

Modeling the Effects of Nutrition with Mixed-Effect Bayesian Network

Jari Turkia, CGI / University of Eastern Finland, jari.turkia@cgi.com

The Effects of Nutrition

I am proposing a Mixed-Effect Bayesian Network (MEBN) as method for modeling the effects of nutrition in both population and personal level. By the effects of nutrition we mean the way people react to different levels of nutrients at their diets. The Bayesian Networks (BN) are directed acyclic graphical models that contain both observed and latent random variables as vertices and edges as indicators of connection. In the setting of nutritional modeling the observed variables are nutrients at person's diet and their corresponding bodily responses, like blood characteristics. The connections between these variables can be very complex and the BNs seem to be intuitively appealing method for modeling such a system.

Besides of the typical correlations at the system, we are interested in the personal variations at the reaction types. Are some people more sensitive to some nutrient than the others? If this information can be systemically quantified, it allows us to predict personal networks of reaction types. This in turn, opens up lots of new applications in personal recommendations and health care.

For capturing these variances we need a dataset that contains several repeated measurements from number of persons. Measurements from each person are correlated with each other and a general method for modeling this correlation is hierarchical, or mixed-effect, model. The same model can be used to estimate the reaction type in both population and personal level.

Previously BNs have been considered mainly for uncorrelated observations (Aussem et al. 2010), but in this work we are using Mixed-Effect Bayesian Network parametrization (Bae et al. 2016) that allows combining the hierarchical mixed-effect modeling and Bayesian Network as a general framework for expressing the system.

This presentation covers first briefly the theory of graphical models and how it can be expanded to correlated observations. Then the it is shown how this modeling can be implemented with Stan and what benefits fully Bayesian modeling can offer in understanding the uncertainty at the model.

Mixed-Effects Bayesian Network

Let us denote the graph of interconnected nutrients and responses with G . We can then formulate the modeling problem as finding the graph G that is most probable given the data

$$P(G|D) \tag{1}$$

By using Bayes' Rule we can be split this probability into proportions of data likelihood with given graph and any prior information we might have about suitable graphs

$$P(G|D) \propto P(D|G)P(G) \tag{2}$$

Now the problem is converted into a search of the maximum likelihood graph for given data. If all the graphs are equally probable then $P(G)$ is a constant and does not affect the search, but it can be also beneficial to use it to guide the search towards meaningful graphs.

Decomposition of the likelihood. Bayesian network factorizes into local distributions according to *local Markov property* stating that a variable X_i is independent of its non-descendants given its parents at the graph. The *global Markov property* states that a variable is independent of all the remaining variables in the graph conditionally on its *Markov blanket* that consists its parents and child nodes at the graph, and additional parents of the child nodes (Bae et al. 2016, Koller and Friedman (2009)). With this decomposition the joint probability of the graph can be calculated with sum and product rules of probability as a product of the independent local graphs G_i . The graph structure depends also on the parameters θ_i describing the dependencies and they should be taken into account at the estimation.

$$P(D|G) = \prod_{i=1}^v P(y_i|pa(y_i), \theta_i, G_i) P(\theta_i|G_i) \quad (3)$$

assuming we have v independent local distributions at the graph. Since the probability of data in the graph depends on the parameters of the local distributions, θ_i , they have to be integrated out from the equation to make the probability of graph independent of any specific choice of parameters.

$$P(D|G) = \prod_{i=1}^v \int P(y_i|pa(y_i), \theta_i) P(\theta_i|G_i) d\theta_i \quad (4)$$

Linear dependency between variables. As we are more interested in the system and less considered about details of any specific nutritional response, we consider it adequate to model the dependency between the nutrients and bodily responses with an approximate linear model. However, simple linear model is not enough, but we need a parametrization that is able to reflect the correlations between observations and express the amount of variability between persons since the data consists of several repeated measurements from different persons.

Generally, local probability distributions can be from exponential family, but in this example we consider only normally distributed response variables. For every local distribution of response variable X_i we denote its parent nodes with $pa(X_i)$. Subset of parent nodes possibly containing personal variance is denoted with $pa(Z_i)$. Matrix V_i is the variance-covariance matrix that expresses the uncertainty of the prediction with as variance of the distribution.

$$P(X_i|pa(X_i)) = N(X_i|pa(X_i)\beta + pa(Z_i)b, V_i) \quad (5)$$

This theory motivates our search for optimal graph with Stan. By decomposing the joint likelihood into local probability distributions according to Markov properties it is possible to find the optimal graph by estimating one local distributions one by one.

