

Genome-Wide Association Study for Leg Disorders in Austrian Braunvieh and Fleckvieh

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EAAP, 2019



Objectives

Mining the **genetic heterogeneity** between Braunvieh and Fleckvieh



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Material

1 Cows

- 1 999 Fleckvieh (FLV), 985 Braunvieh (BSW)

2 SNPs

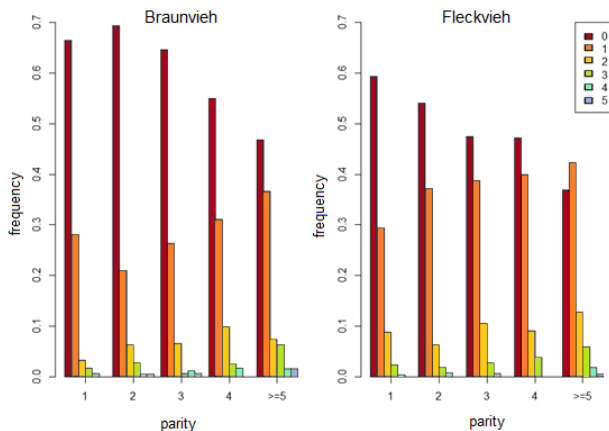
- Geneseek Genomic Profiler HD BeadChip
- 76 932 SNPs
- 74 762 SNPs (MAF 0.01; call rate 99 %)

3 Phenotype

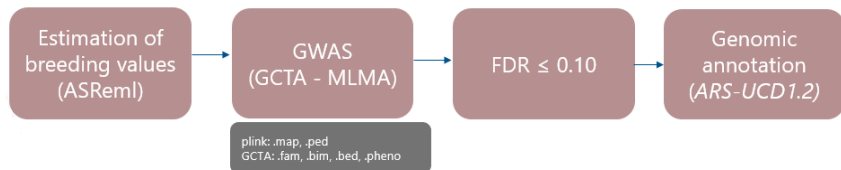
- total number of leg disorders until DIM 300th
- EBV



Material



Methods/genome-wide association study (GWAS)



Methods/EBV prediction

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{Z}_u\mathbf{u} + \mathbf{Z}_v\mathbf{v} + \mathbf{Z}_p\mathbf{p} + \boldsymbol{\varepsilon}$$

\mathbf{y}	total number of leg disorders scored till DIM 300
$\boldsymbol{\beta}$	fixed effects : a general mean, breed (Braunvieh or Fleckvieh), parity (1,2,3,4 or >4), calving year-season, hoof status recording code (four levels)
\mathbf{u}	$\sim N(0, \mathbf{A}\sigma_u^2)$ EBV
\mathbf{v}	$\sim N(0, \mathbf{I}\sigma_v^2)$ random veterinarian effect
\mathbf{p}	$\sim N(0, \mathbf{I}\sigma_p^2)$ random permanent environmental effect
$\boldsymbol{\varepsilon}$	$\sim N(0, \mathbf{I}\sigma_\epsilon^2)$ residual

Methods/GWAS model

$$\mathbf{u} = \mu + \mathbf{X}\mathbf{b} + \mathbf{Z}\mathbf{g} + \mathbf{e}$$

\mathbf{u} EBV

μ general mean

\mathbf{b} additive effect of all single SNP

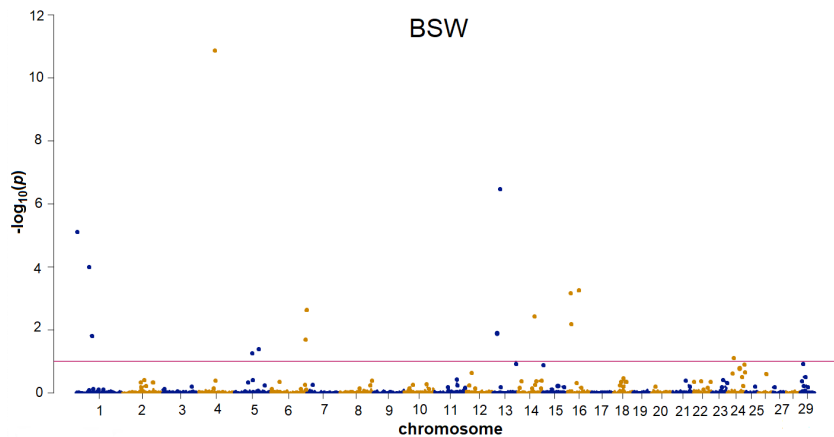
\mathbf{X} design matrix {0, 1, 2}

$\mathbf{g} \sim N(0, \mathbf{G}\sigma_g^2)$ cumulated effect of all remaining SNPs,

