

ΠΑΡΑΡΤΗΜΑ Α (1^ο ΜΑΘΗΜΑΤΟΣ): ΠΑΡΑΔΕΙΓΜΑΤΑ BUGS

1 EXAMPLE 1: BIRTHWEIGHT & ESTRIOL LEVEL

1.1 Model (Bug file)

```
model example1;
const n=31; # n=sample size
var estriol[n], # estriol level of pregnant woman
    birth[n], # birthweight
    mu[n], # regression expected value
    a.star,a,b,tau,s2; # model parameters,
                        # tau = precision, s2=1/tau error variance
data estriol,birth in 'estriol.dat';
inits in 'estriol2.ini';
{
#      definition of likelihood function
#
#      for (i in 1:n) {
#          birth[i]~dnorm( mu[i], tau ); # random component
#          systematic component & link function
#          mu[i]<-a.star+b*(estriol[i]-mean(estriol[]));
#      }
#      prior distributions
#
#      a.star~dnorm( 0, 1.0E-04 ); # normal prior for a
#      b~dnorm( 0, 1.0E-04 ); # normal prior for b
#      tau~dgamma( 1.0E-04 , 1.0E-04 ); # gamma prior for precision
#      s2<-1/tau;
#      a<-a.star-b*mean(estriol[]);
}
```

1.2 Initial Values (estriol2.ini)

```
list(a.star=0.0, b=0.0, tau=1.0)
```

1.3 Data (estriol.dat)

```
7   25
9   25
9   25
12  27
14  27
16  27
16  24
14  30
16  30
16  31
17  30
19  31
21  30
24  28
15  32
16  32
17  32
25  32
27  34
15  34
15  34
15  35
16  35
19  34
18  35
17  36
18  37
20  38
22  40
25  39
```

2 EXAMPLE 2: BEETLES DATASET

2.1 Model (*Bug file*)

```
model beetles;
const
  N = 8;      # number of doses
var
  r[N],p[N],x[N],n[N],alpha,alpha.star,beta,r.hat[N], odds.ratio;
data x, n, r in "beetles.dat";
inits in "beetles.in";
{
  for (i in 1:N) {
    r[i] ~ dbin(p[i], n[i]);
    logit(p[i])   <- alpha.star + beta*(x[i]-mean(x[]));
    r.hat[i] <- p[i]*n[i];  # fitted values
  }
  alpha.star ~ dnorm(0.0, 1.0E-3);
  beta     ~ dnorm(0.0, 1.0E-3);
  alpha    <- alpha.star - beta*mean(x[]);
  odds.ratio <- exp(beta)
}
```

2.2 Initial Values (*beetles.ini*)

```
list(alpha.star=0, beta=0)
```

2.3 Data (*beetles.dat*)

1.6907	59	6
1.7242	60	13
1.7552	62	18
1.7842	56	28
1.8113	63	52
1.8369	59	53
1.8610	62	61
1.8839	60	60

3 EXAMPLE 3: WAIS & SENILITY SYMPTOMS

3.1 Model (*Bug file*)

```
model example3;
const n=54;          # number of observations
var   wais[n],       # wais measurement
      symptom[n],    # symptom binary indicator
      p[n],           # probability of symptom appearance
      alpha,          # alpha parameter
      beta,           # beta parameter
      odds.ratio,     # odds ratio
      x.fifty,         # value of x for p=1/2
      p.meanx;        # fitted probability for x=mean(x)
data wais, symptom in 'wais.dat';
inits in 'wais.ini';
{
# Likelihood Definition
  for (i in 1:n) {
# random component
    symptom[i]~dbern( p[i] );
# systematic component & link function
    logit( p[i] ) <- alpha + beta * wais[i];
  }
# calculation of odds ratio
  odds.ratio<-exp(beta);
# priors for alpha and beta
  alpha~dnorm(0.0, 0.00001);
  beta~dnorm(0.0, 0.00001);
# x.fifty
  x.fifty<- -alpha/beta
# symptom probability for mean of observed wais
  p.meanx<-exp( alpha+beta*mean(wais[]) )/
(1+exp(alpha+beta*mean(wais[])) )
}
```

3.2 Initial Values (*wais.ini*)

```
list( alpha=0.0, beta=0.0 )
```

3.3 Data (*wais.dat*)

```
9      1
13     1
6      1
8      1
10     1
4      1
14     1
8      1
11     1
7      1
9      1
7      1
5      1
14     1
13     0
16     0
10     0
12     0
11     0
14     0
15     0
18     0
7      0
16     0
```

9	0
9	0
11	0
13	0
15	0
13	0
10	0
11	0
6	0
17	0
14	0
19	0
9	0
11	0
14	0
10	0
16	0
10	0
16	0
14	0
13	0
13	0
9	0
15	0
10	0
11	0
12	0
4	0
14	0
20	0

