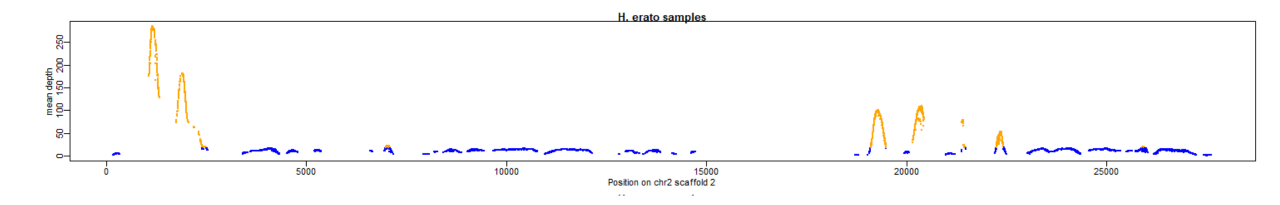
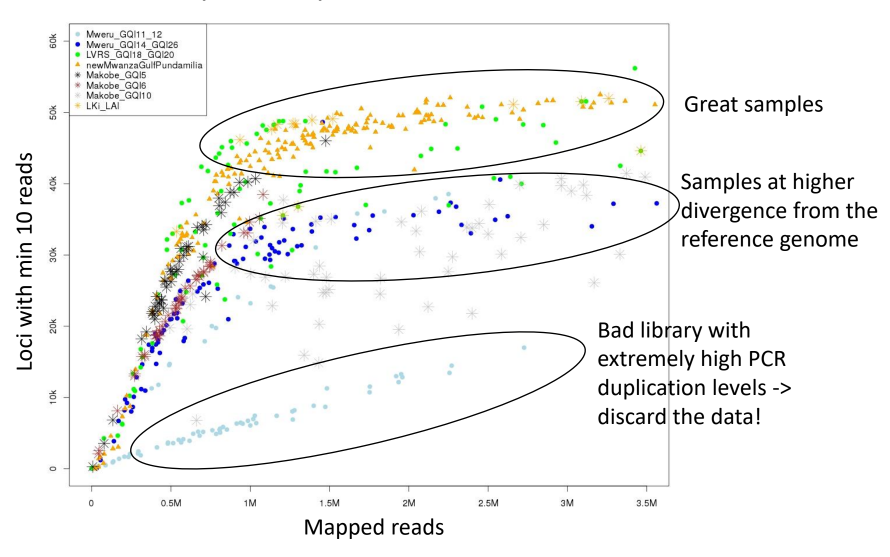
# Detecting high levels of PCR duplication and contamination Joana Meier

