

LaplacesDemon Examples

Statisticat, LLC

Abstract

The **LaplacesDemon** package is a complete environment for Bayesian inference within R. Virtually any probability model may be specified. This vignette is a compendium of examples of how to specify different model forms.

Keywords: Bayesian, Bayesian Inference, Laplace's Demon, LaplacesDemon, R, Statisticat.

LaplacesDemon (Statisticat LLC. 2012), usually referred to as Laplace's Demon, is an R package that is available on CRAN (R Development Core Team 2012). A formal introduction to Laplace's Demon is provided in an accompanying vignette entitled "**LaplacesDemon** Tutorial", and an introduction to Bayesian inference is provided in the "Bayesian Inference" vignette.

The purpose of this document is to provide users of the **LaplacesDemon** package with examples of a variety of Bayesian methods. It is also a testament to the diverse applicability of **LaplacesDemon** to Bayesian inference.

To conserve space, the examples are not worked out in detail, and only the minimum of necessary materials is provided for using the various methodologies. Necessary materials include the form expressed in notation, data (which is often simulated), the Model function, and initial values. The provided data, model specification, and initial values may be copy/pasted into an R file and updated with the LaplacesDemon or (usually) LaplaceApproximation functions. Although many of these examples update quickly, some examples are computationally intensive.

Initial values are usually hard-coded in the examples, though the Parameter-Generating Function (PGF) is also specified. It is recommended to generate initial values with the GIV function according to the user-specified PGF.

Notation in this vignette follows these standards: Greek letters represent parameters, lower case letters represent indices, lower case bold face letters represent scalars or vectors, probability distributions are represented with calligraphic font, upper case letters represent index limits, and upper case bold face letters represent matrices. More information on notation is available at http://www.bayesian-inference.com/notation.

This vignette will grow over time as examples of more methods become included. Contributed examples are welcome. Please send contributed examples or discovered errors in a similar format in an email to software@bayesian-inference.com for review and testing. All accepted

contributions are, of course, credited.

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1. ANCOVA

This example is essentially the same as the two-way ANOVA (see section 3), except that a covariate $X_{,3}$ has been added, and its parameter is δ .

1.1. Form

$$\mathbf{y}_{i} \sim \mathcal{N}(\mu_{i}, \sigma_{1}^{2})$$

$$\mu_{i} = \alpha + \beta[\mathbf{X}_{i,1}] + \gamma[\mathbf{X}_{i,2}] + \delta \mathbf{X}_{i,2}, \quad i = 1, \dots, N$$

$$\epsilon_{i} = \mathbf{y}_{i} - \mu_{i}$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\beta_{j} \sim \mathcal{N}(0, \sigma_{2}^{2}), \quad j = 1, \dots, J$$

$$\beta_{J} = -\sum_{i=1}^{J-1} \beta_{j}$$

$$\gamma_k \sim \mathcal{N}(0, \sigma_3^2), \quad k = 1, \dots, K$$

$$\gamma_K = -\sum_{k=1}^{K-1} \gamma_k$$

$$\delta \sim \mathcal{N}(0, 1000)$$

$$\sigma_m \sim \mathcal{HC}(25), \quad m = 1, \dots, 3$$

1.2. Data

```
N <- 100
J <- 5 #Number of levels in factor (treatment) 1
K <- 3 #Number of levels in factor (treatment) 2</pre>
X \leftarrow matrix(cbind(runif(N,0.5,J+0.49)),round(runif(N,0.5,K+0.49)),
    runif(N,-2,2)), N, 3)
alpha <- runif(1,-1,1)
beta \leftarrow runif(J,-2,2)
beta[J] <- -sum(beta[1:(J-1)])</pre>
gamma <- runif(K,-2,2)</pre>
gamma[J] <- -sum(gamma[1:(K-1)])
delta <- runif(1,-2,2)
y \leftarrow alpha + beta[X[,1]] + gamma[X[,2]] + delta*X[,3] + rnorm(N,0,0.1)
mon.names <- c("LP", "beta[5]", "gamma[3]", "sigma[1]", "sigma[2]", "sigma[3]",
     "s.beta", "s.gamma", "s.epsilon")
parm.names <- as.parm.names(list(alpha=0, beta=rep(0,J-1), gamma=rep(0,K-1),
    delta=0, log.sigma=rep(0,3)))
PGF <- function(Data) return(c(rnormv(1,1,10), rnorm(Data$J-1,0,1),
    rnorm(Data$K-1,0,1), rnormv(1,0,10), log(rhalfcauchy(3,25))))
MyData <- list(J=J, K=K, N=N, PGF=PGF, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
```

```
Model <- function(parm, Data)
    {
     ### Parameters
     alpha <- parm[1]
     beta <- rep(NA,Data$J)
     beta[1:(Data$J-1)] <- parm[2:Data$J]
     beta[Data$J] <- -sum(beta[1:(Data$J-1)]) #Sum-to-zero constraint
     gamma <- rep(NA,Data$K)
     gamma[1:(Data$K-1)] <- parm[grep("gamma", Data$parm.names)]
     gamma[Data$K] <- -sum(gamma[1:(Data$K-1)]) #Sum-to-zero constraint
     delta <- parm[grep("delta", Data$parm.names)]
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])
     ### Log(Prior Densities)</pre>
```

```
alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
beta.prior <- sum(dnorm(beta, 0, sigma[2], log=TRUE))</pre>
gamma.prior <- sum(dnorm(gamma, 0, sigma[3], log=TRUE))</pre>
delta.prior <- dnormv(delta, 0, 1000, log=TRUE)</pre>
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
### Log-Likelihood
mu <- alpha + beta[Data$X[,1]] + gamma[Data$X[,2]] +</pre>
    delta*Data$X[,3]
LL <- sum(dnorm(Data$y, mu, sigma[1], log=TRUE))</pre>
### Variance Components
s.beta <- sd(beta)
s.gamma <- sd(gamma)
s.epsilon <- sd(Data$y - mu)
### Log-Posterior
LP <- LL + alpha.prior + beta.prior + gamma.prior + delta.prior +
    sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, beta[Data$J],</pre>
    gamma[Data$K], sigma, s.beta, s.gamma, s.epsilon),
    yhat=rnorm(length(mu), mu, sigma[1]), parm=parm)
return(Modelout)
}
```

1.4. Initial Values

Initial. Values $\leftarrow c(0, rep(0, (J-1)), rep(0, (K-1)), 0, rep(log(1), 3))$

2. ANOVA, One-Way

When J = 2, this is a Bayesian form of a t-test.

2.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma_1^2)$$

$$\mu_i = \alpha + \beta[\mathbf{x}_i], \quad i = 1, \dots, N$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\beta_j \sim \mathcal{N}(0, \sigma_2^2), \quad j = 1, \dots, J$$

$$\beta_J = -\sum_{j=1}^{J-1} \beta_j$$

$$\sigma_{1:2} \sim \mathcal{HC}(25)$$

```
2.2. Data
```

2.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    alpha <- parm[1]
    beta <- rep(NA,Data$J)</pre>
    beta[1:(Data$J-1)] <- parm[2:Data$J]</pre>
    beta[Data$J] <- -sum(beta[1:(Data$J-1)]) #Sum-to-zero constraint
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    beta.prior <- sum(dnorm(beta, 0, sigma[2], log=TRUE))</pre>
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- alpha + beta[Data$x]</pre>
    LL <- sum(dnorm(Data$y, mu, sigma[1], log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + sigma.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,beta[Data$J],</pre>
          sigma), yhat=rnorm(length(mu), mu, sigma[1]), parm=parm)
    return(Modelout)
    }
```

2.4. Initial Values

```
Initial. Values \leftarrow c(0, rep(0, (J-1)), rep(log(1), 2))
```

3. ANOVA, Two-Way

In this representation, σ^m are the superpopulation variance components, s.beta and s.gamma are the finite-population within-variance components of the factors or treatments, and s.epsilon is the finite-population between-variance component.

3.1. Form

$$\mathbf{y}_{i} \sim \mathcal{N}(\mu_{i}, \sigma_{1}^{2})$$

$$\mu_{i} = \alpha + \beta[\mathbf{X}_{i,1}] + \gamma[\mathbf{X}_{i,2}], \quad i = 1, \dots, N$$

$$\epsilon_{i} = \mathbf{y}_{i} - \mu_{i}$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\beta_{j} \sim \mathcal{N}(0, \sigma_{2}^{2}), \quad j = 1, \dots, J$$

$$\beta_{J} = -\sum_{j=1}^{J-1} \beta_{j}$$

$$\gamma_{k} \sim \mathcal{N}(0, \sigma_{3}^{2}), \quad k = 1, \dots, K$$

$$\gamma_{K} = -\sum_{k=1}^{K-1} \gamma_{k}$$

$$\sigma_{m} \sim \mathcal{HC}(25), \quad m = 1, \dots, 3$$

```
N <- 100
J <- 5 #Number of levels in factor (treatment) 1
K <- 3 #Number of levels in factor (treatment) 2</pre>
X <- matrix(cbind(round(runif(N, 0.5, J+0.49)),round(runif(N,0.5,K+0.49))),</pre>
    N, 2)
alpha <- runif(1,-1,1)
beta <- runif(J,-2,2)
beta[J] <- -sum(beta[1:(J-1)])</pre>
gamma <- runif(K,-2,2)</pre>
gamma[J] <- -sum(gamma[1:(K-1)])
y \leftarrow alpha + beta[X[,1]] + gamma[X[,2]] + rnorm(1,0,0.1)
mon.names <- c("LP", "beta[5]", "gamma[3]", "sigma[1]", "sigma[2]", "sigma[3]",
    "s.beta", "s.gamma", "s.epsilon")
parm.names <- as.parm.names(list(alpha=0, beta=rep(0,J-1), gamma=rep(0,K-1),
    log.sigma=rep(0,3))
PGF <- function(Data) return(c(rnormv(1,0,1000), rnorm(Data$J-1,0,1),
    rnorm(Data$K-1,0,1000), log(rhalfcauchy(3,25))))
MyData <- list(J=J, K=K, N=N, PGF=PGF, X=X, mon.names=mon.names,
```

```
parm.names=parm.names, y=y)
```

3.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    alpha <- parm[1]
    beta <- rep(NA,Data$J)</pre>
    beta[1:(Data$J-1)] <- parm[2:Data$J]</pre>
    beta[Data$J] <- -sum(beta[1:(Data$J-1)]) #Sum-to-zero constraint
     gamma <- rep(NA,Data$K)</pre>
    gamma[1:(Data$K-1)] <- parm[grep("gamma", Data$parm.names)]</pre>
     gamma[Data$K] <- -sum(gamma[1:(Data$K-1)]) #Sum-to-zero constraint</pre>
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
     ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    beta.prior <- sum(dnorm(beta, 0, sigma[2], log=TRUE))</pre>
     gamma.prior <- sum(dnorm(gamma, 0, sigma[3], log=TRUE))</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- alpha + beta[Data$X[,1]] + gamma[Data$X[,2]]</pre>
    LL <- sum(dnorm(Data$y, mu, sigma[1], log=TRUE))</pre>
    ### Variance Components
     s.beta <- sd(beta)
     s.gamma <- sd(gamma)
    s.epsilon <- sd(Data$y - mu)</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + gamma.prior +
         sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, beta[Data$J],</pre>
         gamma[Data$K], sigma, s.beta, s.gamma, s.epsilon),
         yhat=rnorm(length(mu), mu, sigma[1]), parm=parm)
    return(Modelout)
     }
```

3.4. Initial Values

```
Initial. Values \leftarrow c(0, rep(0, (J-1)), rep(0, (K-1)), rep(log(1), 3))
```

4. Approximate Bayesian Computation (ABC)

Approximate Bayesian Computation (ABC), also called likelihood-free estimation, is not a statistical method, but a family of numerical approximation techniques in Bayesian inference. ABC is especially useful when evaluation of the likelihood, $p(\mathbf{y}|\Theta)$ is computationally pro-

hibitive, or when suitable likelihoods are unavailable. The current example is the application of ABC in the context of linear regression. The log-likelihood is replaced with the negative sum of the distance between **y** and **y**^{rep} as the approximation of the log-likelihood. Distance reduces to the absolute difference. Although linear regression has an easily calculated likelihood, it is used as an example due to its generality. This example demonstrates how ABC may be estimated either with MCMC via the LaplacesDemon function or with Laplace Approximation via the LaplaceApproximation function. In this method, a tolerance (which is found often in ABC) does not need to be specified, and the logarithm of the unnormalized joint posterior density is maximized, as usual. The negative and summed distance, above, may be replaced with the negative and summed distance between summaries of the data, rather than the data itself, but this has not been desirable in testing.

4.1. Form

$$\mathbf{y} = \mu + \epsilon$$
$$\mu = \mathbf{X}\beta$$
$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

4.2. Data

```
Model <- function(parm, Data)
    {
     ### Parameters
     beta <- parm[1:Data$J]
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
     ### Log-Likelihood Approximation
     mu <- as.vector(tcrossprod(Data$X, t(beta)))
     epsilon <- Data$y - mu
     sigma <- sd(epsilon)
     LL <- -sum(abs(epsilon))</pre>
```

4.4. Initial Values

Initial.Values <- c(rep(0,J))</pre>

5. ARCH-M(1,1)

5.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{t}^{2}), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} \sim \mathcal{N}(\mu_{T+1}, \sigma_{new}^{2})$$

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1} + \delta \sigma_{t-1}^{2}, \quad t = 1, \dots, (T+1)$$

$$\epsilon_{t} = \mathbf{y}_{t} - \mu_{t}$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\delta \sim \mathcal{N}(0, 1000)$$

$$\sigma_{new}^{2} = \omega + \theta \epsilon_{T}^{2}$$

$$\sigma_{t}^{2} = \omega + \theta \epsilon_{t-1}^{2}$$

$$\omega < -\mathcal{HC}(25)$$

$$\theta \sim \mathcal{U}(0, 1)$$

```
y <- c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91, 2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28, 1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03, -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02, 0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36, 1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70, 0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83, 0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24, 0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40, -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
```

```
0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17
T <- length(y)
mon.names <- c("LP", "ynew", "sigma2.new")</pre>
parm.names <- c("alpha","phi","delta","log.omega","theta")</pre>
PGF <- function(Data) return(c(rnormv(3,0,1000), log(rhalfcauchy(1,25)),
    runif(1)))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,
    y=y)
5.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    alpha <- parm[1]; phi <- parm[2]; delta <- parm[3]</pre>
    omega <- exp(parm[4])</pre>
    parm[5] <- theta <- interval(parm[5], 1e-10, 1-1e-5)</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
    delta.prior <- dnormv(delta, 0, 1000, log=TRUE)</pre>
    omega.prior <- dhalfcauchy(omega, 25, log=TRUE)</pre>
    theta.prior <- dunif(theta, 0, 1, log=TRUE)
    ### Log-Likelihood
    mu <- c(alpha, alpha + phi*Data$y[-Data$T])</pre>
    epsilon <- Data$y - mu
    sigma2 <- c(omega, omega + theta*epsilon[-Data$T]^2)</pre>
    mu <- mu + delta*sigma2
```

sigma2.new <- omega + theta*epsilon[Data\$T]^2</pre>

5.4. Initial Values

Initial. Values $\leftarrow c(rep(0,3), rep(0.5,2))$

6. Autoregression, AR(1)

6.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma^{2}), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} = \alpha + \mu_{T+1}$$

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1}, \quad t = 1, \dots, (T+1)$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\sigma \sim \mathcal{HC}(25)$$

```
y <- c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91, 2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28, 1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03, -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02, 0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36, 1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70, 0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83, 0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24, 0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40, -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19, 0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39, -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14, -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06, -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
```

```
0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17
T <- length(y)
mon.names <- c("LP", "sigma", "ynew")</pre>
parm.names <- c("alpha", "phi", "log.sigma")</pre>
PGF <- function(Data) return(c(rnormv(2,0,1000), log(rhalfcauchy(1,25))))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,
    y=y)
```

6.3. Model

```
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    alpha <- parm[1]; phi <- parm[2]; sigma <- exp(parm[3])</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- c(alpha, alpha + phi*Data$y[-Data$T])</pre>
    ynew <- rnorm(1, alpha + phi*Data$y[Data$T], sigma)</pre>
    LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + phi.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, sigma, ynew),</pre>
          yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

6.4. Initial Values

```
Initial. Values \leftarrow c(rep(0,2), log(1))
```

7. Autoregressive Conditional Heteroskedasticity, ARCH(1,1)

7.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{t}^{2}), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} \sim \mathcal{N}(\mu_{T+1}, \sigma_{new}^{2})$$

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1}, \quad t = 1, \dots, (T+1)$$

$$\epsilon_{t} = \mathbf{y}_{t} - \mu_{t}$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\sigma_{new}^{2} = \omega + \theta \epsilon_{T}^{2}$$

$$\sigma_{t}^{2} = \omega + \theta \epsilon_{t-1}^{2},$$

$$\omega \sim \mathcal{HC}(25)$$

$$\theta \sim \mathcal{U}(0, 1)$$

```
y < -c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91,
    2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28,
    1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03,
    -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02,
    0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36,
    1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70,
    0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83,
    0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24,
    0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40,
    -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
    0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
```

```
-0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17)
T <- length(y)
mon.names <- c("LP", "ynew", "sigma2.new")</pre>
parm.names <- c("alpha","phi","log.omega","theta")</pre>
PGF <-function(Data) return(c(rnormv(2,0,1000), log(rhalfcauchy(1,25)),
    runif(1)))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,</pre>
    y=y)
7.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    alpha <- parm[1]; phi <- parm[2]; omega <- exp(parm[3])</pre>
    parm[4] <- theta <- interval(parm[4], 1e-10, 1-1e-5)</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
    omega.prior <- dhalfcauchy(omega, 25, log=TRUE)</pre>
    theta.prior <- dunif(theta, 0, 1, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- c(alpha, alpha + phi*Data$y[-Data$T])</pre>
    epsilon <- Data$y - mu
    sigma2 <- c(omega, omega + theta*epsilon[-Data$T]^2)</pre>
    sigma2.new <- omega + theta*epsilon[Data$T]^2</pre>
    ynew <- rnormv(1, alpha + phi*Data$y[Data$T], sigma2.new)</pre>
    LL <- sum(dnormv(Data$y, mu, sigma2, log=TRUE))
    ### Log-Posterior
    LP <- LL + alpha.prior + phi.prior + omega.prior + theta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, ynew,</pre>
         sigma2.new), yhat=rnormv(length(mu), mu, sigma2), parm=parm)
    return(Modelout)
    }
```

7.4. Initial Values

```
Initial. Values \leftarrow c(rep(0,2), rep(0.5,2))
```

8. Autoregressive Moving Average, ARMA(1,1)

8.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma^{2}), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} = \alpha + \phi \mathbf{y}_{T} + \theta \epsilon_{T}$$

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1} + \theta \epsilon_{t-1}$$

$$\epsilon_{t} = \mathbf{y}_{t} - \mu_{t}$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\sigma \sim \mathcal{HC}(25)$$

$$\theta \sim \mathcal{N}(0, 1000)$$

```
y \leftarrow c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91,
    2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28,
    1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03,
    -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02,
    0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36,
    1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70,
    0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83,
    0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24,
    0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40,
    -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
    0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17)
T <- length(y)
mon.names <- c("LP", "sigma", "ynew")</pre>
```

```
parm.names <- c("alpha", "phi", "sigma", "theta")</pre>
PGF <- function(Data) return(c(rnormv(2,0,1000), log(rhalfcauchy(1,25)),
    rnormv(1,0,1000)))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,</pre>
8.3. Model
Model <- function(parm, Data)</pre>
     {
    ### Parameters
     alpha <- parm[1]; phi <- parm[2]; theta <- parm[3]</pre>
    sigma <- exp(parm[4])</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    theta.prior <- dnormv(theta, 0, 1000, log=TRUE)
    ### Log-Likelihood
    mu <- c(alpha, alpha + phi*Data$y[-Data$T])</pre>
    epsilon <- Data$y - mu
    mu <- c(mu[1], mu[-1] + theta * epsilon[-Data$T])</pre>
    ynew <- rnorm(1, alpha + phi*Data$y[Data$T] + theta*epsilon[Data$T],</pre>
    LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + phi.prior + sigma.prior + theta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, sigma, ynew),</pre>
         yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

8.4. Initial Values

Initial. Values $\leftarrow c(rep(0,2), 0, log(1))$

9. Beta Regression

9.1. Form

$$\mathbf{y} \sim \mathcal{BETA}(a, b)$$
$$a = \mu \phi$$

$$b = (1 - \mu)\phi$$

$$\mu = \Phi(\beta_1 + \beta_2 \mathbf{x}), \quad \mu \in (0, 1)$$

$$\beta_j \sim \mathcal{N}(0, 10), \quad j = 1, \dots, J$$

$$\phi \sim \mathcal{HC}(25)$$

where Φ is the normal CDF.

9.2. Data

```
\label{eq:continuous_section} $$N <- 100 $$x <- runif(N) $$y <- rbeta(N, (0.5-0.2*x)*3, (1-(0.5-0.2*x))*3) mon.names <- "LP" parm.names <- c("beta[1]","beta[2]","log.phi") $$PGF <- function(Data) return(c(rnormv(2,0,10), log(rhalfcauchy(1,25)))) $$MyData <- list(PGF=PGF, mon.names=mon.names, parm.names=parm.names, x=x) $$y=y$
```

9.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:2]; phi <- exp(parm[3])
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 10, log=TRUE))</pre>
    phi.prior <- dhalfcauchy(phi, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- interval(pnorm(beta[1] + beta[2]*Data$x), 0.001, 0.999,</pre>
         reflect=FALSE)
    a <- mu * phi
    b <- (1 - mu) * phi
    LL <- sum(dbeta(Data$y, a, b, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + phi.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
         yhat=rbeta(length(mu), a, b), parm=parm)
    return(Modelout)
    }
```

9.4. Initial Values

Initial. Values $\leftarrow c(rep(0,2), log(0.01))$

10. Beta-Binomial

10.1. Form

```
\mathbf{y}_i \sim \mathcal{BIN}(\mathbf{n}_i, \pi_i), \quad i = 1, \dots, N
\pi_i \sim \mathcal{BETA}(\alpha, \beta) \in [0.001, 0.999]
```

10.2. Data

```
N <- 20
n <- round(runif(N, 50, 100))
y <- round(runif(N, 1, 10))
mon.names <- "LP"
parm.names <- as.parm.names(list(pi=rep(0,N)))
PGF <- function(Data) return(rbeta(Data$N,1,1))
MyData <- list(N=N, PGF=PGF, mon.names=mon.names, n=n, parm.names=parm.names, y=y)</pre>
```

10.3. Model

10.4. Initial Values

Initial.Values <- c(rep(0.5,N))</pre>

11. Binary Logit

11.1. Form

 $\mathbf{y} \sim \mathcal{BERN}(\eta)$

$$\eta = \frac{1}{1 + \exp(-\mu)}$$
$$\mu = \mathbf{X}\beta$$
$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

11.2. Data

```
data(demonsnacks)
J <- 3
y <- ifelse(demonsnacks$Calories <= 137, 0, 1)
X <- cbind(1, as.matrix(demonsnacks[,c(7,8)]))
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=rep(0,J)))
PGF <- function(Data) return(rnormv(Data$J,0,1000))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names, parm.names=parm.names, y=y)</pre>
```

11.3. Model

11.4. Initial Values

Initial.Values <- rep(0,J)</pre>

12. Binary Log-Log Link Mixture

A weighted mixture of the log-log and complementary log-log link functions is used, where α is the weight. Since the log-log and complementary log-log link functions are asymmetric (as opposed to the symmetric logit and probit link functions), it may be unknown *a priori* whether the log-log or complementary log-log will perform better.

12.1. Form

$$\mathbf{y} \sim \mathcal{BERN}(\eta)$$

$$\eta = \alpha \exp(-\exp(\mu)) + (1 - \alpha)(1 - \exp(-\exp(\mu)))$$

$$\mu = \mathbf{X}\beta$$

$$\alpha \sim \mathcal{U}(0, 1)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

12.2. Data

```
Model <- function(parm, Data)
    {
     ### Parameters
     parm[Data$J+1] <- alpha <- interval(parm[Data$J+1], -700, 700)
     beta <- parm[1:Data$J]
     ### Log(Prior Densities)
     alpha.prior <- dunif(alpha, 0, 1, log=TRUE)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
     ### Log-Likelihood
     mu <- tcrossprod(Data$X, t(beta))
     eta <- alpha*invloglog(mu) + (1-alpha)*invcloglog(mu)
     LL <- sum(dbern(Data$y, eta, log=TRUE))
     ### Log-Posterior
     LP <- LL + alpha.prior + beta.prior
     Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,alpha),</pre>
```

```
yhat=rbern(length(eta), eta), parm=parm)
return(Modelout)
}
```

12.4. Initial Values

Initial.Values <- c(rep(0,J), 0)</pre>

13. Binary Probit

13.1. Form

$$\mathbf{y} \sim \mathcal{BERN}(\mathbf{p})$$

$$\mathbf{p} = \phi(\mu)$$

$$\mu = \mathbf{X}\beta \in [-10, 10]$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

where ϕ is the CDF of the standard normal distribution, and J=3.

13.2. Data

```
data(demonsnacks)
J <- 3
y <- ifelse(demonsnacks$Calories <= 137, 0, 1)
X <- cbind(1, as.matrix(demonsnacks[,c(7,8)]))
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=rep(0,J)))
PGF <- function(Data) return(rnormv(Data$J,0,1000))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names, parm.names=parm.names, y=y)</pre>
```

```
Model <- function(parm, Data)
    {
     ### Parameters
     beta <- parm[1:Data$J]
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
     ### Log-Likelihood
     mu <- tcrossprod(Data$X, t(beta))</pre>
```

13.4. Initial Values

Initial.Values <- rep(0,J)</pre>

14. Binomial Logit

14.1. Form

$$\mathbf{y} \sim \mathcal{BIN}(\mathbf{p}, \mathbf{n})$$

$$\mathbf{p} = \frac{1}{1 + \exp(-\mu)}$$

$$\mu = \beta_1 + \beta_2 \mathbf{x}$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

14.2. Data

```
Model <- function(parm, Data)
{
    ### Parameters
    beta <- parm[1:Data$J]</pre>
```

14.4. Initial Values

Initial.Values <- rep(0,J)</pre>

15. Binomial Probit

15.1. Form

$$\mathbf{y} \sim \mathcal{BIN}(\mathbf{p}, \mathbf{n})$$

$$\mathbf{p} = \phi(\mu)$$

$$\mu = \beta_1 + \beta_2 \mathbf{x} \in [-10, 10]$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

where ϕ is the CDF of the standard normal distribution, and J=2.

15.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:Data$J]
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- beta[1] + beta[2]*Data$x</pre>
    mu <- interval(mu, -10, 10, reflect=FALSE)</pre>
    p <- pnorm(mu)
    LL <- sum(dbinom(Data$y, Data$n, p, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
         yhat=rbinom(length(p), Data$n, p), parm=parm)
    return(Modelout)
    }
```

15.4. Initial Values

Initial.Values <- rep(0,J)</pre>

16. Cluster Analysis, Confirmatory (CCA)

This is a parametric, model-based, cluster analysis, also called a finite mixture model or latent class cluster analysis, where the number of clusters C is fixed. When the number of clusters is unknown, exploratory cluster analysis should be used (see section 17).

16.1. Form

$$\mathbf{Y}_{i,j} \sim \mathcal{N}(\mu_{\theta[i],j}, \sigma_{\theta[i]}^2), \quad i = 1, \dots, N, \quad j = 1, \dots, J$$

$$\theta_i = \text{Max}(\mathbf{p}_{i,1:C})$$

$$\mathbf{p}_{i,c} = \frac{\delta_{i,c}}{\sum_{c=1}^{C} \delta_{i,c}}$$

$$\pi_{1:C} \sim \mathcal{D}(\alpha_{1:C})$$

$$\pi_c = \frac{\sum_{i=1}^{N} \delta_{i,c}}{\sum \delta}$$

$$\alpha_c = 1$$

$$\delta_{i,C} = 1$$

$$\delta_{i,C} = 1$$

$$\delta_{i,c} \sim \mathcal{N}(\log(\frac{1}{C}), 1000) \in [\exp(-10), \exp(10)], \quad c = 1, \dots, (C-1)$$

```
\mu_{c,j} \sim \mathcal{N}(0, \nu_j^2)
\sigma_c \sim \mathcal{HC}(25)
\nu_j \sim \mathcal{HC}(25)
```

```
C <- 3 #Number of clusters
alpha <- rep(1,C) #Prior probability of cluster proportion
# Create a Y matrix
n <- 100; N <- 15 \#Full sample; model sample
J <- 5 #Number of predictor variables
cluster <- rcat(n, rep(1,C))</pre>
centers <- matrix(runif(C*J, 0, 10), C, J)</pre>
Y.Full <- matrix(0, n, J)
for (i in 1:n) {for (j in 1:J)
    {Y.Full[i,j] <- rnorm(1,centers[cluster[i],j],1)}}
mean.temp <- colMeans(Y.Full)</pre>
sigma.temp <- apply(Y.Full,2,sd)</pre>
centers.cs <- (centers - matrix(rep(mean.temp,C), C, J, byrow=TRUE)) /</pre>
     (2 * matrix(rep(sigma.temp,C), C, J, byrow=TRUE))
for (j in 1:J) {Y.Full[,j] <- scale(Y.Full[,j],2)}
#summary(Y.Full)
MySample <- sample(1:n, N)
Y <- Y.Full[MySample,]
mon.names <- c("LP", as.parm.names(list(nu=rep(0,J), pi=rep(0,C),</pre>
    sigma=rep(0,C), theta=rep(0,N))))
parm.names <- as.parm.names(list(log.delta=matrix(0,N,C-1), mu=matrix(0,C,J),</pre>
    log.nu=rep(0,J), log.sigma=rep(0,C)))
PGF <- function(Data) return(c(log(rtrunc(Data$N*(Data$C-1), "norm",
    a=-exp(-10), b=exp(10), log(1/Data\$C), sqrt(1000)),
    rnorm(Data$C*Data$J,0,
    matrix(rhalfcauchy(Data$J,25),Data$C,Data$J,byrow=TRUE)),
    log(rhalfcauchy(Data$J+Data$C,25))))
MyData <- list(C=C, J=J, N=N, PGF=PGF, Y=Y, alpha=alpha,
    mon.names=mon.names, parm.names=parm.names)
16.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    delta <- interval(parm[grep("log.delta", Data$parm.names)], -10, 10)
    parm[grep("log.delta", Data$parm.names)] <- delta</pre>
    delta <- matrix(c(exp(delta), rep(1, Data$N)), Data$N, Data$C)</pre>
    mu <- matrix(parm[grep("mu", Data$parm.names)], Data$C, Data$J)</pre>
```

```
nu <- exp(parm[grep("log.nu",Data$parm.names)])</pre>
pi <- colSums(delta) / sum(delta)</pre>
sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
### Log(Prior Densities)
delta.prior <- sum(dtrunc(delta, "norm", a=exp(-10), b=exp(10),
    mean=log(1/Data$C), sd=sqrt(1000), log=TRUE))
mu.prior <- sum(dnorm(mu, 0, matrix(rep(nu,Data$C), Data$C,</pre>
    Data$J, byrow=TRUE), log=TRUE))
nu.prior <- sum(dhalfcauchy(nu, 25, log=TRUE))</pre>
pi.prior <- ddirichlet(pi, Data$alpha, log=TRUE)</pre>
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
### Log-Likelihood
p <- delta / rowSums(delta)</pre>
theta <- max.col(p)
LL <- sum(dnorm(Data$Y, mu[theta,], sigma[theta], log=TRUE))
### Log-Posterior
LP <- LL + delta.prior + mu.prior + nu.prior + pi.prior +
    sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,nu,pi,sigma,theta),</pre>
    yhat=rnorm(length(mu[theta,]), mu[theta,], sigma[theta]),
    parm=parm)
return(Modelout)
}
```

16.4. Initial Values

Initial. Values $\leftarrow c(\text{runif}(N*(C-1),-1,1), \text{rep}(0,C*J), \text{rep}(0,J), \text{rep}(0,C))$

17. Cluster Analysis, Exploratory (ECA)

In "exploratory cluster analysis", the optimal number of clusters C is unknown before the model update. This is a nonparametric, model-based, infinite mixture model that uses truncated stick-breaking within a truncated Dirichlet process. The user must specify the maximum number of clusters (mixture components), C to explore, where C is discrete, greater than one, and less than the number of records, N. The records in the $N \times J$ matrix \mathbf{Y} are clustered, where J is the number of predictors.

