1 Genomic prediction with whole-genome

2 sequence data in intensely selected pig lines

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15

Abstract

Background

16 Early simulations indicated that whole-genome sequence data (WGS) could improve 17 prediction accuracy and its persistence across generations and breeds. However, 18 results in real datasets have been ambiguous so far. Large data sets that capture most 19 of the genome diversity in a population must be assembled so that allele substitution 20 effects are estimated with higher accuracy. The objectives of this study were to use a 21 large pig dataset to assess the benefits of using WGS for genomic prediction 22 compared to using commercial marker arrays, to identify scenarios in which WGS 23 provides the largest advantage, and to identify potential pitfalls for its effective implementation. 24

Methods

We sequenced 6,931 individuals from seven commercial pig lines with different numerical size. Genotypes of 32.8 million variants were imputed for 396,100 individuals (17,224 to 104,661 per line). We used BayesR to perform genomic prediction for 8 real traits and 9 simulated traits with different genetic architectures. Genomic predictions were performed using either data from a marker array or variants preselected from WGS based on linkage disequilibrium, functional annotation, or association tests. Both single and multi-line training sets were explored.

Results

Using WGS improved prediction accuracy relative to the marker array, provided that training sets were sufficiently large, especially for traits with high heritability and low number of quantitative trait nucleotides. The performance of each set of predictor variants was not robust across traits and lines. The most robust results were obtained

- when preselected variants with statistically significant associations were added to the
 marker array. Under this method, average improvements of prediction accuracy of 2.5
 and 4.2 percentage points were observed in within-line and multi-line scenarios,
 respectively, with training sets of around 80k individuals.
 Conclusions
- 40 Our results evidenced the potential for WGS to improve genomic prediction accuracy
- 41 in intensely selected pig lines. Although the prediction accuracy improvements
- 42 achieved so far were modest at best, we would expect that more robust improvements
- 43 could be attained with a combination of larger training sets and optimised pipelines.

44

Introduction

45 Whole-genome sequence data (WGS) has the potential to empower the 46 identification of causal variants that underlie quantitative traits or diseases [1–4], 47 increase the precision and scope of population genetic studies [5,6], and enhance 48 livestock breeding. Genomic prediction has been successfully implemented in the 49 main livestock species and it has increased the rate of genetic gain [7]. Genomic 50 prediction has provided many benefits such as greater accuracies of genetic 51 evaluations and the reduction of the generational interval in dairy cattle. However, 52 since its early implementations, genomic prediction is typically performed using 53 marker arrays that capture the effects of the (usually unknown) causal variants via 54 linkage disequilibrium. Alternatively, WGS are assumed to contain the causal variants 55 themselves. For this reason, it was hypothesized that such data could further improve 56 prediction accuracy and its persistence across generations and breeds. Early 57 simulations indicated that causal mutations from WGS could increase prediction 58 accuracy [8–13]. One simulation study indicated that the magnitude of prediction 59 accuracy improvement relative to dense marker arrays ranged from 2.5 to 3.7%, with 60 a persistence of over 10 generations [11]. Another one reached improvements of 30% 61 if causal variants with low minor allele frequency could be captured by the WGS [9]. 62 However, benefits could be on the lower end of that range in standard livestock 63 populations due to small effective population sizes and long-term negative selection 64 [10].

During the last few years, there have been several attempts at improving the
accuracy of genomic prediction with the use of WGS in the main livestock species.
Results have been ambiguous so far. When predicting genomic breeding values within
breed or line, some studies found no relevant improvement of prediction accuracy for

WGS compared to marker arrays [14–18]. Other studies found small, and often unstable, improvements (e.g., from 1 to 5% or no improvement depending on prediction method [19–21], or trait-dependent results [21,22]). When predicting genomic breeding values across populations, the identification of causal variants from WGS can improve prediction accuracy [23–26], especially for small populations where initial prediction accuracy was low or that were not included in the training population [23,25–28].

76 One of the most successful strategies to exploit WGS consists in augmenting 77 available marker arrays with preselected variants from WGS based on their 78 association with the trait of interest [29–32]. In some cases, this strategy improved 79 prediction accuracy by up to 9% [31] and 11% [32]. However, it did not improve 80 prediction accuracies in other within-line scenarios [16]. Nevertheless, this shows 81 how identifying causal variants could enhance genomic prediction with WGS. Whole-82 genome sequence data has already been applied in genome-wide association studies 83 (GWAS) to identify variants associated to a variety of traits in livestock [2,33–35], 84 including pigs [36,37]. However, the fine-mapping of causal variants remains 85 challenging due to the pervasive long-range linkage disequilibrium across extremely 86 dense variation.

High accuracy in estimating allele substitution effects and, ideally, the identification of causal variants amongst millions of other variants are important for the usefulness of WGS in research and breeding. This requires large data sets able to capture most of the genome diversity in a population. Despite that low-cost sequencing strategies have been developed, which typically involve sequencing a subset of the individuals in a population at low coverage and then imputing WGS for the remaining individuals [38–40], the cost of generating accurate WGS at this scale,

94 as well as the large computational requirements for the analyses of such datasets, have 95 limited the population sizes or number of populations tested in some of the previous 96 studies. This hinders the interpretation of results across studies, which are very 97 diverse in population structures, sequencing strategies and prediction methodologies 98 used. The largest studies on the use of WGS for genomic prediction to date have been 99 performed in cattle, for which large multi-breed reference panels are available from 100 the 1000 Bull Genomes Project [2,19,33]. This has enabled the imputation of WGS 101 for cattle populations. The lack of such available reference panels has been cited as an 102 important limiting factor for performing similar studies in other species, such as pigs 103 [36].

104 We have previously described our approach to impute WGS in large pedigreed 105 populations without the need for haplotype phased reference panels [41]. Following 106 that strategy, we generated WGS for 396,100 pigs from seven intensely selected lines 107 with diverse genetic backgrounds and numerical size. The objectives of this study 108 were to use this large pig dataset to assess the benefits of using WGS for genomic 109 prediction compared to using commercial marker arrays, to identify scenarios in 110 which WGS provides the largest advantage, and to identify potential pitfalls for its 111 effective implementation.

112

Materials and Methods

Populations and sequencing strategy

We performed whole-genome re-sequencing of 6,931 individuals from seven commercial pig lines (Genus PIC, Hendersonville, TN) with a total coverage of approximately 27,243x. Sequencing effort in each of the seven lines was proportional to population size. Approximately 1.5% (0.9 to 2.1% in each line) of the pigs in each line were sequenced. Most pigs were sequenced at low coverage, with target coverage
of 1 or 2x, but a subset of pigs were sequenced at higher coverage of 5, 15, or 30x.
Thus, the average individual coverage was 3.9x, but the median coverage was 1.5x.
The number of pigs sequenced and at which coverage for each line is summarized in
Table 1.

122 The sequenced pigs and their coverage were selected following a three-part 123 sequencing strategy developed to represent the haplotype diversity in each line. First 124 (1), sires and dams with the highest number of genotyped progeny were sequenced at 125 2x and 1x, respectively. Sires were sequenced at a greater coverage because they 126 contributed with more progeny than dams. Then (2), the individuals with the greatest 127 genetic footprint on the population (i.e., those that carry more of the most common 128 haplotypes) and their immediate ancestors were sequenced at a coverage between 1x 129 and 30x (AlphaSeqOpt part 1; [42]). The sequencing coverage was allocated with an 130 algorithm that maximises the expected phasing accuracy of the common haplotypes 131 from the cumulated family information. Finally (3), pigs that carried haplotypes with 132 low cumulated coverage (below 10x) were sequenced at 1x (AlphaSeqOpt part 2; 133 [43]). Sets (2) and (3) were based on haplotypes inferred from marker array genotypes 134 (GGP-Porcine HD BeadChip; GeneSeek, Lincoln, NE), which were phased and 135 imputed using AlphaPhase [44] and AlphaImpute [45].

Most sequenced pigs and their relatives were also genotyped either at low density (15k markers) using the GGP-Porcine LD BeadChip (GeneSeek) or at high density (80k markers) using the GGP-Porcine HD BeadChip (GeneSeek). Quality control of the marker array data was based on the individuals genotyped at high density. Markers with minor allele frequency below 0.01, call rate below 0.80, or that 141 failed the Hardy-Weinberg equilibrium test were removed. After quality control,

142 38,634 to 43,966 markers remained in each line.

143

Sequencing and data processing

144 Tissue samples were collected from ear punches or tail clippings. Genomic 145 DNA was extracted using Qiagen DNeasy 96 Blood & Tissue kits (Qiagen Ltd., 146 Mississauga, ON, Canada). Paired-end library preparation was conducted using the 147 TruSeq DNA PCR-free protocol (Illumina, San Diego, CA). Libraries for 148 resequencing at low coverage (1 to 5x) were produced with an average insert size of 149 350 bp and sequenced on a HiSeq 4000 instrument (Illumina). Libraries for 150 resequencing at high coverage (15 or 30x) were produced with an average insert size 151 of 550 bp and sequenced on a HiSeq X instrument (Illumina). All libraries were 152 sequenced at Edinburgh Genomics (Edinburgh Genomics, University of Edinburgh, 153 Edinburgh, UK).

154 DNA sequence reads were pre-processed using Trimmomatic [46] to remove 155 adapter sequences from the reads. The reads were then aligned to the reference 156 genome Sscrofall.1 (GenBank accession: GCA 000003025.6) using the BWA-MEM 157 algorithm [47]. Duplicates were marked with Picard 158 (http://broadinstitute.github.io/picard). Single nucleotide polymorphisms (SNPs) and 159 short insertions and deletions (indels) were identified with the variant caller GATK 160 HaplotypeCaller (GATK 3.8.0) [48,49] using default settings. Variant discovery with 161 GATK HaplotypeCaller was performed separately for each individual and then a joint 162 variant set for all the individuals in each population was obtained by extracting the 163 variant positions from all the individuals.

