Hotspots and Causal Inference For Yeast Data

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October 24, 2012

Here we reproduce the analysis of the budding yeast genetical genomics data-set presented in Chaibub Neto et al. (2012). The data represents a cross of a standard yeast laboratory strain, and a wild isolate from a California vineyard (Brem and Kruglyak 2005). It consists of expression measurements on 5,740 transcripts measured on 112 segregant strains with dense genotype data on 2,956 markers. Processing of the expression measurements raw data was done as described in Brem and Kruglyak (2005), with an additional step of converting the processed measurements to normal quantiles by the transformation $\Phi^{-1}[(r_i - 0.5)/112]$, where Φ is the standard normal cumulative density function, and the r_i are the ranks.

The data were provided by Rachel Brem and further edited by Jun Zhu and Bin Zhang (formerly of Sage Bionetworks). Elias Chaibub Neto and Brian Yandell have organized the data and analysis into this R statistical package, qtlyeast, which is available through GITHUB.

We first load the yeast cross object (yeast.orf), and compute the conditional genotype probabilities using Haldane's map function, genotype error rate of 0.0001, and setting the maximum distance between positions at which genotype probabilities were calculated to 2cM.

```
> library(qtlhot)
> library(qtlyeast)
> ## data(yeast.orf) is loaded lazily.
> yeast.orf <- calc.genoprob(yeast.orf, step = 2)</pre>
```

The following command does an genome scan for QTL using R/qtl for all the traits using Haley-Knott regression (Haley and Knott 1992).

```
> scan.orf <- scanone(yeast.orf, pheno.col = seq(nphe(yeast.orf)), method = "hk")
```

To save space, we work with only the genome regions that are above the single trait LOD threshold and within 1.5 LOD of the maximum per chromosome. We do this after we determine the permutation LOD threshold below.

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1 Hotspot Inference

Plan of action: 1. Find Churchill-Doerge 5% LOD threshold 2. Determine hotspot counts relative to LOD threshold (Jansen method) 3. conduct permutation test (using CHTC) 4. report Jansen and Chaibub-Neto results 5. identify hotspots.

1.1 Churchill-Doerge LOD threshold

Since we are using normal scores on the traits, we need only conduct permutation threshold calculation with a normal response. Here we create one trait and then do 1000 permutations. We have saved this as perm.orf.

```
> cross <- yeast.orf
> cross$pheno <- data.frame(norm = rnorm(nind(cross)))
> set.seed(12345)
> perm.orf <- scanone(cross, method = "hk", n.perm = 1000)
> ## data(perm.orf) is loaded lazily.
> summary(perm.orf)

LOD thresholds (1000 permutations)
    lod
5% 3.48
10% 3.08
> lod.thr <- c(summary(perm.orf, alpha = 0.05))
Now we save only the high lods of the scan.orf object to save space.</pre>
```

> highlod.orf <- highlod(scan.orf, lod.thr = lod.thr, drop.lod = 1.5)

This takes considerable time, so we have actually saved the completed scans as object scan.orf. However, the scan.orf object is 203Mb, so we don't keep it in the package. Instead we have saved highlod.orf.

> ## data(highlod.orf) is loaded lazily.

1.2 Hotspots for Yeast Data above LOD threshold

Now we show the hotspots. We can get summary and plot from highlod.orf, but it is sometimes more helpful to first turn it into a hotsize object. We use an arbitrary threshold of 80 traits per hotspot, which is passed along to scanone summary and plot methods, to get some handle on hotspots.

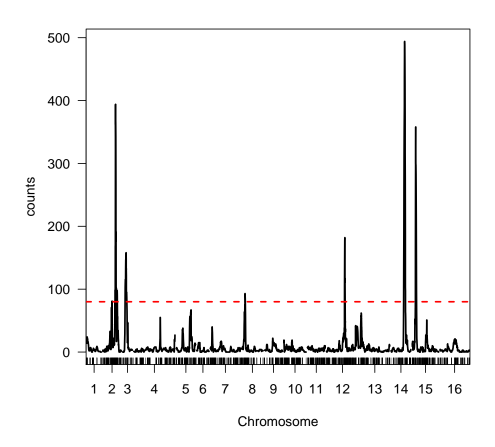
```
> hotsize.orf <- hotsize(highlod.orf, lod.thr = lod.thr)
> summary(hotsize.orf, threshold = 80)
```

hotsize elements: chr pos max.N

LOD threshold: 3.475609

	chr	pos	${\tt max.N}$
YBR154C_chr2@548401	2	224.1	394
c3.loc60	3	60.0	158
NHR001C.4_chr8@111686	8	58.5	93
c12.loc324	12	324.0	182
c14.loc240	14	240.0	494
gOL02_chr15@170945	15	61.1	358

