glmpathcr: An R Package for Ordinal Response Prediction in High-dimensional Data Settings

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Abstract

This paper describes an R package, **glmpathcr**, that provides a function for fitting a penalized continuation ratio model when interest lies in predicting an ordinal response. The function, **glmpathcr** uses the coordinate descent fitting algorithm as implemented in **glmpath** and described by (Park and Hastie 2007a). Methods for extracting all estimated coefficients, extracting non-zero coefficient estimates, obtaining the predicted class, and obtaining the class-specific fitted probabilities have been implemented. Additionally, generic methods from **glmpath** including **summary**, **print**, and **plot** can be applied to a **glmpathcr** object.

Keywords: ordinal response, penalized models, LASSO, L₁ constraint, R.

1. Introduction

High-throughput genomic experiments are frequently conducted for the purpose of examining whether genes are predictive of or significantly associated with phenotype. In many biomedical settings where histopathological or health status data are collected, phenotypic variables are recorded on an ordinal scale. Nevertheless, most often investigators neglect the ordinality of the phenotypic data and rather dichotomize the ordinal class than apply statistical methods suitable for two-class comparisons and predictions. This tendency to analyze ordinal data using dichotomous class methodologies may be due to the lack of available statistical methods and software for modeling an ordinal response in the presence of a high-dimensional covariate space. The approach of collapsing ordinal categories may neglect important information in the study (Armstrong and Sloan 1989).

A variety of statistical modeling procedures, namely, proportional odds, adjacent category, stereotype logit, and continuation ratio models can be used to predict an ordinal response. In this paper, we focus attention to the continuation ratio model because its likelihood can be easily re-expressed such that existing software can be readily adapted and used for model fitting. Suppose for each observation, i = 1, ..., n, the response Y_i belongs to one ordinal class k = 1, ..., K and \mathbf{x}_i represents a p-length vector of covariates. The backward formulation of the continuation ratio models the logit as

$$logit (P(Y = k | Y \le k, \mathbf{X} = \mathbf{x})) = \alpha_k + \beta_k^T \mathbf{x}$$
(1)

whereas the forward formulation models the logit as

$$logit (P(Y = k | Y \ge k, \mathbf{X} = \mathbf{x})) = \alpha_k + \boldsymbol{\beta}_k^T \mathbf{x}.$$
 (2)

Rather than describe both formulations in detail, here we present the backward formulation, which is commonly used when progression through disease states from none, mild, moderate, severe is represented by increasing integer values, and interest lies in estimating the odds of more severe disease compared to less severe disease (Bender and Benner 2000). Therefore for i = 1, ..., n we can construct a vector \mathbf{y}_i from Y_i to represent ordinal class membership, such that $\mathbf{y}_i = (y_{i1}, y_{i2}, ..., y_{iK})^T$, where $y_{ik} = 1$ if the response is in category k and 0 otherwise, so that $n_i = \sum_{k=1}^K y_{ik} = 1$. Using the logit link, the equation representing the conditional probability for class k is

$$\delta_k(\mathbf{x}) = P(Y = k | Y \le k, \mathbf{X} = \mathbf{x}) = \frac{\exp(\alpha_k + \boldsymbol{\beta}^T \mathbf{x})}{1 + \exp(\alpha_k + \boldsymbol{\beta}^T \mathbf{x})}.$$
 (3)

The likelihood for the continuation ratio model is then the product of conditionally independent binomial terms (Cox 1975), which is given by

$$L(\boldsymbol{\beta}|\mathbf{y},\mathbf{x}) = \prod_{i=1}^{n} \delta_2^{y_{i2}} (1 - \delta_2)^{1 - \sum_{k=2}^{K} y_{ik}} \times \dots \times \delta_K^{y_{iK}} (1 - \delta_K)^{1 - y_{iK}}$$
(4)

where here we have simplified our notation by not explicitly including the dependence of the conditional probability δ_k on \mathbf{x} . Further, simplifying our notation to let $\boldsymbol{\beta}$ represent the vector containing both the thresholds $(\alpha_2, \ldots, \alpha_K)$ and the log odds $(\beta_1, \ldots, \beta_p)$ for all K-1 logits, the full parameter vector is

