xs), Type(0.0), Type(0.001) * us_alpha_xs.size(), true); // soft sum-to-z
  val -= bym2_conditional_lpdf(u_alpha_xs, us_alpha_xs, sigma_alpha_xs, phi_alpha_xs, Q_x);
PARAMETER_VECTOR(u_alpha_a);
if(u_alpha_a.size() > 0)
  val += SCALE(AR1(phi_alpha_a), sigma_alpha_a)(u_alpha_a);
```

```
PARAMETER_VECTOR(u_alpha_as);
if(u_alpha_as.size() > 0)
  val += SCALE(AR1(phi_alpha_as), sigma_alpha_as)(u_alpha_as);
PARAMETER_VECTOR(u_alpha_xa);
val -= dnorm(u_alpha_xa, 0.0, sigma_alpha_xa, true).sum();
// * HIV incidence model *
PARAMETER (OmegaT raw);
val -= dnorm(OmegaT_raw, Type(0.0), Type(1.0), true);
Type OmegaT = OmegaT0 + OmegaT_raw * sigma_OmegaT;
PARAMETER(log_betaT);
val -= dnorm(exp(log_betaT), Type(0.0), Type(1.0), true) + log_betaT;
Type betaT = betaT0 + exp(log_betaT) * sigma_betaT;
PARAMETER(log_sigma_lambda_x);
Type sigma_lambda_x(exp(log_sigma_lambda_x));
val -= dnorm(sigma_lambda_x, Type(0.0), Type(1.0), true) + log_sigma_lambda_x;
PARAMETER VECTOR(ui lambda x);
val -= sum(dnorm(ui_lambda_x, 0.0, sigma_lambda_x, true));
// * ANC testing model *
// ANC prevalence and ART coverage random effects
PARAMETER(log_sigma_ancrho_x);
Type sigma_ancrho_x(exp(log_sigma_ancrho_x));
val -= dnorm(sigma_ancrho_x, Type(0.0), Type(2.5), true) + log_sigma_ancrho_x;
PARAMETER(log_sigma_ancalpha_x);
Type sigma_ancalpha_x(exp(log_sigma_ancalpha_x));
val -= dnorm(sigma_ancalpha_x, Type(0.0), Type(2.5), true) + log_sigma_ancalpha_x;
PARAMETER VECTOR(ui anc rho x);
val -= sum(dnorm(ui_anc_rho_x, 0.0, sigma_ancrho_x, true));
PARAMETER_VECTOR(ui_anc_alpha_x);
val -= sum(dnorm(ui_anc_alpha_x, 0.0, sigma_ancalpha_x, true));
// * ART attendance model *
PARAMETER(log_sigma_or_gamma);
Type sigma_or_gamma(exp(log_sigma_or_gamma));
val -= dnorm(sigma_or_gamma, Type(0.0), Type(2.5), true) + log_sigma_or_gamma;
PARAMETER_VECTOR(log_or_gamma);
val -= dnorm(log_or_gamma, 0.0, sigma_or_gamma, true).sum();
```

```
// *** Process model ***
// HIV prevalence time 1
vector<Type> mu_rho(X_rho * beta_rho +
  logit rho offset +
  Z_rho_x * u_rho_x +
  Z rho xs * u rho xs +
  Z_rho_a * u_rho_a +
  Z_rho_as * u_rho_as +
  Z_rho_xa * u_rho_xa);
// paediatric prevalence
vector<Type> rho_15to49f_t1((X_15to49f * vector<Type>(invlogit(mu_rho) * population_t1)) / (X_15to49f
vector<Type> mu_rho_paed(X_paed_rho_ratio * rho_15to49f_t1 + paed_rho_ratio_offset);
mu_rho_paed = logit(mu_rho_paed);
mu_rho += mu_rho_paed;
// ART coverage time 1
vector<Type> mu_alpha(X_alpha * beta_alpha +
  logit_alpha_offset +
  Z_alpha_x * u_alpha_x +
  Z_alpha_xs * u_alpha_xs +
  Z_alpha_a * u_alpha_a +
  Z_alpha_as * u_alpha_as +
  Z_alpha_xa * u_alpha_xa);
vector<Type> rho_t1(invlogit(mu_rho));
