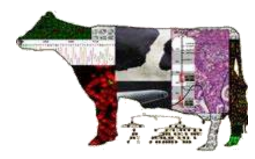
Copy number variation of individual cattle genomes using Next Generation Sequencing





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Creating CNV maps with NGS data

Background and Introduction



Detection within populations

Background and Introduction

Genetic Variation

Sequence

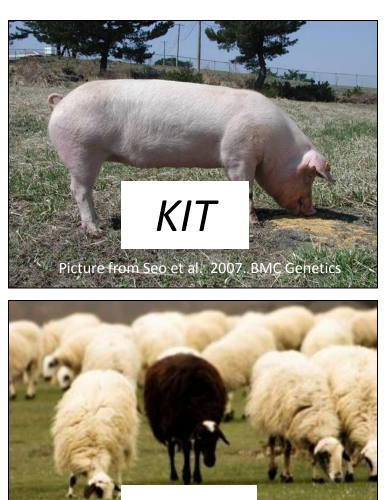
How genomes change over time

- 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.

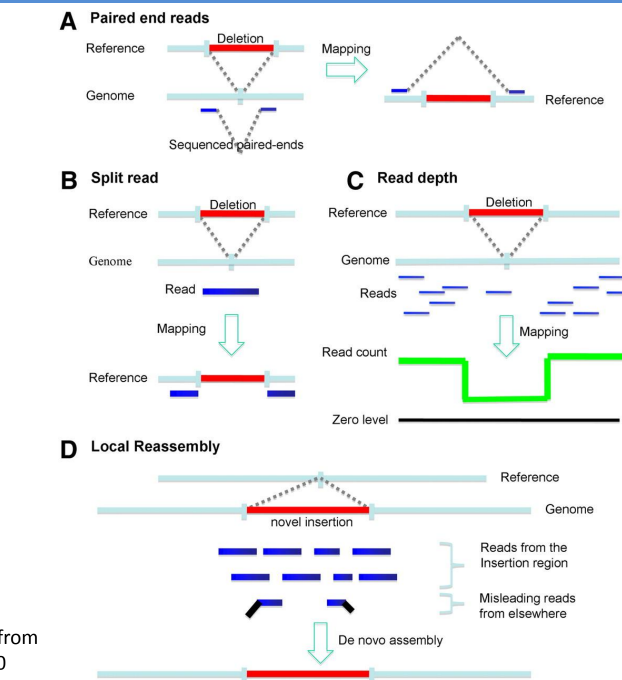
Cytogenetics

CNVs contribute to phenotype



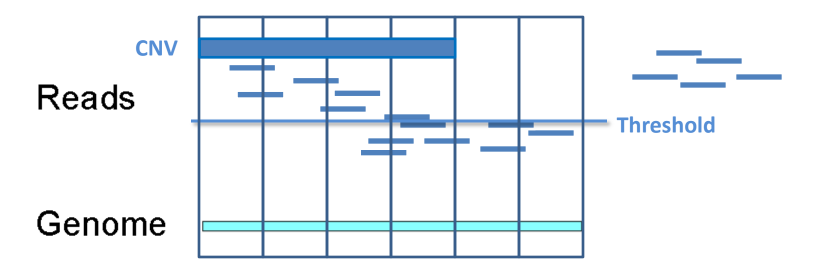


ASIP



Figures Adapted from Snyder et al, 2010

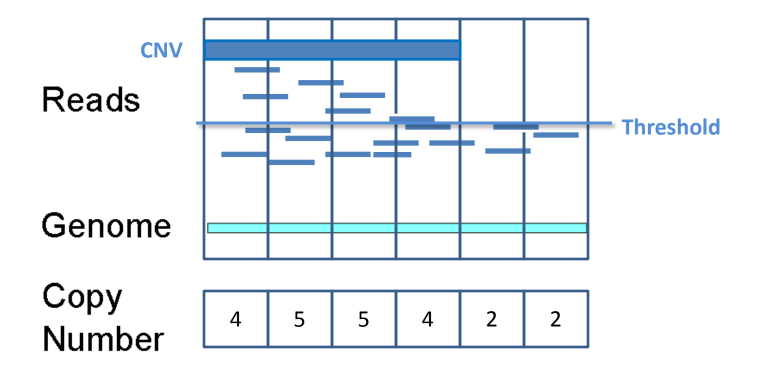
Read Depth (RD) Detection