17.1. Form

$$\mathbf{Y}_{i,j} \sim \mathcal{N}(\mu_{\theta[i],j}, \sigma_{\theta[i]}^2), \quad i = 1, \dots, N, \quad j = 1, \dots, J$$

$$\theta_i \sim \mathcal{CAT}(\pi_{i,1:C})$$

$$\mathbf{p}_{i,c} = \frac{\delta_{i,c}}{\sum_{c=1}^{C} \delta_{i,c}}$$

$$\pi \sim \operatorname{Stick}(\gamma)$$

$$\delta_{i,C} = 1$$

$$\delta_{i,c} \sim \mathcal{N}(\log(\frac{1}{C}), 1000) \in [\exp(-10), \exp(10)], \quad c = 1, \dots, (C-1)$$

$$\mu_{c,j} \sim \mathcal{N}(0, \nu_j^2)$$

$$\alpha \sim \mathcal{HC}(25)$$

$$\beta \sim \mathcal{HC}(25)$$

$$\gamma \sim \mathcal{G}(\alpha, \beta)$$

$$\sigma_c \sim \mathcal{HC}(25)$$

$$\nu_j \sim \mathcal{HC}(25)$$

```
C <- 3 #Number of clusters for simulated DGP
# Create a Y matrix
n <- 100; N <- 15 #Full sample; model sample
J <- 5 #Number of predictor variables
cluster <- round(runif(n,0.5,C+0.49))</pre>
centers <- matrix(runif(C*J, 0, 10), C, J)</pre>
Y.Full <- matrix(0, n, J)
for (i in 1:n) {for (j in 1:J)
    {Y.Full[i,j] <- rnorm(1,centers[cluster[i],j],1)}}
mean.temp <- colMeans(Y.Full)</pre>
sigma.temp <- apply(Y.Full,2,sd)
centers.cs <- (centers - matrix(rep(mean.temp,C), C, J, byrow=TRUE)) /</pre>
     (2 * matrix(rep(sigma.temp,C), C, J, byrow=TRUE))
for (j in 1:J) {Y.Full[,j] <- scale(Y.Full[,j],2)}</pre>
MySample <- sample(1:n, N)</pre>
Y <- Y.Full[MySample,]
C <- 10 #Number of clusters to explore
mon.names <- c("LP", as.parm.names(list(nu=rep(0,J), pi=rep(0,C),</pre>
    sigma=rep(0,C), theta=rep(0,N))))
parm.names <- as.parm.names(list(log.delta=matrix(0,N,C-1),</pre>
    mu=matrix(0,C,J), log.nu=rep(0,J), log.sigma=rep(0,C),
    lambda=rep(0,C-1), log.alpha=0, log.beta=0, log.gamma=0))
PGF <- function(Data) return(c(log(rtrunc(Data$N*(Data$C-1), "norm",
    a=-exp(-10), b=exp(10), log(1/Data$C), sqrt(1000))),
    rnorm(Data$C*Data$J,0,
    matrix(rhalfcauchy(Data$J,25),Data$C,Data$J,byrow=TRUE)),
    log(rhalfcauchy(Data$J+Data$C,25)), runif(Data$C-1,1e-5,1-1e-5),
    log(rhalfcauchy(2,25)), log(rgamma(1,rhalfcauchy(2,25)))))
MyData <- list(C=C, J=J, N=N, PGF=PGF, Y=Y, mon.names=mon.names,
```

parm.names=parm.names)

```
Model <- function(parm, Data)</pre>
    ### Hyperhyperparameters
     alpha <- exp(parm[grep("log.alpha", Data$parm.names)])</pre>
     beta <- exp(parm[grep("log.beta", Data$parm.names)])</pre>
     ### Hyperparameters
     gamma <- exp(parm[grep("log.gamma", Data$parm.names)])</pre>
    nu <- exp(parm[grep("log.nu",Data$parm.names)])</pre>
    ### Parameters
     delta <- interval(parm[grep("log.delta", Data$parm.names)], -10, 10)
     parm[grep("log.delta", Data$parm.names)] <- delta</pre>
     delta <- matrix(c(exp(delta), rep(1, Data$N)), Data$N, Data$C)</pre>
    lambda <- interval(parm[grep("lambda", Data$parm.names)], 1e-5, 1-1e-5)</pre>
    mu <- matrix(parm[grep("mu", Data$parm.names)], Data$C, Data$J)</pre>
    pi <- as.vector(Stick(lambda))</pre>
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    p <- delta / rowSums(delta)</pre>
     theta <- max.col(p)
    ### Log(Hyperhyperprior Densities)
     alpha.prior <- dhalfcauchy(alpha, 25, log=TRUE)</pre>
    beta.prior <- dhalfcauchy(beta, 25, log=TRUE)</pre>
     ### Log(Hyperprior Densities)
     gamma.prior <- dgamma(gamma, alpha, beta, log=TRUE)</pre>
    nu.prior <- sum(dhalfcauchy(nu, 25, log=TRUE))</pre>
    ### Log(Prior Densities)
    delta.prior <- sum(dtrunc(delta, "norm", a=exp(-10), b=exp(10),
         mean=log(1/Data$C), sd=sqrt(1000), log=TRUE))
    mu.prior <- sum(dnorm(mu, 0, matrix(rep(nu,Data$C), Data$C,</pre>
         Data$J, byrow=TRUE), log=TRUE))
    pi.prior <- dStick(pi, gamma, log=TRUE)</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
     theta.prior <- sum(dcat(theta, pi, log=TRUE))</pre>
    ### Log-Likelihood
    LL <- sum(dnorm(Data$Y, mu[theta,], sigma[theta], log=TRUE))
     ### Log-Posterior
    LP <- LL + delta.prior + mu.prior + nu.prior + pi.prior +
         alpha.prior + beta.prior + gamma.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,nu,pi,sigma,theta),</pre>
         yhat=rnorm(length(mu[theta,]), mu[theta,], sigma[theta]),
         parm=parm)
    return(Modelout)
    }
```

17.4. Initial Values

Initial.Values <- c(runif(N*(C-1),-1,1), rep(0,C*J), rep(0,J), rep(0,C), rbeta(C-1,1,2), rep(1,3))

18. Conditional Autoregression (CAR), Poisson

This CAR example is a slightly modified form of example 7.3 (Model A) in Congdon (2003). The Scottish lip cancer data also appears in the WinBUGS (Spiegelhalter, Thomas, Best, and Lunn 2003) examples and is a widely analyzed example. The data \mathbf{y} consists of counts for $i = 1, \ldots, 56$ counties in Scotland. A single predictor \mathbf{x} is provided. The errors, ϵ , are allowed to include spatial effects as smoothing by spatial effects from areal neighbors. The vector ϵ_{μ} is the mean of each area's error, and is a weighted average of errors in contiguous areas. Areal neighbors are indicated in adjacency matrix \mathbf{A} .

18.1. Form

$$\mathbf{y} \sim \mathcal{P}(\lambda)$$

$$\lambda = \exp(\log(\mathbf{E}) + \beta_1 + \beta_2 \mathbf{x} + \epsilon)$$

$$\epsilon \sim \mathcal{N}(\epsilon_{\mu}, \sigma^2)$$

$$\epsilon_{\mu[i]} = \rho \sum_{j=1}^{J} \mathbf{A}_{i,j} \epsilon_j, \quad i = 1, \dots, N$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\rho \sim \mathcal{U}(-1, 1)$$

$$\sigma \sim \mathcal{HC}(25)$$

```
A[2,c(7,10)] \leftarrow 1 #Area 2 is adjacent to areas 7 and 10
A[3,c(6,12)] \leftarrow 1; A[4,c(18,20,28)] \leftarrow 1; A[5,c(1,11,12,13,19)] \leftarrow 1
A[6,c(3,8)] \leftarrow 1; A[7,c(2,10,13,16,17)] \leftarrow 1; A[8,6] \leftarrow 1
A[9,c(1,11,17,19,23,29)] \leftarrow 1; A[10,c(2,7,16,22)] \leftarrow 1
A[11,c(1,5,9,12)] \leftarrow 1; A[12,c(3,5,11)] \leftarrow 1; A[13,c(5,7,17,19)] \leftarrow 1
A[14,c(31,32,35)] \leftarrow 1; A[15,c(25,29,50)] \leftarrow 1
A[16,c(7,10,17,21,22,29)] \leftarrow 1; A[17,c(7,9,13,16,19,29)] \leftarrow 1
A[18,c(4,20,28,33,55,56)] \leftarrow 1; A[19,c(1,5,9,13,17)] \leftarrow 1
A[20,c(4,18,55)] \leftarrow 1; A[21,c(16,29,50)] \leftarrow 1; A[22,c(10,16)] \leftarrow 1
A[23,c(9,29,34,36,37,39)] \leftarrow 1; A[24,c(27,30,31,44,47,48,55,56)] \leftarrow 1
A[25,c(15,26,29)] \leftarrow 1; A[26,c(25,29,42,43)] \leftarrow 1
A[27,c(24,31,32,55)] \leftarrow 1; A[28,c(4,18,33,45)] \leftarrow 1
A[29,c(9,15,16,17,21,23,25,26,34,43,50)] <-1
A[30,c(24,38,42,44,45,56)] \leftarrow 1; A[31,c(14,24,27,32,35,46,47)] \leftarrow 1
A[32,c(14,27,31,35)] \leftarrow 1; A[33,c(18,28,45,56)] \leftarrow 1
A[34,c(23,29,39,40,42,43,51,52,54)] \leftarrow 1; A[35,c(14,31,32,37,46)] \leftarrow 1
A[36,c(23,37,39,41)] \leftarrow 1; A[37,c(23,35,36,41,46)] \leftarrow 1
A[38,c(30,42,44,49,51,54)] \leftarrow 1; A[39,c(23,34,36,40,41)] \leftarrow 1
A[40,c(34,39,41,49,52)] \leftarrow 1; A[41,c(36,37,39,40,46,49,53)] \leftarrow 1
A[42,c(26,30,34,38,43,51)] \leftarrow 1; A[43,c(26,29,34,42)] \leftarrow 1
A[44,c(24,30,38,48,49)] \leftarrow 1; A[45,c(28,30,33,56)] \leftarrow 1
A[46,c(31,35,37,41,47,53)] \leftarrow 1; A[47,c(24,31,46,48,49,53)] \leftarrow 1
A[48,c(24,44,47,49)] \leftarrow 1; A[49,c(38,40,41,44,47,48,52,53,54)] \leftarrow 1
A[50,c(15,21,29)] \leftarrow 1; A[51,c(34,38,42,54)] \leftarrow 1
A[52,c(34,40,49,54)] \leftarrow 1; A[53,c(41,46,47,49)] \leftarrow 1
A[54,c(34,38,49,51,52)] \leftarrow 1; A[55,c(18,20,24,27,56)] \leftarrow 1
A[56,c(18,24,30,33,45,55)] <- 1
mon.names <- c("LP", "sigma")</pre>
parm.names <- as.parm.names(list(beta=rep(0,2), epsilon=rep(0,N), rho=0,
     log.sigma=0))
PGF <- function(Data) return(c(rnormv(2,0,1000), rnorm(Data$N,0,1),
     runif(1,-1,1), log(rhalfcauchy(1,25))))
MyData <- list(A=A, E=E, N=N, PGF=PGF, mon.names=mon.names,
     parm.names=parm.names, x=x, y=y)
18.3. Model
Model <- function(parm, Data)</pre>
     {
     ### Parameters
     beta <- parm[1:2]
     epsilon <- parm[grep("epsilon", Data$parm.names)]</pre>
     rho <- interval(parm[grep("rho", Data$parm.names)], -1, 1)</pre>
     parm[grep("rho", Data$parm.names)] <- rho</pre>
     epsilon.mu <- rho * rowSums(epsilon * Data$A)</pre>
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
     ### Log(Prior Densities)
```

18.4. Initial Values

Initial. Values $\leftarrow c(rep(0,2), rep(0,N), 0, 0)$

19. Conditional Predictive Ordinate

For a more complete introduction to the conditional predictive ordinate (CPO), see the vignette entitled "Bayesian Inference". Following is a brief guide to the applied use of CPO.

To include CPO in any model that is to be updated with MCMC, calculate and monitor the record-level inverse of the likelihood, $InvL_i$ for records i = 1, ..., N. CPO_i is the inverse of the posterior mean of $InvL_i$. The inverse CPO_i , or $ICPO_i$, is the posterior mean of $InvL_i$. $ICPO_i$ larger than 40 can be considered as possible outliers, and higher than 70 as extreme values.

Here, CPO is added to the linear regression example in section 40. In this data, record 6 is a possible outlier, and record 8 is an extreme value.

19.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

```
data(demonsnacks)
y <- log(demonsnacks$Calories)
X <- cbind(1, as.matrix(demonsnacks[,c(7,8,10)]))
J <- ncol(X)</pre>
```

```
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}</pre>
mon.names <- c("LP", "sigma", as.parm.names(list(InvL=rep(0,N))))</pre>
parm.names <- as.parm.names(list(beta=rep(0,J), log.sigma=0))</pre>
PGF <- function(Data) return(c(rnormv(Data$J,0,1000),
     log(rhalfcauchy(1,25))))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names,</pre>
    parm.names=parm.names, y=y)
19.3. Model
Model <- function(parm, Data)</pre>
     {
     ### Parameters
    beta <- parm[1:Data$J]
     sigma <- exp(parm[Data$J+1])</pre>
     ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
     sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta))</pre>
    LL <- dnorm(Data$y, mu, sigma, log=TRUE)
    InvL \leftarrow 1 / exp(LL)
    LL <- sum(LL)
    ### Log-Posterior
    LP <- LL + beta.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma,InvL),</pre>
         yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

19.4. Initial Values

Initial.Values <- c(rep(0,J), log(1))</pre>

20. Contingency Table

The two-way contingency table, matrix \mathbf{Y} , can easily be extended to more dimensions. For this example, it is vectorized as y, and used like an ANOVA data set. Contingency table \mathbf{Y} has \mathbf{J} rows and \mathbf{K} columns. The cell counts are fit with Poisson regression, according to intercept α , main effects β_j for each row, main effects γ_k for each column, and interaction effects $\delta_{j,k}$ for dependence effects. An omnibus (all cells) test of independence is done by estimating two models (one with δ , and one without), and a large enough Bayes Factor indicates a violation of independence when the model with δ fits better than the model without δ . In an ANOVA-like style, main effects contrasts can be used to distinguish rows or groups of rows from each other, as well as with columns. Likewise, interaction effects contrasts can be used

to test independence in groups of $\delta_{j,k}$ elements. Finally, single-cell interactions can be used to indicate violations of independence for a given cell, such as when zero is not within its 95% probability interval.

20.1. Form

$$\mathbf{Y}_{j,k} \sim \mathcal{P}(\lambda_{j,k}), \quad j = 1, \dots, J, \quad k = 1, \dots, K$$

$$\lambda_{j,k} = \exp(\alpha + \beta_j + \gamma_k + \delta_{j,k}), \quad j = 1, \dots, J, \quad k = 1, \dots, K$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\beta_j \sim \mathcal{N}(0, \beta_\sigma^2), \quad j = 1, \dots, J$$

$$\beta_J = -\sum_{j=1}^{J-1} \beta_j$$

$$\beta_\sigma \sim \mathcal{HC}(25)$$

$$\gamma_k \sim \mathcal{N}(0, \gamma_\sigma^2), \quad k = 1, \dots, K$$

$$\gamma_K = -\sum_{k=1}^{K-1} \gamma_k$$

$$\gamma_\sigma \sim \mathcal{HC}(25)$$

$$\delta_{j,k} \sim \mathcal{N}(0, \delta_\sigma^2)$$

$$\delta_{J,K} = -\sum_{j=1}^{J-1} \delta_{-J,-K}$$

$$\delta_\sigma \sim \mathcal{HC}(25)$$

```
J <- 3 #Rows
K <- 3 #Columns
log.alpha <- log(runif(1, 100, 200))
beta <- rnorm(J-1, 3); beta <- c(beta, -sum(beta))
gamma <- rnorm(J-1, 3); gamma <- c(gamma, -sum(gamma))</pre>
delta <- rnorm(J*K-1, 2); delta <- c(delta, -sum(delta))
Y <- matrix(exp(log.alpha), J, K) + matrix(beta, J, K, byrow=TRUE) +
    matrix(gamma, J, K) + matrix(delta, J, K) +
    matrix(rnorm(J*K,0,0.1), J, K)
Y \leftarrow round(Y)
y <- as.vector(Y)
N <- length(y) #Cells
r \leftarrow rep(1:J, N/J)
co \leftarrow rep(1,K)
for (k in 2:K) \{co \leftarrow c(co, rep(k, K))\}
mon.names <- c("LP","beta.sigma","gamma.sigma","delta.sigma")</pre>
```

```
parm.names <- as.parm.names(list(log.alpha=0, beta=rep(0,J-1),</pre>
    gamma=rep(0,K-1), log.b.sigma=0, log.g.sigma=0, log.d.sigma=0,
    delta=rep(0,J*K-1)))
PGF <- function(Data) return(c(log(rnorm(1,mean(Y),1)),
    rnorm(Data$J-1,0,rhalfcauchy(1,5)),
    rnorm(Data$K-1,0,rhalfcauchy(1,5)),
    log(rhalfcauchy(3,5)), rnorm(Data$J*Data$K-1,0,rhalfcauchy(1,5))))
MyData <- list(J=J, K=K, N=N, PGF=PGF, co=co, mon.names=mon.names,
    parm.names=parm.names, r=r, y=y)
20.3. Model
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    beta.sigma <- exp(parm[grep("log.b.sigma", Data$parm.names)])</pre>
    gamma.sigma <- exp(parm[grep("log.g.sigma", Data$parm.names)])</pre>
    delta.sigma <- exp(parm[grep("log.d.sigma", Data$parm.names)])</pre>
    alpha <- exp(parm[grep("log.alpha", Data$parm.names)])</pre>
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    beta <- c(beta, -sum(beta))</pre>
                                      gamma <- parm[grep("gamma", Data$parm.names)]</pre>
                                          delta <- parm[grep("delta", Data$parm.names)]</pre>
    gamma <- c(gamma, -sum(gamma))
    c(delta, -sum(delta))
         delta <- matrix(delta, Data$J, Data$K)</pre>
    ### Log(Hyperprior Densities)
    beta.sigma.prior <- dhalfcauchy(beta.sigma, 25, log=TRUE)</pre>
    gamma.sigma.prior <- dhalfcauchy(gamma.sigma, 25, log=TRUE)</pre>
    delta.sigma.prior <- dhalfcauchy(delta.sigma, 25, log=TRUE)</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    beta.prior <- sum(dnorm(beta, 0, beta.sigma, log=TRUE))</pre>
    gamma.prior <- sum(dnorm(gamma, 0, gamma.sigma, log=TRUE))</pre>
    delta.prior <- sum(dnorm(delta, 0, delta.sigma, log=TRUE))</pre>
    ### Log-Likelihood
    lambda <- exp(alpha + beta[Data$r] + gamma[Data$co] +</pre>
         diag(delta[Data$r,Data$co]))
    LL <- sum(dpois(Data$y, lambda, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + beta.sigma.prior +
         gamma.prior + gamma.sigma.prior + delta.prior +
         delta.sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, beta.sigma,
         gamma.sigma, delta.sigma), yhat=rpois(length(lambda), lambda),
         parm=parm)
```

return(Modelout)

}

20.4. Initial Values

Initial.Values <- c(log(mean(y)), rep(0,J-1), rep(0,K-1), rep(0,3), rep(0,J*K-1))

21. Covariance Separation Strategy

A Seemingly Unrelated Regression (SUR) model is used to provide an example of a flexible way to estimate covariance or precision matrices with the "separation strategy" decomposition of Barnard, McCulloch, and Meng (2000). For more information on SUR models, see section 70.

The most common way of specifying a covariance matrix, such as for the multivariate normal distribution, may be with the conjugate inverse Wishart distribution. Alternatively, the conjugate Wishart distribution is often used for a precision matrix. The Wishart and inverse Wishart distributions, however, do not always perform well, due to only one parameter for variability, and usually in the case of small sample sizes or when its dimension approaches the sample size. There are several alternatives. This example decomposes a covariance matrix into a standard deviation vector and a correlation matrix, each of which are easy to understand (as opposed to setting priors on eigenvalues). A precision matrix may be decomposed similarly, though the separated components are interpreted differently.

Barnard et al. (2000) prefer to update the covariance separation strategy with Gibbs sampling rather than Metropolis-Hastings, though the form presented here works well in testing with Adaptive MCMC.

$$\mathbf{Y}_{t,j} \sim \mathcal{N}_{J}(\mu_{t,j}, \Sigma), \quad t = 1, \dots, T; \quad j = 1, \dots, J$$

$$\mu_{t,1} = \alpha_{1} + \alpha_{2}\mathbf{X}_{t-1,1} + \alpha_{3}\mathbf{X}_{t-1,2}, \quad t = 2, \dots, T$$

$$\mu_{t,2} = \beta_{1} + \beta_{2}\mathbf{X}_{t-1,3} + \beta_{3}\mathbf{X}_{t-1,4}, \quad t = 2, \dots, T$$

$$\Sigma = \mathbf{SRS}$$

$$\alpha_{k} \sim \mathcal{N}(0, 1000), \quad k = 1, \dots, K$$

$$\beta_{k} \sim \mathcal{N}(0, 1000), \quad k = 1, \dots, K$$

$$\mathbf{R}_{i,j} \sim \mathcal{N}(\rho_{\mu}, \rho_{\sigma}^{2}), \quad \mathbf{R}_{i,j} \in [-1, 1], \quad i = 1, \dots, J$$

$$\mathbf{S} = \sigma \mathbf{I}_{J}$$

$$\rho_{\mu} \sim \mathcal{N}(0, 2), \quad \in [-1, 1]$$

$$\rho_{\sigma} \sim \mathcal{HC}(25), \quad \in (0, 1000]$$

$$\sigma_{j} \sim \mathcal{N}(\sigma_{\mu}, \sigma_{\sigma})$$

```
\sigma_{\mu} \sim \mathcal{HN}(1000), \in (0, 1000]
\sigma_{\sigma} \sim \mathcal{HC}(25)
```

```
T <- 20 #Time-periods
year <- c(1935,1936,1937,1938,1939,1940,1941,1942,1943,1944,1945,1946,
    1947, 1948, 1949, 1950, 1951, 1952, 1953, 1954)
IG <- c(33.1,45.0,77.2,44.6,48.1,74.4,113.0,91.9,61.3,56.8,93.6,159.9,
    147.2,146.3,98.3,93.5,135.2,157.3,179.5,189.6)
VG <- c(1170.6,2015.8,2803.3,2039.7,2256.2,2132.2,1834.1,1588.0,1749.4,
    1687.2,2007.7,2208.3,1656.7,1604.4,1431.8,1610.5,1819.4,2079.7,
    2371.6,2759.9)
CG \leftarrow c(97.8, 104.4, 118.0, 156.2, 172.6, 186.6, 220.9, 287.8, 319.9, 321.3, 319.6,
    346.0,456.4,543.4,618.3,647.4,671.3,726.1,800.3,888.9)
IW <- c(12.93,25.90,35.05,22.89,18.84,28.57,48.51,43.34,37.02,37.81,
    39.27,53.46,55.56,49.56,32.04,32.24,54.38,71.78,90.08,68.60)
VW <- c(191.5,516.0,729.0,560.4,519.9,628.5,537.1,561.2,617.2,626.7,
    737.2,760.5,581.4,662.3,583.8,635.2,723.8,864.1,1193.5,1188.9)
CW \leftarrow c(1.8, 0.8, 7.4, 18.1, 23.5, 26.5, 36.2, 60.8, 84.4, 91.2, 92.4, 86.0, 111.1,
    130.6,141.8,136.7,129.7,145.5,174.8,213.5)
J <- 2 #Number of dependent variables
Y <- matrix(c(IG,IW), T, J)
R <- diag(J)
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=rep(0,3), beta=rep(0,3),
    R=diag(J), rho.mu=0, rho.sigma=0, log.sigma=rep(0,J), sigma.mu=0,
    log.sig.sigma=0), uppertri=c(0,0,1,0,0,0,0,0)
PGF <- function(Data) return(c(rnormv(3,0,10), rnormv(3,0,10),
    runif(length(upper.triangle(diag(Data$J), diag=TRUE)), -1, 1),
    rtrunc(1, "norm", a=-1, b=1, mean=0, sd=2),
    log(rhalfcauchy(Data$J+1,25)), rhalfnorm(1, 10),
    log(rhalfcauchy(1,25))))
MyData <- list(J=J, PGF=PGF, T=T, Y=Y, CG=CG, CW=CW, IG=IG, IW=IW, VG=VG,
    VW=VW, mon.names=mon.names, parm.names=parm.names)
21.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Hyperparameters
    rho.mu <- interval(parm[grep("rho.mu", Data$parm.names)], -1, 1)</pre>
    parm[grep("rho.mu", Data$parm.names)] <- rho.mu</pre>
    rho.sigma <- interval(parm[grep("rho.sigma", Data$parm.names)],</pre>
         .Machine$double.eps, 1000)
```

```
parm[grep("rho.sigma", Data$parm.names)] <- rho.sigma</pre>
sigma.mu <- interval(parm[grep("sigma.mu", Data$parm.names)],</pre>
     .Machine$double.eps, 1000)
parm[grep("sigma.mu", Data$parm.names)] <- sigma.mu</pre>
sigma.sigma <- exp(parm[grep("log.sig.sigma", Data$parm.names)])</pre>
### Parameters
alpha <- parm[1:3]</pre>
beta <- parm[4:6]
R <- as.parm.matrix(R, Data$J, parm, Data, a=-1, b=1)</pre>
parm[grep("R", Data$parm.names)] <- upper.triangle(R, diag=TRUE)</pre>
sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
S <- diag(sigma)
Sigma <- as.symmetric.matrix(S %*% R %*% S)
### Log(Hyperprior Densities)
rho.mu.prior <- dtrunc(rho.mu, "norm", a=-1, b=1, mean=0, sd=2,
     log=TRUE)
rho.sigma.prior <- dhalfcauchy(rho.sigma, 25, log=TRUE)</pre>
sigma.mu.prior <- dhalfnorm(sigma.mu, 1000, log=TRUE)</pre>
sigma.sigma.prior <- dhalfcauchy(sigma.sigma, 25, log=TRUE)</pre>
### Log(Prior Densities)
alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
R.prior <- sum(dtrunc(upper.triangle(R, diag=TRUE), "norm",</pre>
     a=-1, b=1, mean=rho.mu, sd=rho.sigma, log=TRUE))
sigma.prior <- sum(dnorm(sigma, sigma.mu, sigma.sigma, log=TRUE))</pre>
### Log-Likelihood
mu <- Data$Y
mu[-1,1] \leftarrow alpha[1] + alpha[2]*Data$CG[-Data$T] +
     alpha[3]*Data$VG[-Data$T]
mu[-1,2] \leftarrow beta[1] + beta[2]*Data$CW[-Data$T] +
    beta[3] *Data$VW[-Data$T]
LL <- sum(dmvn(Data$Y[-1,], mu[-1,], Sigma, log=TRUE))</pre>
### Log-Posterior
LP <- LL + alpha.prior + beta.prior + R.prior + rho.mu.prior +
     rho.sigma.prior + sigma.prior + sigma.mu.prior +
     sigma.sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
     yhat=rmvn(nrow(mu), mu, Sigma), parm=parm)
return(Modelout)
}
```

21.4. Initial Values

```
Initial.Values <- c(rep(0,3), rep(0,3), upper.triangle(R, diag=TRUE), rep(0,2), rep(0,J), rep(1,2))
```

22. Discrete Choice, Conditional Logit

22.1. Form

$$\mathbf{y}_{i} \sim \mathcal{CAT}(\mathbf{p}_{i,1:J}), \quad i = 1, \dots, N, \quad j = 1, \dots, J$$

$$\mathbf{p}_{i,j} = \frac{\phi_{i,j}}{\sum_{j=1}^{J} \phi_{i,j}}$$

$$\phi = \exp(\mu)$$

$$\mu_{i,j} = \beta_{j,1:K} \mathbf{X}_{i,1:K} + \gamma \mathbf{Z}_{i,1:C} \in [-700, 700], \quad j = 1, \dots, (J-1)$$

$$\mu_{i,J} = \gamma \mathbf{Z}_{i,1:C}$$

$$\beta_{j,k} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1)$$

$$\gamma_{c} \sim \mathcal{N}(0, 1000)$$

```
y \leftarrow x01 \leftarrow x02 \leftarrow z01 \leftarrow z02 \leftarrow c(1:300)
y[1:100] <- 1
y[101:200] <- 2
y[201:300] <- 3
x01[1:100] <- rnorm(100, 25, 2.5)
x01[101:200] <- rnorm(100, 40, 4.0)
x01[201:300] \leftarrow rnorm(100, 35, 3.5)
x02[1:100] <- rnorm(100, 2.51, 0.25)
x02[101:200] <- rnorm(100, 2.01, 0.20)
x02[201:300] \leftarrow rnorm(100, 2.70, 0.27)
z01[1:100] <- 1
z01[101:200] <- 2
z01[201:300] <- 3
z02[1:100] <- 40
z02[101:200] <- 50
z02[201:300] <- 100
N <- length(y)
J <- 3 #Number of categories in y
K <- 3 #Number of individual attributes (including the intercept)
C <- 2 #Number of choice-based attributes (intercept is not included)
X \leftarrow \text{matrix}(c(\text{rep}(1,N),x01,x02),N,K) \text{ #Design matrix of individual attrib.}
Z \leftarrow matrix(c(z01,z02),N,C) #Design matrix of choice-based attributes
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=matrix(0,J-1,K), gamma=rep(0,C)))</pre>
PGF <- function(Data) return(c(rnormv((Data$J-1)*Data$K,0,10),
    rnormv(Data$C,0,10)))
MyData <- list(C=C, J=J, K=K, N=N, PGF=PGF, X=X, Z=Z, mon.names=mon.names,
```

parm.names=parm.names, y=y)

22.3. Model

```
Model <- function(parm, Data)</pre>
            ### Parameters
     beta <- matrix(parm[grep("beta", Data$parm.names)], Data$J-1, Data$K)</pre>
     gamma <- parm[grep("gamma", Data$parm.names)]</pre>
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
     gamma.prior <- sum(dnormv(gamma, 0, 1000, log=TRUE))</pre>
     ### Log-Likelihood
     mu <- matrix(rep(tcrossprod(gamma, Data$Z), Data$J), Data$N, Data$J)</pre>
     mu[,-Data$J] <- mu[,-Data$J] + tcrossprod(Data$X, beta)</pre>
     mu <- interval(mu, -700, 700, reflect=FALSE)</pre>
     phi <- exp(mu)</pre>
     p <- phi / rowSums(phi)</pre>
     LL <- sum(dcat(Data$y, p, log=TRUE))</pre>
     ### Log-Posterior
     LP <- LL + beta.prior + gamma.prior
     Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP, yhat=rcat(nrow(p), p),</pre>
         parm=parm)
     return(Modelout)
     }
```

22.4. Initial Values

Initial. Values $\leftarrow c(rep(0,(J-1)*K), rep(0,C))$

23. Discrete Choice, Mixed Logit

$$\mathbf{y}_{i} \sim \mathcal{CAT}(\mathbf{p}_{i,1:J}), \quad i = 1, \dots, N$$

$$\mathbf{p}_{i,j} = \frac{\phi_{i,j}}{\sum_{j=1}^{J} \phi_{i,j}}$$

$$\phi = \exp(\mu)$$

$$\mu_{i,j} = \beta_{j,1:K} \mathbf{X}_{i,1:K} + \gamma \mathbf{Z}_{i,1:C} \in [-700, 700], \quad j = 1, \dots, (J-1)$$

$$\mu_{i,J} = \gamma \mathbf{Z}_{i,1:C}$$

$$\beta_{j,k} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1)$$

$$\gamma_c \sim \mathcal{N}(\zeta_{\mu[c]}, \zeta_{\sigma[c]}^2)$$
$$\zeta_{\mu[c]} \sim \mathcal{N}(0, 1000)$$
$$\zeta_{\sigma[c]} \sim \mathcal{HC}(25)$$

23.2. Data

```
y \leftarrow x01 \leftarrow x02 \leftarrow z01 \leftarrow z02 \leftarrow c(1:300)
y[1:100] < -1
y[101:200] <- 2
y[201:300] <- 3
x01[1:100] <- rnorm(100, 25, 2.5)
x01[101:200] <- rnorm(100, 40, 4.0)
x01[201:300] \leftarrow rnorm(100, 35, 3.5)
x02[1:100] \leftarrow rnorm(100, 2.51, 0.25)
x02[101:200] <- rnorm(100, 2.01, 0.20)
x02[201:300] \leftarrow rnorm(100, 2.70, 0.27)
z01[1:100] <- 1
z01[101:200] <- 2
z01[201:300] <- 3
z02[1:100] <- 40
z02[101:200] <- 50
z02[201:300] <- 100
N <- length(y)
J <- 3 #Number of categories in y
K <- 3 #Number of individual attributes (including the intercept)</pre>
C <- 2 #Number of choice-based attributes (intercept is not included)
X \leftarrow \text{matrix}(c(\text{rep}(1,N),x01,x02),N,K) \text{ $\#$Design matrix of individual attrib.}
Z \leftarrow matrix(c(z01,z02),N,C) #Design matrix of choice-based attributes
mon.names <- c("LP", as.parm.names(list(zeta.sigma=rep(0,C))))</pre>
parm.names <- as.parm.names(list(beta=matrix(0,J-1,K), gamma=rep(0,C),</pre>
     zeta.mu=rep(0,C), log.zeta.sigma=rep(0,C)))
PGF <- function(Data) return(c(rnormv((Data$J-1)*Data$K,0,1000),
    rnorm(Data$N*Data$C,
    matrix(rnormv(Data$C,0,1000), Data$N, Data$C, byrow=TRUE),
    matrix(rhalfcauchy(Data$C,25), Data$N, Data$C, byrow=TRUE)),
    rnormv(Data$C,0,1000), log(rhalfcauchy(Data$C,25))))
MyData <- list(C=C, J=J, K=K, N=N, X=X, Z=Z, mon.names=mon.names,
    parm.names=parm.names, y=y)
```

23.3. Model

```
zeta.mu <- parm[grep("zeta.mu", Data$parm.names)]</pre>
zeta.sigma <- exp(parm[grep("log.zeta.sigma", Data$parm.names)])</pre>
### Log(Prior Densities)
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
gamma.prior <- sum(dnorm(gamma, matrix(zeta.mu, Data$N, Data$C,
     byrow=TRUE), matrix(zeta.sigma, Data$N, Data$C, byrow=TRUE),
log=TRUE))
zeta.mu.prior <- sum(dnormv(zeta.mu, 0, 1000, log=TRUE))</pre>
zeta.sigma.prior <- sum(dhalfcauchy(zeta.sigma, 25, log=TRUE))</pre>
### Log-Likelihood
mu <- matrix(rep(rowSums(gamma * Data$Z),Data$J), Data$N, Data$J)</pre>
mu[,-Data$J] <- tcrossprod(Data$X, beta) + gamma * Data$Z</pre>
mu <- interval(mu, -700, 700, reflect=FALSE)</pre>
phi <- exp(mu)
p <- phi / rowSums(phi)</pre>
LL <- sum(dcat(Data$y, p, log=TRUE))</pre>
### Log-Posterior
LP <- LL + beta.prior + gamma.prior + zeta.mu.prior + zeta.sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,zeta.sigma),</pre>
     yhat=rcat(nrow(p), p), parm=parm)
return(Modelout)
}
```

23.4. Initial Values

Initial. Values $\leftarrow c(rep(0,(J-1)*K), rep(0,N*C), rep(0,C), rep(0,C))$

24. Discrete Choice, Multinomial Probit

$$\mathbf{W}_{i,1:(J-1)} \sim \mathcal{N}_{J-1}(\mu_{i,1:(J-1)}, \Sigma), \quad i = 1, ..., N$$

$$\mathbf{W}_{i,j} \in \begin{cases} [0,10] & \text{if } \mathbf{y}_i = j \\ [-10,0] \end{cases}$$

$$\mu_{1:N,j} = \mathbf{X}\beta_{j,1:K} + \mathbf{Z}\gamma$$

$$\Sigma = \mathbf{U}^T \mathbf{U}$$

$$\beta_{j,k} \sim \mathcal{N}(0,10), \quad j = 1, ..., (J-1), \quad k = 1, ..., K$$

$$\gamma_c \sim \mathcal{N}(0,10), \quad c = 1, ..., C$$

$$\mathbf{U}_{j,k} \sim \mathcal{N}(0,1), \quad j = 1, ..., (J-1), \quad k = 1, ..., (J-1), \quad j \ge k, \quad j \ne k = 1$$

