- 1 Variants at RNF212 and RNF212B are associated with
- 2 recombination rate variation in Soay sheep (Ovis aries).
- Susan E. Johnston<sup>1,\*</sup>, Martin Stoffel<sup>1</sup>, Josephine M. Pemberton<sup>1</sup>

  <sup>1</sup>Institute of Evolutionary Biology, School of Biological Sciences,

  University of Edinburgh, Edinburgh, EH9 3FL, United Kingdom

  \*Corresponding author: Susan.Johnston@ed.ac.uk

# 7 Abstract

8 [to be written]

# 9 Introduction

Meiotic recombination is ubiquitous in sexually-reproducing organisms and is an important 10 driver of genomic diversity in eukaryotes (BARTON and CHARLESWORTH, 1998; OTTO and 11 LENORMAND, 2002). The process of crossing-over ensures the proper disjunction of homol-12 ogous chromosomes during meiosis (HASSOLD and HUNT, 2001), with most species having 13 a minimum requirement of at least one obligate crossover per chromosome pair (OTTO and PAYSEUR, 2019). It also has the advantage of generating novel haplotypes and preventing 15 the accumulation of harmful mutations in the genome, allowing populations to respond to 16 selection faster and to a greater degree (Muller, 1964; Hill and Robertson, 1966; Batta-17 GIN et al., 2016). However, this can come at the cost of increased mutations at crossover 18 sites (HALLDORSSON et al., 2019) and the break up of favourable haplotypes (BARTON and 19 CHARLESWORTH, 1998). Variation in recombination rate is likely to arise across taxa with 20 variation in genome size and chromosome number (STAPLEY et al., 2017); in addition, the 21 22 evolutionary costs and benefits of recombination may vary depending on the selective con-23 text, again leading to the expectation of recombination rate variation within and between populations (Otto and Barton, 2001; Otto and Lenormand, 2002). Indeed, cytogenetic 24 and linkage mapping studies have shown that recombination rates can vary by orders of 25 magnitude within and between chromosomes (Myers et al., 2005), individuals (Kong et al., 26 2004), populations (SAMUK et al., 2020), sexes (LENORMAND and DUTHEIL, 2005) and species 27 (STAPLEY *et al.*, 2017). 28 In mammals, genomic studies in different species (i.e. humans, cattle, sheep, pigs and deer) have shown that recombination rate is often heritable, with a significant proportion of the 30 additive genetic variation underpinned by loci of moderate to large effects (SANDOR et al., 31 2012; Kong et al., 2014; Ma et al., 2015; Kadri et al., 2016; Petit et al., 2017; Johnston et al., 32 2018; JOHNSSON et al., 2020). Genome-wide association studies (GWAS) have shown repeated 34 associations with variants at ring finger protein 212 RNF212 and its paralogue RNF212B, 35 whose associated proteins are essential for the crossover designation process during meiosis (REYNOLDS et al., 2013), as well as other loci with established roles in meiotic processes (e.g. 36

37 HEI10, REC8 and TOP2; SANDOR et al. 2012; Kong et al. 2014; MA et al. 2015; KADRI et al.

38 2016; Petit et al. 2017; Johnston et al. 2018; Johnsson et al. 2020). This heritable, oligogenic

39 architecture implies that recombination rate has the potential to evolve rapidly to selection

40 if variation in recombination rate is associated with individual fitness (Otto and Barton,

41 2001; STAPLEY et al., 2017). Therefore, detailed examination of the genetic architecture of

2 recombination rate is a critical step in understanding how rates are evolving over different

43 evolutionary timescales.

The Soay sheep (Ovis aries) are a Neolithic breed of domestic sheep that have been living 44 wild on the St Kilda archipelago since the Bronze age. These sheep are a model system for 45 studies of ecology and evolution, and have been intensively studied since 1985 (Clutton-46 Brock et al., 2004). A previous study integrated pedigree information with genotypic data 47 from ~39,000 SNPs to characterise autosomal crossover counts (ACCs) in ~3,300 gametes 48 transmitted from parents to their offspring (JOHNSTON et al., 2016). A GWAS showed that a 49 single region on chromosome 6, corresponding to the locus RNF212, explained around 47% 50 of heritable variation in female ACC only. A broader-scale "regional heritability" analysis identified a tentative association in both sexes at a 1.09 Mb region on chromosome 7, con-52 taining the candidate loci REC8 and RNF212B), but this was not significant in the GWAS 53 or in sex-specific regional heritability models. Since then, addition of more SNP genotyped 54 individuals and improvements in pedigree construction methodology (Huisman, 2017) now 55 allows us to characterise recombination rate in a further ~3,900 gametes. Furthermore, ad-56 vances in genotype imputation methods in complex pedigrees has allowed us to increase the 57 SNP dataset for conducting genome-wide association study from ~39,000 to ~417,000 SNPs. 58

59 Here, we revisit our previous analysis to carry out GWAS of male and female recombina-

60 tion rate. Our motivation for this study was to: (a) confirm a sex-limited association with

61 RNF212 on chromosome 6; (b) fine-map the association on chromosome 7 and determine the

62 action and effect size of this locus in both males and females; and (c) to identify any further

3 associations that could not be detected using the previous dataset.

## 64 Materials and Methods

## 65 Study Population.

- 66 Soay sheep living within the Village Bay area on the island of Hirta, St Kilda, Scotland
- 67 (57°49'N, 8°34'W) have been individually studied since 1985 (Clutton-Brock et al., 2004).
- 68 All sheep are ear-tagged at their first capture, including 95% of lambs born within the study
- 69 area, and the majority of animals are recaptured on an annual basis. DNA samples are rou-
- 70 tinely obtained from ear punches and/or blood sampling. All animal work is carried out
- 71 according to UK Home Office procedures and was licensed under the UK Animals (Scientific
- 72 Procedures) Act 1986 (License no. PPL60/4211).

