

# Impact of genomic selection on genetic diversity in 5 European local cattle breeds

EAAP 2023

Renzo Bonifazi, Gwendal Restoux, Pascal Croiseau, Theo Meuwissen, Stephanie Minery, Jack Windig



## Acknowledgments



- Breeding companies for providing pedigree and SNP data
- Gerben de Jong (CRV)
- Mathijs van Pelt (CRV)



TOWARDS IMPROVEMENT OF **RUMINANT** BREEDING  
THROUGH **GENOMIC** AND EPIGENOMIC APPROACHES

- This project has received funding from the European Union's Horizon 2020 Programme for Research & Innovation under grant agreement n°101000226.
- This project is part of EuroFAANG (<https://eurofaang.eu>).

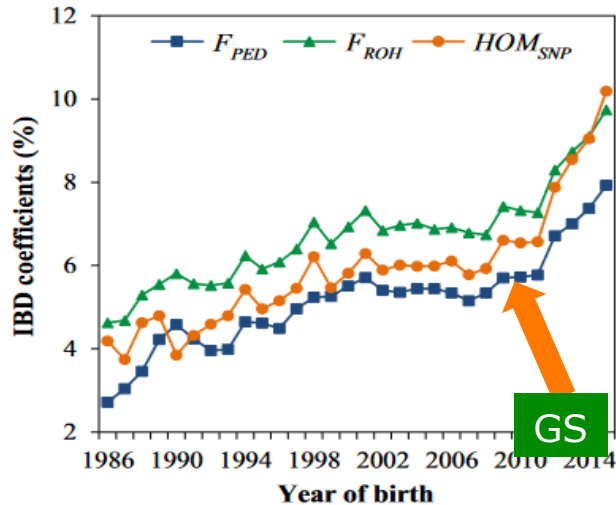
# Genetic diversity in local cattle populations

- Strong selection can lead to high inbreeding rates
  - associated loss of genetic diversity
  - + inbreeding depression
  - + risk expression genetic defect
- Especially in small populations
- Genomic selection (GS) now applied in local breeds
- Main question
  - Did inbreeding rates change after the introduction of GS in 5 local breeds?



# Changes in inbreeding rate after introduction GS

- Genomic selection (GS): starting ~2008 (VanRaden, 2008)
- Larger genetic gains, shorter generation intervals
- Expectation  $\Delta F$  to go down



## Up

- Holstein
  - the Netherlands (Doekes et al. 2018)
  - USA (Forutan et al. 2018; Makanjuola et al. 2020)
  - Poland (Topolski and Jagusiak 2020)
  - France (Doublet et al. 2019)
  - Australia (Scott et al. 2021)
  - Italy (Ablondi 2022)
- Jersey (Makanjuola et al. 2020)
- Finnish Ayrshire (Sarviaho et al. 2023)

## Not up

- Normande
  - Doublet et al. 2019
- Mont Beliarde
  - Lozada-Soto et al. 2021
- Aberdeen Angus
  - Lozada-Soto et al. 2021

# 5 cattle populations



MRY (NL)

Abondance (F)



Tarentaise (F)



Norwegian Red (N)



Vosgienne (F)

