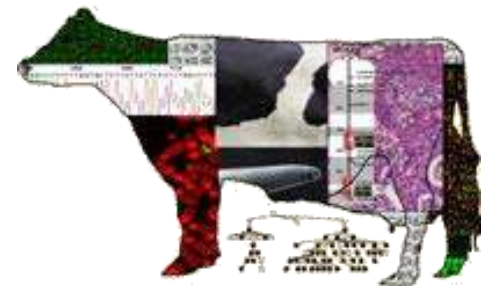


# Copy number variation of individual cattle genomes using Next Generation Sequencing



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# Creating CNV maps with NGS data



Background and Introduction



Individualized CNV maps



Detection within populations



# Background and Introduction

# Genetic Variation

## How genomes change over time

*Sequence*

- Single nucleotide variations – SNP (human 60 million)
- Small insertions/deletions – frameshift, microsatellite, minisatellite
- Mobile Elements – SINE, LINE Transposition (300bp - 6 kb)
- Genomic structural variation (1 kb – 5 Mb)
  - Large-scale Insertions/Deletions (Copy Number Variation: CNV)
  - Segmental Duplications (> 1kb, > 90%)
  - Chromosomal Inversions, Translocations, Fusions.

*Cytogenetics*

# CNVs contribute to phenotype



Picture from Wright et al. 2009. PLoS Genet.

