



# EAAP 2018

69<sup>th</sup> Annual Meeting of the European Federation of  
Animal Science

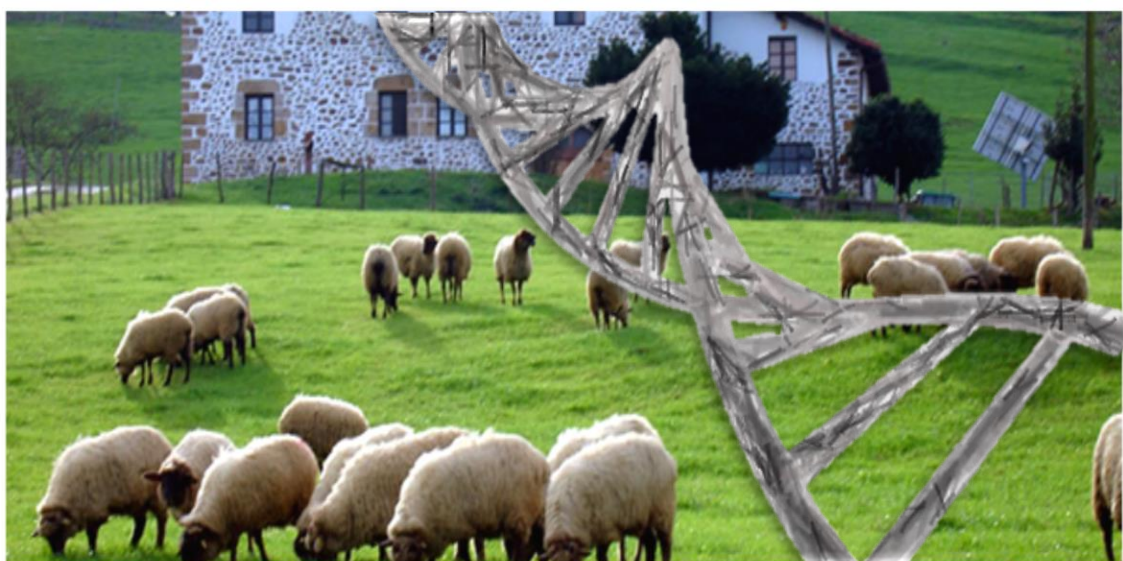
Dubrovnik, Croatia, 27<sup>th</sup> to 31<sup>st</sup> August 2018

## Session 26 Genetics & Genomics

Connectedness and application of breeding schemes for sheep & goats

# Genomic selection on Latxa dairy sheep

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DEPARTAMENTO DE DESARROLLO  
ECONÓMICO Y COMPETITIVIDAD

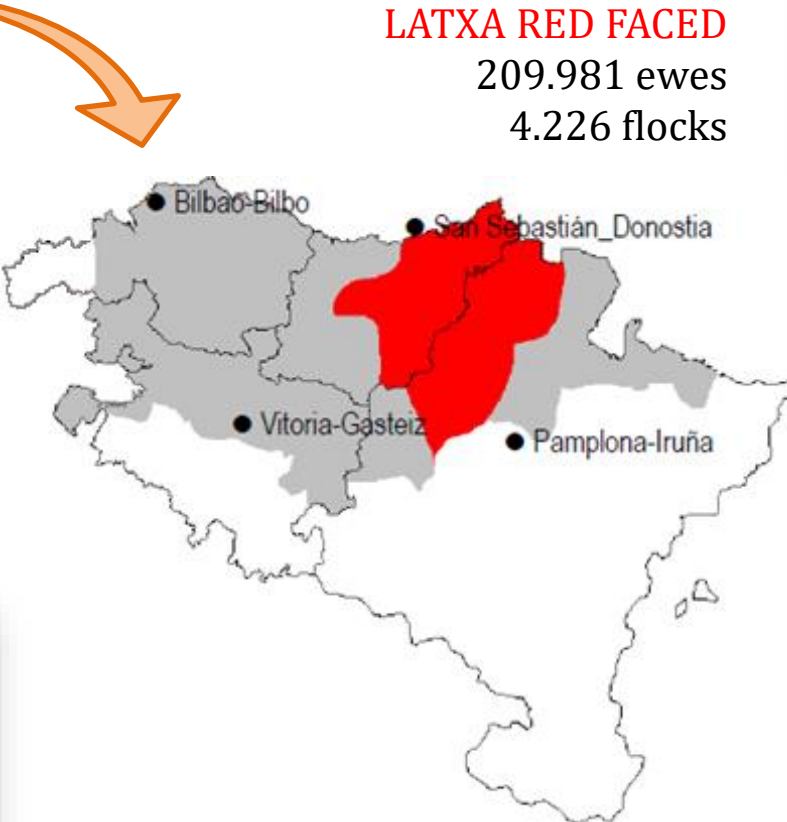


# Breeding program of Latxa breed (1984)

- Dairy sheep breed native from Basque Country and Navarre (Spain)