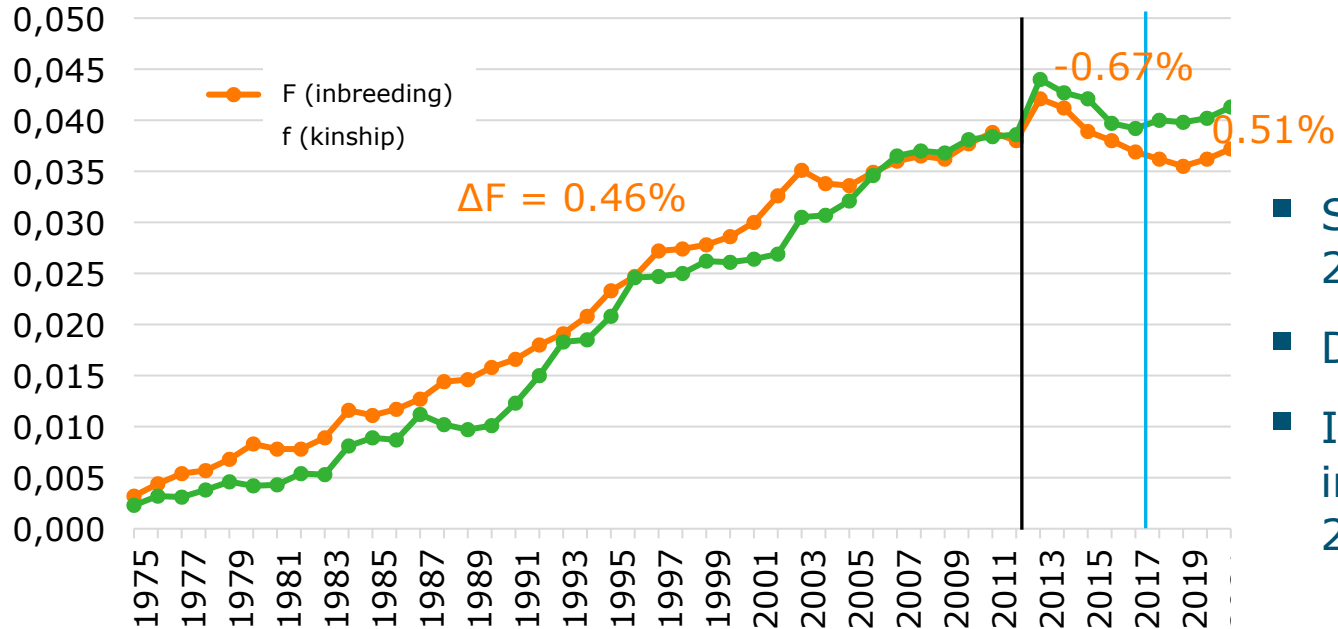


# Some data

<b>Breed</b>	<b>Calf/year</b>	<b>Pedigree</b>	<b>Genotypes</b>	<b>Years analysed</b>	<b>Introduction GS</b>
<b>MRIJ</b>	5,053	205,934	4,645	1975-2021	2018
<b>Norwegian Red</b>	35,701	721,805	193,489	2000-2020	2016
<b>Abondance</b>	25,020	807,387	16,427	2000-2020	2014
<b>Tarentaise</b>	9,514	259,150	8,882	2000-2020	2014
<b>Vosgienne</b>	3,738	99,833	4,466	2000-2020	2014

