# MGLM package vignette

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The analysis of multivariate count data arises in numerous fields including genomics, image analysis, text mining, and sports analytics. The multinomial logit model is limiting due to its restrictive mean-variance structure. Moreover, it assumes that counts of different categories are negatively correlated. Models that allow over-dispersion and possess more flexible positive and/or negative correlation structures offer more realism. We implement four models in the R package MGLM: multinomial logit (MN), Dirichlet multinomial (DM), generalized Dirichlet multinomial (GDM), and negative mutinomial (NegMN). Distribution fitting, regression, hypothesis testing, and variable selection are treated in a unified framework.

The simulated data we plan to analyze is multivariate count data with d categories.

## 1 Distribution fitting

The function MGLMfit fits various multivariate discrete distributions and outputs a list with the maximum likelihood estimate (MLE) and relavent statistics.

When fitting distributions, i.e. no covariates involved, MN is a sub-model of DM, and DM is a sub-model of GDM. MGLMfit outputs the p-value of the likelihood ratio test (LRT) for comparing the fitted model with the most commonly used multinomial model. NegMN model does not have a nesting relationship with any of the other three models. Therefore no LRT is performed when fitting a NegMN distribution.

### 1.1 Multinomial (MN)

We first generate data from a multinomial distribution. Note the multinomial parameter (must be positive) supplied to the 'rmn' function is automatically scaled to be a probability vector.

```
> require(MGLM)
> set.seed(123)
> n <- 200
> d <- 4
> alpha <- rep(1,d)
> m <- 50
> Y <- rmn(n, m, alpha)</pre>
```

Multinomial distribution fitting, although trivial, is implemented.

```
> mnFit <- MGLMfit(Y, dist="MN")</pre>
> print(mnFit)
        estimate
alpha_1
          0.2568 0.03089124
alpha_2
          0.2467 0.03048271
alpha_3
          0.2451 0.03041595
alpha_4
          0.2514 0.03067556
Distribution: Multinomial
Log-likelihood: -1457.788
BIC: 2931.471
AIC: 2921.576
LRT test p value:
Iterations:
```

As a comparison, we fit the DM distribution to the same data set. The results indicate that using a more complex model on the multinomial data shows no advantage.

The DM parameter estimates and their standard errors are both big, indicating possible overfitting by the DM model. This is confirmed by the fact that the p-value of the LRT for comparing MN to DM is close to 1.

### 1.2 Dirichlet-multinomial (DM)

DM is a Dirichlet mixture of multinomials and allows over-dispersion. Same as MN model, it assumes that the counts of any two different categories are negatively correlated. We generate the data from the DM model and fit distribution.

```
> set.seed(123)
> n <- 200
> d <- 4
> alpha <- rep(1, d)
> m <- 50
> Y <- rdirm(n, m, alpha)</pre>
```

BIC: 4043.644

AIC: 4030.451

LRT test p value: <0.0001

Iterations: 4

The estimate is very close to the true value with small standard errors. The LRT shows that the DM model is significantly better than the MN model.

### 1.3 Generalized Dirichlet-multinomial (GDM)

GDM model uses d-2 more parameters than the DM model and allows both positive and negative correlations among categories. DM is a sub-model of GDM. Here we fit a GDM model to the above DM sample.

```
> print(gdmFit)

estimate SE
alpha_1 1.1584741 0.12340343
alpha_2 0.9932931 0.43723945
alpha_3 0.8399666 0.10637444
beta_1 3.7068631 0.22641418
beta_2 1.9793891 0.09464476
beta_3 0.7596409 0.08440347
```

> gdmFit <- MGLMfit(Y, dist="GDM")</pre>

Distribution: Generalized Dirichlet Multinomial

Log-likelihood: -2007.559

BIC: 4046.907 AIC: 4027.117

LRT test p value: <0.0001

Iterations: 27

GDM yields a slightly larger log-likelihood value but a larger BIC, suggesting DM as a preferred model. Now we simulate data from GDM and fit the GDM distribution.

```
> set.seed(124)
> n <- 200
> d <- 4
> alpha <- rep(1, d-1)
> beta <- rep(1, d-1)</pre>
```