vector<Type> alpha_t1(invlogit(mu_alpha));
vector<Type> plhiv_t1(population_t1 * rho_t1);
vector<Type> prop_art_t1(rho_t1 * alpha_t1);
vector<Type> artnum_t1(population_t1 * prop_art_t1);
vector<Type> plhiv 15to49 t1(X 15to49 * plhiv t1);
vector<Type> rho_15to49_t1(plhiv_15to49_t1 / (X_15to49 * population_t1));
vector<Type> alpha_15to49_t1((X_15to49 * artnum_t1) / plhiv_15to49_t1);
vector<Type> mu_lambda_t1(X_lambda * beta_lambda + log_lambda_t1_offset +
  Z_x * \text{vector} < \text{Type} > (\log(\text{rho}_15\text{to}49_{t1}) + \log(1.0 - \text{omega} * \text{alpha}_15\text{to}49_{t1})) +
  Z_lambda_x * ui_lambda_x);
vector<Type> lambda_adult_t1(exp(mu_lambda_t1));
// Add paediatric incidence
vector<Type> lambda_paed_t1(X_paed_lambda_ratio_t1 * rho_15to49f_t1);
vector<Type> lambda_t1(lambda_adult_t1 + lambda_paed_t1);
vector<Type> infections_t1(lambda_t1 * (population_t1 - plhiv_t1));
```

```
// likelihood for household survey data
vector<Type> rho_obs_t1((A_prev * plhiv_t1) / (A_prev * population_t1));
vector<Type> hhs_prev_ll = dbinom(x_prev, n_prev, rho_obs_t1, true);
val -= sum(hhs_prev_ll);
vector<Type> alpha_obs_t1((A_artcov * artnum_t1) / (A_artcov * plhiv_t1));
vector<Type> hhs artcov ll = dbinom(x artcov, n artcov, alpha obs t1, true);
val -= sum(hhs_artcov_ll);
vector<Type> pR_infections_obs_t1(A_recent * infections_t1);
vector<Type> pR_plhiv_obs_t1(A_recent * plhiv_t1);
vector<Type> pR_population_obs_t1(A_recent * population_t1);
vector<Type> pR_lambda_obs_t1(pR_infections_obs_t1 / (pR_population_obs_t1 - pR_plhiv_obs_t1));
vector<Type> pR_rho_obs_t1(pR_plhiv_obs_t1 / pR_population_obs_t1);
vector<Type> pR(1.0 - exp(-(pR_lambda_obs_t1 * (1.0 - pR_rho_obs_t1) / pR_rho_obs_t1 *
  (OmegaT - betaT * ritaT) + betaT)));
val -= dbinom(x_recent, n_recent, pR, true).sum();
// ANC prevalence and ART coverage model
// Note: currently this operates on the entire population vector, producing
         lots of zeros for males and female age groups not exposed to fertility.
//
         It would be more computationally efficient to project this to subset
//
         of female age 15-49 age groups. But I don't know if it would be
//
         meaningfully more efficient.
vector<Type> mu_anc_rho_t1(mu_rho +
  logit_anc_rho_t1_offset +
  X_ancrho * beta_anc_rho +
  Z_ancrho_x * ui_anc_rho_x);
vector<Type> anc_rho_t1(invlogit(mu_anc_rho_t1));
vector<Type> mu_anc_alpha_t1(mu_alpha +
  logit_anc_alpha_t1_offset +
  X_ancalpha * beta_anc_alpha +
  Z_ancalpha_x * ui_anc_alpha_x);
vector<Type> anc_alpha_t1(invlogit(mu_anc_alpha_t1));
// JE NOTE 6 Jan 2022: removed mu_asfr term -- should not use for aggregate ANC.
