



Molecular analysis of White Galloway coat colour variations

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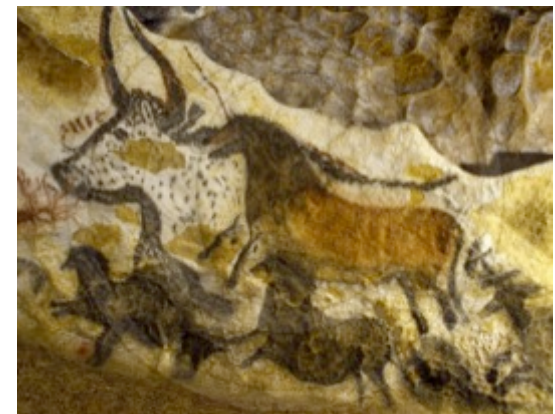
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Coat colour of livestock

- Coat colour variations have always been fascinating for man
- In early mythology and art coat colour of animals played an important role
(Siebel, 1997; Forbis, 1980)
- Coat colour variations have been one of the earliest selection criteria
(Schmutz, 2002)



(www.onlinekunst.de, Das Rind in der Kunst)



(www.lascaux.culture.fr)

Coat colour variations in White Galloway and White Park cattle



wsg

white, well marked

wsü

white, strongly marked

wss

white, mismarked

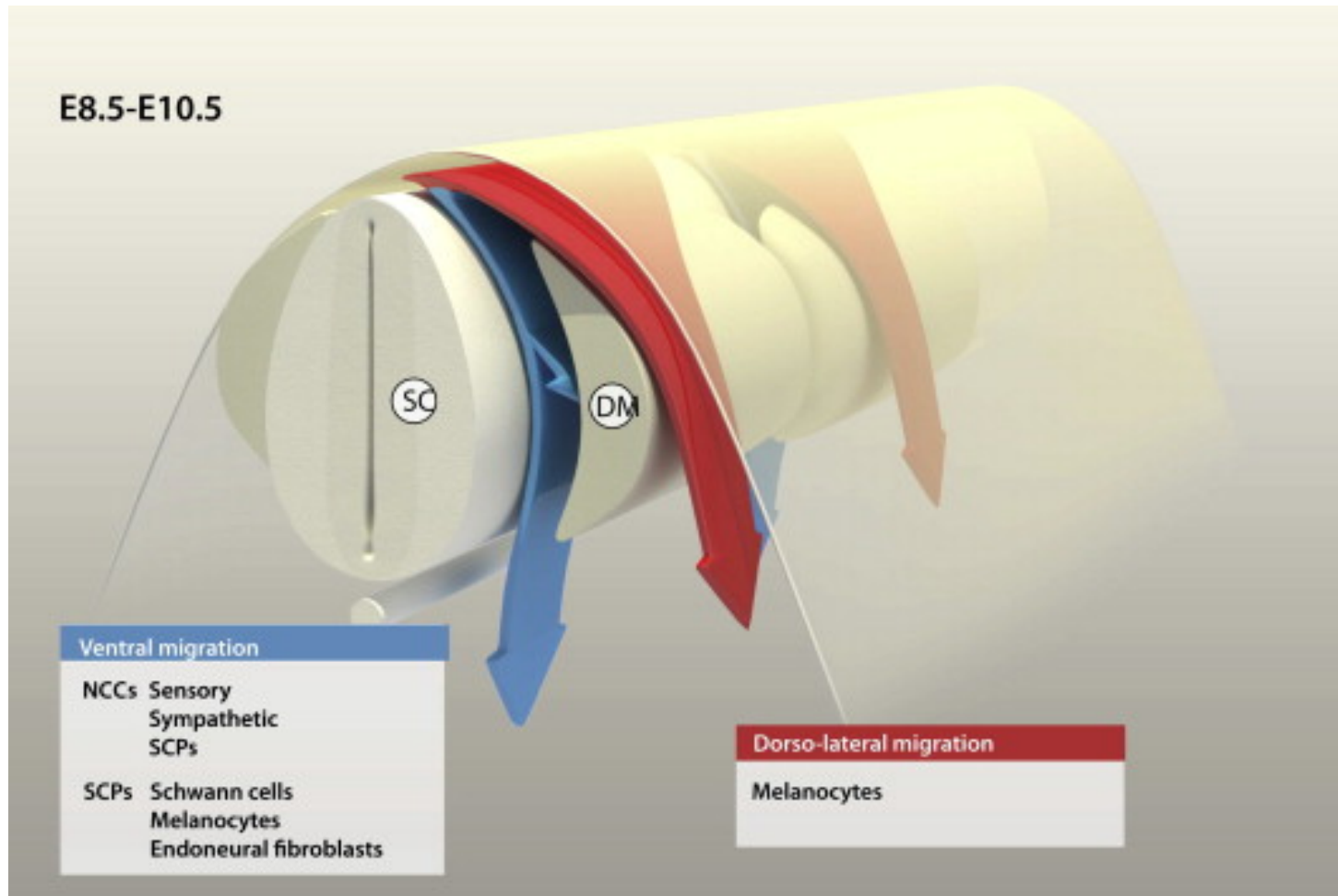
wsch

white, fully black

Aims of the study

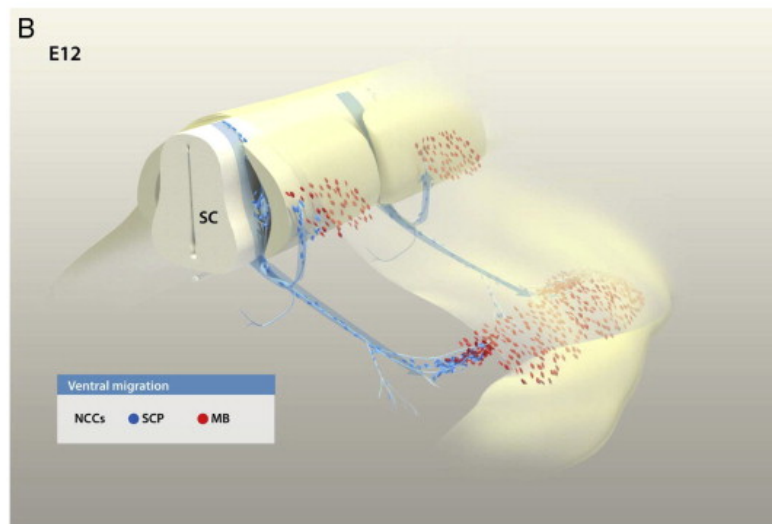
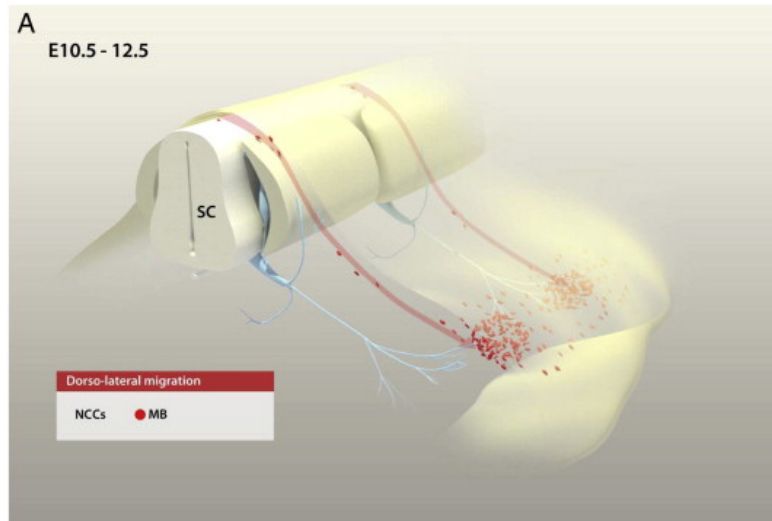
- Elucidation of the genetic background of the different coat colour variations
- Identification of DNA variants associated with the coat colour variations
- Development of a DNA –based test for breeding

Coat colour pathways



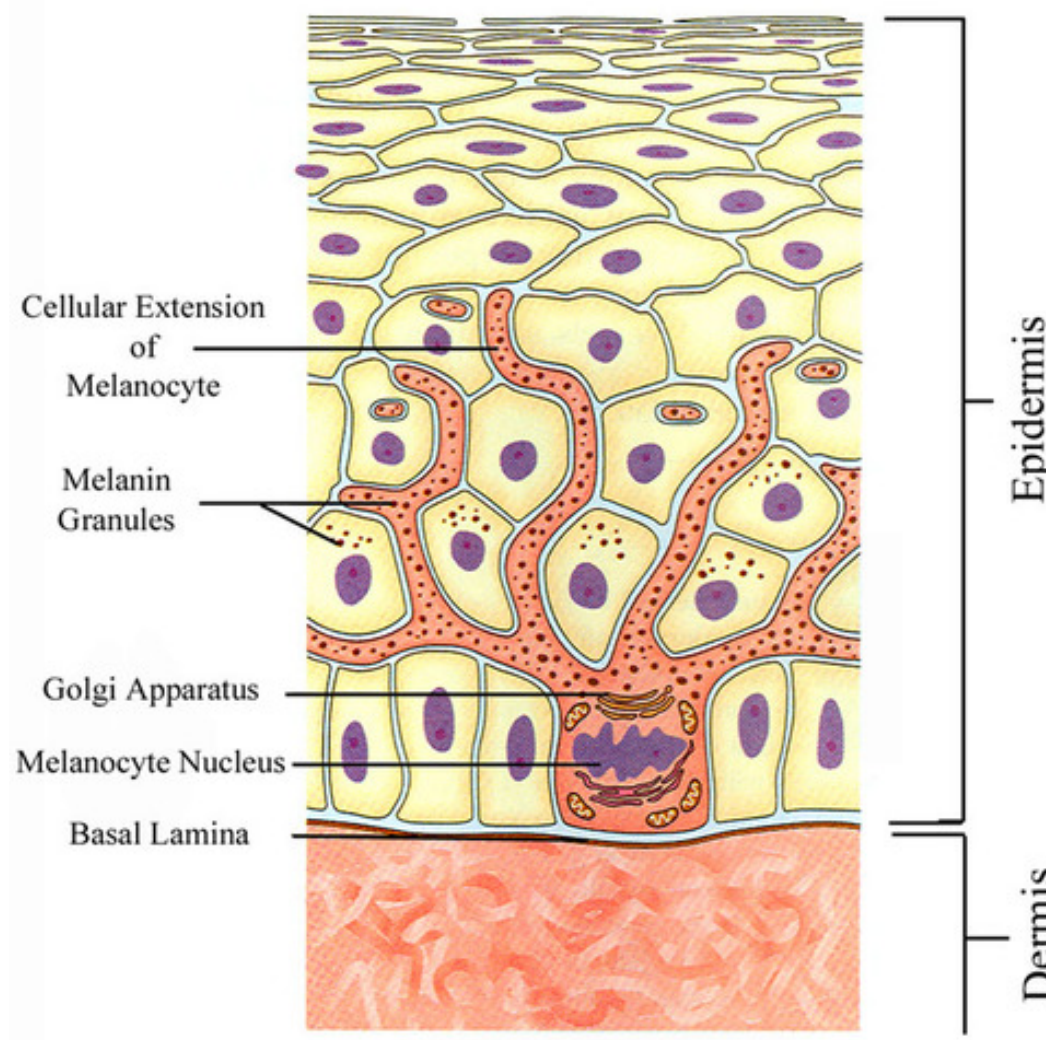
(Ernfors (2010) Exp Cell Res 316, 1397-1407)