$$\boldsymbol{\beta} = (\alpha_2, \beta_{21}, \beta_{22}, \dots, \beta_{2p}, \dots, \alpha_K, \beta_{K,1}, \beta_{K,2}, \dots, \beta_{K,p})^T$$
(5)

which is of length (K-1)(p+1). As can be seen from equation 4, the likelihood can be factored into K-1 independent likelihoods, so that maximization of the independent likelihoods will lead to an overall maximum likelihood estimate for all terms in the model (Bender and Benner 2000). A model consisting of K-1 different β vectors may be overparameterized so to simplify, one commonly fits a constrained continuation model, which includes the K-1 thresholds $(\alpha_2, \ldots, \alpha_K)$ and one common set of p slope parameters, $(\beta_1, \ldots, \beta_p)$. To fit a constrained continuation ratio model, the original dataset can be restructured by forming K-1 subsets, where for classes $k=2,\ldots,K$, the subset contains those observations in the original dataset up to class k. Additionally, for the k^{th} subset, the outcome is dichotomized as y=1 if the ordinal class is k and y=0 otherwise. Furthermore, an indicator is constructed for each subset representing subset membership. Thereafter the K-1 subsets are appended to form the restructured dataset, which represents the K-1 conditionally independent datasets in equation 4. Applying a logistic regression model to this restructured dataset yields an L_1 constrained continuation ratio model.

2. Penalized Models

For datasets where the number of covariates p exceeds the sample size n, the backwards stepwise procedure cannot be undertaken. Furthermore, for any problem using a forward selection procedure the discrete variable inclusion process can exhibit high variance. Moreover, for high-dimensional covariate spaces, the best subset procedure is computationally prohibitive. Two penalized methods, ridge and L_1 penalization, places a penalty on a function of the coefficient estimates, thereby permitting a model fit even for high-dimensional

data Tibshirani (1996, 1997). A generalization of these penalized models can be expressed as,

$$\tilde{\beta} = \arg\min_{\beta} \left(\sum_{i=1}^{n} (y_i - \beta_0 - \sum_{j=1}^{p} x_{ij} \beta_j)^2 + \lambda \sum_{j=1}^{p} |\beta_j|^q \right)$$
 (6)

for $q \geq 0$. When q = 1 we have the an L₁ penalized model, when q = 2 we have ridge regression. Values of $q \in (1,2)$ provide a compromise between the L₁ and ridge penalized models. Because when q > 1 coefficients are no longer set exactly equal to 0, the elastic net penalty was introduced

$$\lambda \sum_{j=1}^{p} (\alpha \beta_j^2 + (1-\alpha)|\beta_j|). \tag{7}$$

3. Implementation

The glmpather package was written in the R programming environment (R Development Core Team 2009) and depends on the glmpath package (Park and Hastie 2007b). Similar to the **Design** package which includes a function cr.setup for restructuring a dataset for fitting a forward continuation ratio model, in this package the model is fit by restructuring the dataset then passing the restructured dataset to a penalized logistic regression fitting function. However, unlike cr.setup which produces an object of class list from which the response and restructured independent variables are extracted and passed to a model fitting algorithm, in the glmpather package the restructuring functions are transparent to the user. Specifically, the **glmpathcr** package fits either a forward or backward (default) penalized constrained continuation ratio model by specification of method="forward" in the glmpather call. The glmpather function restructures the dataset to represent the K-1conditionally independent likelihoods and then fits the penalized continuation ratio model using the glmpath framework. Therefore, the predictor-corrector fitting procedure used by the glmpath function in the glmpath package is used in fitting the penalized continuation ratio model when invoking glmpather. This allows fitting a penalized model for situations where the number of covariates p exceed the sample size n. In addition, methods for extracting the best fitting model from the path using AIC and BIC criteria, obtaining predicted class and fitted class probabilities, and returning coefficient estimates were written in addition to adapting the print, summary, and plot methods from glmpath for a glmpather object.