# MRY inbreeding and kinship levels

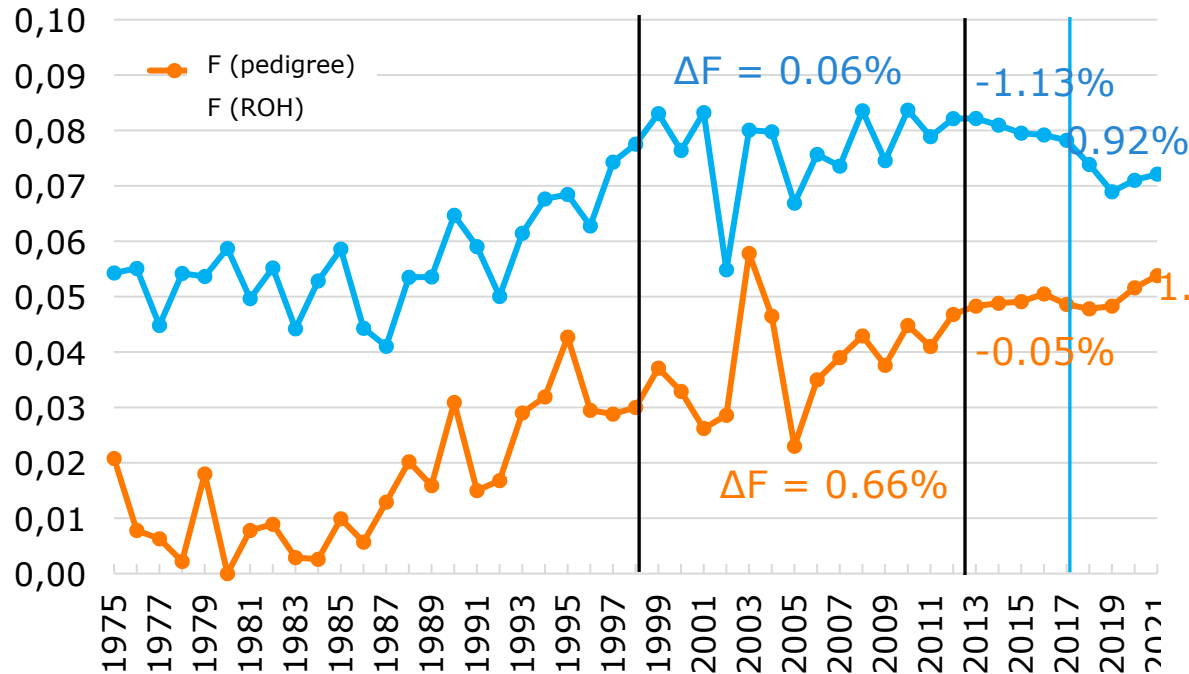
## ■ All animals in pedigree



- Steady increase up to 2013
- Decrease 2013 – 2018
- Increase since introduction GS in 2019

# MRY inbreeding and kinship levels

## ■ Genotyped animals only

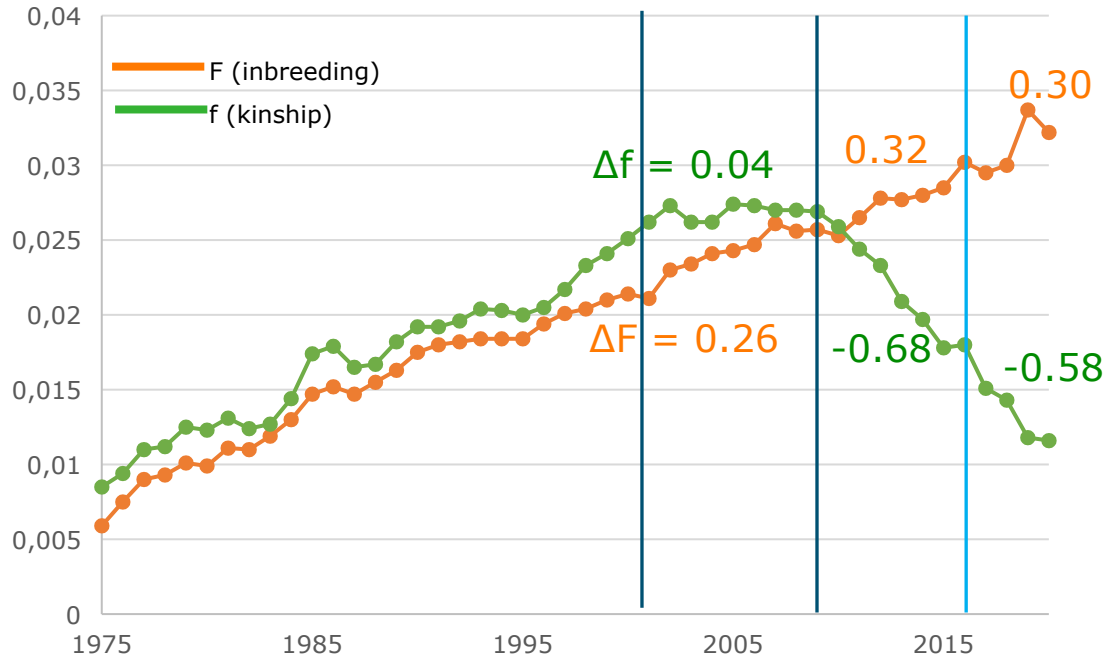


- Steady but irregular increase up to 2013
- Decrease F(ROH) but not F(ped) 2013 – 2018
- Increase since introduction GS in 2019



