

# 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 two-QTL scan
  - $vec(Y) = Xvec(B) + vec(E)$
  - Calculate likelihood at each ordered pair of positions
- Calculate likelihood ratio test statistic

# Multiparental populations



# Challenges in multiparental populations

- Complex patterns of relatedness

*Multivariate random effects*

- Multiple founder lines

*Fixed effect for each founder allele*

# Test procedure

- Perform a two-dimensional two-QTL scan
  - $vec(Y) = Xvec(B) + vec(G) + vec(E)$
  - Calculate likelihood at each ordered pair of positions
- Calculate likelihood ratio test statistic

# Test procedure

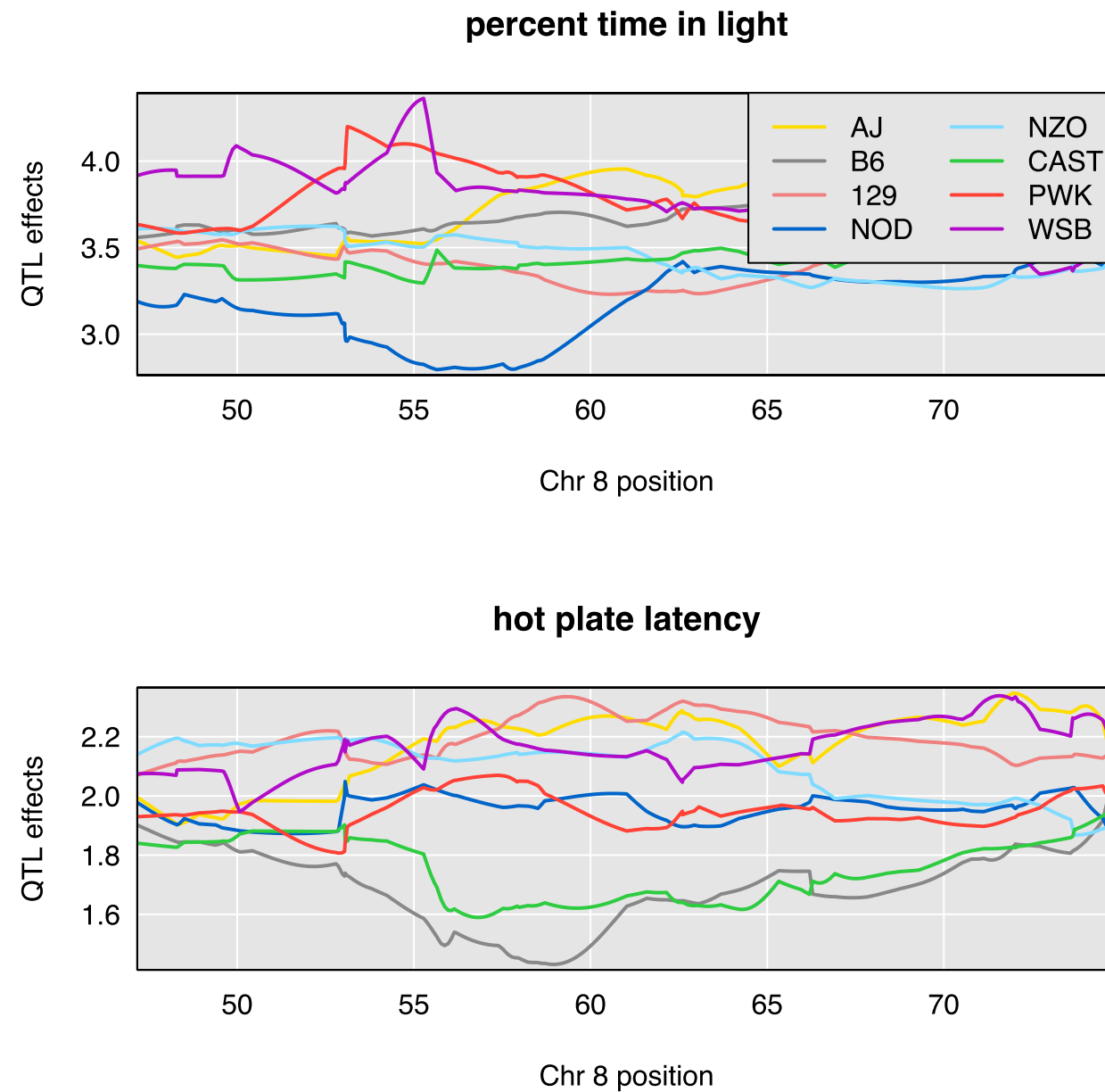
- Test statistic:  $-\log_{10} \max$
- Parametric bootstrap to get a p-value



