

Genetic and genomic selection to reduce boar taint in Danish pigs

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Outline

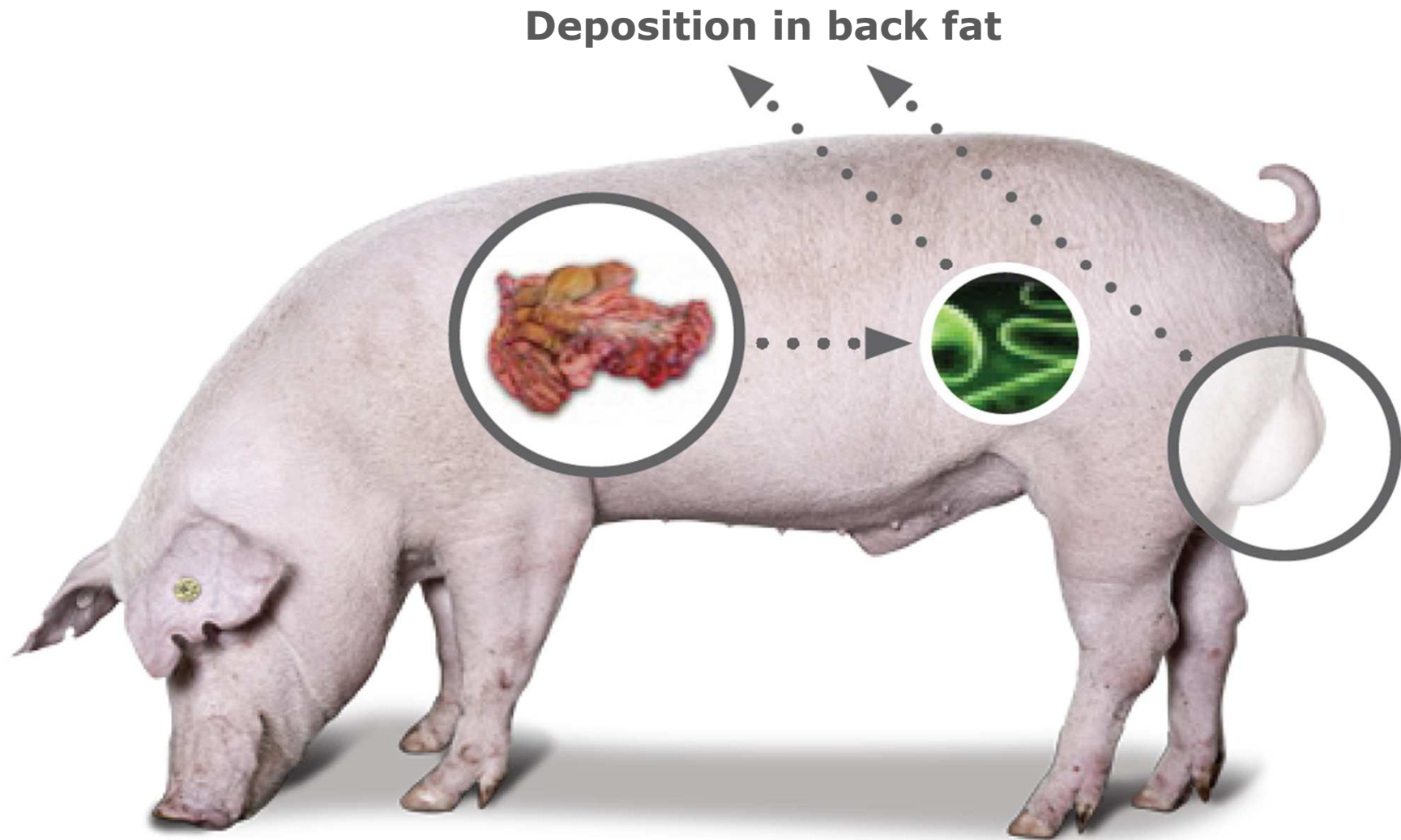
- Introduction
- Performance testing in Denmark
- Genetic associations to production and reproductive traits in Landrace and Duroc
- Results from genome wide associations
- Summary

Introduction

- Stop surgical castration by 2018???



- ☹ animal welfare
- ☹ labour intensive, especially with the use of anesthesia
- ☹ consumer acceptance



Skatole and indole:

- Hindgut microbial metabolism
- Tryptophan is the substrate
- Fecal like odor

Androstenone:

- Produced in testis
- Concentrated and secreted in saliva
- Urine like odor

Odours perceived by humans



So why has nothing happened?