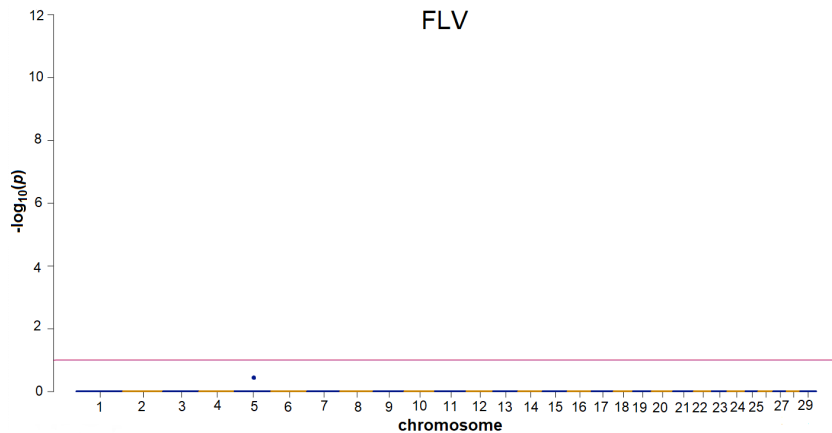
\mathbf{G} corresponds to the genomic covariance matrix between cows
calculated based on SNPs.

$\mathbf{e} \sim N(0, \mathbf{I}\sigma_e^2)$ residual

Results/GWAS



Results/GWAS



Results/GWAS

SNP	Position [bp]**	Additive effect	FDR*	Annotation**
ARS-BFGL-NGS-92033	1:3,303,269	0.018	$7.5 \cdot 10^{-6}$	intergenic, between MIS18A and HUNK
BTA-89698-no-rs	1:43,542,488	0.012	0.0001	intergenic, between DCBLD2 and COL8A1
ARS-BFGL-NGS-6521	1:50,767,507	0.007	0.0150	intergenic, between CBLB and CCDC54
BovineHD0400014448	4:52,028,036	0.029	$1.4 \cdot 10^{-11}$	intergenic, between CAV2 and TES
BovineHD0400014458	4:52,079,221	0.029	$1.4 \cdot 10^{-11}$	
Hapmap48066-BTA-73690	5:61,220,624	0.010	0.0530	intergenic, closest to NEDD1
ARS-BFGL-NGS-85328	5:81,769,685	0.008	0.0430	intergenic, between CCDC91 and PTHLH
ARS-BFGL-NGS-103113	6:115,208,599	0.016	0.0210	intergenic, between ADRA2C and LRPAP1
ARS-BFGL-NGS-100768	6:114,116,280	0.010	0.0020	intron of SORCS2
ARS-BFGL-NGS-25175	13:13,590,662	0.008	0.0120	intergenic, closest to CELF2
ARS-BFGL-NGS-101509	13:23,590,146	0.019	$3.3 \cdot 10^{-7}$	intergenic, between SPAG6 and PIP4K2A
ARS-BFGL-NGS-63852	14:55,768,446	0.010	0.0040	intergenic, between TMEM74 and EMC2
Hapmap55901-rs29024589	16:12,125,227	0.010	0.0007	intergenic between B3GALT2 and GLRX2
ARS-BFGL-NGS-109246	16:12,280,122	0.008	0.0060	intergenic, between UCHL5 and RGS2
Hapmap51828-BTA-38538	16:36,037,389	0.007	0.0006	intergenic, between RGS7 and XCL1
BovineHD2400006669	24:24,273,191	0.002	0.0820	intergenic, between CCDC178 and KLHL14

*False Discovery Rate

**ARS UCD1.2.assembly

Methods/principal components analysis



Methods/Local principal components analysis between breeds/ Mahalanobis distance

$$D_M = \sqrt{\mathbf{d}' \mathbf{V}^{-1} \mathbf{d}}$$

$$\mathbf{d} = [\bar{\epsilon}_{1BSW} - \bar{\epsilon}_{1FLV}, \bar{\epsilon}_{2BSW} - \bar{\epsilon}_{2FLV}, \dots, \bar{\epsilon}_{10BSW} - \bar{\epsilon}_{10FLV}]$$