vector<Type> anc_clients_t1(population_t1 * exp(log_asfr_t1_offset));
vector<Type> anc_plhiv_t1(anc_clients_t1 * anc_rho_t1);
vector<Type> anc_already_art_t1(anc_plhiv_t1 * anc_alpha_t1);
// likelihood for ANC testing observations
vector<Type> anc_rho_obs_t1(A_anc_prev_t1 * anc_plhiv_t1 / (A_anc_prev_t1 * anc_clients_t1));
vector<Type> anc_rho_obs_t1_ll = dbinom(x_anc_prev_t1, n_anc_prev_t1, anc_rho_obs_t1, true);
val -= sum(anc_rho_obs_t1_ll);
vector<Type> anc_alpha_obs_t1(A_anc_artcov_t1 * anc_already_art_t1 / (A_anc_artcov_t1 * anc_plhiv_t1)
vector<Type> anc_alpha_obs_t1_ll = dbinom(x_anc_artcov_t1, n_anc_artcov_t1, anc_alpha_obs_t1, true);
val -= sum(anc_alpha_obs_t1_ll);
```

```
// * ART attendance model *
vector<Type> gamma_art_t1(exp(Xgamma * log_or_gamma + log_gamma_offset));
int cum nb = 0;
for(int i = 0; i < n_nb.size(); i++){</pre>
 Type cum_exp_or_gamma_i = 0.0;
 for(int j = 0; j < n_nb[i]+1; j++)
    cum_exp_or_gamma_i += gamma_art_t1[cum_nb + i + j];
 for(int j = 0; j < n_nb[i]+1; j++)
    gamma_art_t1[cum_nb + i + j] /= cum_exp_or_gamma_i;
  cum_nb += n_nb[i];
vector<Type> prop_art_ij_t1((Xart_idx * prop_art_t1) * (Xart_gamma * gamma_art_t1));
vector<Type> population_ij_t1(Xart_idx * population_t1);
vector<Type> artnum_ij_t1(population_ij_t1 * prop_art_ij_t1);
vector<Type> A_j_t1(A_artattend_t1 * artnum_ij_t1);
vector<Type> sd_A_j_t1(A_artattend_t1 * vector<Type>(population_ij_t1 * prop_art_ij_t1 * (1 - prop_ar
sd_A_j_t1 = sd_A_j_t1.sqrt();
vector<Type> artnum_t1_ll = dnorm(x_artnum_t1, A_j_t1, sd_A_j_t1, true);
val -= sum(artnum t1 ll);
// Calculate model outputs
DATA_SPARSE_MATRIX(A_out);
DATA_SPARSE_MATRIX(A_anc_out);
DATA_INTEGER(calc_outputs);
DATA_INTEGER(report_likelihood)
  if(calc_outputs) {
    vector<Type> population_t1_out(A_out * population_t1);
    vector<Type> plhiv_t1_out(A_out * plhiv_t1);
    vector<Type> rho_t1_out(plhiv_t1_out / population_t1_out);
   vector<Type> artnum_t1_out(A_out * artnum_t1);
   vector<Type> alpha_t1_out(artnum_t1_out / plhiv_t1_out);
    vector<Type> artattend_t1_out(A_out * (A_artattend_mf * artnum_ij_t1));
    vector<Type> artattend_ij_t1_out(A_art_reside_attend * artnum_ij_t1);
    vector<Type> untreated_plhiv_num_t1_out(plhiv_t1_out - artnum_t1_out);
    // Calculate number of PLHIV who attend facility in district i; denominator for artattend
    vector<Type> plhiv_attend_ij_t1((Xart_idx * plhiv_t1) * (Xart_gamma * gamma_art_t1));
    vector<Type> plhiv_attend_t1_out(A_out * (A_artattend_mf * plhiv_attend_ij_t1));
    vector<Type> untreated_plhiv_attend_t1_out(plhiv_attend_t1_out - artattend_t1_out);
    vector<Type> infections_t1_out(A_out * infections_t1);
    vector<Type> lambda_t1_out(infections_t1_out / (population_t1_out - plhiv_t1_out));
    vector<Type> anc_clients_t1_out(A_anc_out * anc_clients_t1);
```

```
vector<Type> anc_plhiv_t1_out(A_anc_out * anc_plhiv_t1);
  vector<Type> anc_already_art_t1_out(A_anc_out * anc_already_art_t1);
  // Note: assuming that:
  // (1) anc_known_pos is equivalent to anc_already_art
  // (2) All ANC attendees are diagnosed and initated on ART.