☺ **Boar taint is heritable**

☹ Definition of “boar taint” is inconsistent

☹ No common reference method

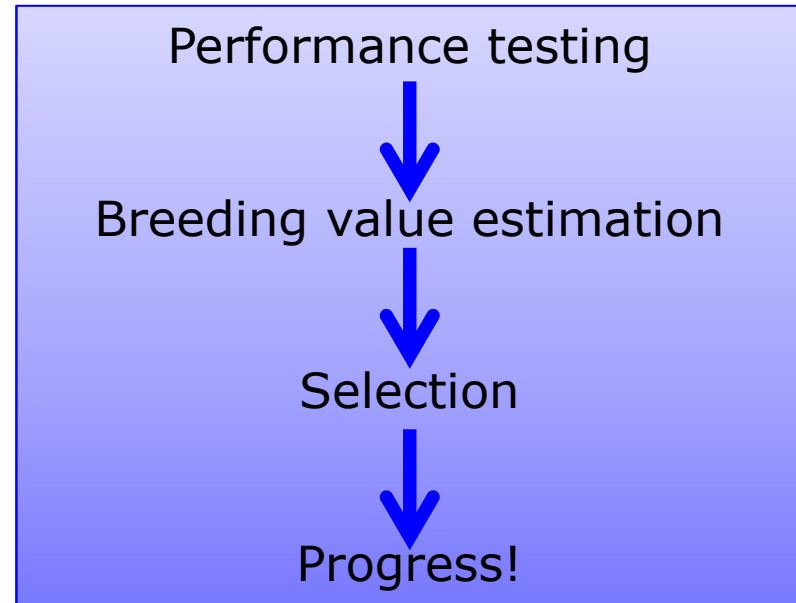
☹ Breeding takes time

☹ Unfavourable correlations to other important traits...?



Objective

- To develop a feasible selection strategy against boar taint in Danish pig breeds



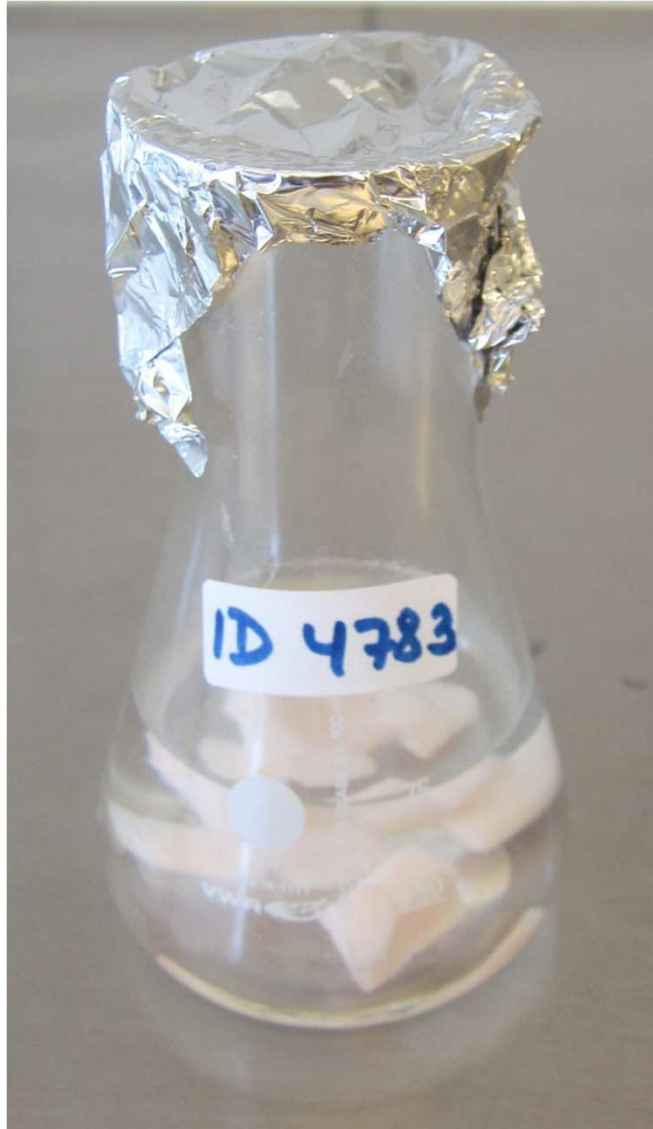
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Performance testing

