Checking SNP and pedigree information of sibs for Mendelian inconsistencies

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Introduction

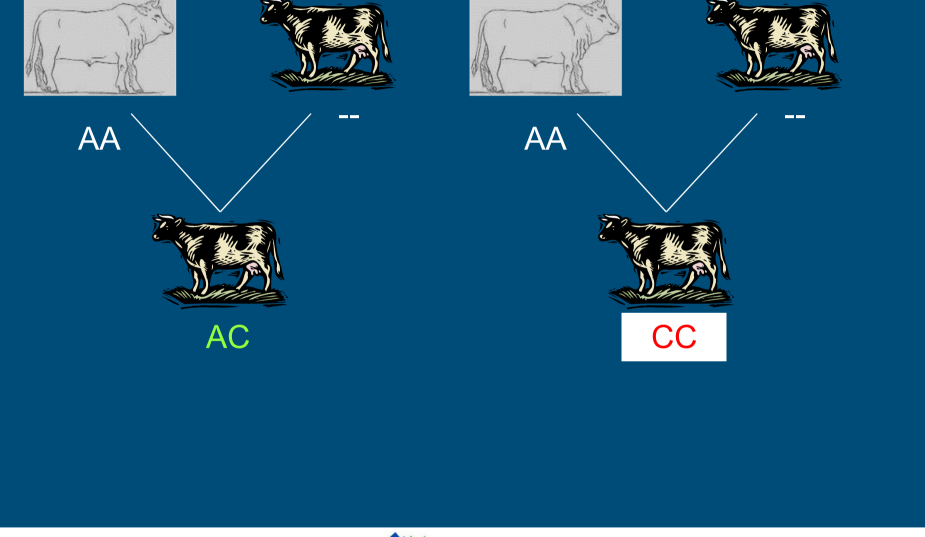
Mendelian Inconsistencies (MI):

- Identify animals with conflicting pedigree and SNP info
- Identified by counting loci with opposing homozygotes between 2 animals
- Identification is straightforward for parent-offspring pairs
 - Expected number of loci with opposing homozygotes = 0





MI – Parent-offspring pair







MI – Paternal half sib pair (dams not genotyped)











AA

CC

What if we don't know the sires genotype?



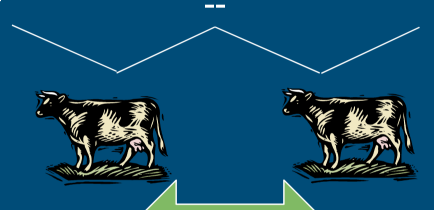


MI – Paternal half sib pair (parents not genotyped)









AA opposing CC

What is the expected number of opposing homozygote loci

for a pair of sibs?

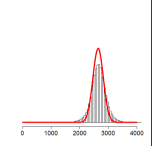




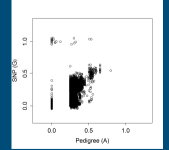


Develop tests to detect MI between sibs, based on:

Realized <u>number of opposing homozygote SNP loci</u>



Comparing <u>pedigree and SNP based relationships</u> between sibs







Two tests developed for sibs - algorithms SIBCOUNT

Count SNP with opposing homozygotes per sib pair

SIBREL

 Compare A (pedigree) with G (SNP) relationships per sib pair

In both tests:
Define empirical thresholds to declare conflicts

Iteratively remove animals that cause most MI





Identify animals causing MI – approach taken

Detect MI for parent-offspring pairs

- Parent-offspring pairs were defined based on pedigree or SNP
- Run SIBCOUNT or SIBREL
- Type I & II error rates for SIB tests were derived by permuting part of the 'cleaned up' data





RobustMilk genotype data

2,078 cows and 211 bulls; 36,884 SNP loci
 Cows from research herds in NL, UK, Ireland & Sweden