  vector<Type> anc_art_new_t1_out(anc_plhiv_t1_out - anc_already_art_t1_out);
  vector<Type> anc known pos t1 out(anc already art t1 out);
  vector<Type> anc_tested_pos_t1_out(anc_plhiv_t1_out - anc_known_pos_t1_out);
  vector<Type> anc_tested_neg_t1_out(anc_clients_t1_out - anc_plhiv_t1_out);
  vector<Type> anc_rho_t1_out(anc_plhiv_t1_out / anc_clients_t1_out);
  vector<Type> anc_alpha_t1_out(anc_already_art_t1_out / anc_plhiv_t1_out);
  REPORT(population_t1_out);
  REPORT(rho_t1_out);
  REPORT(plhiv_t1_out);
  REPORT(alpha_t1_out);
  REPORT(artnum_t1_out);
  REPORT(artattend_t1_out);
  REPORT(artattend ij t1 out);
  REPORT(untreated_plhiv_num_t1_out);
  REPORT(plhiv_attend_t1_out);
  REPORT(untreated_plhiv_attend_t1_out);
  REPORT(lambda t1 out);
  REPORT(infections t1 out);
  REPORT(anc clients t1 out);
  REPORT(anc_plhiv_t1_out);
  REPORT(anc_already_art_t1_out);
  REPORT(anc_art_new_t1_out);
  REPORT(anc_known_pos_t1_out);
  REPORT(anc_tested_pos_t1_out);
  REPORT(anc_tested_neg_t1_out);
  REPORT(anc_rho_t1_out);
  REPORT(anc_alpha_t1_out);
}
if(report_likelihood){
  REPORT(hhs_prev_11);
  REPORT(hhs artcov 11);
  REPORT(artnum t1 ll);
  REPORT(anc_rho_obs_t1_ll);
}
// Adam addition to include reporting of all latent field and hyper-parameter elements
// Hyper-parameter
REPORT(logit_phi_rho_x)
  REPORT(log_sigma_rho_x)
  REPORT(logit_phi_rho_xs)
```

```
REPORT(log_sigma_rho_xs)
REPORT(logit_phi_rho_a)
REPORT(log_sigma_rho_a)
REPORT(logit_phi_rho_as)
REPORT(log_sigma_rho_as)
REPORT(log_sigma_rho_xa)
REPORT(logit_phi_alpha_x)
REPORT(log sigma alpha x)
REPORT(logit_phi_alpha_xs)
REPORT(log_sigma_alpha_xs)
REPORT(logit_phi_alpha_a)
REPORT(log_sigma_alpha_a)
REPORT(logit_phi_alpha_as)
REPORT(log_sigma_alpha_as)
REPORT(log_sigma_alpha_xa)
REPORT(OmegaT_raw)
REPORT(log_betaT)
REPORT(log_sigma_lambda_x)
REPORT(log_sigma_ancrho_x)
REPORT(log_sigma_ancalpha_x)
REPORT(log_sigma_or_gamma)
// Latent field
REPORT(beta_rho)
REPORT(beta alpha)
REPORT(beta_lambda)
REPORT(beta anc rho)
REPORT(beta_anc_alpha)
REPORT(u_rho_x)
REPORT(us_rho_x)
REPORT(u_rho_xs)
REPORT(us_rho_xs)
REPORT(u_rho_a)
REPORT(u_rho_as)
REPORT(u_rho_xa)
REPORT(u_alpha_x)
REPORT(us_alpha_x)
REPORT(u alpha xs)
REPORT(us_alpha_xs)
REPORT(u alpha a)
REPORT(u_alpha_as)
REPORT(u_alpha_xa)
REPORT(ui lambda x)
REPORT(ui anc rho x)
REPORT(ui_anc_alpha_x)
REPORT(log_or_gamma)
return val;
```

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