```
N <- 50
J <- 5 #Categories of y
K <- 8 #Number of columns in design matrix X
C <- 2 #Number of choice-based attributes
X \leftarrow matrix(runif(N*K,-2,2), N, K)
X[,1] <- 1
beta <- matrix(runif((J-1)*K), J-1, K)
gamma <- runif(C)</pre>
Z <- matrix(runif(N*C), N, C) #Design matrix of choice-based attributes
Z[,1] < -1
mu <- tcrossprod(X, beta) + as.vector(tcrossprod(Z, t(gamma)))</pre>
S \leftarrow diag(J-1)
u \leftarrow c(0, rnorm((J-2) + (factorial(J-1) /
     (factorial(J-1-2)*factorial(2))),0,1))
U \leftarrow diag(J-1)
U[upper.tri(U, diag=TRUE)] <- u</pre>
diag(U) <- exp(diag(U))</pre>
Sigma <- t(U) %*% U
Sigma[1,] <- Sigma[,1] <- U[1,]
mu <- tcrossprod(X, beta)</pre>
W \leftarrow rmvn(N, mu, Sigma) + matrix(rnorm(N*(J-1),0,0.1), N, J-1)
y <- max.col(cbind(W,0))</pre>
table(y)
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=matrix(0,(J-1),K), gamma=rep(0,C),</pre>
    U=U, W=matrix(0,N,J-1)), uppertri=c(0,0,1,0))
parm.names <- parm.names[-which(parm.names == "U[1,1]")]</pre>
PGF <- function(Data) {</pre>
    beta <- rnormv((Data$J-1)*Data$K,0,1)</pre>
     gamma <- rnormv(Data$C,0,1)</pre>
    U <- rnorm((Data$J-2) + (factorial(Data$J-1) /</pre>
         (factorial(Data$J-1-2)*factorial(2))),0,1)
    W <- matrix(runif(Data$N*(Data$J-1),-10,0), Data$N, Data$J-1)
    Y <- as.indicator.matrix(Data$y)
    W <- ifelse(Y[,-Data$J] == 1, abs(W), W)
    return(c(beta, gamma, U, as.vector(W)))}
MyData <- list(C=C, J=J, K=K, N=N, PGF=PGF, S=S, X=X, Z=Z,
    mon.names=mon.names, parm.names=parm.names, y=y)
24.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
```

```
beta <- matrix(parm[grep("beta", Data$parm.names)], Data$J-1, Data$K)</pre>
gamma <- parm[grep("gamma", Data$parm.names)]</pre>
u <- c(0, parm[grep("U", Data$parm.names)])</pre>
U <- diag(Data$J-1)</pre>
U[upper.tri(U, diag=TRUE)] <- u</pre>
diag(U) <- exp(diag(U))</pre>
Sigma <- t(U) %*% U
Sigma[1,] <- Sigma[,1] <- U[1,]
W <- matrix(parm[grep("W", Data$parm.names)], Data$N, Data$J-1)
Y <- as.indicator.matrix(Data$y)</pre>
W \leftarrow ifelse(Y[,-c(Data$J)] == 1, interval(W, 0, 10),
     interval(W, -10, 0))
parm[grep("W", Data$parm.names)] <- as.vector(W)</pre>
### Log(Prior Densities)
beta.prior <- sum(dnormv(beta, 0, 10, log=TRUE))</pre>
gamma.prior <- sum(dnormv(gamma, 0, 10, log=TRUE))</pre>
U.prior <- sum(dnorm(u[-length(u)], 0, 1, log=TRUE))</pre>
### Log-Likelihood
mu <- tcrossprod(Data$X, beta) +</pre>
    as.vector(tcrossprod(Data$Z, t(gamma)))
#eta <- exp(cbind(mu,0))</pre>
#p <- eta / rowSums(eta)</pre>
LL <- sum(dmvn(W, mu, Sigma, log=TRUE))
### Log-Posterior
LP <- LL + beta.prior + gamma.prior + U.prior</pre>
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
     yhat=max.col(cbind(rmvn(nrow(mu), mu, Sigma),0)), parm=parm)
return(Modelout)
}
```

24.4. Initial Values

Initial.Values <- GIV(Model, MyData, PGF=TRUE)</pre>

25. Distributed Lag, Koyck

This example applies Koyck or geometric distributed lags to k = 1, ..., K discrete events in covariate \mathbf{x} , transforming the covariate into a $N \times K$ matrix \mathbf{X} and creates a $N \times K$ lag matrix \mathbf{L} .

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1} + \sum_{k=1}^{K} \mathbf{X}_{t,k} \beta \lambda^{\mathbf{L}[t,k]}, \quad k = 1, \dots, K, \quad t = 2, \dots, T$$

$$\mu_{1} = \alpha + \sum_{k=1}^{K} \mathbf{X}_{1,k} \beta \lambda^{\mathbf{L}[1,k]}, \quad k = 1, \dots, K$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\beta \sim \mathcal{N}(0, 1000)$$

$$\lambda \sim \mathcal{U}(0, 1)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\sigma \sim \mathcal{HC}(25)$$

```
y \leftarrow c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91,
    2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28,
    1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03,
    -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02,
    0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36,
    1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70,
    0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83,
    0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24,
    0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40,
    -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
    0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17
```

}

```
T <- length(y)
K <- length(which(x != 0))</pre>
L <- X <- matrix(0, T, K)
for (i in 1:K) {
   X[which(x != 0)[i]:T,i] \leftarrow x[which(x != 0)[i]]
   L[(which(x != 0)[i]):T,i] \leftarrow 0:(T - which(x != 0)[i]))
mon.names <- "LP"
parm.names <- c("alpha","beta","lambda","phi","log.sigma")</pre>
PGF <- function(Data) return(c(rnormv(2,0,1000), runif(1),
   rnormv(1,0,1000), log(rhalfcauchy(1,25))))
MyData <- list(L=L, PGF=PGF, T=T, X=X, mon.names=mon.names,</pre>
   parm.names=parm.names, y=y)
25.3. Model
Model <- function(parm, Data)</pre>
   {
   ### Parameters
   alpha <- parm[1]; beta <- parm[2]</pre>
   parm[3] <- lambda <- interval(parm[3], 0, 1)</pre>
   phi <- parm[4]; sigma <- exp(parm[5])</pre>
   ### Log(Prior Densities)
   alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
   beta.prior <- dnormv(beta, 0, 1000, log=TRUE)
   lambda.prior <- dunif(lambda, 0, 1, log=TRUE)</pre>
   phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
   sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
   ### Log-Likelihood
   mu <- c(alpha, alpha + phi*Data$y[-Data$T]) +</pre>
      rowSums(Data$X * beta * lambda^Data$L)
   LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))</pre>
   ### Log-Posterior
   LP <- LL + alpha.prior + beta.prior + lambda.prior + phi.prior +
      sigma.prior
   Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
      yhat=rnorm(length(mu), mu, sigma), parm=parm)
   return(Modelout)
```

25.4. Initial Values

Initial. Values $\leftarrow c(rep(0,2), 0.5, 0, log(1))$

26. Exponential Smoothing

26.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu_t = \alpha \mathbf{y}_{t-1} + (1 - \alpha)\mu_{t-1}, \quad t = 2, \dots, T$$

$$\alpha \sim \mathcal{U}(0, 1)$$

$$\sigma \sim \mathcal{HC}$$

26.2. Data

```
T <- 10
y <- rep(0,T)
y[1] <- 0
for (t in 2:T) {y[t] <- y[t-1] + rnorm(1,0,0.1)}
mon.names <- c("LP", "sigma")
parm.names <- c("alpha","log.sigma")
PGF <- function(Data) return(c(runif(1), log(rhalfcauchy(1,25))))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names, y=y)</pre>
```

26.3. Model

```
Model <- function(parm, Data)
    {
     ### Parameters
     parm[1] <- alpha <- interval(parm[1], 0, 1)
     sigma <- exp(parm[2])
     ### Log(Prior Densities)
     alpha.prior <- dunif(alpha, 0, 1, log=TRUE)
     sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)
     ### Log-Likelihood
     mu <- y
     mu[-1] <- alpha*Data$y[-1]
     mu[-1] <- mu[-1] + (1 - alpha) * mu[-Data$T]
     LL <- sum(dnorm(Data$y[-1], mu[-Data$T], sigma, log=TRUE))</pre>
```

26.4. Initial Values

Initial. Values \leftarrow c(0.5, log(1))

27. Factor Analysis, Approximate Dynamic

The Approximate Dynamic Factor Analysis (ADFA) model has many names, including the approximate factor model and approximate dynamic factor model. An ADFA is a Dynamic Factor Analysis (DFA) in which the factor scores of the dynamic factors are approximated with principal components. This is a combination of principal components and common factor analysis, in which the factor loadings of common factors are estimated from the data and factor scores are estimated from principal components. This is a two-stage model: principal components are estimated in the first stage and a decision is made regarding how many principal components to retain, and ADFA is estimated in the second stage. For more information on DFA, see section 29.

27.1. Form

$$\mathbf{Y}_{t,j} \sim \mathcal{N}(\mu_{t,j}, \sigma_j^2), \quad t = 2, \dots, T, \quad j = 1, \dots, J$$

$$\mu_{t,j} = \mathbf{F}_{t-1,j} \Lambda$$

$$\Lambda_{p,j} \sim \mathcal{N}(0, 1000), \quad p = 1, \dots, P, \quad j = 1, \dots, J$$

$$\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, J$$

```
T <- 10 #Number of time-periods
J <- 20 #Number of variables
P <- 5 #Number of approximate dynamic factors
Lambda <- matrix(runif(J*P,-1,1), P, J)
Sigma <- matrix(runif(P*P), P, P); diag(Sigma) <- runif(P)*5
Sigma <- as.symmetric.matrix(Sigma); Sigma <- as.positive.definite(Sigma)
F <- rmvn(T, rep(0,P), Sigma)
Y <- tcrossprod(F, t(Lambda))
PCA <- prcomp(Y, scale=TRUE)
F <- PCA$x[,1:P]
mon.names <- c("LP", paste("ynew[", 1:J, "]", sep=""))</pre>
```

27.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    Lambda <- matrix(parm[1:(Data$P*Data$J)], Data$P, Data$J)</pre>
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
     ### Log(Prior Densities)
    Lambda.prior <- sum(dnormv(Lambda, 0, 1000, log=TRUE))</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- rbind(rep(0, Data$J), tcrossprod(F[-Data$T,], t(Lambda)))</pre>
    Sigma <- matrix(sigma, Data$T, Data$J, byrow=TRUE)</pre>
    ynew <- rnorm(Data$J, tcrossprod(F[Data$T,], t(Lambda)), sigma)</pre>
    LL <- sum(dnorm(Data$Y, mu, Sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + Lambda.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,ynew),</pre>
         yhat=rnorm(prod(dim(mu)), mu, Sigma), parm=parm)
    return(Modelout)
    }
```

27.4. Initial Values

Initial.Values <- c(rep(0,P*J), rep(1,J))</pre>

28. Factor Analysis, Confirmatory

Factor scores are in matrix \mathbf{F} , factor loadings for each variable are in vector λ , and \mathbf{f} is a vector that indicates which variable loads on which factor.

$$\mathbf{Y}_{i,m} \sim \mathcal{N}(\mu_{i,m}, \sigma_m^2), \quad i = 1, \dots, N, \quad m = 1, \dots, M$$
$$\mu = \alpha^T + \mathbf{F}_{1:N,\mathbf{f}}\lambda^T$$
$$\mathbf{F}_{i,1:P} \sim \mathcal{N}_P(\gamma, \Omega^{-1}), \quad i = 1, \dots, N$$
$$\alpha_m \sim \mathcal{N}(0, 1000), \quad m = 1, \dots, M$$

```
\lambda_m \sim \mathcal{N}(0, 1000), \quad m = 1, \dots, M
\sigma_m \sim \mathcal{HC}(25), \quad m = 1, \dots, M
\Omega \sim \mathcal{W}_N(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_P
```

```
data(swiss)
Y <- cbind(swiss$Agriculture, swiss$Examination, swiss$Education,
    swiss$Catholic, swiss$Infant.Mortality)
M <- ncol(Y) #Number of variables
N <- nrow(Y) #Number of records
P <- 3 #Number of factors
f \leftarrow c(1,3,2,2,1) #Indicator f for the factor for each variable m
gamma \leftarrow rep(0,P)
S <- diag(P)
mon.names <- "LP"
parm.names <- as.parm.names(list(F=matrix(0,N,P), lambda=rep(0,M),</pre>
    U=diag(P), alpha=rep(0,M), log.sigma=rep(0,M)),
    uppertri=c(0,0,1,0,0))
PGF <- function(Data) return(c(rmvnpc(Data$N, Data$gamma,
    rwishartc(Data$N,Data$S)), rnormv(Data$M,0,1000),
    upper.triangle(rwishartc(Data$N,Data$S), diag=TRUE),
    rnormv(Data$M,0,1000), log(rhalfcauchy(Data$M,25))))
MyData <- list(M=M, N=N, P=P, PGF=PGF, S=S, Y=Y, f=f, gamma=gamma,
    mon.names=mon.names, parm.names=parm.names)
28.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    alpha <- parm[grep("alpha", Data$parm.names)]</pre>
    lambda <- parm[grep("lambda", Data$parm.names)]</pre>
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    F <- matrix(parm[grep("F", Data$parm.names)], Data$N, Data$P)
    U <- as.parm.matrix(U, Data$P, parm, Data, chol=TRUE)</pre>
    diag(U) <- exp(diag(U))</pre>
    ### Log(Prior Densities)
    alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
    lambda.prior <- sum(dnormv(lambda, 0, 1000, log=TRUE))</pre>
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    U.prior <- dwishartc(U, Data$N, Data$S, log=TRUE)</pre>
    F.prior <- sum(dmvnpc(F, Data$gamma, U, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- matrix(alpha, Data$N, Data$M, byrow=TRUE) + F[,Data$f] *</pre>
         matrix(lambda, Data$N, Data$M, byrow=TRUE)
```

28.4. Initial Values

```
Initial.Values <- c(rep(0,N*P), rep(0,M), upper.triangle(S, diag=TRUE),
    rep(0,M), rep(0,M))</pre>
```

29. Factor Analysis, Dynamic

The factor scores in **F** are dynamic with respect to time, and are estimated as in a state space model (SSM) or dynamic linear model (DLM) with constant variance in the state vector. For more information on SSMs, see section 77. For more information on exploratory factor analysis, see section 30.

29.1. Form

$$\mathbf{Y}_{t,j} \sim \mathcal{N}(\mu_{t,j}, \sigma_j^2), \quad t = 2, \dots, T, \quad j = 1, \dots, J$$

$$\mu_{2:T,} = \mathbf{F}_{1:(T-1),\Lambda}$$

$$\mathbf{F}_{1,1:P} \sim \mathcal{N}_P(0, \Omega^{-1})$$

$$\mathbf{F}_{t,1:P} \sim \mathcal{N}_P(\mathbf{F}_{t-1,1:P}, \Omega^{-1}), \quad t = 2, \dots, T$$

$$\Lambda_{p,j} \sim \mathcal{N}(0, 1000), \quad p = 1, \dots, P, \quad j = 1, \dots, J$$

$$\Omega \sim \mathcal{W}_N(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_P$$

$$\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, J$$

```
T <- 10 #Number of time-periods
J <- 20 #Number of time-series
P <- 3 #Number of dynamic factors
Lambda <- matrix(runif(J*P,-1,1), P, J)
Sigma <- matrix(runif(P*P), P, P); diag(Sigma) <- runif(P)*5
Sigma <- as.symmetric.matrix(Sigma); Sigma <- as.positive.definite(Sigma)
F <- rmvn(T, rep(0,P), Sigma)</pre>
```

29.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    F <- matrix(parm[1:(Data$T*Data$P)], Data$T, Data$P)</pre>
    U <- as.parm.matrix(U, Data$P, parm, Data, chol=TRUE)</pre>
     diag(U) <- exp(diag(U)</pre>
    Lambda <- matrix(parm[grep("Lambda", Data$parm.names)], Data$P, Data$J)
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    F.prior <- sum(dmvnpc(F, rbind(rep(0, Data$P), F[-Data$T,]), U,
         log=TRUE))
    U.prior <- dwishartc(U, Data$P+1, Data$S, log=TRUE)</pre>
    Lambda.prior <- sum(dnormv(Lambda, 0, 1000, log=TRUE))</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- rbind(rep(0, Data$J), tcrossprod(F[-Data$T,], t(Lambda)))</pre>
    Sigma <- matrix(sigma, Data$T, Data$J, byrow=TRUE)</pre>
    ynew <- rnorm(Data$J, tcrossprod(F[Data$T,], t(Lambda)), sigma)</pre>
    LL <- sum(dnorm(Data$Y, mu, Sigma, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + F.prior + U.prior + Lambda.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,ynew),</pre>
         yhat=rnorm(prod(dim(mu)), mu, Sigma), parm=parm)
    return(Modelout)
    }
```

29.4. Initial Values

Initial.Values <- c(rep(0,T*P), S[upper.tri(S, diag=TRUE)], rep(0,P*J),
 rep(0,J))</pre>

30. Factor Analysis, Exploratory

Factor scores are in matrix \mathbf{F} and factor loadings are in matrix Λ .

30.1. Form

$$\mathbf{Y}_{i,m} \sim \mathcal{N}(\mu_{i,m}, \sigma_m^2), \quad i = 1, \dots, N, \quad m = 1, \dots, M$$

$$\mu = \alpha^T + \mathbf{F}\Lambda$$

$$\mathbf{F}_{i,1:P} \sim \mathcal{N}_P(\gamma, \Omega^{-1}), \quad i = 1, \dots, N$$

$$\alpha_m \sim \mathcal{N}(0, 1000), \quad m = 1, \dots, M$$

$$\gamma_p = 0, \quad p = 1, \dots, P$$

$$\Lambda_{p,m} \sim \mathcal{N}(0, 1000), \quad p = 1, \dots, P, \quad m = 1, \dots, M$$

$$\Omega \sim \mathcal{W}_N(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_P$$

$$\sigma_m \sim \mathcal{HC}(25), \quad m = 1, \dots, M$$

```
M <- 10 #Number of variables
N <- 20 #Number of records
P <- 3 #Number of factors
alpha <- runif(M)*10</pre>
Lambda <- matrix(runif(M*P,-1,1), P, M)</pre>
U <- diag(P) U[upper.tri(U, diag=TRUE)] <- runif(length(upper.triangle(U, diag=TRUE)))</pre>
F \leftarrow rmvnc(N, rep(0,P), U)
Y <- matrix(alpha, N, M, byrow=TRUE) + tcrossprod(F, t(Lambda))
gamma \leftarrow rep(0,P)
S \leftarrow diag(P)
mon.names <- "LP"
parm.names <- as.parm.names(list(F=matrix(0,N,P), Lambda=matrix(0,P,M),</pre>
    U=diag(P), alpha=rep(0,M), log.sigma=rep(0,M)),
    uppertri=c(0,0,1,0,0))
PGF <- function(Data) return(c(rmvnpc(Data$N, Data$gamma,
    rwishartc(Data$N, Data$S)), rnormv(Data$P*Data$M,0,1000),
    upper.triangle(rwishartc(Data$N, Data$S), diag=TRUE),
    rnormv(Data$M,0,1000), log(rhalfcauchy(Data$M,25))))
MyData <- list(M=M, N=N, P=P, PGF=PGF, S=S, Y=Y, gamma=gamma,
    mon.names=mon.names, parm.names=parm.names)
```

30.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    F <- matrix(parm[grep("F", Data$parm.names)], Data$N, Data$P)
    Lambda <- matrix(parm[grep("Lambda", Data$parm.names)],</pre>
         Data$P, Data$M)
    U <- as.parm.matrix(U, Data$P, parm, Data, chol=TRUE)</pre>
    diag(U) <- exp(diag(U))</pre>
    alpha <- parm[grep("alpha", Data$parm.names)]</pre>
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    F.prior <- sum(dmvnpc(F, Data$gamma, U, log=TRUE))</pre>
     Lambda.prior <- sum(dnormv(Lambda, 0, 1000, log=TRUE))</pre>
    U.prior <- dwishartc(U, Data$N, Data$S, log=TRUE)</pre>
     alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
     ### Log-Likelihood
    mu <- matrix(alpha, Data$N, Data$M, byrow=TRUE) +</pre>
          tcrossprod(F, t(Lambda))
    Sigma <- matrix(sigma, Data$N, Data$M, byrow=TRUE)</pre>
    LL <- sum(dnorm(Data$Y, mu, Sigma, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + F.prior + Lambda.prior + U.prior + alpha.prior +
                                                                                sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
    yhat=rnorm(prod(dim(mu)), mu, Sigma), parm=parm)
    return(Modelout)
    }
```

30.4. Initial Values

```
Initial.Values <- c(rep(0,N*P), rep(0,P*M), upper.triangle(S, diag=TRUE),
    rep(0,M), rep(0,M))</pre>
```

31. Factor Regression

This example of factor regression is constrained to the case where the number of factors is equal to the number of independent variables (IVs) less the intercept. The purpose of this form of factor regression is to orthogonalize the IVs with respect to \mathbf{y} , rather than variable reduction. This method is the combination of confirmatory factor analysis in section 28 and linear regression in section 40.

31.1. Form

$$\mathbf{y} \sim \mathcal{N}(\nu, \sigma_{J+1}^2)$$

$$\nu = \mu \beta$$

$$\mu_{i,1} = 1$$

$$\mu_{i,j+1} = \mu_{i,j}, \quad j = 1, \dots, J$$

$$\mathbf{X}_{i,j} \sim \mathcal{N}(\mu_{i,j}, \sigma_j^2), \quad i = 1, \dots, N, \quad j = 2, \dots, J$$

$$\mu_{i,j} = \alpha_j + \lambda_j \mathbf{F}_{i,j}, \quad i = 1, \dots, N, \quad j = 2, \dots, J$$

$$\mathbf{F}_{i,1:J} \sim \mathcal{N}_{J-1}(0, \Omega^{-1}), \quad i = 1, \dots, N$$

$$\alpha_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\lambda_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1)$$

$$\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, (J+1)$$

$$\Omega \sim \mathcal{W}_N(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_J$$

31.2. Data

```
data(demonsnacks)
N <- nrow(demonsnacks)</pre>
J <- 3
y <- log(demonsnacks$Calories)</pre>
X <- as.matrix(demonsnacks[,c(7,8,10)])</pre>
for (j in 1:J) {X[,j] <- CenterScale(X[,j])}</pre>
S \leftarrow diag(J)
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=rep(0,J), beta=rep(0,J+1),</pre>
    lambda=rep(0,J), log.sigma=rep(0,J+1), F=matrix(0,N,J),
    Omega=diag(J)), uppertri=c(0,0,0,0,0,1))
PGF <- function(Data) return(c(rnormv(Data$J,0,1000),
    rnormv(Data$J+1,0,1000), rnormv(Data$J,0,1000),
    log(rhalfcauchy(Data$J+1,25)), rmvnp(Data$N, rep(0,Data$J), Data$S)))
MyData <- list(J=J, N=N, PGF=PGF, S=S, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
```

31.3. Model

```
Model <- function(parm, Data)
    {
     ### Parameters
     alpha <- parm[grep("alpha", Data$parm.names)]</pre>
```

```
beta <- parm[grep("beta", Data$parm.names)]</pre>
lambda <- parm[grep("lambda", Data$parm.names)]</pre>
sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
F <- matrix(parm[grep("F", Data$parm.names)], Data$N, Data$J)</pre>
Omega <- as.parm.matrix(Omega, Data$J, parm, Data)</pre>
parm[grep("Omega", Data$parm.names)] <- upper.triangle(Omega,</pre>
    diag=TRUE)
### Log(Prior Densities)
alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
lambda.prior <- sum(dnormv(lambda, 0, 1000, log=TRUE))</pre>
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
Omega.prior <- dwishart(Omega, Data$N, Data$S, log=TRUE)</pre>
F.prior <- sum(dmvnp(F, rep(0,Data$J), Omega, log=TRUE))
### Log-Likelihood
mu <- matrix(alpha, Data$N, Data$J, byrow=TRUE) + F *</pre>
    matrix(lambda, Data$N, Data$J, byrow=TRUE)
nu <- tcrossprod(beta, cbind(rep(1,Data$N),mu))</pre>
LL <- sum(dnorm(Data$X, mu, matrix(sigma[1:Data$J], Data$N, Data$J,
    byrow=TRUE)), dnorm(Data$y, nu, sigma[Data$J+1], log=TRUE))
### Log-Posterior
LP <- LL + alpha.prior + beta.prior + lambda.prior + sigma.prior +
    F.prior + Omega.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
    yhat=rnorm(Data$N, nu, sigma[Data$J+1]), parm=parm)
return(Modelout)
}
```

31.4. Initial Values

```
Initial.Values <- c(rep(0,J), rep(0,J+1), rep(0,J), rep(0,J+1), rep(0,N*J), upper.triangle(S, diag=TRUE))
```

32. Gamma Regression

$$\mathbf{y} \sim \mathcal{G}(\lambda \tau, \tau)$$

$$\lambda = \exp(\mathbf{X}\beta)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\tau \sim \mathcal{HC}(25)$$

32.2. Data

32.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    tau <- exp(parm[grep("log.tau", Data$parm.names)])</pre>
    sigma2 <- 1/tau
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    tau.prior <- dhalfcauchy(tau, 25, log=TRUE)</pre>
    ### Log-Likelihood
    lambda <- exp(tcrossprod(Data$X, t(beta)))</pre>
    LL <- sum(dgamma(Data$y, tau*lambda, tau, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + beta.prior + tau.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma2),</pre>
          yhat=rgamma(nrow(lambda), tau*lambda, tau), parm=parm)
    return(Modelout)
    }
```

32.4. Initial Values

Initial.Values <- c(rep(0,J), 1)</pre>

33. GARCH(1,1)

$$\mathbf{y}_t \sim \mathcal{N}(\mu_t, \sigma_t^2), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} \sim \mathcal{N}(\mu_{T+1}, \sigma_{new}^2)$$

$$\mu_t = \alpha + \phi \mathbf{y}_{t-1}, \quad t = 1, \dots, (T+1)$$

$$\epsilon_t = \mathbf{y}_t - \mu_t$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\sigma_{new}^2 = \theta_1 + \theta_2 \epsilon_T^2 + \theta_3 \sigma_T^2$$

$$\sigma_t^2 = \theta_1 + \theta_2 \epsilon_{t-1}^2 + \theta_3 \sigma_{t-1}^2$$

$$\theta_k = \frac{1}{1 + \exp(-\theta_k)}, \quad k = 1, \dots, 3$$

$$\theta_k \sim \mathcal{N}(0, 1000) \in [-10, 10], \quad k = 1, \dots, 3$$

33.2. Data

```
y \leftarrow c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91,
    2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28,
    1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03,
    -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02,
    0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36,
    1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70,
    0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83,
    0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24,
    0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40,
    -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
    0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17
T <- length(y)
```

mon.names <- c("LP", "ynew", "sigma2.new")</pre>

```
parm.names <- c("alpha", "phi", "logit.theta[1]", "logit.theta[2]",</pre>
     "logit.theta[3]")
PGF <- function(Data) return(c(rnormv(2,0,1000),
    log(rhalfcauchy(1,25)), runif(2)))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,
    y=y)
33.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    alpha <- parm[1]; phi <- parm[2]</pre>
    theta <- invlogit(interval(parm[grep("logit.theta",
         Data$parm.names)], -10, 10))
    parm[grep("logit.theta", Data$parm.names)] <- logit(theta)</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
    theta.prior <- sum(dnormv(theta, 0, 1000, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- c(alpha, alpha + phi*Data$y[-Data$T])</pre>
    epsilon <- Data$y - mu
     sigma2 <- c(theta[1], theta[1] + theta[2]*epsilon[-Data$T]^2)</pre>
     sigma2[-1] \leftarrow sigma2[-1] + theta[3]*sigma2[-Data$T]
    sigma2.new <- theta[1] + theta[2]*epsilon[Data$T]^2 +</pre>
         theta[3]*sigma2[Data$T]
    ynew <- rnormv(1, alpha + phi*Data$y[Data$T], sigma2.new)</pre>
    LL <- sum(dnormv(Data$y, mu, sigma2, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + phi.prior + theta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, ynew, sigma2.new),</pre>
         yhat=rnormv(length(mu), mu, sigma2), parm=parm)
    return(Modelout)
    }
33.4. Initial Values
```

34. GARCH-M(1,1)

Initial. Values $\leftarrow c(rep(0,2), rep(0.4,3))$

34.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{t}^{2}), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} \sim \mathcal{N}(\mu_{T+1}, \sigma_{new}^{2})$$

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1} + \delta \sigma_{t-1}^{2}, \quad t = 1, \dots, (T+1)$$

$$\epsilon_{t} = \mathbf{y}_{t} - \mu_{t}$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\phi \sim \mathcal{N}(0, 1000)$$

$$\sigma_{new}^{2} = \omega + \theta_{1} \epsilon_{T}^{2} + \theta_{2} \sigma_{T}^{2}$$

$$\sigma_{t}^{2} = \omega + \theta_{1} \epsilon_{t-1}^{2} + \theta_{2} \sigma_{t-1}^{2}$$

$$\omega \sim \mathcal{HC}(25)$$

$$\theta_{k} \sim \mathcal{U}(0, 1), \quad k = 1, \dots, 2$$

```
y \leftarrow c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91,
    2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28,
    1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03,
    -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02,
    0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36,
    1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70,
    0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83,
    0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24,
    0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40,
    -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
    0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
```

```
0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17
T <- length(y)
mon.names <- c("LP", "ynew", "sigma2.new")</pre>
parm.names <- c("alpha", "phi", "delta", "log.omega", "theta[1]", "theta[2]")</pre>
PGF <- function(Data) return(c(rnormv(3,0,1000),
    log(rhalfcauchy(1,25)), runif(2)))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,</pre>
    y=y)
34.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
     alpha <- parm[1]; phi <- parm[2]; delta <- parm[3]</pre>
     omega <- exp(parm[4])</pre>
    parm[5:6] <- theta <- interval(parm[5:6], 1e-10, 1-1e-5)</pre>
    ### Log(Prior Densities)
    alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dnormv(phi, 0, 1000, log=TRUE)</pre>
     delta.prior <- dnormv(delta, 0, 1000, log=TRUE)</pre>
    omega.prior <- dhalfcauchy(omega, 25, log=TRUE)</pre>
    theta.prior <- sum(dunif(theta, 0, 1, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- c(alpha, alpha + phi*Data$y[-Data$T])</pre>
    epsilon <- Data$y - mu
     sigma2 <- c(omega, omega + theta[1]*epsilon[-Data$T]^2)</pre>
     sigma2[-1] \leftarrow sigma2[-1] + theta[2]*sigma2[-Data$T]
    sigma2.new <- omega + theta[1]*epsilon[Data$T]^2 +</pre>
         theta[2]*sigma2[Data$T]
    mu <- mu + delta*sigma2
    ynew <- rnormv(1, alpha + phi*Data$y[Data$T] + delta*sigma2[Data$T],</pre>
         sigma2.new)
    LL <- sum(dnormv(Data$y, mu, sigma2, log=TRUE))
    ### Log-Posterior
    LP <- LL + alpha.prior + phi.prior + delta.prior + omega.prior +
         theta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, ynew, sigma2.new),</pre>
         yhat=rnormv(length(mu), mu, sigma2), parm=parm)
    return(Modelout)
    }
```

34.4. Initial Values

```
Initial. Values <- c(rep(0,3), rep(0.3,3))
```

35. Geographically Weighted Regression

35.1. Form

$$\mathbf{y}_{i,k} \sim \mathcal{N}(\mu_{i,k}, \tau_{i,k}^{-1}), \quad i = 1, \dots, N, \quad k = 1, \dots, N$$

$$\mu_{i,1:N} = \mathbf{X}\beta_{i,1:J}$$

$$\tau = \frac{1}{\sigma^2} \mathbf{w} \nu$$

$$\mathbf{w} = \frac{\exp(-0.5\mathbf{Z}^2)}{\mathbf{h}}$$

$$\alpha \sim \mathcal{U}(1.5, 100)$$

$$\beta_{i,j} \sim \mathcal{N}(0, 1000), \quad i = 1, \dots, N, \quad j = 1, \dots, J$$

$$\mathbf{h} \sim \mathcal{N}(0.1, 1000) \in [0.1, \infty]$$

$$\nu_{i,k} \sim \mathcal{G}(\alpha, 2), \quad i = 1, \dots, N, \quad k = 1, \dots, N$$

$$\sigma_i \sim \mathcal{HC}(25), \quad i = 1, \dots, N$$

```
crime <- c(18.802, 32.388, 38.426, 0.178, 15.726, 30.627, 50.732,
    26.067, 48.585, 34.001, 36.869, 20.049, 19.146, 18.905, 27.823,
    16.241, 0.224, 30.516, 33.705, 40.970, 52.794, 41.968, 39.175,
    53.711, 25.962, 22.541, 26.645, 29.028, 36.664, 42.445, 56.920,
    61.299, 60.750, 68.892, 38.298, 54.839, 56.706, 62.275, 46.716,
    57.066, 54.522, 43.962, 40.074, 23.974, 17.677, 14.306, 19.101,
    16.531, 16.492)
income <- c(21.232, 4.477, 11.337, 8.438, 19.531, 15.956, 11.252,
    16.029, 9.873, 13.598, 9.798, 21.155, 18.942, 22.207, 18.950,
    29.833, 31.070, 17.586, 11.709, 8.085, 10.822, 9.918, 12.814,
    11.107, 16.961, 18.796, 11.813, 14.135, 13.380, 17.017, 7.856,
    8.461, 8.681, 13.906, 14.236, 7.625, 10.048, 7.467, 9.549,
    9.963, 11.618, 13.185, 10.655, 14.948, 16.940, 18.739, 18.477,
    18.324, 25.873)
housing <- c(44.567, 33.200, 37.125, 75.000, 80.467, 26.350, 23.225,
    28.750, 18.000, 96.400, 41.750, 47.733, 40.300, 42.100, 42.500,
    61.950, 81.267, 52.600, 30.450, 20.300, 34.100, 23.600, 27.000,
    22.700, 33.500, 35.800, 26.800, 27.733, 25.700, 43.300, 22.850,
    17.900, 32.500, 22.500, 53.200, 18.800, 19.900, 19.700, 41.700,
    42.900, 30.600, 60.000, 19.975, 28.450, 31.800, 36.300, 39.600,
    76.100, 44.333)
easting <- c(35.62, 36.50, 36.71, 33.36, 38.80, 39.82, 40.01, 43.75,
    39.61, 47.61, 48.58, 49.61, 50.11, 51.24, 50.89, 48.44, 46.73,
    43.44, 43.37, 41.13, 43.95, 44.10, 43.70, 41.04, 43.23, 42.67,
```

