general HMM for multi-parent populations

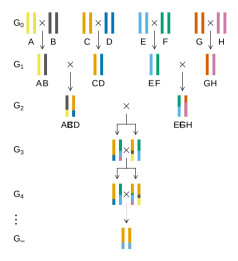
Karl Broman

Biostatistics & Medical Informatics, UW-Madison

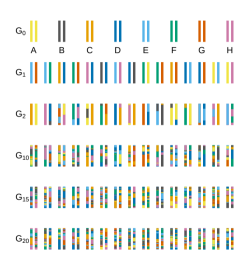
@kwbroman
 kbroman.org
 github.com/kbroman
kbroman.org/Talk_GeneralHMM



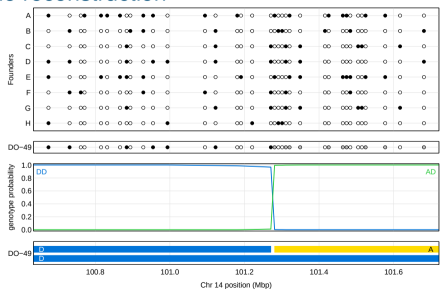
Recombinant Inbred Lines



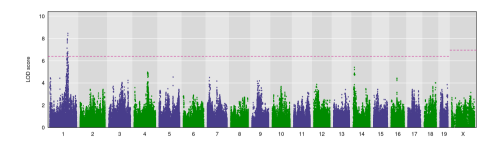
Advanced Intercross Population



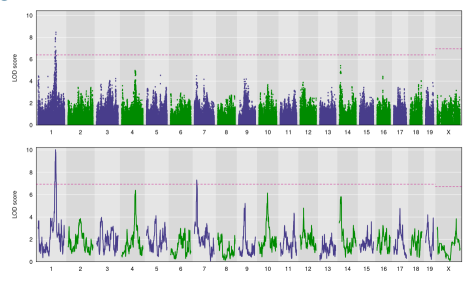
Genome reconstruction



QTL genome scan



QTL genome scan



Summary

► bioRxiv manuscript: doi:10/gswx

Slides: kbroman.org/Talk_GeneralHMM



bioRxiv manuscript: doi.org/gswx

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