# 73 Genotype and Pedigree Data.

- 74 All sheep were genotyped at 51,135 SNP markers on the Illumina Ovine SNP50 BeadChip
- 75 (Kijas et al., 2009). Quality control was carried out using the check.marker function in
- 76 GenABEL v1.8-0 with the following thresholds: SNP minor allele frequency (MAF) > 0.001,
- 77 SNP locus genotyping success > 0.99, individual sheep genotyping success > 0.95; a total
- 78 of 7,700 sheep and 39,398 SNPs remained. Pedigree relationships were previously inferred
- 79 using data from 438 SNP loci in the R package Sequoia v1.02 (Huisman, 2017) and from field
- 80 observations between mothers and their offspring born within the study area. Autosomal
- 81 SNP genotypes were used to calculate the  $\hat{F}_{III}$  genomic inbreeding coefficients using GCTA
- 82 v1.90.2 (Yang et al., 2011).
- 83 A further 189 individuals were genotyped at 430,702 polymorphic SNPs on the Ovine In-
- 84 finium HD SNP BeadChip for imputation of genotypes into individuals typed on the 50K
- 85 chip (see JOHNSTON et al. 2016 for method and individual selection criteria and STOFFEL et al.
- 86 2020 for full imputation method for this dataset). Briefly, SNP genotypes from the HD Chip
- 87 were imputed into the SNP50 Chip typed individuals using AlphaImpute v1.98 (HICKEY et al.,
- 88 2012; Antolín et al., 2017) resulting in a dataset with 7,691 individuals genotyped at 417,373

- 89 SNPs, with a mean genotyping rate per individual of 99.5% (range 94.8% 100%). As there
- 90 were no X-chromosome SNPs in common between the two SNP Chips, only the SNP50
- 91 Chip SNPs were used for association analyses on the X chromosome. SNP positions for both
- 92 chips were are known relative to sheep genome assembly Oar v3.1 (GenBank assembly ID
- 93 GCA 000298735.1).

# 94 Estimation of autosomal crossover counts and linkage maps.

- 95 Autosomal crossover positions were estimated using an identical protocol to that outlined in
- 96 JOHNSTON et al. (2016). Briefly, the method uses Ovine SNP50 BeadChip SNP data from a
- 97 focal individual's parents, mate and offspring to characterise crossovers that occurred in the
- 98 gametes transmitted from the focal individual to its offspring. Crossovers were determined
- 99 using the software CRI-MAP v2.504a (GREEN et al., 1990). Here, we estimated ACC for a
- 100 further 3,908 gametes, leading to a dataset of ACC for 7,235 gametes transmitted from 1,632
- 101 unique focal individuals. Because of differences in the SNP quality control between analyses,
- 102 we used 37,853 SNPs in common with the previous recombination rate study. Simulation
- 103 studies showed that this method will identify >99% of crossovers, meaning that ACC es-
- 104 timated using this method can be used as a proxy for individual recombination rate. The
- 105 method also provides linkage maps for each chromosome, allowing us to update the existing
- 106 map for Soay sheep to include information from a much larger number of meioses.

#### 107 Animal Models

- 108 A restricted maximum likelihood (REML) animal model approach (HENDERSON, 1975) was
- 109 used to estimate components of phenotypic variance for ACC, including the additive genetic
- 110 effect (i.e. heritability). A genomic relatedness matrix (GRM) was constructed for all poly-
- 111 morphic autosomal SNPs from the 50K chip using GCTA v1.90.2 (YANG et al., 2011) and
- 112 was adjusted with the argument --grm-adj 0, which assumes a similar frequency spectra of
- 113 genotyped and causal loci. Related individuals were not removed from the GRM as there is
- 114 high relatedness in the study population, and there are were no significant parental effects

or common environment effects on individual recombination rate (JOHNSTON et al., 2016). 115 116 Animal models were constructed for both sexes combined, females only and males only, and were run in ASReml-R v4 (BUTLER et al., 2009) in R v3.6.2. Fixed effects included sex (in 117 the combined model) and the genomic inbreeding coefficient  $\hat{F}_{III}$ ; initial models fit age in 118 years fit as a linear covariate, but as it was not significant, it was removed from the final mod-119 els (Figure 1). The significance of fixed effects was tested using Wald tests. Random effects 120 included the additive genetic effect (as estimated using the GRM) and identity of the focal 121 122 individual (i.e. "permanent environment") effect to account for repeated measures. Despite not being significant, the fixed effect  $\hat{F}_{III}$  and the random permanent environment effect were 123 retained in all models to account for potential underestimation of ACC due to long runs of 124 homozygosity and pseudoreplication, respectively. Significance of the additive genetic effect 125 was determined by dropping the additive genetic effect from each model and comparing with 126 the full model using a likelihood ratio test distributed as  $\chi_1^2$ . Bivariate models examining 127 the genetic correlation  $r_A$  between male and female ACC were run using the CORGH error 128 129 structure in ASReml-R (correlation with heterogeneous variances). Models were set with  $r_A$ to be unconstrained. To test whether  $r_A$  was significantly different from 0 and 1, the model 130 was compared to models with  $r_A$  fixed at a value of 0 or 0.999 using likelihood ratio tests.

#### 132 Genome-wide association studies.

Genome-wide association studies of ACC with the imputed SNP dataset were carried out us-133 ing function rGLS in the package RepeatABEL v1.1.31 (RÖNNEGÅRD et al., 2016) in R v3.6.2. 134 This approach fits both repeated measures and the GRM, the latter accounting for population 135 structure. Association statistics were corrected for any further inflation by dividing by the 136 genomic control parameter  $\lambda$ , calculated as the median observed  $\chi_1^2$  statistic divided by the 137 median  $\chi_1^2$  expected under a null distribution. The significance threshold at  $\alpha=0.05$  was 138 calculated by Stoffel et al. (2020) as  $P < 1.28 \times 10^{-06}$ , using a 'simpleM' approach using 139 linkage disequilibrium information to determine the effective number of independent tests 140 (GAO et al., 2008). 141

142 For the most highly-associated SNPs, genotype effect sizes were estimated by fitting the SNP 143 genotype as a three level factor in the animal models described above. Then, in separate models, the proportion of phenotypic variance explained by significant regions was then 144 determined using a regional heritability approach. For these models, GRMs were constructed 145 as above for the 20 SNPs from the SNP50 chip spanning the highest association SNP (i.e. 146 147 10 SNPs from either side of the association). In one case where the association was at the 148 end of the chromosome, the last 20 SNPs on that chromosome were used to construct the 149 GRM. These regional GRMs were then fit as additional random effects in the animal models described above, with their significance determined using likelihood ratio tests distributed as 150 151  $\chi_1^2$ .