- 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

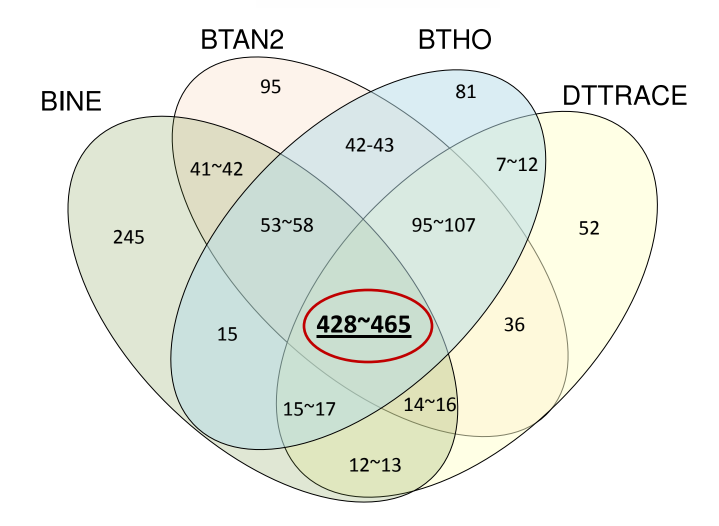
Individualized CNV maps

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

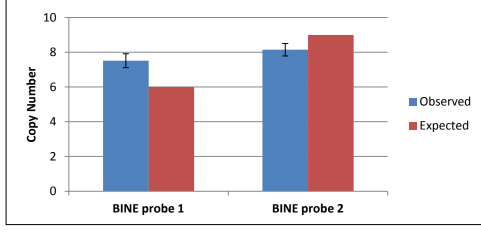
CNV events more diverse than in humans



Experimental validation

	Hereford	Angus	Holstein
28A8	.4		



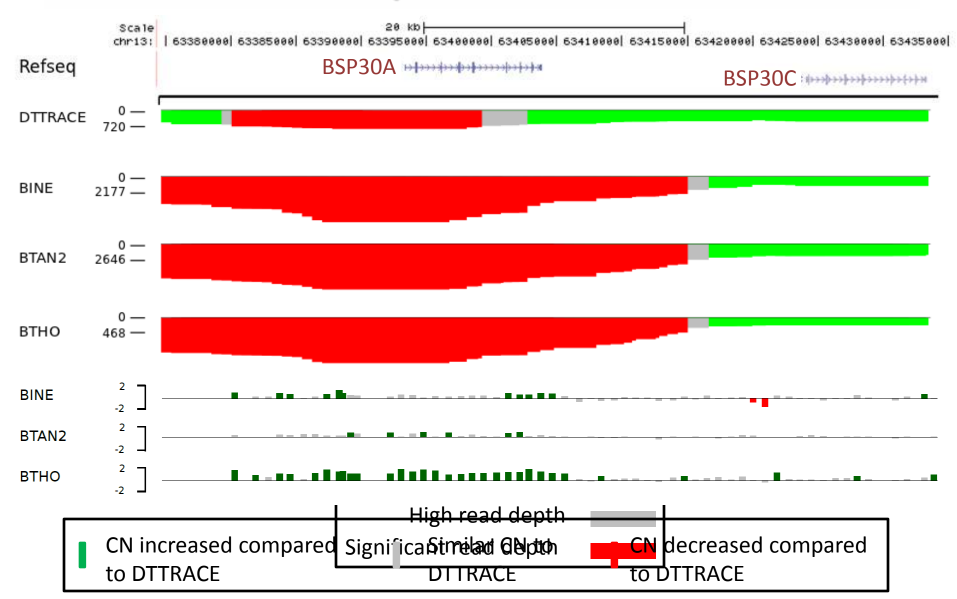


~55% agreement with BAC FISH

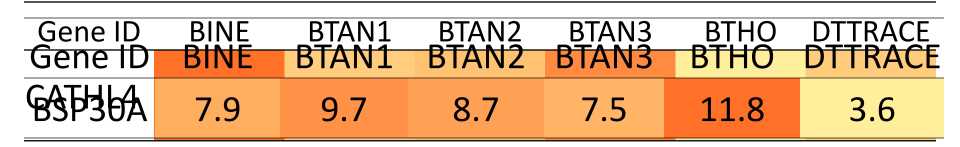
82% agreement with qPCR assays

• 8% false discovery compared to aCGH

BSP30A is duplicated in all breeds



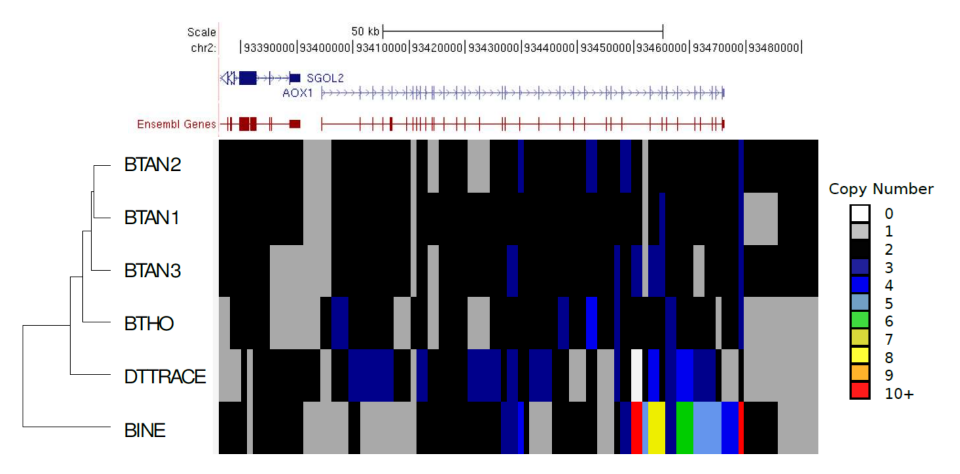
CN estimates give better resolution



Copy Number Gradient

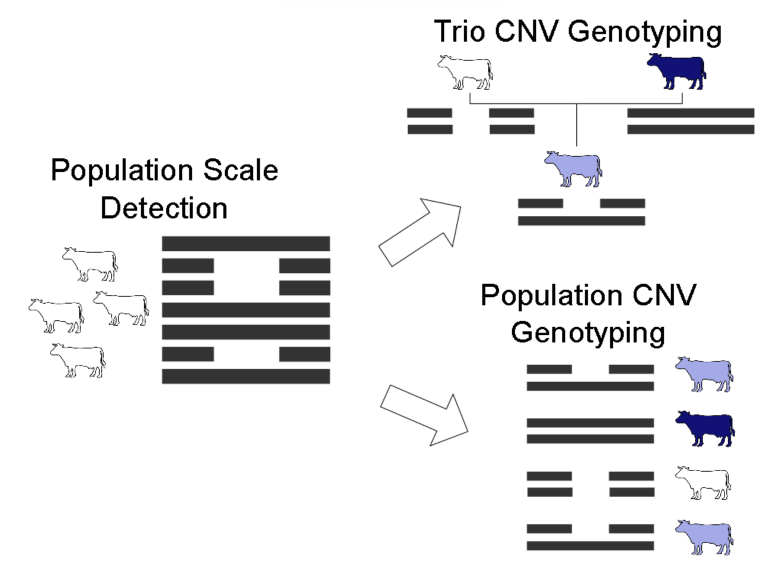


Breed differences can be highlighted