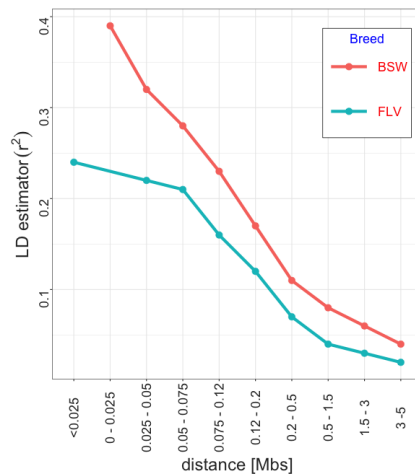
$\bar{\epsilon}_1, \bar{\epsilon}_2, \dots, \bar{\epsilon}_{10}$ - eigenvectors

\mathbf{V} - pooled covariance matrix of all eigenvectors

$$T = \frac{n_B n_F}{n_B + n_F} \cdot \frac{n_B + n_F - 11}{10(n_B + n_F - 2)} \cdot \mathbf{d}' \mathbf{V}^{-1} \mathbf{d} \sim F_{10, n_B + n_F - 11}$$

n_x - number of cows representing each breed

Comparison of LD patterns



Methods/Differences in pairwise LD structure between breeds

$$S = \left[\sum_{i=1}^n [(v_{i21} + v_{i22}) - (v_{i12} + v_{i21})]^2 \right] + \left[\sum_{i=1}^n [(v_{i11} + v_{i12}) - (v_{i21} + v_{i22})]^2 \right]$$

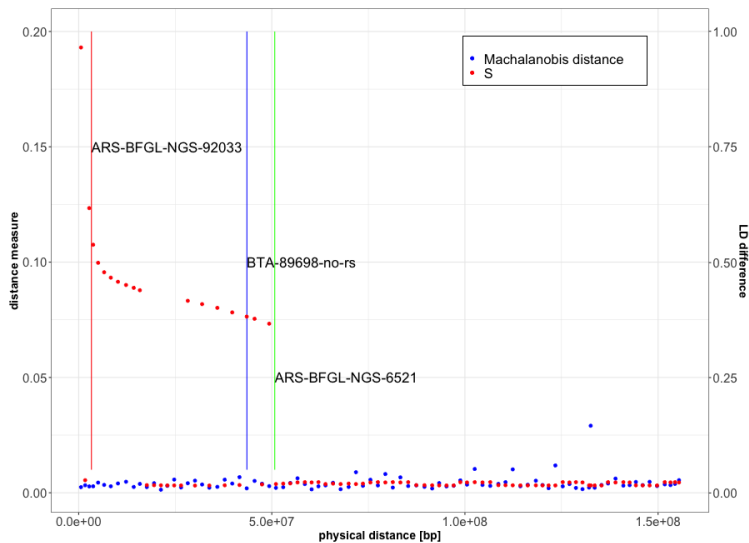
S – general measure of variability

v_{ij} – product of linkage disequilibrium covariance matrix of breed_{*i*}
and the vector of eigenvectors corresponding to breed_{*j*}

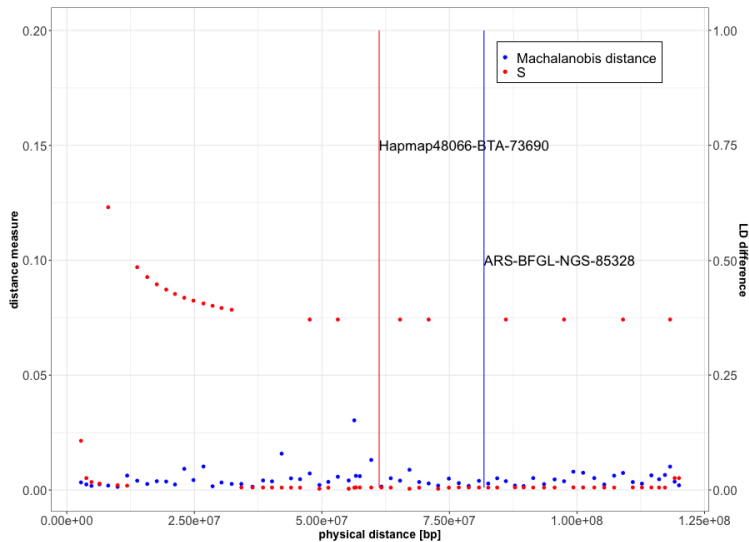
(Garcia, 2012)