Estimating the Hierarchical Local Distributions with Stan

Let us elaborate still how variance V is calculated and how this calculation can be implemented with Stan. Consider the probability distribution of response variable y_i with possible parameters θ_i and subgraph G_i

$$P(y_i|X, \theta_i, G_i) = N(X\beta + Zb, V) \quad (6)$$

Besides the coefficients β and b , we are also interested in their variances and covariances, defined by

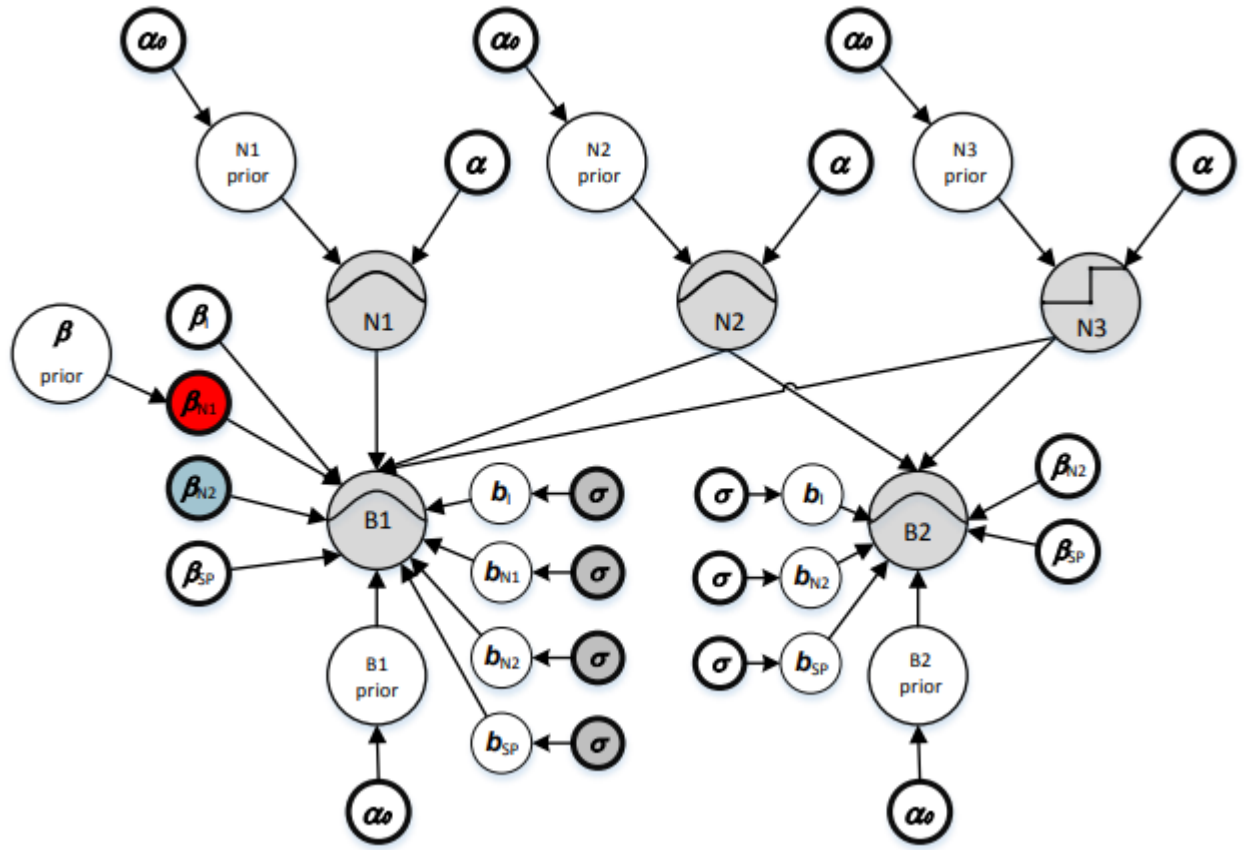


Figure 1: Bayesian Network of nutrients and responses with all parameters and hyperparameters

