Uncovering gene regulatory relationships from knockdown expression data using BayesKnockdown

William Chad Young, Ka Yee Yeung, and Adrian E. Raftery Department of Statistics (WCY and AER) and Institute of Technology (KYY) University of Washington

This document illustrates the use of the BayesKnockdown R package to calculate posterior probabilities of relationships between a single predictor and multiple potential targets. The package was developed specifically for gene expression datasets in the form of knockdown experiments, but can be applied more generally to other over-expression data and to infer differential expression.

1 Posterior Probabilities

Given a predictor x and a set of possible targets y, the BayesKnockdown function can be invoked to estimate the posterior probabilities of a relationship between x and each individual target in y [2]. The BayesKnockdown function allows specification of a prior probability of regulation via the prior argument, and it can be a constant for all targets or unique to each target. This is useful particularly when an informative prior is available to incorporate additional knowledge. The prior is set to 0.5 by default, which corresponds to an uninformative prior.

Additionally, the method allows specification of Zellner's g-prior via the g argument [3]. The g-prior specifies the expected strength of the signal relative to noise, with larger values corresponding to a larger expected signal. It is recommended that g be set to a value between 1 and the number of observations in the data. The default value is \sqrt{n} , which we have found to be a good compromise between the extremes.

1.1 Simulated Data Example

As a simple example of using the BayesKnockdown function, we generate random data for the knockdown gene as well as the potential targets. We then introduce a relationship between the knockdown gene and target number 3. The BayesKnockdown function takes this data and produces the posterior probability of a relationship between x and each target. Figure 1 shows the posterior probabilities calculated for each target.

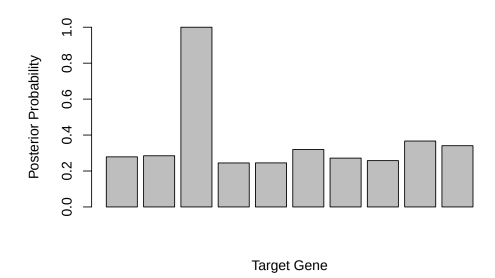


Figure 1: Bar plot showing the posterior probabilities of a relationship between the knockdown gene and each target in simulated data. Gene 3 is the only true relationship.

1.2 Knockdown Data Example

A more realistic example uses data from the National Institute of Health (NIH) Library of Integrated Network-based Cellular Signatures (LINCS) program (http://lincsproject.org) [1]. The aim of this program is to generate genetic and molecular signatures of cells in response to various perturbations. To support this endeavor, many large datasets have been made available, including proteomic and imaging data.

The LINCS L1000 data capture gene expression levels of 1,000 genes in human cell lines under a variety of conditions. The lincs.kd data is a 21 by 27 matrix containing data from knockdown experiments targeting gene PPARG in cell line A375. Cell line A375 is a human skin melanoma cell line with over 100,000 experiments in the L1000 data. The first row is the expression levels of PPARG in the 27 experiments targeting PPARG for knockdown, while the other 20 rows are a subset of the measured genes in the same experiments. The data have been normalized to account for differences in the experimental settings, as described in [2]. The full LINCS L1000 data is available at http://lincscloud.org.

Given the L1000 data, the BayesKnockdown function can be invoked to calculate the posterior probabilities of a relationship between gene PPARG and the other genes in the dataset. In this case, we specify a prior probability of 0.0005, reflecting the belief that there are very few relationships relative to the total possible number. Figure 2 shows the range of values returned for the different target genes.

```
> data(lincs.kd);
> kdResult <- BayesKnockdown(lincs.kd[1,], lincs.kd[-1,], prior=0.0005);</pre>
> kdResult;
        ATF1
                 SERPINE1
                                  CEBPA
                                                MUC1
                                                              EZH2
                                                                          SNX13
0.9959445881 0.0271544446 0.0007644977 0.0090443118 0.9637199199 0.0504674678
      ELOVL6
                    CASC3
                                 MRPL12
                                               KIF2C
                                                             BCL7B
0.0002019284 0.8210955027 0.0048439740 0.9986842921 0.9997559484 0.0005646142
        NET1
                   ATP1B1
                                  H2AFV
                                             TIMM17B
                                                            ZNF586
0.0002563962 0.0046399699 0.0222102583 0.0004304130 0.0153763269 0.0003694639
       CDK19
                   SFMBT1
0.9990571273 0.9053986408
> barplot(kdResult, names.arg="", xlab="Target Gene",
          ylab="Posterior Probability", ylim=c(0,1));
```

1.3 ExpressionSet Example

The BayesKnockdown.es function allows calculation of posterior probabilities using an ExpressionSet object from the bioBase library. The function works similarly to the BayesKnockdown function, except that one of the features of the ExpressionSet is identified to be the predictor variable, and all other features are used as response variables.

```
> library(Biobase);
> data(sample.ExpressionSet);
```

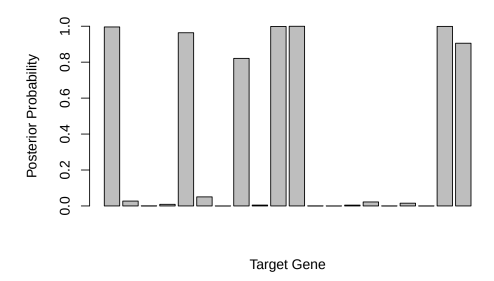


Figure 2: Bar plot showing the posterior probabilities of a relationship between the knockdown gene PPARG and each target in LINCS L1000 data.

```
> subset <- sample.ExpressionSet[1:10,];</pre>
> BayesKnockdown.es(subset, "AFFX-MurIL10_at");
 AFFX-MurIL2_at
                 AFFX-MurIL4_at
                                  AFFX-MurFAS_at
                                                   AFFX-BioB-5_at
                                                                    AFFX-BioB-M_at
      0.3418659
                       0.3361832
                                        0.7430095
                                                         0.3940327
                                                                         0.4110827
 AFFX-BioB-3_at
                 AFFX-BioC-5_at
                                  AFFX-BioC-3_at AFFX-BioDn-5_at
                                        0.3181790
                                                         0.7516404
      0.6523990
                       0.3267071
```

2 2-Class Data

The BayesKnockdown.diffExp function tests for differential expression in a set of variables between two experimental conditions. In gene expression data, this often takes the form of comparing the effects of a drug perturbation compared to a baseline. Of interest is the set of genes which show different expression levels between the two conditions. The BayesKnockdown.diffExp function takes two matrices of observations for a set of variables, one matrix for each condition, and gives posterior probabilities that the variables are different between the two conditions.

As an example, we generate two random datasets for 10 genes, corresponding to different experimental conditions. The first has 25 observations and the second has 30. We add an offset for gene 3 in the second dataset, reflecting a change of expression between the two conditions. The BayesKnockdown.diffExp function produces posterior probabilities for each gene reflecting how likely they are to be expressed differently between the two conditions.

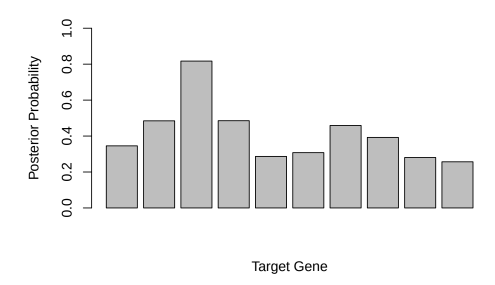


Figure 3: Bar plot showing the posterior probabilities that each gene is differentially expressed between two conditions. Gene 3 is the only gene which is actually differentially expressed.

Figure 3 shows the posterior probabilities that each gene is differentially expressed between the two conditions.

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