

Status as of: 2023-03-24

Form GENO

DESCRIPTION OF NATIONAL GENOMIC EVALUATION SYSTEMS

Country (or countries)	SLOVENIJA		
Main trait group ^a .	CONFORMATION (sta, usu)		
NOTE. Only one trait group per			
form!			
Breed(s)	BSW		
Trait definition(s) and unit(s) of	Stature (cm), Udder Support (1-9)		
measurement			
Attach an appendix if needed			
Source of genotypes (chips used)	Weatherbys Scientific Bovine VersaSNP 50K		
	Illumina Bovine SNP50 BeadChip		
	GeneSeek Genomic Profiler v4		
Imputation method for missing			
genotypes			
Propagation of genomic			
information to non-genotyped			
descendants and ancestors			
Animals included in reference	All genotyped animals with MACE.		
population (males, females,			
countries included, total number)			
Source of phenotypic data (DYD,	De-regressed predicted genetic merit (DPGM) is calculated as:		
de-regressed proofs, national	DPGM = EBV/REL. In the genomic model, DPGM are weighted		
EBVs and/or MACE evaluations)	by REL _{dau} / (1 - REL _{dau}) , where REL _{dau} is reliability from		
	daughters.		
Other criteria (data edits) for			
inclusion of records			
Criteria for extension of records			
(if applicable)			
Sire categories	All		
Genomic model (linear, Bayesian,	An iterative, nonlinear model with heavy-tailed prior for marker		
polygenic effect, genotypes or	effects analogous to Bayes A is used. Base population allele		
haplotypes)	frequencies are subtracted from genotypes, and a polygenic		
	effect (poly) with 10% of additive variance is fit in the model:		
	DPGM = mean + ∑genotypes*effects + poly + error.		
Blending of direct genomic value	The evaluation model includes international (MACE) evaluation		
(DGV) with traditional EBV	of ancestors, and consequently the foreign information for non-		
	genotyped ancestors is included		
Environmental effects in the	Schotyped directors is included		
genetic evaluation model			
Senetic evaluation model]		



Adjustment for heterogeneous			
variance in evaluation model			
Computation of genomic reliability	Reliability of direct genomic value is computed from the traditional daughter equivalents plus the genomic daughter equivalents, which differ for each animal depending on its average genomic relationship to the reference population. Final reliability is computed by selection index using reliabilities of DGV, traditional PGM, and subset PGM.		
Blending of foreign/Interbull	Calculation of DGV includes foreign information from a subset		
information in evaluation	of the current Interbull evaluation for the genotyped animals.		
Genetic parameters in the evaluation	Use Appendix GENO for heritability/genetic variance estimates; for multiple-trait genetic evaluations, provide genetic correlation estimates between traits separately. Use also appendices PR, CO, BCO, SM, LO, CA, as applicable, if you participate in the international genetic evaluations of Interbull		
Expression of genetic evaluations If standardized (e.g. RBV), give standardization formula in the appendix	GEBV		
Definition of genetic reference base	The same as used in the national genetic evaluation.		
Labeling of genomic evaluations			
Criteria for official publication of			
evaluations			
Number of evaluations /	There will be 3 evaluations per year. The release dates would be		
publications per year	two weeks after the official release of Interbull evaluations.		
Use in total merit index			
Anticipated changes in the near future			
Key reference on methodology applied	VanRaden, P.M. Efficient methods to compute genomic predictions. J. Dairy Sci. 91(11):4414–4423. 2008.		
	VanRaden, P.M., Van Tassell, C.P., Wiggans, G.R., Sonstegard, T.S., Schnabel, R.D., Taylor, J.F., and Schenkel, F.S. <i>Invited review:</i> Reliability of genomic predictions for North American Holstein bulls. J. Dairy Sci. 92(1):16–24. 2009. Wiggans, G.R., VanRaden, P.M., Bacheller, L.R., Tooker, M.E., Hutchison, J.L., Cooper, T.A., and Sonstegard, T.S. <u>Selection and management of DNA markers for use in genomic evaluation</u> . J. Dairy Sci. 93(5):2287-2292. 2010.		
Key organization: name, address, phone, fax, e-mail, web site	University of Ljubljana, Biotechnical Faculty, Department of Animal Science, Groblje 3, 1230 Domžale, Slovenija		
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^aEither: 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 Al'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	7
this trait group:	
If including foreign reference bulls:	
4-yr old de-regressed MACE EBVs,	
OR	
Current de-regressed MACE EBVs	
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): SLOVE		ENIA					
Main trait group: C		CONFORMATION					
Breed (repeat as necessary):		BSW					
Trait	Definition		ITB ^a	h ^{2b}	Genetic variance ^b	Official proof standardisation formula ^c	
sta	Stature		X	0.4416	7.409	a=0 b=1.8760 c=12 d=100	
usu	Udder Support		X	0.1948	0.342	a=0 b=0.2994 c=12 d=100	

^aIndicate, with X, traits that are submitted to Interbull for international genetic evaluations.

^bIf 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.

^cExpressed 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.