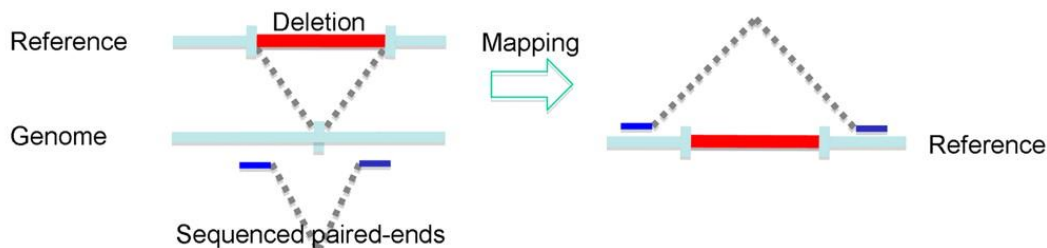


Picture from Seo et al. 2007. BMC Genetics

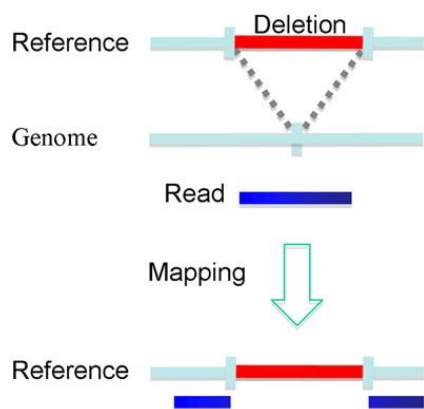


ASIP

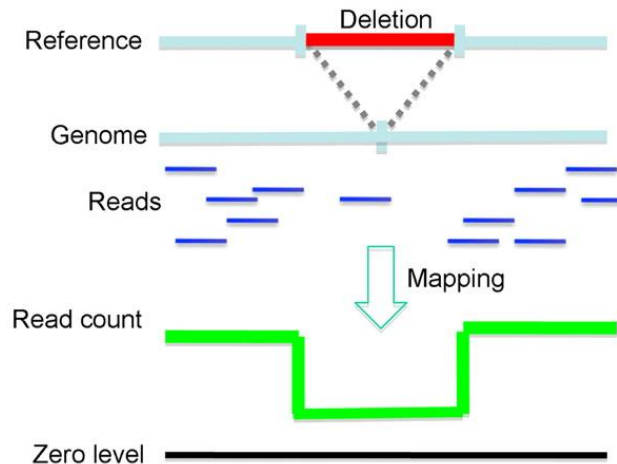
### A Paired end reads



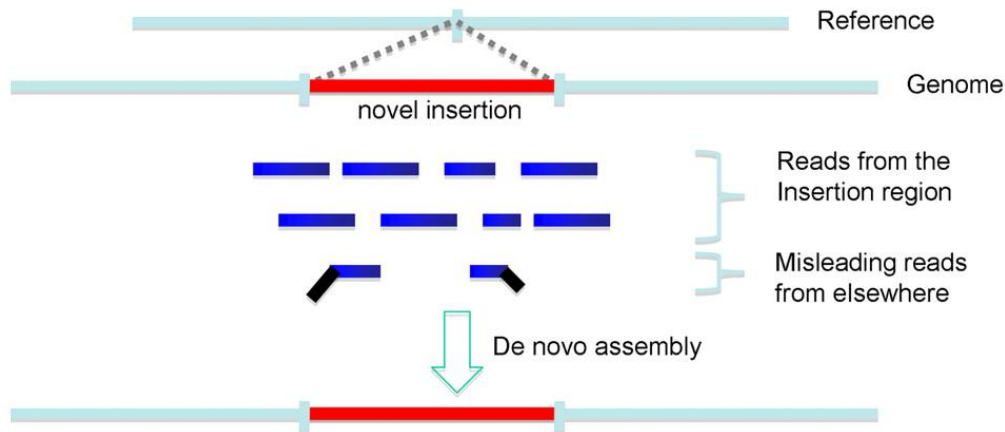
### B Split read



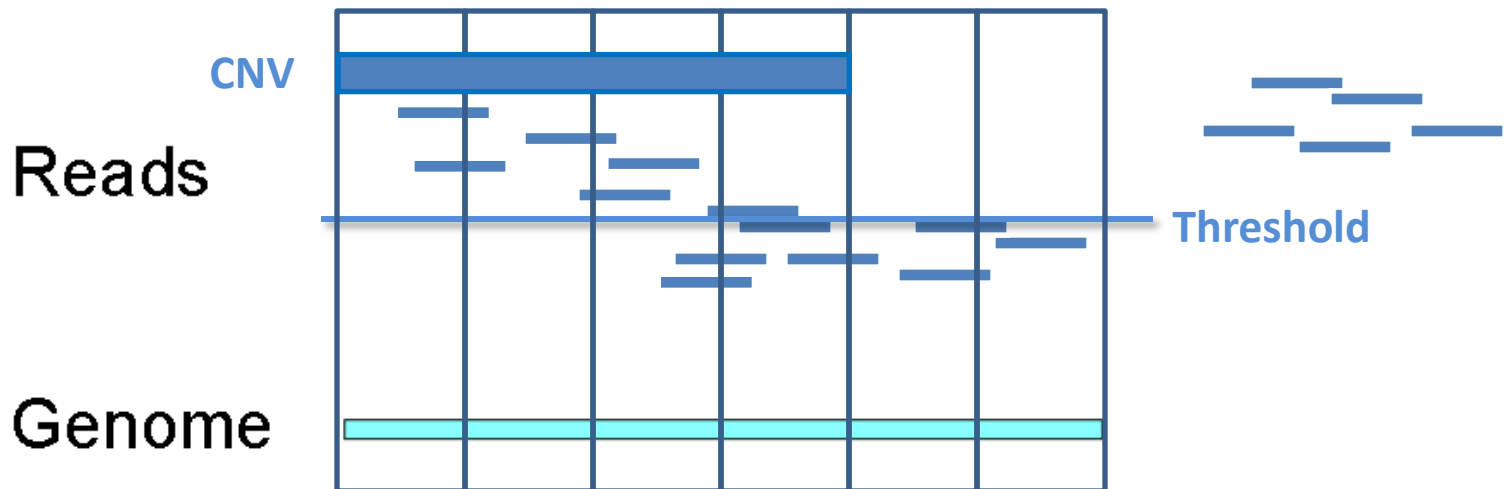
### C Read depth



### D Local Reassembly

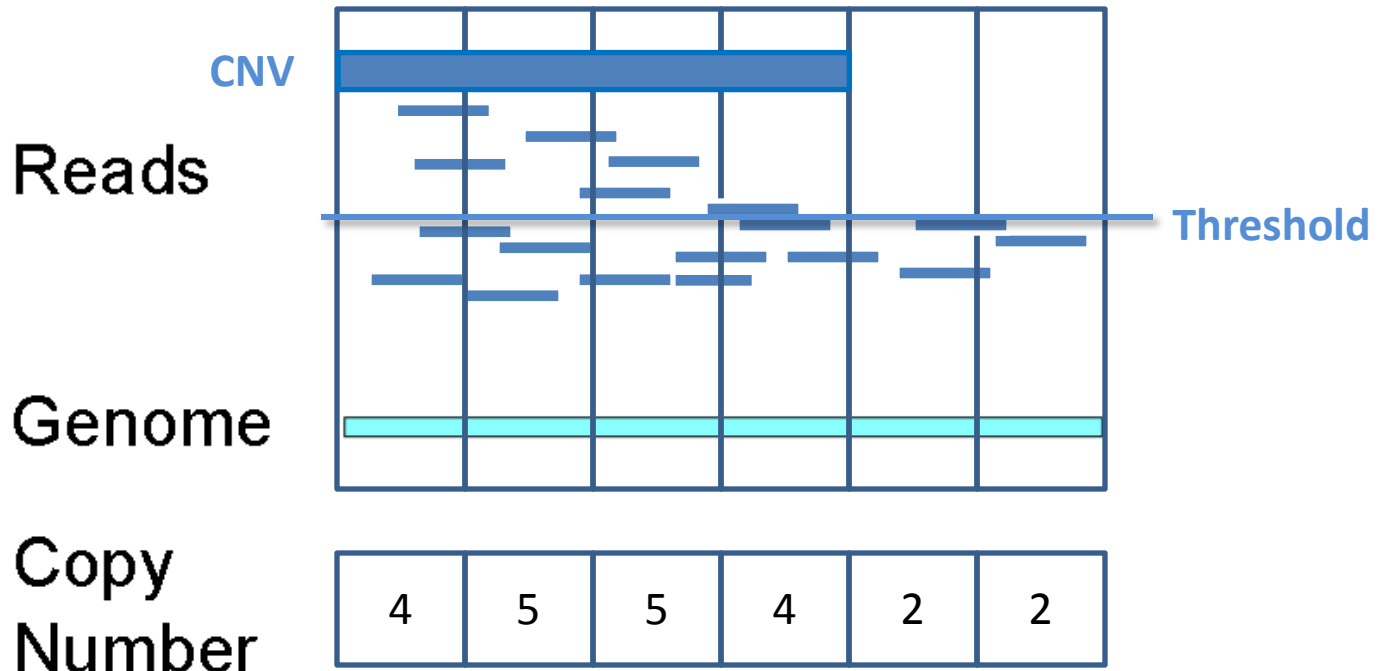


# Read Depth (RD) Detection



- Align reads to reference genome (used mrsFAST)
- Determine average RD and set Threshold
  - Threshold value: average + 4 Stdev
  - Normalized for GC bias
- Section genome into windows and call CNVs
- Analysis can be done in SD regions

# RD provides genome-wide Copy Number (CN)



Advantages of this approach:

- Assign CN to Gene Regions
- CN is not relative to other samples
- Works on all reference assemblies