164 We extracted the read counts supporting each allele directly from the aligned reads stored in the BAM files using a pile-up function to avoid biases towards the 165 166 reference allele introduced by GATK when applied on low-coverage WGS [50]. That 167 pipeline the tool pysam (version 0.13.0; https://github.com/pysamuses 168 developers/pysam), which is a wrapper around htslib and the samtools package [51]. 169 We extracted the read counts for all biallelic variant positions, after filtering variants 170 in potential repetitive regions (defined as variants that had mean depth values 3 times 171 greater than the average realized coverage) with VCFtools [52]. This amounted to a 172 total of 55.6 million SNP (19.6 to 31.1 million within each line) and 10.2 million 173 indels (4.1 to 5.6 million within each line). A more complete description of the 174 variation across the lines is provided in [53].

175

Genotype imputation

176 Genotypes were jointly called, phased and imputed for a total of 483,353 177 pedigree-related individuals using the 'hybrid peeling' method implemented in 178 AlphaPeel [54,55]. This method used all the available marker array and WGS. Imputation was performed separately for each line using complete multi-generational 179 180 pedigrees, which encompassed from 21,129 to 122,753 individuals each (Table 1). 181 We have previously published reports on the accuracy of imputation in the same 182 populations using this method [41]. The estimated average individual-wise dosage 183 correlation was 0.94 (median: 0.97). Individuals with low predicted imputation 184 accuracy were removed before further analyses. An individual was predicted to have 185 low imputation accuracy if itself or all of its grandparents were not genotyped with a 186 marker array or if it had a low degree of connectedness to the rest of the population. 187 These criteria were based on the analysis of simulated and real data on imputation

188 accuracy [41]. A total of 396,100 individuals remained, with each line comprising 189 between 17,224 and 104,661 individuals (Table 1). The expected average individual-190 wise dosage correlation of the remaining individuals was 0.97 (median: 0.98) 191 according to our previous estimates. We also excluded from the analyses variants with a minor allele frequency lower than 0.023, as their estimated variant-wise dosage 192 193 correlations was lower than 0.90 [41]. After imputation, 32.8 million variants (14.5 to 194 19.9 million within each line) remained for downstream analyses, out of which 9.9 195 million segregated across all seven lines.

196

Traits

197 We analysed data of 8 traits that are commonly included in selection 198 objectives of pig breeding programmes: average daily gain (ADG, g), backfat 199 thickness (BFT, mm), loin depth (LD, mm), average daily feed intake (ADFI, kg), 200 feed conversion ratio (FCR), total number of piglets born (TNB), litter weight at 201 weaning (LWW, kg), and return to oestrus 7 days after weaning (RET, binary trait). 202 Most pigs with records were born during the 2008–2020 period. Breeding values were 203 estimated by line with a linear mixed model that included polygenic and non-genetic 204 (as relevant for each trait) effects. Deregressed breeding values (dEBV) were obtained 205 following the method by VanRaden and Wiggans [56]. Only individuals in which the 206 trait was directly measured were retained for further analyses. The number of records 207 for each trait used in the analyses of each line is detailed in Table 2.

208

209 Simulated traits

To assist in the interpretation of results, we also created 9 simulated traits with different numbers of quantitative trait nucleotides (QTN; 100, 1,000 or 10,000 QTN)

and heritability levels (h²; 0.10, 0.25 or 0.50). Positions of the QTN were sampled 212 213 randomly amongst all variants called across all lines. Because QTN were sampled 214 from all variants, some QTN were fixed in some of the lines while segregating in 215 others. There were only negligible differences in the number of segregating OTN per 216 line (53 to 61, 531 to 583, or 5375 to 6058, respectively). Marker effects of the OTN 217 were sampled from a gamma distribution with shape=2 and scale=5. After a polygenic 218 term was calculated for each individual using these marker effects, residual terms 219 were sampled from a normal distribution with a variance parameter adjusted to 220 produce the desired heritability level. The number of records for the simulated traits is 221 detailed in Table 2. In these simulations, we used the imputed genotypes as real 222 genotypes and, therefore, implicitly cancelled any errors that might arise from the 223 processing of the sequencing reads and genotype imputation.

224

Training and testing sets

225 We split the individuals in each population into training and testing sets. The 226 testing sets were defined as those individuals from full-sib families from the last 227 generation of the pedigree (i.e., individuals that did not have any progeny of their 228 own). Only families with a minimum of 5 full-sibs were considered. The training set 229 was defined as all those individuals that had a pedigree coefficient of relationship 230 lower than 0.5 with any individual of the testing set. This design was chosen to mimic 231 a realistic situation in which breeding companies evaluate the selection candidates 232 available in the selection nucleus at any given time.

233

Genome-wide association study

To assess whether variants from the WGS could provide a finer mapping of causal variants than marker array data, and to provide an association-based criterion to preselect variants for the genomic prediction tests, we performed a GWAS for each trait and line. This step included only the individuals in the training set. We fitted a univariate linear mixed model that accounted for the genomic relationship matrix as:

$$\mathbf{y} = \mathbf{x}_i \boldsymbol{\beta}_i + \mathbf{u} + \mathbf{e},$$

240 where y is the vector of dEBV, \mathbf{x}_i is the vector of genotypes for the *i*th SNP coded as 0 and 2 if homozygous for either allele or 1 if heterozygous, β_i is the additive effect 241 of the *i*th SNP on the trait, $\mathbf{u} \sim N(0, \sigma_{u}^{2}\mathbf{K})$ is the vector of polygenic effects with the 242 covariance matrix equal to the product of the polygenic additive variance σ_{μ}^2 and a 243 genomic relationship matrix **K**, and **e** is a vector of uncorrelated residuals. Due to 244 245 computational limitations, the genomic relationship matrix **K** was calculated using 246 only imputed SNP genotypes in the marker array regardless of whether the association study involves the SNPs in the marker array or the variants in WGS. We used the 247 248 FastLMM software [57,58] to fit the model.

We used the same p-value threshold ($p < 10^{-6}$) for both marker array and for 249 250 sequence associations, because while the WGS contains many more variants, they are 251 also expected to be in higher linkage disequilibrium. This threshold was based on 252 Bonferroni's multiple test correction assuming that the markers from the marker array 253 were independent. For the simulated traits, we defined genomic regions that contained 254 significant associations and assessed whether or not they contained a QTN. These 255 regions were defined by overlapping 500-kb segments centered on the significant 256 markers.

257

Genomic prediction in within-line scenarios

258 To test whether variants from the WGS could provide greater prediction 259 accuracy than the marker array, we tested genomic prediction using variants from the 260 marker array, from the WGS, or combining them. The marker array data (referred to 261 as 'Chip') was set as the benchmark for prediction accuracy. It contained all ~40k 262 variants in the marker array. For the sequence-based predictors, we preselected sets of 263 variants because currently available methods for genomic prediction are not yet 264 capable of handling datasets as large as the complete WGS. We tested different 265 alternative strategies for preselecting the predictor variants:

LDTags. Tag variants retained after pruning based on linkage disequilibrium.
 Variants were removed so that no pairs of SNPs with r²>0.1 remained in any 10 Mb window (windows slid by 2,000 variants) using Plink 1.9 [59]. The number
 of predictor variants preselected by this method was on average of 30k variants
 (range: 5k to 80k).

Top40k. Variants preselected based on GWAS analyses. To mimic the number of variants in Chip, we preselected the variants with the lowest p-value (not necessarily below the significance threshold) in each of consecutive non-overlapping 55-kb windows along the genome. In addition, to test the impact of variant density on prediction accuracy, we preselected 10k, 25k, 75k, or 100k predictor variants following the same criterion.

ChipPlusSign. Variants preselected based on GWAS analyses as in Top40k, but
 only significant variants (p≤10⁻⁶) were preselected and merged with those in
 Chip. When a 55-kb window contained more than one significant variant, only
 that with the lowest p-value was selected as a proxy, in order to reduce the

preselection of multiple significant SNPs tagging the same causal variant. On
average, 309 significant variants were identified per trait and line (range: 23 to
1083; Table 3). These significant variants were merged with those in Chip.

Functional. Variants that were annotated as loss-of-function or missense according to Ensembl Variant Effect Predictor (Ensembl VEP; version 97, July 2019) [60]. The most severe predicted consequence type for each variant was retrieved. The number of predictor variants preselected by this method was on average of 35k variants (range: 27k to 40k).

• *Rand40k*. The same number of predictor variants as in Chip, chosen randomly.

Genomic prediction was performed by fitting a univariate model with BayesR [61,62], with a mixture of normal distributions as the prior for variant effects, including one distribution that sets the variant effects to zero. The model was:

$\mathbf{y} = \mathbf{1}\boldsymbol{\mu} + \mathbf{X}\boldsymbol{\beta} + \mathbf{e},$

293 where **y** is the vector of dEBV, **1** is a vector of ones, μ is the general mean, **X** is a 294 matrix of genotypes, β is a vector of variant effects, and e is a vector of uncorrelated 295 residuals. The prior variance of the variant effects in β had four components with variances $\sigma_1^2 = 0$, $\sigma_2^2 = 0.0001\sigma_g^2$, $\sigma_3^2 = 0.001\sigma_g^2$, or $\sigma_4^2 = 0.01\sigma_g^2$, where σ_g^2 is the 296 297 total genetic variance. We used a uniform and almost uninformative prior for the 298 mixture distribution. We used a publicly available implementation of BayesR 299 (https://github.com/syntheke/bayesR; accessed on 30 April 2021), with default 300 settings. Prediction accuracy was calculated in the testing set as the correlation 301 between the genomic estimated breeding value and the dEBV. Bias of the prediction 302 accuracy was calculated as the regression coefficient of the dEBV on the genomic 303 estimated breeding values.