# Application

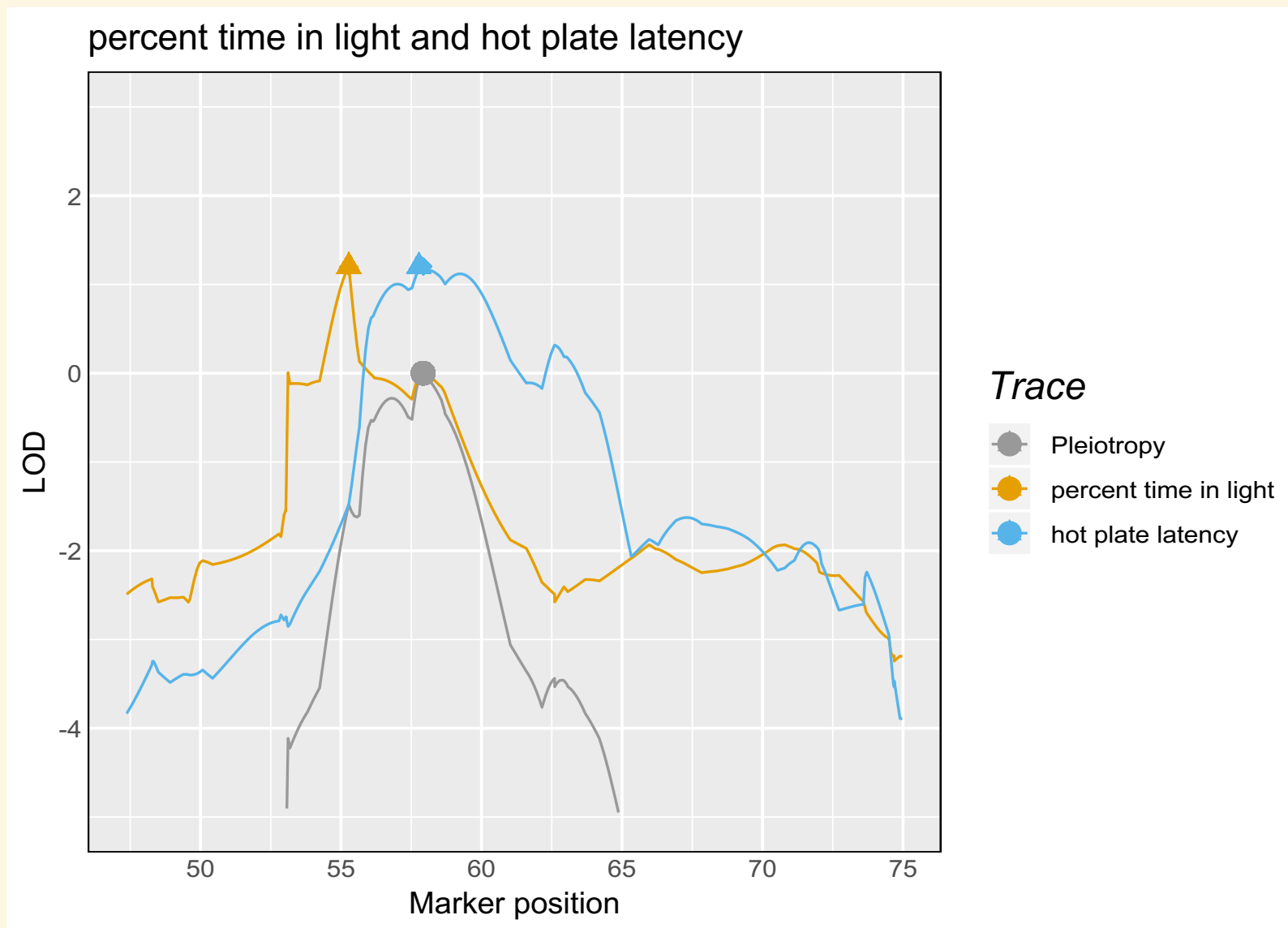
- Logan, et al. (2013) and Recla, et al. (2014) studied 261 Diversity Outbred mice
- Measured about two dozen behavioral traits
- Two traits map to Chr 8:
  - "hot plate latency" (57 cM)
  - "percent time in light" (55 cM)

# Allele effects plots



# Profile LOD

$$\text{profile LOD}_{\text{trait 1}}(\lambda_1) = \max_{\lambda_2} \text{LOD}(\lambda_1, \lambda_2)$$



