



Phenotypic and genetic effects of recessive haplotypes on yield, longevity, and fertility

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ABSTRACT

Phenotypes from the August 2015 US national genetic evaluation were used to compute phenotypic effects of 18 recessive haplotypes in Ayrshire ($n = 1$), Brown Swiss ($n = 5$), Holstein ($n = 10$), and Jersey ($n = 2$) cattle on milk, fat, and protein yields, somatic cell score (SCS), single-trait productive life (PL), daughter pregnancy rate (DPR), heifer conception rate (HCR), and cow conception rate (CCR). The haplotypes evaluated were Ayrshire haplotype 1, Brown Swiss haplotypes 1 and 2, spinal dysmyelination, spinal muscular atrophy, Weaver Syndrome, brachyspina, Holstein cholesterol deficiency, Holstein haplotypes 1 to 5, bovine leukocyte adhesion deficiency, complex vertebral malformation, mulefoot (syndactyly), and Jersey haplotypes 1 and 2. When causal variants are unknown and tests are based only on single nucleotide polymorphism haplotypes, it can sometimes be difficult to accurately determine carrier status. For example, 2 Holstein haplotypes for cholesterol deficiency have the same single nucleotide polymorphism genotype, but only one of them carries the causative mutation. Genotyped daughters of carrier bulls included in the analysis ranged from 8 for Weaver Syndrome to 17,869 for Holstein haplotype 3. Lactation records preadjusted for nongenetic factors and direct genomic values (DGV) were used to estimate phenotypic and genetic effects of recessive haplotypes, respectively. We found no phenotypic or genetic differences between carriers and noncarriers of Ayrshire or Brown Swiss defects. Several associations were noted for Holstein haplotypes, including fat and HCR for Holstein haplotype 0 carriers; milk, protein, SCS, PL, and fertility for Holstein haplotype 1; protein, PL, CCR, and HCR for Holstein haplotype 2; milk, protein, and fertility for Holstein haplotype 4; and protein yield and DPR for Holstein haplotype 5. There were no differences among bovine leukocyte adhesion deficiency carriers, but complex vertebral malformation affected fat

yield and mulefoot carriers had higher SCS and lower PL DGV. Jersey haplotype 1 was not associated with any phenotypic effects, but we noted significant differences among DGV for fat, protein, PL, DPR, CCR, and HCR. Jersey haplotype 2 was associated only with lower phenotypic CCR. Effects of the recessive haplotypes on other traits studied generally were small even when significant. Almost \$11 million of economic losses per year due to reduced fertility and perinatal calf death in the US population can be avoided by selecting mate pairs that will not produce affected embryos. Carrier animals may continue to be selected if the merit of their favorable alleles exceeds the loss from their recessive alleles, but carrier bulls can be generally avoided without reducing the average genetic merit of the sires available for mating.

Key words: genomic evaluation, phenotypic effects, recessive disorders

INTRODUCTION

The rapid growth in the number of genotyped dairy cattle, which recently surpassed 1 million in the United States (CDCB, 2015), has resulted in the identification of several recessive disorders (e.g., Adams et al., 2012; Cooper et al., 2013; Daetwyler et al., 2014; McClure et al., 2014) and permitted the determination of carrier status of genotyped animals using haplotypes in place of laboratory tests (Cole et al., 2013). Five haplotypes affecting fertility in Holsteins (HH1–HH5) have been identified using the method of VanRaden et al. (2011b), and the causative variants now are known for HH1 (Adams et al., 2012), HH3 (Daetwyler et al., 2014; McClure et al., 2014), HH4 (Fritz et al., 2013), and HH5 (Schütz et al., 2016). The same method was used to determine carrier status for Holstein haplotypes that track bovine leukocyte adhesion deficiency (**BLAD**; Shuster et al., 1992), brachyspina (Charlier et al., 2012), complex vertebral malformation (**CVM**; Agerholm et al., 2001), deficiency of uridine monophosphate synthase (**DUMPS**; Shanks et al., 1984), and mulefoot (syndactyly; Duchesne et al., 2006); results are now routinely reported by the Council on Dairy

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Cattle Breeding (Reynoldsburg, OH). A new Holstein recessive, cholesterol deficiency (**CD**), was identified by Kipp et al. (2015), but assignment of carrier status is complicated by the presence of harmful and normal variants that have identical SNP haplotypes with a new mutation in one family member a few generations back. The recent discovery of the causal mutation for the Holstein haplotype for cholesterol deficiency (**HCD**) in the apolipoprotein B (*APOB*) gene (Charlier, 2016; Menzi et al., 2016; Schütz et al., 2016) has resulted in the development of an exact test for carrier status, which is available from some laboratories and is included on some SNP chips. Haplotype tests are available for Brown Swiss haplotypes 1 and 2 (**BH1** and **BH2**), and the causal variant for BH2 has been identified in the *TUBD1* gene (Schwarzenbacher et al., 2016). Brown Swiss haplotype tests for spinal dysmyelination (**SDM**; Thomsen et al., 2010), spinal muscular atrophy (**SMA**; Krebs et al., 2007), and Weaver Syndrome (McClure et al., 2013; Kunz et al., 2016) also are provided, but those tests do not directly include the causative mutation. Causal variants are known for SDM, SMA, and Weaver Syndrome, but the tests currently available on commercial SNP are based on haplotypes. An exact test of the loss-of-function mutation is available for Jersey haplotype 1 (Sonstegard et al., 2013), and a haplotype test is available for Jersey haplotype 2 (VanRaden et al., 2014); a haplotype that affects conception rate in Ayrshires (**AH1**) also is reported (Cooper et al., 2014). The status of each haplotype is reported for all animals receiving genomic evaluations and that information can be used when making mating and culling decisions. Sonstegard et al. (2013) showed that the concordance of haplotype with gene tests generally was very good, ranging from 94.4 (SDM) to 100% (DUMPS).

