Testing pleiotropy in multiparental populations

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Introduction

- 10,000+ traits with RNA sequencing and mass spectrometry
- Multiparental populations offer high-resolution QTL mapping
- New analysis tools, such as a pleiotropy test for multiparental populations, are needed

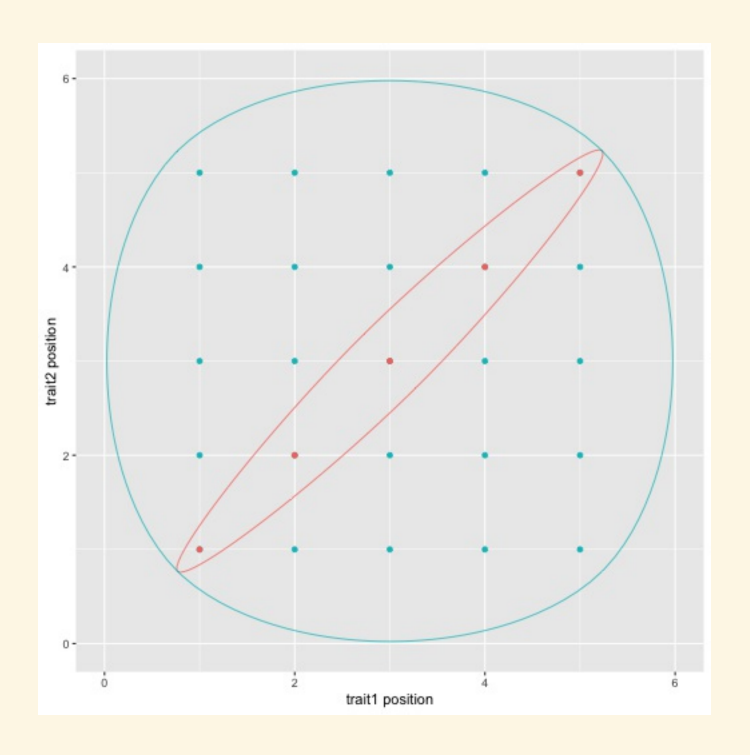
Jiang and Zeng (1995) test

- Two-parent crosses
- Applies to two traits that co-map
- H_0 : Pleiotropy
- H_A: Two separate QTL