# Test results

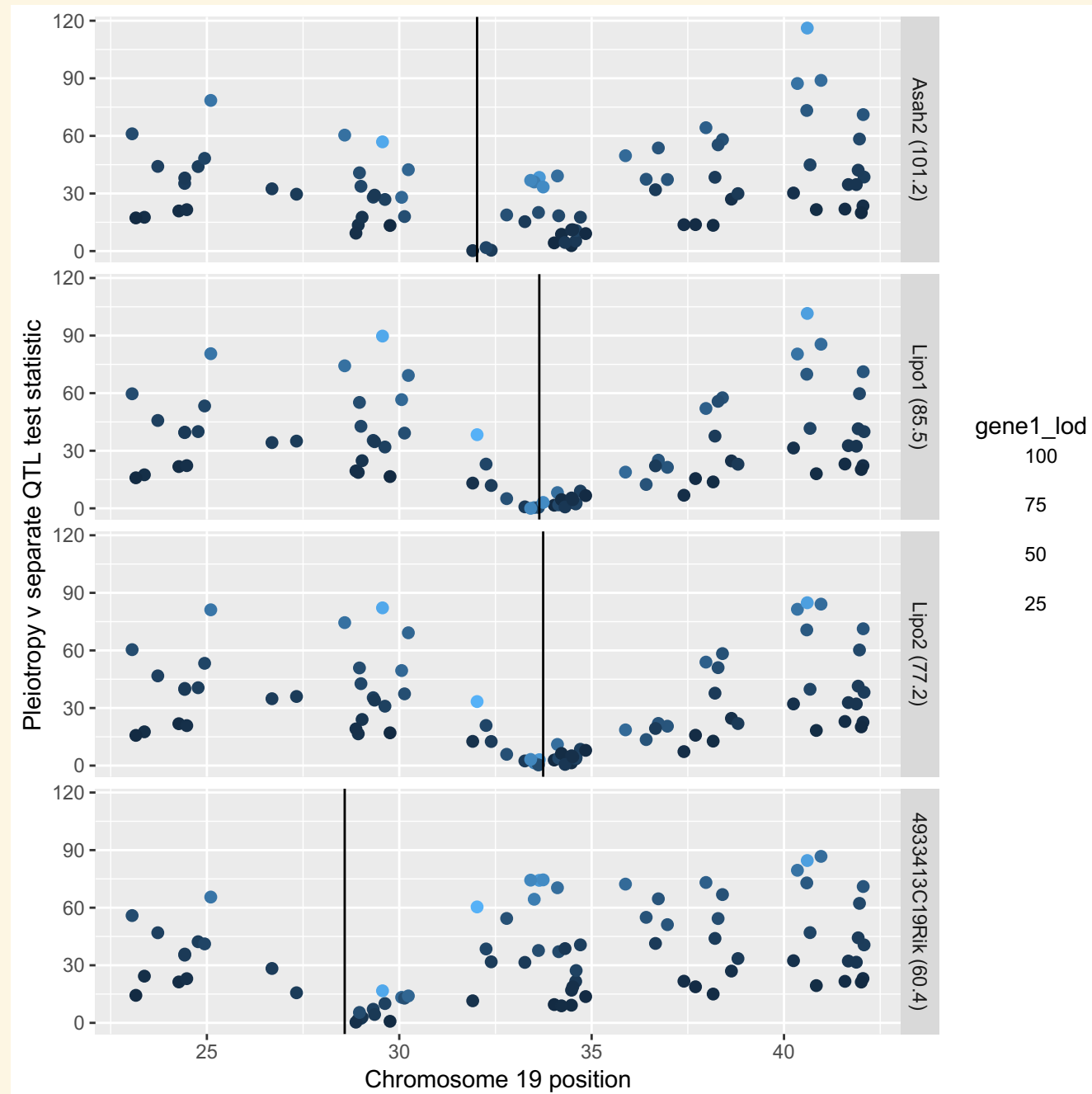
- $\log_{10} \Lambda = 1.2$
- $p = 0.11$  (1000 bootstrap samples)