> plot(hotsize.orf)



This shows hotspots, but there is no way yet to assess their significance. To do that, we must run some further permutations across all the traits together, preserving their correlation structure. This takes even more time, so we will do it offline and show the results.

> abline(h = 80, 1wd = 2, col = "red", 1ty = 2)

2 Causal Inference

Current efforts in systems genetics have focused on the development of statistical approaches that aim to disentangle causal relationships among molecular phenotypes in segregating populations. Model selection criterions, such as the AIC and BIC, have been widely used for this purpose, in spite of being unable to quantify the uncertainty associated with the model selection call. We illustrate analysis of the Brem and Kruglyak (2005 PNAS) data using software implemented in R/qtlhot.

In order to evaluate the precision of the causal predictions made by the methods we used validated causal relationships extracted from a data-base of 247 knock-out experiments in yeast (Hughes et al. 2000, Zhu et al. 2008). In each of these experiments, one gene was knocked-out, and the expression levels of the remainder genes in control and knocked-out strains were interrogated for differential expression. The set of differentially expressed genes form the knock-out signature (ko-signature) of the knocked-out gene (ko-gene), and show direct evidence of a causal effect of the ko-gene on the ko-signature genes.

Next, we load a yeast annotation data.frame, yeast.annot, that provides the orf, gene symbol, and chromosome location (in both Mb and cM) of each one of the 5,740 transcripts. (This information will be needed to determine which ko-genes show significant QTLs.) Next, we load a yeast annotation (derived from the YEAST R package) data.frame, yeast.annot, that provides ORFs, gene names, chromosome, and position in Mb and cM.

```
> ## data(yeast.annot) is loaded lazily.
> head(yeast.annot)
```

```
gene chr
                          Mb.pos
                                    cM.pos
         orf
3952 YAL001C
              TFC3
                      1 0.151168 102.4066
3951 YAL002W
              VPS8
                      1 0.143709 101.3745
3950 YAL003W
              EFB1
                      1 0.142176 101.1623
1330 YAL005C
                      1 0.141433 101.0595
              SSA1
3934 YAL007C
              ERP2
                      1 0.138347 100.1245
3933 YALOO8W FUN14
                      1 0.136916 100.1245
```

Next, we load the list of ko-signatures derived from the knock-out experiments in Hughes et al. (2000) and Zhu et al. (2008). We show below the first knock-out signature.

```
> ## data(ko.list) is loaded lazily.
> length(ko.list)
[1] 247
> ko.list[[1]]
 [1] "YAR073W"
                  "YBL013W"
                               "YBL032W"
                                            "YBL042C"
                                                                      "YBL064C"
                                                         "YBL054W"
                                                                      "YBR186W"
 [7] "YBR013C"
                  "YBR054W"
                               "YBR072W"
                                            "YBR126C"
                                                         "YBR155W"
[13] "YCL030C"
                               "YDL234C"
                                            "YDL244W"
                  "YDL038C"
                                                         "YDR001C"
                                                                      "YDR018C"
```

```
"YDR077W"
                               "YDR085C"
[19]
     "YDR055W"
                                             "YDR399W"
                                                          "YDR518W"
                                                                       "YDR533C"
[25] "YDR534C"
                  "YER055C"
                               "YER062C"
                                             "YFL014W"
                                                          "YFL030W"
                                                                       "YFL058W"
[31] "YGL156W"
                               "YGL187C"
                                            "YGL234W"
                                                          "YGR032W"
                                                                       "YGR043C"
                  "YGL162W"
[37] "YGR138C"
                  "YGR161C"
                               "YGR171C"
                                             "YGR213C"
                                                          "YGR250C"
                                                                       "YHL040C"
                                            "YHR216W"
[43] "YHR087W"
                  "YHR096C"
                               "YHR104W"
                                                          "YIL125W"
                                                                       "YJL034W"
[49] "YJL054W"
                  "YJL116C"
                               "YJR151C"
                                            "YKL029C"
                                                          "YKL090W"
                                                                       "YKL097W.A"
[55] "YKL163W"
                  "YKL165C"
                               "YKR061W"
                                            "YLL019C"
                                                          "YLL060C"
                                                                       "YLR120C"
[61] "YLR121C"
                  "YLR142W"
                               "YLR178C"
                                            "YLR194C"
                                                          "YLR350W"
                                                                       "YLR359W"
[67] "YML130C"
                  "YML131W"
                               "YMRO40W"
                                             "YMR090W"
                                                          "YMR173W"
                                                                       "YMR181C"
[73]
    "YMR300C"
                  "YNL112W"
                               "YNL134C"
                                            "YNL160W"
                                                          "YNL220W"
                                                                       "YOL151W"
[79] "YOLO31C"
                  "YOR173W"
                               "YOR289W"
                                             "YOR338W"
                                                          "YOR382W"
                                                                       "YPL088W"
                                                          "YFR024C.A" "YOL053C.A"
[85] "YPL277C"
                  "YPR156C"
                               "YARO75W"
                                             "YDR243C"
```