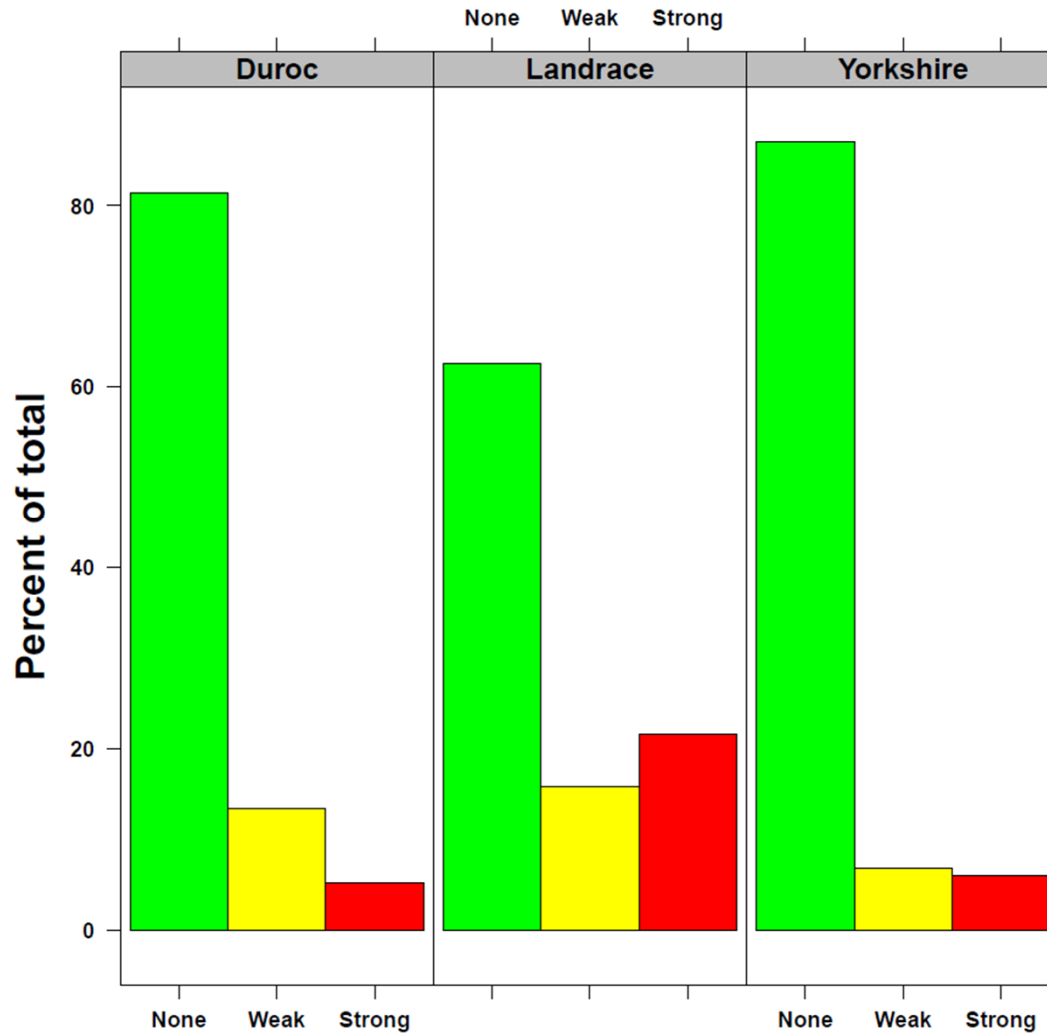
- Boar taint trait is recorded at test-station
 - All boars, i.e. AI- and slaughter boars
- Human-nose-score
 - Slaughter boars only – trained panel at DC-Ringsted
 - Photometric determination of skatoleEq
- Boar taint on live AI-boars via biopsy
 - Androstenone, skatole and indole

Human nose score



- Procedure:
 - 100 mL sample bottle
 - 5 g lard (medium size)
 - 75 mL boiling hot water
 - Stand for 2 min
- Scale:
 - 0 = no boar taint,
 - 1 = weak boar taint
 - 2 = strong boar taint
- Cat. 2 = Threshold

Human-nose score



Prevalence:

Duroc: 5.3%

Landrace: 21.7%

Yorkshire: 6.2%

Heritability of HNS

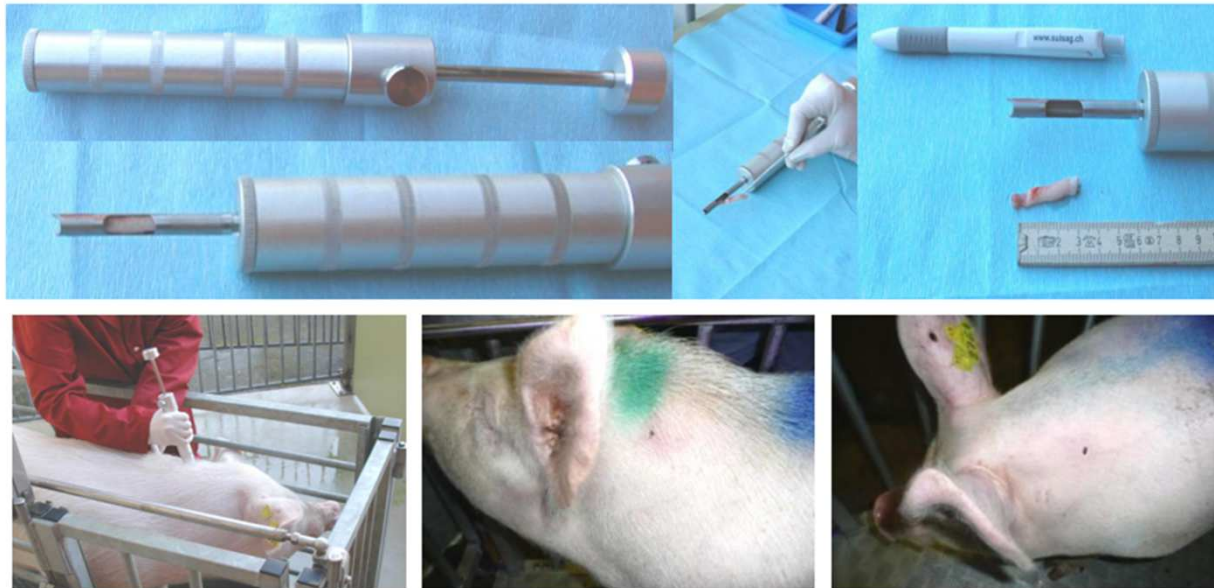
Unitrait model:

$$\text{HNS} = \underbrace{\text{Sektion*Panel} + \text{Lwgt} + \text{Age}}_{\text{(Fixed effects)}} + \underbrace{\text{Slweek} + \text{Animal} + \text{e}}_{\text{(Random effects)}}$$

Breed	Estimate
Duroc	0.08 (0.05)
Landrace	0.16 (0.07)
Yorkshire	0.09 (0.05)

Biopsy procedure

- Biopsy-device from SUIISAG
 - Baes et al. *Animal*. 2013. 7:714-20
- ~2700 AI-boars without complications
 - Animal care protocol is needed



Heritabilities of BT-compounds

Multitrait model, $i = \{\text{Log(Ska)}, \text{Log(Ind)}, \text{Log(And)}\}$:

$$\text{Log(BT}_i) = \underbrace{\text{Sektion} + \text{Lwgt} + \text{Age}}_{\text{(Fixed effects)}} + \underbrace{\text{Animal} + e}_{\text{(Random effects)}}$$

Duroc:

	Log(Ska)	Log(Ind)	Log(And)
Log(Ska)	0.22 (0.07)		
Log(Ind)	0.78 (0.20)	0.23 (0.07)	
Log(And)	0.28 (0.25)	0.21 (0.24)	0.48 (0.09)

Yorkshire:

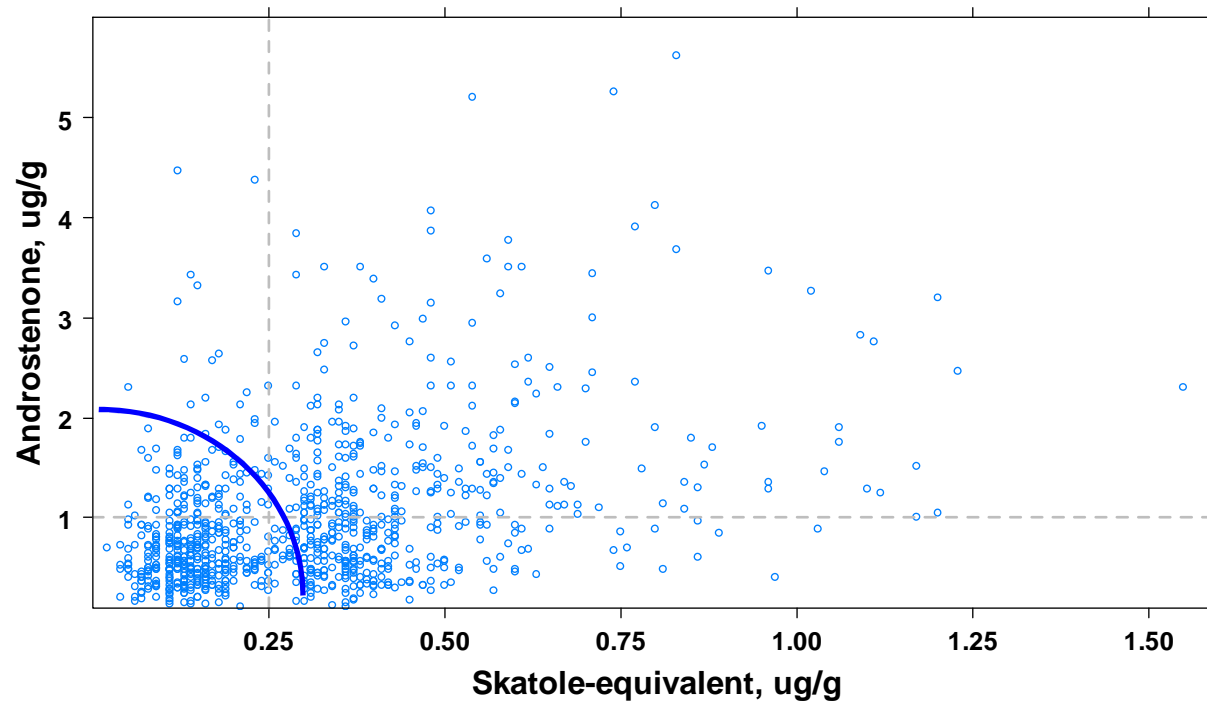
	Log(Ska)	Log(Ind)	Log(And)
Log(Ska)	0.37 (0.14)		
Log(Ind)	0.85 (0.33)	0.19 (0.11)	
Log(And)	0.18 (0.26)	0.58 (0.38)	0.53 (0.14)