$$V = ZDZ' + R \quad (7)$$

where R is a variance-covariance matrix of residuals and D is a variance-covariance matrix of personal, or random-effects

$$D = \mathcal{T}C\mathcal{T}' \quad (8)$$

There \mathcal{T} is a diagonal matrix of personal effect variances and C is correlation matrix that can be divided into Cholesky decompositions as

$$C = LL' \quad (9)$$

and with L we can define the personal effects as

$$b = Lu, u \sim N(0, I) \quad (10)$$

as we assume for now that personal random-effects are drawn from Normal distribution.

This is implemented in Stan as follows

```
transformed parameters {
  // ...
  // Create diagonal matrix from sigma_b and premultiply it with L
  D = diag_pre_multiply(sigma_b, L);

  // Group-level effects are generated by multiplying D with z that has standard normal distribution
  for(j in 1:J)
    b[j] = D * z[j];
}
```

The actual model with Normal distribution having the linear mixed-effect likelihood is defined below. Notice that instead of matrix V , in Stan we are using vectors `group` and scalar `sigma_e`.

```
model {
  // ...
  // Standard normal prior for random effects
  for (j in 1:J)
    z[j] ~ normal(0,1);

  // Likelihood
  // - link function (identity function for Normal dist.) for typical correlation
  mu = temp_Intercept + Xc * beta;

  // - add personal (group) effects
  for (i in 1:N)
  {
    mu[i] = mu[i] + Z[i] * b[group[i]];
  }

  // Y and mu are vectors, sigma_e is a scalar that is estimated for whole vector
  Y ~ normal(mu, sigma_e);
}
```

Constructing the Population Level Graph of Nutritional Effects

The dataset in this example comes from Sysdimet study (Lankinen et al. 2011). We have picked few interesting variables indicating person's diet, blood test results and personal information, like gender and medication.

There exist plenty of general algorithms for constructing BNs, but for this special case we can constrain the search to biologically plausible reaction graphs. We assume that all possible graphs are directed bipartite graphs with nutrients and personal information as root nodes and blood tests as targets.

```
# Read the data description
datadesc <- mebn.load_datadesc("Data description.xlsx")

# Read the actual data matching the description
sysdimet <- read.csv(file="data\\SYSDIMET_diet.csv", sep=";", dec=",")

# Define how to iterate through the graph
assumedpredictors <- datadesc[datadesc$Order==100,]
assumedtargets <- datadesc[datadesc$Order==200,]
```

Pruning the edges. Construction of the graph starts from fully connected graph, but for gaining nutritional knowledge of significant connections and also for efficient factorization of the BN likelihood, it is necessary to prune out the insignificant connections at the graph. For this we use shrinkage prior on beta coefficients to push the insignificant coefficients towards zero. Especially, we use regularized horseshoe prior (Piironen and Vehtari 2017a) that allows specifying the number of non-zero coefficients for each target. In the nutritional setting this provides a way for specifying prior knowledge about the relevant nutrients for each response. For now, we approximate that one third of the predictive nutrients are relevant, but finer approximation will be done based on previous nutritional research.

If the shrinkage prior does not shrink the coefficients to exactly zero, we are pruning out the insignificant connections with following test. Notice that in the population level graph we are keeping the connections that have large variance between persons even though they are not typically relevant. Personal variance means that the connection is relevant for someone.

To assure that this pruning does not affect predicting accuracy of the model a projection method (Piironen and Vehtari 2017b) could be also used here. In projection approach the edges are removed if it does not affect the distance from the true model measured with KL-divergence.

```
my.RanefTest <- function(localsummary, PredictorId)
{
  abs(localsummary$fixef[PredictorId]) > 0.01 ||
  localsummary$ranef_sd[PredictorId] > 0.01
}
```

Constructing the graph. The data structure of the graph is based on iGraph package. The process of BN construction starts by adding a node for every observed variable at the dataset.

```
# Add data columns describing random variables as nodes to the graph
# - initial_graph is iGraph object with only nodes and no edges
initial_graph <- mebn.new_graph_with_randomvariables(datadesc)
```

The joint distribution of the Bayesian Network is decomposed as local distributions, one for each target variable. These local distributions correspond to hierarchical regression models that are estimated with Stan and the resulting MCMC sampling is cached to files.