LATXA BLACK FACED  
from BC  
120.792 ewes  
3.742 flocks



LATXA RED FACED  
209.981 ewes  
4.226 flocks

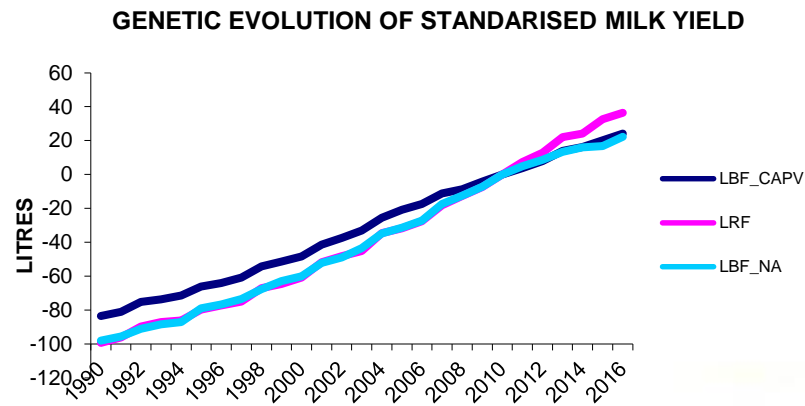


LATXA BLACK FACED  
from Navarre  
100.088 ewes  
985 flocks



# Breeding program of Latxa breed (1984)

- Dairy sheep breed native from Basque Country and Navarre (Spain)
- Selection based on pure breed
- Breeding goals:
  - Milk yield
  - Milk composition: fat and protein content
  - Udder morphology
- Annual genetic improvement of 3.5-4% in milk yield (Ugarte, 2017)



# GS in dairy sheep

**Lacaune** (Baloche et al., 2014; Barillet et al., 2014)

- Accuracy gain of GS over parent average
  - Milk yield: 0.15
  - Other characters: 0.10-0.20
- Reduction of AI rams by 20-40 %

**Lacaune:** GS was implemented in 2015

**Manech and Basco-Béarnaise** (Legarra et al., 2014)

- Increase in accuracy in single-breed predictions
  - MTN: 0.11   MTR: 0.16   BB: 0.06
- Multiple-breed pooled predictions
  - No increase in accuracy of genomic evaluations

**Manech and Basco-Béarnaise:** GS was implemented in 2017

# Objective

**Latxa** (Legarra et al., 2014)

- Hardly understandable results → No conclusive
  - Negative accuracy for CS
  - Contradictory results between breeds
- Probably due to the data structure
  - Weak relationship between training and validation group
  - Reduced number of genotyped rams

**Latxa:** Further studies are required

→ Ascertain the benefit of including molecular information  
into genetic evaluations of the Latxa sheep



SGTAACACACTCTGTAAACCTAACGATTAACTTGATCCACTGATTCAACGTACCGTAACGAACGTACCAATTGAGACTAAATATAACGTACCCATTAAAGA  
TAACGAACGTATCATTAAGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTACCAATTGAGACTAAATATAACGTACCCATTAAAGAAGCTACCGTA  
AATGATAACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACGTATCAATTGAGACTAAATATAACGTACCCATTAAAGA  
AGAGCTACCGTGCACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACGTATCAATTGAGACTAAATATAACGTACCCATTAAAGA  
AGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACAGTAACACACTCTGTTAACGAAACGTATCAATTGAGACTAAATATAACGTACCCATTAAAGA  
CACTGATTCAACGTTAACGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGCTCTGTTAACCTTAAGATTACTTGATCCACT

# Materials & Methods

## Latxa Red Faced



**illumina®**  
OvineSNP50  
Genotyping BeadChip

**Data 1984-2017**

**Genealogical** 150.164

**Phenotypical** 392.295

**Nº ♀ in data** 133.230

**Nº flocks** 325

**Nº ♂ in data** 1835

**Genotyped** 429

**SNPs** 39.593

# Materials & Methods

## Criterion:

### Accuracy difference

$$R_{\text{validation}} = \frac{\text{cov(EBV,TEBV)}}{V_{\text{EBV}} V_{\text{TEBV}}}$$



$$R^2_{\text{validation}} = R^2(\text{EBV}, \text{TEBV}) \frac{1+k}{\bar{EDC}} \quad k = \frac{4-h^2}{h^2} \quad (\text{Mäntysaari et al., 2010})$$