```
41.21, 39.32, 41.09, 38.3, 41.31, 39.36, 39.72, 38.29, 36.60,
    37.60, 37.13, 37.85, 35.95, 35.72, 35.76, 36.15, 34.08, 30.32,
    27.94, 27.27, 24.25, 25.47, 29.02)
northing <- c(42.38, 40.52, 38.71, 38.41, 44.07, 41.18, 38.00, 39.28,
    34.91, 36.42, 34.46, 32.65, 29.91, 27.80, 25.24, 27.93, 31.91,
    35.92, 33.46, 33.14, 31.61, 30.40, 29.18, 28.78, 27.31, 24.96,
    25.90, 25.85, 27.49, 28.82, 30.90, 32.88, 30.64, 30.35, 32.09,
    34.08, 36.12, 36.30, 36.40, 35.60, 34.66, 33.92, 30.42, 28.26,
    29.85, 28.21, 26.69, 25.71, 26.58)
N <- length(crime)
J <- 3 #Number of predictors, including the intercept
X <- matrix(c(rep(1,N), income, housing),N,J)</pre>
D <- as.matrix(dist(cbind(northing,easting), diag=TRUE, upper=TRUE))</pre>
Z <- D / sd(as.vector(D))</pre>
y \leftarrow matrix(0,N,N); for (i in 1:N) {for (k in 1:N) {y[i,k] <- crime[k]}}
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=0, beta=matrix(0,N,J), log.h=0,
    log.nu=matrix(0,N,N), log.sigma=rep(0,N)))
PGF <- function(Data) return(c(runif(1,1.5,100),
    rnormv(Data$N*Data$J,0,1000),
    log(rtrunc(1, "normv", a=0.1, b=Inf, mean=0.1, var=1000)),
    log(rgamma(Data$N*Data$N,runif(1,1.5,100),2)),
    log(rhalfcauchy(Data$N,25))))
MyData <- list(J=J, N=N, PGF=PGF, X=X, Z=Z, latitude=northing,
    longitude=easting, mon.names=mon.names, parm.names=parm.names, y=y)
35.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    parm[1] <- alpha <- interval(parm[1], 1.5, 100)</pre>
    beta <- matrix(parm[grep("beta", Data$parm.names)], Data$N, Data$J)</pre>
    h \leftarrow \exp(parm[2+(N*J)]) + 0.1
    nu <- exp(matrix(parm[grep("log.nu", Data$parm.names)],</pre>
         Data$N, Data$N))
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    alpha.prior <- dunif(alpha, 1.5, 100, log=TRUE)</pre>
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    h.prior <- dtrunc(h, "normv", a=0.1, b=Inf, mean=0.1, var=1000,
         log=TRUE)
    nu.prior <- sum(dgamma(nu, alpha, 2, log=TRUE))</pre>
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    w \leftarrow \exp(-0.5 * Data$Z^2) / h
```

35.4. Initial Values

36. Inverse Gaussian Regression

36.1. Form

$$\mathbf{y} \sim \mathcal{N}^{-1}(\mu, \lambda)$$

$$\mu = \exp(\mathbf{X}\beta) + C$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\lambda \sim \mathcal{HC}(25)$$

where C is a small constant, such as 1.0E-10.

parm.names=parm.names, y=y)

36.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:Data$J]
    lambda <- exp(parm[Data$J+1])</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    lambda.prior <- dhalfcauchy(lambda, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- exp(tcrossprod(Data$X, t(beta))) + 1.0E-10</pre>
    LL <- sum(dinvgaussian(Data$y, mu, lambda, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + beta.prior + lambda.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,lambda),</pre>
         yhat=rinvgaussian(length(mu), mu, lambda), parm=parm)
    return(Modelout)
    }
```

36.4. Initial Values

Initial.Values <- c(rep(0,J), log(1))</pre>

37. Kriging

This is an example of universal kriging of \mathbf{y} given \mathbf{X} , regression effects β , and spatial effects ζ . Euclidean distance between spatial coordinates (longitude and latitude) is used for each of $i=1,\ldots,N$ records of \mathbf{y} . An additional record is created from the same data-generating process to compare the accuracy of interpolation. For the spatial component, ϕ is the rate of spatial decay and κ is the scale. κ is often difficult to identify, so it is set to 1 (Gaussian), but may be allowed to vary up to 2 (Exponential). In practice, ϕ is also often difficult to identify. While Σ is spatial covariance, spatial correlation is $\rho = \exp(-\phi \mathbf{D})$. To extend this to a large data set, consider the predictive process kriging example in section 38.

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma_1^2)$$

$$\mu = \mathbf{X}\beta + \zeta$$

$$\mathbf{y}^{new} = \mathbf{X}\beta + \sum_{i=1}^{N} \left(\frac{\rho_i}{\sum \rho} \zeta_i\right)$$

```
\rho = \exp(-\phi \mathbf{D}^{new})^{\kappa}
\zeta \sim \mathcal{N}_N(\zeta_{\mu}, \Sigma)
\Sigma = \sigma_2^2 \exp(-\phi \mathbf{D})^{\kappa}
\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, 2
\sigma_j \sim \mathcal{HC}(25) \in [0.1, 10], \quad j = 1, \dots, 2
\phi \sim \mathcal{U}(1, 5)
\zeta_{\mu} = 0
\kappa = 1
```

```
N <- 20
longitude <- runif(N+1,0,100)</pre>
latitude <- runif(N+1,0,100)</pre>
D <- as.matrix(dist(cbind(longitude,latitude), diag=TRUE, upper=TRUE))
Sigma < -10000 * exp(-1.5 * D)
zeta <- as.vector(apply(rmvn(1000, rep(0,N+1), Sigma), 2, mean))</pre>
beta <- c(50,2)
X \leftarrow matrix(runif((N+1)*2,-2,2),(N+1),2); X[,1] \leftarrow 1
mu <- as.vector(tcrossprod(X, t(beta)))</pre>
y <- mu + zeta
longitude.new <- longitude[N+1]; latitude.new <- latitude[N+1]</pre>
Xnew \leftarrow X[N+1,]; ynew \leftarrow y[N+1]
longitude <- longitude[1:N]; latitude <- latitude[1:N]</pre>
X \leftarrow X[1:N,]; y \leftarrow y[1:N]
D <- as.matrix(dist(cbind(longitude,latitude), diag=TRUE, upper=TRUE))</pre>
D.new <- sqrt((longitude - longitude.new)^2 + (latitude - latitude.new)^2)</pre>
mon.names <- c("LP", "ynew")</pre>
parm.names <- as.parm.names(list(zeta=rep(0,N), beta=rep(0,2),
     sigma=rep(0,2), phi=0))
PGF <- function(Data) return(c(rmvn(1, rep(0, Data$N),
    rhalfcauchy(1,25)^2 *exp(-runif(1,1,5)*Data$D)),
    rnormv(2,0,1000), log(rhalfcauchy(2,25)), runif(1,1,5)))
MyData <- list(D=D, D.new=D.new, latitude=latitude, longitude=longitude,
    N=N, PGF=PGF, X=X, Xnew=Xnew, mon.names=mon.names,
    parm.names=parm.names, y=y)
37.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
```

```
beta <- parm[grep("beta", Data$parm.names)]</pre>
zeta <- parm[grep("zeta", Data$parm.names)]</pre>
kappa <- 1
sigma <- interval(parm[grep("sigma", Data$parm.names)], 0.1, 10)</pre>
phi <- interval(parm[grep("phi", Data$parm.names)], 1, 5)</pre>
parm[grep("phi", Data$parm.names)] <- phi</pre>
Sigma <- sigma[2] * exp(-phi * Data$D)^kappa
### Log(Prior Densities)
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
zeta.prior <- dmvn(zeta, rep(0, Data$N), Sigma, log=TRUE)
sigma.prior <- sum(dhalfcauchy(sigma - 1, 25, log=TRUE))</pre>
phi.prior <- dunif(phi, 1, 5, log=TRUE)</pre>
### Interpolation
rho <- exp(-phi * Data$D.new)^kappa</pre>
ynew <- rnorm(1, sum(beta * Data$Xnew) + sum(rho / sum(rho) * zeta),</pre>
    sigma)
### Log-Likelihood
mu <- tcrossprod(Data$X, t(beta)) + zeta</pre>
LL <- sum(dnorm(Data$y, mu, sigma[1], log=TRUE))
### Log-Posterior
LP <- LL + beta.prior + zeta.prior + sigma.prior + phi.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,ynew),</pre>
    yhat=rnorm(length(mu), mu, sigma[1]), parm=parm)
return(Modelout)
}
```

37.4. Initial Values

Initial. Values \leftarrow c(rep(0,N), rep(0,2), rep(1,2), 1)

38. Kriging, Predictive Process

The first K of N records in \mathbf{y} are used as knots for the parent process, and the predictive process involves records $(K+1), \ldots, N$. For more information on kriging, see section 37.

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma_1^2)$$

$$\mu_{1:K} = \mathbf{X}_{1:K,1:J}\beta + \zeta$$

$$\mu_{(K+1):N} = \mathbf{X}_{(K+1):N,1:J}\beta + \sum_{p=1}^{N-K} \frac{\lambda_{p,1:K}}{\sum_{q=1}^{N-K} \lambda_{q,1:K}} \zeta^T$$

$$\lambda = \exp(-\phi \mathbf{D}_P)^{\kappa}$$

$$\mathbf{y}^{new} = \mathbf{X}\boldsymbol{\beta} + \sum_{k=1}^{K} \left(\frac{\rho_k}{\sum \rho} \zeta_k\right)$$

$$\rho = \exp(-\phi \mathbf{D}^{new})^{\kappa}$$

$$\zeta \sim \mathcal{N}_K(0, \Sigma)$$

$$\Sigma = \sigma_2^2 \exp(-\phi \mathbf{D})^{\kappa}$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, 2$$

$$\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, 2$$

$$\phi \sim \mathcal{N}(0, 1000) \in [0, \infty]$$

$$\kappa = 1$$

```
N <- 100
K <- 30 #Number of knots
longitude <- runif(N+1,0,100)</pre>
latitude <- runif(N+1,0,100)</pre>
D <- as.matrix(dist(cbind(longitude,latitude), diag=TRUE, upper=TRUE))</pre>
Sigma < -10000 * exp(-1.5 * D)
zeta <- as.vector(apply(rmvn(1000, rep(0,N+1), Sigma), 2, mean))</pre>
beta <- c(50,2)
X \leftarrow matrix(runif((N+1)*2,-2,2),(N+1),2); X[,1] \leftarrow 1
mu <- as.vector(tcrossprod(X, t(beta)))</pre>
y <- mu + zeta
longitude.new <- longitude[N+1]; latitude.new <- latitude[N+1]</pre>
Xnew <- X[N+1,]; ynew <- y[N+1]</pre>
longitude <- longitude[1:N]; latitude <- latitude[1:N]</pre>
X \leftarrow X[1:N,]; y \leftarrow y[1:N]
D <- as.matrix(dist(cbind(longitude[1:K],latitude[1:K]), diag=TRUE,
    upper=TRUE))
D.P <- matrix(0, N-K, K)</pre>
for (i in (K+1):N) {
    D.P[K+1-i,] <- sqrt((longitude[1:K] - longitude[i])^2 +</pre>
          (latitude[1:K] - latitude[i])^2)}
D.new <- sqrt((longitude[1:K] - longitude.new)^2 +</pre>
     (latitude[1:K] - latitude.new)^2)
mon.names <- c("LP", "sigma[1]", "sigma[2]", "ynew")</pre>
parm.names <- as.parm.names(list(zeta=rep(0,K), beta=rep(0,2),</pre>
     sigma=rep(0,2), log.phi=0))
PGF <- function(Data) return(c(rmvn(1, rep(0, Data$K),
    rhalfcauchy(1,25)^2 *exp(-runif(1,1,5)*Data$D)),
     rnormv(2,0,1000), log(rhalfcauchy(2,25)), runif(1,1,5)))
MyData <- list(D=D, D.new=D.new, D.P=D.P, K=K, N=N, PGF=PGF, X=X,
    Xnew=Xnew, latitude=latitude, longitude=longitude,
```

mon.names=mon.names, parm.names=parm.names, y=y)

38.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    zeta <- parm[grep("zeta", Data$parm.names)]</pre>
    kappa <- 1
    sigma <- interval(parm[grep("sigma", Data$parm.names)], 0, Inf)</pre>
    parm[grep("sigma", Data$parm.names)] <- sigma</pre>
    phi <- exp(parm[grep("log.phi", Data$parm.names)])</pre>
    Sigma <- sigma[2] *sigma[2] * exp(-phi * Data$D)^kappa
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    zeta.prior <- dmvn(zeta, rep(0, Data$K), Sigma, log=TRUE)</pre>
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    phi.prior <- dunif(phi, 1, 5, log=TRUE)</pre>
    ### Interpolation
    rho <- exp(-phi * Data$D.new)^kappa</pre>
     ynew <- rnorm(1, sum(beta * Data$Xnew) + sum(rho / sum(rho) * zeta),</pre>
         sigma)
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta))</pre>
    mu[1:Data$K] <- mu[1:Data$K] + zeta</pre>
    lambda <- exp(-phi * Data$D.P)^kappa</pre>
    mu[(Data$K+1):Data$N] <- mu[(Data$K+1):Data$N] +</pre>
         rowSums(lambda / rowSums(lambda) *
         matrix(zeta, Data$N - Data$K, Data$K, byrow=TRUE))
    LL <- sum(dnorm(Data$y, mu, sigma[1], log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + beta.prior + zeta.prior + sigma.prior + phi.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma,ynew),</pre>
         yhat=rnorm(length(mu), mu, sigma[1]), parm=parm)
    return(Modelout)
    }
```

38.4. Initial Values

```
Initial. Values \leftarrow c(rep(0,K), c(mean(y), 0), rep(0,2), log(1))
```

39. Laplace Regression

This linear regression specifies that \mathbf{y} is Laplace-distributed, where it is usually Gaussian

or normally-distributed. It has been claimed that it should be surprising that the normal distribution became the standard, when the Laplace distribution usually fits better and has wider tails (Kotz, Kozubowski, and Podgorski 2001). Another popular alternative is to use the t-distribution (see Robust Regression in section 69), though it is more computationally expensive to estimate, because it has three parameters. The Laplace distribution has only two parameters, location and scale like the normal distribution, and is computationally easier to fit. This example could be taken one step further, and the parameter vector β could be Laplace-distributed. Laplace's Demon recommends that users experiment with replacing the normal distribution with the Laplace distribution.

39.1. Form

$$\mathbf{y} \sim \mathcal{L}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

39.2. Data

39.3. Model

```
Model <- function(parm, Data)
    {
     ### Parameters
     beta <- parm[1:Data$J]
     sigma <- exp(parm[Data$J+1])
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
     sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)
     ### Log-Likelihood</pre>
```

Initial.Values <- c(rep(0,J), log(1))</pre>

40. Linear Regression

40.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

40.2. Data

```
Model <- function(parm, Data)
{</pre>
```

40.4. Initial Values

Initial.Values <- c(rep(0,J), log(1))</pre>

41. Linear Regression, Frequentist

By eliminating prior probabilities, a frequentist linear regression example is presented. Although frequentism is not endorsed here, the purpose of this example is to illustrate how the **LaplacesDemon** package can be used for Bayesian or frequentist inference.

41.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$
$$\mu = \mathbf{X}\beta$$

41.3. Model

41.4. Initial Values

Initial.Values <- c(rep(0,J), log(1))</pre>

42. Linear Regression, Hierarchical Bayesian

42.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{N}(\gamma, \delta), \quad j = 1, \dots, J$$

$$\gamma \sim \mathcal{N}(0, 1000)$$

$$\delta \sim \mathcal{HC}(25)$$

$$\sigma \sim \mathcal{HC}(\tau)$$

$$\tau \sim \mathcal{HC}(25)$$

```
data(demonsnacks)
y <- log(demonsnacks$Calories)
X <- cbind(1, as.matrix(demonsnacks[,c(7,8,10)]))
J <- ncol(X)</pre>
```

```
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}</pre>
mon.names <- c("LP","delta","sigma","tau")</pre>
parm.names <- as.parm.names(list(beta=rep(0,J), gamma=0, log.delta=0,</pre>
     log.sigma=0, log.tau=0))
PGF <- function(Data) return(c(rnormv(Data$J,0,1000), rnormv(1,0,1000),
    log(rhalfcauchy(3,25))))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names,</pre>
    parm.names=parm.names, y=y)
42.3. Model
Model <- function(parm, Data)</pre>
    ### Hyperparameters
     gamma <- parm[Data$J+1]</pre>
    delta <- exp(parm[Data$J+2])</pre>
    tau <- exp(parm[Data$J+4])</pre>
    ### Parameters
    beta <- parm[1:Data$J]</pre>
    sigma <- exp(parm[Data$J+3])</pre>
    ### Log(Hyperprior Densities)
    gamma.prior <- dnormv(gamma, 0, 1000, log=TRUE)</pre>
    delta.prior <- dhalfcauchy(delta, 25, log=TRUE)</pre>
     tau.prior <- dhalfcauchy(tau, 25, log=TRUE)</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, gamma, delta, log=TRUE))</pre>
    sigma.prior <- dhalfcauchy(sigma, tau, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta))</pre>
    LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + gamma.prior + delta.prior + sigma.prior +
         tau.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,delta,sigma,tau),</pre>
         yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
42.4. Initial Values
```

43. Linear Regression, Multilevel

Initial. Values $\leftarrow c(rep(0,J), 0, rep(1,3))$

43.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^{2})$$

$$\mu_{i} = \mathbf{X}\beta_{\mathbf{m}[i], 1:J}$$

$$\beta_{g,1:J} \sim \mathcal{N}_{J}(\gamma, \Omega^{-1}), \quad g = 1, \dots, G$$

$$\Omega \sim \mathcal{W}_{J+1}(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_{J}$$

$$\gamma_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

where **m** is a vector of length N, and each element indicates the multilevel group (g = 1, ..., G) for the associated record.

43.2. Data

```
N <- 30
J <- 2 ### Number of predictors (including intercept)
G <- 2 ### Number of Multilevel Groups
X <- matrix(rnorm(N,0,1),N,J); X[,1] <- 1</pre>
Sigma <- matrix(runif(J*J,-1,1),J,J)</pre>
diag(Sigma) <- runif(J,1,5)</pre>
Sigma <- as.positive.definite(Sigma)</pre>
gamma <- runif(J,-1,1)</pre>
beta <- matrix(NA,G,J)
for (g in 1:G) {beta[g,] <- rmvn(1, gamma, Sigma)}</pre>
m <- rcat(N, rep(1/G,G)) ### Multilevel group indicator</pre>
y \leftarrow rowSums(beta[m,] * X) + rnorm(N,0,0.1)
S \leftarrow diag(J)
mon.names <- c("LP", "sigma")</pre>
parm.names <- as.parm.names(list(beta=matrix(0,G,J), log.sigma=0,</pre>
     gamma=rep(0,J), U=S), uppertri=c(0,0,0,1))
PGF <- function(Data) return(c(rmvnpc(Data$G, rnormv(Data$J,0,100),
    rwishartc(Data$J+1, Data$S)), log(rhalfcauchy(1,25)),
    rnormv(Data$J,0,100),
    upper.triangle(rwishartc(Data$J+1, Data$S), diag=TRUE)))
MyData <- list(G=G, J=J, N=N, PGF=PGF, S=S, X=X, m=m, mon.names=mon.names,
    parm.names=parm.names, y=y)
```

```
Model <- function(parm, Data)
    {
     ### Parameters
    beta <- matrix(parm[1:(Data$G * Data$J)], Data$G, Data$J)
     gamma <- parm[grep("gamma", Data$parm.names)]</pre>
```

```
sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
U <- as.parm.matrix(U, Data$J, parm, Data, chol=TRUE)</pre>
diag(U) <- exp(diag(U))</pre>
### Log(Prior Densities)
U.prior <- dwishartc(U, Data$J+1, Data$S, log=TRUE)</pre>
beta.prior <- sum(dmvnpc(beta, gamma, U, log=TRUE))</pre>
gamma.prior <- sum(dnormv(gamma, 0, 100, log=TRUE))</pre>
sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
### Log-Likelihood
mu <- rowSums(beta[Data$m,] * Data$X)</pre>
LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))</pre>
### Log-Posterior
LP <- LL + U.prior + beta.prior + gamma.prior + sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, sigma),</pre>
     yhat=rnorm(length(mu), mu, sigma), parm=parm)
return(Modelout)
}
```

43.4. Initial. Values

```
Initial.Values <- c(rep(0,G*J), log(1), rep(0,J),
    upper.triangle(S, diag=TRUE))</pre>
```

44. Linear Regression with Full Missingness

With 'full missingness', there are missing values for both the dependent variable (DV) and at least one independent variable (IV). The 'full likelihood' approach to full missingness is excellent as long as the model is identifiable. When it is not identifiable, imputation may be done in a previous stage. In this example, matrix α is for regression effects for IVs, vector β is for regression effects for the DV, vector γ is for missing values for IVs, and δ is for missing values for the DV.

$$\mathbf{y}^{imp} \sim \mathcal{N}(\nu, \sigma_J^2)$$
 $\mathbf{X}^{imp} \sim \mathcal{N}(\mu, \sigma_{-J}^2)$
 $\nu = \mathbf{X}^{imp} \beta$
 $\mu = \mathbf{X}^{imp} \alpha$
 $\mathbf{y}^{imp} = \begin{cases} \delta & \text{if } \mathbf{y}^{mis} \\ \mathbf{y}^{obs} \end{cases}$
 $\mathbf{X}^{imp} = \begin{cases} \gamma & \text{if } \mathbf{X}^{mis} \end{cases}$

```
\alpha_{j,l} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1), \quad l = 1, \dots, (J-1)
\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J
\gamma_m \sim \mathcal{N}(0, 1000), \quad m = 1, \dots, M
\delta_p \sim \mathcal{N}(0, 1000), \quad p = 1, \dots, P
\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, J
```

```
N <- 100
J <- 5
X \leftarrow matrix(runif(N*J,-2,2),N,J); X[,1] \leftarrow 1 \#Design matrix X
M \leftarrow matrix(round(runif(N*J)-0.45),N,J); M[,1] \leftarrow 0 \#Missing indicators
X <- ifelse(M == 1, NA, X) #Simulated X gets missings according to M
beta.orig <- runif(J,-2,2)
y <- as.vector(tcrossprod(X, t(beta.orig)) + rnorm(N,0,0.1))
y[sample(1:N, round(N*.05))] <- NA
m <- ifelse(is.na(y), 1, 0) #Missing indicator for vector y
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=matrix(0,J-1,J-1),</pre>
    beta=rep(0,J),
    gamma=rep(0,sum(is.na(X))),
    delta=rep(0,sum(is.na(y))),
    log.sigma=rep(0,J)))
PGF <- function(Data) return(c(rnormv((Data$J-1)*(Data$J-1),0,10),
    rnormv(Data$J,0,10),
    rnormv(sum(is.na(Data$X)),0,10),
    rnormv(sum(is.na(Data$y)),mean(Data$y, na.rm=TRUE),1),
    log(rhalfcauchy(Data$J,25))))
MyData <- list(J=J, N=N, PGF=PGF, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
44.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    alpha <- matrix(parm[grep("alpha", Data$parm.names)], Data$J-1,</pre>
         Data$J-1)
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    gamma <- parm[grep("gamma", Data$parm.names)]</pre>
    delta <- parm[grep("delta", Data$parm.names)]</pre>
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
```

```
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
gamma.prior <- sum(dnormv(gamma, 0, 1000, log=TRUE))</pre>
delta.prior <- sum(dnormv(delta, 0, 1000, log=TRUE))</pre>
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
### Log-Likelihood
mu <- X.imputed <- Data$X
X.imputed[which(is.na(X.imputed))] <- gamma</pre>
y.imputed <- Data$y</pre>
y.imputed[which(is.na(y.imputed))] <- delta</pre>
for (j in 2:Data$J) {mu[,j] <- tcrossprod(X.imputed[,-j],</pre>
     t(alpha[,(j-1)]))}
nu <- tcrossprod(X.imputed, t(beta))</pre>
LL <- sum(dnorm(X.imputed[,-1], mu[,-1],
    matrix(sigma[1:(Data$J-1)], Data$N, Data$J-1), log=TRUE),
     dnorm(y.imputed, nu, sigma[Data$J], log=TRUE))
### Log-Posterior
LP <- LL + alpha.prior + beta.prior + gamma.prior + delta.prior +
     sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
     yhat=rnorm(length(nu), nu, sigma[Data$J]), parm=parm)
return(Modelout)
}
```

44.4. Initial Values

```
Initial.Values <- c(rep(0, (J-1)^2), rep(0,J), rep(0,sum(is.na(X))),
    rep(0,sum(is.na(y))), rep(1,J))</pre>
```

45. Linear Regression with Missing Response

This is an introductory example to missing values using data augmentation with auxiliary variables. The dependent variable, or response, has both observed values, \mathbf{y}^{obs} , and missing values, \mathbf{y}^{mis} . The α vector is for missing value imputation, and enables the use of the full-likelihood by augmenting te state with these auxiliary variables. In the model form, M is used to denote the number of missing values, though it is used as an indicator in the data.

$$\mathbf{y}^{imp} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mathbf{y}^{imp} = \begin{cases} \alpha & \text{if } \mathbf{y}^{mis} \\ \mathbf{y}^{obs} & \\ \mu = \mathbf{X}\beta & \\ \alpha_m \sim \mathcal{N}(0, 1000), \quad m = 1, \dots, M \\ \beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J \end{cases}$$

 $\sigma \sim \mathcal{HC}(25)$

45.2. Data

}

```
data(demonsnacks)
N <- nrow(demonsnacks)</pre>
J <- ncol(demonsnacks)</pre>
y <- log(demonsnacks$Calories)</pre>
y[sample(1:N, round(N*0.05))] <- NA
M \leftarrow ifelse(is.na(y), 1, 0)
X <- cbind(1, as.matrix(demonsnacks[,c(1,3:10)]))</pre>
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}</pre>
mon.names <- c("LP", "sigma")</pre>
parm.names <- as.parm.names(list(alpha=rep(0,sum(M)), beta=rep(0,J),</pre>
     log.sigma=0))
PGF <- function(Data) return(c(rnorm(sum(Data$M),mean(y,na.rm=TRUE),1),
    rnormv(Data$J,0,1000), log(rhalfcauchy(1,25))))
MyData <- list(J=J, M=M, PGF=PGF, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
45.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
     alpha <- parm[1:sum(Data$M)]</pre>
    beta <- parm[sum(Data$M)+1:Data$J]</pre>
     sigma <- exp(parm[sum(Data$M)+Data$J+1])</pre>
    ### Log(Prior Densities)
     alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE)</pre>
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
     sigma.prior <- dgamma(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta))</pre>
    y.imputed <- Data$y
    y.imputed[which(is.na(Data$y))] <- alpha</pre>
    LL <- sum(dnorm(y.imputed, mu, sigma, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, sigma),</pre>
          yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
```

45.4. Initial Values

Initial. Values $\leftarrow c(rep(0, sum(M)), rep(0, J), log(1))$

46. Linear Regression with Missing Response via ABB

The Approximate Bayesian Bootstrap (ABB), using the ABB function, is used to impute missing values in the dependent variable (DV), or response, given a propensity score. In this example, vector α is used to estimate propensity score η , while vector β is for regression effects, and vector γ has the monitored missing values. For more information on ABB, see the ABB function.

46.1. Form

$$\mathbf{y}^{imp} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mathbf{y}^{imp} = \begin{cases} \gamma & \text{if } \mathbf{y}^{mis} \\ \mathbf{y}^{obs} & \text{if } \mathbf{y}^{mis} \end{cases}$$

$$\mu = \mathbf{X}\beta$$

$$\gamma \sim p(\mathbf{y}^{obs}|\mathbf{y}^{obs}, \mathbf{y}^{mis}, \eta)$$

$$\eta = \frac{1}{1 + \exp(-\nu)}$$

$$\nu = \mathbf{X}\alpha$$

$$\alpha_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

```
data(demonsnacks)
N <- nrow(demonsnacks)
J <- ncol(demonsnacks)
y <- log(demonsnacks$Calories)
y[sample(1:N, round(N*0.05))] <- NA
M <- ifelse(is.na(y), 1, 0)
X <- cbind(1, as.matrix(demonsnacks[,c(1,3:10)]))
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}
mon.names <- c("LP","sigma",paste("gamma[",1:sum(is.na(y)),"]",sep=""))
parm.names <- as.parm.names(list(alpha=rep(0,J), beta=rep(0,J), log.sigma=0))
PGF <- function(Data) return(c(rnormv(Data$J,0,10), rnormv(Data$J,0,10), log(rhalfcauchy(1,25))))
MyData <- list(J=J, M=M, PGF=PGF, X=X, mon.names=mon.names,</pre>
```

```
parm.names=parm.names, y=y)
```

46.3. Model

```
Model <- function(parm, Data)</pre>
     ### Parameters
     alpha <- parm[1:Data$J]
    beta <- parm[Data$J+1:Data$J]</pre>
     sigma <- exp(parm[2*Data$J+1])</pre>
     ### Log(Prior Densities)
     alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
     sigma.prior <- dgamma(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    y.imputed <- Data$y
    mu <- tcrossprod(Data$X, t(beta))</pre>
    nu <- as.vector(tcrossprod(Data$X, t(alpha)))</pre>
    eta <- invlogit(nu)</pre>
    breaks <- as.vector(quantile(eta, probs=c(0,0.2,0.4,0.6,0.8,1)))</pre>
    B <- matrix(breaks[-length(breaks)], length(Data$y), 5, byrow=TRUE)
    z <- rowSums(eta >= B)
    for (i in 1:5) {
         if(any(is.na(Data$y[which(z == i)]))) {
              imp <- unlist(ABB(Data$y[which(z == i)]))</pre>
              y.imputed[which({z == i} & is.na(Data$y))] <- imp}}</pre>
    gamma <- y.imputed[which(is.na(Data$y))]</pre>
     LL <- sum(dbern(Data$M, eta, log=TRUE),
         dnorm(y.imputed, mu, sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma,gamma),</pre>
         yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

46.4. Initial Values

47. Linear Regression with Power Priors

Power priors (Ibrahim and Chen 2000) are a class of informative priors when relevant historical data is available. Power priors may be used when it is desirable to take historical data into account while analyzing similar, current data. Both the current data, \mathbf{y} and \mathbf{X} , and historical data, \mathbf{y}_h and \mathbf{X}_h , are included in the power prior analysis, where h indicates historical data.

Each data set receives its own likelihood function, though the likelihood of the historical data is raised to an exponential power, $\alpha \in [0, 1]$. In this example, α is a constant.

47.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mathbf{y}_h \sim \mathcal{N}(\mu_h, \sigma^2)^{\alpha}$$

$$\mu = \mathbf{X}\beta$$

$$\mu_h = \mathbf{X}_h\beta$$

$$\alpha = 0.5$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

47.2. Data

```
N <- 100
J <- 5 #Number of predictors, including the intercept
X \leftarrow Xh \leftarrow matrix(1,N,J)
for (j in 2:J) {
    X[,j] \leftarrow rnorm(N,runif(1,-3,3),runif(1,0.1,1))
    Xh[,j] \leftarrow rnorm(N,runif(1,-3,3),runif(1,0.1,1))
beta.orig <- runif(J,-3,3)</pre>
e <- rnorm(N,0,0.1)
yh <- as.vector(tcrossprod(beta.orig, Xh) + e)</pre>
y <- as.vector(tcrossprod(beta.orig, X) + e)
mon.names <- c("LP", "sigma")</pre>
parm.names <- as.parm.names(list(beta=rep(0,J), log.sigma=0))</pre>
PGF <- function(Data) return(c(rnormv(Data$J,0,1000),
    log(rhalfcauchy(1,25))))
MyData <- list(alpha=0.5, J=J, PGF=PGF, X=X, Xh=Xh, mon.names=mon.names,
    parm.names=parm.names, y=y, yh=yh)
```

```
Model <- function(parm, Data)
    {
     ### Parameters
     beta <- parm[1:Data$J]
     sigma <- exp(parm[Data$J+1])
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
```

Initial.Values <- c(rep(0,J), log(1))</pre>

48. LSTAR.

This is a Logistic Smooth-Threshold Autoregression (LSTAR), and is specified with a transition function that includes γ as the shape parameter, \mathbf{y} as the transition variable, θ as the location parameter, and d as the delay parameter.

