Genomic Prediction from Whole Genome Sequence in Livestock: the 1000 Bull Genomes Project

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ABSTRACT: Advantages of using whole genome sequence data to predict genomic estimated breeding values (GEBV) include better persistence of accuracy of GEBV across generations and more accurate GEBV across breeds. The 1000 Bull Genomes Project provides a database of whole genome sequenced key ancestor bulls, that can be used for imputing sequence variant genotypes into reference sets for genomic prediction. Run 3.0 included 429 sequences, with 31.8 million variants detected. BayesRC, a new method for genomic prediction, addresses some of the challenges associated with using the sequence data, and takes advantage of biological information. In a dairy data set, predictions using BayesRC and imputed sequence data from the 1000 Bull Genomes data were 2% more accurate than with 800k data, and we could demonstrate the method was able to identify causal mutations in some cases. Further improvements will come from more accurate imputation of sequence variant genotypes and improved biological information.

Keywords: genomic prediction, whole genome sequence, biological information

Introduction

Genomic prediction of breeding values is increasingly widely used in livestock breeding programs for dairy, beef, sheep, chickens and pigs (eg. Wiggans et al. 2011, Saatchi et al. 2012, Daetwyler et al. 2012, Hawken et al. 2014, Wellman et al. 2013). To date, most of these predictions have been based on arrays of approximately 50,000 SNP. The advantage of using whole genome sequence data for genomic prediction over such SNP arrays arises from the fact that with the sequence data, the actual casual mutations responsible for trait variation should now be in the data set. This could have at least three benefits:

Better persistence of accuracy of genomic 1) predictions over generations, and less erosion of accuracy in genomic predictions for individuals that are less related to the reference set. It has become clear that much of the accuracy of current genomic breeding values, based on 50,000 DNA markers, in fact derives from prediction of the effect of large chromosome segments that segregate within fairly closely related animals (eg. Habier et al. 2010). As a result, the accuracy of the prediction equation rapidly decays over generations as large chromosome segments break up due to recombination. If the causal mutations were identified using the sequence data, and the prediction equation was based on their effects, accuracy would persist over many more generations, and in more distantly related Macleod et al. (2014) and Meuwissen and animals. Goddard (2010) could clearly demonstrate this in simulated sequence data.

2) Higher accuracy of genomic predictions. One issue with the current SNP arrays (eg. 50k, 800k for cattle) is that the SNP have been selected to have a high minor allele frequency. This means that the SNP arrays are less likely to have SNP in linkage disequilibrium with casual mutations where one of the alleles is at low frequency in the population. If this variation from rare alleles could be captured with the whole genome sequence data, and exploited in genomic predictions, accuracy of genomic breeding value may be able to be improved in the order of 2-30%, depending on trait (Druet et al. 2014). It should be pointed out that in populations with small effective population size, such as Holsteins, the gain in accuracy from using sequence data is likely to be limited, as the current 50k SNP arrays capture a large proportion of the genetic variance. For example for milk production, Haile-Mariam et al. (2013), and Jensen et al. (2012), the additive genetic variance captured by the 50k SNP was 80-90%. However, for some traits like fertility and survival, the genetic variance captured by SNP arrays is only $\sim 60\%$, perhaps because there are more rare variants affecting these traits (Haile-Mariam et al. 2013). For these traits gains in accuracy from using sequence data might be higher.

3) More accurate genomic predictions across breeds. For a breed like Holstein dairy cattle, large reference populations have been assembled, leading to high accuracies of genomic predictions. For other breeds, assembling such large populations is challenging, so using genomic information across breeds would be appealing. However, the accuracy of genomic predictions across breeds with the Bovine 50k array is close to zero, and this improves slightly when the 800k array is used (Erbe et al. 2012). With the whole genome sequence data, at least the causative mutations which do segregate across breeds could be captured and this information used in multi-breed genomic predictions. A multi-breed reference population would be required to achieve this. A multi-breed reference would also benefit from the fact that linkage disequilibrium across breeds is lower than that within breeds, so that causative mutations could be mapped more precisely (eg Raven et al. 2014). So multi-breed reference combined with sequence data should be the best approach to achieve the potential benefits from sequence data stated above.