# Norwegian Red inbreeding and kinship levels

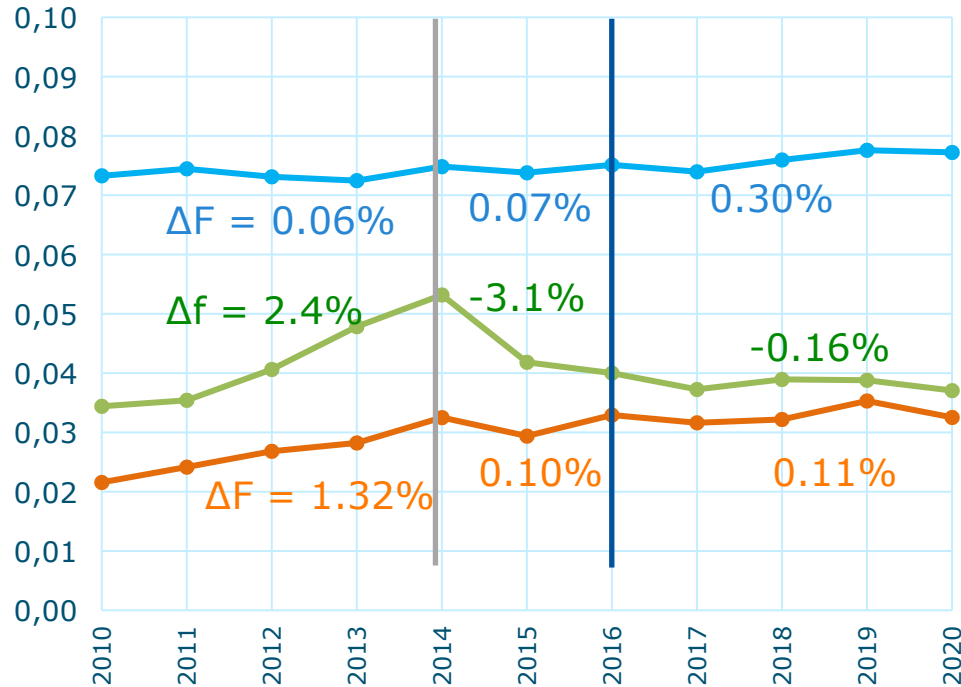
## ■ All animals in pedigree



- Steady increase of inbreeding
- Kinship decreases since 2009
- Inbreeding continues to increase since introduction GS in 2016
- Kinship continues to decrease

# Norwegian Red inbreeding and kinship levels

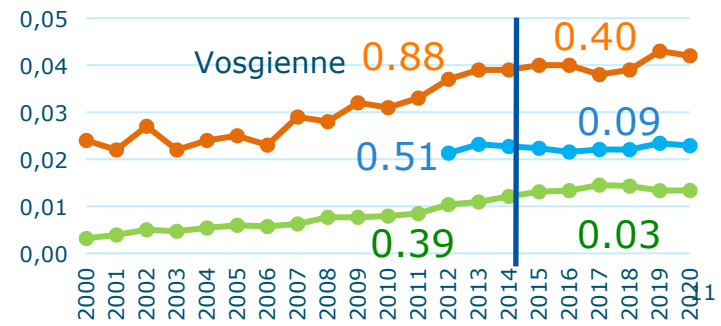
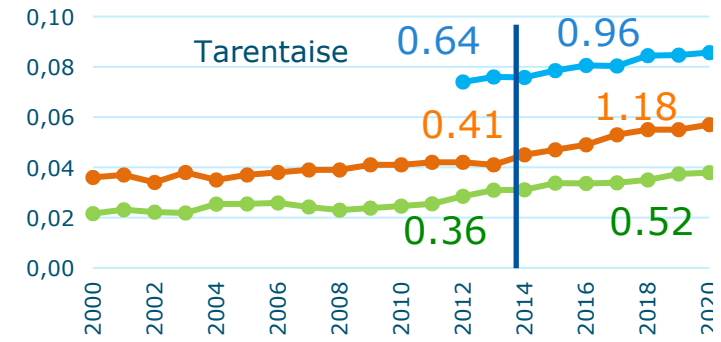
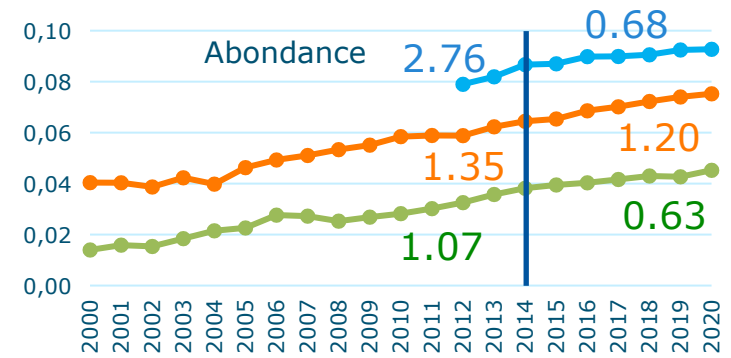
## ■ Genotyped animals only