4. Example

The glmpather package includes a filtered microarray dataset diabetes in which asymptomatic males not previously diagnosed with Type II diabetes were enrolled and subsequently were cross-classified as either normal controls (N=8), having impaired fasting glucose (N=7), or as Type II diabetics (N=9) based on a fasting glucose intolerance test. From the code below we can see that the classification variable is stored as y in the first column of the diabetes data.frame; all subsequent columns are the 11,066 Illumina probes having no negative expression values. In fitting the model we can extract the covariates into an object x and the ordinal outcome into the object y. The code for fitting a backward (default) continuation ratio model is given by

As with glmpath model objects, methods such as summary and plot can be applied to glmpathcr model objects, which are helpful for selecting the step at which to select the final model from the solution path.

> summary(fit)

> fit <- glmpathcr(x,y)</pre>

```
Df
               Deviance
                            AIC
                                      BIC
Step 1
         3 5.248273e+01 58.48273 62.01690
Step 2
         4 2.334393e+01 33.34393 39.23419
         5 2.047337e+01 30.47337 36.36364
Step 5
Step 7
        6 2.013650e+01 32.13650 39.20482
Step 10 7 1.587273e+01 29.87273 38.11911
Step 14
       8 7.939954e+00 23.93995 33.36438
Step 17
       9 7.530910e+00 25.53091 36.13339
Step 19 10 7.327910e+00 27.32791 39.10845
Step 21 11 7.055827e+00 29.05583 42.01442
Step 24 12 6.608876e+00 30.60888 44.74552
Step 28 13 3.583362e+00 29.58336 44.89806
Step 31 14 3.176651e+00 31.17665 47.66940
Step 34 15 2.491539e+00 32.49154 50.16235
Step 38 16 1.803196e+00 33.80320 52.65206
Step 41 17 1.718769e+00 35.71877 55.74568
Step 46 18 4.362168e-01 36.43622 57.64119
Step 48 18 9.654998e-02 36.09655 57.30152
Step 54 18 1.079936e-02 38.01080 60.39382
Step 55 19 1.014368e-02 38.01014 60.39317
```

```
Step 56 19 6.701543e-03 38.00671 60.38973
Step 57 19 6.698089e-03 40.00670 63.56777
Step 58 20 6.638869e-03 40.00664 63.56772
Step 61 21 4.759556e-03 42.00476
                                  66.74389
Step 62 22 4.746810e-03 44.00475 69.92193
Step 65 23 4.429264e-03 46.00443 73.09967
Step 67 24 4.098176e-03 48.00410 76.27739
Step 68 24 3.733398e-03 48.00373 76.27703
Step 69 23 3.581801e-03 46.00358 73.09882
Step 70 22 3.121123e-03 44.00312 69.92031
Step 71 21 3.063389e-03 42.00306 66.74219
Step 72 20 3.021411e-03 40.00302 63.56410
Step 73 19 2.718144e-03 38.00272 60.38574
Step 74 18 2.470063e-03 36.00247 57.20744
Step 75 17 2.349117e-03 34.00235 54.02926
Step 77 17 2.208164e-03 34.00221 54.02912
Step 78 18 2.104337e-03 36.00210 57.20707
Step 79 19 2.089033e-03 38.00209 60.38511
Step 80 20 2.057374e-03 40.00206 63.56313
Step 82 21 1.984043e-03 42.00198 66.74111
Step 83 22 1.952043e-03 44.00195
                                  69.91914
Step 84 22 1.943284e-03 44.00194 69.91913
Step 85 22 1.666121e-03 46.00167
                                  73.09690
Step 87 23 1.475796e-03 46.00148 73.09671
Step 89 24 1.419090e-03 48.00142 76.27471
Step 90 25 1.392713e-03 50.00139 79.45274
Step 91 25 1.372528e-03 50.00137 79.45272
Step 92 25 1.318666e-03 50.00132 79.45266
Step 94 26 1.248220e-03 52.00125 82.63065
Step 95 27 1.221184e-03 56.00122 88.98673
Step 96 28 1.167649e-03 56.00117 88.98667
Step 97 28 1.161679e-03 56.00116 88.98667
Step 98 27 1.134997e-03 54.00113 85.80859
Step 99 27 1.130143e-03 54.00113 85.80858
Step 100 27 1.121460e-03 54.00112 85.80857
Step 101 27 1.107686e-03 54.00111 85.80856
Step 102 28 1.102366e-03 56.00110 88.98661
Step 104 29 1.043914e-03 58.00104 92.16460
Step 105 30 1.008973e-03 60.00101 95.34262
Step 106 31 9.025276e-04 62.00090 98.52057
Step 107 32 8.992031e-04 64.00090 101.69862
Step 109 33 8.588539e-04 66.00086 104.87664
Step 110 34 8.523875e-04 68.00085 108.05468
Step 111 35 8.410635e-04 70.00084 111.23273
Step 112 35 8.397137e-04 70.00084 111.23272
Step 114 35 7.787702e-04 70.00078 111.23266
Step 115 36 7.567549e-04 72.00076 114.41069
```