# High sequencing depth and heterozygosity indicate collapsed paralogous regions

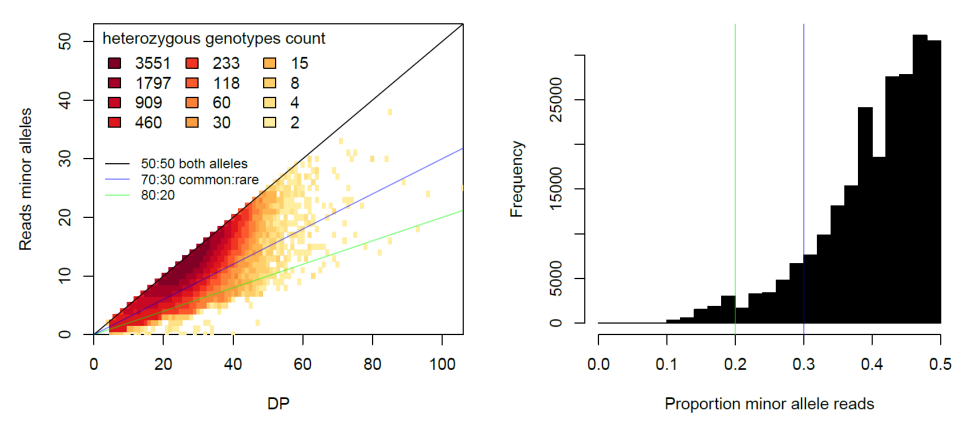


# Checking PCR duplication levels of single-end sequenced RAD/GBS/UCE libraries

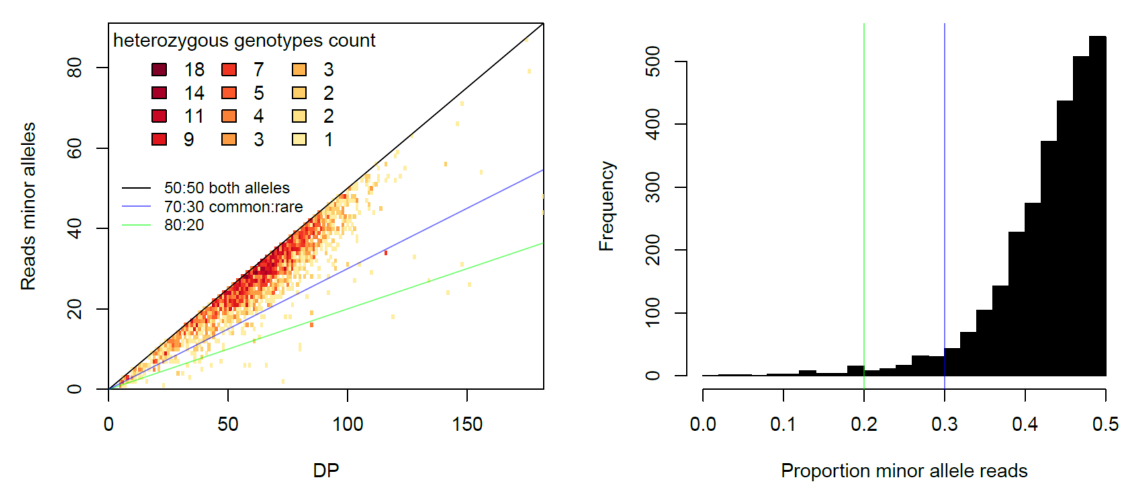


If reads truly independent sequences of the same genotype (from different cells of the same individual), heterozygotes are expected to have a roughly equal number of reads supporting each allele.

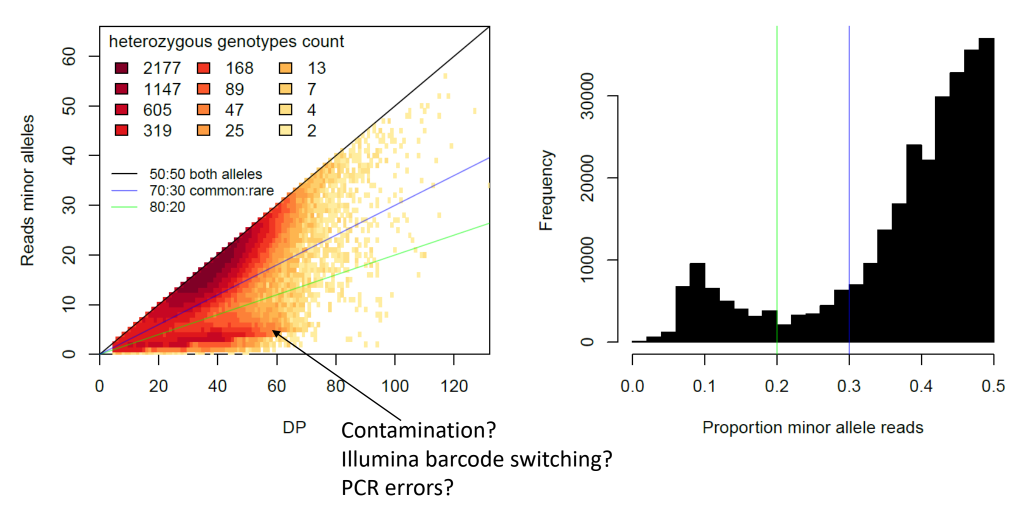
#### Example from a good wgs sample



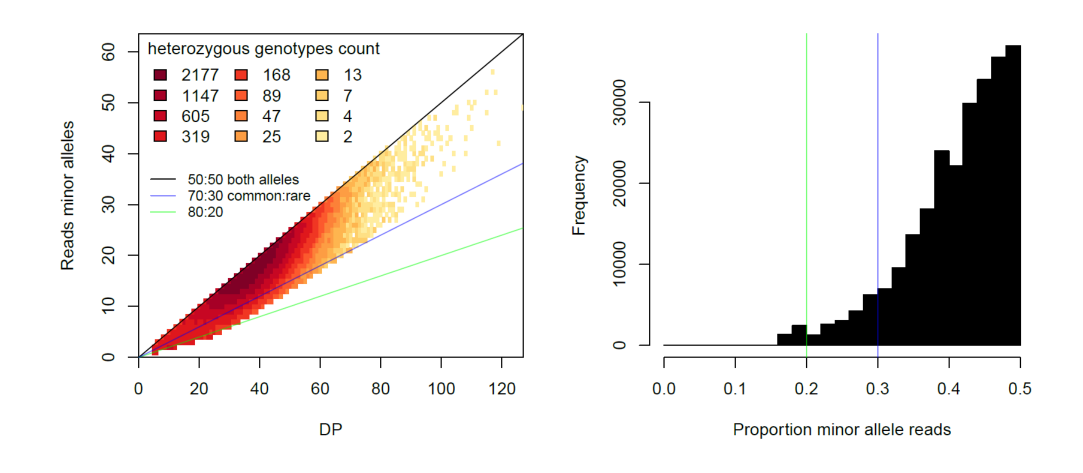
#### **Example from a good RAD library**