In order to identify potential candidate genes for ACC, sheep gene IDs SNPs in the broad 152 153 associated regions were extracted from Ensembl (gene build ID Oar v3.1.100) using the function getBM in the R package biomaRt v2.42.1 (Durinck et al., 2009) in R v3.6.2. Gene 154 orthologues in humans (Homo sapiens), cattle (Bos taurus), mouse (Mus musculus) and rat 155 (Rattus norvegicus) were extracted using the function getLDS. The associated gene ontology 156 (GO annotations) information for all genes and orthologues were then extracted, again using 157 the getBM function. All gene names, phenotype descriptions, GO terms and definitions were 158 159 then queried for terms associated with meiosis and recombination, using the R command grep 160 with the text strings meio and recombin.

#### 161 Data Availability

- 162 Raw data will be publicly archived. Code for the analysis is archived at
- 163 https://github.com/susjoh/Soay Recombination 2020.

## 164 Results

## 165 Variation and heritability of autosomal crossover count.

166 Males had higher recombination rates than females, with 7.24 more autosomal crossovers observed per gamete (SE = 0.18. Wald  $\chi_1^2$  = 1613.6, P < 0.001; Figure 1). There was no 167 association with ACC and age or the inbreeding coefficient  $\hat{F}_{III}$  (P > 0.05; Figure 1). Females 168 had higher phenotypic variance than males (female  $V_P = 32.41$ , male  $V_P = 24.162$ ; Table 1). 169 ACC was heritable in both sexes ( $h^2 = 0.148$ ) and in males and females separately ( $h^2 = 0.148$ ) 170 0.118 and 0.181, respectively; Table 1); likelihood ratio tests confirmed that including the 171 additive genetic effect significantly improved the animal models ( $\chi_1^2 = 139.4$ , 19.72 and 119.9 172 for both sexes, males, and females, respectively; P < 0.001). The permanent environment 173 174 contribution to phenotypic variance was not significant in any models (P > 0.05), with the remaining variance attributed to residual effects (Table 1). Bivariate models of ACC between 175 males and females showed a positive genetic correlation ( $r_A = 0.555$ , SE = 0.138) that was 176 significantly different from both  $r_A$  = 0 and 1 ( $\chi_1^2$  = 11.95 and 17.91, respectively; P < 177 0.001). Linkage map lengths from the ACC analysis in this study were strongly correlated 178 with the previous maps (linear regression, adjusted  $R^2 > 0.999$ , Figure S1). The updated map 179 180 had 15,787 unique centiMorgan positions compared with 12,359 in the previous study; it is is 181 provided in Table S2 and Figure S2.

#### 182 Genome-wide Association Study.

There was a strong association between SNPs at the sub-telomeric region of chromosome 6 and female ACC (*oar3\_OAR6\_116402578*, P = 2.45 × 10<sup>-17</sup>; Figures 2 & S4; Tables 2 & S3). This locus explained 45.9% of the heritable variation in female ACC, with a difference of 5 crossovers per gamete between the two homozygotes (Figure S3, Table 2). There was no significant association with this locus and male ACC, confirming that its action is likely to be sex-limited. The same SNP showed the highest genome-wide association in the previous study

189 (JOHNSTON et al., 2016) and occurs 25.2 kb from the putative location of RNF212. It should 190 be noted that RNF212 is not annotated on the Oar v3.1 sheep genome due to a gap in the sequence, but is predicted to be positioned between 116,427,802 and 116,446,904 bp (John-191 192 STON et al., 2016); RNF212 is present in this position on the Oar rambouillet v1.0 genome 193 assembly (GenBank assembly ID GCA 002742125.1). A second association was observed in both sexes on chromosome 7 (oar3\_OAR7\_21368818 and oar3\_OAR7\_21347355,  $P = 6.11 \times 10^{-2}$ 194  $10^{-13}$ ,  $1.83 \times 10^{-10}$  and  $7.20 \times 10^{-7}$  in both sexes combined, females and males, respectively); 195 the association statistic in males was also identical for locus oar3 OAR7 21116299 (Figures 2 196 197 & S5; Tables 2 & S3). The action of this locus was similar between males and females, with a difference of ~4.5 crossovers per gamete between the two homozygotes (Figure S3, Table 198 199 2) and explaining between 19.7% and 24.8% of the heritable variation in ACC. These SNPs spanned the candidate locus RNF212B (21,241,831 to 21,273,165 bp). GO term analysis indi-200 201 cated that both RNF212 and RNF212B were the closest genes to each hit that were associated 202 with meiotic processes (Table S4; Figures S4 & S5).