Step 117 37 6.878607e-04 74.00069 117.58868 Step 118 38 6.849152e-04 76.00068 120.76673

> plot(fit, xvar = "step", type = "bic")

BIC

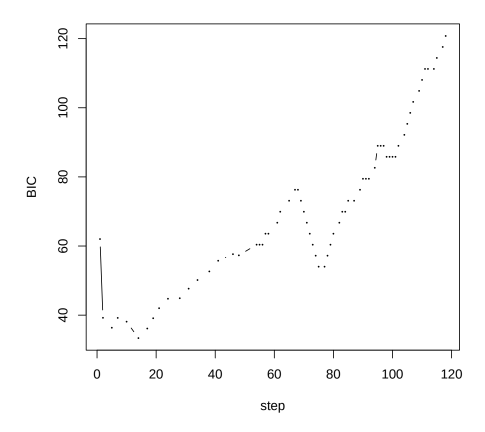


Figure 1: Plot of regularization path for glmpather object using simulated dataset, data.

Note that when plotting, the horizontal axis can be norm, lambda, or step, however extractor functions for glmpathcr generally require the step to be selected, so we have selected xvar = "step" in this example. The vertical axis can be coefficients, aic or bic. As one can see, there is a multitude of models fit from one call to glmpathcr. To faciliate extraction of best fitting models using commonly used criterion, the model.select function can be used. The model.select function extracts the best fitting model from the solution path, where the which parameter allows one to select either AIC or by default, BIC.

```
> BIC.step <- model.select(fit)
> BIC.step
```

```
> AIC.step <- model.select(fit)
> AIC.step
```

[1] 14

In this example, the minimum BIC corresponds to a 8 degree of freedom model.

The coef function returns all estimated coefficients for a glmpathcr fitted model, where the model selected is indicated by step number, s. The nonzero.coef function returns only those non-zero coefficient estimates for a selected model.

```
> coefficients<-coef(fit, s=BIC.step)
> sum(coefficients!=0)
[1] 8
```

> nonzero.coef(fit, s=BIC.step)

```
Intercept ILMN_1705116 ILMN_1733757 ILMN_1758311 ILMN_1759232
18.3204861060 0.0334955980 -0.0003751318 0.0022607450 -0.0347775942
ILMN_2100437 cp1 cp2
-0.0001679020 2.0164341452 -2.0164341455
```