Distribution: Generalized Dirichlet Multinomial

Log-likelihood: -1820.616

BIC: 3673.021 AIC: 3653.231

LRT test p value: <0.0001

Iterations: 24

### 1.4 Negative multinomial (NegMN)

NegMN model is a multivariate extension to the negative binomial model. It assumes positive correlation among the counts. To generate data from NegMN model and fit the NegMN distribution,

```
> set.seed(1220)
> n <- 100
> d <- 4
> p <- 5
> prob <- rep(0.2, d)
> beta <- 10
> Y <- rnegmn(n, prob, beta)
> negmnFit <- MGLMfit(Y, dist="NegMN")</pre>
> print(negmnFit)
      estimate
p_1 0.1881512 0.009583840
p_2 0.1943109 0.009837429
p_3 0.1915110 0.009722206
p_4 0.1961775 0.009914205
phi 12.3139348 2.266096911
Distribution: Negative Multinomial
Log-likelihood: -1104.579
BIC: 2232.184
AIC: 2219.158
LRT test p value: NA
Iterations: 4
```

## 2 Regression

In regression, the  $n \times p$  covariate matrix X is similar to that used in the glm function. The response should be a  $n \times d$  count matrix. Unlike estimating a parameter vector  $\beta$  in GLM, we need to estimate a parameter matrix B when the responses are multivariate. The dimension of the parameter matrix depends on the model:

```
    MN: p × (d - 1)
    DM: p × d
    GDM: p × 2(d - 1)
    NegMN: p × (d + 1)
```

The GDM model provides the most flexibility, but also requires most parameters. When using function MGLMreg to run regression, we can pick the model by specifying the option dist="MN", "DM", "GDM" or "NegMN".

The rows  $B_{j,\cdot}$  of the parameter matrix correspond to covariates. By default, the function output the Wald test statistics and p-values for testing  $H_0: B_{j,\cdot} = \mathbf{0}$  vs  $H_a: B_{j,\cdot} \neq \mathbf{0}$ . If specifying the option LRT=TRUE, the function also outputs LRT statistics and p-values.

Next we demonstrate that model mis-specification results in failure in hypothesis testing. We simulate response data from the GDM model. Covariates  $X_1$  and  $X_2$  have no effect and  $X_3$ ,  $X_4$ ,  $X_5$  have impact on the response.

```
> set.seed(1234)
> n <- 200
> p <- 5
> d <- 4
> X <- matrix(runif(p*n), n, p)
> alpha <- matrix(c(0.6, 0.8, 1), p, d-1, byrow=TRUE)
> alpha[c(1,2),] <- 0
> Alpha <- exp(X%*%alpha)
> beta <- matrix(c(1.2, 1, 0.6), p, d-1, byrow=TRUE)
> beta[c(1,2),] <- 0
> Beta <- exp(X%*%beta)
> m <- runif(n, min=0, max=25) + 25
> Y <- rgdirm(n, m, Alpha, Beta)</pre>
```

We fit various regression models and test significance of covariates.

### 2.1 Multinomial regression

```
X2 0.5430639 0.4227301 0.2465230

X3 0.3332517 0.5176055 0.2218513

X4 0.3568425 0.4867224 0.5654272

X5 -0.3024545 0.1925076 0.3237132
```

#### Hypothesis test:

wald value Pr(>wald)
X1 24.63244 1.842859e-05
X2 21.99680 6.533133e-05
X3 23.10908 3.832310e-05
X4 25.07475 1.489470e-05
X5 49.37327 1.086326e-10

Distribution: Multinomial Log-likelihood: -2194.448

BIC: 4468.371 AIC: 4418.896 Iterations: 5

The Wald test shows that all predictors are significantly different from 0, including the null predictors  $X_1$  and  $X_2$ .

### 2.2 Dirichlet-multinomial regression

```
> dmReg <- MGLMreg(Y~0+X, dist="DM")
> print(dmReg)
```

Call: MGLMreg(formula = Y ~ 0 + X, dist = "DM")

#### Coefficients:

Col\_1 Col\_2 Col\_3 Col\_4
X1 0.1541366 -0.1182637 -0.1883392 -0.01317229
X2 0.1832642 0.1420338 -0.1833943 -0.33388600
X3 1.1431456 1.2548276 1.0926352 0.81125874
X4 0.3927028 0.5454214 0.5900146 0.13113218
X5 0.2263497 0.6601081 0.9395988 0.48703415

#### Hypothesis test:

wald value Pr(>wald)
X1 3.349794 5.010817e-01
X2 7.845339 9.741075e-02
X3 25.497386 3.995529e-05
X4 8.735121 6.807217e-02
X5 23.136042 1.189431e-04

Distribution: Dirichlet Multinomial

Log-likelihood: -1683.961

BIC: 3473.889 AIC: 3407.922 Iterations: 7 Again, Wald test declares all predictors as significant.