Coat colour pathways

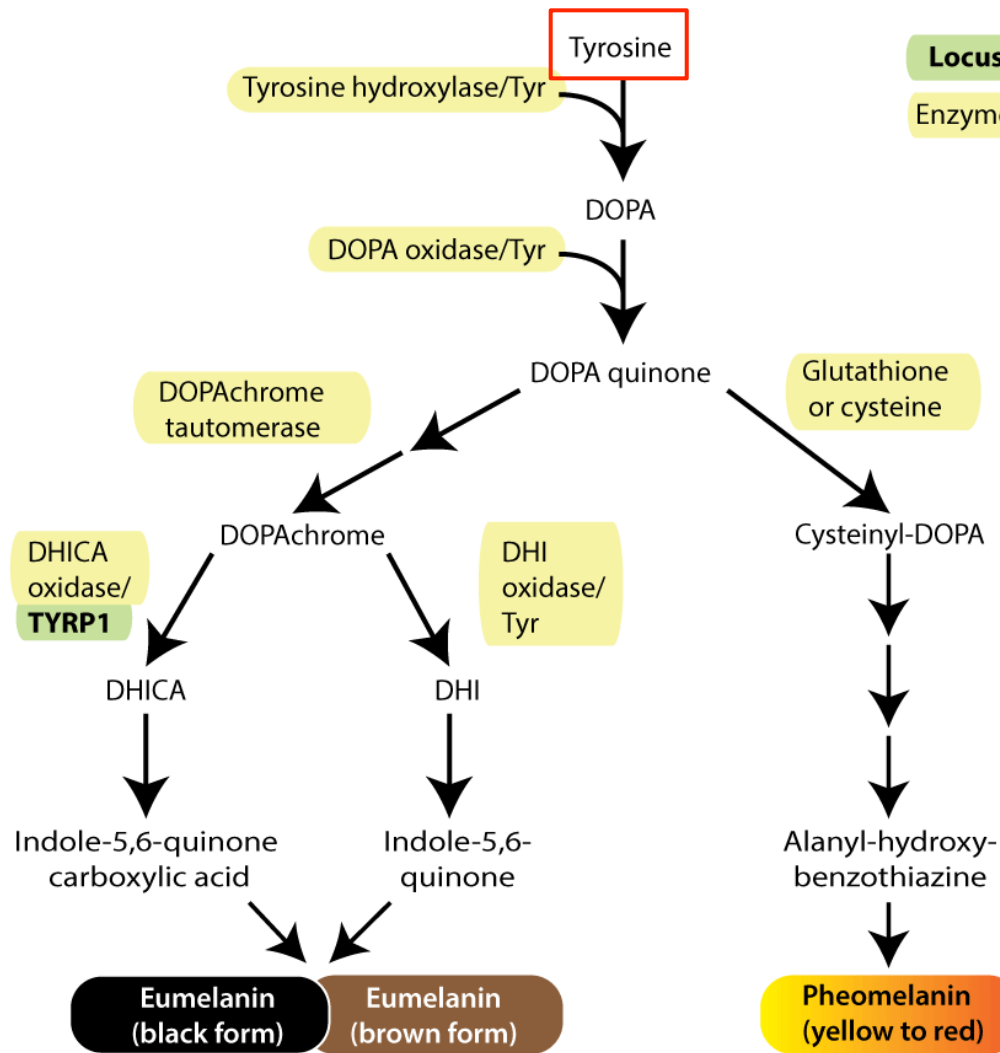


(Ernfors (2010) Exp Cell Res 316, 1397-1407)

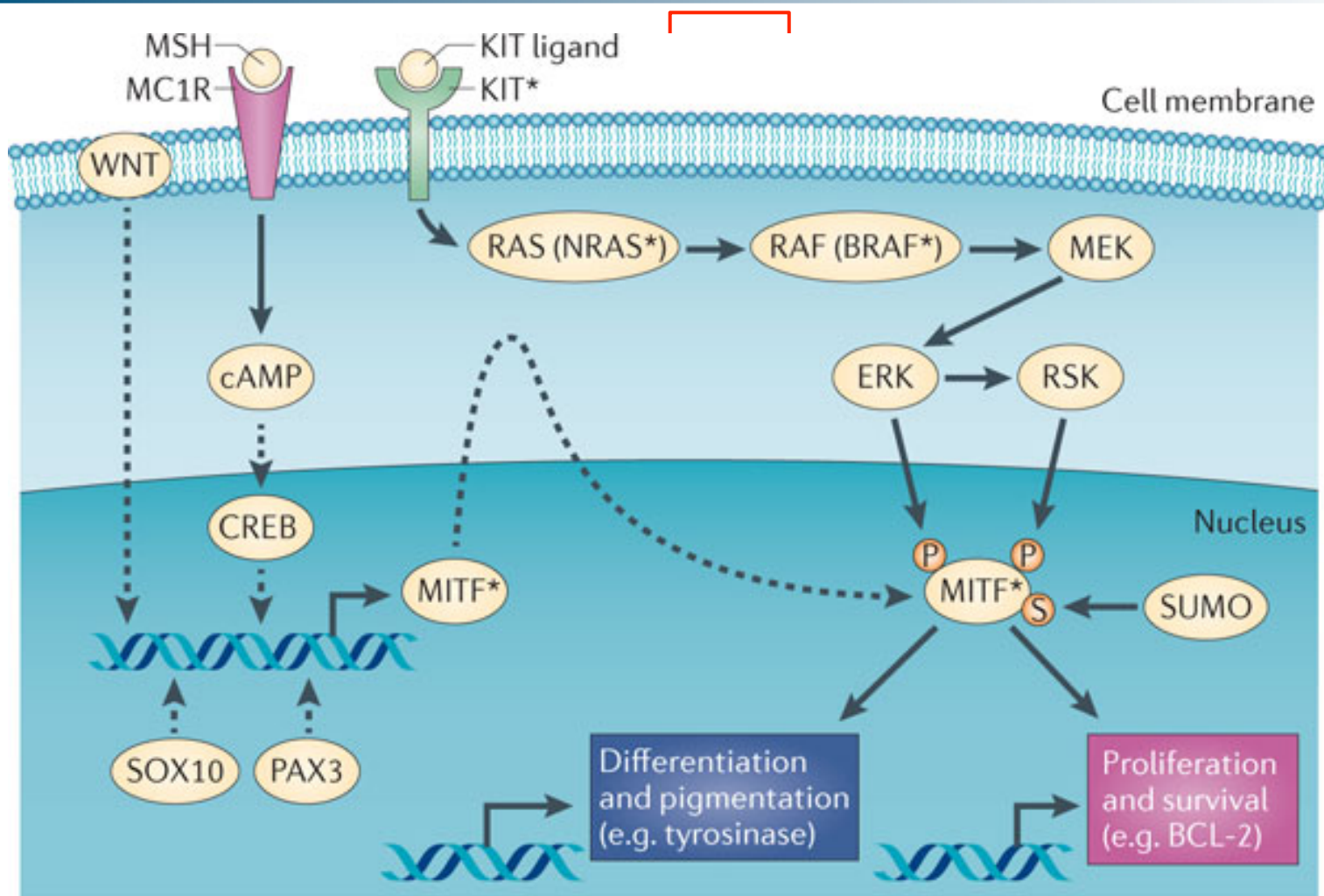
Coat colour pathways



Coat colour pathways



Coat colour pathways



Coat colour genetics

- Different genes regulate
 - Melanin synthesis
 - Timing of pigmentation
 - Pigment deposition in hairs
 - Pigment distribution in the skin
- Gene interactions modify basic colours
- > 120 genes for coat colours are known in mice