304 It has been noted that using the same reference individuals for preselecting 305 variants through GWAS and for training the predictive equation can reduce prediction 306 accuracy and bias the predicted breeding values [16,63]. To account for that, we 307 reanalysed some of the scenarios after splitting the training set into two exclusive 308 subsets, one for GWAS to preselect the predictor variants and one for training the 309 predictive equation. The GWAS subset was defined by randomly selecting either 10% 310 or 50% of the individuals in the original training set. Those individuals were excluded 311 from the subset used for training the predictive equation afterwards.

312

Genomic prediction in multi-line scenarios

313 We considered multi-line scenarios in which the training set consisted of 314 merging the training sets that had been defined for each line. All analyses were 315 performed as for the within-line scenarios but with a line effect. In the multi-line 316 scenarios, all SNPs from the marker array that passed quality control and were 317 imputed for at least one line were included in the baseline (referred to as 'ML-Chip'). 318 For ease of computation, the strategies for preselection of predictor variants from WGS were applied only to the subset of 9.9 million variants that had been called and 319 320 imputed in all seven lines. Thus, we defined the predictor sets 'ML-Top40k' and 321 'ML-ChipPlusSign' by preselecting variants following the same criteria as in within-322 line scenarios, but using a multi-line GWAS analyses with line effect instead. For 323 ML-ChipPlusSign, 60 to 7247 significant variants were identified per trait (Table 3) 324 and merged with those in ML-Chip. For comparison purposes, prediction accuracy 325 was calculated for the testing set of each individual line.

326

Results

Prediction accuracy within line

327 Whole-genome sequence data can improve prediction accuracy of marker 328 array data when there is a sufficiently large training set and if an appropriate set of 329 predictor variants is preselected. Figure 1 shows the prediction accuracy for the case 330 with the largest training set using different sets of predictor variants. In this case, all 331 tested sets of variants from the WGS, except for LDTags, yielded increases of 332 prediction accuracy that ranged from +2.0% to +9.2%. Using WGS also reduced bias 333 relative to Chip in some scenarios. However, the performance across predictor 334 variants set was not robust for the most part, and differed for each trait and line 335 (Additional File 1), often leading to no improvements of prediction accuracy or even 336 reduced prediction accuracy relative to Chip. One stable feature of the results was 337 LDTags showing a noticeable decrease in prediction accuracy in most traits and lines.

338 The size of the training set was one of the main factors that determined the 339 capacity of predictor variants from the WGS to improve the baseline prediction 340 accuracy of Chip. Figures 2 and 3 show the difference in prediction accuracy of 341 Top40k and ChipPlusSign with respect to the baseline of Chip against the number of 342 phenotypic records available in the training set. We observed large variability for the 343 difference in prediction accuracy, especially when the training set was small. This 344 variability was larger in Top40k than in ChipPlusSign, in a way that shrinkage of 345 variation as the training set was larger was more noticeable in ChipPlusSign. Gains in 346 prediction accuracy were low-to-moderate in the most favourable cases. In the most 347 unfavourable ones we observed large losses in prediction accuracy for Top40k but 348 more restrained losses for ChipPlusSign with moderate training set sizes. For both 349 sets of predictor variants, there was a positive trend that supported the need for large

training sets. This trend was clearer in ChipPlusSign than in Top40k, because of the
apparent lower robustness of the latter. Results for the other sets of predictor variants
are provided in Additional File 2.

353 The genetic architecture of the traits was also related to the success of WGS for improving prediction accuracy. As the true genetic architecture of real complex 354 355 traits is mostly unknown, we used simulated traits to show that traits with high 356 heritability and low number of QTN were more likely to show larger improvements in 357 predictive performances. With Top40k (Figure 4), heritability seemed to be the main 358 factor that affected the expected improvement with large training sets (from null improvements when $h^2=0.1$ to improvements of approximately 0.05 when $h^2=0.5$, 359 regardless of number of QTN, with a training set of 92k individuals). With 360 361 ChipPlusSign (Figure 5), the expected improvements with the same training set (92k 362 individuals) were not only greater in magnitude but depended on both heritability and number of OTN (from null improvements when $h^2=0.1$ to improvements of 363 approximately 0.03 to 0.10 when $h^2=0.5$ with a number of 100 to 10k OTN. 364 respectively). Results confirmed the trends observed for the real traits (Figures 4 and 365 366 5); for instance, the higher robustness of ChipPlusSign compared to Top40k.

We observed diminishing returns when we increased the density of the predictor variants. Increasing the number of predictor variants from the 40k in Top40k to 75k selected in the same way yielded small improvements in prediction accuracy compared to Top40k, but increases up to 100k variants provided smaller or null additional gains (Additional File 3).

372 Splitting the original training set into two exclusive subsets, one for the 373 GWAS-based preselection of the variants and one for the training of the predictive 374 equation did not improve the prediction accuracy (Additional File 4). For

375 ChipPlusSign, this strategy reduced the bias but prediction accuracy decreased too,

376 probably because of the smaller subset available for training the predictive equation.

377

Prediction accuracy in multi-line scenarios

378 The performance of genomic predictors trained with multi-line datasets was 379 systematically lower than in the within-line scenarios (Additional File 5). 380 Nonetheless, the ML-ChipPlusSign predictor variants in general increased prediction 381 accuracy relative to ML-Chip (Figure 6). The increase in genomic prediction accuracy 382 for each line was largely dependent on the number of individuals of each line in the 383 training set. Therefore, the greatest improvements were achieved for the largest lines. 384 However, in the multi-line scenarios we observed increases of prediction accuracy for 385 some traits and lines for which no improvements were observed in the within-line 386 scenarios (Figure 7). In contrast, results for ML-Top40k were not robust (Additional 387 File 6).

388

Association tests

389 First, we assessed the performance of GWAS using the simulated traits. Table 390 4 shows the number of regions with significant associations that were detected using 391 either Chip or WGS, and whether they contained zero, one or multiple true QTN. The 392 WGS allowed the detection of a much larger proportion of true OTN than the Chip, 393 especially for the traits with high heritability and with large population sizes. The 394 most favourable scenarios for identifying regions that contained unequivocally a 395 single QTN with WGS were those in which the trait was controlled by a low number 396 of true QTN. However, even though the genetic architecture was very simple and 397 consisted of additive effects alone, the regions with significant associations only 398 captured a small fraction of the QTN that segregated within each line. Moreover, 399 using WGS also increased the number of regions with significant associations that 400 contained no QTN, which could therefore be considered as false positives. Some of 401 the selected regions contained multiple OTN, which could indicate either a 'hit by 402 chance' or an inability to disentangle multiple causal variants. While false positives 403 also occur with Chip, their incidence was more severe with the WGS, especially for 404 traits with a large number of QTN. Large population sizes further aggravated the 405 inflation of genome-wide p-values.

406 Despite this, with the real traits we found that GWAS using WGS can 407 contribute to a better understanding of the genetic mechanisms that underlie the traits 408 of interest. To illustrate this, we examined the GWAS results for BFT in line A, for 409 which a large number of phenotypic records were available. Figure 8 shows the 410 results for chromosome 1 as an example, while Additional File 7 shows the results for 411 six genomic regions of interest. The main genomic regions and candidate genes 412 associated to BFT detected with Chip in the same genetic lines studied here were 413 reported elsewhere [64]. We will use the candidate genes reported there to refer to the 414 genomic regions with significant associations. Using Chip, we identified 6 genomic regions ($p < 10^{-6}$). Using WGS (with a more stringent significance threshold of $p < 10^{-9}$ 415 416 to focus on the most significant associations), we confirmed 3 of these genomic regions that co-located to candidate genes MC4R, DOLK, and DGKI or PTN. 417 418 However, the most associated variants in each of these genomic regions located 419 outside the coding region of these putative causal genes. These signals sometimes had 420 very strong evidence of association for some variants that were relatively distant from 421 our candidate functional gene, which could cast doubts about the fine-mapping of the 422 causal mutation. The region at SSC18, 9-13 Mb, contained two candidate genes

423 DGKI and PTN, but the WGS revealed significantly associated variants within DGKI 424 and none within *PTN*, despite that the strongest associations were away from both 425 genes at 10.5-11 Mb. Using the WGS we also detected 24 additional genomic regions that contained candidate genes such as CYB5R4, IGF2, and LEPR. These genes were 426 427 previously detected in other lines using the Chip but not in this one [64], sometimes 428 because there were no markers for the associated region in Chip (SSC2, 0-4 Mb). The 429 region at SSC1, 52.5-53.5 Mb, showed many significant variants that encompassed 430 not only the previously identified candidate gene CYB5R4, but also MRAP2 431 (annotated with functions on feeding behavior and energy homeostasis). In contrast, 432 candidate gene LEPR was located within the region at SSC6, 146.5–147.0 Mb, where 433 many significant variants were located, although the most significant variants were 434 not in the coding regions of the gene. Using the WGS we also identified additional 435 candidate genes that had not been previously detected in any of the lines, such as 436 CYP24A1 (annotated with functions on fatty acid omega-oxidation and vitamin D 437 metabolism; not shown). For many of the other genomic regions, it was difficult to 438 pinpoint a candidate gene with the available information or there were no annotated 439 genes.