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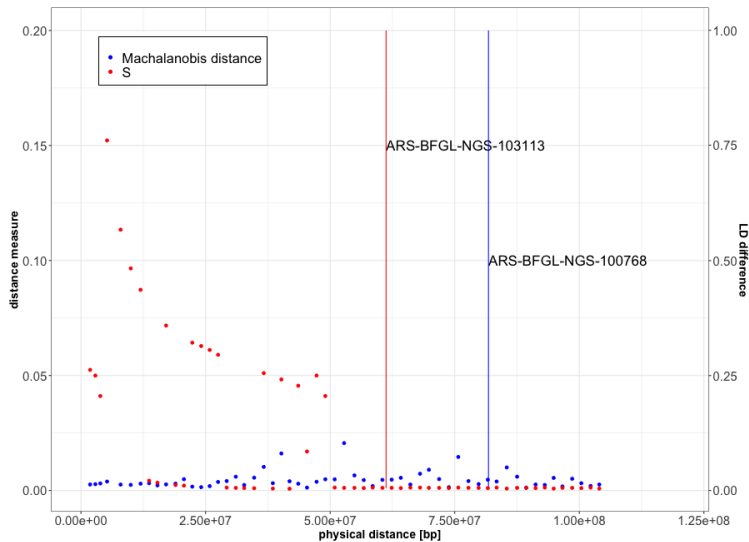
Results/PCA/BTA1