Molecular genetics of bovine coat colours

- 8 genes and corresponding alleles have been described:
 - **KIT** (Mast/stem cell growth factor receptor): Colour-sided, White spotting
 - **KITLG** (KIT ligand): Roan
 - **MC1R** (Melanocortin 1 receptor): Extension
 - **MITF** (Microphthalmia-associated transcription factor): Dominant White, Piebaldism
 - **TYR** (Tyrosinase): Albinism
 - **ASIP** (Agouti signaling protein): Agouti
 - **PMEL** (Premelanosome protein): Dilution
 - **TYRP1** (Tyrosinase-related protein 1): Brown



(Quelle: Durkin et al., 2012, S. Schmutz, 2010, RSH eG)

Identification of causative genes in White Galloway cattle

- Candidate gene analysis *KITLG*, *KIT*, *TYR* and *MITF*
 - 3 animals of each phenotype
 - Exons, splice-donor/acceptor-sites, UTR
 - Identification of functional SNPs
 - Genotyping of all animals (n=179)
- Genotyping of the *MC1R* locus
 - E^D, E⁺, e
 - n=184

Results of candidate gene-analysis

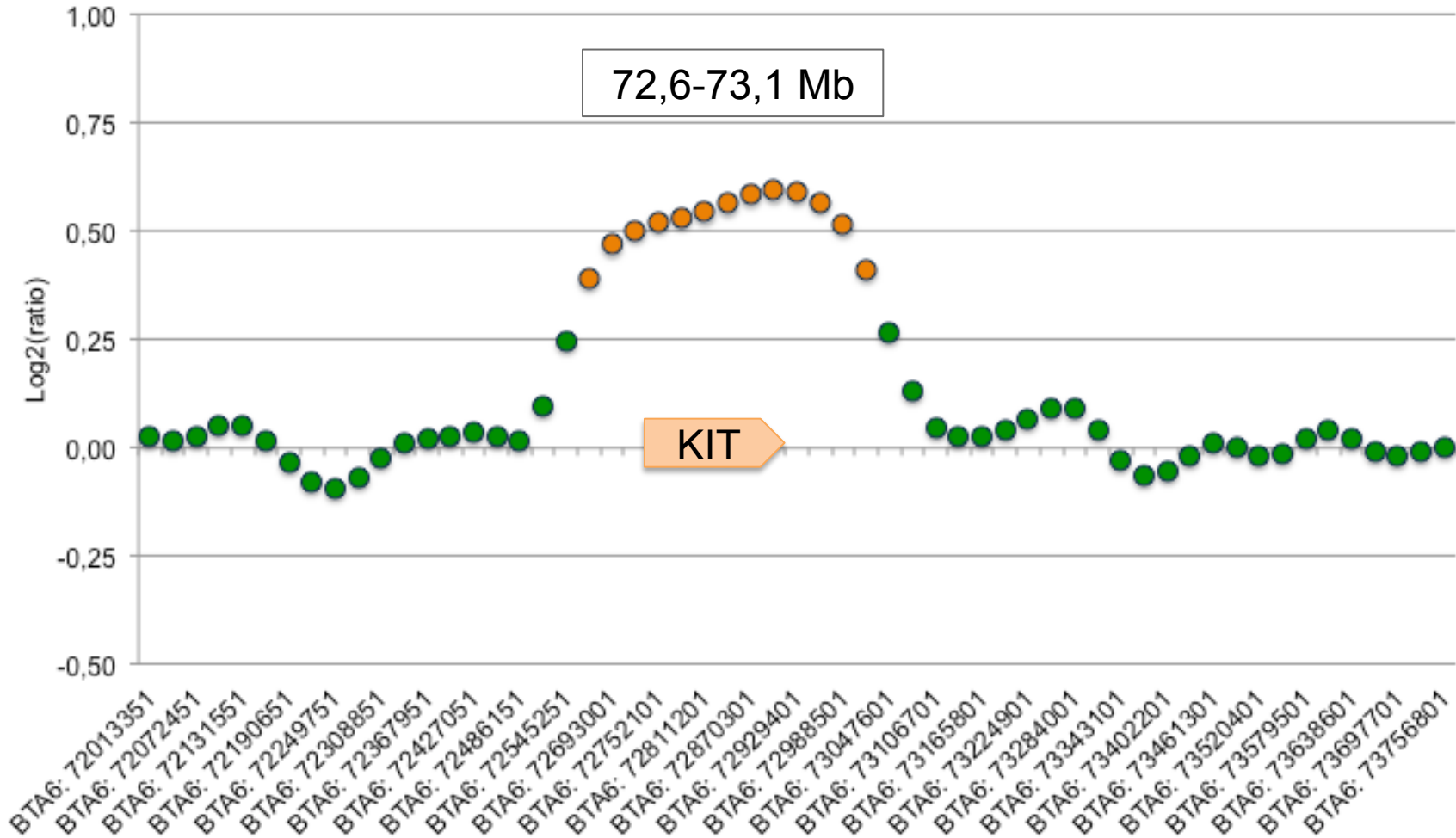
| Gene | Position | Polymorphism | Protein | Genotype | Number | | | |
|--------------|----------|-------------------|---------|----------|-------------|-------------|------------|------------|
| | | | | | wsch (n=27) | wsg (n=102) | wsü (n=10) | wss (n=40) |
| <i>KITLG</i> | 5' UTR | g.221755T>C | | T/T | 12 | 60 | 4 | 24 |
| | | | | T/C | 12 | 38 | 4 | 13 |
| | | | | C/C | 3 | 3 | 2 | 3 |
| | 5' UTR | g.221761G>A | | G/G | 12 | 60 | 4 | 24 |
| | | | | G/A | 12 | 38 | 4 | 13 |
| | | | | A/A | 3 | 3 | 2 | 3 |
| | Exon 7 | g.45568C>A (roan) | p.A193D | C/C | 27 | 102 | 10 | 40 |
| | | | | C/A | 0 | 0 | 0 | 0 |
| | | | | A/A | 0 | 0 | 0 | 0 |
| <i>TYR</i> | Exon 1 | g.66288G>C | p.R255P | G/G | 17 | 84 | 7 | 33 |
| | | | | G/C | 10 | 17 | 3 | 6 |
| | | | | C/C | 0 | 1 | 0 | 1 |
| <i>KIT</i> | Exon 5 | g.71877602T>C | p.M258T | T/T | 0 | 0 | 0* | 0* |
| | | | | T/C | 7 | 14 | 2* | 10* |
| | | | | C/C | 20 | 88 | 7* | 29* |
| <i>MITF</i> | Exon 1 | g.136G>A | p.M1I | G/G | 4* | 6* | 2* | 6* |
| | | | | G/A | 9* | 6* | 4* | 6* |
| | | | | A/A | 0* | 0* | 3* | 0* |