For estimating the typical graph, we iterate the hierarchical Stan-model through all the assumed predictors and targets. The result is an iGraph object that contains a directed bipartite graph with relating nutrients as predictors and blood test results as targets.

```
sysdimet_graph <- mebn.typical_graph(reaction_graph = initial_graph,
                                     inputdata = sysdimet,
                                     predictor_columns = assumedpredictors,
                                     assumed_targets = assumedtargets,
                                     group_column = "SUBJECT_ID",
                                     local_estimation = mebn.sampling,
                                     local_model_cache = "models",
                                     stan_model_file = "mebn\\BLMM_rhs.stan",
                                     edge_significance_test = my.Raneftest,
                                     normalize_values = TRUE,
                                     use_regularization = TRUE)
```

```
## [1] "Loading models\\fshdl_blmm.rds"
## [1] "Loading models\\fsldl_blmm.rds"
## [1] "Loading models\\fsins_blmm.rds"
## [1] "Loading models\\fskol_blmm.rds"
## [1] "Loading models\\fpgluk_blmm.rds"
```

Now “sysdimet_graph” is a Mixed Effect Bayesian Network for whole sample population. We can store it in GEXF-format and load to visualization for closer inspection. The visual nature of the Bayesian Networks, and the graphical models in general, provide a useful framework for creating visualizations. Comparing to schematic figure 1, the coefficients and latent variables are removed and denoted by different colors. Blue edge denotes typically negative correlation, red edge denotes positive correlation and gray shade denotes the amount of personal variation at the correlation.

It can be seen from the visualization that many of the nutrients affect to several responses. This multiresponse effect is not yet implemented to the model, but should be accounted. It allows, for example, to gain knowledge about responses that have missing measurements for some persons, but have known connections to other responses and predictors.

Inference

The graph gives an overview and we can query some of the most interesting reactions. We can, for example, investigate most significant typical reactions by querying largest beta coefficients

```
allnodes <- V(sysdimet_graph)
beta <- allnodes[allnodes$type=="beta"]

typical_effects<-data.frame(matrix(NA, nrow=length(beta), ncol=0))
typical_effects$name <- beta$name
typical_effects$value <- beta$value

largest_typical_negative <- typical_effects[order(typical_effects$value),]
largest_typical_positive <- typical_effects[order(-typical_effects$value),]
```

Largest typically negative effects

```
##               name               value
## 24 beta_kolestrolilaakitys_fsldl -0.031008304
## 68 beta_kolestrolilaakitys_fskol -0.013418128
## 12          beta_hhydr_fshdl -0.006197030
## 3          beta_energia_fshdl -0.005957983
## 45          beta_sukupuoli_fsins -0.004618822
```