# Power study with local expression QTL

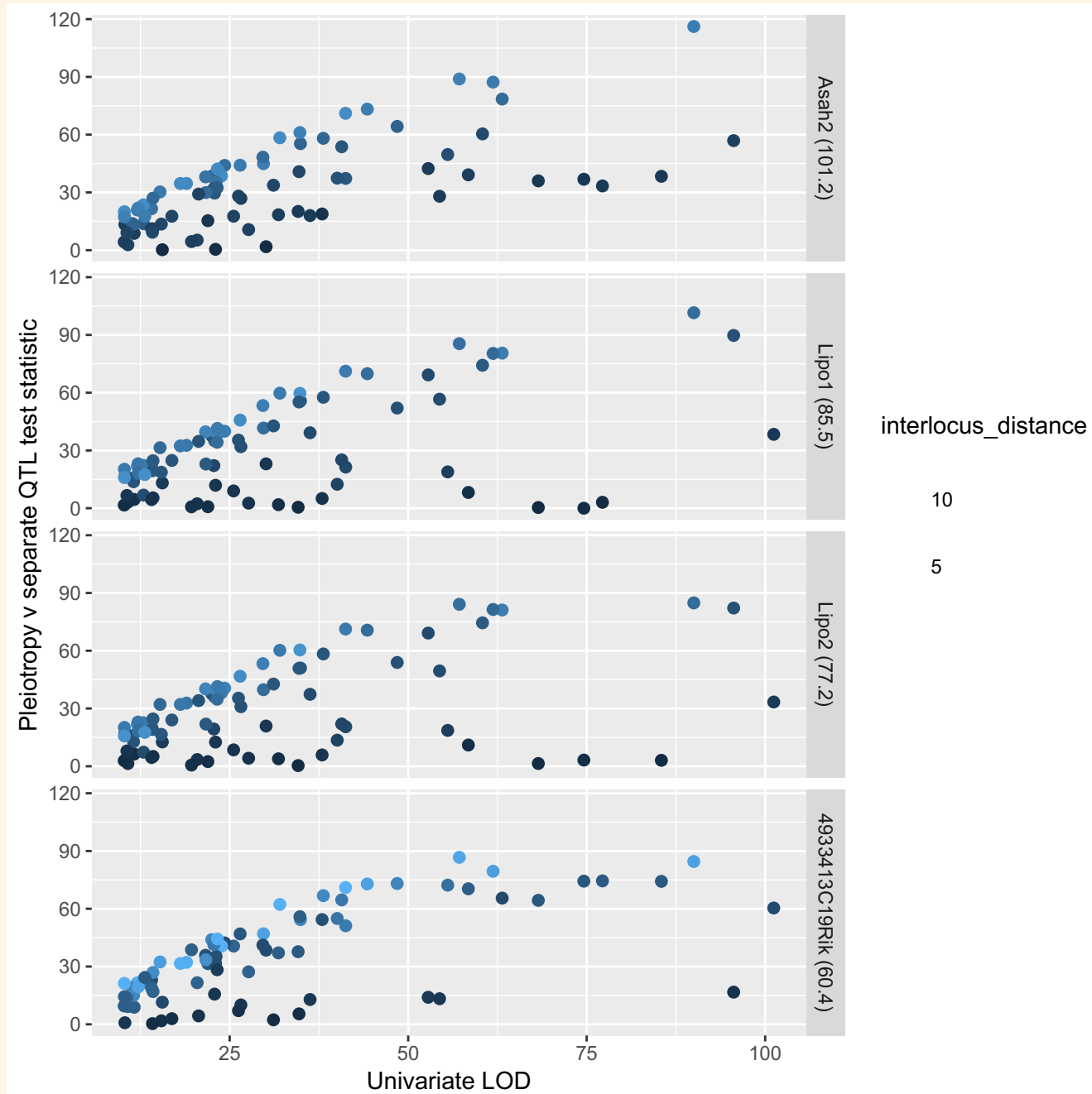
# Power study: Design

- Examine power when I know true eQTL locations
  - Interlocus distance
  - Univariate LOD
- 80 local eQTL on Chr 19 ( $\text{LOD} > 7.2$ )
- 4 strong, centrally located, local eQTL chosen as "anchor" traits
  - Asah2, Lipo1, Lipo2, 4933413C19Rik
- Pairwise pleiotropy tests each involving one anchor and one of 79 other traits

# Power study: Interlocus distance



# Power study: Univariate LOD





# Power study: Conclusions

- $\uparrow$  Interlocus distance  $\rightarrow$   $\uparrow$  Pleiotropy test statistics
- $\uparrow$  Univariate LOD  $\rightarrow$   $\uparrow$  Pleiotropy test statistics

# qtl2pleio R package

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- Functions for d-variate, d-QTL scan & profile LOD plots
- Uses C++ for matrix calculations (via Rcpp and RcppEigen)
- Uses [gemma2](#) R implementation of GEMMA EM algorithm for multivariate random effects
- Unit tests, vignettes, and version control

Thank you!

# References

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

Logan, R. W, R. F. Robledo, et al. (2013). "High-precision genetic mapping of behavioral traits in the diversity outbred mouse population". In: *Genes, Brain and Behavior* 12.4, pp. 424-437.

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: *Mammalian genome* 25.5-6, pp. 211-222.