Next, we determine which of the 247 ko-genes also showed a significant QTL in our data set, according to a permutation test (Churchill and Doerge 1994) aiming to control GWER < 0.05. For each one of the ko-genes with a significant QTL, that is, with LOD score above lod.thr = 3.48, the function GetCandReg returns the ko-gene's chromosome (phys.chr) and physical position in cM (phys.pos), as well as, the LOD score (peak.lod) at the peak position (peak.pos), and the chromosome where the peak is located (peak.chr). In total, we observed 135 ko-genes with significant QTLs. These ko-genes are our candidate regulators. We show below the information on the first 10 candidate regulators. Note that some ko-genes map to the same chromosome where they are physically located, while other map to different chromosomes.

```
> cand.reg <- GetCandReg(highlod.orf, yeast.annot, names(ko.list))
> dim(cand.reg)
[1] 135
          6
> head(cand.reg)
      gene phys.chr phys.pos peak.chr
                                           peak.pos peak.lod
  YMR282C
                  13 473.2316
2
                                    14 236.0138450 3.692560
3
  YER017C
                   5 152.3216
                                    14 238.0138450 6.597231
7
  YER069W
                  5 211.7280
                                     3
                                        54.0140660 3.975861
                                          0.9067482 3.569372
  YORO58C
                  15 188.5460
                                     8
```

7 227.0394

13 235.2625

10 YGL017W

14 YMR055C

Genes that map to positions close to their physical locations are said to map in *cis* (local-linkages). Genes that map to positions away from their physical locations are said to map in *trans* (distal-linkages). There is no unambiguous way the determine how close a gene needs to map to its physical location in order to be classified as cis. Our choice is to classify a gene as cis if the 1.5-LOD support interval (Manichaikul et al. 2006) around the LOD peak contains the gene's physical location, and if the LOD score at its physical location is higher the the LOD threshold. The function GetCisCandReg determines which of the candidate regulators map in cis.

7 221.6439074 5.894020

13 246.0276440 5.578000

```
> cis.cand.reg <- GetCisCandReg(highlod.orf, cand.reg)
> dim(cis.cand.reg)

[1] 28 7
> head(cis.cand.reg)
```

	gene	phys.chr	phys.pos	peak.pos	peak.lod	<pre>peak.pos.lower</pre>	<pre>peak.pos.upper</pre>
10	YGL017W	7	227.0394	221.6439	5.894020	210.0037	230.7461
14	YMR055C	13	235.2625	246.0276	5.578000	144.0276	260.0276
16	S YMR275C	13	467.3183	460.0276	5.508846	420.0276	472.0276
48	3 YLR342W	12	402.5087	402.5087	8.666742	400.0196	410.0196
6:	YNLO21W	14	278.5199	242.0138	4.359721	222.0138	280.0138
63	YORO38C	15	179.1709	174.0012	5.060326	156.0012	184.0012

We see that only 28, out of the 135 candidate regulators, show cis-linkages. (The additional columns peak.pos.lower and peak.pos.upper show, respectively, the lower and upper bounds of the 1.5-LOD support interval around peak.pos.)