440

Discussion

Our results evidenced the potential for WGS to improve genomic prediction accuracy in intensely selected pig lines, provided that the training sets are large enough. Improvements achieved so far were modest at best. On one hand, these modest improvements indicated that the strategies that we tested were likely suboptimal. On the other hand, the positive trend for the largest training sets indicated that we might have not reached the critical mass of data that is needed to leverage the 447 potential of WGS, especially in scenarios where genomic prediction with marker arrays is already yielding high accuracy. The results from several traits and lines with 448 449 different training set sizes and the use of simulated phenotypes allowed us to identify 450 the most favourable scenarios for genomic prediction with WGS. We will discuss (1) 451 the prediction accuracy that we achieved with WGS compared to commercial marker 452 array data and the scenarios in which WGS may become beneficial, and (2) the 453 potential pitfalls for its effective implementation and the need for an optimised 454 strategy.

455

Prediction accuracy with whole-genome sequence data

456 We compared the genomic prediction accuracy of the current marker array 457 (Chip) with sets of preselected sequence variants in a way that the number of variants 458 remained similar across sets. Improvements of prediction accuracy can be limited if 459 current marker arrays are already sufficiently dense to capture a large proportion of 460 the genetic variance in intensely selected livestock populations. These populations 461 typically have small effective population size [10,19]. Nevertheless, modest improvements have been achieved under certain scenarios. In our study, the most 462 463 robust results were obtained for the ChipPlusSign set, where variants that showed 464 statistically significant associations to the trait were preselected and added to the 465 information from the marker array. This is consistent with previous reports that showed an improvement of prediction accuracy under similar approaches [29-32]. 466 467 We added 23 to 1083 significant variants to those in Chip in different scenarios. In the 468 most successful ones, at least around 200 significant variants were added and average 469 improvements of prediction accuracy of 2.5 percentage points were observed with 470 training sets of around 80k individuals. In other instances, however, additions of a

471 larger number of variants have been proposed. The addition of 1623 variants 472 (preselected as the combination of 3-5 variants for each of the top QTL per trait and 473 breed) to a 50k array increased prediction reliability (accuracy squared) by up to 5 474 percentage points in Nordic cattle [29]. Adding the 16k SNPs with largest estimated 475 effects to a 60k array increased prediction reliability on average by 2.7 (up to 4.8) 476 percentage points in Holstein cattle [30]. For the custom 50k array for Hanwoo cattle, 477 it has been reported that adding at least around 12k SNPs (3k for each of four traits) 478 improved prediction accuracy by up to ~6 percentage points [32]. The addition of 479 ~400 variants preselected by GWAS with regional heritability mapping to a 50k array increased prediction accuracy by 9 percentage points in sheep [31]. In other cases in 480 481 Nordic cattle, however, the addition of ~1500 variants preselected by GWAS to a 54k 482 panel produced negligible improvements in the prediction of traits with low 483 heritability [65].

484 Preselecting an entirely new set of predictor variants from WGS, as in 485 Top40k, proved more challenging than ChipPlusSign. In Top40k, we preselected the 486 variants with the lowest p-value in each of consecutive non-overlapping 55-kb 487 windows along the genome. This strategy did not perform much differently from just 488 taking random variants from these windows, as in Rand40k. One possible reason for 489 these results is that at this variant density, random variants effectively tag the same 490 associations as Top40k thanks to linkage disequilibrium. Denser sets of predictor 491 variants provided only small further improvements of prediction accuracy with 492 diminishing returns.

The modest performance of ChipPlusSign and Top40k could also be a consequence of the difficulty for fine-mapping causal variants through GWAS with WGS. Theoretically, the identification of all causal variants associated with a trait

496 should enable the improvement of prediction accuracy [12]. Even though WGS allows 497 the detection of a very large number of associations, problems such as false positives 498 or p-value inflation also become more severe in a way that added noise might offset 499 the detected signal. For instance, results in cattle showed that GWAS with WGS did 500 not detect clearer associated regions relative to marker arrays and failed to capture 501 QTL for genomic prediction [14], as the effect of potential QTL were spread across 502 multiple variants. Therefore, WGS performed better with simple genetic architectures 503 (i.e., traits with low number of QTN). This is consistent with expectations and 504 simulation results [8] that indicated that the benefit of using WGS for genomic 505 prediction would be limited by the number and size of OTN. When there are many 506 QTN with small effects it becomes much more difficult to properly estimate their 507 effects accurately. Therefore, for largely polygenic traits (as most traits of interest in 508 livestock production), training sets need to be very large before WGS can increase 509 prediction accuracy [8].

510 The advantage of using WGS might be limited by the current training set 511 sizes, especially in scenarios where marker arrays are already yielding high prediction 512 accuracy [14,20]. Multi-line training sets could be particularly beneficial with the use 513 of WGS because they allow a larger training set with low pairwise relationship degree 514 among individuals. Previous simulations suggested that WGS might be the most 515 beneficial with multi-breed reference panels [66], especially for numerically small 516 populations. Our results with a multi-line training set indicated that WGS can improve 517 prediction accuracy in scenarios that are less optimised than within-line genomic 518 prediction. The average improvements of prediction accuracy of 4.2 percentage points 519 were observed for the populations that contributed around 80k individuals to the 520 training set. However, in general those predictions were still less accurate than using 521 variants preselected under within-line training sets. In our multi-line scenarios we 522 only used variation that segregated across all seven lines. We observed that 523 population-specific variation accounted only for small fractions of genetic variance 524 [53] and it seems unlikely that they would contribute much to prediction accuracy 525 across breeds. Another possible obstacle is the differences in the allele substitution 526 effects of the causal mutations across breeds. This can be caused by differences in 527 allele frequency, contributions of non-additive effects and different genetic 528 backgrounds, or even gene-by-environment interactions among others [24,67].

529 We observed low robustness of genomic prediction with WGS across traits 530 and lines, and drops in prediction accuracy in those scenarios where genomic 531 prediction with WGS failed. Regarding bias, we did not observe a systematic increase 532 for ChipPlusSign despite using the same individuals for variant preselection and for 533 training the predictors [16,63]. When we split the training set into two subsets, one for GWAS-based variant preselection and the other for training of the predictive 534 535 equations, we did not observe any improvement in accuracy or bias. One hypothesis is 536 that both subsets belong to the same population and therefore retained similar interrelationship degrees (i.e., they are not strictly independent sets of individuals). 537 538 Moreover, the reduction in individuals available for training the predictors negatively 539 affected prediction accuracy.

We did not directly test persistence of prediction accuracy, but previous studies with real data found no higher persistence of prediction accuracy for WGS, not even with low degree of relationship between training and testing sets [14]. We would expect such obstacles to persistence of accuracy until causal variants can be successfully identified.

545

Suboptimal strategy and pitfalls

546 The use of WGS for genomic prediction can only be reached after many other 547 steps are completed to produce the genotypes at whole-genome level. Each of these steps has its pitfalls. It is unavoidable that the success of using of WGS is sensitive 548 549 not only to the prediction methodology itself but also to the strategy followed until 550 genotyping. This strategy includes the choice of which individuals to sequence, the 551 bioinformatics pipeline to call variants, the imputation of the WGS and choice of variant filters. When combined with the multiplicity of prediction methods and the 552 553 preselection of predictor variants (which is unavoidable with current datasets, 554 predictive methodologies and computational capacities), there are many options and 555 variables in the whole process that can affect the final result and that are not yet well 556 understood. Therefore, a much greater effort for optimising such strategies is required. Here we tested relatively simple approaches to see how they performed with large 557 558 WGS datasets. We have discussed what in our opinion are the main pitfalls of our 559 approach for selection of the individuals to sequence [55] and the biases that may 560 appear during processing of sequencing reads [50] elsewhere, and therefore here we will focus on imputation of WGS and its use for genomic prediction. 561

562

563 *Imputation accuracy*

It is widely recognized that imputation from marker arrays to WGS from very few sequenced individuals can introduce genotyping errors and that genotype uncertainty can be high [17,21,68,69]. The accuracy of the imputed WGS is one of the main factors that may limit its performance for genomic prediction. In a simulation study, van den Berg et al. [17] quantified the impact of imputation errors on

569 prediction accuracy and showed that prediction accuracy decreases as errors 570 accumulate, especially in the testing set.

571 Imputation of WGS is particularly challenging because typically we have to 572 impute a very large number of variants for a very large number of individuals from 573 very few sequenced individuals. We assessed the imputation accuracy of our approach 574 elsewhere [41,55] and recommended that $\sim 2\%$ of the population should be sequenced. 575 In our study, line D was the line where prediction accuracy with Top40k performed 576 the worst, mostly performing below Chip predictors. In this line, only 0.9% of the 577 individuals in the population had been sequenced and therefore lower imputation 578 accuracy could be expected. Although there was not enough evidence for establishing 579 a link between these two features (sequencing effort and prediction accuracy), we 580 recommend cautious design of a sequencing strategy that is suited to the intended 581 imputation method [55].

582 Prediction accuracy could be improved by accounting for genotype uncertainty 583 of the imputed WGS. For that, it could be advantageous to use allele dosages rather 584 than best-guess genotypes [69], although most current implementations cannot handle 585 such information.

586

587 *Preselection of predictor variants*

Simply using WGS to increase the number of markers does not improve prediction accuracy [18,21,24]. Due to the large dimensionality of WGS, there is a need to remove uninformative variants [24,31,66,68,70]. Predictor variants must be causal or at least informative of the causal variants, which depends on the distance between the markers and the causal variants [13]. For this reason, variants that are in weak linkage disequilibrium with causal mutations have a 'dilution' effect, i.e., they 594 add noise and limit prediction accuracy [24,31,70]. However, if too stringent filters 595 are applied during preselection of predictor variants, there is a risk of removing true 596 causal variants, and that would debilitate persistence of accuracy across generations 597 and across populations [66,71]. For instance, the impact of removing predictor variants with low minor allele frequency can vary depending on the minor allele 598 599 frequency of the causal variants as well as the distance between predictor and causal 600 variants [13]. Losing causal or informative variants would negatively affect multi-line 601 or multi-breed prediction.