Largest typically positive effects

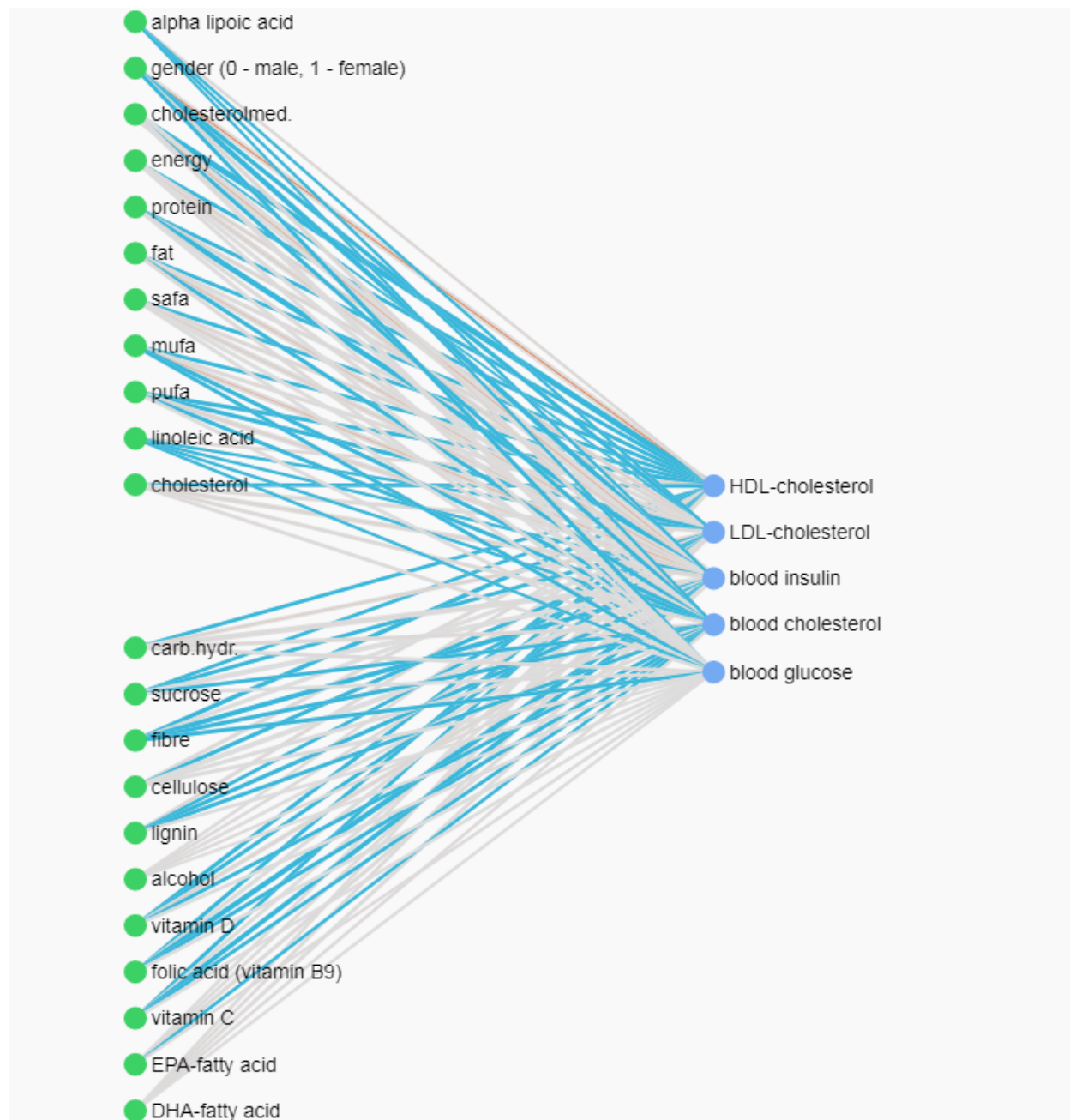


Figure 2:

```
##              name      value
## 1      beta_sukupuoli_fshdl 0.018007452
## 46 beta_kolestrolilaakitys_fsins 0.008752224
## 51              beta_mufa_fsins 0.007505062
## 49              beta_rasva_fsins 0.006085272
## 47      beta_energia_fsins 0.004064071
```

It can be inspected that females have typically higher levels of HDL cholesterol and cholesterol medication, besides of lowering the cholesterol levels, also raises blood insuling level.

What we are really interested in, though, are the variations between persons. For this, we can query the largest variances of random-effects..

```
b_sigma <- allnodes[allnodes$type=="b_sigma"]

personal_variances<-data.frame(matrix(NA, nrow=length(b_sigma), ncol=0))
personal_variances$name <- b_sigma$name
personal_variances$value <- b_sigma$value

largest_personal_variance <- personal_variances[order(-personal_variances$value),]
head(largest_personal_variance, 10)
```

```
##              name      value
## 47      b_sigma_energia_fsins 0.16384585
## 3      b_sigma_energia_fshdl 0.10823367
## 25      b_sigma_energia_fsldl 0.10494613
## 48      b_sigma_prot_fsins 0.09991203
## 50      b_sigma_safa_fsins 0.09930752
## 69      b_sigma_energia_fskol 0.09913437
## 57      b_sigma_sakkar_fsins 0.09774721
## 52      b_sigma_pufa_fsins 0.09769463
## 51      b_sigma_mufa_fsins 0.09746815
## 46 b_sigma_kolestrolilaakitys_fsins 0.09725983
```

So, there seems to be large variance in how the energy intake affects to insulin and cholesterol levels.