Note that the glmpathcr function fits a penalized constrained continuation ratio model; therefore for K classes, there will be K-1 intercepts representing the cutpoints between adjacent classes. In this package, the nomenclature for these cutpoints is to use "cpk" where $k=1,\ldots,K-1$. In this dataset, K=3 so the intercepts are cp1 and cp2 with Intercept being an offset. The probe having the largest absolute coefficient estimate, ILMN 1759232, corresponds to the insulin receptor substrate 1 (IRS1) gene which is biologically meaningful. Continuation ratio models predicts conditional probabilities so a new method to extract the fitted probabilities and predicted class was created. The predict and fitted functions are equivalent, and return either the predicted class or the fitted probabilities from the penalized continuation ratio model for a glmpather object. The user is required to supply the fitted glmpather model object, a data matrix news that is either the same as the training data or an independent dataset having the same number and order of covariates as the training data, a vector newy that provides the class labels of the ordinal response. These functions extract the fitted values for the best fitting model using the BIC criteria by default, which can be changed to extracting the best fitting AIC model by supplying which="AIC". By default, the predicted class is output. If one desired the fitted class-specific probabilities from the model, the type="probs" argument should be supplied.

```
> pred <- predict(fit)
> table(pred, y)
```

	У					
pred	control	impaired	fasting	glucose	type 2	diabetes
control	8			0		0
impaired fasting glucose	0			7		0
type 2 diabetes	0			0		9

```
> pred <- predict(fit, type="probs")
> pred
```

	control	impaired	fasting glucose	type 2 diabetes
[1,]	8.178419e-01	_	0.17830887	0.0038492509
[2,]	9.239276e-01		0.07464256	0.0014297981
[3,]	8.947141e-01		0.10324488	0.0020410069
[4,]	9.530268e-01		0.04611633	0.0008568896
[5,]	8.835483e-01		0.11416679	0.0022848811
[6,]	9.070107e-01		0.09121017	0.0017791192
[7,]	8.672968e-01		0.13005260	0.0026506120
[8,]	8.784051e-01		0.11919565	0.0023992189
[9,]	7.140319e-02		0.76841557	0.1601812429
[10,]	2.703789e-01		0.68655530	0.0430657810
[11,]	5.059698e-02		0.74290859	0.2064944315
[12,]	9.145857e-02		0.77758826	0.1309531692
[13,]	5.568355e-02		0.75132840	0.1929880471
[14,]	8.375911e-02		0.77530691	0.1409339750
[15,]	1.423227e-01		0.77016247	0.0875147921
[16,]	3.278482e-04		0.12689759	0.8727745668
[17,]	4.987756e-04		0.15400425	0.8454969777
[18,]	1.249965e-04		0.08046247	0.9194125349
[19,]	1.588154e-04		0.09020377	0.9096374103
[20,]	1.239401e-04		0.08013625	0.9197398128
[21,]	1.891746e-04		0.09801511	0.9017957106
[22,]	8.823833e-04		0.19912988	0.7999877347
[23,]	6.318904e-05		0.05792144	0.9420153755
[24,]	1.585017e-04		0.09011893	0.9097225676

For illustrative purposes, a forward continuation ratio model can be fit using the syntax

```
> fit <- glmpathcr(x, y, method="forward")</pre>
```

As before, the parameter estimates corresponding to the model attaining the minimum BIC can be extracted using the following code.

2	<i>T</i>					
pred	${\tt control}$	${\tt impaired}$	fasting	glucose	type	$2 \ {\tt diabetes}$
control	8			1		0
impaired fasting glucose	0			6		0
type 2 diabetes	0			0		9

Summary

Herein we have described the **glmpathcr** package which works in conjunction with the **glmpath** package in the R programming environment. The package provides methods for fitting either a forward or backward penalized continuation ratio model. Moreover, the likelihood-based penalized continuation ratios models have been demonstrated to have good performance when applied to microarray gene expression datasets (Archer and Williams 2012) in comparison to corresponding penalized Bayesian continuation ratio models (Kiiveri 2008). A similar package, **glmnetcr**, which uses the **glmnet** fitting algorithm for fitting a penalized constrained continuation ratio model has also been developed and is available for download from the Comprehensive R Archive Network. Functions for extracting coefficients, extracting non-zero coefficients, and obtaining fitted probabilities and predicted class in the **glmnetcr** package follow those in **glmpathcr** and both packages have similar performance (Archer and Williams 2012). Therefore either the **glmpathcr** or **glmnetcr** package should be helpful when predicting an ordinal response for datasets where the number of covariates exceeds the number of available samples.

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