602 A popular strategy to preselect the predictor variants is based on association 603 tests. Genome-wide association studies on WGS are expected to confirm associations 604 that were already detected with marker arrays and identify novel associations (e.g., 605 [36,72]). However, preliminary inspection of our GWAS results for the real traits 606 showed that the added noise could easily offset the added information and fine-607 mapping remains challenging. Multi-breed GWAS [4] and meta-analyses [73] are 608 suitable alternatives for GWAS to accommodate much larger population sizes and for 609 combining results of populations with diverse genetic backgrounds. Multi-breed 610 GWAS can be more efficient to identify informative variants than single-breed 611 GWAS, which may benefit even prediction within lines [74]. Because the signal of 612 some variants may go undetected for some traits but not for other correlated traits, 613 combining GWAS information of several traits can also help identifying weak or 614 moderate associations [25]. We did not test whether combining the significant 615 markers from the different single-trait GWAS yielded greater improvements in 616 prediction accuracy [29,32]. Multi-trait GWAS models could be more suited for that 617 purpose [72,75]. To improve fine-mapping, other GWAS models that incorporate

618 biological information have been proposed (e.g., functional annotation [76] or 619 metabolomics [77]).

620 There have been other suggested methods that may improve variant 621 preselection for genomic prediction. VanRaden et al. [30] suggested that preselecting 622 variants based on the genetic variance that they contribute rather than the significance 623 of the association could be more advantageous, as the former would indirectly 624 preselect variance with higher minor allele frequency. Other authors proposed 625 preselection of variants using statistics that do not depend on GWAS, such as the 626 fixation index (F_{ST}) score between groups of individuals with high and low phenotype 627 values [70], as an alternative to avoid the negative impact of spurious associations.

628 Preselecting predictor variants based on functional annotation was not useful, 629 as it reduced prediction accuracy in several traits and lines. Previous studies showed 630 that subsets of variants based on functionality either did not improve or reduced 631 prediction accuracy [20] and that adding preselected variants from coding regions to 632 marker arrays produced lower prediction accuracy than just adding the same number 633 of variants without considering functional classification [32]. A plausible explanation 634 is that functional variants are enriched for lower minor allele frequency, which can be 635 less informative for prediction [13]. Furthermore, functional annotation does not 636 necessarily capture true effects, and the method we used is biased towards protein-637 coding variants, which may lead to an underrepresentation of functional non-coding 638 variants that may explain a large fraction of quantitative trait variance. Xiang et al. 639 [78] found that expression QTL and non-coding variants explained more variation in 640 quantitative traits in cattle than protein-coding functional variants. When functional 641 annotation is not considered, intergenic variants are more likely to be preselected by 642 chance. Such variants tend to be more common and widespread across populations,

and therefore can act as tag variants and capture much larger fractions of trait variance[53].

Another popular strategy to reduce the number of variants is to prune variants based on linkage disequilibrium (LDTags). This strategy performed very poorly in our populations. Other studies reported different outcomes, where pruning for $r^2>0.9$ provided positive results [18,21]. It is possible that this was in part due to the stringent threshold ($r^2>0.1$) that we used in order to retain only a small number of variants.

650

651 New models and methods

652 It is also likely that models, methods, and their implementations need to be 653 improved to handle the complexity of WGS and to efficiently estimate marker effects 654 of so many variants with high accuracy, among other features. This is a very active 655 area of research and multiple novel methodologies have been proposed over the last 656 vears. Some examples are a combination of subsampling and Gibbs sampling [79]. and a model that simultaneously fits a GBLUP term for a polygenic effect and a 657 658 BayesC term for variants with large effects selected by the model (BayesGC) [26]. 659 Testing alternative models and methods for genomic prediction was out of the scope 660 of this report. However, together with refinements in the preselection of predictor 661 variants, it remains an interesting avenue for further optimisation of the analysis 662 pipeline.

Some of the most promising methods are designed to incorporate prior biological information into the models. One of such methods is BayesRC [23], which extends BayesR by assigning flatter prior distributions to classes of variants that are more likely to be causal [19,22]. Similarly, GFBLUP [80] could be used to incorporate prior biological information from either QTL databases or GWAS as 668 genomic features [21,35,68]. The model MBMG [27], which fits two genomic 669 relationship matrices according to prior biological information, has also been 670 proposed for multi-breed scenarios to improve genomic prediction in small 671 populations. Haplotype-based prediction methods could provide greater prediction 672 accuracy with WGS than SNP-based methods in pigs [81] and cattle [82]. These 673 methods reduce the number of model dimensions. However, the uptake of such 674 methods has been limited so far due to their greater complexity, for example, to define 675 haplotype blocks.

676

Conclusion

677 Our results evidenced the potential for WGS to improve genomic prediction 678 accuracy in intensely selected pig lines. The performance of each set of predictor 679 variants was not robust across traits and lines and the improvements that we achieved 680 so far were modest at best. The most robust results were obtained when variants that 681 showed statistically significant associations to the trait were preselected and added to 682 the marker array. With this method, average improvements of prediction accuracy of 683 2.5 and 4.2 percentage points were observed in within-line and multi-line scenarios, 684 respectively, with training sets of around 80k individuals. We would expect that a 685 combination of larger training sets and improved pipelines could help achieve greater 686 improvements of prediction accuracy. The robustness of the whole strategy for 687 generating WGS at the population level must be carefully stress-tested and further 688 optimised.

689

Ethics approval and consent to participate

- 690 The samples used in this study were derived from the routine breeding activities of
- 691 PIC.

Consent for publication

692 Not applicable.

Availability of data and material

- 693 The software packages AlphaSeqOpt, AlphaPhase, AlphaImpute and AlphaPeel are
- 694 available from the AlphaGenes website (http://www.alphagenes.roslin.ed.ac.uk). The
- 695 datasets generated and analysed in this study are derived from the PIC breeding
- 696 programme and not publicly available.

Competing interests

- 697 The authors declare that they have no competing interests. BDV, CYC, and WOH are
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Authors' contributions

- RRF, GG and JMH designed the study; CYC assisted in preparing the datasets; RRF,
- AW and MJ performed the analyses; RRF wrote the first draft; AW, CYC, BDV,

- 706 WHO, GG and JMH assisted in the interpretation of the results and provided
- comments on the manuscript. All authors read and approved the final manuscript.

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- 710

References

- 1. Pasaniuc B, Rohland N, McLaren PJ, Garimella K, Zaitlen N, Li H, et al.
 Extremely low-coverage sequencing and imputation increases power for genomewide association studies. Nat Genet. 2012;44:631–5.
- 2. Daetwyler HD, Capitan A, Pausch H, Stothard P, van Binsbergen R, Brondum RF,
 et al. Whole-genome sequencing of 234 bulls facilitates mapping of monogenic and
 complex traits in cattle. Nat Genet. 2014;46:858–65.
- 3. Nicod J, Davies RW, Cai N, Hassett C, Goodstadt L, Cosgrove C, et al. Genomewide association of multiple complex traits in outbred mice by ultra-low-coverage
 sequencing. Nat Genet. 2016;48:912–8.
- 4. Sanchez M-P, Govignon-Gion A, Croiseau P, Fritz S, Hozé C, Miranda G, et al.
 Within-breed and multi-breed GWAS on imputed whole-genome sequence variants
 reveal candidate mutations affecting milk protein composition in dairy cattle. Genet
 Sel Evol. 2017;49:68.
- 5. Das A, Panitz F, Gregersen VR, Bendixen C, Holm L-E. Deep sequencing of
 Danish Holstein dairy cattle for variant detection and insight into potential loss-offunction variants in protein coding genes. BMC Genomics. 2015;16:1043.
- 6. Gudbjartsson DF, Helgason H, Gudjonsson SA, Zink F, Oddson A, Gylfason A, et
 al. Large-scale whole-genome sequencing of the Icelandic population. Nat Genet.
 2015;47:435–44.
- 730 7. VanRaden PM. Symposium review: How to implement genomic selection. J Dairy731 Sci. 2020;103:5291–301.
- 8. Clark SA, Hickey JM, van der Werf JH. Different models of genetic variation andtheir effect on genomic evaluation. Genet Sel Evol. 2011;43:18.
- 9. Druet T, Macleod IM, Hayes BJ. Toward genomic prediction from whole-genome
 sequence data: impact of sequencing design on genotype imputation and accuracy of
 predictions. Heredity. 2014;112:39–47.