**KIT*: wsü (n=9), wss (n=39); *MITF*: wsch (n=13), wsg (n=12), wsü (n=9), wss (n=12)

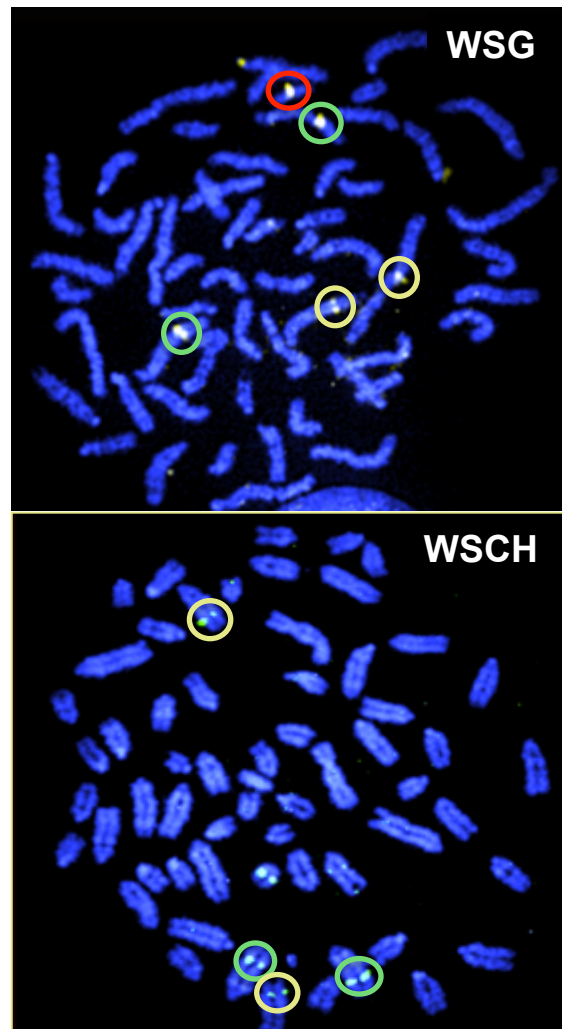
Results of genotyping

- *KITLG, TYR, KIT, and MITF*
 - SNPs within the candidate genes do not show association with White Galloway coat colours
- *MC1R*
 - 182 black Galloway cattle: e/e, E^D/E^D; e/e, E^D/E⁺
 - 2 red Galloway cattle: e/e, E^{+/E⁺}