#### 203 Discussion

204 This study revisited a previous analysis investigating the genetic architecture of ACC in Soay 205 sheep, with more than double the number of gametes and more than ten times the number of SNP loci. We confirmed that ACC is ~1.27 times higher in males than in females, and is 206 not associated with age or inbreeding in neither males nor females. We identified two candi-207 208 date genes associated with ACC: RNF212 on chromosome 6, which is associated with female 209 recombination rate only; and its paralogue RNF212B on chromosome 7, which is associated 210 with recombination rate in both sexes. Both loci have repeatedly been associated with recombination rate variation in mammals, with all studies conducted to date implicating either one 211 or both of these loci; these include humans (Kong et al., 2008, 2014), cattle (Sandor et al., 212 2012; Ma et al., 2015; Kadri et al., 2016), pigs (Johnsson et al., 2020), deer (Johnston et al., 213 2018) and domestic sheep (Petit et al., 2017). Functional studies in mice indicate that the 214 protein RNF212 is essential for crossover formation during meiosis, and that it has a dosage 215 216 sensitive effect (REYNOLDS et al., 2013). We observe an additive effect of alleles on ACC at 217 both loci in this study, suggesting that a similar dosage sensitive effect is the mechanism driving rate differences in Soay sheep. 218 219 The current study confirms the previous association at RNF212 and does not gain any novel 220 insights into the nature of its association with ACC. However, the observation at RNF212B 221 builds on tentative evidence of an association in this genomic region. In the 2016 study, a 222 genome-wide regional heritability analysis identified an association in both sexes combined 223 in a 1.09Mb region, which contained RNF212B and another candidate locus, meiotic recombination protein REC8. However, there was no corresponding association using GWAS and 224 225 there was no significant effect observed in the sex-specific regional heritability analyses, providing limited insight into the action of this region on ACC variation. In this study, we have 226 227 shown that adding a larger number of gametes and SNPs has increased the power to detect 228 an association at RNF212B, and indicating that ACC variation is more likely to be attributed 229 to RNF212B than REC8 (although see further discussion below). The MAF of RNF212B is relatively low (0.082), suggesting that lower sample sizes in the 2016 study were not sufficient 230

to detect the effects of rare loci within the population. Using the SNP50 Chip alone, the increased number of gametes in current study would shown a significant on chromosome 7 in females (but not in males; Figure S5), however, the highest association would be around 345Kb away from the that observed in the current study. Our work shows that increasing marker density using imputation is likely to be an increasingly important approach in conducting GWAS in natural populations.

Soay sheep have extensive linkage disequilibrium (LD) throughout the genome due to recent 237 population bottlenecks, a small effective population size and a high prevalence of inbreeding 238 (CLUTTON-BROCK et al., 2004; STOFFEL et al., 2020). An advantage of this is that there is a 239 higher change of typing SNP loci that are in LD with causal loci, and that genotype imputa-240 tion can be carried out with high accuracy. However, a disadvantage is that high LD within 241 242 the population makes it difficult to separate the effects of linked loci that also contribute to phenotypic variation in the population. The associated regions on both chromosomes 6 and 243 7 extend over several megabases, with several other loci associated with meiotic processes 244 occurring within these regions (Figures S4 & S5, Table S4); we cannot rule out that these 245 loci also affect variation in ACC in Soay sheep. Functional validation of the role of RNF212 246 and RNF212B is not possible within this wild system as experimental manipulation and in-247 vasive sampling are prohibited. However, further studies in domestic sheep (e.g. Petit et al. 248 249 2017) and related species may further elucidate the role of loci within these regions and the relative importance of regulatory and protein-coding variation in driving recombination rate 250

Overall, this study has highlighted the merits of genotype imputation and increasing sample sizes to determine the genetic architecture of recombination rate variation in a natural system. Identifying candidate loci and their sex-specific effect sizes provides a stepping stone for future studies investigating the evolution of recombination rates within this and related systems, through modelling the temporal dynamics of these loci, their origin and their association with individual fitness.

251

variation.

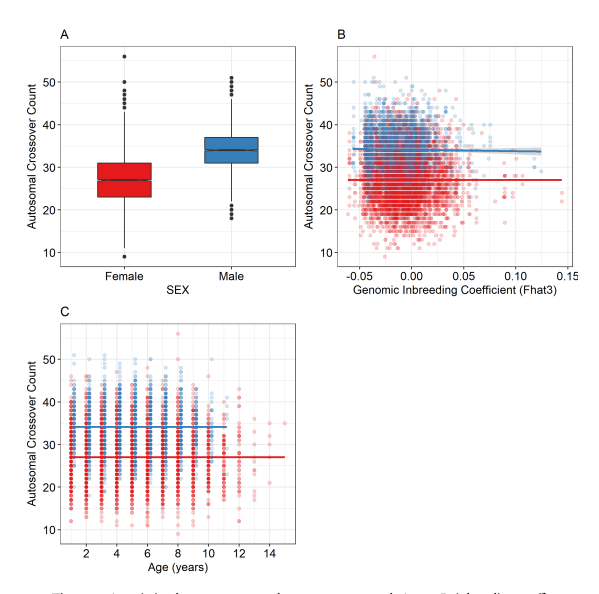
# 258 Acknowledgements

259 We thank Jill Pilkington, Ian Stevenson and all Soay sheep project members and volunteers for collection of data and samples. Camillo Bérénos, Philip Ellis and Hannah Lemon prepared 260 DNA samples for Ovine SNP50 BeadChip genotyping and Jisca Huisman assisted with the 261 pedigree construction. This work has made extensive use of the resources provided by the 262 Edinburgh Compute and Data Facility (http://www.ecdf.ed.ac.uk/). Permission to work on 263 St Kilda is granted by The National Trust for Scotland, and logistical support is provided 264 265 by QinetiQ and Eurest. The Soay sheep project has been supported by grants from the UK Natural Environment Research Council, with the SNP genotyping funded by a European 266 267 Research Council Advanced Grant to JMP. SEJ is supported by a Royal Society University Research Fellowship. 268

# 269 Author Contributions

- 270 S.E.J and J.M.P. conceived the study. J.M.P organised the collection of samples. M.S. con-
- 271 ducted the genotype imputation. S.E.J. analysed the data and wrote the paper. All authors
- 272 contributed to revisions.