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Danish Landrace

- Boar taint data (SABRE)
 - Skatole(equivalents) – 6000 boars
 - Androstenone – 500 pairs of full sibs



Genetic parameters for Landrace

Bivariate model:

$$\text{Log(Skatole)} = \text{HYS} + \text{SLwgt} + \text{SLage} + \text{Litter} + \text{Animal} + e$$

$$\text{Log(Andro)} = \text{Herd} + \text{SLwgt} + \text{SLage} + \text{Litter} + \text{Animal} + e$$

Trait	h^2	r_g	r_p
Log(skatole)	0.33 (0.05)	0.37 (0.12)	0.26 (0.03)
Log(Androstenone)	0.59 (0.14)		

Production traits: Landrace

- Multi-trait model for performance traits and boar taint compounds

Trait _x	$r_{g(x, \text{Log}(\text{skatole}))}$	$r_{g(x, \text{Log}(\text{androstenone}))}$
ADG100	-0.04 (0.08)	0.10 (0.11)
Feed conversion ratio	0.18 (0.19)	-0.04 (0.26)
Meat percentage	-0.20 (0.07)	-0.18 (0.10)

- Low genetic correlations and largely favorable!

Production traits: Duroc

➤ Results from fitting bivariate models

Trait _x	$r_{g(x, \text{ADG})}$	$r_{g(x, \text{Meat percentage})}$
Log(skatole)	-0.28 (0.22)	-0.01 (0.18)
Log(Indole)	-0.24 (0.20)	-0.29 (0.18)
Log(Androstenone)	-0.17 (0.11)	-0.25 (0.10)
HNS	0.28 (0.33)	-0.33 (0.28)

➤ In both breeds low genetic correlations

➤ Consistent with Dutch results

➤ Windig et al. (J. Anim. Sci. 2012.90:2120–2129)

Litter size data: Landrace

- Litter size traits: TBN and LP5
 - Full and half sib females to the BT boars
 - 1. parity sows and pure bred litters

Trait	No	Mean	SD	Min	Max
TBN	35715	13.6	3.80	1.0	28.0
LP5	34991	10.5	3.45	0.00	22.0

Litter size and BT: Landrace

- Heritability on the diag. with genetic correlations on the off diag.


Trait		TNB		LP5		Log(skatole)	Log(androstenone)
		Sire	Dam	Sire	Dam		
TNB	Sire	0.02 (0.01)					
	Dam	0.36 (0.14)	0.09 (0.01)				
LP5	Sire	0.70 (0.10)	0.17 (0.12)	0.02 (0.01)			
	Dam	0.43 (0.05)	0.58 (0.15)	0.38 (0.13)	0.06 (0.01)		
Log(skatole)		0.05 (0.22)	0.06 (0.11)	-0.11 (0.18)	0.03 (0.13)	0.33 (0.04)	
Log(androstenone)		-0.20 (0.27)	-0.14 (0.15)	-0.40 (0.22)	-0.20 (0.17)	0.41 (0.14)	0.59 (0.13)

- In general, weak genetic correlations between BT compounds and service-sire fertility

Littersize and BT: Duroc

- Bivariate models for TNB and boar taint

	TNB(dam)	TNB(S-sire)	Log(Andro)	Log(Ska)
TNB(dam)	0.10 (0.01)			
TNB(S-sire)	0.30 (0.11)	0.04 (0.01)		
Log(Andro)	-0.08 (0.16)	-0.13 (0.19)	0.47 (0.08)	



Model for semen and BT

- Combine Rep-model₍₁₎ with models_(2, 3) for boar taint compounds

$$\begin{bmatrix} \mathbf{y}_1 \\ \mathbf{y}_2 \\ \mathbf{y}_3 \end{bmatrix} = \begin{bmatrix} \mathbf{X}_1 & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{X}_2 & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{X}_3 \end{bmatrix} \begin{bmatrix} \mathbf{b}_1 \\ \mathbf{b}_2 \\ \mathbf{b}_3 \end{bmatrix} + \begin{bmatrix} \mathbf{Z}_{pe} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{0} \end{bmatrix} \begin{bmatrix} \mathbf{p} \\ \mathbf{0} \\ \mathbf{0} \end{bmatrix} \\ + \begin{bmatrix} \mathbf{0} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{Z}_{c_2} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{Z}_{c_3} \end{bmatrix} \begin{bmatrix} \mathbf{0} \\ \mathbf{c}_2 \\ \mathbf{c}_3 \end{bmatrix} + \begin{bmatrix} \mathbf{Z}_{a_1} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{Z}_{a_2} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{Z}_{a_3} \end{bmatrix} \begin{bmatrix} \mathbf{a} \\ \mathbf{a} \\ \mathbf{a} \end{bmatrix}$$