48.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma^{2}), \quad t = 1, \dots, T$$

$$\mu_{t} = \pi_{t}(\alpha_{1} + \phi_{1}\mathbf{y}_{t-1}) + (1 - \pi_{t})(\alpha_{2} + \phi_{2}\mathbf{y}_{t-1}), \quad t = 2, \dots, T$$

$$\pi_{t} = \frac{1}{1 + \exp(-(\gamma(\mathbf{y}_{t-d} - \theta)))}$$

$$\alpha_{j} \sim \mathcal{N}(0, 1000) \in [\mathbf{y}_{min}, \mathbf{y}_{max}], \quad j = 1, \dots, 2$$

$$\phi_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, 2$$

$$\gamma \sim \mathcal{HC}(25)$$

$$\theta \sim \mathcal{U}(\mathbf{y}_{min}, \mathbf{y}_{max})$$

$$\pi_{1} \sim \mathcal{U}(0.001, 0.999)$$

$$\sigma \sim \mathcal{HC}(25)$$

```
y <- c(26.73, 26.75, 26.24, 25.94, 27.40, 26.14, 23.99, 23.08, 22.55, 20.64, 23.28, 24.92, 25.07, 26.53, 28.14, 30.10, 27.43, 27.24, 23.96, 25.85, 26.76, 26.05, 26.79, 26.69, 29.89, 29.09, 23.84,
```

```
24.87, 24.47, 22.85, 22.05, 22.82, 22.99, 21.60, 20.32, 20.80,
    19.78, 19.87, 18.78, 19.64, 20.00, 21.51, 21.49, 21.96, 22.58,
    21.22, 22.34, 22.76, 18.37, 17.50, 17.55, 12.14, 4.76, 3.75,
    2.05, 2.69, 3.85, 4.72, 5.00, 3.31, 3.02, 3.15, 2.50,
    3.33, 3.95, 4.00, 3.86, 3.87, 3.51, 3.19, 2.39, 2.33,
    2.57, 2.80, 2.43, 2.43, 2.10, 2.31, 2.21, 2.11, 2.10,
    1.70, 1.35, 1.83, 1.55, 1.63, 1.91, 2.14, 2.41, 2.06,
    1.87, 2.11, 2.28, 2.26, 2.03, 2.06, 2.08, 1.91, 1.95,
    1.56, 1.44, 1.60, 1.77, 1.77, 1.95, 2.01, 1.65, 1.87,
    2.01, 1.84, 1.94, 1.93, 1.93, 1.75, 1.73, 1.80, 1.74,
    1.80, 1.75, 1.67, 1.60, 1.61, 1.55, 1.56, 1.57, 1.55,
    1.56, 1.57, 1.69, 1.66, 1.74, 1.64, 1.65, 1.62, 1.54,
    1.58, 1.49, 1.41, 1.42, 1.37, 1.45, 1.31, 1.37, 1.26,
    1.35, 1.41, 1.29, 1.28, 1.23, 1.08, 1.03, 1.00, 1.04,
    1.04, 0.92, 0.96, 0.90, 0.85, 0.78, 0.73, 0.59, 0.54,
    0.53, 0.41, 0.46, 0.52, 0.42, 0.42, 0.43, 0.43, 0.35,
    0.35, 0.35, 0.42, 0.41, 0.41, 0.50, 0.83, 0.96, 1.38,
    1.62, 1.26, 1.48, 1.39, 1.20, 1.10, 1.02, 0.95, 1.00,
    1.07, 1.14, 1.14, 1.10, 1.05, 1.08, 1.16, 1.42, 1.52,
    1.60, 1.69, 1.62, 1.29, 1.46, 1.43, 1.50, 1.46, 1.40,
    1.34, 1.41, 1.38, 1.38, 1.46, 1.73, 1.84, 1.95, 2.01,
    1.90, 1.81, 1.60, 1.84, 1.72, 1.83, 1.81, 1.78, 1.80,
    1.70, 1.70, 1.66, 1.67, 1.69, 1.66, 1.56, 1.47, 1.64,
    1.71, 1.66, 1.65, 1.60, 1.61, 1.61, 1.53, 1.48, 1.40,
    1.47, 1.53, 1.39, 1.41, 1.42, 1.46, 1.46, 1.33, 1.16)
T <- length(y)
mon.names <- c("LP", "sigma", "ynew", paste("pi[", 1:T, "]", sep=""),
     "pi.new")
parm.names <- as.parm.names(list(alpha=rep(0,2), phi=rep(0,2), log.gamma=0,
    theta=0, pi=0, log.sigma=0))
PGF <- function(Data) return(c(rnormv(2,0,10), rnormv(2,0,10),
    log(rhalfcauchy(1,25)), runif(1,min(Data$y),max(Data$y)),
    runif(1,0.001,0.999), log(rhalfcauchy(1,25))))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names,</pre>
    y=y)
48.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    parm[1:2] <- alpha <- interval(parm[1:2], min(Data$y), max(Data$y))</pre>
    parm[3:4] <- phi <- interval(parm[3:4], -1, 1)
    gamma <- exp(parm[5])</pre>
    parm[6] <- theta <- interval(parm[6], min(Data$y), max(Data$y))</pre>
    parm[7] <- pi <- interval(parm[7], 0.001, 0.999)</pre>
```

```
sigma <- exp(parm[8])</pre>
### Log(Prior Densities)
alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
phi.prior <- sum(dnormv(phi, 0, 1000, log=TRUE))</pre>
gamma.prior <- dhalfcauchy(gamma, 25, log=TRUE)</pre>
theta.prior <- dunif(theta, min(Data$y), max(Data$y), log=TRUE)</pre>
pi.prior <- dunif(pi, 0.001, 0.999, log=TRUE)</pre>
sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
### Log-Likelihood
pi \leftarrow c(pi, 1 / (1 + exp(-(gamma*(Data$y[-Data$T]-theta)))))
pi.new <- 1 / (1 + exp(-(gamma*(Data$y[Data$T]-theta))))</pre>
mu <- pi * c(alpha[1], alpha[1] + phi[1]*Data$y[-Data$T]) +</pre>
     (1-pi) * c(alpha[2], alpha[2] + phi[2]*Data$y[-Data$T])
ynew <- rnorm(1, pi.new * (alpha[1] + phi[1]*Data$y[Data$T]) +</pre>
     (1-pi.new) * (alpha[2] + phi[2] *Data$y[Data$T]), sigma)
LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))
### Log-Posterior
LP <- LL + alpha.prior + phi.prior + gamma.prior + theta.prior +
    pi.prior + sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma,ynew,pi,pi.new),</pre>
     yhat=rnorm(length(mu), mu, sigma), parm=parm)
return(Modelout)
}
```

Initial. Values $\leftarrow c(rep(min(y), 4), log(1), mean(y), 0.5, log(1))$

49. MANCOVA

Since this is a multivariate extension of ANCOVA, please see the ANCOVA example in section 1 for a univariate introduction.

$$\mathbf{Y}_{i,1:J} \sim \mathcal{N}_{K}(\mu_{i,1:J}, \Sigma), \quad i = 1, \dots, N$$

$$\mu_{i,k} = \alpha_{k} + \beta_{k,\mathbf{X}[i,1]} + \gamma_{k,\mathbf{X}[i,1]} + \mathbf{X}_{1:N,3:(C+J)} \delta_{k,1:C}$$

$$\epsilon_{i,k} = \mathbf{Y}_{i,k} - \mu_{i,k}$$

$$\alpha_{k} \sim \mathcal{N}(0, 1000), \quad k = 1, \dots, K$$

$$\beta_{k,l} \sim \mathcal{N}(0, \sigma_{1}^{2}), \quad l = 1, \dots, (L-1)$$

$$\beta_{1:K,L} = -\sum_{l=1}^{L-1} \beta_{1:K,l}$$

$$\gamma_{k,m} \sim \mathcal{N}(0, \sigma_2^2), \quad m = 1, \dots, (M-1)$$

$$\gamma_{1:K,M} = -\sum_{m=1}^{M-1} \beta_{1:K,m}$$

$$\delta_{k,c} \sim \mathcal{N}(0, 1000)$$

$$\Omega \sim \mathcal{W}_{K+1}(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_K$$

$$\Sigma = \Omega^{-1}$$

$$\sigma_{1:J} \sim \mathcal{HC}(25)$$

```
C <- 2 #Number of covariates
J <- 2 #Number of factors (treatments)</pre>
K <- 3 #Number of endogenous (dependent) variables
L <- 4 #Number of levels in factor (treatment) 1
M <- 5 #Number of levels in factor (treatment) 2
N <- 100
X \leftarrow \text{matrix}(\text{cbind}(\text{round}(\text{runif}(N, 0.5, L+0.49)), \text{round}(\text{runif}(N, 0.5, M+0.49)),
     runif(C*N,0,1)), N, J + C)
alpha <- runif(K,-1,1)
beta <- matrix(runif(K*L,-2,2), K, L)
beta[,L] <- -rowSums(beta[,-L])</pre>
gamma <- matrix(runif(K*M,-2,2), K, M)</pre>
gamma[,M] <- -rowSums(gamma[,-M])</pre>
delta <- matrix(runif(K*C), K, C)</pre>
Y <- matrix(NA,N,K)
for (k in 1:K) {
    Y[,k] \leftarrow alpha[k] + beta[k,X[,1]] + gamma[k,X[,2]] +
     tcrossprod(delta[k,], X[,-c(1,2)]) + rnorm(1,0,0.1)
S \leftarrow diag(K)
mon.names <- c("LP", "s.o.beta", "s.o.gamma", "s.o.epsilon",
     as.parm.names(list(s.beta=rep(0,K), s.gamma=rep(0,K),
     s.epsilon=rep(0,K))))
parm.names <- as.parm.names(list(alpha=rep(0,K), beta=matrix(0,K,(L-1)),</pre>
    gamma=matrix(0,K,(M-1)), delta=matrix(0,K,C), U=diag(K),
     log.sigma=rep(0,2)), uppertri=c(0,0,0,0,1,0))
PGF <- function(Data) return(c(rnormv(Data$K,0,1000),
    rnorm(Data$K*(Data$L-1),0,rhalfcauchy(1,25)),
    rnorm(Data$K*(Data$M-1),0,rhalfcauchy(1,25)),
    rnormv(Data$K*Data$C,0,1000),
    upper.triangle(rwishartc(Data$K+1,Data$S), diag=TRUE),
     log(rhalfcauchy(2,25))))
MyData <- list(C=C, J=J, K=K, L=L, M=M, N=N, PGF=PGF, S=S, X=X, Y=Y,
     mon.names=mon.names, parm.names=parm.names)
```

```
Model <- function(parm, Data)</pre>
    ### Parameters
     alpha <- parm[grep("alpha", Data$parm.names)]</pre>
    beta <- matrix(c(parm[grep("beta", Data$parm.names)], rep(0,Data$K)),</pre>
    Data$K, Data$L)
    beta[,Data$L] <- -rowSums(beta[,-Data$L])</pre>
     gamma <- matrix(c(parm[grep("gamma", Data$parm.names)],</pre>
          rep(0,Data$K)), Data$K, Data$M)
    gamma[,Data$M] <- -rowSums(gamma[,-Data$M])</pre>
    delta <- matrix(parm[grep("delta", Data$parm.names)], Data$K, Data$C)</pre>
    U <- as.parm.matrix(U, Data$K, parm, Data, chol=TRUE)</pre>
     diag(U) <- exp(diag(U))</pre>
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
     alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
    beta.prior <- sum(dnorm(beta, 0, sigma[1], log=TRUE))</pre>
     gamma.prior <- sum(dnorm(gamma, 0, sigma[2], log=TRUE))</pre>
     delta.prior <- sum(dnormv(delta, 0, 1000, log=TRUE))</pre>
    U.prior <- dwishartc(U, Data$K+1, Data$S, log=TRUE)</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- matrix(0,Data$N,Data$K)</pre>
    for (k in 1:Data$K) {
         mu[,k] \leftarrow alpha[k] + beta[k,Data$X[,1]] + gamma[k,Data$X[,2]] +
          tcrossprod(Data$X[,-c(1,2)], t(delta[k,]))}
    LL <- sum(dmvnpc(Data$Y, mu, U, log=TRUE))
    ### Variance Components, Omnibus
    s.o.beta <- sd(as.vector(beta))</pre>
     s.o.gamma <- sd(as.vector(gamma))</pre>
     s.o.epsilon <- sd(as.vector(Data$Y - mu))</pre>
    ### Variance Components, Univariate
     s.beta <- apply(beta,1,sd)</pre>
     s.gamma <- apply(gamma,1,sd)</pre>
     s.epsilon <- apply(Data$Y - mu,2,sd)</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + gamma.prior + delta.prior +
          U.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, s.o.beta, s.o.gamma,
          s.o.epsilon, s.beta, s.gamma, s.epsilon),
          yhat=rmvnpc(nrow(mu), mu, U), parm=parm)
    return(Modelout)
    }
```

49.4. Initial Values

Initial.Values <- c(rep(0,K), rep(0,K*(L-1)), rep(0,K*(M-1)),
 rep(0,C*K), upper.triangle(S, diag=TRUE), rep(0,2))</pre>

50. MANOVA

Since this is a multivariate extension of ANOVA, please see the two-way ANOVA example in section 3 for a univariate introduction.

50.1. Form

$$\mathbf{Y}_{i,1:J} \sim \mathcal{N}_{K}(\mu_{i,1:J}, \Omega^{-1}), \quad i = 1, \dots, N$$

$$\mu_{i,k} = \alpha_{k} + \beta_{k,\mathbf{X}[i,1]} + \gamma_{k,\mathbf{X}[i,1]}$$

$$\epsilon_{i,k} = \mathbf{Y}_{i,k} - \mu_{i,k}$$

$$\alpha_{k} \sim \mathcal{N}(0, 1000), \quad k = 1, \dots, K$$

$$\beta_{k,l} \sim \mathcal{N}(0, \sigma_{1}^{2}), \quad l = 1, \dots, (L-1)$$

$$\beta_{1:K,L} = -\sum_{l=1}^{L-1} \beta_{1:K,l}$$

$$\gamma_{k,m} \sim \mathcal{N}(0, \sigma_{2}^{2}), \quad m = 1, \dots, (M-1)$$

$$\gamma_{1:K,M} = -\sum_{m=1}^{M-1} \beta_{1:K,m}$$

$$\Omega \sim \mathcal{W}_{K+1}(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_{K}$$

$$\sigma_{1:J} \sim \mathcal{HC}(25)$$

```
for (k in 1:K) {
    Y[,k] \leftarrow alpha[k] + beta[k,X[,1]] + gamma[k,X[,2]] + rnorm(1,0,0.1)
S \leftarrow diag(K)
mon.names <- c("LP", "s.o.beta", "s.o.gamma", "s.o.epsilon",
    as.parm.names(list(s.beta=rep(0,K), s.gamma=rep(0,K),
    s.epsilon=rep(0,K))))
parm.names <- as.parm.names(list(alpha=rep(0,K), beta=matrix(0,K,(L-1)),</pre>
    gamma=matrix(0,K,(M-1)), U=diag(K), log.sigma=rep(0,2)),
    uppertri=c(0,0,0,1,0))
PGF <- function(Data) return(c(rnormv(Data$K,0,1000),
    rnorm(Data$K*(Data$L-1),0,rhalfcauchy(1,25)),
    rnorm(Data$K*(Data$M-1),0,rhalfcauchy(1,25)),
    upper.triangle(rwishartc(Data$K+1,Data$S), diag=TRUE),
    log(rhalfcauchy(2,25))))
MyData <- list(J=J, K=K, L=L, M=M, N=N, PGF=PGF, S=S, X=X, Y=Y,
    mon.names=mon.names, parm.names=parm.names)
50.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    alpha <- parm[grep("alpha", Data$parm.names)]</pre>
    beta <- matrix(c(parm[grep("beta", Data$parm.names)], rep(0,Data$K)),</pre>
         Data$K, Data$L)
    beta[,Data$L] <- -rowSums(beta[,-Data$L])</pre>
    gamma <- matrix(c(parm[grep("gamma", Data$parm.names)],</pre>
         rep(0,Data$K)), Data$K, Data$M)
    gamma[,Data$M] <- -rowSums(gamma[,-Data$M])</pre>
    U <- as.parm.matrix(U, Data$K, parm, Data, chol=TRUE)</pre>
    diag(U) <- exp(diag(U))</pre>
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
    beta.prior <- sum(dnorm(beta, 0, sigma[1], log=TRUE))</pre>
    gamma.prior <- sum(dnorm(gamma, 0, sigma[2], log=TRUE))</pre>
    U.prior <- dwishartc(U, Data$K+1, Data$S, log=TRUE)</pre>
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- matrix(0,Data$N,Data$K)</pre>
    for (k in 1:Data$K) {
         mu[,k] <- alpha[k] + beta[k,Data$X[,1]] + gamma[k,Data$X[,2]]}</pre>
    LL <- sum(dmvnpc(Data$Y, mu, U, log=TRUE))
    ### Variance Components, Omnibus
    s.o.beta <- sd(as.vector(beta))</pre>
    s.o.gamma <- sd(as.vector(gamma))</pre>
```

50.4. Initial Values

```
Initial.Values <- c(rep(0,K), rep(0,K*(L-1)), rep(0,K*(M-1)),
     upper.triangle(S, diag=TRUE), rep(0,2))</pre>
```

51. Mixture Model, Finite

This finite mixture model (FMM) imposes a multilevel structure on each of the J regression effects in β , so that mixture components share a common residual standard deviation, ν_j . Identifiability is gained at the expense of some shrinkage.

$$\mathbf{y} \sim \mathcal{N}(\mu_{1:N,m}, \sigma^2)$$

$$\mu_{1:N,m} = \mathbf{X}\beta_{m,1:J}, \quad m = 1, \dots, M$$

$$\beta_{m,j} \sim \mathcal{N}(0, \nu_j^2), \quad j = 1, \dots, J$$

$$\nu_j \sim \mathcal{HC}(25)$$

$$\sigma \sim \mathcal{HC}(25)$$

$$\pi_{1:M} \sim \mathcal{D}(\alpha_{1:M})$$

$$\pi_m = \frac{\sum_{i=1}^N \delta_{i,m}}{\sum \delta}$$

$$\mathbf{p}_{i,m} = \frac{\delta_{i,m}}{\sum_{m=1}^M \delta_{i,m}}$$

$$\delta_{i,m} = \exp(\mathbf{X}\delta_{i,m}), \quad m = 1, \dots, (M-1)$$

$$\delta_{1:N,M} = 1$$

$$\delta_{i,m} \sim \mathcal{N}(0, 1000) \in [-10, 10], \quad m = 1, \dots, (M-1)$$

 $\alpha_m = 1$

51.2. Data

```
M <- 2 #Number of mixtures
alpha <- rep(1,M) #Prior probability of mixing probabilities
data(demonsnacks)
N <- nrow(demonsnacks)</pre>
y <- log(demonsnacks$Calories)</pre>
X \leftarrow cbind(1, as.matrix(demonstracks[,c(7,8,10)]))
J \leftarrow ncol(X)
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}</pre>
mon.names <- c("LP", as.parm.names(list(pi=rep(0,M), sigma=0)))</pre>
parm.names <- as.parm.names(list(beta=matrix(0,M,J), log.nu=rep(0,J),</pre>
    log.delta=matrix(0,N,M-1), log.sigma=0))
PGF <- function(Data) return(c(rnorm(Data$M*Data$J, 0,
    matrix(rhalfcauchy(Data$J,25), Data$M, Data$J, byrow=TRUE)),
    log(rhalfcauchy(Data$J,25)),
    log(rtrunc(Data$N*(Data$M-1),"norm",a=exp(-10),b=exp(10),
    mean=log(1/Data$M), sd=sqrt(1000))), log(rhalfcauchy(1,25))))
MyData <- list(J=J, M=M, N=N, PGF=PGF, X=X, alpha=alpha,
    mon.names=mon.names, parm.names=parm.names, y=y)
51.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- matrix(parm[grep("beta", Data$parm.names)], Data$M, Data$J)</pre>
    delta <- interval(parm[grep("log.delta", Data$parm.names)], -10, 10)</pre>
    parm[grep("log.delta", Data$parm.names)] <- delta</pre>
    delta <- matrix(c(exp(delta), rep(1, Data$N)), Data$N, Data$M)
    pi <- colSums(delta) / sum(delta)</pre>
    nu <- exp(parm[grep("log.nu", Data$parm.names)])</pre>
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnorm(beta, 0, matrix(rep(nu, Data$M), Data$M,</pre>
         Data$J, byrow=TRUE), log=TRUE))
    delta.prior <- sum(dtrunc(delta, "norm", a=exp(-10), b=exp(10),
         mean=log(1/Data$M), sd=sqrt(1000), log=TRUE))
    pi.prior <- ddirichlet(pi, Data$alpha, log=TRUE)</pre>
    nu.prior <- sum(dhalfcauchy(nu, 25, log=TRUE))</pre>
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    p <- delta / rowSums(delta)</pre>
```

mu <- tcrossprod(Data\$X, beta)</pre>

51.4. Initial Values

Initial. Values \leftarrow c(runif(M*J), rep(0,J), runif(N*(M-1),-1,1), 0)

52. Mixture Model, Infinite

This infinite mixture model (IMM) imposes a multilevel structure on each of the J regression effects in β , so that mixture components share a common residual standard deviation, ν_j . The infinite number of mixture components is truncated to a finite number, and the user specifies the maximum number to explore, M, where M is discrete, greater than one, and less than the number of records, N. A truncated stick-breaking process within a truncated Dirichlet process defines the nonparametric mixture component selection.

$$\mathbf{y} \sim \mathcal{N}(\mu_{1:N,m}, \sigma^2)$$

$$\mu_{1:N,m} = \mathbf{X}\beta_{m,1:J}, \quad m = 1, \dots, M$$

$$\beta_{m,j} \sim \mathcal{N}(0, \nu_j^2), \quad j = 1, \dots, J$$

$$\nu_j \sim \mathcal{HC}(25)$$

$$\sigma \sim \mathcal{HC}(25)$$

$$\theta_i \sim \mathcal{CAT}(\pi_{i,1:C})$$

$$\pi \sim \text{Stick}(\gamma)$$

$$\mathbf{p}_{i,m} = \frac{\delta_{i,m}}{\sum_{m=1}^{M} \delta_{i,m}}$$

$$\delta_{i,m} = \exp(\mathbf{X}\delta_{i,m}), \quad m = 1, \dots, (M-1)$$

$$\delta_{1:N,M} = 1$$

$$\delta_{i,m} \sim \mathcal{N}(0, 1000) \in [-10, 10], \quad m = 1, \dots, (M-1)$$

$$\alpha \sim \mathcal{HC}(25)$$

```
\iota \sim \mathcal{HC}(25)
\gamma \sim \mathcal{G}(\alpha, \iota)
```

52.2. Data

```
M <- 3 \#Maximum number of mixtures to explore
data(demonsnacks)
N <- nrow(demonsnacks)</pre>
y <- log(demonsnacks$Calories)</pre>
X \leftarrow cbind(1, as.matrix(demonstracks[,c(7,8,10)]))
J \leftarrow ncol(X)
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}</pre>
mon.names <- c("LP", as.parm.names(list(pi=rep(0,M), sigma=0)))</pre>
parm.names <- as.parm.names(list(beta=matrix(0,M,J), log.nu=rep(0,J),</pre>
    log.delta=matrix(0,N,M-1), log.sigma=0, lambda=rep(0,M-1),
    log.alpha=0, log.iota=0, log.gamma=0))
PGF <- function(Data) return(c(rnorm(Data$M*Data$J, 0,
    matrix(rhalfcauchy(Data$J,25), Data$M, Data$J, byrow=TRUE)),
    log(rhalfcauchy(Data$J,25)),
    log(rtrunc(Data$N*(Data$M-1),"norm",a=exp(-10),b=exp(10),
    mean=log(1/Data$M), sd=sqrt(1000))), log(rhalfcauchy(1,25)),
    runif(Data$M-1,1e-5,1-1e-5), log(rhalfcauchy(2,25)),
    log(rgamma(1,rhalfcauchy(1,25),rhalfcauchy(1,25)))))
MyData <- list(J=J, M=M, N=N, PGF=PGF, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
52.3. Model
Model <- function(parm, Data)</pre>
    ### Hyperhyperparameters
    alpha <- exp(parm[grep("log.alpha", Data$parm.names)])</pre>
    iota <- exp(parm[grep("log.iota", Data$parm.names)])</pre>
    ### Hyperparameters
    gamma <- exp(parm[grep("log.gamma", Data$parm.names)])</pre>
    nu <- exp(parm[grep("log.nu",Data$parm.names)])</pre>
    ### Parameters
    beta <- matrix(parm[grep("beta", Data$parm.names)], Data$M, Data$J)</pre>
    delta <- interval(parm[grep("log.delta", Data$parm.names)], -10, 10)</pre>
    parm[grep("log.delta", Data$parm.names)] <- delta</pre>
    delta <- matrix(c(exp(delta), rep(1, Data$N)), Data$N, Data$M)</pre>
```

lambda <- interval(parm[grep("lambda", Data\$parm.names)], 1e-5, 1-1e-5)</pre>

sigma <- exp(parm[grep("log.sigma", Data\$parm.names)])</pre>

pi <- as.vector(Stick(lambda))</pre>

p <- delta / rowSums(delta)</pre>

```
theta <- max.col(p)
### Log(Hyperhyperprior Densities)
alpha.prior <- dhalfcauchy(alpha, 25, log=TRUE)</pre>
iota.prior <- dhalfcauchy(iota, 25, log=TRUE)
### Log(Hyperprior Densities)
gamma.prior <- dgamma(gamma, alpha, iota, log=TRUE)</pre>
nu.prior <- sum(dhalfcauchy(nu, 25, log=TRUE))</pre>
### Log(Prior Densities)
beta.prior <- sum(dnorm(beta, 0, matrix(rep(nu, Data$M), Data$M,
    Data$J, byrow=TRUE), log=TRUE))
delta.prior <- sum(dtrunc(delta, "norm", a=exp(-10), b=exp(10),
    mean=log(1/Data$M), sd=sqrt(1000), log=TRUE))
pi.prior <- dStick(pi, gamma, log=TRUE)</pre>
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
theta.prior <- sum(dcat(theta, pi, log=TRUE))</pre>
### Log-Likelihood
mu <- tcrossprod(Data$X, beta)</pre>
mu <- diag(mu[,theta])</pre>
LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))
### Log-Posterior
LP <- LL + beta.prior + delta.prior + pi.prior + nu.prior +</pre>
    sigma.prior + alpha.prior + iota.prior + gamma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,pi,sigma),</pre>
    yhat=rnorm(length(mu), mu, sigma), parm=parm)
return(Modelout)
}
```

52.4. Initial Values

```
Initial.Values <- c(runif(M*J), rep(0,J), runif(N*(M-1),-1,1), 0,
    rbeta(M-1,1,2), rep(0,3))</pre>
```

53. Mixture Model, Poisson-Gamma

$$\mathbf{y} \sim \mathcal{P}(\lambda)$$

$$\lambda \sim \mathcal{G}(\alpha \mu, \alpha)$$

$$\mu = \exp(\mathbf{X}\beta)$$

$$\alpha \sim \mathcal{HC}(25)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

53.2. Data

```
Model <- function(parm, Data)</pre>
    ### Hyperparameters
     alpha <- exp(parm[grep("log.alpha", Data$parm.names)])</pre>
    ### Parameters
    beta <- parm[grep("beta", Data$parm.names)]</pre>
     lambda <- exp(parm[grep("log.lambda", Data$parm.names)])</pre>
    mu <- exp(tcrossprod(Data$X, t(beta)))</pre>
    ### Log(Hyperprior Densities)
    alpha.prior <- dhalfcauchy(alpha, 25, log=TRUE)</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    lambda.prior <- sum(dgamma(lambda, alpha*mu, alpha, log=TRUE))</pre>
    ### Log-Likelihood
    LL <- sum(dpois(Data$y, lambda, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + lambda.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
         yhat=rpois(length(lambda), lambda), parm=parm)
    return(Modelout)
    }
```

53.4. Initial Values

```
Initial. Values \leftarrow c(0, rep(0,J), rep(0,N))
```

54. Multinomial Logit

54.1. Form

$$\mathbf{y}_{i} \sim \mathcal{CAT}(\mathbf{p}_{i,1:J}), \quad i = 1, \dots, N$$

$$\mathbf{p}_{i,j} = \frac{\phi_{i,j}}{\sum_{j=1}^{J} \phi_{i,j}}, \quad \sum_{j=1}^{J} \mathbf{p}_{i,j} = 1$$

$$\phi = \exp(\mu)$$

$$\mu_{i,J} = 0, \quad i = 1, \dots, N$$

$$\mu_{i,j} = \mathbf{X}_{i,1:K} \beta_{j,1:K} \in [-700, 700], \quad j = 1, \dots, (J-1)$$

$$\beta_{i,k} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1), \quad k = 1, \dots, K$$

54.2. Data

```
y <- x01 <- x02 <- c(1:300)
y[1:100] < -1
y[101:200] <- 2
y[201:300] <- 3
x01[1:100] <- rnorm(100, 25, 2.5)
x01[101:200] <- rnorm(100, 40, 4.0)
x01[201:300] \leftarrow rnorm(100, 35, 3.5)
x02[1:100] <- rnorm(100, 2.51, 0.25)
x02[101:200] \leftarrow rnorm(100, 2.01, 0.20)
x02[201:300] \leftarrow rnorm(100, 2.70, 0.27)
N <- length(y)
J <- 3 #Number of categories in y
K <- 3 #Number of predictors (including the intercept)</pre>
X \leftarrow matrix(c(rep(1,N),x01,x02),N,K)
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=matrix(0,J-1,K)))</pre>
PGF <- function(Data) return(rnormv((Data$J-1)*Data$K,0,1000))
MyData <- list(J=J, K=K, N=N, PGF=PGF, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
```

```
Model <- function(parm, Data)
    {
     ### Parameters
    beta <- matrix(parm, Data$J-1, Data$K)
     ### Log(Prior Densities)</pre>
```

Initial.Values <- c(rep(0,(J-1)*K))</pre>

55. Multinomial Logit, Nested

$$\mathbf{y}_{i} \sim \mathcal{CAT}(\mathbf{P}_{i,1:J}), \quad i = 1, \dots, N$$

$$\mathbf{P}_{1:N,1} = \frac{\mathbf{R}}{\mathbf{R} + \exp(\alpha \mathbf{I})}$$

$$\mathbf{P}_{1:N,2} = \frac{(1 - \mathbf{P}_{1:N,1})\mathbf{S}_{1:N,1}}{\mathbf{V}}$$

$$\mathbf{P}_{1:N,3} = \frac{(1 - \mathbf{P}_{1:N,1})\mathbf{S}_{1:N,2}}{\mathbf{V}}$$

$$\mathbf{R}_{1:N} = \exp(\mu_{1:N,1})$$

$$\mathbf{S}_{1:N,1:2} = \exp(\mu_{1:N,2:3})$$

$$\mathbf{I} = \log(\mathbf{V})$$

$$\mathbf{V}_{i} = \sum_{k=1}^{K} \mathbf{S}_{i,k}, \quad i = 1, \dots, N$$

$$\mu_{1:N,1} = \mathbf{X}\iota \in [-700, 700]$$

$$\mu_{1:N,2} = \mathbf{X}\beta_{2,1:K} \in [-700, 700]$$

$$\iota = \alpha\beta_{1,1:K}$$

$$\alpha \sim \mathcal{EXP}(1) \in [0, 2]$$

$$\beta_{i,k} \sim \mathcal{N}(0,1000), \quad j = 1, \dots, (J-1) \quad k = 1, \dots, K$$

where there are J=3 categories of \mathbf{y} , K=3 predictors, \mathbf{R} is the non-nested alternative, \mathbf{S} is the nested alternative, \mathbf{V} is the observed utility in the nest, α is effectively 1 - correlation and has a truncated exponential distribution, and ι is a vector of regression effects for the isolated alternative after α is taken into account. The third alternative is the reference category.

```
y < -x01 < -x02 < -c(1:300)
y[1:100] <- 1
y[101:200] <- 2
y[201:300] <- 3
x01[1:100] <- rnorm(100, 25, 2.5)
x01[101:200] <- rnorm(100, 40, 4.0)
x01[201:300] \leftarrow rnorm(100, 35, 3.5)
x02[1:100] \leftarrow rnorm(100, 2.51, 0.25)
x02[101:200] <- rnorm(100, 2.01, 0.20)
x02[201:300] \leftarrow rnorm(100, 2.70, 0.27)
N <- length(y)
J <- 3 #Number of categories in y
K <- 3 #Number of predictors (including the intercept)</pre>
X \leftarrow matrix(c(rep(1,N),x01,x02),N,K)
mon.names <- c("LP", as.parm.names(list(iota=rep(0,K))))</pre>
parm.names <- as.parm.names(list(alpha=0, beta=matrix(0,J-1,K)))</pre>
PGF <- function(Data) return(c(rtrunc(1, "exp", a=0, b=2, rate=1),
    rnormv((Data$J-1)*Data$K,0,1)))
MyData <- list(J=J, K=K, N=N, PGF=PGF, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
55.3. Model
Model <- function(parm, Data)</pre>
    ### Hyperparameters
     alpha.rate <- 1
     ### Parameters
    parm[1] <- alpha <- interval(parm[1],0,2)</pre>
     beta <- matrix(parm[grep("beta", Data$parm.names)], Data$J-1, Data$K)</pre>
     ### Log(Prior Densities)
     alpha.prior <- dtrunc(alpha, "exp", a=0, b=2, rate=alpha.rate,
         log=TRUE)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- P <- matrix(0, Data$N, Data$J)</pre>
     iota <- alpha * beta[1,]
    mu[,1] <- tcrossprod(Data$X, t(iota))</pre>
    mu[,2] <- tcrossprod(Data$X, t(beta[2,]))</pre>
```

Initial.Values <- c(0.5, rep(0.1, (J-1)*K))