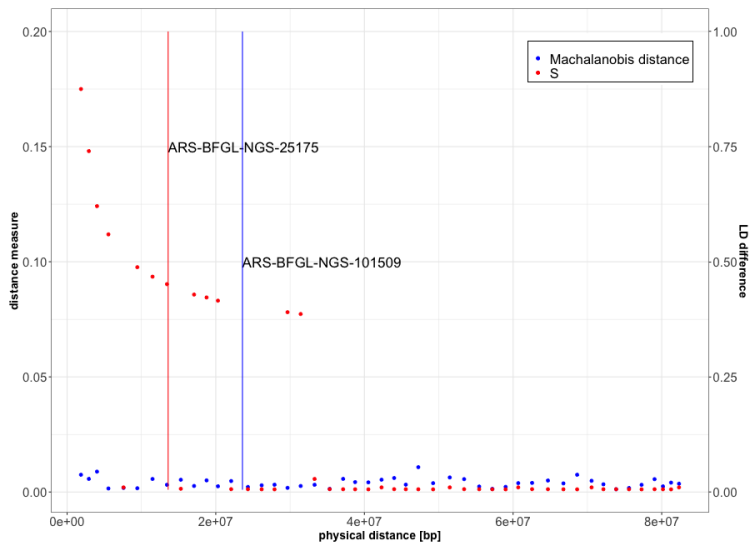
Results/PCA/BTA5



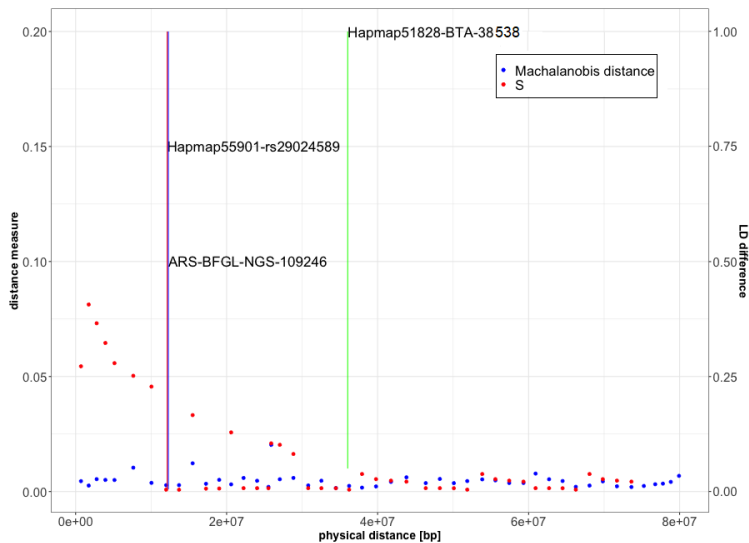
Results/PCA/BTA6



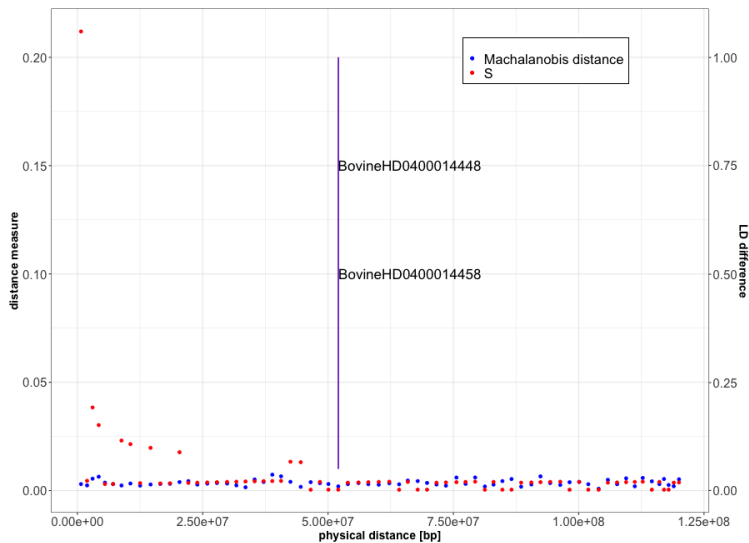
Results/PCA/BTA13



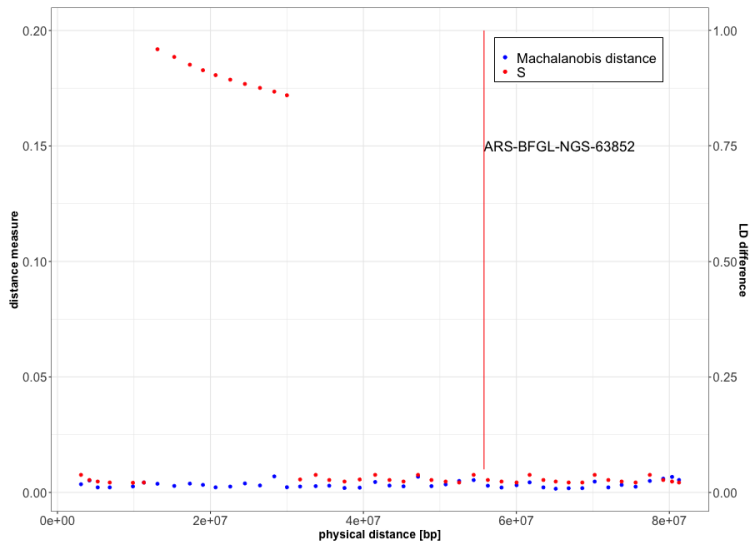
Results/PCA/BTA16



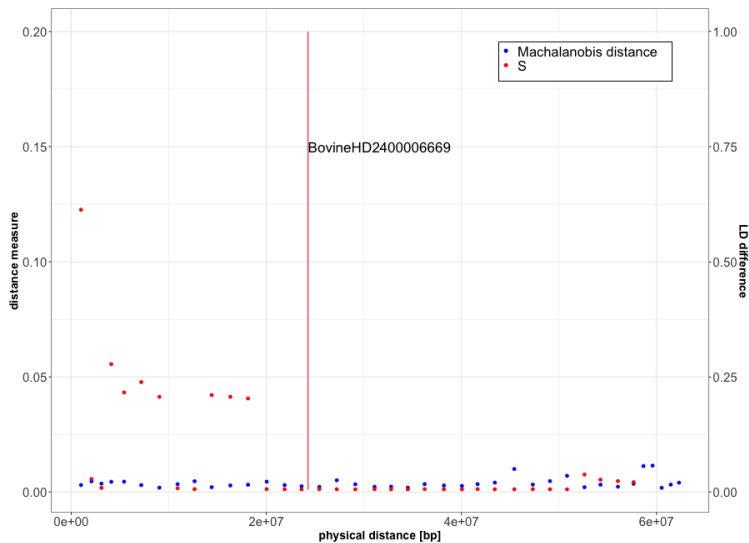
Results/PCA/BTA4



Results/PCA/BTA14



Results/PCA/BTA24



Conclusions

- 1 Genetic heterogeneity in the number of leg disorders between Fleckvieh and Braunvieh
- 2 Differences in LD structure between Fleckvieh and Braunvieh in some genomic regions
- 3 Differences in LD structure partially explain genetic heterogeneity

Acknowledgements

- 1 NCN grant No. 2015/19/B/NZ9/03725 as well as by the Efficient Cow and the Gene2Farm projects
- 2 Biostatistics group, Department of Genetics, Wrocław University of Environmental and Life Sciences
 - Magdalena Fraszczak
 - Joanna Szyda
 - Tomasz Suchocki
- 3 ZuchtData, Vienna, Austria
 - Christa Egger-Danner,
 - Hermann Schwarzenbacher