# 273 Figures and Tables



**Figure 1:** Association between autosomal crossover count and: A. sex; B. inbreeding coefficient  $\hat{F}_{III}$ ; and C. age in years. Plots were made using the raw data. Points are red for females and blue for males. Lines in B. and C. are general additive model smoothing parameters as fit by the plotting library ggplot2 v3.3.2 (WICKHAM, 2016) in R v3.6.2. Points in plot C are slightly jittered between the sexes on the x-axis to more clearly visualise the sex differences with age. Sample sizes are provided in Table 1.

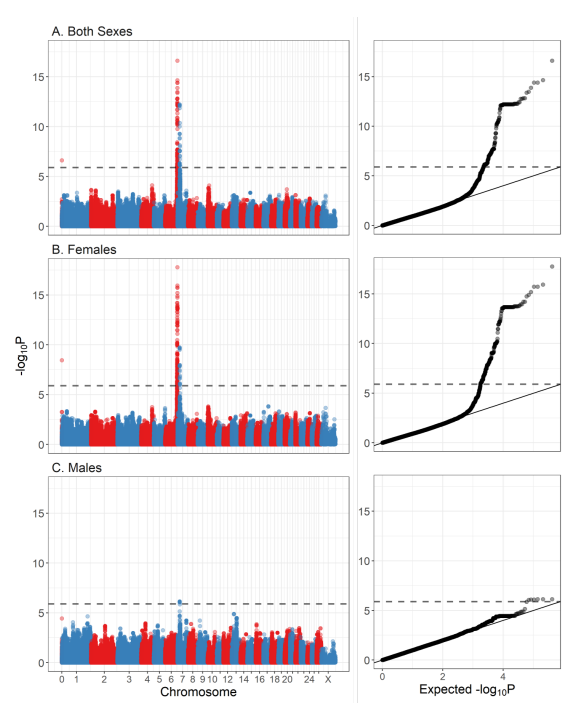


Figure 2: Genome-wide association plots of autosomal crossover counts for A. both sexes combined, B. females only and C. males only. The dashed line is the significant threshold equivalent to  $\alpha < 0.05$ . The left panels show the association for individual SNPs relative to their genomic position, with points colour-coded by chromosome. The right plots show the distribution of the observed  $-log_{10}P$  values against those expected under a null distribution. All association statistics have been corrected for genomic control using the  $\lambda$  parameter. All results are provided in Table S3. Sample sizes are provided in Table 1.

**Table 1:** Data information and animal model results for autosomal crossover count (ACC) in Soay sheep. Numbers in parentheses are the standard error unless otherwise stated.  $N_{OBS}$  is the observed number of gametes, with numbers in parentheses the number of unique focal individuals.  $N_{xovers}$  are the total number of crossovers in the dataset. The mean ACC and the variance  $(V_{OBS})$  were calculated from the raw data.  $V_P$  is phenotypic variance.  $h^2$ ,  $pe^2$  and  $e^2$  are the narrow-sense heritability, the permanent environment effect, and the residual effect, respectively; all are calculated as the proportion of  $V_P$ .

Analysis	$N_{OBS}$	$N_{xovers}$	Mean	$V_{OBS}$	$V_P$	$h^2$	$pe^2$	$e^2$
Both Sexes	7626	227646	29.851	41.32	29.514	0.148	0.025	0.827
	(1483)				(0.588)	(0.019)	(0.013)	(0.014)
Males	3043	103914	34.149	23.974	24.162	0.118	0	0.882
	(434)				(0.72)	(0.032)	(0.024)	(0.02)
Females	4583	123732	26.998	32.439	32.41	0.181	0	0.819
	(1049)				(0.797)	(0.024)	(0.017)	(0.017)

Table 2: The most highly associated SNPs from GWAS of ACC in Soay sheep. MAF is the minor allele frequency (corresponding to the A allele for each locus). Sex indicates if the model was run in both sexes combined, or in females or males only. P is the P-value for the association statistic after correction using genomic control  $\lambda$ . Effect sizes are provided for animal models where the genotype fit as a fixed factor, and are given relative to the model intercept of 0 at genotype AA. Prop.  $V_A$  is the proportion of the additive genetic variance explained by the genomic region containing the SNP locus. All values in parentheses are standard errors. Full GWAS results are provided in Table S3. Effect sizes and associated sample sizes for each genotype are shown in Figure S3.

SNP Information	MAF	Sex	P	Effect AG	Effect GG	Prop. $V_A$
oar3_OAR6_116402578	0.275	Both Sexes	$2.45 \times 10^{-17}$	2.201	3.837	0.399
Chr. 6				(0.332)	(0.361)	(0.118)
Position 116,402,578		Females	$1.68 \times 10^{-18}$	3.038	4.985	0.459
Candidate Gene: RNF212				(0.397)	(0.423)	(0.115)
		Males	0.0317	0.168	0.884	0.049
				(0.577)	(0.607)	(0.067, ns)
oar3_OAR7_21368818*	0.082	Both Sexes	$6.11 \times 10^{-13}$	-2.383	-4.520	0.242
Chr. 7				(0.922)	(0.952)	(0.102)
Position 21,368,818		Females	$1.83 \times 10^{-10}$	-2.705	-4.757	0.197
Candidate Gene: RNF212B				(1.152)	(1.194)	(0.09)
		Males	$7.20 \times 10^{-07}$	-2.479	-4.394	0.248
				(1.454)	(1.447)	(0.145)

<sup>\*</sup> This association was identical to hits at SNP loci oar3\_OAR7\_21347355 at position 21,347,355 (in both sexes, males and females) and oar3\_OAR7\_21116299 at position 21,116,299 (in males).