For each one of the 135 candidate ko-genes, we determined which other genes also co-mapped to the same QTL of the ko-gene. The co-mapping genes represent the putative targets of a ko-gene. The function GetCoMappingTraits returns a list with the putative targets of each ko-gene. A gene is included in the putative target list of a ko-gene when its LOD peak is greater than lod.thr and the 1.5 LOD support interval around the peak contains the location of the ko-gene's QTL.

```
> comap.targets <- GetCoMappingTraits(highlod.orf, cand.reg)</pre>
> summary(sapply(comap.targets, length))
                 Median
                            Mean 3rd Qu.
   Min. 1st Qu.
                                             Max.
    1.0
           63.5
                                   479.0
                   188.0
                           236.9
                                            569.0
> comap.targets[[7]]
[1] "YDL013W" "YGL254W" "YML069W" "YMR247C"
> length(unlist(comap.targets))
[1] 31975
```

The number of targets vary from ko-gene to ko-gene (from 1 to 569). We illustrate with the putative targets of one ko-gene (YMR275C) with 4 putative targets. In total, the 135 candidate regulators have 31975 targets.

Next, we use the function FitAllTests to fit the causality tests of each candidate regulator ko-gene (pheno1) to its putative targets (pheno2). We use the candidate regulator's QTL (Q.chr and Q.pos) as a causal anchor. This function fits: the AIC and BIC model selection criterions (Schadt et al. 2005); the AIC- and BIC-based versions of the joint, parametric and

non-parametric CMST tests (Chaibub Neto et al. 2012); and the CIT test (Millstein et al. 2009). We do not run it here because this step can take a few hours, as we perform a total of 31975 tests for each of the 9 approaches. The function JoinTestOutputs joins together the outputs of the 135 separate fits of the FitAllTests function.

```
> set.seed(123456789) # we fix a seed because cit uses bootstrap
> for (k in 1 : 135) {
>
    cat("trait=", k, "\n")
    out <- FitAllTests(cross = yeast.orf,
>
                       pheno1 = cand.reg[k, 1],
                       pheno2 = comap.targets[[k]],
                       Q.chr = cand.reg[k, 4],
                       Q.pos = cand.reg[k, 5])
    save(out, file=paste("output_ko_validation", cand.reg[k, 1], "RData",
>
         sep = "."), compress = TRUE)
+
> }
> ko.tests <- JoinTestOutputs(comap.targets, file = "output_ko_validation")</pre>
```

We are now using the Benjamini-Hochberg adjustment for the non-parametric CMST tests. Therefore to get the adjusted values, we do the following:

```
> ## data(ko.tests) is loaded lazily.
> adj.ko.tests <- p.adjust.np(ko.tests)</pre>
```

After loading the joined results we use the function PrecTpFpMatrix to summarize the performance of the different methods in terms of "biologically validated" true positives, false positives and precision, of the inferred causal relations. Since we already have the results of the knock-out experiments (recall that ko.list holds the ko-signatures of the ko-genes), we define a true positive as a statistically significant causal relation between a ko-gene and a putative target gene, when the putative target gene belongs to the ko-signature of the ko-gene. Similarly, we define a false positive as a statistically significant causal relation between a ko-gene and a putative target gene when the target gene doesn't belong to the ko-signature. (For the AIC and BIC methods, that do not provide a p-value measuring the significance of the causal call, we simply use the detected causal relations in the computation of true and false positives). The "validated precision", is computed as the ratio of true positives by the sum of true and false positives. The PrecTpFpMatrix computes these measures to both all ko-genes, and to cis ko-genes only. The argument alpha sets the significant levels at each the summaries are computed. Since this takes awhile, we have also stored roc.aux as a data object in the package.

```
> ## data(roc.aux) is loaded lazily.
```