56. Multinomial Probit

56.1. Form

$$\mathbf{W}_{i,1:(J-1)} \sim \mathcal{N}_{J-1}(\mu_{i,1:(J-1)}, \Sigma), \quad i = 1, \dots, N$$

$$\mathbf{W}_{i,j} \in \begin{cases} [0,10] & \text{if } \mathbf{y}_i = j \\ [-10,0] \end{cases}$$

$$\mu_{1:N,j} = \mathbf{X}\beta_{j,1:K}$$

$$\Sigma = \mathbf{U}^T \mathbf{U}$$

$$\beta_{j,k} \sim \mathcal{N}(0,10), \quad j = 1, \dots, (J-1), \quad k = 1, \dots, K$$

$$\mathbf{U}_{i,k} \sim \mathcal{N}(0,1), \quad j = 1, \dots, (J-1), \quad k = 1, \dots, (J-1), \quad j \ge k, \quad j \ne k = 1$$

```
N <- 50
J <- 5 #Categories of y
K <- 8 #Number of columns in design matrix X
X <- matrix(runif(N*K,-2,2), N, K)
X[,1] <- 1
beta <- matrix(runif((J-1)*K), J-1, K)
mu <- tcrossprod(X, beta)</pre>
```

```
S \leftarrow diag(J-1)
u \leftarrow c(0, rnorm((J-2) + (factorial(J-1) /
     (factorial(J-1-2)*factorial(2))),0,1))
U \leftarrow diag(J-1)
U[upper.tri(U, diag=TRUE)] <- u</pre>
diag(U) <- exp(diag(U))</pre>
Sigma <- t(U) %*% U
Sigma[1,] <- Sigma[,1] <- U[1,]
mu <- tcrossprod(X, beta)</pre>
W \leftarrow rmvn(N, mu, Sigma) + matrix(rnorm(N*(J-1),0,0.1), N, J-1)
y <- max.col(cbind(W,0))
table(y)
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=matrix(0,(J-1),K),</pre>
    U=U, W=matrix(0,N,J-1)), uppertri=c(0,1,0))
parm.names <- parm.names[-which(parm.names == "U[1,1]")]</pre>
PGF <- function(Data) {</pre>
    beta <- rnormv((Data$J-1)*Data$K,0,1)</pre>
    U <- rnorm((Data$J-2) + (factorial(Data$J-1) /</pre>
          (factorial(Data$J-1-2)*factorial(2))),0,1)
    W <- matrix(runif(Data$N*(Data$J-1),-10,0), Data$N, Data$J-1)
    Y <- as.indicator.matrix(Data$y)</pre>
    W <- ifelse(Y[,-Data$J] == 1, abs(W), W)
    return(c(beta, U, as.vector(W)))}
MyData <- list(J=J, K=K, N=N, PGF=PGF, S=S, X=X, mon.names=mon.names,
    parm.names=parm.names, y=y)
56.3. Model
Model <- function(parm, Data)</pre>
    {
    ### Parameters
    beta <- matrix(parm[grep("beta", Data$parm.names)], Data$J-1, Data$K)
    u <- c(0, parm[grep("U", Data$parm.names)])</pre>
    U <- diag(Data$J-1)</pre>
    U[upper.tri(U, diag=TRUE)] <- u</pre>
    diag(U) <- exp(diag(U))</pre>
    Sigma <- t(U) %*% U
    Sigma[1,] <- Sigma[,1] <- U[1,]
    W <- matrix(parm[grep("W", Data$parm.names)], Data$N, Data$J-1)
    Y <- as.indicator.matrix(Data$y)</pre>
    W \leftarrow ifelse(Y[,-c(Data$J)] == 1, interval(W, 0, 10),
         interval(W, -10, 0))
    parm[grep("W", Data$parm.names)] <- as.vector(W)</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 10, log=TRUE))</pre>
```

Initial.Values <- GIV(Model, MyData, PGF=TRUE)</pre>

57. Multivariate Binary Probit

57.1. Form

$$\mathbf{Z}_{i,1:J} \sim \mathcal{N}_{J}(\mu_{i,1:J}, \Sigma), \quad i = 1, \dots, N$$

$$\mathbf{Z}_{i,j} \in \begin{cases} [0,10] & \text{if } \mathbf{y}_{i} = j \\ [-10,0] & \text{} \end{cases}$$

$$\mu_{1:N,j} = \mathbf{X}\beta_{j,1:K}$$

$$\Sigma \sim \mathcal{IW}_{J+1}(\mathbf{S}^{-1}), \quad \mathbf{S} = \mathbf{I}_{J}, \quad \Sigma[1,1] = 1$$

$$\beta_{j,k} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, (J-1), \quad k = 1, \dots, K$$

$$\beta_{J,k} = -\sum_{j=1}^{J-1} \beta_{j,k}$$

$$\mathbf{Z}_{i,j} \sim \mathcal{N}(0, 1000) \in [-10, 10]$$

```
N <- 30
J <- 3 #Number of binary dependent variables
K <- 3 #Number of columns to be in design matrix X
Y <- matrix(round(runif(N*J)),N,J)
X <- matrix(1,N, K)
for (k in 2:K) {X[,k] <- rnorm(N, runif(1,-3,3), runif(1,0.1,3))}
S <- diag(J)</pre>
```

```
mon.names <- "LP"
sigma.temp <- as.parm.names(list(Sigma=diag(J)), uppertri=1)</pre>
parm.names <- c(sigma.temp[2:length(sigma.temp)],</pre>
     as.parm.names(list(beta=matrix(0,(J-1),K), Z=matrix(0,N,J))))
MyData <- list(J=J, K=K, N=N, S=S, X=X, Y=Y, mon.names=mon.names,
    parm.names=parm.names)
57.3. Model
Model <- function(parm, Data)</pre>
     {
    ### Parameters
    beta <- matrix(parm[grep("beta", Data$parm.names)], Data$J, Data$K)</pre>
    u <- c(0, parm[grep("U", Data$parm.names)])</pre>
    U <- diag(Data$J)</pre>
    U[upper.tri(U, diag=TRUE)] <- u</pre>
    diag(U) <- exp(diag(U))</pre>
    Sigma <- t(U) %*% U
    Sigma[1,] <- Sigma[,1] <- U[1,]
    W <- matrix(parm[grep("W", Data$parm.names)], Data$N, Data$J)</pre>
    W \leftarrow ifelse(Y == 1, interval(W, 0, 10),
         interval(W, -10, 0))
    parm[grep("W", Data$parm.names)] <- as.vector(W)</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    U.prior <- sum(dnorm(u[-length(u)], 0, 1, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, beta)</pre>
    LL <- sum(dmvn(W, mu, Sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + U.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
         yhat=1*(rmvn(prod(nrow(mu)), mu, Sigma) >= 0), parm=parm)
    return(Modelout)
    }
```

57.4. Initial Values

Initial.Values <- GIV(Model, MyData, PGF=TRUE)</pre>

58. Multivariate Laplace Regression

58.1. Form

$$\mathbf{Y}_{i,k} \sim \mathcal{L}_K(\mu_{i,k}, \Sigma), \quad i = 1, \dots, N; \quad k = 1, \dots, K$$

$$\mu_{i,k} = \mathbf{X}_{1:N,k} \beta_{k,1:J}$$

$$\Sigma = \Omega^{-1}$$

$$\Omega \sim \mathcal{W}_{K+1}(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_K$$

$$\beta_{k,j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

58.2. Data

```
N <- 100
J <- 6 #Number of columns in design matrix
K <- 3 #Number of DVs
X <- matrix(runif(N*J),N,J); X[,1] <- 1</pre>
Y <- mu <- tcrossprod(X, matrix(rnorm(J*K),K,J))
Sigma <- matrix(runif(K*K),K,K); diag(Sigma) <- runif(K,1,K)</pre>
Sigma <- as.symmetric.matrix(Sigma)</pre>
for (i in 1:N) {Y[i,] <- colMeans(rmvn(1000, mu[i,], Sigma))}</pre>
S \leftarrow diag(K)
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=matrix(0,K,J), U=diag(K)),</pre>
    uppertri=c(0,1)
PGF <- function(Data) return(c(rnormv(Data$K*Data$J,0,1),
    upper.triangle(rwishartc(Data$K+1,Data$S), diag=TRUE)))
MyData <- list(J=J, K=K, N=N, PGF=PGF, S=S, X=X, Y=Y, mon.names=mon.names,
    parm.names=parm.names)
```

```
Model <- function(parm, Data)
    {
    ### Parameters
    beta <- matrix(parm[grep("beta", Data$parm.names)], Data$K, Data$J)
    U <- as.parm.matrix(U, Data$K, parm, Data, chol=TRUE)
    diag(U) <- exp(diag(U))
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
    U.prior <- dwishart(U, Data$K+1, Data$S, log=TRUE)
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, beta)
    LL <- sum(dmvlc(Data$Y, mu, U, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + U.prior</pre>
```

58.4. Initial Values

Initial.Values <- c(rep(0,J*K), upper.triangle(S, diag=TRUE))</pre>

59. Multivariate Regression

59.1. Form

$$\mathbf{Y}_{i,k} \sim \mathcal{N}_K(\mu_{i,k}, \Omega^{-1}), \quad i = 1, \dots, N; \quad k = 1, \dots, K$$
$$\mu_{i,k} = \mathbf{X}_{1:N,k} \beta_{k,1:J}$$
$$\Omega \sim \mathcal{W}_{K+1}(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_K$$
$$\beta_{k,j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

59.2. Data

```
N <- 100
J <- 6 #Number of columns in design matrix
K <- 3 #Number of DVs
X <- matrix(runif(N*J),N,J); X[,1] <- 1</pre>
Y <- mu <- tcrossprod(X, matrix(rnorm(J*K),K,J))
Sigma <- matrix(runif(K*K),K,K); diag(Sigma) <- runif(K,1,K)</pre>
Sigma <- as.symmetric.matrix(Sigma)</pre>
for (i in 1:N) {Y[i,] <- colMeans(rmvn(1000, mu[i,], Sigma))}</pre>
S \leftarrow diag(K)
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=matrix(0,K,J), U=diag(K)),</pre>
    uppertri=c(0,1))
PGF <- function(Data) return(c(rnormv(Data$K*Data$J,0,1),</pre>
    upper.triangle(rwishartc(Data$K+1,Data$S), diag=TRUE)))
MyData <- list(J=J, K=K, N=N, PGF=PGF, S=S, X=X, Y=Y, mon.names=mon.names,
    parm.names=parm.names)
```

```
Model <- function(parm, Data)
{</pre>
```

```
### Parameters
beta <- matrix(parm[grep("beta", Data$parm.names)], Data$K, Data$J)</pre>
U <- as.parm.matrix(U, Data$K, parm, Data, chol=TRUE)</pre>
diag(U) <- exp(diag(U))</pre>
### Log(Prior Densities)
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
U.prior <- dwishartc(U, Data$K+1, Data$S, log=TRUE)</pre>
### Log-Likelihood
mu <- tcrossprod(Data$X, beta)</pre>
LL <- sum(dmvnpc(Data$Y, mu, U, log=TRUE))
### Log-Posterior
LP <- LL + beta.prior + U.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
     yhat=rmvnpc(nrow(mu), mu, U), parm=parm)
return(Modelout)
}
```

Initial.Values <- c(rep(0,J*K), upper.triangle(S, diag=TRUE))</pre>

60. Negative Binomial Regression

This example was contributed by Jim Robison-Cox.

60.1. Form

$$\mathbf{y} \sim \mathcal{NB}(\mu, \kappa)$$

$$p = \frac{\kappa}{\kappa + \mu}$$

$$\mu = \exp(\mathbf{X}\beta)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\kappa \sim \mathcal{HC}(25) \in (0, \infty]$$

```
N <- 100
J <- 5 #Number of predictors, including the intercept
kappa.orig <- 2
beta.orig <- runif(J,-2,2)
X <- matrix(runif(J*N,-2, 2), N, J); X[,1] <- 1
mu <- exp(tcrossprod(X, t(beta.orig)) + rnorm(N))
p <- kappa.orig / (kappa.orig + mu)</pre>
```

```
y <- rnbinom(N, size=kappa.orig, mu=mu)
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=rep(0,J), kappa=0))</pre>
PGF <- function(Data) return(c(rnormv(Data$J,0,1000),
    log(rhalfcauchy(1,25))))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names,</pre>
    parm.names=parm.names, y=y)
60.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:Data$J]</pre>
    parm[Data$J + 1] <- kappa <- interval(parm[Data$J + 1],</pre>
          .Machine$double.xmin, Inf)
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    kappa.prior <- dhalfcauchy(kappa, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- as.vector(exp(tcrossprod(Data$X, t(beta))))</pre>
    #p <- kappa / (kappa + mu)</pre>
    LL <- sum(dnbinom(Data$y, size=kappa, mu=mu, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + kappa.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
         yhat=rnbinom(length(mu), size=kappa, mu=mu), parm=parm)
    return(Modelout)
    }
```

60.4. Initial Values

```
Initial.Values <- c(rep(0,J), 1)</pre>
```

61. Normal, Multilevel

This is Gelman's school example (Gelman, Carlin, Stern, and Rubin 2004). Note that **LaplacesDemon** is slower to converge than WinBUGS through the **R2WinBUGS** package (Gelman 2011), an R package on CRAN. This example is very sensitive to the prior distributions. The recommended, default, half-Cauchy priors with scale 25 on scale parameters overwhelms the likelihood, so uniform priors are used.

61.1. Form

$$\mathbf{y}_{j} \sim \mathcal{N}(\theta_{j}, \sigma_{j}^{2}), \quad j = 1, \dots, J$$

$$\theta_{j} \sim \mathcal{N}(\theta_{\mu}, \theta_{\sigma}^{2})$$

$$\theta_{\mu} \sim \mathcal{N}(0, 1000000)$$

$$\theta_{\sigma[j]} \sim \mathcal{N}(0, 1000)$$

$$\sigma \sim \mathcal{U}(0, 1000)$$

61.2. Data

```
J <- 8
y <- c(28.4, 7.9, -2.8, 6.8, -0.6, 0.6, 18.0, 12.2)
sd <- c(14.9, 10.2, 16.3, 11.0, 9.4, 11.4, 10.4, 17.6)
mon.names <- "LP"
parm.names <- as.parm.names(list(theta=rep(0,J), theta.mu=0, theta.sigma=0))
PGF <- function(Data) return(c(rnorm(Data$J, rnormp(1,0,1E-6), runif(1,0,1000)), rnormp(1,0,1E-6), runif(1,0,1000)))
MyData <- list(J=J, PGF=PGF, mon.names=mon.names, parm.names=parm.names, sd=sd, y=y)</pre>
```

```
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    theta.mu <- parm[Data$J+1]</pre>
    theta.sigma <- interval(parm[Data$J+2], .Machine$double.eps, Inf)</pre>
    parm[Data$J+2] <- theta.sigma
    ### Parameters
    theta <- parm[1:Data$J]</pre>
    ### Log(Hyperprior Densities)
    theta.mu.prior <- dnormp(theta.mu, 0, 1.0E-6, log=TRUE)
    theta.sigma.prior <- dunif(theta.sigma, 0, 1000, log=TRUE)
    ### Log(Prior Densities)
    theta.prior <- sum(dnorm(theta, theta.mu, theta.sigma, log=TRUE))
    sigma.prior <- sum(dunif(Data$sd, 0, 1000, log=TRUE))</pre>
    ### Log-Likelihood
    LL <- sum(dnorm(Data$y, theta, Data$sd, log=TRUE))
    ### Log-Posterior
    LP <- LL + theta.prior + theta.mu.prior + theta.sigma.prior +
         sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
```

```
yhat=rnorm(length(theta), theta, Data$sd), parm=parm)
return(Modelout)
}
```

61.4. Initial Values

Initial.Values <- c(rep(mean(y), J), mean(y), 1)</pre>

62. Ordinal Logit

62.1. Form

$$\mathbf{y}_{i} \sim \mathcal{CAT}(P_{i,1:J})$$

$$P_{,J} = 1 - Q_{,(J-1)}$$

$$P_{,j} = |Q_{,j} - Q_{,(j-1)}|, \quad j = 2, \dots, (J-1)$$

$$P_{,1} = Q_{,1}$$

$$Q = \frac{1}{1 + \exp(\mu)}$$

$$\mu_{,j} = \delta_{j} - \mathbf{X}\beta, \quad \in [-5, 5]$$

$$\beta_{k} \sim \mathcal{N}(0, 1000), \quad k = 1, \dots, K$$

$$\delta_{j} \sim \mathcal{N}(0, 1) \in [(j-1), j] \in [-5, 5], \quad j = 1, \dots, (J-1)$$

```
data(demonsnacks)
N <- nrow(demonsnacks)
J <- 3 #Number of categories in y
X <- as.matrix(demonsnacks[,c(1,3:10)])
K <- ncol(demonsnacks) #Number of columns in design matrix X
y <- log(demonsnacks$Calories)
y <- ifelse(y < 4.5669, 1, ifelse(y > 5.5268, 3, 2)) #Discretize
for (k in 1:K) {X[,k] <- CenterScale(X[,k])}
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=rep(0,K), delta=rep(0,J-1)))
PGF <- function(Data)
    {
      delta <- rnorm(Data$J-1,0,1)
      delta <- delta[order(delta)]
      return(c(rnormv(Data$K,0,10), delta))
    }
MyData <- list(J=J, K=K, N=N, PGF=PGF, X=X, mon.names=mon.names,</pre>
```

```
parm.names=parm.names, y=y)
```

62.3. Model

```
Model <- function(parm, Data)</pre>
     ### Parameters
     beta <- parm[1:Data$K]</pre>
     delta <- interval(parm[-(1:Data$K)], -5, 5)</pre>
     delta <- delta[order(delta)]</pre>
     parm[-(1:Data$K)] <- delta</pre>
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
     delta.prior <- sum(dtrunc(delta, "norm", a=-5, b=5, log=TRUE,
         mean=0, sd=1)
     ### Log-Likelihood
     mu <- matrix(delta, Data$N, Data$J-1, byrow=TRUE) -</pre>
          matrix(tcrossprod(Data$X, t(beta)), Data$N, Data$J-1)
     P <- Q <- invlogit(mu)</pre>
     P[,-1] \leftarrow abs(Q[,-1] - Q[,-(Data$J-1)])
     P <- cbind(P, 1 - Q[,(Data$J-1)])</pre>
     LL <- sum(dcat(Data$y, P, log=TRUE))</pre>
     ### Log-Posterior
     LP <- LL + beta.prior + delta.prior
     Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP, yhat=rcat(nrow(P), P)</pre>
         parm=parm)
     return(Modelout)
```

62.4. Initial Values

Initial.Values <- c(rep(0,K), seq(from=-1, to=1, len=(J-1)))</pre>

63. Ordinal Probit

63.1. Form

$$\mathbf{y}_{i} \sim \mathcal{CAT}(P_{i,1:J})$$

$$P_{,J} = 1 - Q_{,(J-1)}$$

$$P_{,j} = |Q_{,j} - Q_{,(j-1)}|, \quad j = 2, \dots, (J-1)$$

$$P_{,1} = Q_{,1}$$

```
Q = \phi(\mu)
\mu_{,j} = \delta_j - \mathbf{X}\beta, \quad \in [-5, 5]
\beta_k \sim \mathcal{N}(0, 1000), \quad k = 1, \dots, K
\delta_j \sim \mathcal{N}(0, 1) \in [(j-1), j] \in [-5, 5], \quad j = 1, \dots, (J-1)
```

63.2. Data

```
data(demonsnacks)
N <- nrow(demonsnacks)</pre>
J <- 3 #Number of categories in y
X <- as.matrix(demonsnacks[,c(1,3:10)])</pre>
K <- ncol(demonsnacks) #Number of columns in design matrix X
y <- log(demonsnacks$Calories)</pre>
y \leftarrow ifelse(y < 4.5669, 1, ifelse(y > 5.5268, 3, 2)) #Discretize
for (k in 1:K) {X[,k] <- CenterScale(X[,k])}</pre>
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=rep(0,K), delta=rep(0,J-1)))</pre>
PGF <- function(Data)</pre>
    {
    delta <- rnorm(Data$J-1,0,1)</pre>
    delta <- delta[order(delta)]</pre>
    return(c(rnormv(Data$K,0,10), delta))
MyData <- list(J=J, K=K, N=N, PGF=PGF, X=X, mon.names=mon.names,
parm.names=parm.names, y=y)
63.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:Data$K]</pre>
    delta <- interval(parm[-(1:Data$K)], -5, 5)</pre>
    delta <- delta[order(delta)]</pre>
    parm[-(1:Data$K)] <- delta</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    delta.prior <- sum(dtrunc(delta, "norm", a=-5, b=5, log=TRUE,
         mean=0, sd=1)
    ### Log-Likelihood
    mu <- matrix(delta, Data$N, Data$J-1, byrow=TRUE) -</pre>
         matrix(tcrossprod(Data$X, t(beta)), Data$N, Data$J-1)
    P <- Q <- pnorm(mu)
```

 $P[,-1] \leftarrow abs(Q[,-1] - Q[,-(Data$J-1)])$

Initial.Values <- c(rep(0,K), seq(from=-1, to=1, len=(J-1)))</pre>

64. Panel, Autoregressive Poisson

64.1. Form

$$\mathbf{Y} \sim \mathcal{P}(\Lambda)$$

$$\Lambda_{1:N,1} = \exp(\alpha + \beta \mathbf{x})$$

$$\Lambda_{1:N,t} = \exp(\alpha + \beta \mathbf{x} + \rho \log(\mathbf{Y}_{1:N,t-1})), \quad t = 2, \dots, T$$

$$\alpha_i \sim \mathcal{N}(\alpha_{\mu}, \alpha_{\sigma}^2), \quad i = 1, \dots, N$$

$$\alpha_{\mu} \sim \mathcal{N}(0, 1000)$$

$$\alpha_{\sigma} \sim \mathcal{HC}(25)$$

$$\beta \sim \mathcal{N}(0, 1000)$$

$$\rho \sim \mathcal{N}(0, 1000)$$

```
N <- 10
T <- 10
alpha <- rnorm(N,2,0.5)
rho <- 0.5
beta <- 0.5
x <- runif(N,0,1)
Y <- matrix(NA,N,T)
Y[,1] <- exp(alpha + beta*x)
for (t in 2:T) {Y[,t] <- exp(alpha + beta*x + rho*log(Y[,t-1]))}
Y <- round(Y)
mon.names <- c("LP","alpha.sigma")
parm.names <- as.parm.names(list(alpha=rep(0,N), alpha.mu=0,</pre>
```

```
log.alpha.sigma=0, beta=0, rho=0))
PGF <- function(Data) return(c(rnorm(Data$N, rnormv(1,0,1000),
    rhalfcauchy(1,25)), rnormv(1,0,1000), log(rhalfcauchy(1,25)),
    rnormv(2,0,1000)))
MyData <- list(N=N, PGF=PGF, T=T, Y=Y, mon.names=mon.names,
    parm.names=parm.names, x=x)
64.3. Model
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    alpha.mu <- parm[Data$N+1]</pre>
    alpha.sigma <- exp(parm[Data$N+2])</pre>
    ### Parameters
    alpha <- parm[1:Data$N]</pre>
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    rho <- parm[grep("rho", Data$parm.names)]</pre>
    ### Log(Hyperprior Densities)
    alpha.mu.prior <- dnormv(alpha.mu, 0, 1000, log=TRUE)
    alpha.sigma.prior <- dhalfcauchy(alpha.sigma, 25, log=TRUE)</pre>
    ### Log(Prior Densities)
    alpha.prior <- sum(dnorm(alpha, alpha.mu, alpha.sigma, log=TRUE))</pre>
    beta.prior <- dnormv(beta, 0, 1000, log=TRUE)</pre>
    rho.prior <- dnormv(rho, 0, 1000, log=TRUE)</pre>
    ### Log-Likelihood
    Lambda <- Data$Y
    Lambda[,1] <- exp(alpha + beta*x)</pre>
    Lambda[,2:Data$T] <- exp(alpha + beta*Data$x +</pre>
         rho*log(Data$Y[,1:(Data$T-1)]))
    LL <- sum(dpois(Data$Y, Lambda, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + alpha.prior + alpha.mu.prior + alpha.sigma.prior +
         beta.prior + rho.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,alpha.sigma),</pre>
         yhat=rpois(prod(dim(Lambda)), parm=parm)
    return(Modelout)
    }
64.4. Initial Values
```

65. Poisson Regression

Initial. Values <- c(rep(0,N), 0, log(1), 0, 0)

65.1. Form

```
\mathbf{y} \sim \mathcal{P}(\lambda)
\lambda = \exp(\mathbf{X}\beta)
\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J
```

65.2. Data

65.3. Model

65.4. Initial Values

```
Initial.Values <- rep(0,J)</pre>
```

66. Polynomial Regression

In this univariate example, the degree of the polynomial is specified as D. For a more robust extension to estimating nonlinear relationships between \mathbf{y} and \mathbf{x} , see penalized spline regression in section $\ref{eq:condition}$?

66.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\mathbf{X}_{i,d} = \mathbf{x}_i^{d-1}, \quad d = 1, \dots, (D+1)$$

$$\mathbf{X}_{i,1} = 1$$

$$\beta_d \sim \mathcal{N}(0, 1000), \quad d = 1, \dots, (D+1)$$

$$\sigma \sim \mathcal{HC}(25)$$

66.2. Data

```
Model <- function(parm, Data)
   {
    ### Parameters
   beta <- parm[grep("beta", Data$parm.names)]
   sigma <- exp(parm[grep("log.sigma", Data$parm.names)])
   ### Log(Prior Densities)
   beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
   sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)
   ### Log-Likelihood
   X <- matrix(Data$x, Data$N, Data$D)
   for (d in 2:Data$D) {X[,d] <- X[,d]^d}
   X <- cbind(1,X)
   mu <- tcrossprod(X, t(beta))
   LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))</pre>
```

Initial.Values <- c(rep(0,D+1), log(1))</pre>

67. Proportional Hazards Regression, Weibull

Although the dependent variable is usually denoted as \mathbf{t} in survival analysis, it is denoted here as \mathbf{y} so Laplace's Demon recognizes it as a dependent variable for posterior predictive checks. This example does not support censoring, but it will be included soon.

67.1. Form

$$\mathbf{y}_i \sim \mathcal{WEIB}(\gamma, \mu_i), \quad i = 1, \dots, N$$

$$\mu = \exp(\mathbf{X}\beta)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\gamma \sim \mathcal{G}(1, 0.001)$$

67.2. Data

```
Model <- function(parm, Data)
{</pre>
```

67.4. Initial Values

```
Initial.Values <- c(rep(0,J), log(1))</pre>
```

68. Revision, Normal

This example provides both an analytic solution and numerical approximation of the revision of a normal distribution. Given a normal prior distribution (α) and data distribution (β) , the posterior (γ) is the revised normal distribution. This is an introductory example of Bayesian inference, and allows the user to experiment with numerical approximation, such as with MCMC in LaplacesDemon. Note that, regardless of the data sample size N in this example, Laplace Approximation is inappropriate due to asymptotics since the data (β) is perceived by the algorithm as a single datum rather than a collection of data. MCMC, on the other hand, is biased only by the effective number of samples taken of the posterior.

68.1. Form

$$\alpha \sim \mathcal{N}(0, 10)$$
$$\beta \sim \mathcal{N}(1, 2)$$
$$\gamma = \frac{\alpha_{\sigma}^{-2}\alpha + N\beta_{\sigma}^{-2}\beta}{\alpha_{\sigma}^{-2} + N\beta_{\sigma}^{-2}}$$

68.2. Data

```
N <- 10
mon.names <- c("LP","gamma")
parm.names <- c("alpha","beta")
PGF <- function(Data) return(c(rnorm(1,0,10), rnorm(1,1,2)))
MyData <- list(N=N, PGF=PGF, mon.names=mon.names, parm.names=parm.names)</pre>
```

68.3. Model

```
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    alpha.mu <- 0
    alpha.sigma <- 10
    beta.mu <- 1
    beta.sigma <- 2
    ### Parameters
    alpha <- parm[1]
    beta <- parm[2]
    ### Log(Prior Densities)
    alpha.prior <- dnorm(alpha, alpha.mu, alpha.sigma, log=TRUE)</pre>
    ### Log-Likelihood Density
    LL <- dnorm(beta, beta.mu, beta.sigma, log=TRUE)
    ### Posterior
    gamma <- (alpha.sigma^-2 * alpha + N * beta.sigma^-2 * beta) /</pre>
         (alpha.sigma^-2 + N * beta.sigma^-2)
    ### Log(Posterior Density)
    LP <- LL + alpha.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,gamma),</pre>
         yhat=rnorm(1, beta.mu, beta.sigma), parm=parm)
    return(Modelout)
    }
```

68.4. Initial Values

```
Initial.Values <- c(0,0)</pre>
```

69. Robust Regression

By replacing the normal distribution with the Student t distribution, linear regression is often called robust regression. As an alternative approach to robust regression, consider Laplace regression (see section 39).

69.1. Form

$$\mathbf{y} \sim t(\mu, \sigma^{2}, \nu)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

$$\nu \sim \mathcal{HC}(25)$$

69.2. Data

```
Model <- function(parm, Data)
    {
     ### Parameters
     beta <- parm[1:Data$J]
     sigma <- exp(parm[Data$J+1])
     nu <- exp(parm[Data$J+2])
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
     sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)
     nu.prior <- dhalfcauchy(nu, 25, log=TRUE)
     ### Log-Likelihood</pre>
```

Initial. Values $\leftarrow c(rep(0,J), log(1), log(5))$

70. Seemingly Unrelated Regression (SUR)

The following data was used by Zellner (1962) when introducing the Seemingly Unrelated Regression methodology. This model uses the conjugate Wishart distribution for precision in a multivariate normal distribution. See section 21 for a non-Wishart alternative that is more flexible and converges much faster.

70.1. Form

$$\mathbf{Y}_{t,k} \sim \mathcal{N}_{K}(\mu_{t,k}, \Omega^{-1}), \quad t = 1, \dots, T; \quad k = 1, \dots, K$$

$$\mu_{1,t} = \alpha_{1} + \alpha_{2} \mathbf{X}_{t-1,1} + \alpha_{3} \mathbf{X}_{t-1,2}, \quad t = 2, \dots, T$$

$$\mu_{2,t} = \beta_{1} + \beta_{2} \mathbf{X}_{t-1,3} + \beta_{3} \mathbf{X}_{t-1,4}, \quad t = 2, \dots, T$$

$$\Omega \sim \mathcal{W}_{K+1}(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_{K}$$

$$\alpha_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\beta_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

where J=3, K=2, and T=20.

```
IW \leftarrow c(12.93,25.90,35.05,22.89,18.84,28.57,48.51,43.34,37.02,37.81,
    39.27,53.46,55.56,49.56,32.04,32.24,54.38,71.78,90.08,68.60)
VW <- c(191.5,516.0,729.0,560.4,519.9,628.5,537.1,561.2,617.2,626.7,
    737.2,760.5,581.4,662.3,583.8,635.2,723.8,864.1,1193.5,1188.9)
CW \leftarrow c(1.8, 0.8, 7.4, 18.1, 23.5, 26.5, 36.2, 60.8, 84.4, 91.2, 92.4, 86.0, 111.1,
    130.6,141.8,136.7,129.7,145.5,174.8,213.5)
J <- 2 #Number of dependent variables
Y <- matrix(c(IG,IW), T, J)
S \leftarrow diag(J)
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=rep(0,3), beta=rep(0,3),
    U=diag(2), uppertri=c(0,0,1))
PGF <- function(Data) return(c(rnormv(3,0,1000), rnormv(3,0,1000),
    upper.triangle(rwishartc(Data$J+1,Data$S), diag=TRUE)))
MyData <- list(J=J, PGF=PGF, S=S, T=T, Y=Y, CG=CG, CW=CW, IG=IG, IW=IW,
    VG=VG, VW=VW, mon.names=mon.names, parm.names=parm.names)
```

```
Model <- function(parm, Data)</pre>
     {
    ### Parameters
     alpha <- parm[1:3]
    beta <- parm[4:6]
    U <- as.parm.matrix(U, Data$J, parm, Data, chol=TRUE)</pre>
    diag(U) <- exp(diag(U))</pre>
    ### Log(Prior Densities)
    alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    U.prior <- dwishartc(U, Data$J+1, Data$S, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- Data$Y
    mu[-1,1] \leftarrow alpha[1] + alpha[2]*Data$CG[-Data$T] +
          alpha[3]*Data$VG[-Data$T]
    mu[-1,2] \leftarrow beta[1] + beta[2]*Data$CW[-Data$T] +
         beta[3] *Data$VW[-Data$T]
    LL <- sum(dmvnpc(Data$Y[-1,], mu[-1,], U, log=TRUE))
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior + U.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
          yhat=rmvnpc(nrow(mu), mu, U), parm=parm)
    return(Modelout)
    }
```

Initial.Values <- c(rep(0,3), rep(0,3), upper.triangle(S, diag=TRUE))</pre>

71. Simultaneous Equations

This example of simultaneous equations uses Klein's Model I (Kleine 1950) regarding economic fluctations in the United States in 1920-1941 (\mathbf{N} =22). Usually, this example is modeled with 3-stage least squares (3SLS), excluding the uncertainty from multiple stages. By constraining each element in the instrumental variables matrix $\nu \in [-10, 10]$, this example estimates the model without resorting to stages. The dependent variable is matrix \mathbf{Y} , in which $\mathbf{Y}_{1,1:N}$ is \mathbf{C} or Consumption, $\mathbf{Y}_{2,1:N}$ is \mathbf{I} or Investment, and $\mathbf{Y}_{3,1:N}$ is $\mathbf{W}\mathbf{p}$ or Private Wages. Here is a data dictionary:

A = Time Trend measured as years from 1931

C = Consumption

G = Government Nonwage Spending

I = Investment

K = Capital Stock

P = Private (Corporate) Profits

T = Indirect Business Taxes Plus Neg Exports

Wg = Government Wage Bill

Wp = Private Wages

X = Equilibrium Demand (GNP)

See Kleine (1950) for more information.