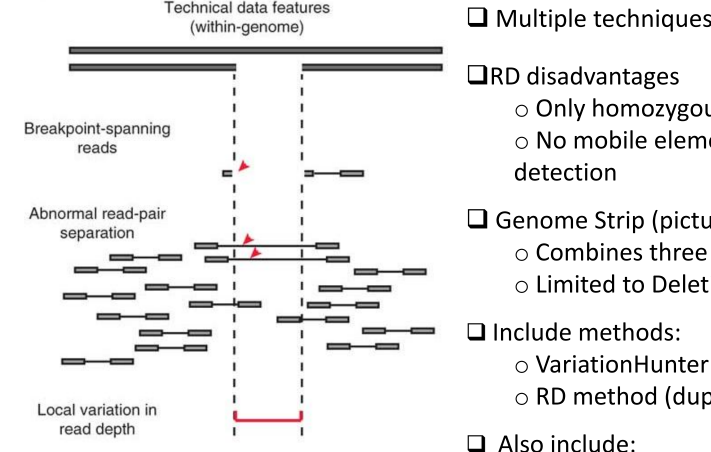


Detection within Populations

Expanding the sample size allows high resolution detection



Improve analysis with multiple methods



Images adapted from Handsaker et al. 2010

Multiple techniques increase detection

- Only homozygous deletions \circ No mobile element insertion (MEI)
- Genome Strip (pictured left)
 - Combines three techniques
 - Limited to Deletions
 - \circ VariationHunter v2 (MEI detection) • RD method (duplication detection)
- SNP array data ○ CGH array data

Summary

Selected suitable detection strategy

Created high resolution individual maps

Transitioning to population scale analysis

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