Many of these recent recessives were discovered because no homozygous embryos survive, causing reduced fertility (e.g., VanRaden et al., 2011b; Fritz et al., 2013), but limited information exists in the literature on the effects of recessive disorders on other phenotypes, such as yield and longevity of heterozygous females. Some recessive defects, such as DUMPS and HCD, are known to cause differing levels of lactic acid or cholesterol in the blood or milk when heterozygous, and may cause other phenotypic effects. Hoeschele and Meinert (1990) found that daughters of Brown Swiss bulls carrying the Weaver recessive had higher production than daughters of noncarriers, and carrier cows had significantly higher milk and fat production than noncarriers. A within-family analysis of sons and grandsons of Skokie Sensation Ned, the originator of the DUMPS disorder, found that heterozygotes had significantly higher genetic merit for milk and protein yields and economic

merit (Shanks and Greiner, 1992). Powell et al. (1996) reported that daughters of Holstein carrier bulls for BLAD had significantly lower protein yields ($P = 0.02$) and higher protein concentrations ($P = 0.10$) than daughters of noncarriers, but the magnitudes of the differences were small. Nielsen et al. (2003) found that fertility was depressed for Holstein cows carrying CVM-affected calves.

Any recessive which results in the death of calves following their birth and early rearing period, such as bovine hereditary zinc deficiency (Yuzbasiyan-Gurkan and Bartlett, 2006), is particularly problematic because the economic impact is much greater than that of recessives that cause early embryonic loss. Recessives that cause loss later in gestation are also expensive if the cow is culled or the next lactation is delayed. The purpose of the current study was to characterize the phenotypic and genetic effects of 18 recessive disorders in the Ayrshire, Brown Swiss, Holstein, and Jersey breeds on 8 yield and fitness traits in the US dairy cattle population and to estimate economic losses.

MATERIALS AND METHODS

Data

The data used in our analysis consisted of phenotypes and direct genomic values (**DGV**) from the August 2015 US national genetic evaluation for genotyped daughters of carrier bulls for 18 recessive haplotypes. The number of animals included in the analysis ranged from 8 for BHW to 17,869 for HH3. The 8 traits included in this study were milk, fat, and protein yields, SCS, single-trait productive life (**PL**), daughter pregnancy rate (**DPR**), heifer conception rate (**HCR**), and cow conception rate (**CCR**). Cows had a record in the phenotypic analysis for each lactation in the database for every trait but PL and HCR, which had one (lifetime) value, and only animals with phenotypes for all traits were included in the analysis. Animals had only one record each in the analysis of DGV because cows have only one breeding value for each trait. Genotypes for all cows were imputed to the common set of 60,671 SNP used for US genomic evaluations in August 2015 with findhap.f90 version 3 (VanRaden et al., 2011a). The haplotyping strategy was based on the methods described in detail in VanRaden et al. (2011a) and Sonstegard et al. (2013), and refined as discussed by Null and VanRaden (2016). Those SNP were selected based on performance criteria, such as minor allele frequencies, parent-progeny conflicts, call rates, and correlations with other SNP (Wiggans et al., 2010). Descriptive information for each recessive in this study

Table 1. Description of the recessive haplotypes evaluated for phenotypic effects on yield, longevity, and fertility