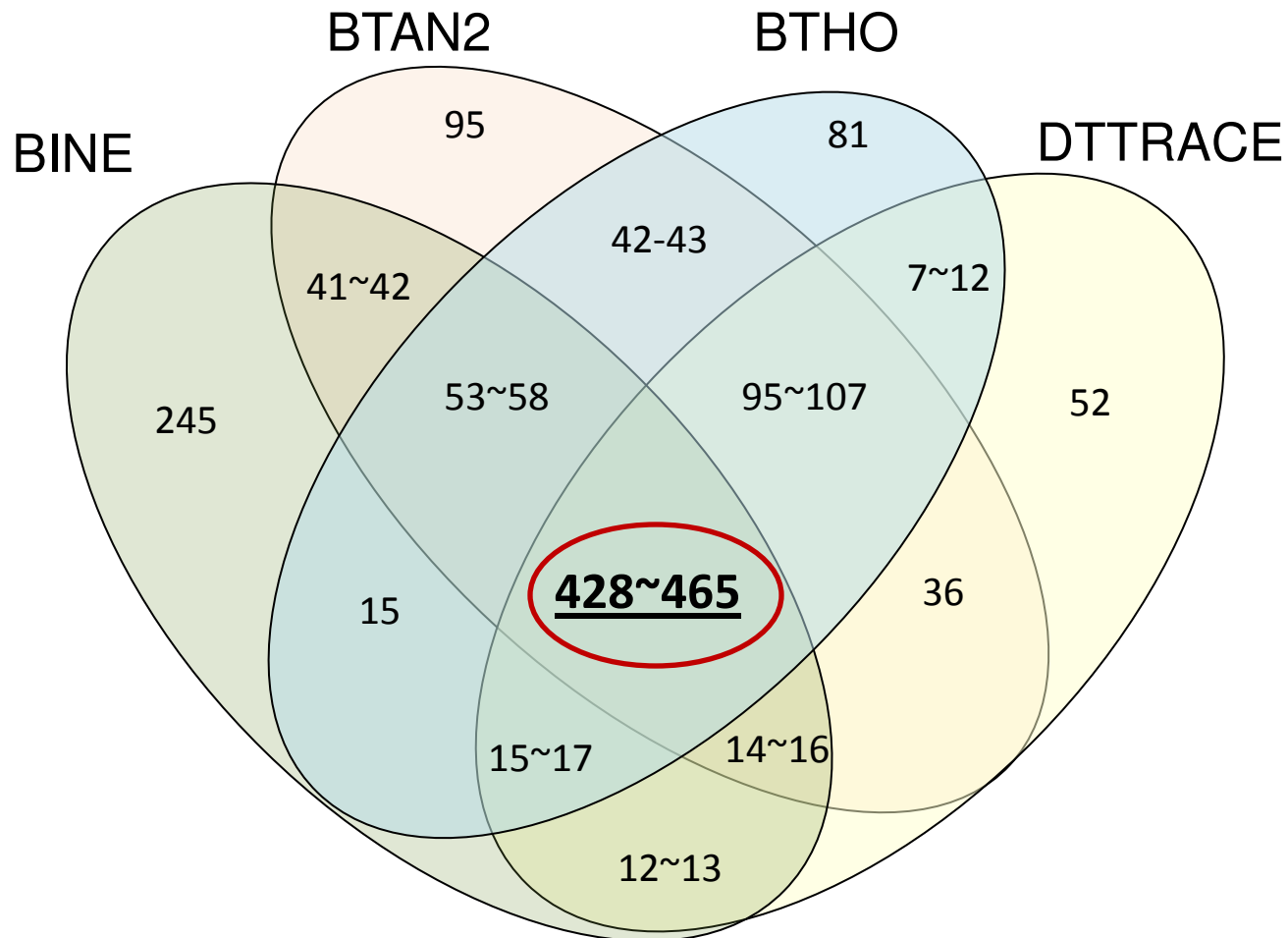
# Individualized CNV maps

# Animals Selected

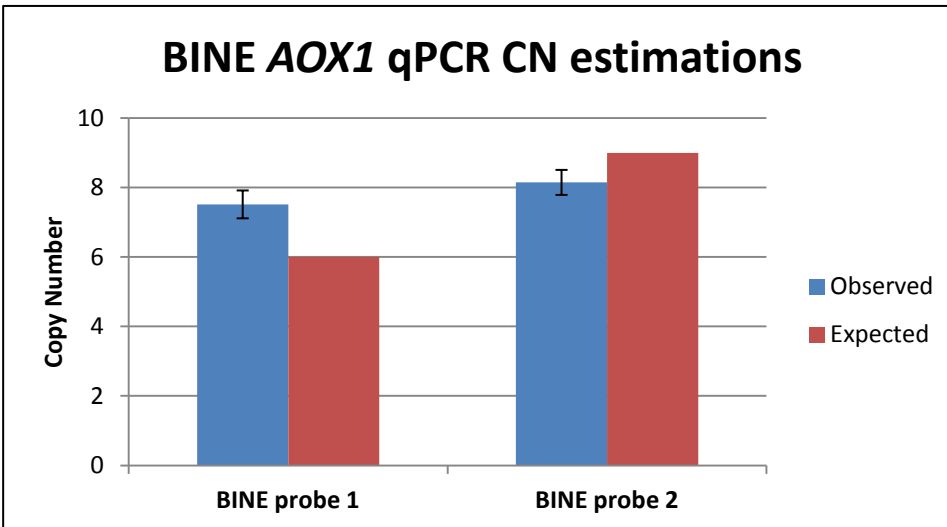
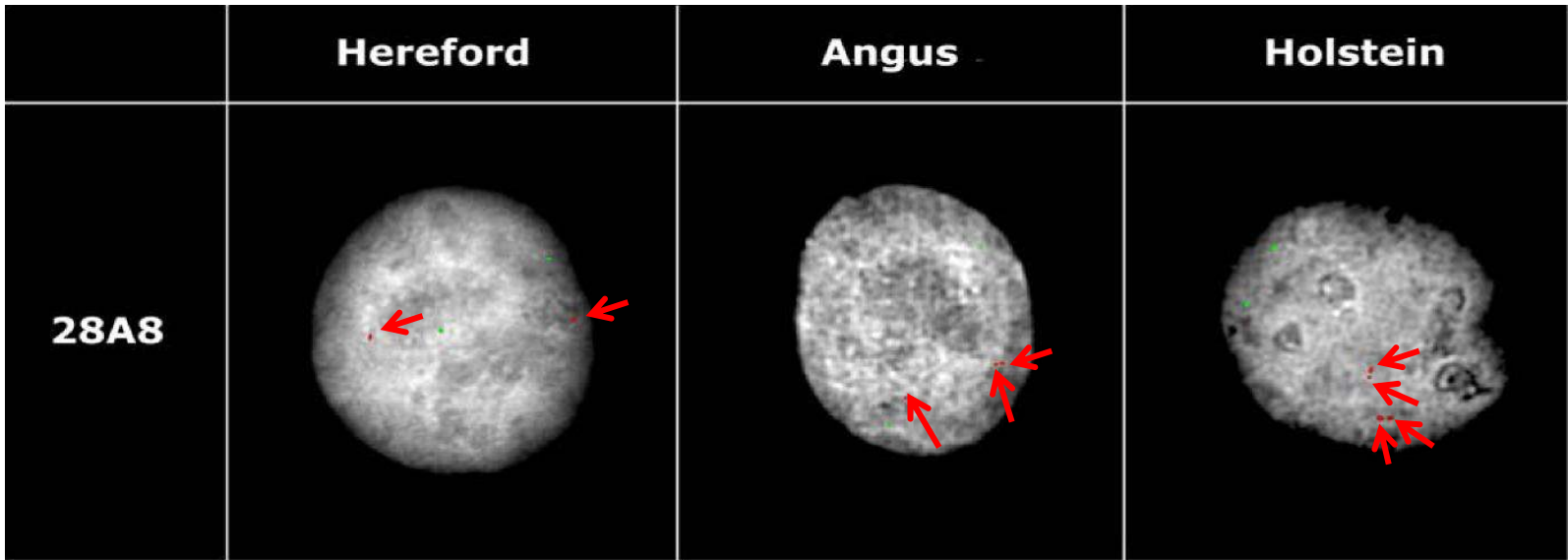
Animal Abbreviation	Breed	Coverage
BINE	Nelore	High
BTAN1	Angus	High
BTAN2	Angus	High
BTAN3	Angus	High
BTHO	Holstein	Low
DTTRACE	Hereford	Low

