

Form BEEF

DATA COLLECTION

Country (or countries)	UK
Trait name	Carcass weight (CWE)
Breed(s)	Simmental and Charolais (taken from across breed evaluation including on phenotypes from 89 other breeds)
Trait definition	The weight of the body of an animal, after the removal of the animal's head, hide, feet/legs, thoracic organs, internal fats, and abdominal organs, dressed according to a defined specification (three specifications exist in the UK namely Standard Specification, EC Reference Specification and UK Specification) followed by chilling (thus also known as cold carcass weight). Cold carcass weight is approximately 2% less than hot carcass weight.
Method and frequency of measurement	One measurement per animal
Who does the performance recording?	ABP food group (there are 6 other abattoirs but we are only supplying ABP data)
Method of collecting data	Carcass is weighed
Which animals get recorded?	Any animal slaughtered at an ABP processing plant
Is birthday recorded?	Yes
Is day of recording available?	Yes
Are the data adjusted and/or selected? If yes please describe the methodology applied	<p>All breeds are adjusted to a common variance by scaling the records on an individual for each trait using the following formula where i is the appropriate breed and sex and j is the appropriate sex but breed type=2.</p> $\text{Scaled phenotype} = \text{Average}(i) + [(\text{phenotype} - \text{average}(i)) * (\text{std}(j)/\text{std}(i))]$ <p>Breed types are defined as 1= Dairy, 2=Native beef, 3=Continental beef, 4=Other</p>
Time period for inclusion of records	02/01/2001-19/10/2018
Criteria (data edits) for inclusion of records	<p>Remove duplicates</p> <p>killdate supplied by the abattoir must be within 10 days of BCMS</p> <p>Sex must be recorded</p> <p>Age at slaughter must be recorded and $365 < \text{slage} < 1095$</p> <p>Carcass weight > 0kg</p> <p>Conformation and fat class recorded and valid EUROP classification</p> <p>Must be a heifer, steer or young bull</p> <p>Dam age > 540 days at calving</p> <p>Dam must be present in BCMS</p>

	<p>Dam birthdate missing Sire and maternal grandsire unknown Carcass weight within $\pm 3sd$ (for sire-breed, sex and age group) Average daily carcass gain $\pm 3sd$ (for sire-breed, sex and age group) BCMS birth recorded Died in the same herd as born No location change from birth herd Last BCMS movement was not death Finishing herd could not be assigned to be death herd Must have spent >60days in finishing herd BCMS kill location was not a slaughter house Birth herd year season must contain > 3 animals Kill herd year season must contain > 3 animals</p>
<p>Is embryo transfer applied? How are ET animals been identified?¹ Is recipient mother ID recorded?</p>	<p>ET animals removed</p>
<p>How do you treat incomplete data?</p>	<p>Whole record removed if any of the above criteria are not met</p>
MODEL	
<p>Model used for genetic evaluation^{2a}</p>	<p>MT-AM-FR</p>
<p>Environmental effects^{2b}</p>	<p>BirthHYS, 242666 (F) Slage (R) Sex (F) Killsite, 7 (F) KillHYS (F) Sex*KillHYS, 182750 (F) Killsite*killHYS, 137750 (F) Dam age (R) Once bred heifer, 3 (F) Dampercentedairy (R) Slage² (R) Het 1 (R) Het 2 (R) Het 3 (R) Het 4 (R) Het 5 (R) Het 6 (R) Rec 1 (R) Rec 2 (R) Rec 3 (R) Rec 4 (R) Rec 5 (R) Rec 6 (R)</p>
<p>Use of genetic groups and relationships</p>	<p>A standard relationship matrix is formed based on available pedigree information. Genetic groups are based on the animal's breed type. Breed types are defined as Dairy, Continental beef, Native beef,</p>