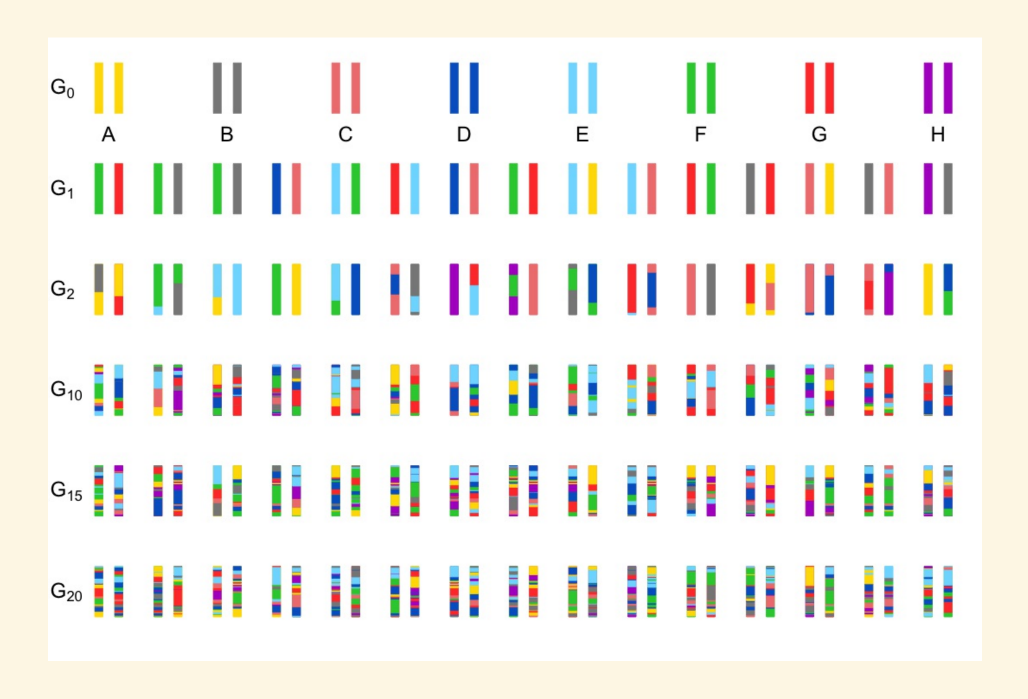
Jiang and Zeng (1995) test

- Perform a two-dimensional QTL scan
- Y = XB + E
- Calculate likelihood ratio test statistic

Jiang and Zeng (1995) test



Multiparental populations



Challenges in multiparental populations

- Complex patterns of relatedness
- Multiple founder lines
- Determining statistical significance

Challenges in multiparental populations

Complex patterns of relatedness

• Multiple founder lines

Determining statistical significance

Test procedure

Model:

$$vec(Y) = Xvec(B) + vec(G) + vec(E)$$

Calculate likelihoods

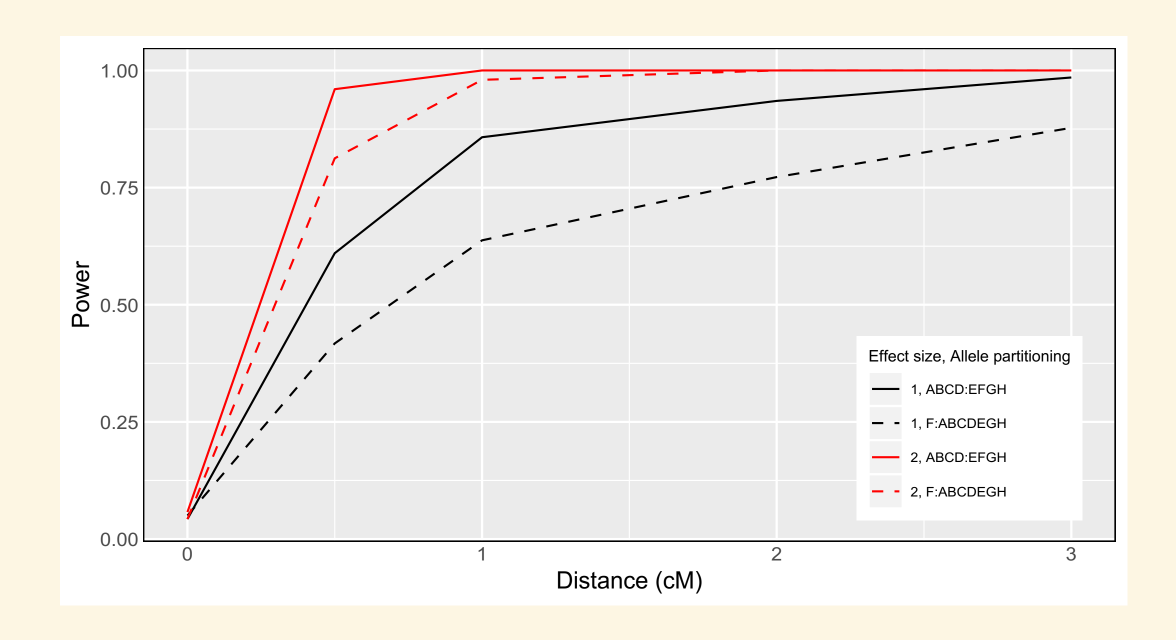
Test procedure

Test statistic:

$$-\log \frac{\text{max(likelihood under pleiotropy)}}{\text{max(likelihood for separate QTL)}}$$