- F(ROH) stable, slight increase since introduction GS
- Kinship decrease after 2014, stable after 2016
- F(ped) increase but less after 2014

# French breeds

- Steady increase in all levels
- After introduction GS (2014)
  - Abondance
    - Less steep
    - Especially F(ROH)
  - Tarentaise
    - Steeper
    - especially F(Ped)
  - Vosgienne
    - Clearly less steep
    - Especially F(ROH)

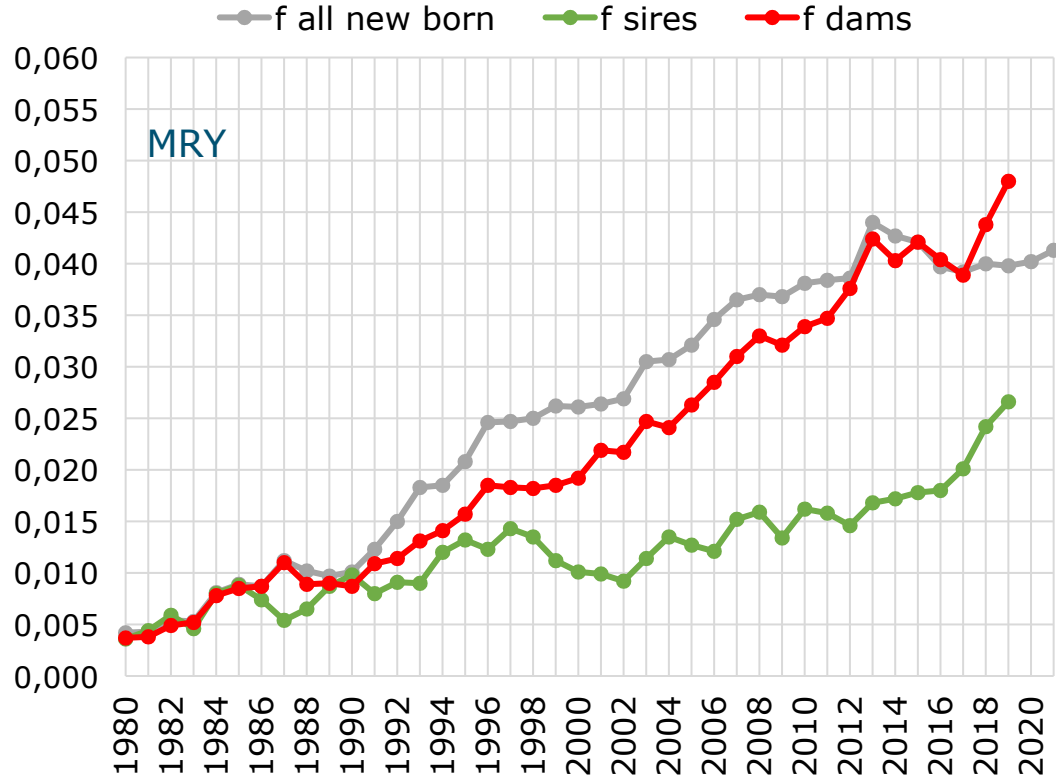


# Change in inbreeding and kinship rates

Breed	$F_{ped}$	$F_{ROH}$	$f$
MRIJ all	Stable/Up	-	Stable/Up
MRIJ genotyped	UP	UP	Stable/Up
NR all	Stable	-	Down/up
NR genotyped	Down/Stable	UP	Down/up
Abondance	Down	Down	Down
Tarentaise	Up	Up	Up
Vosgienne	Down	Down	Down

- Mixed pattern: rates can go
  - up
  - or down
  - or remain stable
- after introduction of GS

# Genetic management can reduce kinship



- MRIJ and Norwegian Red both use optimal contributions to select sires
- After introduction of OC a clear reduction in kinship

# Summary

- Inbreeding rates ( $\Delta F_{gen}$ ):
  - **Increased** (MRIJ, Tarentaise)
  - **More or less stable** (Norwegian Red)
  - **Decreased** (Abondance, Vosgienne)
- Genomic Selection:
  - shorter generation intervals
  - screening more individuals
  - preselection of less sires

## Main take-home message

- **genetic management** of local breeds more important than implementation of genomic selection per se