Even for large breeds such as Holstein, across breed information can be very valuable, for instance for traits difficult to record such as feed efficiency, or for causal mutations at low frequency and therefore difficult to characterize in the major breeds.

In fact it becomes clear from the above that making best use of the sequence information in genomic prediction is actually a QTL mapping problem – the aim of both becomes to identify causative mutations.

In this paper, we review the whole genome sequence data that is available, in cattle at least, from the 1000 Bull Genomes Project, discuss challenges with using such data in genomic predictions and how they can be overcome. We then present some preliminary results for accuracy of genomic predictions using sequence data.

The 1000 bull genomes data set

A large reference set of animals with phenotypes and genotypes is required to estimate the genomic prediction equation. These reference sets must be large (10,000s) to achieve accurate genomic predictions, given the typical architecture of complex traits (large numbers of mutations of small effects). Such large numbers of animals are unlikely to be sequenced in the near future. An alternative strategy is to sequence key ancestors of the population, then impute the genotypes for the sequence variants into much larger reference sets with phenotypes and SNP array genotypes. The 1000 Bull Genomes Project is building this database of sequenced key ancestor bulls for the bovine research community (Daetwyler et al. 2014). Run3.0 of the project included 429 full genome sequences of key ancestors from 15 breeds, sequenced at an average of 10.5 fold coverage, Table 1. There were 31.8 million filtered sequence variants detected in the sequences, including 29.1 million SNP and 1.7 million insertion-deletions (note that filtering does mean that some causative mutations, particularly those at very low frequency, may be filtered from the data set, and that we have only called SNP and Indel, see for example Dolezal et al. these proceedings and Guldbrandtsen et al. these proceedings for calling of other types of variant). Various quality control steps were applied to determine both the accuracy of calling variants in the sequence data, and the accuracy of calling genotypes at the variants. This included assessing the rate of opposing homozygote genotypes for sire-son pairs, and assessing the agreement between sequence genotypes and genotypes from an 800k SNP array in the sequenced bulls. This was high, at 98.8%. The variants were annotated into different functional classes, including intergenic and intragenic, synonymous and non-synonymous, and other classes, Table 2A.B. There was a considerable difference in allele frequency spectrum between the sequence variants, and the variants on the 800k Bovine HD Array, with many more sequence variants at low frequency. Missense (nonsynonymous) variants were at more extreme allele frequencies than synonymous variants.

 Table 1.
 Breeds and number per breed sequenced in

 Run 3.0 of 1000 Bull Genomes Project.

Breed	Number
	sequenced
Holstein	122
Jersey	26
Simmental	87
Angus	54
Swedish Reds	16
Piedmontese	2
Limousin	25
Hereford	1
Guelph Composite	9
Finnish Aryshire	17
Charolais	8
Brown Swiss	43
Belgian Blue	10
Beef Booster	8
All	429

Table 2A.	Annotation	of SNP	from	Run	3.0	of the	1000
Bull Geno	mes Project.						

renomes Project.	
Annotation	Number
intergenic_variant	19277503
intron_variant	7587343
upstream_gene_variant	1007825
downstream_gene_variant	879304
missense_variant	119236
synonymous_variant	109805
3_prime_UTR_variant	68408
splice_region_variant	22713
5_prime_UTR_variant	12800
stop_gained	3083
splice_donor_variant	2391
non_coding_exon_variant	9894
splice_acceptor_variant	1629
initiator_codon_variant	235
stop_lost	105
coding_sequence_variant	184
stop_retained_variant	74
mature_miRNA_variant	273
nc_transcript_variant	70
Total	29102875

An interesting alternative approach to the strategy adopted by the 1000 Bull Genomes Project (sequencing key ancestors at moderate fold coverage) would be to use the same sequencing effort to sequence a very large number of individuals at very low fold coverage. This strategy has been very successful in inbred plants like rice (Huang et al. 2010). However simulation suggests in outbred species (with heterozygote genotypes) a lower limit of 6 fold coverage is necessary to achieve accurate imputation of sequence variant genotypes into animals genotyped with SNP arrays (Druet et al. 2014).