Parametric bootstrap to get p-value

Test characteristics



Diversity Outbred Mice

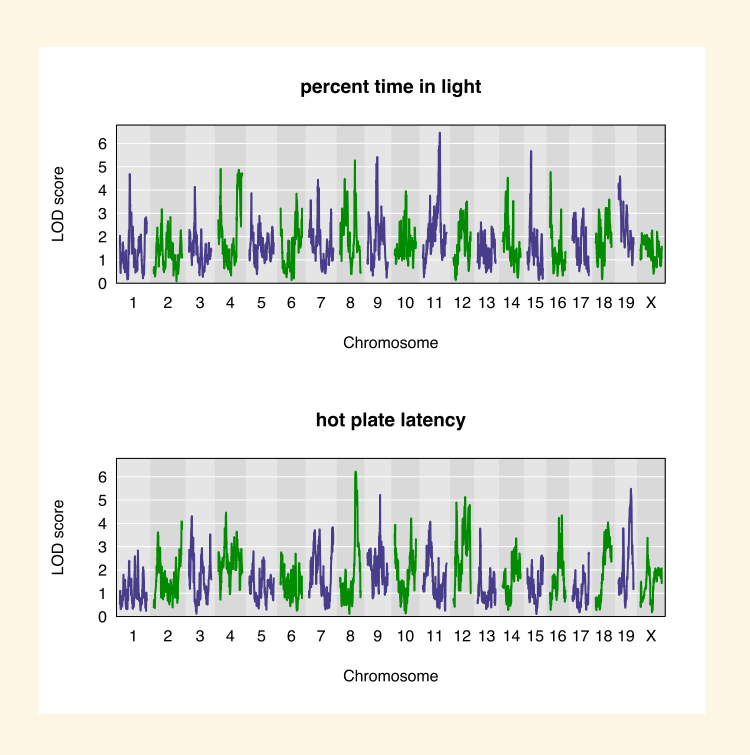
Multiparental population arising from 8 inbred lines



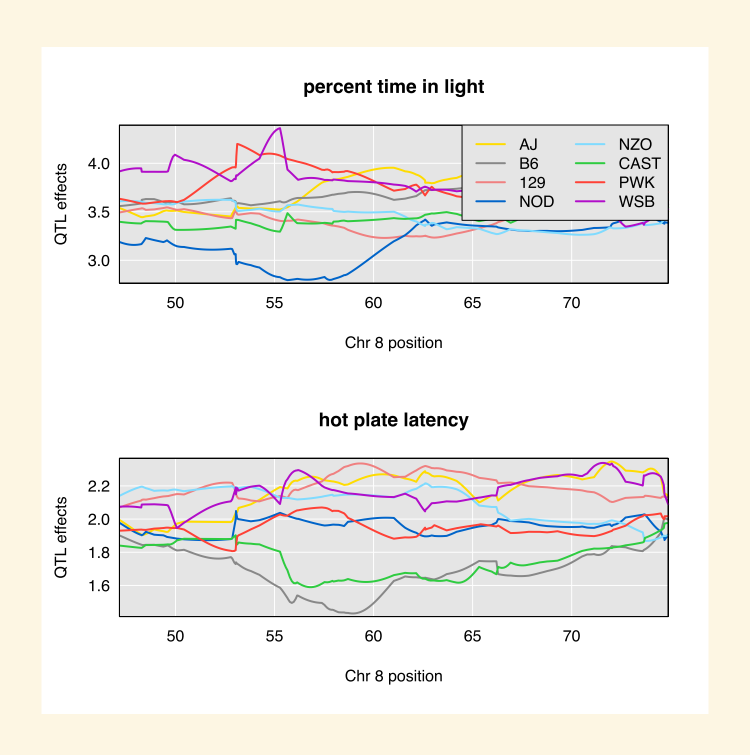
Application

- Logan, et al. (2013) and Recla, et al. (2014) studied 261 Diversity Outbred mice
- Two traits map to Chr 8:
 - "hot plate latency" (57 cM)
 - "percent time in light" (55 cM)