#### 274 References

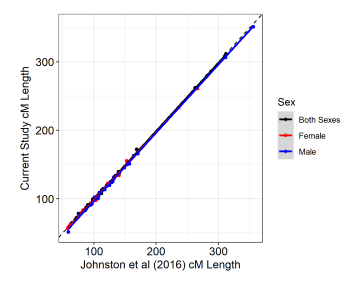
- 275 Antolín, R., C. Nettelblad, G. Gorjanc, D. Money, and J. M. Hickey, 2017 A hybrid
- 276 method for the imputation of genomic data in livestock populations. Genetics Selection
- 277 Evolution : 1–17.
- 278 BARTON, N. H., and B. CHARLESWORTH, 1998 Why sex and recombination? Science 74:
- 279 187-195.
- 280 BATTAGIN, M., G. GORJANC, A.-M. FAUX, S. E. JOHNSTON, and J. M. HICKEY, 2016 Effect of
- 281 manipulating recombination rates on response to selection in livestock breeding programs.
- 282 Genet. Select. Evol. 48: 44.
- 283 BUTLER, D. G., B. R. CULLIS, A. R. GILMOUR, and B. J. GOGEL, 2009 Mixed Models for S
- language Environments: ASReml-R reference manual.
- 285 CLUTTON-BROCK, T., J. PEMBERTON, T. N. COULSON, I. R. STEVENSON, and A. D. C. MAC-
- 286 Coll, 2004 The sheep of St Kilda. In Soay Sheep: Dynamics and Selection in an Island
- 287 Population.2. Cambridge University Press, 17–51.
- DURINCK, S., P. T. SPELLMAN, E. BIRNEY, and W. HUBER, 2009 Mapping identifiers for the inte-
- gration of genomic datasets with the R/Bioconductor package biomaRt. Nature protocols
- **4**: 1184–1191.
- 291 GAO, X., J. STARMER, and E. R. MARTIN, 2008 A multiple testing correction method for
- 292 genetic association studies using correlated single nucleotide polymorphisms. Genetic epi-
- 293 demiology **32**: 361–369.
- 294 Green, P., K. Falls, and S. Crooks, 1990 Documentation for CRIMAP, version 2.4. Washing-
- 295 ton University School of Medicine.
- 296 HALLDORSSON, B. V., G. PALSSON, O. A. STEFANSSON, H. JONSSON, M. T. HARDARSON, et al.,
- 297 2019 Characterizing mutagenic effects of recombination through a sequence-level genetic
- 298 map. Science **363**: eaau1043.

- 299 HASSOLD, T., and P. HUNT, 2001 To err (meiotically) is human: the genesis of human aneu-
- 300 ploidy. Nat. Rev. Genet. 2: 280-291.
- 301 HENDERSON, C. R., 1975 Best linear unbiased estimation and prediction under a selection
- 302 model. Biometrics **31**: 423–447.
- 303 HICKEY, J. M., B. P. KINGHORN, B. TIER, J. H. J. V. D. WERF, and M. A. CLEVELAND, 2012
- A phasing and imputation method for pedigreed populations that results in a single-stage
- genomic evaluation. Genet Sel Evol 44: 9.
- 306 HILL, W. G., and A. ROBERTSON, 1966 The effect of linkage on limits to artificial selection.
- 307 Genet. Res. 8: 269-294.
- 308 Huisman, J., 2017 Pedigree reconstruction from SNP data: parentage assignment, sibship
- 309 clustering and beyond. Mol. Ecol. Resour. 17: 1009–1024.
- 310 JOHNSSON, M., A. WHALEN, R. ROS-FREIXEDES, G. GORJANC, C.-Y. CHEN, et al., 2020 Genetics
- of recombination rate variation in the pig. bioRxiv.
- 312 JOHNSTON, S. E., C. BÉRÉNOS, J. SLATE, and J. M. PEMBERTON, 2016 Conserved genetic ar-
- 313 chitecture underlying individual recombination rate variation in a wild population of soay
- 314 sheep (Ovis aries). Genetics 203: 583–598.
- 315 JOHNSTON, S. E., J. HUISMAN, and J. M. PEMBERTON, 2018 A genomic region containing
- 316 REC8 and RNF212B is associated with individual recombination rate variation in a wild
- population of red deer (Cervus elaphus). BioRxiv.
- 318 KADRI, N. K., C. HARLAND, P. FAUX, N. CAMBISANO, L. KARIM, et al., 2016 Coding and
- noncoding variants in HFM1, MLH3, MSH4, MSH5, RNF212, and RNF212B affect re-
- 320 combination rate in cattle. Genome research **26**: 1323–1332.
- 321 Kijas, J. W., D. Townley, B. P. Dalrymple, M. P. Heaton, J. F. Maddox, et al., 2009 A
- 322 genome wide survey of SNP variation reveals the genetic structure of sheep breeds. PLoS
- 323 ONE 4: e4668.
- 324 Kong, A., J. Barnard, D. F. Gudbjartsson, G. Thorleifsson, G. Jonsdottir, et al., 2004
- Recombination rate and reproductive success in humans. Nat. Genet. **36**: 1203–1206.