Before we show plots, here are some preliminary plot settings in a simple plot routine that will be used repeatedly for figures.

```
> plots <- function(roc.aux, elements) {</pre>
    par(mfrow = c(1,3))
    par(mar=c(5, 4.1, 4, 2) + 0.1)
    myplot(roc.aux[[elements[1]]], "Number of true positives", "(a)")
    myplot(roc.aux[[elements[2]]], "Number of false positives", "(b)")
    myplot(roc.aux[[elements[3]]], "Precision", "(c)")
+
+ }
> myplot <- function(sum.type, sum.label, main = "") {</pre>
    ymax <- max(sum.type)</pre>
    my.pch \leftarrow c(1, 21, 24, 23, 25, 2, 5, 6, 8)
    xaxis \leftarrow seq(0.01, 0.10, by=0.01)
    yaxis <- seq(0, ymax,length.out = length(xaxis))</pre>
    plot(xaxis, yaxis, type = "n", ylab = sum.label, cex = 1.5,
         xlab = "target level", cex.axis = 1.5,
+
         cex.lab = 1.7, main = main, cex.main = 2)
    for (k in 1 : 9) {
+
      lines(xaxis, sum.type[k,], type="b", lwd=2, pch=my.pch[k], cex=1.5,
             col = "black", bg = "black")
    }
+ }
```

Below we reproduce Figure 8 of Chaibub Neto et al. (2012). This figure presents the number of inferred true positives, number of inferred false positives and the prediction precision across varying significance levels for each one of the methods. The results were computed using all 135 ko-gene/putative target lists. Next, we reproduce Figure 9 of Chaibub Neto et al. (2012). This figure was generated using the results of the 27 cis ko-gene/putative targets lists.

3 References

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- Zhu J., B. Zhang, E. N. Smith, B. Drees, R. B. Brem, L. Kruglyak, R. E. Bumgarner, E. E. Schadt, 2008 Integrating large-scale functional genomic data to dissect the complexity of yeast regulatory networks. Nature Genetics 40: 854-861.

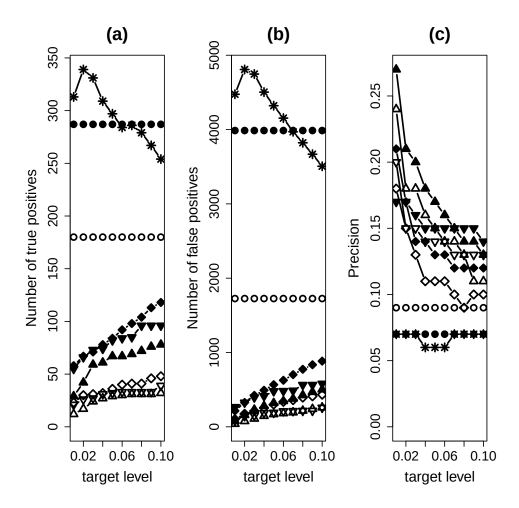


Figure 1: Reproduction of Figure 8 on Chaibub Neto et al. 2012. Target significance level by overall (a) number of true positives, (b) number of false positives and (c) precision across all 135 ko-gene/putative target lists. Asterisk represents the CIT. Empty and filled symbols represent, respectively, AIC- and BIC-based methods. Diamonds: parametric CMST. Point-down triangles: non-parametric CMST. Point-up triangles: joint-parametric CMST. Circles: AIC and BIC.

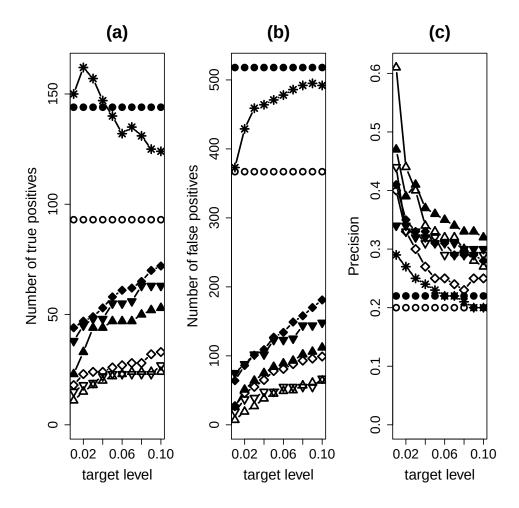


Figure 2: Reproduction of Figure 9 on Chaibub Neto et al. 2012. Target significance level by overall (a) number of true positives, (b) number of false positives and (c) precision restricted to 27 cis ko-gene/putative target lists. Asterisk represents the CIT. Empty and filled symbols represent, respectively, AIC- and BIC-based methods. Diamonds: parametric CMST. Point-down triangles: non-parametric CMST. Point-up triangles: joint-parametric CMST. Circles: AIC and BIC.