71.1. Form

$$\mathbf{Y} \sim \mathcal{N}_{3}(\mu, \Omega^{-1})$$

$$\mu_{1,1} = \alpha_{1} + \alpha_{2}\nu_{1,1} + \alpha_{4}\nu_{2,1}$$

$$\mu_{1,i} = \alpha_{1} + \alpha_{2}\nu_{1,i} + \alpha_{3}\mathbf{P}_{i-1} + \alpha_{4}\nu_{2,i}, \quad i = 2, ..., N$$

$$\mu_{2,1} = \beta_{1} + \beta_{2}\nu_{1,1} + \beta_{4}\mathbf{K}_{1}$$

$$\mu_{2,i} = \beta_{1} + \beta_{2}\nu_{1,i} + \beta_{3}\mathbf{P}_{i-1} + \beta_{4}\mathbf{K}_{i}, \quad i = 2, ..., N$$

$$\mu_{3,1} = \gamma_{1} + \gamma_{2}\nu_{3,1} + \gamma_{4}\mathbf{A}_{1}$$

$$\mu_{3,i} = \gamma_{1} + \gamma_{2}\nu_{3,i} + \gamma_{3}\mathbf{X}_{i-1} + \gamma_{4}\mathbf{A}_{i}, \quad i = 2, ..., N$$

$$\mathbf{Z}_{j,i} \sim \mathcal{N}(\nu_{j,i}, \sigma_{j}^{2}), \quad j = 1, ..., 3$$

$$\nu_{j,1} = \pi_{j,1} + \pi_{j,3}\mathbf{K}_{1} + \pi_{j,5}\mathbf{A}_{1} + \pi_{j,6}\mathbf{T}_{1} + \pi_{j,7}\mathbf{G}_{1}, \quad j = 1, ..., 3$$

$$\nu_{j,i} = \pi_{j,1} + \pi_{j,2}\mathbf{P}_{i-1} + \pi_{j,3}\mathbf{K}_{i} + \pi_{j,4}\mathbf{X}_{i-1} + \pi_{j,5}\mathbf{A}_{i} + \pi_{j,6}\mathbf{T}_{i} + \pi\mathbf{G}_{i}, \quad i = 1, ..., N, \quad j = 1, ..., 3$$

$$\alpha_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, ..., 4$$

$$\beta_{j} \sim \mathcal{N}(0, 1000), \quad j = 1, ..., 4$$

$$\gamma_{i} \sim \mathcal{N}(0, 1000), \quad j = 1, ..., 4$$

```
\pi_{j,i} \sim \mathcal{N}(0, 1000) \in [-10, 10], \quad j = 1, \dots, 3, \quad i = 1, \dots, N
\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, 3
\Omega \sim \mathcal{W}_4(\mathbf{S}), \quad \mathbf{S} = \mathbf{I}_3
```

71.2. Data

Model <- function(parm, Data)</pre>

{

```
N <- 22
A \leftarrow c(-11, -10, -9, -8, -7, -6, -5, -4, -3, -2, -1, 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10)
C \leftarrow c(39.8,41.9,45,49.2,50.6,52.6,55.1,56.2,57.3,57.8,55,50.9,45.6,46.5,
     48.7,51.3,57.7,58.7,57.5,61.6,65,69.7)
G \leftarrow c(2.4,3.9,3.2,2.8,3.5,3.3,3.3,4,4.2,4.1,5.2,5.9,4.9,3.7,4,4.4,2.9,4.3,
     5.3,6.6,7.4,13.8)
I \leftarrow c(2.7, -0.2, 1.9, 5.2, 3, 5.1, 5.6, 4.2, 3, 5.1, 1, -3.4, -6.2, -5.1, -3, -1.3, 2.1, 2,
     -1.9, 1.3, 3.3, 4.9)
K \leftarrow c(180.1,182.8,182.6,184.5,189.7,192.7,197.8,203.4,207.6,210.6,215.7,
     216.7,213.3,207.1,202,199,197.7,199.8,201.8,199.9,201.2,204.5)
P \leftarrow c(12.7, 12.4, 16.9, 18.4, 19.4, 20.1, 19.6, 19.8, 21.1, 21.7, 15.6, 11.4, 7, 11.2,
     12.3,14,17.6,17.3,15.3,19,21.1,23.5)
T \leftarrow c(3.4,7.7,3.9,4.7,3.8,5.5,7,6.7,4.2,4,7.7,7.5,8.3,5.4,6.8,7.2,8.3,6.7,
    7.4,8.9,9.6,11.6)
Wg \leftarrow c(2.2,2.7,2.9,2.9,3.1,3.2,3.3,3.6,3.7,4,4.2,4.8,5.3,5.6,6,6.1,7.4,
     6.7, 7.7, 7.8, 8, 8.5
Wp \leftarrow c(28.8, 25.5, 29.3, 34.1, 33.9, 35.4, 37.4, 37.9, 39.2, 41.3, 37.9, 34.5, 29, 28.5,
     30.6,33.2,36.8,41,38.2,41.6,45,53.3)
X \leftarrow c(44.9,45.6,50.1,57.2,57.1,61,64,64.4,64.5,67,61.2,53.4,44.3,45.1,
    49.7,54.4,62.7,65,60.9,69.5,75.7,88.4)
year <- c(1920,1921,1922,1923,1924,1925,1926,1927,1928,1929,1930,1931,1932,</pre>
     1933, 1934, 1935, 1936, 1937, 1938, 1939, 1940, 1941)
Y <- matrix(c(C,I,Wp),3,N, byrow=TRUE)
Z <- matrix(c(P, Wp+Wg, X), 3, N, byrow=TRUE)</pre>
S <- diag(nrow(Y))</pre>
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=rep(0,4), beta=rep(0,4),
     gamma=rep(0,4), pi=matrix(0,3,7), log.sigma=rep(0,3),
     Omega=diag(3)), uppertri=c(0,0,0,0,0,1))
PGF <- function(Data) return(c(rnormv(4,0,10), rnormv(4,0,10),
    rnormv(4,0,10), rnormv(3*7,0,10), log(rhalfcauchy(3,25)),
     upper.triangle(rwishartc(nrow(Data$S)+1,Data$S), diag=TRUE)))
MyData <- list(A=A, C=C, G=G, I=I, K=K, N=N, P=P, PGF=PGF, S=S, T=T, Wg=Wg,
     Wp=Wp, X=X, Y=Y, Z=Z, mon.names=mon.names, parm.names=parm.names)
71.3. Model
```

```
### Parameters
alpha <- parm[1:4]; beta <- parm[5:8]; gamma <- parm[9:12]
pi <- matrix(interval(parm[grep("pi", Data$parm.names)],-10,10), 3, 7)</pre>
parm[grep("pi", Data$parm.names)] <- as.vector(pi)</pre>
sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
Omega <- as.parm.matrix(Omega, nrow(Data$S), parm, Data)</pre>
parm[grep("Omega", Data$parm.names)] <- upper.triangle(Omega,</pre>
    diag=TRUE)
### Log(Prior Densities)
alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
gamma.prior <- sum(dnormv(gamma, 0, 1000, log=TRUE))</pre>
pi.prior <- sum(dnormv(pi, 0, 1000, log=TRUE))</pre>
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
Omega.prior <- dwishart(Omega, nrow(Data$S)+1, Data$S, log=TRUE)</pre>
### Log-Likelihood
mu <- nu <- matrix(0,3,Data$N)</pre>
for (i in 1:3) {
    nu[i,1] <- pi[i,1] + pi[i,3]*Data$K[1] + pi[i,5]*Data$A[1] +</pre>
         pi[i,6]*Data$T[1] + pi[i,7]*Data$G[1]
    nu[i,-1] <- pi[i,1] + pi[i,2]*Data$P[-Data$N] +</pre>
         pi[i,3]*Data$K[-1] + pi[i,4]*Data$X[-Data$N] +
         pi[i,5]*Data$A[-1] + pi[i,6]*Data$T[-1] +
         pi[i,7]*Data$G[-1]}
LL <- sum(dnorm(Data$Z, nu, matrix(sigma, 3, Data$N), log=TRUE))
mu[1,1] \leftarrow alpha[1] + alpha[2]*nu[1,1] + alpha[4]*nu[2,1]
mu[1,-1] \leftarrow alpha[1] + alpha[2]*nu[1,-1] +
    alpha[3]*Data$P[-Data$N] + alpha[4]*nu[2,-1]
mu[2,1] \leftarrow beta[1] + beta[2]*nu[1,1] + beta[4]*Data$K[1]
mu[2,-1] \leftarrow beta[1] + beta[2]*nu[1,-1] +
    beta[3]*Data$P[-Data$N] + beta[4]*Data$K[-1]
mu[3,1] <- gamma[1] + gamma[2]*nu[3,1] + gamma[4]*Data$A[1]</pre>
mu[3,-1] <- gamma[1] + gamma[2]*nu[3,-1] +
    gamma[3]*Data$X[-Data$N] + gamma[4]*Data$A[-1]
LL <- LL + sum(dmvnp(t(Data$Y), t(mu), Omega, log=TRUE))</pre>
### Log-Posterior
LP <- LL + alpha.prior + beta.prior + gamma.prior + pi.prior +
    sigma.prior + Omega.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
    yhat=rmvnp(ncol(mu), t(mu), Omega), parm=parm)
return(Modelout)
}
```

```
Initial. Values \leftarrow c(rep(0,4), rep(0,4), rep(0,4), rep(0,3*7), rep(0,3),
```

upper.triangle(S, diag=TRUE))

72. Space-Time, Dynamic

This approach to space-time or spatiotemporal modeling applies kriging to a stationary spatial component for points in space $s=1,\ldots,S$ first at time t=1, where space is continuous and time is discrete. Vector ζ contains these spatial effects. Next, SSM (State Space Model) or DLM (Dynamic Linear Model) components are applied to the spatial parameters $(\phi, \kappa, \text{ and } \lambda)$ and regression effects (β) . These parameters are allowed to vary dynamically with time $t=2,\ldots,T$, and the resulting spatial process is estimated for each of these time-periods. When time is discrete, a dynamic space-time process can be applied. The matrix Θ contains the dynamically varying stationary spatial effects, or space-time effects. Spatial coordinates are given in longitude and latitude for $s=1,\ldots,S$ points in space and measurements are taken across discrete time-periods $t=1,\ldots,T$ for $\mathbf{Y}_{s,t}$. The dependent variable is also a function of design matrix \mathbf{X} (which may also be dynamic, but is static in this example) and dynamic regression effects matrix $\beta_{1:J,1:T}$. For more information on kriging, see section 37. For more information on SSMs or DLMs, see section 77. To extend this to a large spatial data set, consider incorporating the predictive process kriging example in section 38.

72.1. Form

$$\mathbf{Y}_{s,t} \sim \mathcal{N}(\mu_{s,t}, \sigma_{1}^{2}), \quad s = 1, \dots, S, \quad t = 1, \dots, T$$

$$\mu_{s,t} = \mathbf{X}_{s,1:J}\beta_{1:J,t} + \Theta_{s,t}$$

$$\Theta_{s,t} = \frac{\sum_{s,s,t}}{\sum_{r=1}^{S} \sum_{r,s,t}} \Theta_{s,t-1}, \quad s = 1, \dots, S, \quad t = 2, \dots, T$$

$$\Theta_{s,1} = \zeta_{s}$$

$$\zeta \sim \mathcal{N}_{S}(0, \Sigma_{1:S,1:S,1})$$

$$\Sigma_{1:S,1:S,t} = \lambda_{t}^{2} \exp(-\phi_{t}\mathbf{D})^{\kappa[t]}$$

$$\sigma_{1} \sim \mathcal{HC}(25)$$

$$\beta_{j,1} \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, 2$$

$$\beta_{1,t} \sim \mathcal{N}(\beta_{1,t-1}, \sigma_{2}^{2}), \quad t = 2, \dots, T$$

$$\beta_{2,t} \sim \mathcal{N}(\beta_{2,t-1}, \sigma_{3}^{2}), \quad t = 2, \dots, T$$

$$\phi_{1} \sim \mathcal{H}\mathcal{N}(1000)$$

$$\phi_{t} \sim \mathcal{N}(\phi_{t-1}, \sigma_{4}^{2}) \in [0, \infty], \quad t = 2, \dots, T$$

$$\kappa_{1} \sim \mathcal{H}\mathcal{N}(1000)$$

$$\kappa_{t} \sim \mathcal{N}(\kappa_{t-1}, \sigma_{5}^{2}) \in [0, \infty], \quad t = 2, \dots, T$$

$$\lambda_{1} \sim \mathcal{H}\mathcal{N}(1000)$$

$$\lambda_{t} \sim \mathcal{N}(\lambda_{t-1}, \sigma_{6}^{2}) \in [0, \infty], \quad t = 2, \dots, T$$

```
S <- 20
T <- 10
longitude <- runif(S,0,100)</pre>
latitude <- runif(S,0,100)</pre>
D <- as.matrix(dist(cbind(longitude, latitude), diag=TRUE, upper=TRUE))
beta \leftarrow matrix(c(50,2), 2, T)
phi <- rep(1,T); kappa <- rep(1.5,T); lambda <- rep(10000,T)
for (t in 2:T) {
    beta[1,t-1] \leftarrow beta[1,t-1] + rnorm(1,0,1)
    beta[2,t-1] \leftarrow beta[2,t-1] + rnorm(1,0,0.1)
    phi[t] \leftarrow phi[t-1] + rnorm(1,0,0.1)
    if(phi[t] < 0.001) phi[t] <- 0.001
    kappa[t] \leftarrow kappa[t-1] + rnorm(1,0,0.1)
    lambda[t] \leftarrow lambda[t-1] + rnorm(1,0,1000)
Sigma <- array(0, dim=c(S,S,T))</pre>
for (t in 1:T) {
    Sigma[ , ,t] <- lambda[t] * exp(-phi[t] * D)^kappa[t]}</pre>
zeta <- as.vector(apply(rmvn(1000, rep(0,S), Sigma[ , ,1]), 2, mean))</pre>
Theta <- matrix(zeta,S,T)</pre>
for (t in 2:T) {for (s in 1:S) {
     Theta[,t] \leftarrow sum(Sigma[,s,t] / sum(Sigma[,s,t]) * Theta[,t-1]) \} 
X \leftarrow matrix(runif(S*2,-2,2),S,2); X[,1] \leftarrow 1
mu <- tcrossprod(X, t(beta))</pre>
Y \leftarrow mu + Theta + matrix(rnorm(S*T,0,0.1),S,T)
mon.names <- c("LP", as.parm.names(list(sigma=rep(0,6))))</pre>
parm.names <- as.parm.names(list(zeta=rep(0,S), beta=matrix(0,2,T),</pre>
    log.phi=rep(0,T), log.kappa=rep(0,T), log.lambda=rep(0,T),
    log.sigma=rep(0,6))
PGF <- function(Data) return(c(rmvn(1, rep(0,Data$S),
    rhalfnorm(1,sqrt(1000))^2 *
     exp(-rhalfnorm(1,sqrt(1000))*Data$D)^rhalfnorm(1,sqrt(1000))),
    rnormv(2*Data$T,0,1000), log(rhalfnorm(Data$T,sqrt(1000))),
    log(rhalfnorm(Data$T,sqrt(1000))), log(rhalfnorm(Data$T,sqrt(1000))),
     log(rhalfcauchy(6,25)))
MyData <- list(D=D, PGF=PGF, S=S, T=T, X=X, Y=Y, latitude=latitude,
     longitude=longitude, mon.names=mon.names, parm.names=parm.names)
72.3. Model
Model <- function(parm, Data)</pre>
     ### Parameters
    beta <- matrix(parm[grep("beta", Data$parm.names)], 2, Data$T)</pre>
    zeta <- parm[grep("zeta", Data$parm.names)]</pre>
    phi <- exp(parm[grep("log.phi", Data$parm.names)])</pre>
```

```
kappa <- exp(parm[grep("log.kappa", Data$parm.names)])</pre>
lambda <- exp(parm[grep("log.lambda", Data$parm.names)])</pre>
sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
Sigma <- array(0, dim=c(Data$S, Data$T))</pre>
for (t in 1:Data$T) {
    Sigma[ , ,t] <- lambda[t]^2 * exp(-phi[t] * Data$D)^kappa[t]}</pre>
### Log(Prior Densities)
beta.prior <- sum(dnormv(beta[,1], 0, 1000, log=TRUE),</pre>
    dnorm(beta[,-1], beta[,-Data$T], matrix(sigma[2:3], 2,
    Data$T-1), log=TRUE))
zeta.prior <- dmvn(zeta, rep(0,Data$S), Sigma[ , , 1], log=TRUE)</pre>
phi.prior <- sum(dhalfnorm(phi[1], sqrt(1000), log=TRUE),</pre>
    dtrunc(phi[-1], "norm", a=0, b=Inf, mean=phi[-Data$T],
    sd=sigma[4], log=TRUE))
kappa.prior <- sum(dhalfnorm(kappa[1], sqrt(1000), log=TRUE),</pre>
    dtrunc(kappa[-1], "norm", a=0, b=Inf, mean=kappa[-Data$T],
    sd=sigma[5], log=TRUE))
lambda.prior <- sum(dhalfnorm(lambda[1], sqrt(1000), log=TRUE),</pre>
    dtrunc(lambda[-1], "norm", a=0, b=Inf, mean=lambda[-Data$T],
    sd=sigma[6], log=TRUE))
sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
### Log-Likelihood
mu <- tcrossprod(Data$X, t(beta))</pre>
Theta <- matrix(zeta, Data$S, Data$T)</pre>
for (t in 2:Data$T) {
    for (s in 1:Data$S) {
         Theta[,t] \leftarrow Sigma[,s,t] / sum(Sigma[,s,t]) * Theta[,t-1]\}
mu <- mu + Theta
LL <- sum(dnorm(Data$Y, mu, sigma[1], log=TRUE))
### Log-Posterior
LP <- LL + beta.prior + zeta.prior + sum(phi.prior) +</pre>
    sum(kappa.prior) + sum(lambda.prior) + sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma),</pre>
    yhat=rnorm(prod(dim(mu)), mu, sigma[1]), parm=parm)
return(Modelout)
}
```

72.4. Initial Values

```
Initial.Values <- c(rep(0,S), rep(c(mean(Y),0),T), log(rep(1,T)), log(rep(1,T)), rep(1,T), log(rep(1,6)))
```

73. Space-Time, Nonseparable

This approach to space-time or spatiotemporal modeling applies kriging both to the stationary

spatial and temporal components, where space is continuous and time is discrete. Matrix Ξ contains the space-time effects. Spatial coordinates are given in longitude and latitude for s = 1, ..., S points in space and measurements are taken across time-periods t = 1, ..., T for $\mathbf{Y}_{s,t}$. The dependent variable is also a function of design matrix \mathbf{X} and regression effects vector β . For more information on kriging, see section 37. This example uses a nonseparable, stationary covariance function in which space and time are separable only when $\psi = 0$. To extend this to a large space-time data set, consider incorporating the predictive process kriging example in section 38.

73.1. Form

$$\mathbf{Y}_{s,t} \sim \mathcal{N}(\mu_{s,t}, \sigma_1^2), \quad s = 1, \dots, S, \quad t = 1, \dots, T$$

$$\mu = \mathbf{X}\beta + \Xi$$

$$\Xi \sim \mathcal{N}_{ST}(\Xi_{\mu}, \Sigma)$$

$$\Sigma = \sigma_2^2 \exp\left(-\frac{\mathbf{D}_S}{\phi_1}^{\kappa} - \frac{\mathbf{D}_T}{\phi_2}^{\lambda} - \psi \frac{\mathbf{D}_S}{\phi_1}^{\kappa} \frac{\mathbf{D}_T}{\phi_2}^{\lambda}\right)$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\phi_k \sim \mathcal{U}(1, 5), \quad k = 1, \dots, 2$$

$$\sigma_k \sim \mathcal{HC}(25), \quad k = 1, \dots, 2$$

$$\psi \sim \mathcal{HC}(25)$$

$$\Xi_{\mu} = 0$$

$$\kappa = 1, \quad \lambda = 1$$

```
S <- 10
T <- 5
longitude <- runif(S,0,100)</pre>
latitude <- runif(S,0,100)</pre>
D.S <- as.matrix(dist(cbind(rep(longitude,T),rep(latitude,T)), diag=TRUE,
     upper=TRUE))
D.T <- as.matrix(dist(cbind(rep(1:T,each=S),rep(1:T,each=S)), diag=TRUE,
     upper=TRUE))
Sigma \leftarrow 10000 * exp(-D.S/3 - D.T/2 - 0.2*(D.S/3)*(D.T/2))
Xi <- as.vector(apply(rmvn(1000, rep(0,S*T), Sigma), 2, mean))</pre>
Xi <- matrix(Xi,S,T)</pre>
beta <- c(50,2)
X \leftarrow matrix(runif(S*2,-2,2),S,2); X[,1] \leftarrow 1
mu <- as.vector(tcrossprod(X, t(beta)))</pre>
Y \leftarrow mu + Xi
mon.names <- c("LP", "psi", "sigma[1]", "sigma[2]")</pre>
```

```
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    Xi.mu <- rep(0,Data$S*Data$T)</pre>
     ### Parameters
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    Xi <- parm[grep("Xi", Data$parm.names)]</pre>
    kappa <- 1; lambda <- 1
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    phi <- interval(parm[grep("phi", Data$parm.names)], 1, 5)</pre>
    parm[grep("phi", Data$parm.names)] <- phi</pre>
    psi <- exp(parm[grep("log.psi", Data$parm.names)])</pre>
    Sigma <- sigma[2]*sigma[2] * exp(-(Data$D.S / phi[1])^kappa -
         (Data$D.T / phi[2])^lambda -
         psi*(Data$D.S / phi[1])^kappa * (Data$D.T / phi[2])^lambda)
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    Xi.prior <- dmvn(Xi, Xi.mu, Sigma, log=TRUE)</pre>
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    phi.prior <- sum(dunif(phi, 1, 5, log=TRUE))</pre>
    psi.prior <- dhalfcauchy(psi, 25, log=TRUE)</pre>
    ### Log-Likelihood
    Xi <- matrix(Xi, Data$S, Data$T)</pre>
    mu <- as.vector(tcrossprod(Data$X, t(beta))) + Xi</pre>
    LL <- sum(dnorm(Data$Y, mu, sigma[1], log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + beta.prior + Xi.prior + sigma.prior + phi.prior + psi.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,psi,sigma),</pre>
         yhat=rnorm(prod(dim(mu)), mu, sigma[1]), parm=parm)
    return(Modelout)
    }
```

Initial. Values $\leftarrow c(rep(0,S*T), mean(Y), 0, rep(1,2), rep(0,2), 0)$

74. Space-Time, Separable

This introductory approach to space-time or spatiotemporal modeling applies kriging both to the stationary spatial and temporal components, where space is continuous and time is discrete. Vector ζ contains the spatial effects and vector θ contains the temporal effects. Spatial coordinates are given in longitude and latitude for s = 1, ..., S points in space and measurements are taken across time-periods t = 1, ..., T for $\mathbf{Y}_{s,t}$. The dependent variable is also a function of design matrix \mathbf{X} and regression effects vector β . For more information on kriging, see section 37. This example uses separable space-time covariances, which is more convenient but usually less appropriate than a nonseparable covariance function. To extend this to a large space-time data set, consider incorporating the predictive process kriging example in section 38.

74.1. Form

$$\mathbf{Y}_{s,t} \sim \mathcal{N}(\mu_{s,t}, \sigma_1^2), \quad s = 1, \dots, S, \quad t = 1, \dots, T$$

$$\mu_{s,t} = \mathbf{X}_{s,1:J}\beta + \zeta_s + \Theta_{s,t}$$

$$\Theta_{s,1:T} = \theta$$

$$\theta \sim \mathcal{N}_N(\theta_\mu, \Sigma_T)$$

$$\Sigma_T = \sigma_3^2 \exp(-\phi_2 \mathbf{D}_T)^\lambda$$

$$\zeta \sim \mathcal{N}_N(\zeta_\mu, \Sigma_S)$$

$$\Sigma_S = \sigma_2^2 \exp(-\phi_1 \mathbf{D}_S)^\kappa$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, 2$$

$$\sigma_k \sim \mathcal{HC}(25), \quad k = 1, \dots, 3$$

$$\phi_k \sim \mathcal{U}(1, 5), \quad k = 1, \dots, 2$$

$$\zeta_\mu = 0$$

$$\theta_\mu = 0$$

$$\kappa = 1, \quad \lambda = 1$$

```
S <- 20
T <- 10
longitude <- runif(S,0,100)
latitude <- runif(S,0,100)
D.S <- as.matrix(dist(cbind(longitude,latitude), diag=TRUE, upper=TRUE))</pre>
```

```
Sigma.S \leftarrow 10000 * exp(-1.5 * D.S)
zeta <- as.vector(apply(rmvn(1000, rep(0,S), Sigma.S), 2, mean))</pre>
D.T <- as.matrix(dist(cbind(c(1:T),c(1:T)), diag=TRUE, upper=TRUE))
Sigma.T <- 10000 * exp(-3 * D.T)
theta <- as.vector(apply(rmvn(1000, rep(0,T), Sigma.T), 2, mean))
Theta <- matrix(theta,S,T,byrow=TRUE)</pre>
beta <- c(50,2)
X \leftarrow matrix(runif(S*2,-2,2),S,2); X[,1] \leftarrow 1
mu <- as.vector(tcrossprod(X, t(beta)))</pre>
Y <- mu + zeta + Theta + matrix(rnorm(S*T,0,0.1),S,T)
mon.names <- c("LP", "sigma[1]", "sigma[2]", "sigma[3]")
parm.names <- as.parm.names(list(zeta=rep(0,S), theta=rep(0,T),</pre>
    beta=rep(0,2), phi=rep(0,2), log.sigma=rep(0,3)))
PGF <- function(Data) return(c(rmvn(1, rep(0,Data$S),
    rhalfcauchy(1,25)^2 * \exp(-\text{runif}(1,1,5)*\text{Data$D.S})),
     rmvn(1, rep(0,Data$T), rhalfcauchy(1,25)^2 *
     \exp(-\text{runif}(1,1,5)*\text{Data}D.T)), \operatorname{rnormv}(2,0,1000), \operatorname{runif}(2,1,5),
     log(rhalfcauchy(3,25))))
MyData <- list(D.S=D.S, D.T=D.T, PGF=PGF, S=S, T=T, X=X, Y=Y,
     latitude=latitude, longitude=longitude, mon.names=mon.names,
    parm.names=parm.names)
74.3. Model
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    zeta.mu <- rep(0,Data$S)</pre>
    theta.mu <- rep(0,Data$T)
    ### Parameters
    beta <- parm[grep("beta", Data$parm.names)]</pre>
    zeta <- parm[grep("zeta", Data$parm.names)]</pre>
    theta <- parm[grep("theta", Data$parm.names)]</pre>
    kappa <- 1; lambda <- 1
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    phi <- interval(parm[grep("phi", Data$parm.names)], 1, 5)</pre>
    parm[grep("phi", Data$parm.names)] <- phi</pre>
    Sigma.S <- sigma[2]^2 * exp(-phi[1] * Data$D.S)^kappa
    Sigma.T <- sigma[3]^2 * exp(-phi[2] * Data$D.T)^lambda</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    zeta.prior <- dmvn(zeta, zeta.mu, Sigma.S, log=TRUE)</pre>
    theta.prior <- dmvn(theta, theta.mu, Sigma.T, log=TRUE)
     sigma.prior <- sum(dhalfcauchy(25, log=TRUE))</pre>
    phi.prior <- sum(dunif(phi, 1, 5, log=TRUE))</pre>
    ### Log-Likelihood
```

Initial. Values <-c(rep(0,S), rep(0,T), rep(0,2), rep(1,2), rep(0,3))

75. Spatial Autoregression (SAR)

The spatial autoregressive (SAR) model in this example uses are aldata that consists of first-order neighbors that were specified and converted from point-based data with longitude and latitude coordinates.

75.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta + \phi \mathbf{z}$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\phi \sim \mathcal{U}(-1, 1)$$

$$\sigma \sim \mathcal{HC}(25)$$

```
z <- as.vector(apply(Z, 1, sum))</pre>
mon.names <- c("LP", "sigma")</pre>
parm.names <- as.parm.names(list(beta=rep(0,J), phi=0, log.sigma=0))</pre>
PGF <- function(Data) return(c(rnormv(Data$J,0,1000), runif(1,-1,1),
    log(rhalfcauchy(1,25))))
MyData <- list(J=J, PGF=PGF, X=X, latitude=latitude, longitude=longitude,
    mon.names=mon.names, parm.names=parm.names, y=y, z=z)
75.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:Data$J]</pre>
    parm[Data$J+1] <- phi <- interval(parm[Data$J+1], -1, 1)</pre>
     sigma <- exp(parm[Data$J+2])</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    phi.prior <- dunif(phi, -1, 1, log=TRUE)</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta)) + phi*Data$z</pre>
    LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + phi.prior + sigma.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma),</pre>
         yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

75.4. Initial Values

Initial. Values $\leftarrow c(rep(0,J), 0.5, log(1))$

76. STARMA(1,1)

The data in this example of a space-time autoregressive moving average (STARMA) are coordinate-based, and the adjacency matrix \mathbf{A} is created from K nearest neighbors. Otherwise, an adjacency matrix may be specified as usual for areal data. Spatial coordinates are given in longitude and latitude for $s=1,\ldots,S$ points in space and measurements are taken across time-periods $t=1,\ldots,T$ for $\mathbf{Y}_{s,t}$.

76.1. Form

```
\mu_{s,t} = \mathbf{X}_{s,t}\beta + \phi \mathbf{W} \mathbf{1}_{s,t-1} + \theta \mathbf{W} \mathbf{2}_{s,t-1}, \quad s = 1, \dots, S, \quad t = 2, \dots, T
\mathbf{W} \mathbf{1} = \mathbf{V} \mathbf{Y}
\mathbf{W} \mathbf{2} = \mathbf{V} \epsilon
\epsilon = \mathbf{Y} - \mu
\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J
\phi \sim \mathcal{U}(-1, 1)
\sigma \sim \mathcal{HC}(25)
\theta \sim \mathcal{N}(0, 1000)
```

where **V** is an adjacency matrix that is scaled so that each row sums to one, β is a vector of regression effects, ϕ is the autoregressive space-time parameter, σ is the residual variance, and θ is the moving average space-time parameter.

```
S <- 100
T <- 10
K <- 5 #Number of nearest neighbors
latitude <- runif(S,0,100)</pre>
longitude <- runif(S,0,100)</pre>
X1 <- matrix(runif(S*T,-2,2), S, T)</pre>
X2 <- matrix(runif(S*T,-2,2), S, T)</pre>
for (t in 2:T) {
    X1[,t] \leftarrow X1[,t-1] + runif(S,-0.1,0.1)
    X2[,t] \leftarrow X2[,t-1] + runif(S,-0.1,0.1)
beta.orig <- runif(3,-2,2); phi.orig <- 0.8; theta.orig <- 1
epsilon <- matrix(rnorm(S*T,0,0.1), S, T)
Z <- matrix(rnorm(S*T,0,0.1), S, T)</pre>
D <- as.matrix(dist(cbind(longitude, latitude), diag=TRUE, upper=TRUE))
A \leftarrow exp(-D)
A \leftarrow ifelse(D == 0, max(D), A)
A <- apply(A, 1, rank)
A <- ifelse(A <= K, 1, 0)
V <- A / rowSums(A) #Scaled matrix
V <- ifelse(is.nan(V), 1/ncol(V), V)</pre>
Y <- beta.orig[1] + beta.orig[2]*X1 + beta.orig*X2
W1 <- tcrossprod(V, t(Y))
Y <- Y + phi.orig*cbind(rep(0,S), W1[,-T])
W2 <- tcrossprod(V, t(epsilon))
Y <- Y + theta.orig*cbind(rep(0,S), W2[,-T])
mon.names <- "LP"
parm.names <- as.parm.names(list(beta=rep(0,3), phi=0, log.sigma=0,
    theta=0))
PGF <- function(Data) return(c(rnormv(3,0,1000), runif(1,-1,1),
```

```
log(rhalfcauchy(1,25)), rnormv(1,0,1000)))
MyData <- list(K=K, PGF=PGF, S=S, T=T, V=V, X1=X1, X2=X2, Y=Y,
    latitude=latitude, longitude=longitude, mon.names=mon.names,
    parm.names=parm.names)
76.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:3]
    parm[4] <- phi <- interval(parm[4], -1, 1)
    sigma <- exp(parm[5])</pre>
    theta <- parm[6]
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    phi.prior <- dunif(phi, -1, 1, log=TRUE)</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    theta.prior <- dnormv(theta, 0, 1000, log=TRUE)
    ### Log-Likelihood
    W1 <- tcrossprod(Data$V, t(Data$Y))</pre>
    mu <- beta[1] + beta[2]*Data$X1 + beta[3]*Data$X2 +</pre>
         phi*cbind(rep(0, Data$S), W1[,-Data$T])
    epsilon <- Data$Y - mu
    W2 <- tcrossprod(Data$V, t(epsilon))
    mu <- mu + theta*cbind(rep(0, Data$S), W2[,-Data$T])</pre>
    LL <- sum(dnorm(Data$Y[,-1], mu[,-1], sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + phi.prior + sigma.prior + theta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
         yhat=rnorm(prod(dim(mu)), mu, sigma), parm=parm)
    return(Modelout)
```

76.4. Initial Values

}

Initial. Values $\leftarrow c(rep(0,3), 0, 1, 0)$

77. State Space Model (SSM), Linear Regression

The data is presented so that the time-series is subdivided into three sections: modeled $(t = 1, ..., T_m)$, one-step ahead forecast $(t = T_m + 1)$, and future forecast $[t = (T_m + 2), ..., T]$. Note that Dyn must also be specified for the SAMWG and SMWG MCMC algorithms.

77.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{1}^{2}), \quad t = 1, \dots, T_{m}$$

$$\mathbf{y}_{t}^{new} \sim \mathcal{N}(\mu_{t}, \sigma_{1}^{2}), \quad t = (T_{m} + 1), \dots, T$$

$$\mu_{t} = \alpha + \mathbf{x}_{t}\beta_{t}, \quad t = 1, \dots, T$$

$$\alpha \sim \mathcal{N}(0, 1000)$$

$$\beta_{1} \sim \mathcal{N}(0, 1000)$$

$$\beta_{t} \sim \mathcal{N}(\beta_{t-1}, \sigma_{2}^{2}), \quad t = 2, \dots, T$$

$$\sigma_{j} \sim \mathcal{HC}(25), \quad j = 1, \dots, 2$$

77.2. Data

```
T <- 20
T.m < -14
beta.orig \leftarrow x \leftarrow rep(0,T)
for (t in 2:T) {
     beta.orig[t] \leftarrow beta.orig[t-1] + rnorm(1,0,0.1)
     x[t] \leftarrow x[t-1] + rnorm(1,0,0.1)
y <-10 + beta.orig*x + rnorm(T,0,0.01)
y[(T.m+2):T] \leftarrow NA
mon.names <- rep(NA, (T-T.m))</pre>
for (i in 1:(T-T.m)) mon.names[i] <- paste("mu[",(T.m+i),"]", sep="")</pre>
parm.names <- as.parm.names(list(alpha=0, beta=rep(0,T),</pre>
     log.sigma=rep(0,2))
PGF <- function(Data) return(c(rnormv(1,0,1000), rnormv(Data$T,0,1000),
     log(rhalfcauchy(2,25))))
MyData <- list(PGF=PGF, T=T, T.m=T.m, mon.names=mon.names,</pre>
     parm.names=parm.names, x=x, y=y)
Dyn <- matrix(paste("beta[",1:T,"]",sep=""), T, 1)</pre>
```

77.4. Initial Values

Initial.Values <- rep(0,T+3)</pre>

78. State Space Model (SSM), Local Level

The local level model is the simplest, non-trivial example of a state space model (SSM). As such, this version of a local level SSM has static variance parameters.

78.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{1}^{2}), \quad t = 1, \dots, T$$

$$\mu_{t} \sim \mathcal{N}(\mu_{t-1}, \sigma_{2}^{2}), \quad t = 2, \dots, T$$

$$\mu_{1} \sim \mathcal{N}(0, 1000)$$

$$\sigma_{j} \sim \mathcal{HC}(25), \quad j = 1, \dots, 2$$