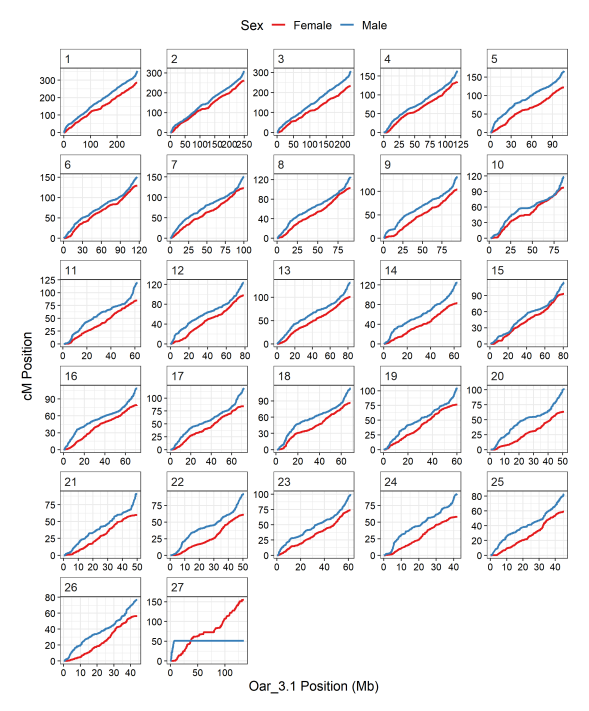
- 326 Kong, A., G. Thorleifsson, M. L. Frigge, G. Masson, D. F. Gudbjartsson, et al., 2014
- 327 Common and low-frequency variants associated with genome-wide recombination rate.
- 328 Nat. Genet. 46: 11-6.
- 329 Kong, A., G. Thorleifsson, H. Stefansson, G. Masson, A. Helgason, et al., 2008 Se-
- quence variants in the RNF212 gene associate with genome-wide recombination rate. Sci-
- 331 ence **319**: 1398–1401.
- 332 LENORMAND, T., and J. DUTHEIL, 2005 Recombination difference between sexes: a role for
- haploid selection. PLoS biology **3**: e63.
- 334 MA, L., J. R. O'CONNELL, P. M. VANRADEN, B. SHEN, A. PADHI, et al., 2015 Cattle Sex-Specific
- Recombination and Genetic Control from a Large Pedigree Analysis. PLoS Genet. 11:
- 336 e1005387.
- 337 Muller, H., 1964 The relation of recombination to mutational advance. Mutat. Res. 1: 2-9.
- 338 Myers, S., L. Bottolo, C. Freeman, G. McVean, and P. Donnelly, 2005 A fine-scale map
- of recombination rates and hotspots across the human genome. Science 310: 321–4.
- 340 Otto, S. P., and N. H. Barton, 2001 Selection for recombination in small populations.
- 341 Evolution **55**: 1921–1931.
- 342 Otto, S. P., and T. Lenormand, 2002 Resolving the paradox of sex and recombination. Nat.
- 343 Rev. Genet. 3: 252–261.
- 344 Otto, S. P., and B. A. Payseur, 2019 Crossover Interference: Shedding Light on the Evolu-
- tion of Recombination. Annual review of genetics **53**: 19–44.
- 346 Petit, M., J.-M. Astruc, J. Sarry, L. Drouilhet, S. Fabre, et al., 2017 Variation in Recom-
- bination Rate and Its Genetic Determinism in Sheep Populations. Genetics 207: 767–784.
- 348 REYNOLDS, A., H. QIAO, Y. YANG, J. K. CHEN, N. JACKSON, et al., 2013 RNF212 is a dosage-
- sensitive regulator of crossing-over during mammalian meiosis. Nat. Genet. 45: 269–78.
- 350 RÖNNEGÅRD, L., S. E. McFarlane, A. Husby, T. Kawakami, H. Ellegren, et al., 2016 Increas-
- ing the power of genome wide association studies in natural populations using repeated
- measures evaluation and implementation. Methods in Ecology and Evolution 7: 792–799.

- 353 SAMUK, K., B. MANZANO-WINKLER, K. R. RITZ, and M. A. F. NOOR, 2020 Natural Selection
- 354 Shapes Variation in Genome-wide Recombination Rate in <em>Drosophila pseudoob-
- 355 scura < /em > . Current Biology 30: 1517–1528.e6.
- 356 SANDOR, C., W. LI, W. COPPIETERS, T. DRUET, C. CHARLIER, et al., 2012 Genetic variants in
- 357 REC8, RNF212, and PRDM9 influence male recombination in cattle. PLoS Genet. 8: 1–13.
- 358 STAPLEY, J., P. G. D. FEULNER, S. E. JOHNSTON, A. W. SANTURE, and C. M. SMADJA, 2017
- 359 Variation in recombination frequency and distribution across Eukaryotes: patterns and
- 360 processes. Philos. Trans. R. Soc. Lond., B, Biol. Sci. 372: 20160455.
- 361 Stoffel, M., S. Johnston, J. Pilkington, and J. Pemberton, 2020 Genetic architecture and
- lifetime dynamics of inbreeding depression in a wild mammal. bioRxiv.
- 363 Wickham, H., 2016 ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York.
- 364 YANG, J., S. H. LEE, M. E. GODDARD, and P. M. VISSCHER, 2011 GCTA: a tool for genome-
- wide complex trait analysis. Am. J. Hum. Genet. 88: 76–82.

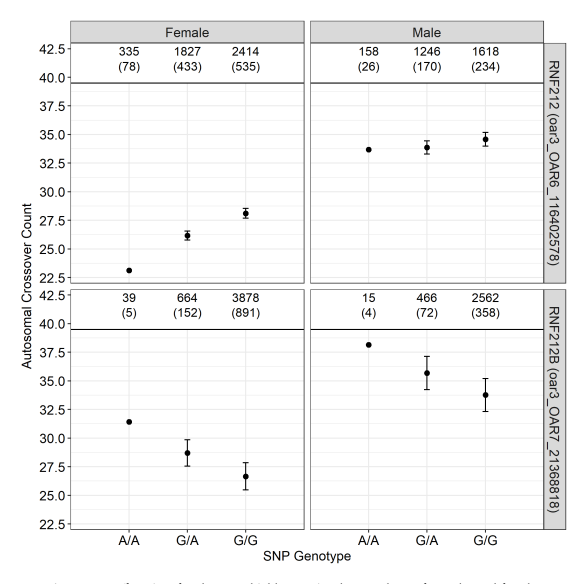
# 366 Supplementary Information



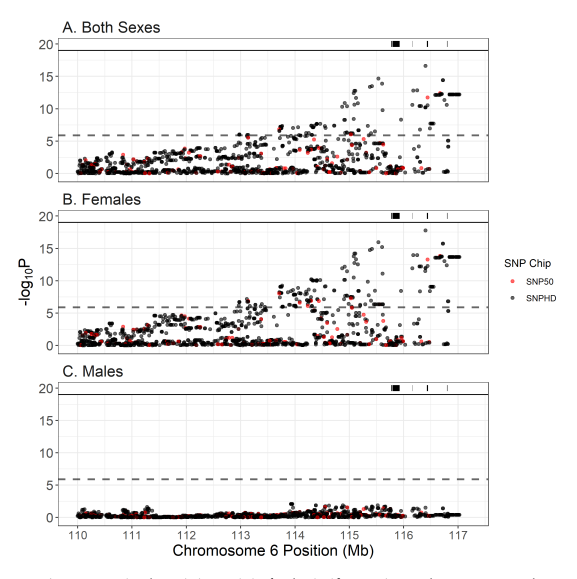
**Figure S1:** Correlation between linkage map lengths determined in the previous study (JOHNSTON *et al.*, 2016) and in the current study. Each point indicates a chromosome, and lines indicate linear regressions. Points are coloured by sex-averaged or sex-specific map types. The black dashed line indicates a perfect correlation with slope = 1 and intercept = 0