4 EXAMPLE 4: BREAST CANCER & AGE AT 1st BIRTH

4.1 Model (*Bug file*)

```
model example4;
const n=4;          # number of cells
var   status[n],    # study group (1=case, 0=control)
      age[n],       # age at 1st birth (0=age<30, 1=age>29)
      counts[n],     # cell counts
      lambda[n],     # expected number of cells
      mu,            # constant parameter
      a,             # status effect
      b,             # age effect
      ab,            # interaction between status and age
      odds.ratio;   # odds ratio
data status, age, counts in 'breast.dat';
inits in 'breast.ini';
{
#      Likelihood Definition
#      for (i in 1:n) {
#          random component
#          counts[i]~dpois( lambda[i] );
#          systematic component & link function
#          log( lambda[i] ) <- mu + a*status[i] +b*age[i]+
ab*status[i]*age[i];
#      }
#      calculation of odds ratio
#      odds.ratio<-exp(ab);
#      priors for model parameters
#      mu~dnorm(0.0, 0.00001);
#      a~dnorm(0.0, 0.00001);
#      b~dnorm(0.0, 0.00001);
#      ab~dnorm(0.0, 0.00001);
}
```

4.2 Initial Values (*breast.ini*)

```
list(mu=0,a=0,b=0,ab=0)
```

4.3 Data (*breast.dat*)

```
1 1 683
1 0 2537
0 1 1498
0 0 8747
```

5 EXAMPLE 5: 2x2x2 COMMON ODDS RATIO ESTIMATION: PASSIVE SMOKING & CANCER ADJUSTING FOR SMOKING STATUS

5.1 ANALYSIS 1: DIFFERENT RISK PER SMOKING STATUS

5.1.1 Model (Bug File)

```
model ex5notcommonOR;
const n=8,
      p=4;
var
  smoking[n],
  status[n],
  passive[n],
  counts[n],
  b[2,p],
  lambda[n],
  or[2];
data smoking, status, passive, counts in 'smoke.dat';
inits in 'smoke.ini';
{
#
#      model for 1st table (nonsmokers)
  for (i in 1:4) {
    counts[i]~dpois( lambda[i] );
    log(lambda[i])<- b[1,1] + b[1,2]*status[i] +
b[1,3]*passive[i] + b[1,4]*status[i]*passive[i];
  }
#
#      model for 2nd table (smokers)
  for (i in 5:8) {
    counts[i]~dpois( lambda[i] );
    log(lambda[i])<- b[2,1] + b[2,2]*status[i] +
b[2,3]*passive[i] + b[2,4]*status[i]*passive[i];
  }
#
#      priors
  for (i in 1:2){
    for (j in 1:p){
      b[i,j]~dnorm(0.0, 1.0E-04)
    }
  }
#
#      odds ratios
  or[1]<-exp(b[1,4])
  or[2]<-exp(b[2,4])
}
```

5.1.2 Initial Values (smoke.ini)

```
list(b=c(0,0,0,0,0,0,0,0))
```

5.1.3 Data (smoke.dat)

```
0 1 1 120
0 1 0 111
0 0 1 80
0 0 0 155
1 1 1 161
1 1 0 117
1 0 1 130
1 0 0 124
```

5.2 ANALYSIS 2: COMMON RISK PER SMOKING STATUS

5.2.1 Model (Bug File)

```
model ex5commonOR;
const n=8,
      p=4;
var
  smoking[n],
  status[n],
  passive[n],
  counts[n],
  b[2,p-1],
  ab,
  lambda[n],
  or;
data smoking, status, passive, counts in 'smoke.dat';
inits in 'smoke2.ini';
{
#
#      model for 1st table (nonsmokers)
  for (i in 1:4) {
    counts[i]~dpois( lambda[i] );
    log(lambda[i])<- b[1,1] + b[1,2]*status[i] +
b[1,3]*passive[i] + ab*status[i]*passive[i];
  }
#
#      model for 2nd table (smokers)
  for (i in 5:8) {
    counts[i]~dpois( lambda[i] );
    log(lambda[i])<- b[2,1] + b[2,2]*status[i] +
b[2,3]*passive[i] + ab*status[i]*passive[i];
  }
#
#      priors
  for (i in 1:2){
    for (j in 1:p-1){
      b[i,j]~dnorm(0.0, 1.0E-04);
    }
  }
  ab~dnorm(0.0, 1.0E-04);
#
#      odds ratios
  or<-exp(ab)
}
```

5.2.2 Initial Values (smoke2.ini)

```
list(b=c(0,0,0,0,0,0),ab=0)
```