Non-existing residual covariances (co betw and BT compounds

- Standard assumptions

$$\mathbf{p} \sim N(\mathbf{0}, \mathbf{I} \otimes \mathbf{K}_p); \mathbf{c} \sim N(\mathbf{0}, \mathbf{I} \otimes \mathbf{C}); \mathbf{a} \sim N(\mathbf{0}, \mathbf{A} \otimes \mathbf{G}); \mathbf{e} \sim N(\mathbf{0}, \mathbf{I} \otimes \mathbf{R})$$

Semen and BT

➤ Key genetic correlations

Trait _x	$r_{g(x, \text{Log}(\text{skatole}))}$	$r_{g(x, \text{Log}(\text{androstenone}))}$
Volume	0.01 (0.13)	0.02 (0.18)
Concentration	-0.11 (0.13)	-0.24 (0.16)
Total sperm	-0.17 (0.13)	-0.13 (0.18)
Functional sperm	-0.16 (0.17)	-0.10 (0.18)

➤ High standard errors on genetic correlations

Semen quality and BT

- Multi-trait liability model for binary semen quality traits

Trait _x	h ²	r _g (x, Log(skatole))	r _g (x, Log(androstenone))
Normal/abnormal	0.08 (0.02)	0.09 (0.20)	-0.39 (0.25)
Motility low/high	0.21 (0.03)	-0.08 (0.15)	-0.38 (0.19)

- Low heritabilities for semen quality traits
- Weak genetic correlations

Semen and BT: Duroc

- Bivariate repeatability models

Trait _x	h ²	r _{g(x, Log(Ska))}	r _{g(x, Log(Ind))}	r _{g(x, Log(And))}
Motility	0.14 (0.02)	-0.41 (0.22)	-0.25 (0.22)	-0.06 (0.13)
Volume	0.29 (0.02)	0.15 (0.20)	0.10 (0.19)	0.15 (0.11)

- Again, weak genetic correlations

Testosterone and semen

- Low Test. Line (LTL): 28 ng/ml testosterone
- High Test. Line (HTL): 44 ng/ml testosterone

Traits ^a	LTL ^b	SE	HTL ^b	SE	<i>P</i> -value ^c
DSP/g, millions	2.89 ±	2.47	2.72 ±	1.50	0.55
TDSP, billions	1.62 ±	0.17	1.42 ±	0.11	0.34
SPM/g, millions	12.65 ±	1.08	11.87 ±	0.65	0.54
TTS, billions	7.08 ±	0.76	6.23 ±	0.46	0.34

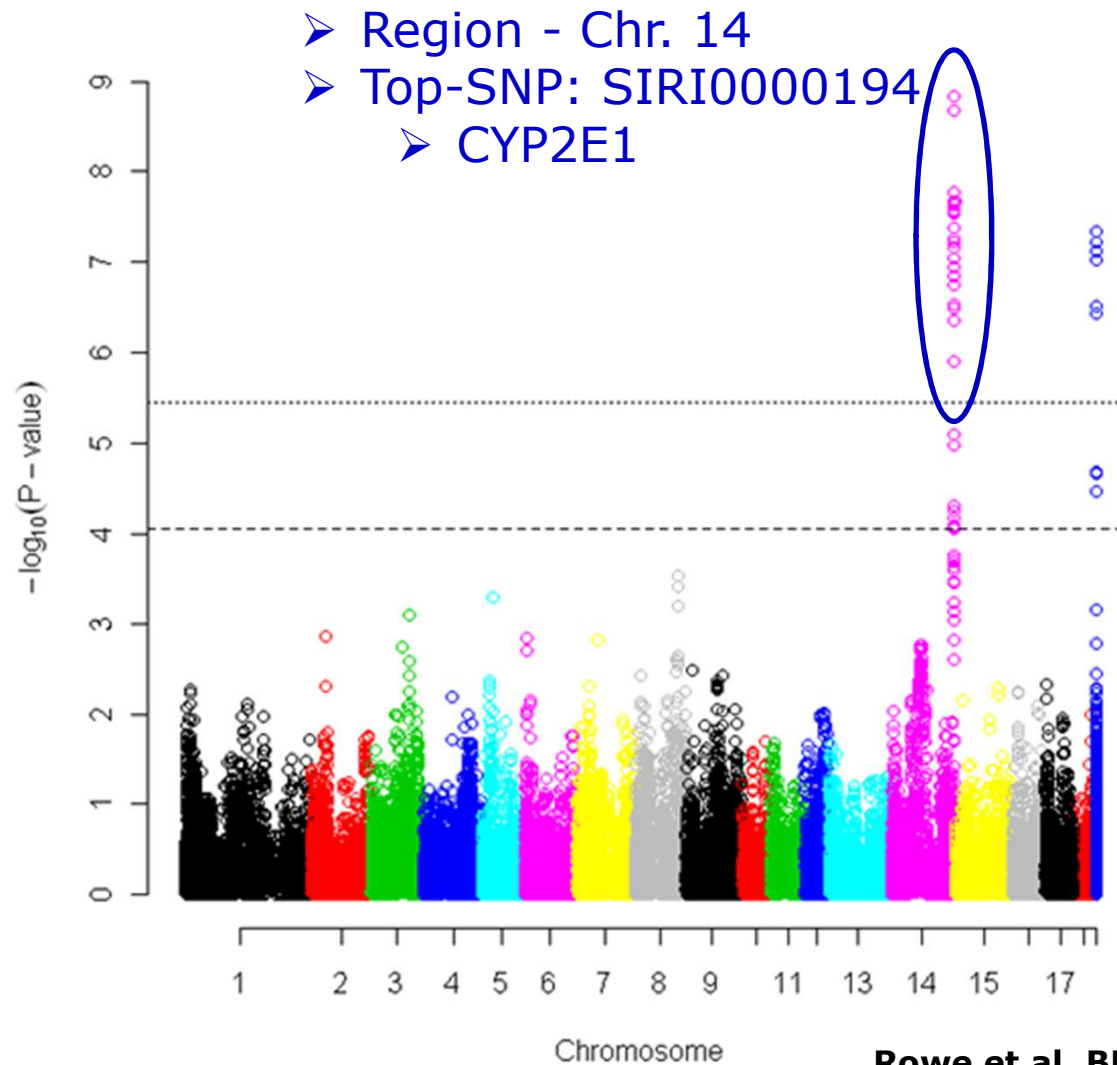
Walker et al. (J. Anim. Sci. 2004. 82:2259–2263)