Relevance of these estimations can be inspected from their posterior distributions.

```
library("bayesplot")

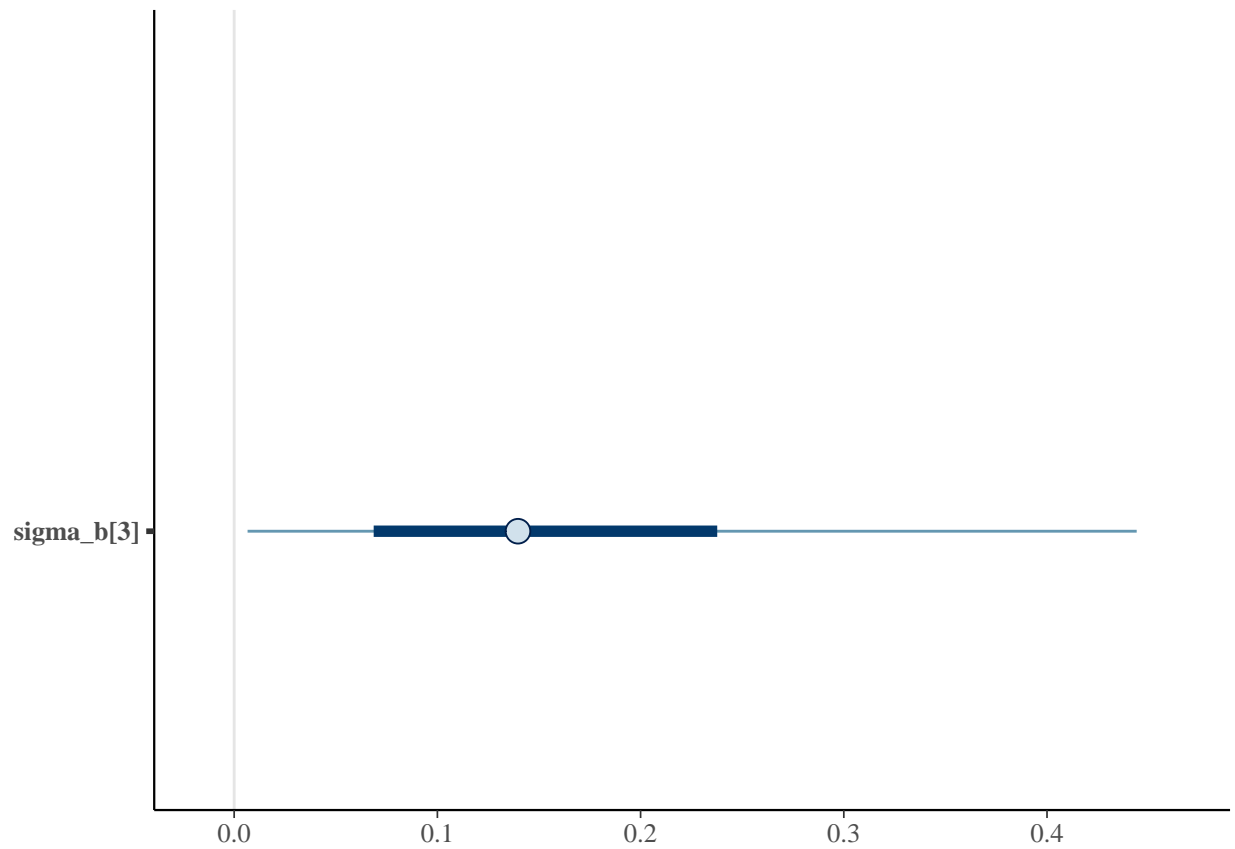
# Local distribution for fsins
fsins_blmm <- mebn.get_localfit("fsins")

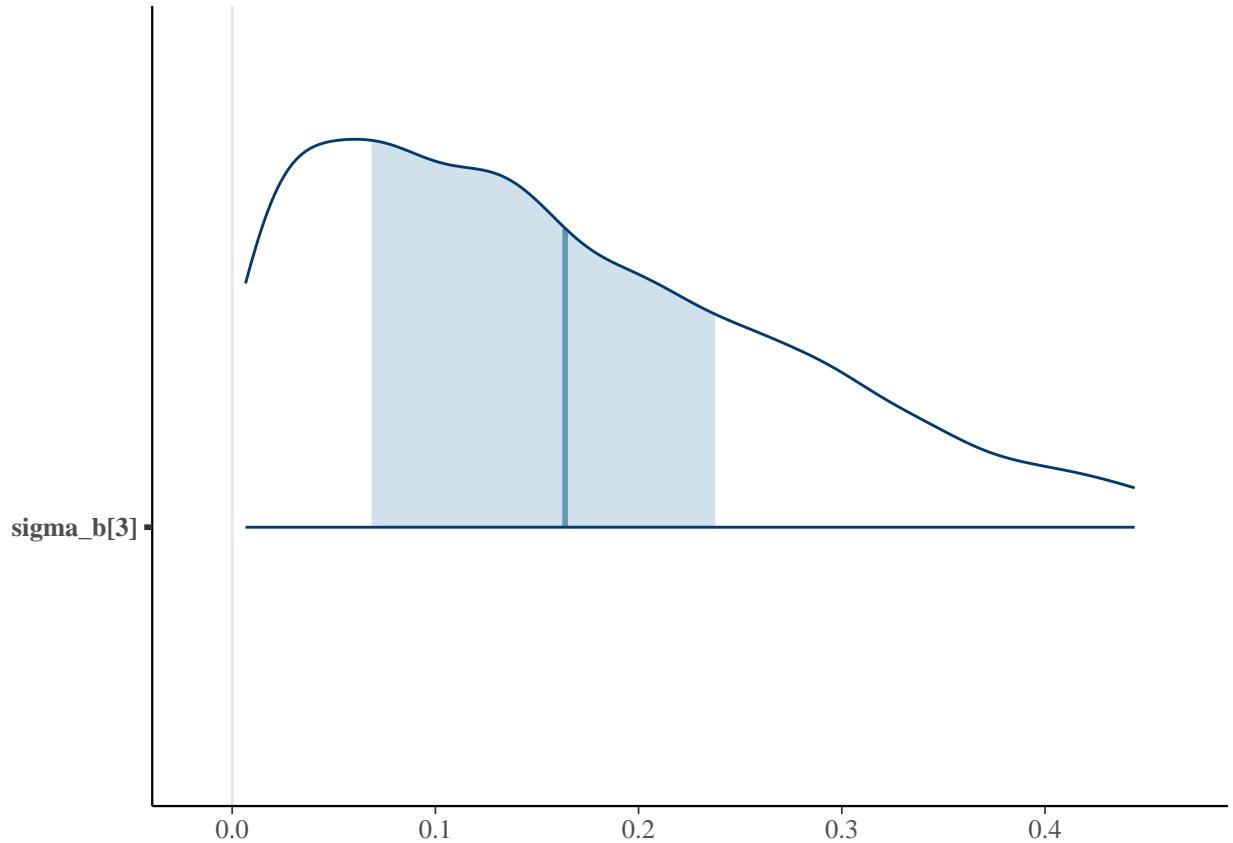
## [1] "Loading models\\fsins_blmm.rds"

# Index of predictor 'energia'
id <- match("energia", datadesc$Name)

posterior <- as.array(fsins_blmm)

mcmc_intervals(posterior, pars = c(paste0("sigma_b[",id,"]")), prob_outer = 0.95)
```



This shows that there exists personal variance in how energy intake affects blood insulin levels as the 95% credible interval is above zero. The wide probability distribution shows that exact estimate is quite uncertain and we would need more observations or stricter prior information for more precise estimation. One can also observe the difference between typical choices of Bayesian point estimation; upper interval chart points at the posterior median, and the lower chart shows posterior mean and MAP estimate at high point of the probability.

Conclusion

Bayesian Networks offer an appealing way to model the system of nutritional effects. By incorporating the mixed-effect modeling to local probability distributions we can estimate the effects in both population and personal level. Furthermore, the fully Bayesian estimation of the distributions allow direct means for including prior information at the model and also addressing the uncertainty of the estimates.

After estimating the personal variances it is possible next to predict this kind of personal reaction graph for a new person. This enables interesting applications in personal recommendations and personal health care.

References

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