Challenges with using sequencing data in genomic prediction

There are at least three challenges for dealing with whole genome sequence data for genomic predictions; the very large number of variants, achieving accurate imputation of sequence variant genotypes into animals genotyped with SNP arrays, and choosing a method to estimate the genomic prediction equation that makes best use of the sequence data.

From Run 3.0 of the 1000 Bull Genomes Project, 31.8 million variants were detected. These variants were either SNP or short insertion deletions - other types of genetic variants which could potentially be causative mutations are more difficult to detect, for example novel mobile genetic element insertions (eg. Guldbrandtsen et al., this proceedings). While 31.8 million variants could readily be used in genome wide association studies (GWAS), because the analysis is highly parallelizable, for genomic prediction methods this presents a significant challenge. One option is to use biological information to prioritise or filter variants. This biological information comes in two forms, sites in the genome where variants are more likely to have an effect on any trait, for example coding regions or regulatory regions, and gene sets in which mutations are more likely to affect specific traits, for example genes expressed in mammary gland for milk production in dairy cattle. For the first type of information, analysis of genome annotations for enrichment of GWAS hits (eg. significant SNP) has identified coding regions, particularly missense mutations and regions upstream and downstream of genes as enriched for trait associated variants in both humans and cattle (Kindt et al. 2013, Koufariotis et al. 2014). More recently, in human studies, regulatory regions (which are often in these upstream and downstream regions), have been shown to be enriched for GWAS hits for disease traits (Maurano et al. 2012). For the second type of information, "Atlases" of bovine gene expression provide a means of identifying potentially important gene sets. The atlases identify genes which are differential expressed across tissues, using either digital gene expression tags or RNA Sequence data (Harhay et al. 2010, Chamberlain these proceedings respectively). The utility of both types of information for genomic prediction is demonstrated later in this paper.

Once a subset of sequence variants has been identified for further analysis, the next step towards genomic prediction is imputation of these genotypes into reference populations, typically already genotyped with SNP arrays. A major challenge here is accurate imputation of rare variants. To demonstrate this, we used cross validation within the 1000 bull genomes data set to assess the accuracy of imputation. Randomly chosen subsets of sequenced Holstein animals had their sequence variant genotypes reduced to the 777K genotypes on the Bovine High density SNP array (there were 625,000 of these that were polymorphic and passed quality control). Beagle4.0 (Browning and Browning 2014) was used to impute full sequence variant genotypes in these animals (using all other sequenced animals as a reference). While the accuracy was reasonable for variants with minor allele frequency greater than 5%, below this, accuracy of imputation rapidly declined, Figure 1. This is in agreement with other studies on the same data (eg. Van Binsbergen et al. 2014). Use of pedigree information only slightly improved accuracy of imputing rare variants (Figure 1), but other imputation programs may perform better with rare variants. Alternative methods are being developed to better utilise multiple breed information in imputation (Bouwman and Veerkamp these proceedings, Daetwyler et al. 2014). Improving the imputation accuracy of sequence variant genotypes, particularly rare variants, is necessary before the full advantage of sequence data for genomic predictions can be realized. Imputation of rare variants is likely to be important, because for example non-synonymous variants tend to have low minor allele frequencies (Figure 1).