- 10. MacLeod IM, Hayes BJ, Goddard ME. The Effects of Demography and LongTerm Selection on the Accuracy of Genomic Prediction with Sequence Data.
 Genetics. 2014;198:1671–84.
- 740 11. Meuwissen T, Goddard M. Accurate Prediction of Genetic Values for Complex
 741 Traits by Whole-Genome Resequencing. Genetics. 2010;185:623–31.
- 742 12. Pérez-Enciso M, Rincón JC, Legarra A. Sequence- vs. chip-assisted genomic
 743 selection: accurate biological information is advised. Genet Sel Evol. 2015;47:43.
- 13. van den Berg I, Boichard D, Guldbrandtsen B, Lund MS. Using Sequence
 Variants in Linkage Disequilibrium with Causative Mutations to Improve AcrossBreed Prediction in Dairy Cattle: A Simulation Study. G3 GenesGenomesGenetics.
 2016;6:2553–61.
- 14. van Binsbergen R, Calus MPL, Bink MCAM, van Eeuwijk FA, Schrooten C,
 Veerkamp RF. Genomic prediction using imputed whole-genome sequence data in
 Holstein Friesian cattle. Genet Sel Evol. 2015;47:71.
- 15. Calus MPL, Bouwman AC, Schrooten C, Veerkamp RF. Efficient genomic
 prediction based on whole-genome sequence data using split-and-merge Bayesian
 variable selection. Genet Sel Evol. 2016;48:49.
- 16. Veerkamp RF, Bouwman AC, Schrooten C, Calus MPL. Genomic prediction
 using preselected DNA variants from a GWAS with whole-genome sequence data in
 Holstein–Friesian cattle. Genet Sel Evol. 2016;48:95.
- 17. van den Berg I, Bowman PJ, MacLeod IM, Hayes BJ, Wang T, Bolormaa S, et al.
 Multi-breed genomic prediction using Bayes R with sequence data and dropping
 variants with a small effect. Genet Sel Evol. 2017;49:70.
- 18. Frischknecht M, Meuwissen THE, Bapst B, Seefried FR, Flury C, Garrick D, et al.
 Short communication: Genomic prediction using imputed whole-genome sequence
 variants in Brown Swiss Cattle. J Dairy Sci. 2018;101:1292–6.
- 19. Hayes BJ, MacLeod IM, Daetwyler HD, Bowman PJ, Chamberlain AJ, Vander
 Jagt CJ, et al. Genomic prediction from whole genome sequence in livestock: the
 1000 Bull Genomes Project. Proc 10th World Congr Genet Appl Livest Prod
 WCGALP. Vancouver, BC, Canada; 2014. p. 183.
- Periode 20. Heidaritabar M, Calus MPL, Megens H-J, Vereijken A, Groenen MAM,
 Bastiaansen JWM. Accuracy of genomic prediction using imputed whole-genome
 sequence data in white layers. J Anim Breed Genet. 2016;133:167–79.
- 21. Song H, Ye S, Jiang Y, Zhang Z, Zhang Q, Ding X. Using imputation-based
 whole-genome sequencing data to improve the accuracy of genomic prediction for
 combined populations in pigs. Genet Sel Evol. 2019;51:58.
- 22. Zhang C, Kemp RA, Stothard P, Wang Z, Boddicker N, Krivushin K, et al.
 Genomic evaluation of feed efficiency component traits in Duroc pigs using 80K,
 650K and whole-genome sequence variants. Genet Sel Evol. 2018;50:14.

MacLeod IM, Bowman PJ, Vander Jagt CJ, Haile-Mariam M, Kemper KE,
Chamberlain AJ, et al. Exploiting biological priors and sequence variants enhances
QTL discovery and genomic prediction of complex traits. BMC Genomics.
2016;17:144.

24. Raymond B, Bouwman AC, Schrooten C, Houwing-Duistermaat J, Veerkamp RF.
Utility of whole-genome sequence data for across-breed genomic prediction. Genet
Sel Evol. 2018;50:27.

783 25. Xiang R, MacLeod IM, Daetwyler HD, de Jong G, O'Connor E, Schrooten C, et

al. Genome-wide fine-mapping identifies pleiotropic and functional variants that
 predict many traits across global cattle populations. Nat Commun. 2021;12:860.

786 26. Meuwissen T, van den Berg I, Goddard M. On the use of whole-genome sequence
787 data for across-breed genomic prediction and fine-scale mapping of QTL. Genet Sel
788 Evol. 2021;53:19.

- 789 27. Raymond B, Bouwman AC, Wientjes YCJ, Schrooten C, Houwing-Duistermaat J,
 790 Veerkamp RF. Genomic prediction for numerically small breeds, using models with
- 791 pre-selected and differentially weighted markers. Genet Sel Evol. 2018;50:49.

792 28. Moghaddar N, Brown DJ, Swan AA, Gurman PM, Li L, Werf JH. Genomic
793 prediction in a numerically small breed population using prioritized genetic markers
794 from whole-genome sequence data. J Anim Breed Genet. 2021;

Provide Structure
29. Brøndum RF, Su G, Janss L, Sahana G, Guldbrandtsen B, Boichard D, et al.
Quantitative trait loci markers derived from whole genome sequence data increases
the reliability of genomic prediction. J Dairy Sci. 2015;98:4107–16.

30. VanRaden PM, Tooker ME, O'Connell JR, Cole JB, Bickhart DM. Selecting
sequence variants to improve genomic predictions for dairy cattle. Genet Sel Evol.
2017;49:32.

31. Al Kalaldeh M, Gibson J, Duijvesteijn N, Daetwyler HD, MacLeod I, Moghaddar
N, et al. Using imputed whole-genome sequence data to improve the accuracy of
genomic prediction for parasite resistance in Australian sheep. Genet Sel Evol.
2019;51:32.

32. Lopez BIM, An N, Srikanth K, Lee S, Oh J-D, Shin D-H, et al. Genomic
Prediction Based on SNP Functional Annotation Using Imputed Whole-Genome
Sequence Data in Korean Hanwoo Cattle. Front Genet. 2021;11:603822.

33. Hayes BJ, Daetwyler HD. 1000 Bull Genomes Project to Map Simple and
Complex Genetic Traits in Cattle: Applications and Outcomes. Annu Rev Anim
Biosci. 2019;7:89–102.

34. Sanchez M-P, Guatteo R, Davergne A, Saout J, Grohs C, Deloche M-C, et al.
Identification of the ABCC4, IER3, and CBFA2T2 candidate genes for resistance to
paratuberculosis from sequence-based GWAS in Holstein and Normande dairy cattle.
Genet Sel Evol. 2020;52:14.

- 815 35. Yang R, Xu Z, Wang Q, Zhu D, Bian C, Ren J, et al. Genome-wide association
- study and genomic prediction for growth traits in yellow-plumage chicken usinggenotyping-by-sequencing. Genet Sel Evol. 2021;53:82.

818 36. Yan G, Liu X, Xiao S, Xin W, Xu W, Li Y, et al. An imputed whole-genome
819 sequence-based GWAS approach pinpoints causal mutations for complex traits in a
820 specific swine population. Sci China Life Sci. 2021;

- 37. Yang R, Guo X, Zhu D, Tan C, Bian C, Ren J, et al. Accelerated deciphering of
 the genetic architecture of agricultural economic traits in pigs using a low-coverage
 whole-genome sequencing strategy. GigaScience. 2021;10:giab048.
- 38. Li Y, Sidore C, Kang HM, Boehnke M, Abecasis GR. Low-coverage sequencing:
 Implications for design of complex trait association studies. Genome Res.
 2011;21:940–51.
- 39. Hickey JM. Sequencing millions of animals for genomic selection 2.0. J Anim
 Breed Genet. 2013;130:331–2.
- 40. Hickey JM, Gorjanc G, Cleveland MA, Kranis A, Jenko J, Mésázros G, et al.
 Sequencing Millions of Animals for Genomic Selection 2.0. Proc 10th World Congr
 Genet Appl Livest Prod WCGALP. Vancouver, BC, Canada; 2014. p. 377.
- 41. Ros-Freixedes R, Whalen A, Chen C-Y, Gorjanc G, Herring WO, Mileham AJ, et
 al. Accuracy of whole-genome sequence imputation using hybrid peeling in large
 pedigreed livestock populations. Genet Sel Evol. 2020;52:17.
- 42. Gonen S, Ros-Freixedes R, Battagin M, Gorjanc G, Hickey JM. A method for the
 allocation of sequencing resources in genotyped livestock populations. Genet Sel
 Evol. 2017;49:47.
- 43. Ros-Freixedes R, Gonen S, Gorjanc G, Hickey JM. A method for allocating lowcoverage sequencing resources by targeting haplotypes rather than individuals. Genet
 Sel Evol. 2017;49:78.
- 44. Hickey JM, Kinghorn BP, Tier B, Wilson JF, Dunstan N, van der Werf JH. A
 combined long-range phasing and long haplotype imputation method to impute phase
 for SNP genotypes. Genet Sel Evol. 2011;43:12.
- 45. Hickey JM, Kinghorn BP, Tier B, van der Werf JH, Cleveland MA. A phasing
 and imputation method for pedigreed populations that results in a single-stage
 genomic evaluation. Genet Sel Evol. 2012;44:9.
- 847 46. Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for Illumina
 848 sequence data. Bioinformatics. 2014;30:2114–20.
- 47. Li H. Aligning sequence reads, clone sequences and assembly contigs with BWAMEM. arXiv. 2013;1303.3997v1 [q bio.GN].

48. DePristo MA, Banks E, Poplin R, Garimella KV, Maguire JR, Hartl C, et al. A
framework for variation discovery and genotyping using next-generation DNA
sequencing data. Nat Genet. 2011;43:491–8.

49. Poplin R, Ruano-Rubio V, DePristo MA, Fennell TJ, Carneiro MO, Van der Auwera GA, et al. Scaling accurate genetic variant discovery to tens of thousands of samples. bioRxiv. 2018;10.1101/201178.

