Testing pleiotropy vs. separate QTL in multiparental populations

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Abstract

- High-dimensional phenotypes & multiparental populations inform complex trait architecture
- New analysis tools needed
- Developed a test of pleiotropy vs. separate QTL for multiparental populations
- Found evidence for two separate OTL affecting three behavioral traits in a 3 cM region in 261 Diversity Outbred mice
- Shared methods in an R package, gt12pleio <https://github.com/fboehm/qtl2pleio>

Behavioral genetics

- Logan et al. (2013) and Recla et al. (2014) genotyped and phenotyped 261 Diversity Outbred mice
- Identified \emph{Hydin} as the chromosome 8 gene affecting "hot plate latency" at 57 cM
- Identified chromosome 8 QTLs for "percent time in light" and "distance traveled in light" at 55 cM
- Motivated us to ask if *Hydin* also affects "percent time in light" and "distance traveled in light"

3 pairwise tests

- "percent time in light" & "hot plate latency": p = 0.109
- "distance traveled in light" & "hot plate latency": p = 0.108
- "percent time in light" & "distance traveled in light": p = 0.871

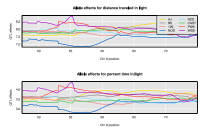
Introduction

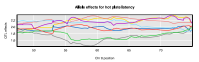
- Experimentalists can now measure tens of thousands of phenotypes with RNA sequencing and mass spectrometry
- Multiparental populations enable high-resolution QTL mapping
- Together, high-dimensional phenotypes and multiparental populations can inform complex trait
- New analysis tools, such as our test of pleiotropy vs. separate QTL, are needed

Background

- Jiang and Zeng (1995) developed a pleiotropy vs. separate OTL test for two-parent crosses
- Applies to two traits that map to a single genomic region
- Pleiotropy is the null hypothesis
- Separate QTL is the alternative hypothesis
- Perform a two-dimensional QTL scan
- Calculate likelihood ratio test statistic

Allele effects plots





Conclusions

- Evidence for two separate QTL affecting the 3 traits • 1 QTL affects both "distance traveled in light" and
- "percent time in light"
- Second QTL contains *Hydin* and affects "hot plate latency'

Future directions

- Examine expression data from Keller et al. (2018)
- Expression QTL hotspot dissection
- · Statistical power studies

Challenges

- Relatedness: Multivariate polygenic random effects
- Eight founder lines: 8 fixed effects
- Test statistic calibration: Parametric bootstrap test

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<https://github.com/fboehm/qtl2pleio>

Test procedure

Fit the model:

$$Y = XB + G + E \$$
 for each ordered pair of markers

$$G \sim MN(0,K,V_g)$$

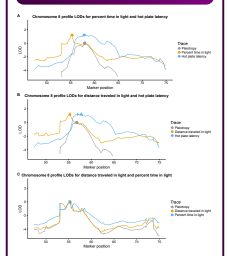
$$E \sim MN(0,I,V_e)$$

- X contains allele probabilities
- B contains allele effects
- Test statistic:

$$-\log_{10} rac{\max L_0(B,\Sigma,\omega_1)}{\max L_A(B,\Sigma,\omega_1,\omega_2)}$$

Parametric bootstrap to get p-value

Profile LOD plots for pairwise



References

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