Reference animal

# CNV events more diverse than in humans



# Experimental validation

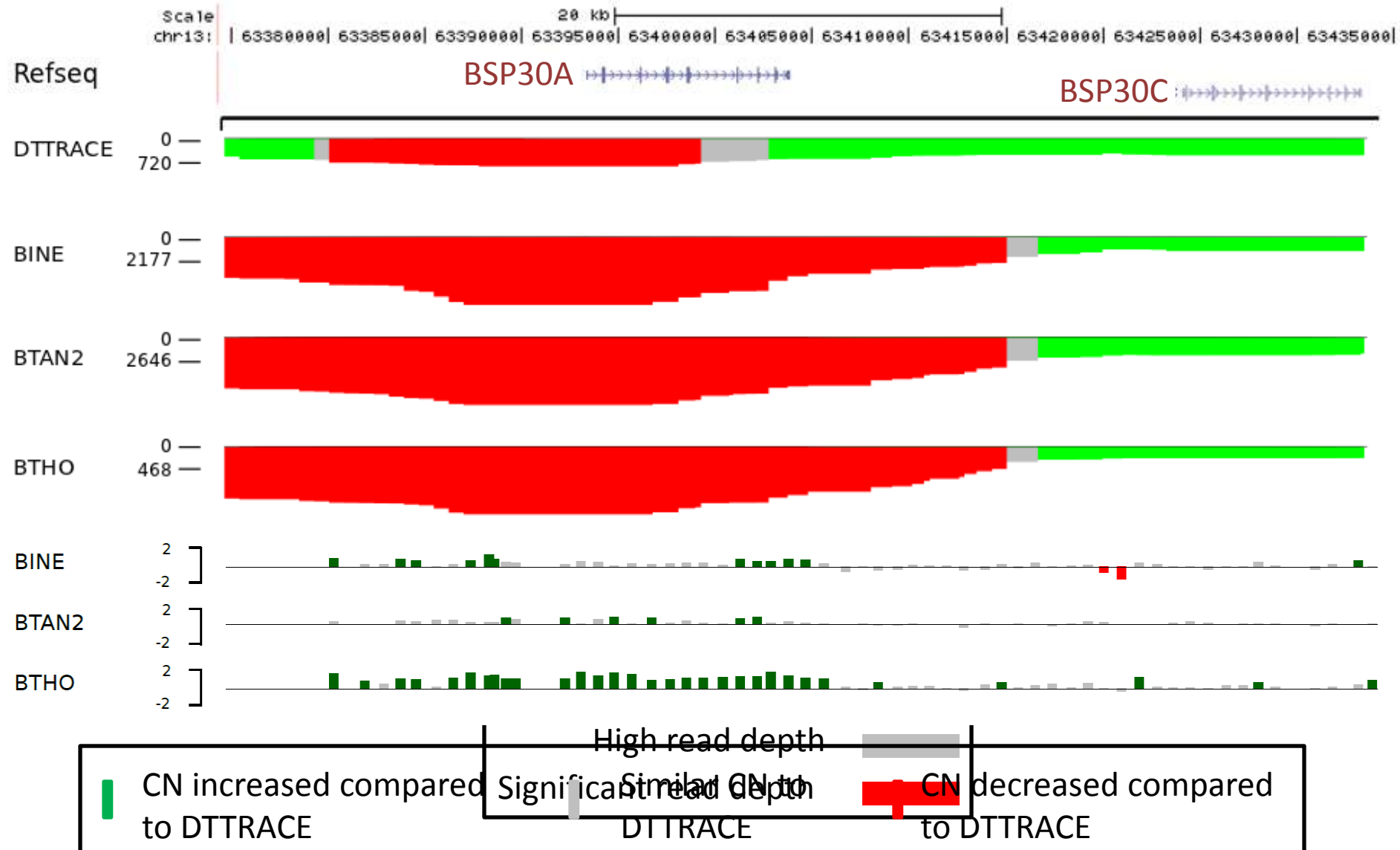


- ~55% agreement with BAC FISH

- 82% agreement with qPCR assays

- 8% false discovery compared to aCGH

# BSP30A is duplicated in all breeds



# CN estimates give better resolution

Gene ID	BINE	BTAN1	BTAN2	BTAN3	BTHO	DTTRACE
<del>CATH14</del> BSP30A	7.9	9.7	8.7	7.5	11.8	3.6

Copy Number Gradient