The final challenge to be addressed is choice of statistical method to derive the genomic prediction equation that will make best use of the sequence data. Best linear unbiased prediction methods (BLUP as described by Meuwissen et al. 2001, or it's mathematical equivalent, GBLUP, Habier et al. 2007), are unlikely to make the best use of sequence data for two reasons. The first is that the prior used in these methods assumes all variants have an effect. One of the main attractions of using sequence data is to identify the causal variants, and remove all other variants from the prediction equation. The BLUP methods cannot arrive at this solution, as every variant will have a predicted non-zero effect derived from a single normal distribution. Another problem with BLUP methodologies is that the severe shrinkage imposed means that the effect of a causative mutation is rarely captured by a single variant, rather the effect is split across several or many SNP (eg. Verbyla et al. 2009). So we require methods that allow a proportion of the variants to have zero effect, and preferably a variable degree of shrinkage, such that moderate to large effects of causal mutations are not smeared across multiple variants that are in moderate LD with the QTL. Methods which meet both these criteria include BayesB (Meuwissen et al. 2001), and BayesR (Erbe et al. 2012). Both BayesB and BayesR have been tested on simulated sequence data (Clark et al. 2011 and Macleod et al. 2014), and lead to both higher genomic prediction accuracy and greater persistence of accuracy over generations than BLUP methods in those simulations. A drawback of the Bayesian methods however is that they are typically implemented using Gibbs sampling, or Metropolis Hastings algorithms in the case of BayesB, and such sampling algorithms are costly in computing time with large number of variants, (eg. Van Binsbergen, this proceedings). Alternatives to sampling such as expectationmaximisation algorithms have been proposed for both BayesB (Shepherd et al. 2010), and BayesR (Wang et al. these proceedings). These revised Bayesian methods can decrease computer processing time by up to 10 fold.

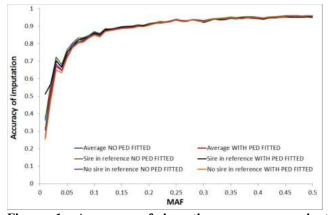


Figure 1. Accuracy of imputing sequence variant genotypes by minor allele frequency (MAF), for Chromosome 26. The accuracy of imputing genotypes was assessed by cross validation, where sets of 25 Holstein animals were removed from the sequence dataset, and the genotypes for these animals were reduced to the genotypes that were on the Bovine HD array, then imputed all the sequence variants using Beagle4. The correlation between the real sequence variants and imputed variants were then calculated. Pedigree information was either included in the imputation (with ped fitted) or not included (no ped fitted).

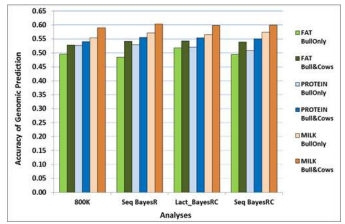


Figure 2. Accuracy of genomic prediction in the validation group of Red Holstein (Dutch origin) using either Australian Bulls only (Holstein and Jersey) or Australian Bulls and Cows (Holstein and Jersey) reference sets. The proxy for accuracy was the correlation between de-regressed MACE proofs and genomic predictions.

Results from genomic prediction with sequence data, with and without biological knowledge

The accuracy of genomic prediction using imputed sequence data was assessed using a reference population of 16,214 Holstein and Jersey cows and bulls (3049, 770 Holstein and Jersey bulls, and 8478, 3917 Holstein and Jersey cows, respectively). The phenotypes were daughter trait deviations for bulls, and trait deviations for cows, for milk volume, fat kg and protein kg. Phenotypes were weighted according to Garrick et al. (2009).

The validation population was 873 Dutch Red Holsteins, with phenotypes represented by de-regressed Interbull proofs on the Australian scale.

Genotypes were either imputed 632,003 SNP genotypes from the Illumina Bovine HD array, or imputed 1,674,245 sequence variant genotypes, where the variants were within genes or +/- 2kb from gene start and stop positions (a crude filtering of variants on biological information). Beagle 4.0 was used for imputation and one of all pairs of variants in complete linkage disequilibrium (r^2 >0.99) were pruned out.

Prediction equations were derived using either BayesR (Erbe et al. 2012), modified to use weights (Kemper et al. these proceedings), or a new method (BayesRC) that incorporates biological information, (MacLeod et al. these proceedings). BayesR assumes variant effects come from one of four normal distributions, one with zero variance (zero effect), and the other three with increasing variance. BayesRC is a modified version of BayesR (Erbe et al. 2012, Kemper et al. these proceedings). The key modification in BayesRC requires that all variants used in the analysis are first allocated to one of several "classes" based on prior knowledge (for example nonsynonymous mutations versus synonymous mutations), and the frequency with which SNP effects come from one of the four normal distributions is estimated separately for each class of SNP. This allows, for instance, the possibility that non-synonymous SNPs have a non-zero effect more often than synonymous SNPs (details in MacLeod et al. these Biological information, used to define proceedings). classes in BayesRC, was from two sources