→ BLUP genetic evaluation

Rams in training group				Rams in validation group			
Nº genotyped	Nº nongenotyped	Birth years	Average EDC	Nº genotyped	Birth years	Average EDC	
184	1158	1984-2011	79.76	93	2012-2014	40.26	

# Validation group

## Results

Model	Bias	Slope	R
CS	$32.64 \pm 2.13$	$0.60 \pm 0.11$	$0.41 \pm 0.06$
GS	$14.09 \pm 4.33$	$0.52 \pm 0.08$	$0.44 \pm 0.05$
GS - CS	$-18.55 \pm 3.07$	$-0.08 \pm 0.06$	<b><math>0.03 \pm 0.03</math></b>

R(Cl,Gl): 0.89

R(CS,GS): 0.97



7.6%

## Previous results (Legarra et al., 2014)

Model	Bias	Slope	R
CS	$145.81 \pm 51.72$	$-0.08 \pm 0.39$	$-0.05 \pm 0.26$
GS	$91.05 \pm 36.72$	$0.33 \pm 0.27$	$0.26 \pm 0.20$
GS - CS	$-50.34 \pm 29.12$	$0.37 \pm 0.22$	<b><math>0.30 \pm 0.13</math></b>

# Families

GGTAAACACACTCTGTAAACCTAACGATTAAGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGACTAAATTAACGTACCATTAAGA  
AACGAAACGTATCAATTAGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGACTAAATTAACGTACCATTAAGAGCTACCGTA  
AATGATAAACAGTAACACACTCTGTAAACCTAACGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGACTAAAT  
AGAGCTACCGTGCACAGTAACACACTCTGTAAACCTAACGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGACTAAAT  
AGTAACACACTCTGTAAACCTAACGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGACTAAATTAACGTACCATTAAGA  
CACTGATTCAACGTTAACGATTACTTGATCCACTGATTCAACGTACCGTAACGAACGTATCAATTGAGCTTCTGTAAACCTAACGATTACTTGATCCACT

Model	50 lambs selected	Nº fathers
CS	44	42
GS		46



# Conclusions

7.6% (?) increase in accuracy for genomic evaluations

Lower than values reported in other breeds

- Interesting genotyping strategies
- Study of the practical implementation



**ESKERRIK ASKO  
MUCHAS GRACIAS  
THANK YOU  
MERCI BEAUCOUP  
HVALA**

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## 2018 results

Breed	Model	Bias	Slope	Accuracy
LBF	Pedigree	$8.02 \pm 22.97$	$0.96 \pm 0.17$	$0.59 \pm 0.08$
	Genomic	$104.84 \pm 8.17$	$0.62 \pm 0.17$	$0.43 \pm 0.09$
	Genomic-pedigree	$96.82 \pm 18.05$	$-0.34 \pm 0.13$	$-0.15 \pm 0.07$
LRF	Pedigree	$32.64 \pm 2.13$	$0.60 \pm 0.11$	$0.47 \pm 0.07$
	Genomic	$14.09 \pm 4.33$	$0.52 \pm 0.08$	$0.50 \pm 0.06$
	Genomic-pedigree	$-18.55 \pm 3.07$	$-0.08 \pm 0.06$	$0.03 \pm 0.04$

## Previous results (Legarra et al., 2014)

Breed	Model	Bias	Slope	Accuracy
LBF	Pedigree	$57.47 \pm 27.94$	$0.33 \pm 0.28$	$0.34 \pm 0.28$
	Genomic	$70.54 \pm 32.02$	$0.19 \pm 0.32$	$0.19 \pm 0.31$
	Genomic-pedigree	$14.77 \pm 15.38$	$-0.15 \pm 0.16$	$-0.18 \pm 0.13$
LRF	Pedigree	$145.81 \pm 51.72$	$-0.08 \pm 0.39$	$-0.05 \pm 0.26$
	Genomic	$91.05 \pm 36.72$	$0.33 \pm 0.27$	$0.26 \pm 0.20$
	Genomic-pedigree	$-50.34 \pm 29.12$	$0.37 \pm 0.22$	$0.30 \pm 0.13$