- 857 50. Ros-Freixedes R, Battagin M, Johnsson M, Gorjanc G, Mileham AJ, Rounsley
 858 SD, et al. Impact of index hopping and bias towards the reference allele on accuracy
 859 of genotype calls from low-coverage sequencing. Genet Sel Evol. 2018;50:64.
- 51. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, et al. The Sequence
 Alignment/Map format and SAMtools. Bioinformatics. 2009;25:2078–9.
- 52. Danecek P, Auton A, Abecasis G, Albers CA, Banks E, DePristo MA, et al. The
 variant call format and VCFtools. Bioinformatics. 2011;27:2156–8.
- 864 53. Ros-Freixedes R, Valente B, Chen C-Y, Herring WO, Gorjanc G, Hickey JM, et
 865 al. Rare and population-specific functional variants across pig lines. bioRxiv
 866 [Internet]. 2022; Available from: https://doi.org/10.1101/2022.02.01.478603
- 54. Whalen A, Ros-Freixedes R, Wilson DL, Gorjanc G, Hickey JM. Hybrid peeling
 for fast and accurate calling, phasing, and imputation with sequence data of any
 coverage in pedigrees. Genet Sel Evol. 2018;50:67.
- 870 55. Ros-Freixedes R, Whalen A, Gorjanc G, Mileham AJ, Hickey JM. Evaluation of
 871 sequencing strategies for whole-genome imputation with hybrid peeling. Genet Sel
 872 Evol. 2020;52:18.
- 56. VanRaden PM, Wiggans GR. Derivation, Calculation, and Use of National
 Animal Model Information. J Dairy Sci. 1991;74:2737–46.
- 57. Lippert C, Listgarten J, Liu Y, Kadie CM, Davidson RI, Heckerman D. FaST
 linear mixed models for genome-wide association studies. Nat Methods. 2011;8:833–
 5.
- 58. Widmer C, Lippert C, Weissbrod O, Fusi N, Kadie C, Davidson R, et al. Further
 Improvements to Linear Mixed Models for Genome-Wide Association Studies. Sci
 Rep. 2015;4:6874.
- 59. Chang CC, Chow CC, Tellier LC, Vattikuti S, Purcell SM, Lee JJ. Secondgeneration PLINK: rising to the challenge of larger and richer datasets. GigaScience.
 2015;4.
- 60. McLaren W, Gil L, Hunt SE, Riat HS, Ritchie GRS, Thormann A, et al. The
 Ensembl Variant Effect Predictor. Genome Biol. 2016;17:122.
- 886 61. Erbe M, Hayes BJ, Matukumalli LK, Goswami S, Bowman PJ, Reich CM, et al.
 887 Improving accuracy of genomic predictions within and between dairy cattle breeds
 888 with imputed high-density single nucleotide polymorphism panels. J Dairy Sci.
 889 2012;95:4114–29.

- 890 62. Moser G, Lee SH, Hayes BJ, Goddard ME, Wray NR, Visscher PM. Simultaneous
- 891 Discovery, Estimation and Prediction Analysis of Complex Traits Using a Bayesian
- 892 Mixture Model. Haley C, editor. PLOS Genet. 2015;11:e1004969.

63. MacLeod IM, Bolormaa S, Schrooten C, Goddard ME, Daetwyler H. Pitfalls of
pre-selecting subsets of sequence variants for genomic prediction. Proc 22nd Conf
Assoc Adv Anim Breed Genet AAABG. Townsville, Queensland, Australia; 2017. p.
141–4.

- 64. Gozalo-Marcilla M, Buntjer J, Johnsson M, Batista L, Diez F, Werner CR, et al.
 Genetic architecture and major genes for backfat thickness in pig lines of diverse
 genetic backgrounds. Genet Sel Evol. 2021;53:76.
- 65. Gebreyesus G, Lund MS, Sahana G, Su G. Reliabilities of Genomic Prediction for
 Young Stock Survival Traits Using 54K SNP Chip Augmented With Additional
 Single-Nucleotide Polymorphisms Selected From Imputed Whole-Genome
 Sequencing Data. Front Genet. 2021;12:667300.
- 66. Iheshiulor OOM, Woolliams JA, Yu X, Wellmann R, Meuwissen THE. Withinand across-breed genomic prediction using whole-genome sequence and single
 nucleotide polymorphism panels. Genet Sel Evol. 2016;48:15.
- 67. Legarra A, Garcia-Baccino CA, Wientjes YCJ, Vitezica ZG. The correlation of
 substitution effects across populations and generations in the presence of non-additive
 functional gene action. PREPRINT. 2021;
- 68. Sarup P, Jensen J, Ostersen T, Henryon M, Sørensen P. Increased prediction
 accuracy using a genomic feature model including prior information on quantitative
 trait locus regions in purebred Danish Duroc pigs. BMC Genet. 2016;17:11.
- 69. Pausch H, MacLeod IM, Fries R, Emmerling R, Bowman PJ, Daetwyler HD, et al.
 Evaluation of the accuracy of imputed sequence variant genotypes and their utility for
 causal variant detection in cattle. Genet Sel Evol. 2017;49:24.
- 70. Ling AS, Hay EH, Aggrey SE, Rekaya R. Dissection of the impact of prioritized
 QTL-linked and -unlinked SNP markers on the accuracy of genomic selection. BMC
 Genomic Data. 2021;22:26.
- 71. Fragomeni BO, Lourenco DAL, Masuda Y, Legarra A, Misztal I. Incorporation of
 causative quantitative trait nucleotides in single-step GBLUP. Genet Sel Evol.
 2017;49:59.
- 72. Bolormaa S, Swan AA, Stothard P, Khansefid M, Moghaddar N, Duijvesteijn N,
 et al. A conditional multi-trait sequence GWAS discovers pleiotropic candidate genes
 and variants for sheep wool, skin wrinkle and breech cover traits. Genet Sel Evol.
 2021;53:58.
- 73. van den Berg I, Xiang R, Jenko J, Pausch H, Boussaha M, Schrooten C, et al.
 Meta-analysis for milk fat and protein percentage using imputed sequence variant
 genotypes in 94,321 cattle from eight cattle breeds. Genet Sel Evol. 2020;52:37.

- 929 74. van den Berg I, Boichard D, Lund MS. Sequence variants selected from a multi-
- breed GWAS can improve the reliability of genomic predictions in dairy cattle. Genet
 Sel Evol. 2016;48:83.

932 75. Yoshida GM, Yáñez JM. Multi-trait GWAS using imputed high-density
933 genotypes from whole-genome sequencing identifies genes associated with body traits
934 in Nile tilapia. BMC Genomics. 2021;22:57.

- 76. Yang J, Fritsche LG, Zhou X, Abecasis G. A Scalable Bayesian Method for
 Integrating Functional Information in Genome-wide Association Studies. Am J Hum
 Genet. 2017;101:404–16.
- 77. Li J, Mukiibi R, Wang Y, Plastow GS, Li C. Identification of candidate genes and
 enriched biological functions for feed efficiency traits by integrating plasma
 metabolites and imputed whole genome sequence variants in beef cattle. BMC
 Genomics. 2021;22:823.
- 78. Xiang R, Berg I van den, MacLeod IM, Hayes BJ, Prowse-Wilkins CP, Wang M,
 et al. Quantifying the contribution of sequence variants with regulatory and
 evolutionary significance to 34 bovine complex traits. Proc Natl Acad Sci.
 2019;116:19398–408.
- 946 79. Xavier A, Xu S, Muir W, Rainey KM. Genomic prediction using subsampling.
 947 BMC Bioinformatics. 2017;18:191.

80. Edwards SM, Sørensen IF, Sarup P, Mackay TFC, Sørensen P. Genomic
Prediction for Quantitative Traits Is Improved by Mapping Variants to Gene Ontology
Categories in *Drosophila melanogaster*. Genetics. 2016;203:1871–83.

81. Bian C, Prakapenka D, Tan C, Yang R, Zhu D, Guo X, et al. Haplotype genomic
prediction of phenotypic values based on chromosome distance and gene boundaries
using low-coverage sequencing in Duroc pigs. Genet Sel Evol. 2021;53:78.

- 82. Li H, Zhu B, Xu L, Wang Z, Xu L, Zhou P, et al. Genomic Prediction Using LDBased Haplotypes Inferred From High-Density Chip and Imputed Sequence Variants
 in Chinese Simmental Beef Cattle. Front Genet. 2021;12:665382.
- 957
- 958

Figures

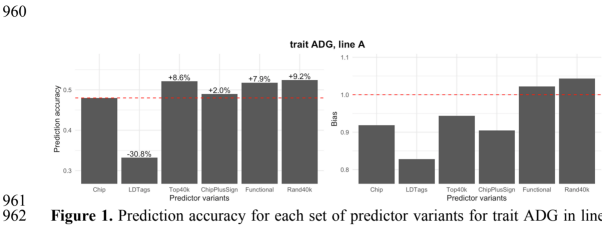
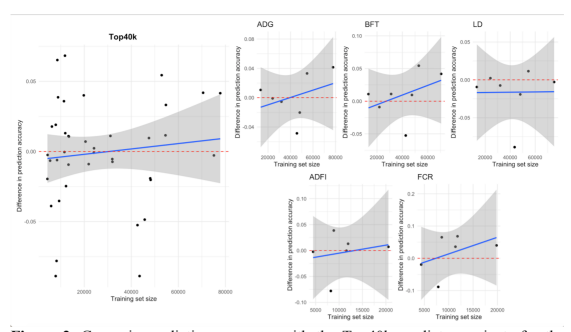


Figure 1. Prediction accuracy for each set of predictor variants for trait ADG in line

963 A. Left: Correlation (left). Dashed line at value of Chip as a reference. Values indicate

964 relative difference to Chip. Right: Bias. Dashed line at the ideal value.

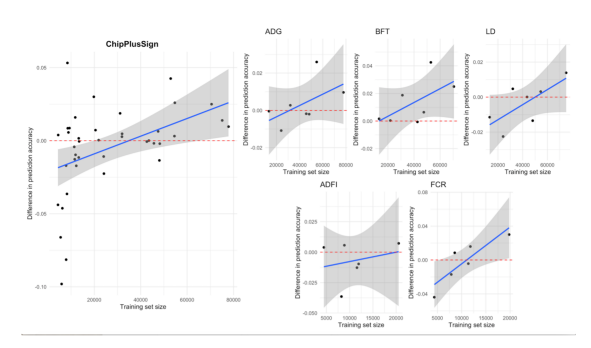


966 967

Figure 2. Genomic prediction accuracy with the Top40k predictor variants for the real traits. The difference of prediction accuracy between Top40k and Chip is shown,

969 for all traits and lines (left) or by trait (right). Red dashed line at 'no difference'.