- $r_p(\text{testosterone}; \text{TTS}) = 0.12$
 - Ren et al. (Reprod. Dom. Anim. 2009. 44:913–919)

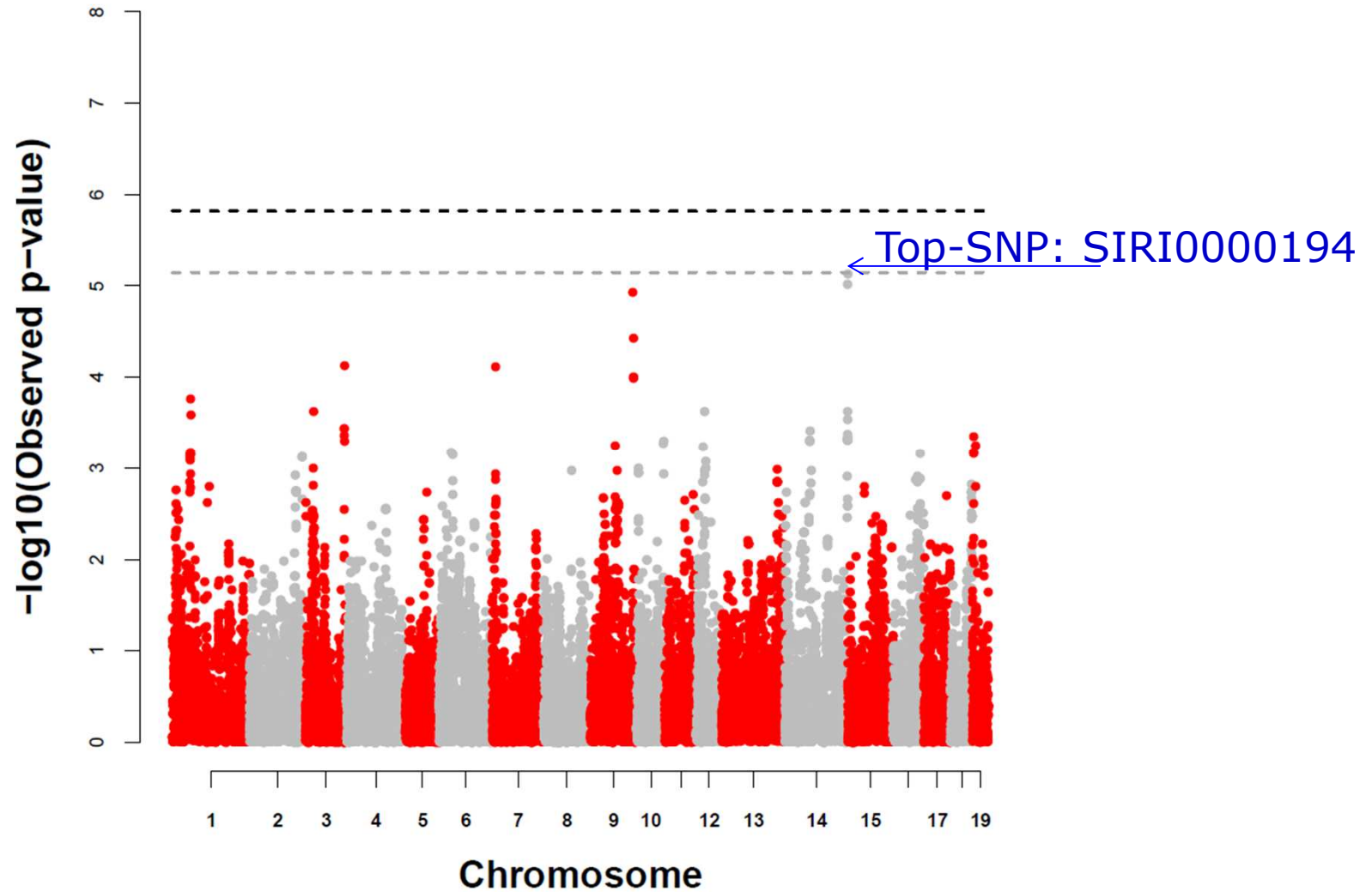
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- Concluding remarks