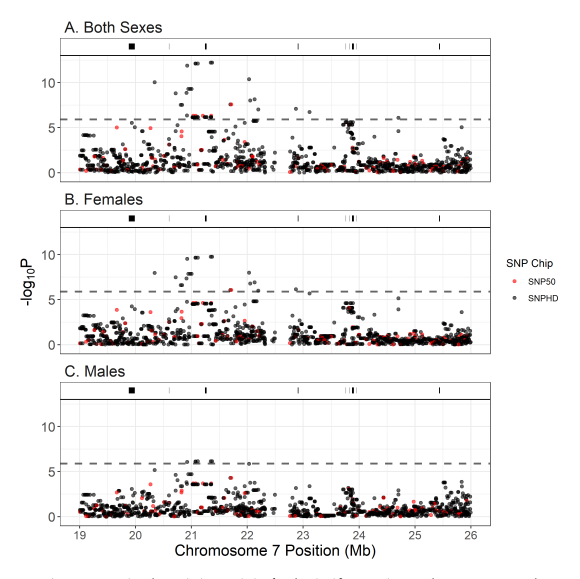
**Figure S2:** Sex-specific linkage maps for each sheep chromosome plotted relative to the sheep genome assembly Oar\_v3.1. The underlying map data is provided in Table S2. Chromosome 27 is the X chromosome, with the short map segment in male sheep corresponding to the pseudoautosomal region (PAR).



**Figure S3:** Effect sizes for the most highly associated SNPs shown for males and females, as calculated from animal models including the SNP genotype as a fixed factor. Genotype AA is the model intercept. Numbers indicate the number of observations at each genotype, with the number of unique focal individuals in parentheses.



**Figure S4:** Regional association statistics for the significant region on chromosome 6. Each point represents a single SNP locus, coloured by their origin on the SNP50 (red) or SNPHD Chips (black, i.e. imputed genotypes). The positions of genes associated with meiotic processes are shown as black bars at the top of each plot (Gene information in Table S4).



**Figure S5:** Regional association statistics for the significant region on chromosome 7. Each point represents a single SNP locus, coloured by their origin on the SNP50 (red) or SNPHD Chips (black, i.e. imputed genotypes). The positions of genes associated with meiotic processes are shown as black bars at the top of each plot (Gene information in Table S4).

**Table S1:** Fixed effects in animal models of autosomal crossover count. Models were run in both sexes, males and females. All Wald statistics were equivalent to a  $\chi^2$  test with 1 degree of freedom.

Model	Effect	Estimate	SE	Z Ratio	Wald Statistic	P
Both Sexes	(Intercept)	26.966	0.106	253.849	112596.078	0.0000
	Fhat3	1.148	3.692	0.311	0.097	0.7557
	Sex (Male)	7.236	0.180	40.161	1613.634	0.0000
Males	(Intercept)	34.178	0.144	236.710	61239.042	0.0000
	Fhat3	-3.402	5.872	-0.579	0.336	0.5623
Females	(Intercept)	26.970	0.113	238.303	59962.846	0.0000
	Fhat3	0.853	4.610	0.185	0.034	0.8533

Table S2: Linkage map information for the current study. Order is the order of markers on the chromosome. SNP.Name is the Ovine SNP50 BeadChip identifier. Chr is the chromosome number, where 27 is the X chromosome. cMPosition, r, cMdiff are the sex-averaged centiMorgan position, recombination fraction and centiMorgan difference with the following locus, respectively; similarly this is provided for Female and Male maps. Oar3\_Chr and Oar3\_Pos are the relative chromosome number and position relative to the sheep genome (Oar\_v3.1), cMPosition.2016, cMPosition.Female.2016 and cMPosition.Male.2016 indicate the centiMorgan positions as determined by the previous study (JOHNSTON et al., 2016).

[File: Table\_S2\_Linkage\_Map.txt]

**Table S3:** Association statistics for genome-wide association studies of autosomal chromosome counts. Sex indicates analyses for both sexes, females only and males only. SNP.Name is the Ovine SNP50 BeadChip identifier. Chromosome and Position (bp) are given relative to the sheep genome Oar\_v3.1. A1 and A2 reference and alternate allele at each SNP. effB is the slope of the effect of allele A2, with the standard error se\_effB. Chi2.1df and P1df is the association chi-squared statistic and associated P-value, respectively, before correction with genomic control  $\lambda$ . Pc1df is the corrected P-value after genomic control. Exp is the corresponding P-value for that SNP locus assuming a null distribution of P-values (see Figure 2). Q.2 is the minor allele frequency. Cumu is the cumulative genomic distance.

[File: Table\_S3\_ACC\_GWAS.txt]

Table S4: Gene positions in the GWAS significant regions obtained using biomaRt v2.42.1. ensembl\_gene\_id is the Ensembl identifier for the sheep gene. external\_gene\_name is the sheep gene name. chromosome\_name, start\_position and end\_position are the chromosome number, start and stop positions for each gene, respectively. Meiotic indicates whether this gene and/or its orthologues in other species have GO terms associated with meiotic processes. Orthos indicates the unique gene names associated with orthologues of each gene.

[File: Table\_S4\_Gene\_List\_Table.txt]