	Other
Genetic parameters in the model ³	See appendix I
Adjustment for heterogeneous variance in evaluation model	<p>Heterosis and Recombination coefficients were calculated from the breed type proportions of the animal's sire and dam and the formulae are as follows:</p> $\text{[heterosis]}_{ij} = ((\text{[sire]}_i * \text{[dam]}_j) + (\text{[sire]}_j * \text{[dam]}_i)) / 100$ $\text{[recombination]}_{ij} = ((\text{[sire]}_i * \text{[sire]}_j) + (\text{[dam]}_i * \text{[dam]}_j)) / 100$ <p>Breed types are defined as 1= Dairy, 2=Native beef, 3=Continental beef, 4=Other</p> <p>Het 1 and Rec 1 = breed types 1 and 2 Het 2 and Rec 2 = breed types 1 and 3 Het 3 and Rec 3 = breed types 1 and 4 Het 4 and Rec 4 = breed types 2 and 3 Het 5 and Rec 5 = breed types 2 and 4 Het 6 and Rec 6 = breed types 3 and 4</p> <p>These 12 terms are then fitted as covariates in the model</p>
System validation	Pre evaluation data quality checks and formation of contemporary groups. Genetic evaluation undertaken using MiX99. Post evaluation checks include adjusting for the genetic base, quality assurance checks and data summaries.
Definition of genetic reference base	Genetic reference animals are those animals that are born in 2010.
Next base change	
Assessment of index quality (computation of reliability, connection)	Reliabilities are computed using the mix99 software using the Mistztal and Wiggans (1988) calculation method.
PUBLICATION	
Expression of genetic evaluations	EBVs are produced and published on http://egenes.co.uk/carcassdata/ on behalf of AHDB
Criteria per official publication of evaluations	None
Number of evaluations / publications per year	Schedule to be arranged (expected 2 or 3 times a year)
Anticipated changes in the near future	None at this stage
Key reference on methodology applied	The mix99 software package is used for the genetic evaluations (Lidauer and Stranden, 1999; Vuori et al., 2006)
Key organization:	Scotland's Rural College (SRUC)
Contact person, address, phone,	contact persons : Abbygail Wells and Karolina Kaseja

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- 1) Use Appendix II BEEF for sample ID of ET animals
- 2a) Use abbreviation listed in the attached list of abbreviation to define the type of model.
- 2b) Use abbreviation for most common effects as listed in the attached list of abbreviation indicating, also, if the effect is treated as random (R) or fixed (F).
- 3) Use Appendix I BEEF for heritability/genetic variance estimates.

Form BEEF

Appendix I BEEF

Parameters used in genetic evaluation

Country: United Kingdom

Main trait group: Carcass weight (CWE)

Breed: Charolais and Simmental

Trait ⁽¹⁾	Definition	h_d^2	h_m^2	$r_{g(d,m)}$	c^2	σ_p^2
CWE	Carcass weight	0.42	-	-	-	610.1

h_d^2 : direct heritability; h_m^2 : maternal heritability; $r_{g(d,m)}$: genetic correlation between direct and maternal effects; c^2 : repeatability of (maternal) permanent environmental effects; σ_p^2 : phenotypic variance.

1) If you have more than one trait provides the correlations between traits.

Genetic, residual, and phenotypic correlations from bivariate combinations of net weight, fat, age, and average daily carcass gain (ADCG)

	Net weight	Conformation	Fat	Age
Genetic correlations				
Net weight				
Conformation	0.49 (0.035)			
Fat	-0.38 (0.040)	-0.51 (0.036)		
Age	0.03 (0.039)	-0.10 (0.039)	-0.04 (0.038)	
ADCG	0.86 (0.011)	0.49 (0.033)	-0.36 (0.038)	-0.42 (0.031)
Residual correlations				
Net weight				
Conformation	0.26 (0.021)			
Fat	0.27 (0.025)	0.19 (0.026)		
Age	0.24 (0.032)	0.10 (0.032)	0.10 (0.033)	
ADCG	0.88 (0.006)	0.23 (0.023)	0.26 (0.027)	-0.08 (0.035)
Phenotypic correlations				
Net weight				
Conformation	0.35 (0.006)			
Fat	-0.01 (0.006)	-0.11 (0.007)		
Age	0.13 (0.007)	-0.003 (0.007)	0.03 (0.008)	
ADCG	0.87 (0.002)	0.34 (0.006)	-0.02 (0.007)	-0.27 (0.007)

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Appendix II BEEF

Sample of ET animal IDs