```
MyData <- list(PGF=PGF, T=T, T.m=T.m, mon.names=mon.names,</pre>
     parm.names=parm.names, y=y)
Dyn <- matrix(paste("mu[",1:T,"]",sep=""), T, 1)</pre>
78.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    mu <- parm[1:Data$T]</pre>
     sigma <- exp(parm[-c(1:Data$T)])</pre>
    ### Log(Prior Densities)
    mu.prior <- sum(dnormv(mu[1], 0, 1000, log=TRUE),</pre>
         dnorm(mu[-1], mu[-Data$T], sigma[2], log=TRUE))
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    LL <- sum(dnorm(Data$y[1:Data$T.m], mu[1:Data$T.m], sigma[1],
         log=TRUE))
    yhat <- rnorm(length(mu), c(mu[1], rnorm(Data$T-1, mu[-Data$T],</pre>
         sigma[2])), sigma[1]) #One-step ahead
                                                      ### Log-Posterior
    LP <- LL + mu.prior + sigma.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=mu[(Data$T.m+1):Data$T],</pre>
         yhat=yhat, parm=parm)
    return(Modelout)
    }
```

Initial.Values <- rep(0,T+2)</pre>

79. State Space Model (SSM), Local Linear Trend

The local linear trend model is a state space model (SSM) that extends the local level model to include a dynamic slope parameter. For more information on the local level model, see section 78. This example has static variance parameters.

79.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{1}^{2}), \quad t = 1, \dots, T$$

$$\mu_{t} \sim \mathcal{N}(\mu_{t-1} + \delta_{t-1}, \sigma_{2}^{2}), \quad t = 2, \dots, T$$

$$\mu_{1} \sim \mathcal{N}(0, 1000)$$

$$\delta_{t} \sim \mathcal{N}(\delta_{t-1}, \sigma_{3}^{2}), \quad t = 2, \dots, T$$

```
\delta_1 \sim \mathcal{N}(0, 1000)
\sigma_j \sim \mathcal{HC}(25), \quad j = 1, \dots, 3
```

```
79.2. Data
```

```
T <- 20
T.m < -14
mu.orig <- delta.orig <- rep(0,T)
for (t in 2:T) {
     delta.orig[t] \leftarrow delta.orig[t-1] + rnorm(1,0,0.1)
    mu.orig[t] \leftarrow mu.orig[t-1] + delta.orig[t-1] + rnorm(1,0,1)
y <- mu.orig + rnorm(T,0,0.1)
y[(T.m+2):T] \leftarrow NA
mon.names <- rep(NA, (T-T.m))
for (i in 1:(T-T.m)) mon.names[i] <- paste("yhat[",(T.m+i),"]", sep="")</pre>
parm.names <- as.parm.names(list(mu=rep(0,T), delta=rep(0,T),</pre>
    log.sigma=rep(0,3))
PGF <- function(Data) return(c(rnormv(Data$T,0,10),
     rnormv(Data$T,0,10), log(rhalfcauchy(3,25))))
MyData <- list(PGF=PGF, T=T, T.m=T.m, mon.names=mon.names,</pre>
    parm.names=parm.names, y=y)
79.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    mu <- parm[1:Data$T]</pre>
    delta <- parm[Data$T+(1:Data$T)]</pre>
    sigma <- exp(parm[2*Data$T+c(1:3)])</pre>
    ### Log(Prior Densities)
    mu.prior <- sum(dnormv(mu[1], 0, 1000, log=TRUE),</pre>
         dnorm(mu[-1], mu[-Data$T]+delta[-Data$T], sigma[2],
         log=TRUE))
    delta.prior <- sum(dnormv(delta[1], 0, 1000, log=TRUE),</pre>
         dnorm(delta[-1], delta[-Data$T], sigma[3], log=TRUE))
     sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
    ### Log-Likelihood
    LL <- sum(dnorm(Data$y[1:Data$T.m], mu[1:Data$T.m], sigma[1],
         log=TRUE))
    yhat <- rnorm(length(mu), c(mu[1], rnorm(Data$T-1, mu[-Data$T],</pre>
```

sigma[2])), sigma[1]) #One-step ahead

LP <- LL + mu.prior + delta.prior + sigma.prior</pre>

Modelout <- list(LP=LP, Dev=-2*LL, Monitor=mu[(Data\$T.m+1):Data\$T],</pre>

Log-Posterior

```
yhat=yhat, parm=parm)
return(Modelout)
}
```

Initial.Values <- rep(0,T*2+3)</pre>

80. State Space Model (SSM), Stochastic Volatility (SV)

80.1. Form

$$\mathbf{y} \sim \mathcal{N}(0, \sigma^{2})$$

$$\sigma^{2} = \frac{1}{\exp(\theta)}$$

$$\beta = \exp(\mu/2)$$

$$\theta_{1} \sim \mathcal{N}(\mu + \phi(\alpha - \mu), \tau)$$

$$\theta_{t} \sim \mathcal{N}(\mu + \phi(\theta_{t-1} - \mu), \tau), \quad t = 2, \dots, T$$

$$\alpha \sim \mathcal{N}(\mu, \tau)$$

$$\phi \sim \mathcal{U}(-1, 1)$$

$$\mu \sim \mathcal{N}(0, 10)$$

$$\tau \sim \mathcal{HC}(25)$$

80.2. Data

```
T <- 20
y <- rep(10,T); epsilon <- rnorm(T,0,1)
for (t in 2:T) {y[t] <- 0.8*y[t-1] + epsilon[t-1]}
mon.names <- c("LP","tau", paste("sigma2[",1:T,"]",sep=""))
parm.names <- as.parm.names(list(theta=rep(0,T), alpha=0, phi=0, mu=0, log.tau=0))
PGF <- function(Data) return(c(rnormv(Data$T,0,10), rnormv(1,rnorm(1,0,10),rhalfcauchy(1,25)), runif(1,-1,1), rnormv(1,0,10), log(rhalfcauchy(1,25))))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names, y=y)
Dyn <- matrix(paste("theta[",1:T,"]",sep=""), T, 1)</pre>
```

```
Model <- function(parm, Data)
{</pre>
```

```
### Parameters
theta <- parm[1:Data$T]
alpha <- parm[Data$T+1]</pre>
parm[Data$T+2] <- phi <- interval(parm[Data$T+2], -1, 1)</pre>
mu <- parm[Data$T+3]</pre>
tau <- exp(parm[Data$T+4])</pre>
### Log(Prior Densities)
alpha.prior <- dnormv(alpha, mu, tau, log=TRUE)</pre>
theta.prior <- sum(dnormv(theta[1], mu + phi*(alpha-mu), tau,
     log=TRUE), dnormv(theta[-1], mu + phi*(theta[-Data$T]-mu), tau,
     log=TRUE))
phi.prior <- dunif(phi, -1, 1, log=TRUE)
mu.prior <- dnormv(mu, 0, 10, log=TRUE)</pre>
tau.prior <- dhalfcauchy(tau, 25, log=TRUE)</pre>
### Log-Likelihood
beta \leftarrow exp(mu / 2)
sigma2 <- 1 / exp(theta)</pre>
LL <- sum(dnormv(Data$y, 0, sigma2, log=TRUE))
### Log-Posterior
LP <- LL + alpha.prior + theta.prior + phi.prior + mu.prior +
     tau.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, tau, sigma2),</pre>
     yhat=rnormv(length(Data$y), 0, sigma2), parm=parm)
return(Modelout)
}
```

80.4. Initial Values

Initial.Values <- rep(0,T+4)</pre>

81. Threshold Autoregression (TAR)

81.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\nu_{t}, \sigma^{2}), \quad t = 1, \dots, T$$

$$\mathbf{y}^{new} = \alpha_{2} + \phi_{2}\mathbf{y}_{T}$$

$$\nu_{t} = \begin{cases} \alpha_{1} + \phi_{1}\mathbf{y}_{t-1}, & t = 1, \dots, T & \text{if } t \geq \theta \\ \alpha_{2} + \phi_{2}\mathbf{y}_{t-1}, & t = 1, \dots, T & \text{if } t < \theta \end{cases}$$

$$\alpha_{j} \sim \mathcal{N}(0, 1000) \in [-1, 1], \quad j = 1, \dots, 2$$

$$\phi_{j} \sim \mathcal{N}(0, 1000), \in [-1, 1], \quad j = 1, \dots, 2$$

$$\theta \sim \mathcal{U}(2, T - 1)$$

$\sigma \sim \mathcal{HC}(25)$

81.2. Data

```
y \leftarrow c(0.02, -0.51, -0.30, 1.46, -1.26, -2.15, -0.91, -0.53, -1.91,
    2.64, 1.64, 0.15, 1.46, 1.61, 1.96, -2.67, -0.19, -3.28,
    1.89, 0.91, -0.71, 0.74, -0.10, 3.20, -0.80, -5.25, 1.03,
    -0.40, -1.62, -0.80, 0.77, 0.17, -1.39, -1.28, 0.48, -1.02,
    0.09, -1.09, 0.86, 0.36, 1.51, -0.02, 0.47, 0.62, -1.36,
    1.12, 0.42, -4.39, -0.87, 0.05, -5.41, -7.38, -1.01, -1.70,
    0.64, 1.16, 0.87, 0.28, -1.69, -0.29, 0.13, -0.65, 0.83,
    0.62, 0.05, -0.14, 0.01, -0.36, -0.32, -0.80, -0.06, 0.24,
    0.23, -0.37, 0.00, -0.33, 0.21, -0.10, -0.10, -0.01, -0.40,
    -0.35, 0.48, -0.28, 0.08, 0.28, 0.23, 0.27, -0.35, -0.19,
    0.24, 0.17, -0.02, -0.23, 0.03, 0.02, -0.17, 0.04, -0.39,
    -0.12, 0.16, 0.17, 0.00, 0.18, 0.06, -0.36, 0.22, 0.14,
    -0.17, 0.10, -0.01, 0.00, -0.18, -0.02, 0.07, -0.06, 0.06,
    -0.05, -0.08, -0.07, 0.01, -0.06, 0.01, 0.01, -0.02, 0.01,
    0.01, 0.12, -0.03, 0.08, -0.10, 0.01, -0.03, -0.08, 0.04,
    -0.09, -0.08, 0.01, -0.05, 0.08, -0.14, 0.06, -0.11, 0.09,
    0.06, -0.12, -0.01, -0.05, -0.15, -0.05, -0.03, 0.04, 0.00,
    -0.12, 0.04, -0.06, -0.05, -0.07, -0.05, -0.14, -0.05, -0.01,
    -0.12, 0.05, 0.06, -0.10, 0.00, 0.01, 0.00, -0.08, 0.00,
    0.00, 0.07, -0.01, 0.00, 0.09, 0.33, 0.13, 0.42, 0.24,
    -0.36, 0.22, -0.09, -0.19, -0.10, -0.08, -0.07, 0.05, 0.07,
    0.07, 0.00, -0.04, -0.05, 0.03, 0.08, 0.26, 0.10, 0.08,
    0.09, -0.07, -0.33, 0.17, -0.03, 0.07, -0.04, -0.06, -0.06,
    0.07, -0.03, 0.00, 0.08, 0.27, 0.11, 0.11, 0.06, -0.11,
    -0.09, -0.21, 0.24, -0.12, 0.11, -0.02, -0.03, 0.02, -0.10,
    0.00, -0.04, 0.01, 0.02, -0.03, -0.10, -0.09, 0.17, 0.07,
    -0.05, -0.01, -0.05, 0.01, 0.00, -0.08, -0.05, -0.08, 0.07,
    0.06, -0.14, 0.02, 0.01, 0.04, 0.00, -0.13, -0.17)
T <- length(y)
mon.names <- c("LP", "sigma", "ynew")</pre>
parm.names <- as.parm.names(list(alpha=rep(0,2), phi=rep(0,2), theta=0,
    log.sigma=0))
PGF <- function(Data) return(c(rtrunc(4, "norm", a=-1, b=1, mean=0,
    sd=sqrt(1000)), runif(1,2,Data$T-1), log(rhalfcauchy(1,25))))
MyData <- list(PGF=PGF, T=T, mon.names=mon.names, parm.names=parm.names, y=y)
81.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
```

parm[1:2] <- alpha <- interval(parm[1:2], -1, 1)

```
parm[3:4] <- phi <- interval(parm[3:4], -1, 1)
parm[5] <- theta <- interval(parm[5], 2, Data$T-1)</pre>
sigma <- exp(parm[6])</pre>
### Log(Prior Densities)
alpha.prior <- sum(dtrunc(alpha, "norm", a=-1, b=1, mean=0,
     sd=sqrt(1000), log=TRUE))
phi.prior <- sum(dtrunc(phi, "norm", a=-1, b=1, mean=0,
    sd=sqrt(1000), log=TRUE))
alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))</pre>
phi.prior <- sum(dnormv(phi, 0, 1000, log=TRUE))</pre>
theta.prior <- dunif(theta, 2, Data$T-1, log=TRUE)</pre>
sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
### Log-Likelihood
mu <- matrix(0, Data$T, 2)</pre>
mu[,1] <- c(alpha[1], alpha[1] + phi[1]*Data$y[-Data$T])</pre>
mu[,2] \leftarrow c(alpha[2], alpha[2] + phi[2]*Data$y[-Data$T])
nu <- ifelse(1:Data$T < theta, mu[,1], mu[,2])</pre>
ynew <- rnorm(1, alpha[2] + phi[2]*Data$y[Data$T], sigma)</pre>
LL <- sum(dnorm(Data$y, nu, sigma, log=TRUE))
### Log-Posterior
LP <- LL + alpha.prior + phi.prior + theta.prior + sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma,ynew),</pre>
     yhat=rnorm(length(nu), nu, sigma), parm=parm)
return(Modelout)
}
```

81.4. Initial Values

Initial. Values $\leftarrow c(rep(0,4), T/2, log(1))$

82. TARCH(1)

In this TARCH example, there are two regimes, one for positive residuals in the previous time-period, and the other for negative. The TARCH parameters are the θ vector.

82.1. Form

$$\mathbf{y}_{t} \sim \mathcal{N}(\mu_{t}, \sigma_{t}^{2}), \quad t = 2, \dots, T$$

$$\sigma_{t}^{2} = \omega + \theta_{1} \delta_{t-1} \epsilon_{t-1}^{2} + \theta_{2} (1 - \delta_{t-1}) \epsilon_{t-1}^{2}, \quad t = 2, \dots, T$$

$$\delta_{t} = \begin{cases} 1 & \text{if } \epsilon_{t} > 0 \\ 0 & \\ \epsilon = \mathbf{y} - \mu \end{cases}$$

$$\mu_{t} = \alpha + \phi \mathbf{y}_{t-1}, \quad t = 2, \dots, T$$

```
\alpha \sim \mathcal{N}(0, 1000)
\phi \sim \mathcal{U}(-1, 1)
\omega \sim \mathcal{HC}(25)
\theta_j \sim \mathcal{U}(0, 1), \quad j = 1, \dots, 2
```

82.2. Data

82.3. Model

```
Model <- function(parm, Data)</pre>
    ### Parameters
    alpha <- parm[1]</pre>
    parm[2] <- phi <- interval(parm[2], -1, 1)</pre>
    omega <- exp(parm[3])</pre>
    parm[4:5] <- theta <- interval(parm[4:5], 0.001, 0.999)</pre>
    ### Log(Prior Densities)
     alpha.prior <- dnormv(alpha, 0, 1000, log=TRUE)</pre>
    phi.prior <- dunif(phi, -1, 1, log=TRUE)
    omega.prior <- dhalfcauchy(omega, 25, log=TRUE)</pre>
    theta.prior <- sum(dunif(theta, 0, 1, log=TRUE))</pre>
    ### Log-Likelihood
    mu <- alpha + c(0, Data$y[-Data$T]) * phi</pre>
    epsilon <- Data$y - mu
    delta \leftarrow (epsilon > 0) * 1
    sigma2 \leftarrow omega + theta[1] * c(0,delta[-Data$T]) *
         c(0, epsilon[-Data$T]^2)
    sigma2[-1] <- sigma2[-1] + theta[2] * (1 - delta[-Data$T]) *
          epsilon[-Data$T]^2
     sigma2.new <- omega + theta[1] * delta[Data$T] * epsilon[Data$T]^2 +
         theta[2] * (1 - delta[Data$T]) * epsilon[Data$T]^2
```

82.4. Initial Values

Initial. Values $\leftarrow c(0, 0, 1, 0.5, 0.5)$

83. Variable Selection, BAL

This approach to variable selection is one of several forms of the Bayesian Adaptive Lasso (BAL). The lasso applies shrinkage to exchangeable scale parameters, γ , for the regression effects, β .

83.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{L}(0, \gamma_j), \quad j = 1, \dots, J$$

$$\gamma_j \sim \mathcal{G}^{-1}(\delta, \tau), \quad \in [0, \infty]$$

$$\delta \sim \mathcal{HC}(25)$$

$$\tau \sim \mathcal{HC}(25)$$

$$\sigma \sim \mathcal{HC}(25)$$

83.2. Data

83.3. Model

```
Model <- function(parm, Data)</pre>
     {
    ### Hyperhyperparameters
    delta <- exp(parm[2*Data$J+1])</pre>
    tau <- exp(parm[2*Data$J+2])</pre>
    ### Hyperparameters
     gamma <- interval(parm[Data$J+1:Data$J], 0, Inf)</pre>
    ### Parameters
    beta <- parm[1:Data$J]</pre>
     sigma <- exp(parm[2*Data$J+3])</pre>
    ### Log(Hyperhyperprior Densities)
    delta.prior <- dhalfcauchy(delta, 25, log=TRUE)</pre>
    tau.prior <- dhalfcauchy(tau, 25, log=TRUE)</pre>
    ### Log(Hyperprior Densities)
     gamma.prior <- sum(dinvgamma(gamma, delta, tau, log=TRUE))</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dlaplace(beta, 0, gamma, log=TRUE))</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta))</pre>
    LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + gamma.prior + delta.prior + tau.prior +
          sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, sigma),</pre>
          yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

83.4. Initial Values

```
Initial. Values \leftarrow c(rep(0,J), rep(0,J), rep(1,3))
```

84. Variable Selection, RJ

This example uses the RJ (Reversible-Jump) algorithm of the LaplacesDemon function for variable selection and Bayesian Model Averaging (BMA). Other MCMC algorithms will not perform variable selection with this example, as presented. This is an example of variable selection in a linear regression. The only difference between the following example, and the example of linear regression (40), is that RJ specifications are also included for the RJ

algorithm, and that the RJ algorithm must be used.

84.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

84.2. Data

```
N <- 1000
J <- 100 #Number of predictors, including the intercept
X <- matrix(1,N,J)</pre>
for (j in 2:J) {X[,j] \leftarrow rnorm(N,runif(1,-3,3),runif(1,0.1,1))}
beta.orig <- runif(J,-3,3)</pre>
zero <- sample(2:J, round(J*0.9)) #Assign most parameters to be zero
beta.orig[zero] <- 0</pre>
e < rnorm(N,0,0.1)
y <- as.vector(tcrossprod(beta.orig, X) + e)
mon.names <- c("LP", "sigma")</pre>
parm.names <- as.parm.names(list(beta=rep(0,J), log.sigma=0))</pre>
PGF <- function(Data) return(c(rnormv(Data$J,0,1),</pre>
    log(rhalfcauchy(1,25))))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names,</pre>
    parm.names=parm.names, y=y)
### Reversible-Jump Specifications bin.n <- J-1 #Maximum allowable model size
bin.p <- 0.4 #Most probable size: bin.p x bin.n is binomial mean and median
parm.p \leftarrow rep(1/J,J+1)
selectable=c(0, rep(1,J-1), 0)
```

84.3. Model

```
Model <- function(parm, Data)
    {
     ### Parameters
     beta <- parm[1:Data$J]
     sigma <- exp(parm[Data$J+1])
     ### Log(Prior Densities)
     beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))
     sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)
     ### Log-Likelihood
     mu <- tcrossprod(Data$X, t(beta))
     LL <- sum(dnorm(Data$y, mu, sigma, log=TRUE))</pre>
```

Initial.Values <- GIV(Model, MyData, PGF=TRUE)</pre>

85. Variable Selection, SSVS

This example uses a modified form of the random-effects (or global adaptation) Stochastic Search Variable Selection (SSVS) algorithm presented in O'Hara and Sillanpaa (2009), which selects variables according to practical significance rather than statistical significance. Here, SSVS is applied to linear regression, though this method is widely applicable. For J variables, each regression effects vector β_j is conditional on γ_j , a binary inclusion variable. Each β_j is a discrete mixture distribution with respect to $\gamma_j = 0$ or $\gamma_j = 1$, with precision 100 or $\beta_{\sigma} = 0.1$, respectively. As with other representations of SSVS, these precisions may require tuning.

With other representations of SSVS, each γ_j is Bernoulli-distributed, though this would be problematic in Laplace's Demon, because γ_j is discrete, rather than continuous. To keep γ in the monitors, a beta density is placed on each prior δ_j , with parameters 1 and 2, and experimentation is encouraged. Each δ_j is constrained to the interval [0.01, 0.99], and rounded to γ_j . Note that $\lfloor x + 0.5 \rfloor$ means to round x. The prior for δ can be manipulated to influence sparseness.

When the goal is to select the best model, each $\mathbf{X}_{1:N,j}$ is retained for a future run when the posterior mean of $\gamma_j \geq 0.5$. When the goal is model-averaging, the results of this model may be used directly, which would please L. J. Savage, who said that "models should be as big as an elephant" (Draper 1995).

85.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$(\beta_j | \gamma_j) \sim (1 - \gamma_j) \mathcal{N}(0, 0.01) + \gamma_j \mathcal{N}(0, \beta_\sigma^2) \quad j = 1, \dots, J$$

$$\beta_\sigma \sim \mathcal{HC}(25)$$

$$\gamma_j = \lfloor \delta_j + 0.5 \rfloor, \quad j = 1, \dots, J$$

$$\delta_j \sim \mathcal{B}(1, 2) \in [0.01, 0.99], \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

85.2. Data

```
data(demonsnacks)
J <- ncol(demonsnacks)</pre>
y <- log(demonsnacks$Calories)</pre>
X <- cbind(1, as.matrix(demonsnacks[,c(1,3:10)]))</pre>
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}</pre>
mon.names <- c("LP", "min.beta.sigma", "sigma",</pre>
     as.parm.names(list(gamma=rep(0,J))))
parm.names <- as.parm.names(list(beta=rep(0,J), delta=rep(0,J),</pre>
     log.beta.sigma=0, log.sigma=0))
PGF <- function(Data) return(c(rnorm(Data$J,0,1), rbeta(Data$J,1,2),
    log(rhalfcauchy(2,25))))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names,</pre>
    parm.names=parm.names, y=y)
85.3. Model
```

```
Model <- function(parm, Data)</pre>
    ### Hyperparameters
    beta.sigma <- exp(parm[grep("log.beta.sigma", Data$parm.names)])</pre>
    ### Parameters
    beta <- parm[1:Data$J]</pre>
     delta <- interval(parm[grep("delta", Data$parm.names)],-100,100)
    parm[grep("delta", Data$parm.names)] <- delta</pre>
    gamma <- round(delta)</pre>
    beta.sigma <- ifelse(gamma == 0, 0.1, beta.sigma)
     sigma <- exp(parm[grep("log.sigma", Data$parm.names)])</pre>
    ### Log(Hyperprior Densities)
    beta.sigma.prior <- sum(dhalfcauchy(beta.sigma, 25, log=TRUE))</pre>
     ### Log(Prior Densities)
    beta.prior <- sum(dnorm(beta, 0, beta.sigma, log=TRUE))</pre>
    delta.prior <- sum(dbeta(delta, 1, 2, log=TRUE))</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- tcrossprod(beta, Data$X)</pre>
    LL <- sum(dnorm(y, mu, sigma, log=TRUE))
    ### Log-Posterior
    LP <- LL + beta.prior + beta.sigma.prior + delta.prior + sigma.prior</pre>
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP, min(beta.sigma),</pre>
         sigma, gamma), yhat=rnorm(length(mu), mu, sigma), parm=parm)
    return(Modelout)
    }
```

Initial. Values $\leftarrow c(rep(0,J), rep(0,J), log(1), log(1))$

86. Vector Autoregression, VAR(1)

86.1. Form

$$\mathbf{Y}_{t,j} \sim \mathcal{N}(\mu_{t,j}, \sigma_j^2), \quad t = 1, \dots, T, \quad j = 1, \dots, J$$

$$\mu_{t,j} = \alpha_j + \Phi_{1:J,j} \mathbf{Y}_{t-1,j}$$

$$\mathbf{y}_j^{new} = \alpha_j + \Phi_{1:J,j} \mathbf{Y}_{T,j}$$

$$\alpha_j \sim \mathcal{N}(0, 1000)$$

$$\sigma_j \sim \mathcal{HC}(25)$$

$$\Phi_{i,k} \sim \mathcal{N}(0, 1000), \quad i = 1, \dots, J, \quad k = 1, \dots, J$$

86.2. Data

86.3. Model

```
Model <- function(parm, Data)
    {
    ### Parameters
    alpha <- parm[1:Data$J]
    Phi <- matrix(parm[grep("Phi", Data$parm.names)], Data$J, Data$J)
    sigma <- exp(parm[grep("log.sigma", Data$parm.names)])
    ### Log(Prior Densities)
    alpha.prior <- sum(dnormv(alpha, 0, 1000, log=TRUE))
    Phi.prior <- sum(dnormv(Phi, 0, 1000, log=TRUE))
    sigma.prior <- sum(dhalfcauchy(sigma, 25, log=TRUE))</pre>
```

```
### Log-Likelihood
mu <- matrix(alpha,Data$T,Data$J,byrow=TRUE)
    mu[-1,] <- mu[-1,] + tcrossprod(Data$Y[-Data$T,], Phi)
Sigma <- matrix(sigma, Data$T, Data$J, byrow=TRUE)
ynew <- rnorm(Data$J, alpha + as.vector(crossprod(Phi, Data$Y[Data$T,])),
    sigma)
LL <- sum(dnorm(Data$Y, mu, Sigma, log=TRUE))
### Log-Posterior
LP <- LL + alpha.prior + Phi.prior + sigma.prior
Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,ynew),
    yhat=rnorm(prod(dim(mu)), mu, Sigma), parm=parm)
return(Modelout)
}</pre>
```

86.4. Initial Values

Initial.Values <- c(colMeans(Y), rep(0,J*J), rep(log(1),J))</pre>

87. Weighted Regression

It is easy enough to apply record-level weights to the likelihood. Here, weights are applied to the linear regression example in section 40.

87.1. Form

$$\mathbf{y} \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \mathbf{X}\beta$$

$$\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J$$

$$\sigma \sim \mathcal{HC}(25)$$

87.2. Data

```
data(demonsnacks)
N <- nrow(demonsnacks)
J <- ncol(demonsnacks)
y <- log(demonsnacks$Calories)
X <- cbind(1, as.matrix(demonsnacks[,c(1,3:10)]))
for (j in 2:J) {X[,j] <- CenterScale(X[,j])}
w <- c(rep(1,5), 0.2, 1, 0.01, rep(1,31))
w <- w * (sum(w) / N)
mon.names <- c("LP","sigma")
parm.names <- as.parm.names(list(beta=rep(0,J), log.sigma=0))
PGF <- function(Data) return(c(rnormy(Data$J,0,1000),</pre>
```

```
log(rhalfcauchy(1,25))))
MyData <- list(J=J, PGF=PGF, X=X, mon.names=mon.names,</pre>
    parm.names=parm.names, w=w, y=y)
87.3. Model
Model <- function(parm, Data)</pre>
    ### Parameters
    beta <- parm[1:Data$J]</pre>
     sigma <- exp(parm[Data$J+1])</pre>
    ### Log(Prior Densities)
    beta.prior <- sum(dnormv(beta, 0, 1000, log=TRUE))</pre>
    sigma.prior <- dhalfcauchy(sigma, 25, log=TRUE)</pre>
    ### Log-Likelihood
    mu <- tcrossprod(Data$X, t(beta))</pre>
    LL <- sum(w * dnorm(Data$y, mu, sigma, log=TRUE))</pre>
    ### Log-Posterior
    LP <- LL + beta.prior + sigma.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=c(LP,sigma),</pre>
         yhat=rnorm(length(mu), mu, sigma), parm=parm)
```

return(Modelout)

Initial.Values <- c(rep(0,J), log(1))</pre>

88. Zero-Inflated Poisson (ZIP)

88.1. Form

}

$$\mathbf{y} \sim \mathcal{P}(\Lambda_{1:N,2})$$

$$\mathbf{z} \sim \mathcal{BERN}(\Lambda_{1:N,1})$$

$$\mathbf{z}_i = \begin{cases} 1 & \text{if } \mathbf{y}_i = 0 \\ 0 & \end{cases}$$

$$\Lambda_{i,2} = \begin{cases} 0 & \text{if } \Lambda_{i,1} \ge 0.5 \\ \Lambda_{i,2} & \end{cases}$$

$$\Lambda_{1:N,1} = \frac{1}{1 + \exp(-\mathbf{X}_1 \alpha)}$$

$$\Lambda_{1:N,2} = \exp(\mathbf{X}_2 \beta)$$

return(Modelout)

}

```
\alpha_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J_1
\beta_j \sim \mathcal{N}(0, 1000), \quad j = 1, \dots, J_2
```

```
88.2. Data
N <- 1000
J1 <- 4
J2 <- 3
X1 <- matrix(runif(N*J1,-2,2),N,J1); X1[,1] <- 1</pre>
X2 \leftarrow matrix(runif(N*J2,-2,2),N,J2); X2[,1] \leftarrow 1
alpha <- runif(J1,-1,1)</pre>
beta <- runif(J2,-1,1)
p <- invlogit(tcrossprod(X1, t(alpha)) + rnorm(N,0,0.1))</pre>
mu <- round(exp(tcrossprod(X2, t(beta)) + rnorm(N,0,0.1)))</pre>
y \leftarrow ifelse(p > 0.5, 0, mu)
z \leftarrow ifelse(y == 0, 1, 0)
mon.names <- "LP"
parm.names <- as.parm.names(list(alpha=rep(0,J1), beta=rep(0,J2)))</pre>
PGF <- function(Data) return(c(rnormv(Data$J1,0,5), rnormv(Data$J2,0,5)))
MyData <- list(J1=J1, J2=J2, N=N, PGF=PGF, X1=X1, X2=X2,</pre>
    mon.names=mon.names, parm.names=parm.names, y=y, z=z)
88.3. Model
Model <- function(parm, Data)</pre>
    {
     ### Parameters
    parm[1:Data$J1] <- alpha <- interval(parm[1:Data$J1], -5, 5)</pre>
    beta <- parm[Data$J1+1:Data$J2]</pre>
    parm[Data$J1+1:Data$J2] <- beta</pre>
    ### Log(Prior Densities)
    alpha.prior <- sum(dnormv(alpha, 0, 5, log=TRUE))</pre>
    beta.prior <- sum(dnormv(beta, 0, 5, log=TRUE))</pre>
    ### Log-Likelihood
    Lambda <- matrix(NA, Data$N, 2)</pre>
    Lambda[,1] <- invlogit(tcrossprod(Data$X1, t(alpha)))</pre>
    Lambda[,2] <- exp(tcrossprod(Data$X2, t(beta)))</pre>
    Lambda[which(Lambda[,1] >= 0.5),2] <- 0
    LL <- sum(dbern(Data$z, Lambda[,1], log=TRUE),
          dpois(Data$y, Lambda[,2], log=TRUE))
    ### Log-Posterior
    LP <- LL + alpha.prior + beta.prior
    Modelout <- list(LP=LP, Dev=-2*LL, Monitor=LP,</pre>
          yhat=rpois(nrow(Lambda), Lambda[,2]), parm=parm)
```

Initial.Values <- GIV(Model, MyData, n=10000)</pre>

References

- Barnard J, McCulloch R, Meng X (2000). "Modeling Covariance Matrices in Terms of Standard Deviations and Correlations, with Application to Shrinkage." *Statistica Sinica*, **10**, 1281–1311.
- Congdon P (2003). Applied Bayesian Modelling. John Wiley & Sons, West Sussex, England.
- Draper D (1995). "Assessment and Propagation of Model Uncertainty." *Journal of the Royal Statistical Society*, **B 57**(1), 45–97.
- Gelman A (2011). R2WinBUGS: Running WinBUGS and OpenBUGS from R / S-PLUS. R package version 2.1-18, URL http://cran.r-project.org/web/packages/R2WinBUGS/index.html.
- Gelman A, Carlin J, Stern H, Rubin D (2004). Bayesian Data Analysis. 2nd edition. Chapman & Hall, Boca Raton, FL.
- Ibrahim J, Chen M (2000). "Power Prior Distributions for Regression Models." *Statistical Science*, **15**, 46–60.
- Kleine L (1950). Economic Fluctuations in the United States 1921-1940. John Wiley & Sons, New York, New York.
- Kotz S, Kozubowski T, Podgorski K (2001). The Laplace Distribution and Generalizations: A Revisit with Applications to Communications, Economics, Engineering, and Finance. Birkauser, Boston.
- O'Hara R, Sillanpaa M (2009). "A Review of Bayesian Variable Selection Methods: What, How and Which." *Journal of Bayesian Analysis*, **4**(1), 85–118.
- R Development Core Team (2012). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL http://www.R-project.org.
- Spiegelhalter D, Thomas A, Best N, Lunn D (2003). WinBUGS User Manual, Version 1.4. MRC Biostatistics Unit, Institute of Public Health and Department of Epidemiology and Public Health, Imperial College School of Medicine, UK.
- Statisticat LLC (2012). LaplacesDemon: Complete Environment for Bayesian Inference. R package version 13.01.07, URL http://cran.r-project.org/web/packages/LaplacesDemon/index.html.
- Zellner A (1962). "An Efficient Method of Estimating Seemingly Unrelated Regression Equations and Tests for Aggregation Bias." *Journal of the American Statistical Association*, **57**, 348–368.

Affiliation:

 $\begin{array}{c} {\rm Statisticat,\ LLC} \\ {\rm Farmington,\ CT} \end{array}$

E-mail: software@bayesian-inference.com

URL: http://www.bayesian-inference.com/software