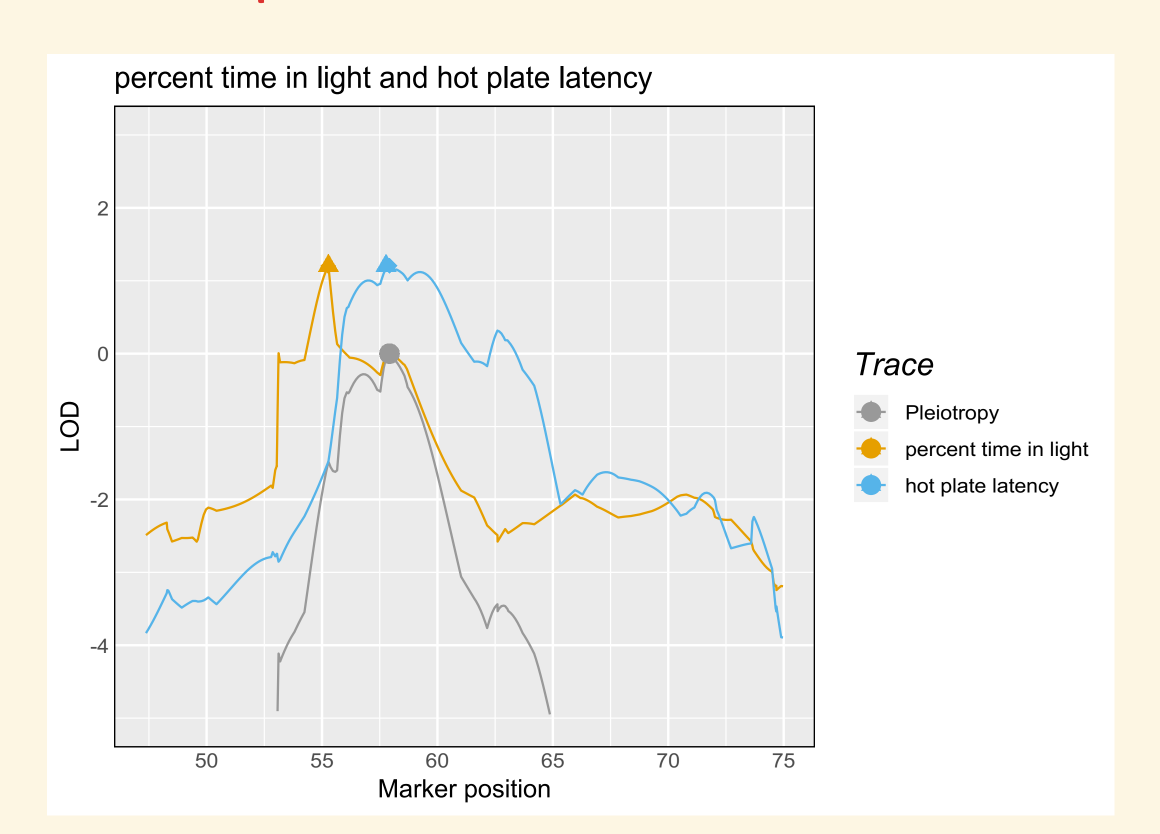
QTL scan results



Allele effects plots



Profile LOD plot



Test results

• p = 0.11 (1000 bootstrap samples)

Conclusions

- Weak evidence for two separate QTL
 - One QTL affects "distance traveled in light"
 - Second QTL affects "hot plate latency"

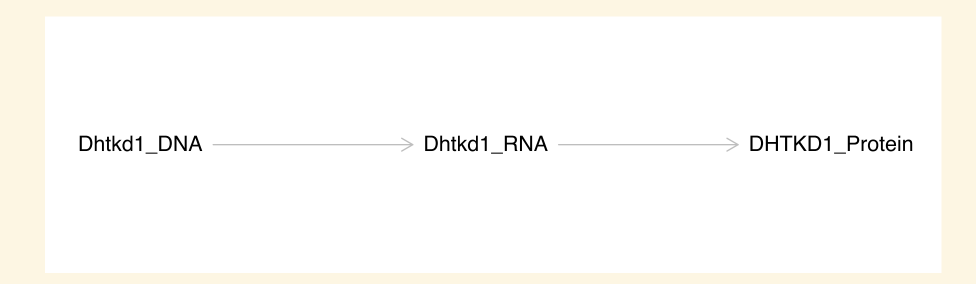
Mediation analysis and pleiotropy testing

Mediation analysis

• Is one molecule an intermediate in a causal pathway?