891 genotyped mother – daughter pairs

1,448 genotyped father – daughter pairs

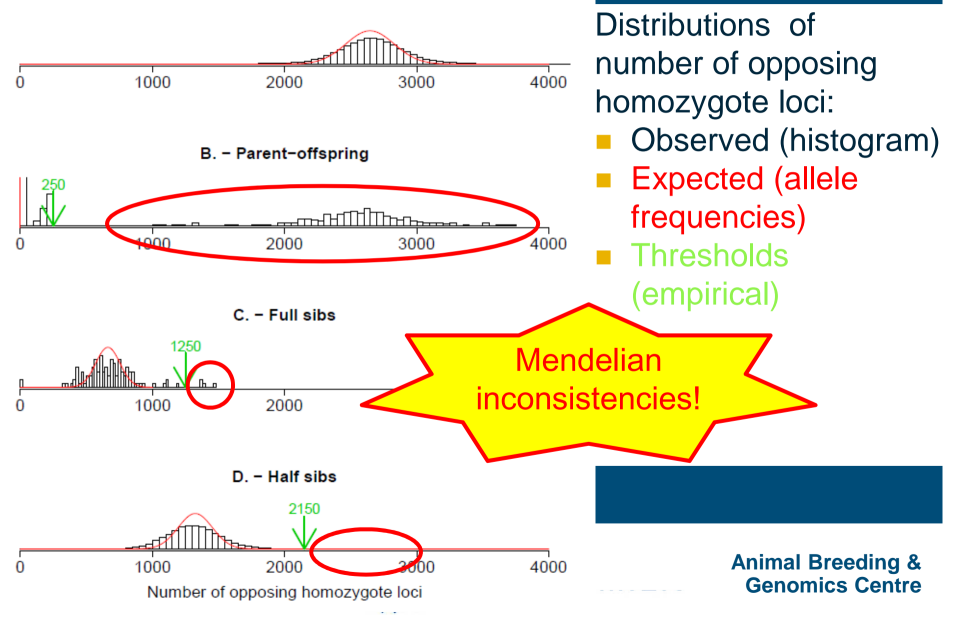
508 genotyped animals without genotyped parent





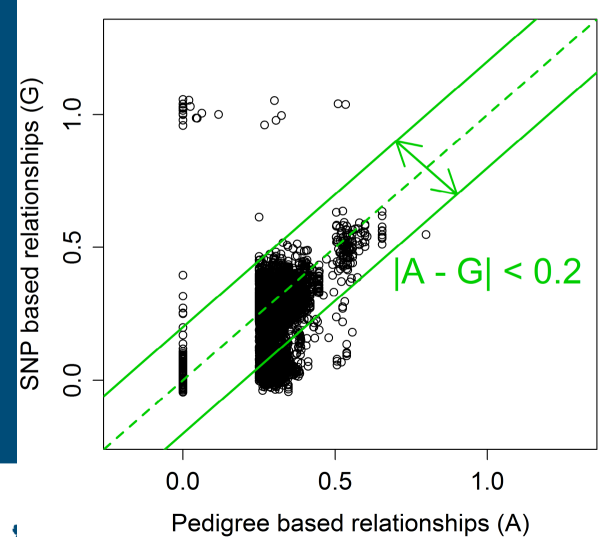
SIBCOUNT

A. – Unrelated animals



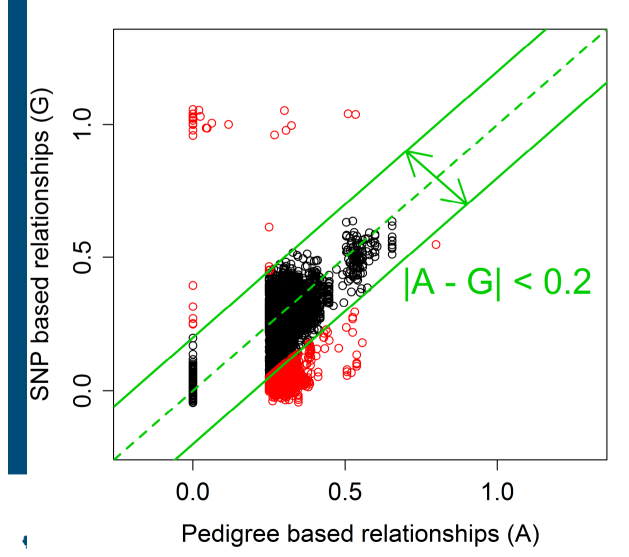
SIBREL: relationships & thresholds

SIBREL



SIBREL: relationships & thresholds

SIBREL



<u>Results</u>

Removed animals:

223 based on parent-offspring inconsistencies
31 (34) based on SIBCOUNT (SIBREL)

After permuting 10% of the data:
Incorrectly deleted animals (Type I error)
Equally low for SIBCOUNT & SIBREL (~ 0.5%)

Incorrectly non-deleted animals (Type II error)
 SIBCOUNT (5.2%) << SIBREL (9.8%)





Conclusions

 SIBCOUNT was better able to identify animals causing MI

• Lower Type II error rate

Test for opposing homozygotes is: Powerful tool to detect MI between sib pairs Easy to implement

Easy to implement







Develop innovative and practical breeding tools for improved dairy products from more robust dairy cows



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