Whole Genome Sequencing



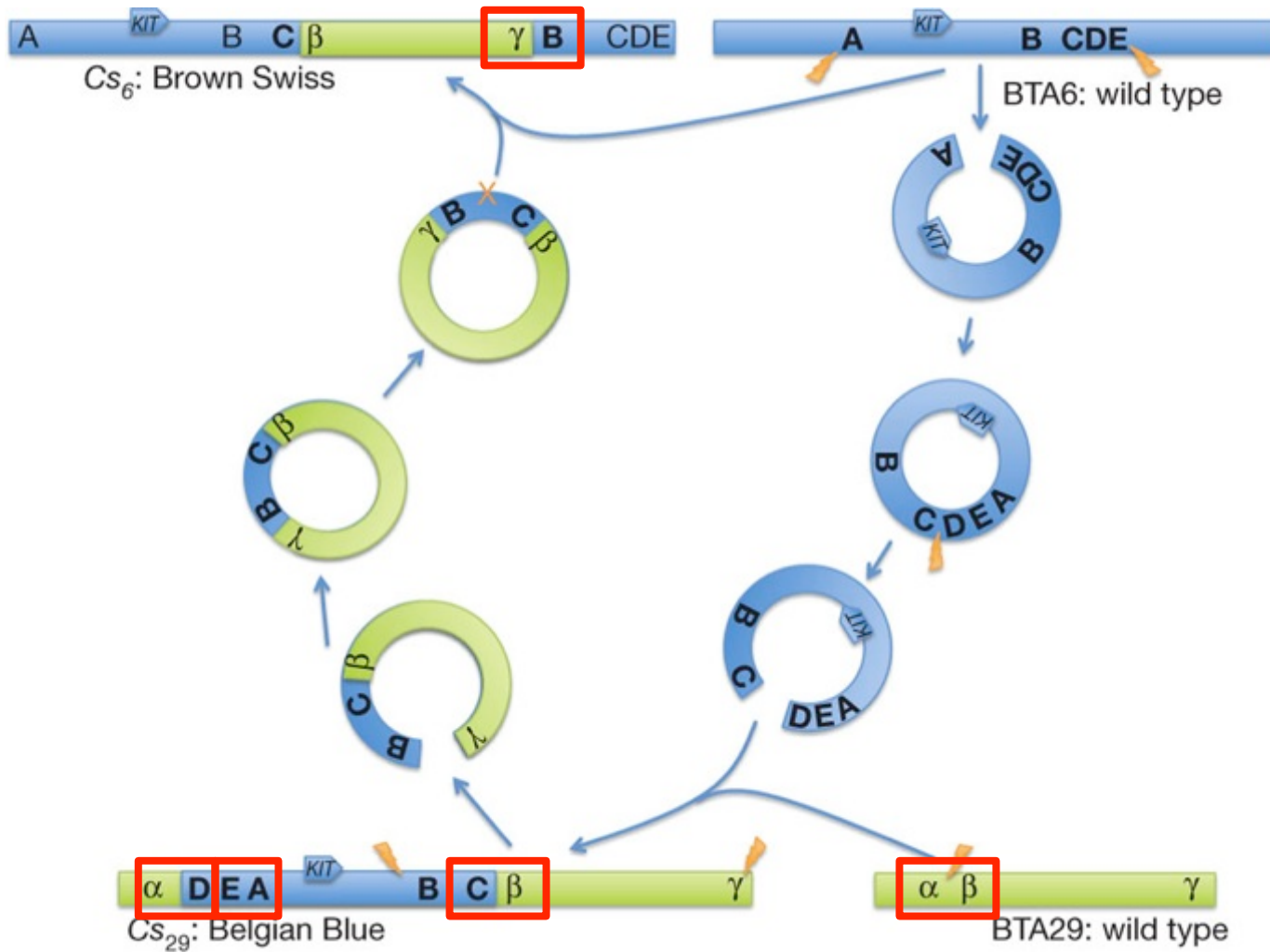
FISH analysis of White Galloway cattle



BAC harbouring *KIT* gene

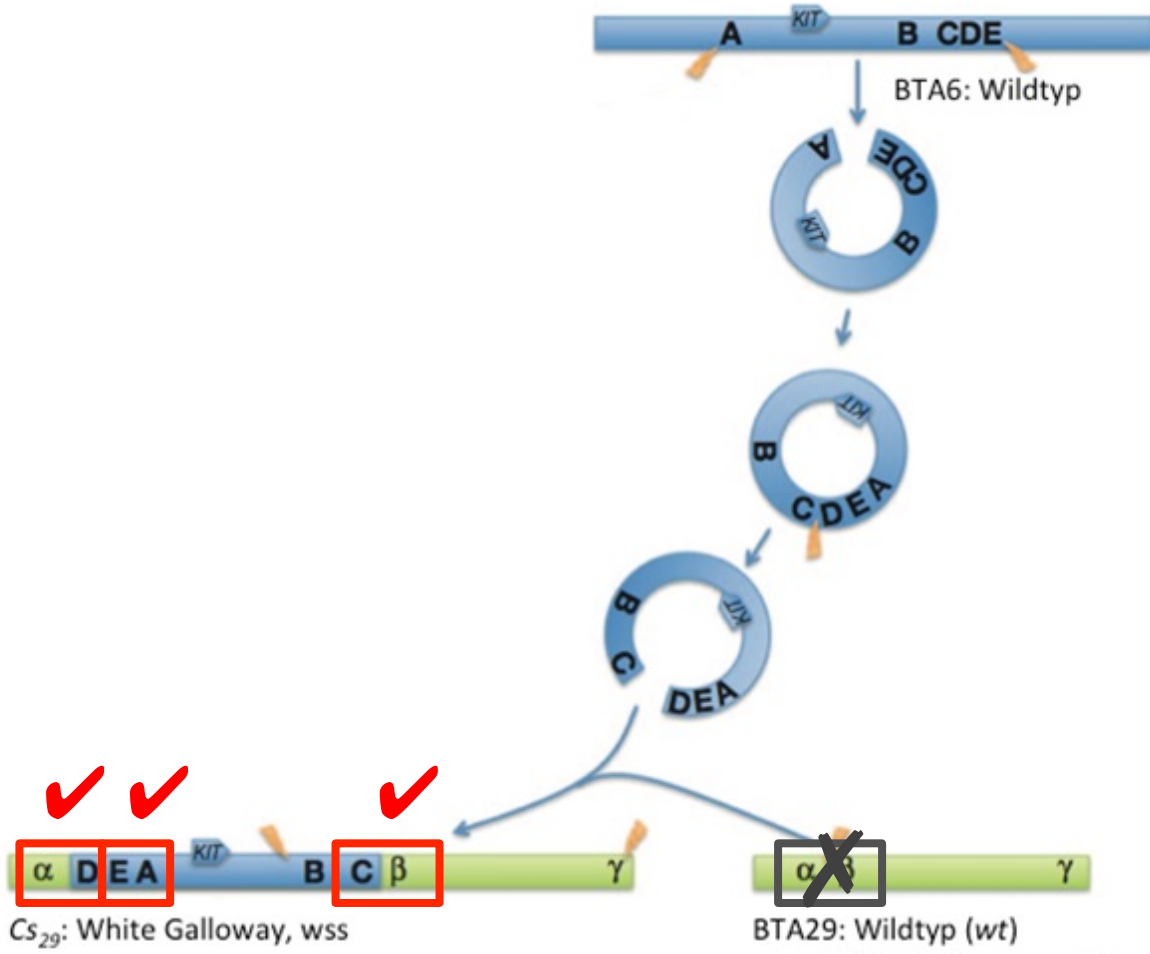
- Signals on
 - wildtype BTA6 (green)
 - translocation BTA29 (red)
 - BTA3 (yellow)
- Signals on
 - wildtype BTA6 (green)
 - BTA3 (yellow)

KIT gene translocation



(Durkin et al., 2012)