### Generalized Dirichlet-multinomial Regression

```
> gdmReg <- MGLMreg(Y~0+X, dist="GDM")</pre>
> print(gdmReg)
```

Call: MGLMreg(formula = Y ~ 0 + X, dist = "GDM")

#### Coefficients:

```
alpha_Col_1 alpha_Col_2 alpha_Col_3 beta_Col_1 beta_Col_2 beta_Col_3
  -0.2839174 0.19050322 0.31570552 -0.4002027 0.6846506 0.4675918
X2 -0.2091710 0.39554111 0.01442559 -0.5082543 0.5526452 -0.1714096
   1.0901404 1.24378465 1.17178640 1.3049550 1.5375262 0.9160329
    0.2968186 \quad 0.40533348 \quad 0.80908676 \quad 0.4878379 \quad 0.5736954 \quad 0.3058887
X4
    X5
```

#### Hypothesis test:

```
wald value
                Pr(>wald)
Х1
    9.424379 1.510802e-01
    4.865683 5.611523e-01
X3 26.828718 1.559064e-04
X4 12.607034 4.971848e-02
X5 38.637877 8.427803e-07
```

Distribution: Generalized Dirichlet Multinomial

Log-likelihood: -1676.399

BIC: 3511.748 AIC: 3412.798 Iterations: 21

When using the correct model, Wald test is able to differentiate the null effects from the significant ones. GDM regression yields the highest log-likelihood and smallest BIC.

### Negative multinomial regression

```
> negReg <- MGLMreg(Y~0+X, dist="NegMN", regBeta=FALSE)
> print(negReg)
Call: MGLMreg(formula = Y ~ 0 + X, dist = "NegMN", regBeta = FALSE)
```

### Coefficients: \$alpha

```
Col_1
```

```
Col_2
                          Col_3
                                    Col 4
X1 0.24360582 -0.21636792 -0.15652499 -0.03137138
X2 0.06189622 -0.05588488 -0.23263498 -0.47977959
X4 -0.17618094 -0.04382845 0.03478118 -0.53012642
X5 -0.60797439 -0.11582939 0.01293699 -0.31585689
```

#### \$phi

13.77531

### Hypothesis test:

```
wald value Pr(>wald)
X1 24.99903 5.033252e-05
X2 24.90077 5.267448e-05
X3 29.32830 6.704198e-06
X4 28.18693 1.143075e-05
X5 60.50179 2.275444e-12
```

Distribution: Negative Multinomial

Log-likelihood: -2908.896

BIC: 5929.056 AIC: 5859.792 Iterations: 15

Again, the Wald test declares all preditors to be significant.

#### 2.5 Prediction

We can use the fitted model to make prediction. The output of the prediction function is the probabilities of the d categories. This helps answer questions such as whether certain features increase the probability of observing category j. Take the fitted GDM model as an example:

### 3 Sparse regression

Regularization is an important tool for model selection and improving the risk property of the estimates. In the package, we implemented three types of penalties on the paramter matrix B:

- $\bullet$  select by entries
- select by rows
- select by rank

The function MGLMtune finds the optimal tuning parameter with smallest BIC and outputs the estimate using the chosen tuning parameter. The output from MGLMtune is a list containing the solution path and the final estimate. Users can either provide a vector of tuning parameters with option lambdas

or specify the number of grid points via option ngridpt and let the function decide the default tuning parameters. The function MGLMsparsereg computes the regularized estimate at a given tuning parameter value lambda.

We generate the data from the DM model, with row sparsity, and show how each penalty type works.

```
> set.seed(118)
> n <- 100
> p <- 10
> d <- 5
> m <- rbinom(n, 200, 0.8)
> X <- matrix(rnorm(n*p),n, p)
> alpha <- matrix(0, p, d)
> alpha[c(1,3, 5), ] <- 1
> Alpha <- exp(X%*%alpha)
> Y <- rdirm(size=m, alpha=Alpha)</pre>
```

### 3.1 Select by entries

Degrees of freedom: 31 Lambda: 4.858822

Iterations: 41

### 3.2 Select by rows

Since the rows of the parameter matrix correspond to predictors, selecting by rows performs variable selection at the predictor level.

### 3.3 Select by singular values

Nuclear norm regularization encourages low rank in the regularized estimate.

Distribution: Dirichlet Multinomial

Log-likelihood: -1492.063

BIC: 3070.422 AIC: 3021.604

Degrees of freedom: 18.7391

Lambda: 37.53776
Iterations: 32