Mediation analysis example

- QTL near Dhtkd1 gene affects Dhtkd1 RNA levels and DHTKD1 protein levels
- Is DHTKD1 an intermediate between QTL and DHTKD1 protein levels?



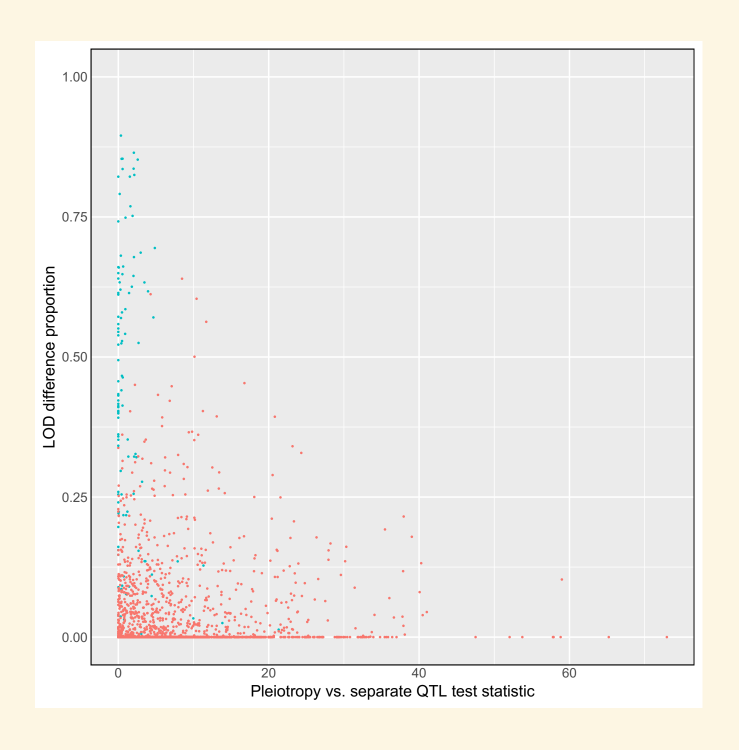
Mediation analysis methods

- LOD score with and without conditioning on candidate intermediate
- LOD difference
- LOD difference proportion