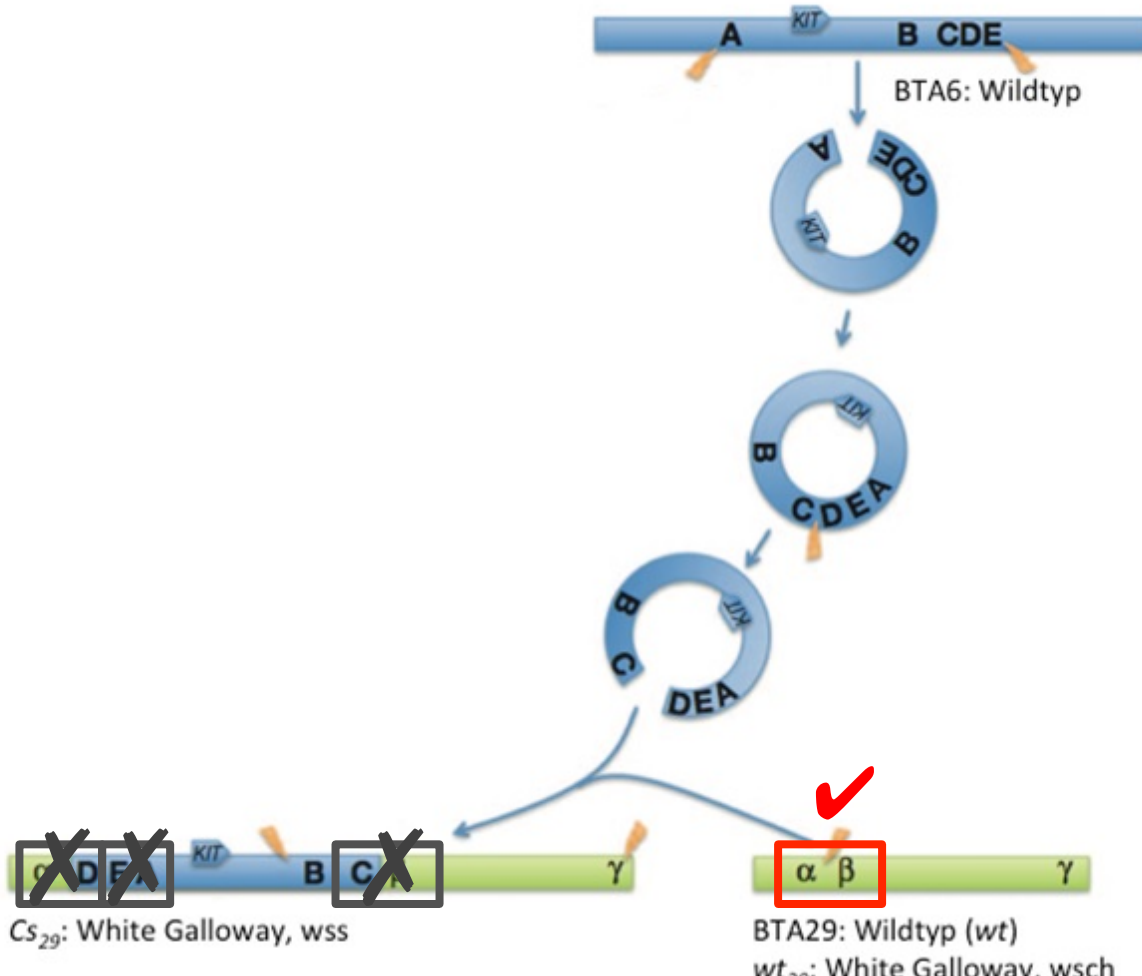
Genotype of mismarked White Galloway (wss)



homozygous
(Cs₂₉/Cs₂₉)



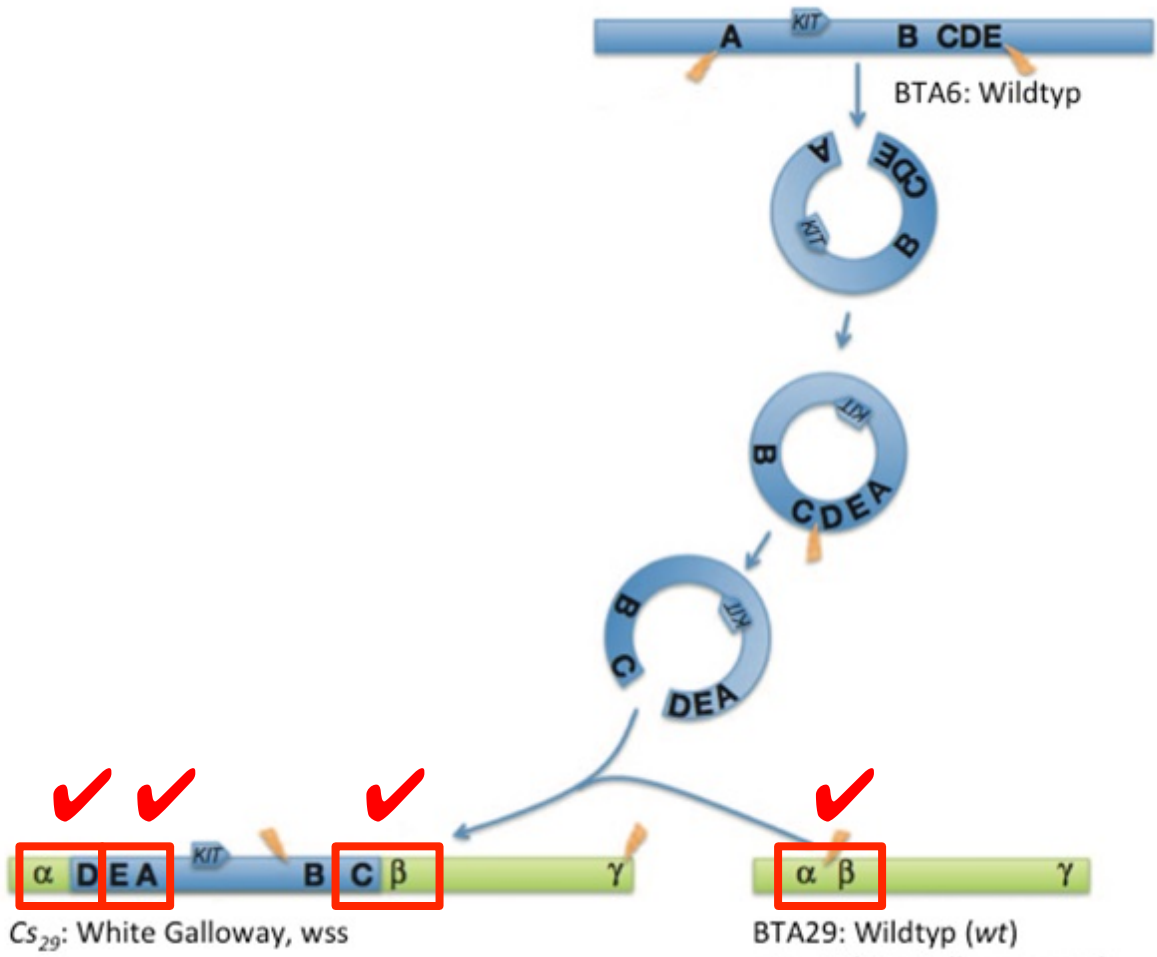
Genotype of fully black White Galloway (wsch)



homozygous
(wt_{29}/wt_{29})