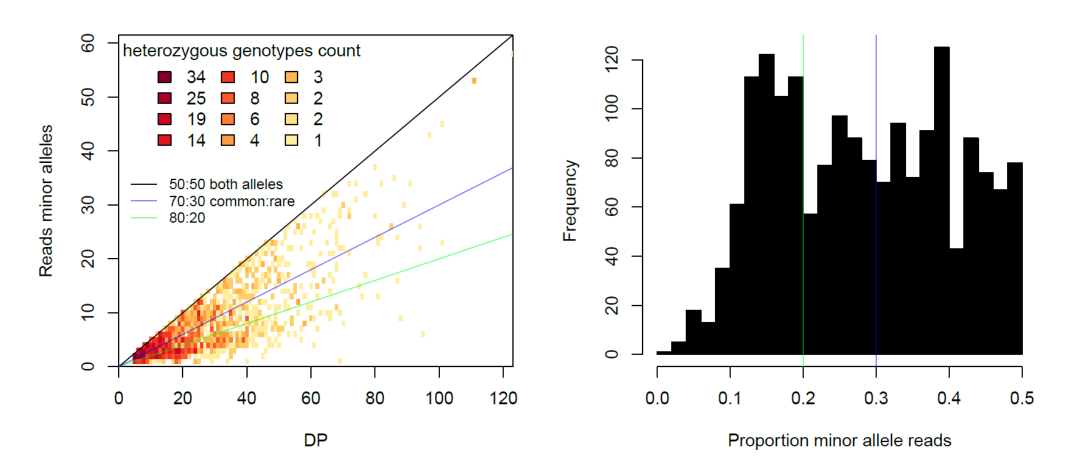
#### **Example from a contaminated wgs sample**



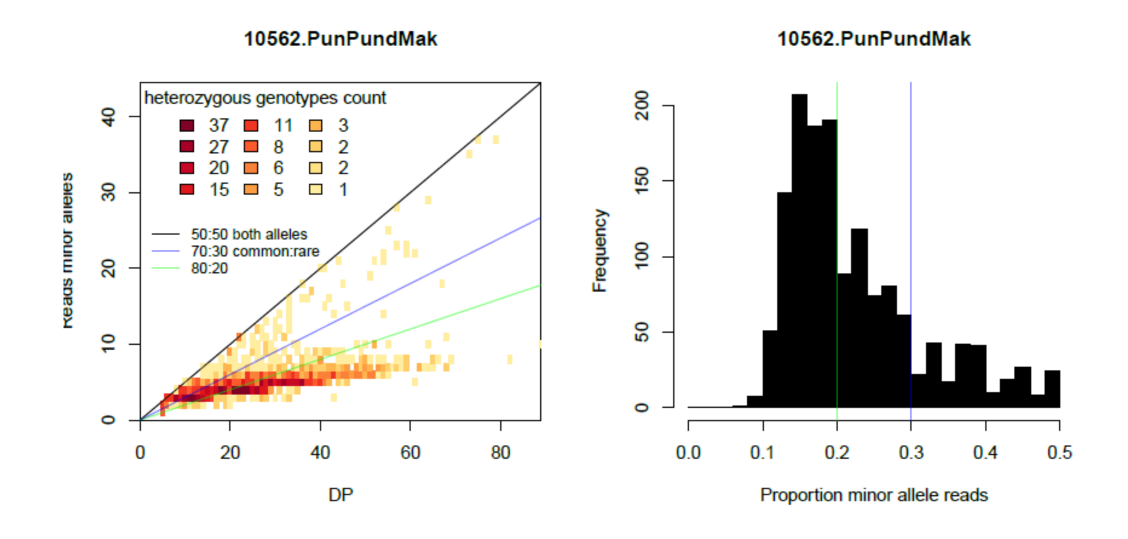
# AllelicBalance.py to remove heterozygote positions failing a binomial test



#### **Example from a very bad RAD library**



This individual looked ok but when looking only at sites with mac 1 across all individuals, PCR errors became very obvious



### How to distinguish the different sources?

- Illumina barcode switching: Affects samples that are most divergent (wrongly assigned reads are most likely to cause SNPs) and samples that do not have good DNA quality, allelic balance should be better in singletons
- PCR errors in libraries with PCR duplicates: allelic balance worst in singletons, usually decreased heterozygosity due to allelic dropout, generally high variance of reads supporting each allele even at high sequencing depth -> make sure to remove singletons
- **Contamination**: increased heterozygosity expected, may appear to be a hybrid, no signal in singletons