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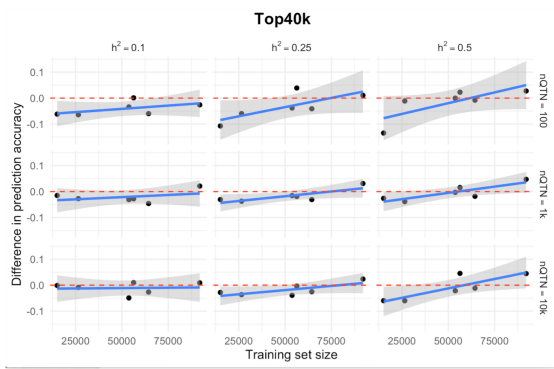


973 Figure 3. Genomic prediction accuracy with the ChipPlusSign predictor variants for

974 the real traits. The difference of prediction accuracy between ChipPlusSign and Chip
975 is shown, for all traits and lines (left) or by trait (right). Red dashed line at 'no
976 difference'.

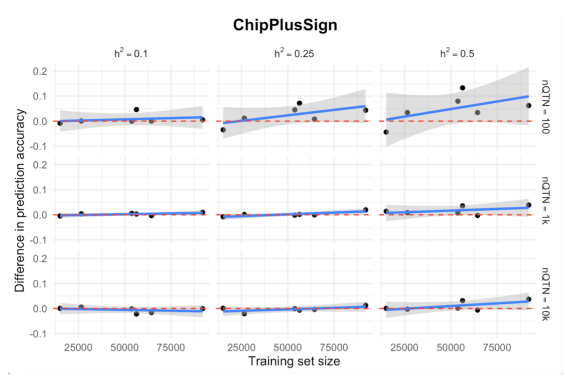
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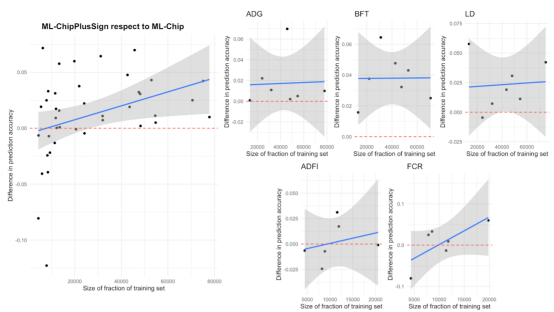
979
980 Figure 4. Genomic prediction accuracy with the Top40k predictor variants for the
981 simulated traits. The difference of prediction accuracy between Top40k and Chip is
982 shown by heritability (h²) and number of quantitative trait nucleotides (nQTN) of the
983 simulated traits. Red dashed line at 'no difference'.

984



986 987 Figure 5. Genomic prediction accuracy with the ChipPlusSign predictor variants for the simulated traits. The difference of prediction accuracy between ChipPlusSign and 988 Chip is shown by heritability (h^2) and number of quantitative trait nucleotides (nQTN) 989 990 of the simulated traits. Red dashed line at 'no difference'.

991



993
 994 Figure 6. Genomic prediction accuracy with the ML-ChipPlusSign predictor variants

for the real traits. The difference of prediction accuracy between ML-ChipPlusSign
and ML-Chip is shown, for all traits and lines (left) or by trait (right). Red dashed line
at 'no difference'.

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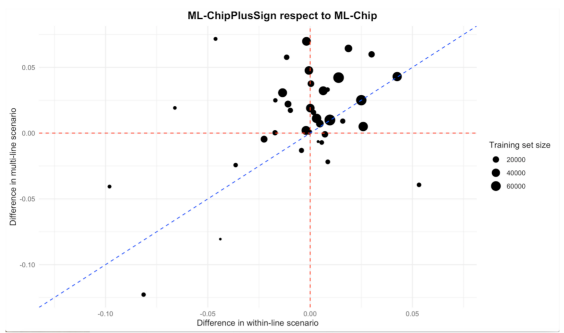
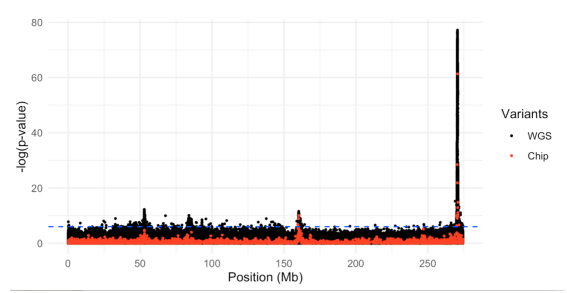


Figure 7. Comparison of the difference in genomic prediction accuracy in the multi-

line scenarios (between ML-ChipPlusSign and ML-Chip) and in the within-line
scenarios (between ChipPlusSign and Chip). Red dashed line at 'no difference'. Blue
dashed line is the bisector.



1007
1008Figure 8. Genome-wide association study results for trait BFT in line A. Only1009chromosome 1 is displayed as an example. In red, results for the variants in the1010marker array (Chip); in black, results for the whole-genome sequence data (WGS).1011The blue dashed line indicates significance threshold with Bonferroni's multiple test1012correction assuming that the markers from the marker arrays were independent (p-1013value $\leq 10^{-6}$).

Tables

Line	Individuals sequenced	Indiv	viduals cov	sequer erage	Individuals used in analyses		
		1x	2x	5x	15-30x	Pedigree	Imputed
А	1,856	1,044	649	73	90	122,753	104,661
В	1,366	685	545	44	92	88,964	76,230
С	1,491	628	728	54	81	84,420	66,608
D	731	362	311	16	42	79,981	60,474
Е	760	394	274	27	65	50,797	41,573
F	381	193	137	16	35	35,309	29,330
G	445	217	176	15	37	21,129	17,224

1015 **Table 1.** Number of sequenced pigs and pigs with imputed data.

	Trait	Α	В	С	D	E	F	G
	ADG	88,342	64,285	56,173	51,061	35,423	26,335	15,452
	BFT	80,146	62,027	55,233	47,509	34,527	23,872	15,268
	LD	85,233	64,141	56,026	48,509	35,495	26,453	15,274
	ADFI	21,960	9,525	9,062	12,256	12,444	4,105*	4,851
	FCR	21,200	9,217	8,654	12,044	12,316	4,016*	4,754
	TNB	13,581	10,721	9,626	7,729*	6,506*	-	3,230*
	LWW	-	9,112	7,251	-	-	-	2,813*
	RET	-	6,978	6,327	-	-	-	1,669*
_	Simulated	104,661	76,230	66,608	60,474	41,573	29,330	17,224

1017 **Table 2.** Number of phenotypic records per trait and line.

1018 ADG average daily gain, BFT backfat thickness, LD loin depth, ADFI average daily

1020 weight at weaning, *RET* return to oestrus 7 days after weaning.

1021 *Included in multi-line scenarios, but excluded in within-line scenarios because of the

1022 limited size of the testing set.

¹⁰¹⁹ feed intake, FCR feed conversion ratio, TNB total number of piglets born, LWW litter

1024 **Table 3.** Number of significant variants from the whole-genome sequence data that

Trait	A	В	С	D	Ε	F	G	Multi-line
ADG	646	581	424	498	279	219	143	4731
BFT	1083	758	664	518	1030	218	237	6149
LD	633	579	458	518	222	215	43	7247
ADFI	145	224	169	23	183	-	119	767
FCR	198	224	162	95	56	-	134	1369
TNB	71	117	161	-	-	-	-	248
LWW	-	32	73	-	-	-	-	480
RET	-	184	31	-	-	-	-	60

1025 were added to the marker array in ChipPlusSign.

1026 ADG average daily gain, BFT backfat thickness, LD loin depth, ADFI average daily

1027 feed intake, FCR feed conversion ratio, TNB total number of piglets born, LWW litter

1028 weight at weaning, *RET* return to oestrus 7 days after weaning.

1030 Table 4. Number of significantly associated genomic regions in the genome-wide

1031 association study for the simulated phenotypes that contained 0, 1 or 2 or more

h ²	nQTN	Line	Chip		Whole-genome sequence			
		size	0 QTN	1 QTN	0 QTN	1 QTN	≥2 QTN	
0.10	100	27k	4	1	8	6	0	
		56k	11	3	19	19	0	
		92k	10	7	44	19	0	
	1k	27k	1	0	4	0	1	
		56k	1	0	16	3	1	
		92k	1	0	283	9	0	
	10k	27k	1	0	1	0	0	
		56k	0	0	16	2	1	
		92k	2	0	186	17	12	
0.25	100	27k	11	6	26	15	1	
		56k	22	8	44	28	3	
		92k	20	7	90	34	1	
	1k	27k	0	0	8	1	3	
		56k	3	0	34	15	6	
		92k	6	0	692	49	16	
	10k	27k	0	0	2	0	0	
		56k	0	0	90	9	22	
		92k	4	0	564	56	164	
0.50	100	27k	18	9	24	24	1	
		56k	30	13	116	41	3	
		92k	17	9	425	44	1	
	1k	27k	6	0	22	9	6	
		56k	5	1	238	59	32	
		92k	11	1	903	169	120	
	10k	27k	0	0	4	0	0	
		56k	0	0	360	77	172	
		92k	10	0	379	116	508	

1032 quantitative trait nucleotides (QTN).