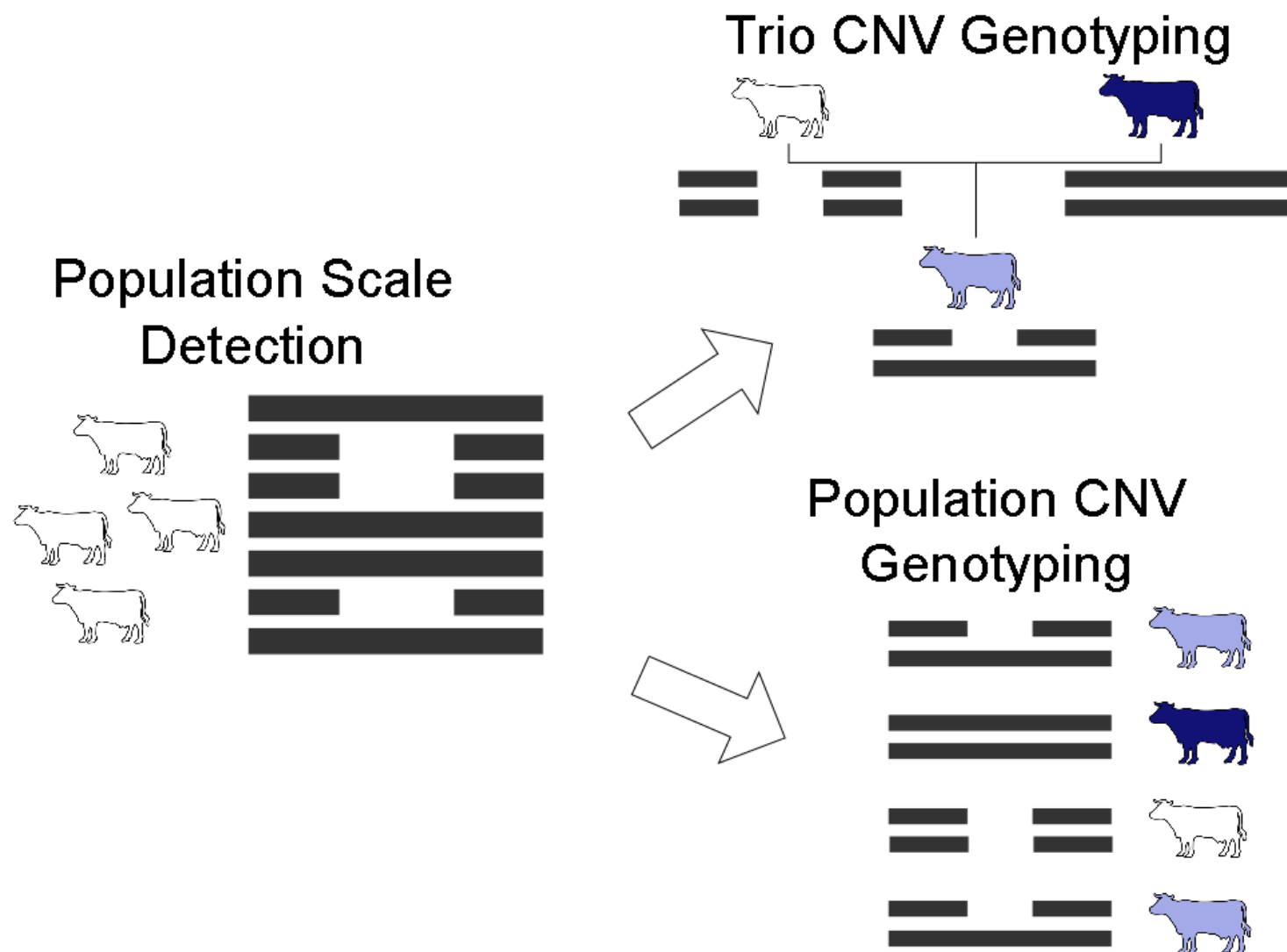




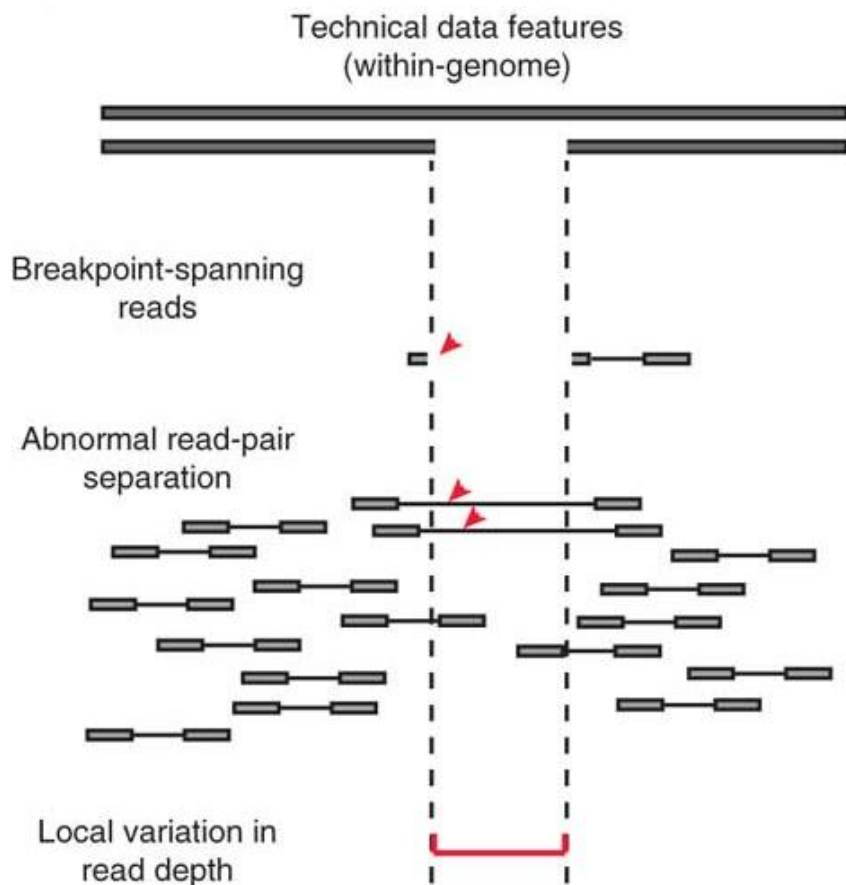
# Detection within Populations



# Expanding the sample size allows high resolution detection



# Improve analysis with multiple methods



- ❑ Multiple techniques increase detection
- ❑ RD disadvantages
  - Only homozygous deletions
  - No mobile element insertion (MEI) detection
- ❑ Genome Strip (pictured left)
  - Combines three techniques
  - Limited to Deletions
- ❑ Include methods:
  - VariationHunter v2 (MEI detection)
  - RD method (duplication detection)
- ❑ Also include:
  - SNP array data
  - CGH array data

# Summary



**Selected suitable detection strategy**



**Created high resolution individual maps**



**Transitioning to population scale analysis**

# Acknowledgements

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