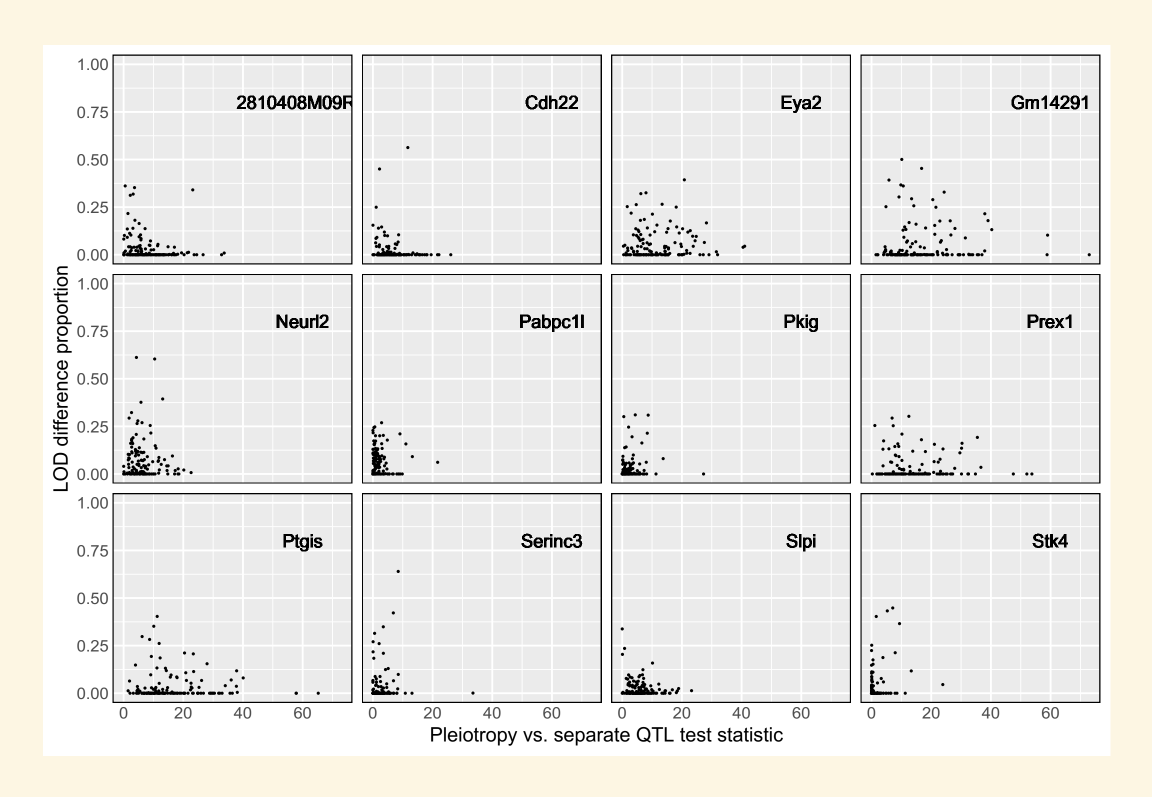
Dissecting an eQTL hotspot

- Keller, et al. (2018) identified 147 nonlocal traits that map to Chr 2 (163.5 Mb to 167.5 Mb)
- expression levels mediate 88 of 147 traits
- Do other local genes act as intermediates?

Dissecting an eQTL hotspot



Dissecting an eQTL hotspot



qtl2pleio R package development

Design elements

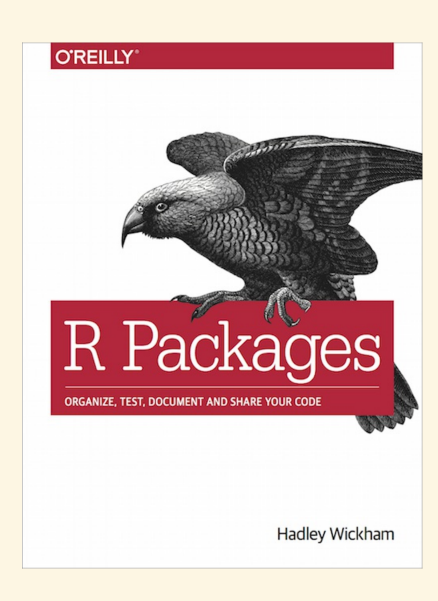
- Written mostly in R
 - Some calculations in C++, via Rcpp package
 - C++ Eigen library and RcppEigen for matrix algebra
 - User interacts only with R functions

Design elements

- Tutorial vignettes demonstrate functionality
- Unit tests included via
 R package
- Documentation with R package
- Package website with
 R package

R packages book

• Hadley Wickham's "R packages" book as a guide



Open-source R package

- Hosted on github
- Install via R package
- commits publicly visible
- Travis CI for testing after every commit

Binder integration

- Enhances reproducibility
- Uses Docker images
- Downloads R packages for a specified date (or commit on Github)

Binder integration: 2 new files

- install.R
 - o contains code to install R packages
- runtime.txt
 - o contains one line of text to indicate date for CRAN packages

Future directions

Contact information

- frederick.boehm@gmail.com
- https://fboehm.us/
- qtl2pleio R package: https://github.com/fboehm/qtl2pleio
- Biorxiv pre-print: http://bit.ly/boehm-biorxiv

References

Jiang, C. and Z. Zeng (1995). "Multiple trait analysis of genetic mapping for quantitative trait loci." In: 140.3, pp. 1111-1127.

Keller, M. P, D. M. Gatti, et al. (2018). "Genetic Drivers of Pancreatic Islet Function". In: pp. genetics-300864.

Logan, R. W, R. F. Robledo, et al. (2013). "Highprecision genetic mapping of behavioral traits in the diversity outbred mouse population". In: 12.4, pp. 424-437.

References

Recla, J. M, R. F. Robledo, et al. (2014). "Precise genetic mapping and integrative bioinformatics in Diversity Outbred mice reveals Hydin as a novel pain gene". In:

25.5-6, pp. 211-222.