Genotype of well marked and strongly marked White Galloway (wsg, wsü)



heterozygous
(Cs₂₉ / wt₂₉)



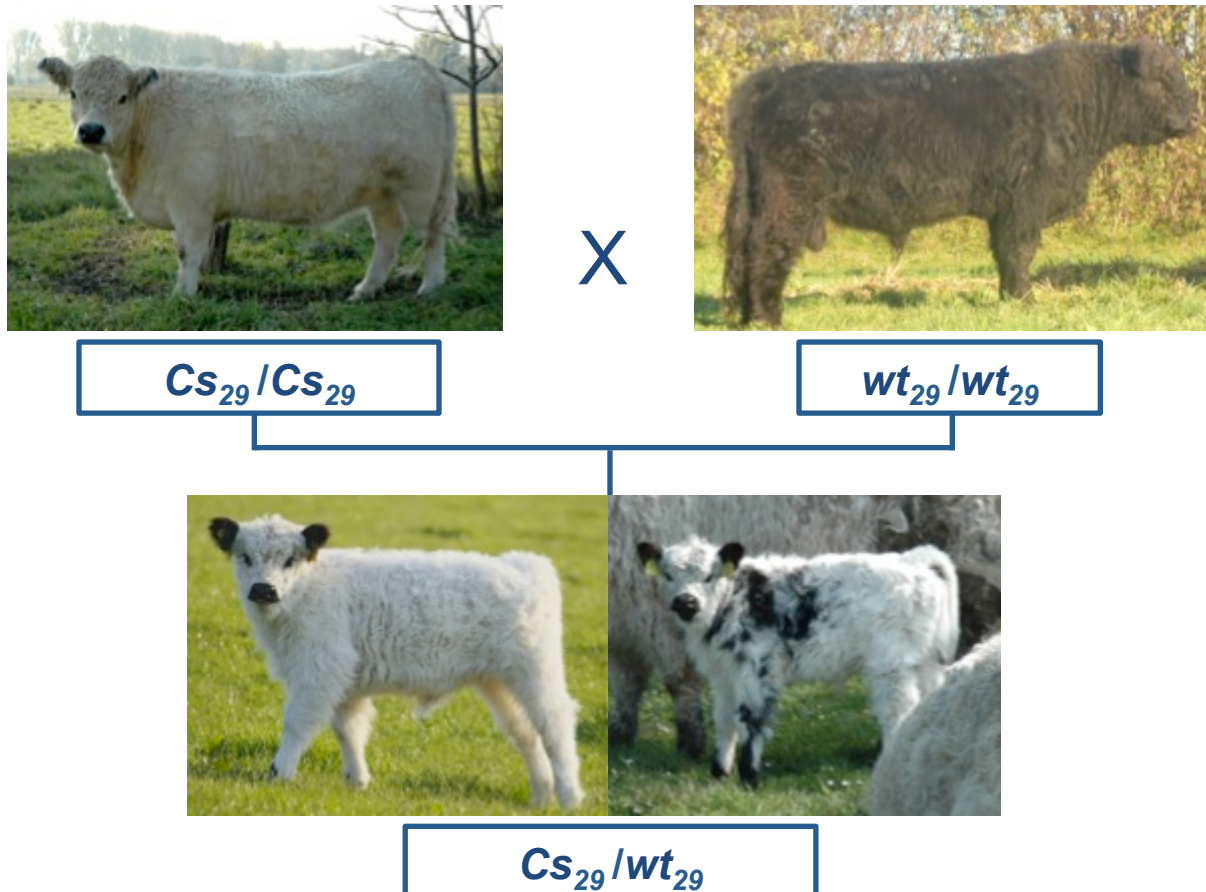
BTA29: Wildtyp (wt)
wt₂₉: White Galloway, wsch

Translocation breakpoint analysis

| White Galloway Phenotype | n | BTA29 (wt_{29}) | BTA29 (Cs_{29}) | | BTA6 (Cs_6) | |
|----------------------------------------|---------|---------------------|---------------------|---------|-----------------|---------------|
| | (total) | $\alpha\beta^*)$ | $\alpha D^*)$ | $EA^*)$ | $C\beta^*)$ | $\gamma B^*)$ |
| well marked (wsg)/strngly marked (wsü) | 162 | 162 | 162 | 162 | 162 | 0 |
| mismarked (wss) | 59 | 0 | 59 | 59 | 59 | 0 |
| fully black (wsch) | 42 | 42 | 0 | 0 | 0 | 0 |

Experimental matings

- 18 matings of White Galloway (wss x wsch)



Conclusions

- Coat colours are inherited in a Mendelian fashion
- Coat colours are caused by a duplication and translocation of the *KIT* gene on chromosome 29 (BTA29)
- Coat colours depend on the number of translocated *KIT* gene copies (dosage effect)
- Development of well marked (wsg) and strongly marked (wsü) phenotypes is still unclear



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White Galloway Breeders

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Sigrid Beny

Beatrix Heim

Hans-Heinrich Mahnke

Andrea u. Christian Wätjen

Hans-Hermann Kähler

Thomas Roth

Imke Möller

Guido Bernasconi

u. v. a.

White Park Breeders

Arche Warder

Familie Derjong

Uwe Müller-Hesse

Gut Deutsch-Nienhof

Wildpark Schorfheide

Lawrence Alderson