### Form GENO

# DESCRIPTION OF NATIONAL GENOMIC EVALUATION SYSTEMS

Country (or countries)	Wallon Region of Belgium
Main trait group <sup>a</sup> .	Female Fertility
NOTE. Only one trait group per	,
form!	
Breed(s)	Belgian Blue
Trait definition(s) and unit(s) of	- Age at first artificial insemination (days)
measurement	- Age at first calving (days)
Attach an appendix if needed	- Interval between first and second calving (days)
Source of genotypes (chips used)	Illumina BovineSNP10K, Illumina Bovine SNP50K, Illumina
	BovineHD, Illumina Bovine MD
Imputation method for missing	Beagle3 (allelic dosage)
genotypes	
Propagation of genomic	The propagation is performed during the Single-step GBLUP
information to non-genotyped	to all animals related to genotyped animals and their relatives.
descendants and ancestors	
Animals included in reference	All genotyped animals included in extracted pedigree : 31222
population (males, females,	animals
countries included, total number)	
Source of phenotypic data (DYD,	Phenotypes
de-regressed proofs, national	- Age at first artificial insemination : data provided by farmer
EBVs and/or MACE evaluations)	or by inseminator
	- Age at first caving and interval between calving : from
	birth declaration
Other criteria (data edits) for	- Age at first artificial insemination : between 11 and 27
inclusion of records	months of age
	- Age at first calving : between 20 and 36 month of age
	- Interval between first and second calving : between 310
	and 570 days
	At least data from 10 animals in a given herd * year.
Criteria for extension of records	N/A
(if applicable)	
Sire categories	All sires
Genomic model (linear,	Single step GBLUP, multiple trait animal model, use of allelic
Bayesian, polygenic effect,	dosages.
genotypes or haplotypes)	
Blending of direct genomic value	Performed during the Single Step GBLUP method
(DGV) with traditional EBV	
Environmental effects in the	Herd * Year (F)
genetic evaluation model	Year * Month (F)
Adjustment for heterogeneous	No adjustment
variance in evaluation model	

Computation of genomic reliability Blending of foreign/Interbull	Genomic reliability is computed as polygenic reliability plus a gain provided by genomics. This gain expresses a number of 'daughter equivalent' that genomic brings compared to polygenic. It is calculated in two different ways, and it is always the smaller of the two that is kept. First gain is obtained by mean theorical genomic reliabilities minus mean theoretical polygenic reliabilities. Second gain is obtained by cross-validation. Younger animals are removed from dataset and the predictive ability is tested with and without genotypes. Difference between genomic predictive ability and polygenic predictive ability is used to get this second gain. No blending. Only local information			
information in evaluation				
Genetic parameters in the evaluation	See Appendix GENO			
Expression of genetic evaluations If standardized (e.g. RBV), give standardization formula in the appendix	Standardized breeding values: animals from genetic reference base are put to a mean of 100 and a standard error of 10			
Definition of genetic reference	All phenotyped animals born 4 years before the current year			
base	Base changes every year			
Labeling of genomic evaluations	GRBV			
Criteria for official publication of evaluations	Genotyped animals Sires : GREL $\ge$ 30 % Females : GREL $\ge$ 15 %			
Number of evaluations /	3			
publications per year				
Use in total merit index	No			
Anticipated changes in the near future	No			
Key reference on methodology applied	Legarra A., Christensen O., Aguilar I. and Misztal I. [2014] Single Step, a general approach for genomic selection. Livestock Science 166, 54-65			
Key organization: name, address, phone, fax, e-mail, web site	Organisation responsible for genetic evaluations and computing centre:			
	Elevéo asbl			
	R&D Department - Genetic Evaluation Unit			
	Rue des Champs Elysées 4			
	B-5590 Ciney			
	+32 (0)83 23.06.32.			
	evalgen@awegroupe.be			
	WEB site for publication of sire breeding values: <u>http://www.eleveo.be</u>			

<sup>a</sup>Either: Production (e.g. milk, fat, protein), Conformation, Health (e.g. mastitis resistance, milk somatic cell, resistance to diseases other than mastitis), Longevity, Calving (e.g. stillbirth, calving ease), Female fertility (e.g. non-return rate, interval between reproductive events, number of AI's, heat strength), Workability (e.g. milking speed, temperament), Beef production, Efficiency (e.g. body weight, energy balance, body conditioning score), or Other traits.

#### System Validation

Approximate number of test bulls for	Cross-validation study : Younger animals are removed from dataset
this trait group:	and the predictive ability is tested with and without genotypes.
If including foreign reference bulls:	
4-yr old de-regressed MACE EBVs,	Genetic trends
OR	
Current de-regressed MACE EBVs	Correlations between consecutive evaluations
If including foreign test bulls (type	
of proof 21 or 22), provide the	
reason.	
If using a truncation $\neq$ 4 years,	
provide the reason.	
If applying an age cutoff for test	
bulls $\neq$ (YYYY-8), provide the	
reason	

## **Appendix GENO**

#### Parameters used in genetic/genomic evaluation

Country (or countries):	Belgium (Wall		lloon R	egion)		
Main trait group:	Aain trait group: Female Fertili		ity			
Breed (repeat as necessary):		Belgian Blue				
Trait	Definition		ITB <sup>a</sup>	h <sup>2b</sup>	Genetic variance <sup>b</sup>	Official proof standardisation formula <sup>c</sup>
Age at first artificial insemination				0,24	1286	
Age at first calving				0,22	1488	
Interval between first and second calving				0,05	132.9	

<sup>a</sup>Indicate, with X, traits that are submitted to Interbull for international genetic evaluations.

<sup>b</sup>If repeated records are treated as separate traits, provide heritability estimates and genetic variances separately for each trait, as well as for all traits pooled, i.e. for the trait submitted to Interbull.

<sup>c</sup>Expressed as follows: StandEval=((eval-a)/b)\*c+d where a=mean of the base adjustment, b=standard deviation of the base, c=standard deviation of expression (include sign if scale is reversed), and d=base of expression.