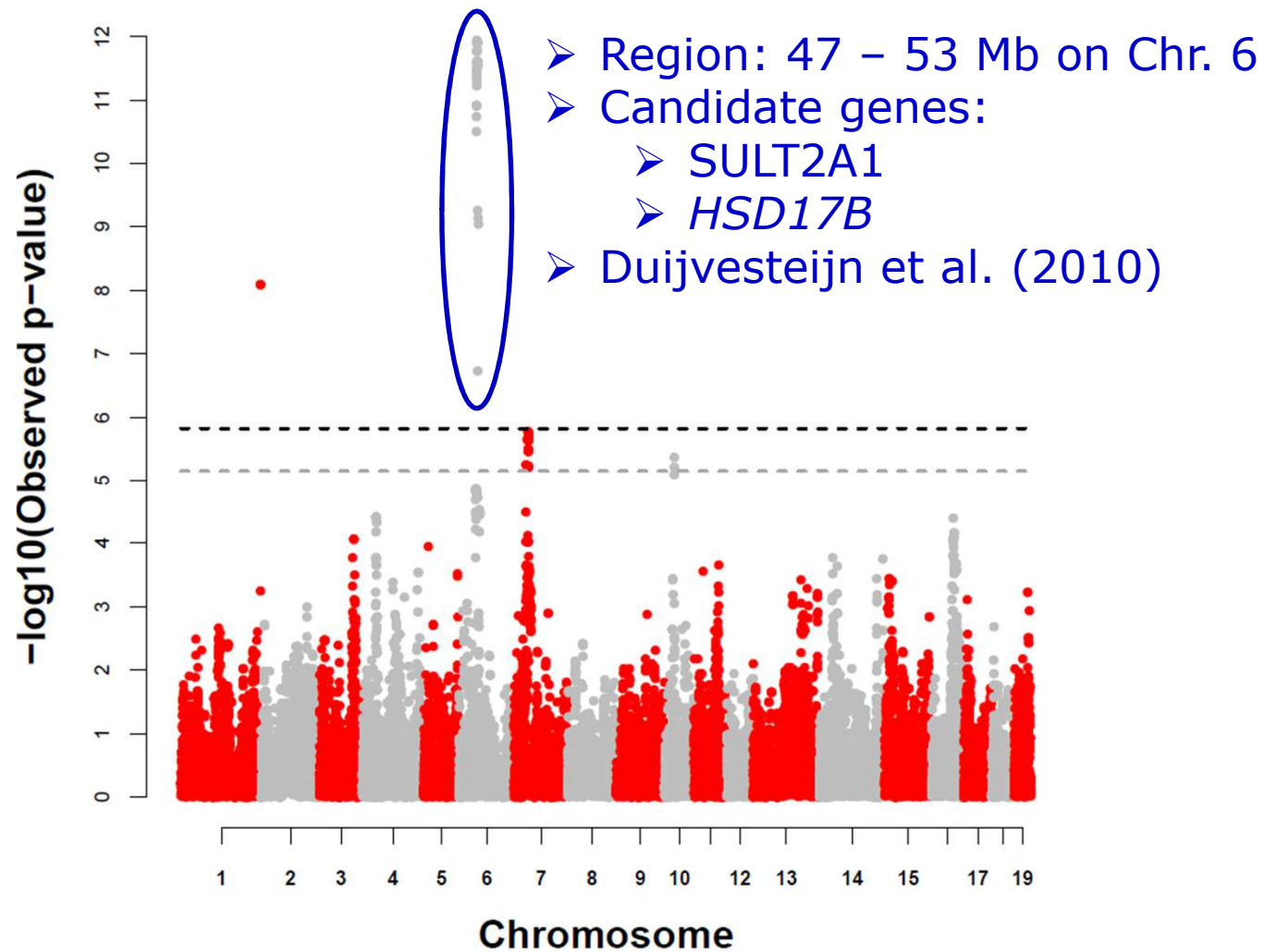
GWAS for skatole in Landrace



GWAS for skatole in Duroc



GWAS for androstenedione in Duroc



Summary

- Selection against boar taint will have
 - Minimal impact on production traits
 - Minimal impact on litter size traits
 - Minimal impact on semen traits

- GWAS results points to previous mapped regions

- Breeding is an option
 - Effective online sorting and grading of carcasses
 - Determine the economic weight of the trait
 - Phenotyping costs must be dramatically reduced
 - Breeding will never guarantee 100% of carcasses, being perceived as free of boar taint