1) Genome annotations. The three classes in BayesRC were then non-synonymous variants, (Table 2A,B), variants in upstream and downstream regions and annotated microRNAs (Table 2A,B), or 800k (variants on the BovineHD array and not in the two classes above). This analysis was called **Seq_BayesRC**,

2) Micro-array experiments to identify a set of genes in mammary gland that were found to play a role in milk synthesis, (Vander Jagt 2012). The classes here were 1) non synonymous coding variants in differentially expressed genes in the micro-array experiments 2) all other variants in genes differentially expressed in lactation experiments, and 3) all other sequence variants. This method was called **Lact_BayesRC**.

We assessed the accuracy of prediction of BayesR using either the 800k genotypes ("**800k**"), using the imputed subset of sequence data ("**Seq BayesR**") and **Seq_BayesRC** and **Lact_ BayesRC**. The proxy for accuracy was the correlation of genomic predicted EBV and de-regressed proof in the 876 Red Holstein bulls.

The largest increase in accuracy for all traits was as a result of including cows in the reference population (\sim 5%), Figure 2. Using sequence variants gave a 2% increase in accuracy over the 800k genotypes, averaged over traits. The BayesRC method did not give noticeably higher accuracy of genomic prediction in this data set, but identification of a relevant gene set could be further

improved by including information from RNAseq atlases (Chamberlain et al. these proceedings).

in 3.0 of the 1000 Bull Genor	nes Project.
Annotation	Number
Intergenic	1135727
intron_variant	460164
upstream_gene_variant	64332
downstream_gene_variant	57775
3_prime_UTR_variant	4823
frameshift_variant	1585
splice_region_variant	1207
inframe_deletion	998
5_prime_UTR_variant	834
inframe_insertion	349
splice_acceptor_variant	160
splice_donor_variant	120
missense_variant	88
non_coding_exon_variant	406
coding_sequence_variant	59
nc_transcript_variant	19
stop_gained	8
mature_miRNA_variant	45
stop_lost	1
Total	1728700

Table 2B.Annotation of Insertion deletions (Indel)from Run 3.0 of the 1000 Bull Genomes Project.

The BayesRC method did lead to more precise mapping of QTL effects (MacLeod et al. these proceedings), which is critical for persistence of accuracy of genomic predictions over generations. To assess the potential of BayesRC to identify causative mutations, we first investigated whether or not two previously identified mutations with effects on milk production traits were identified by the method. These were a mutation in the DGAT1 gene with an effect on fat%, fat kg, and milk volume (Grisart et al. 2002), and a mutation in the promoter of the PAEP gene (previously called Beta Lactoglobulin) with effects on protein percentage (Braunschweig and Leeb 2006). Both mutations were more clearly identified in Lact BayesRC, much more clearly than when BayesR was used, Figures 2 and 3 in MacLeod et al. these proceedings. This is encouraging, especially since there are four 800k SNP in high linkage disequilibrium $(r^2 \sim 0.75)$ with the DGAT1 causative mutation, but these SNP received a lower posterior probability in the BayesRC analysis, indicating that causative mutations can be identified even when there are other variants in relatively high linkage disequilibrium.

Conclusion

Genomic prediction with whole genome sequence data is now possible for cattle. The 1000 Bull Genomes Project provides a database of 31 million variant genotypes in key ancestor bulls that can be imputed into reference populations genotyped with SNP arrays, and genomic prediction methods that can deal with such large data sets are under development. One such method, BayesRC, takes advantage of biological information in genomic prediction from sequence data. In a dairy data set, predictions using BayesR or BayesRC and imputed sequence data from the 1000 bull genomes were 2% more accurate than from an 800k data set, and we could demonstrate BayesRC was able to identify causal mutations in some cases. Further improvements in accuracy of genomic prediction from sequence data, and particularly persistence of accuracy of these predictions across generations, will come from more accurate imputation of sequence variant genotypes, larger data sets, and improved biological information on sites and gene sets that are more likely to harbor mutations affecting quantitative traits.

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