Breed ¹	Haplo ²	OMIA ID ³	Functional/gene name ⁴	Haplotype frequency (%)	BTA	Location (bp) ⁵	Timing ⁶	Bulls ⁷	Cows ⁸
AY	AH1	001934	PIRM/ <i>UBE3B</i>	13.0	17	65,921,497	W	15	32
BS	BH1	001825	—	6.67	7	42,811,272 – 47,002,161	E	42	300
	BH2	001939	<i>TUBD1</i>	7.78	19	10,617,246 – 11,726,846	B, W	50	508
	BHD	001247	SDM/ <i>SPAST</i>	2.19	11	14,742,058	W	13	97
	BHM	000939	SMA/ <i>KDSR (FVT1)</i>	3.61	24	62,118,139 – 62,156,760	W	17	77
	BHW	000827	Weaver/ <i>PNPLA8</i>	1.56	4	49,616,352 – 49,738,691	W	5	8
HO	HCD	001965	Cholesterol deficiency/ <i>APOB</i>	2.5	11	77,953,380 – 78,040,118	W	334	11,333
	HH0	000151	Brachyspina/ <i>FANCI</i>	2.76	21	21,184,869 – 21,188,198	E, B	325	9,150
	HH1	000001	<i>APAF1</i>	1.92	5	63,150,400	E	233	5,749
	HH2	001823	—	1.66	1	94,860,836 – 96,553,339	E	221	8,312
	HH3	001824	<i>SMC2</i>	2.95	8	95,410,507	E	438	17,869
	HH4	001826	<i>GART</i>	0.37	1	1,277,227	E	54	1,218
	HH5	001941	<i>TFB1M</i>	2.22	9	92,350,052 – 93,910,957	E	434	12,587
	HHB	000595	BLAD/ <i>ITGB2</i>	0.25	1	145,119,004	W	14	131
	HHC	001340	CVM/ <i>SLC35A3</i>	1.37	3	43,412,427	E, B	106	1,762
	HHM ⁹	000963	Mulefoot/ <i>LRP4</i>	0.07	15	77,663,790 – 77,701,209	B	5	36
JE	JH1	001697	<i>CWC15</i>	12.10	15	15,707,169	E	344	13,300
	JH2	001942	—	1.3	26	8,812,759 – 9,414,082	E	58	1,279

¹AY = Ayrshire, BS = Brown Swiss, HO = Holstein, and JE = Jersey.

²The recessives analyzed include the fertility haplotypes in each breed; haplotypes for spinal dysmyelination (BHD), spinal muscular atrophy (BHM), and Weaver (BHW) in Brown Swiss; and haplotypes for bovine leukocyte adhesion deficiency (HHB), brachyspina (HH0), complex vertebral malformation (HHC), and mulefoot (syndactyly; HHM) in Holsteins (http://aipl.arsusda.gov/reference/recessive_haplotypes_ARR-G3.html).

³Online Mendelian Inheritance in Animals (OMIA) identification number for *Bos taurus* (National Center for Biotechnology Information species code 9913).

⁴A blank value indicates that the functional mutation is not currently known.

⁵When the causal variant is known (e.g., AH1), this is the location of the associated SNP. When a candidate gene is known, but not the causal variant, the location includes the coordinates of that gene. When no causal variant or candidate gene is known, this column contains the coordinates of the haplotype used to track the recessive.

⁶Timing of embryonic loss or calf death for homozygous individuals: B = calf death at or shortly following birth, E = embryonic loss or abortion, and W = calf death is weeks or months following birth. In some cases, losses may occur in more than one time period.

⁷The number of carrier bulls with genotyped daughters that had yield deviations in the August 2015 US national genetic evaluation.

⁸The number of genotyped cows with yield deviations in the August 2015 US national genetic evaluation.

⁹Mulefoot (syndactyly) is not lethal in that it does not actually kill affected calves, but affected animals are always culled and there is a resulting economic loss associated with the condition.

Table 2. The number of cows and bulls genotyped and the number of SNP markers on the chips used for genotyping

Chip name	SNP count	Ayrshire		Brown Swiss		Holstein		Jersey	
		Bulls	Cows	Bulls	Cows	Bulls	Cows	Bulls	Cows
Imputed genotype	0	0	35	0	332	0	4,920	0	340
Illumina BovineSNP50 BeadChip, Version 1 ¹	54,001	16	0	5,359	85	32,221	19,158	4,680	899
Illumina BovineSNP50 BeadChip, Version 2 ¹	54,609	336	154	8,072	118	30,231	28,478	2,613	776
Illumina Bovine3K BeadChip ¹	2,900	0	3	10	455	2,094	46,241	68	9,358
Illumina BovineHD BeadChip ¹	777,962	516	9	12	0	969	503	27	16
Affymetrix AxiomBOS1 BeadChip ²	648,875	0	0	0	0	19	0	0	0
Illumina BovineLD BeadChip ¹	6,909	6	915	9	517	3,182	151,566	149	9,714
GeneSeek Genomic Profiler ³	8,762	2	54	106	448	9,902	38,578	1,096	12,287
GeneSeek Genomic Profiler-HD ³	77,068	371	420	370	153	7,545	13,551	587	944
GeneSeek Genomic Profiler-Super LD ³	19,809	53	332	936	553	20,698	56,092	2,674	16,621
Zoetis Low Density ⁴	11,410	0	0	18	75	1,069	104,657	121	9,641
Zoetis Medium Density ⁴	56,955	1	559	0	2	491	1,756	20	2
EuroG10K ¹	9,072	0	0	0	0	87	692	0	0
Illumina BovineLD BeadChip, Version 1.1 ¹	6,912	0	0	0	14	946	11,798	6	1,708
GeneSeek Genomic Profiler LD Version 3 ³	26,151	86	637	301	769	22,953	56,976	3,672	19,688
Zoetis Low Density Version 2 ⁴	17,619	0	1	8	259	4,232	181,631	155	12,884
Zoetis Medium Density Version 2 ⁴	60,914	0	0	2	0	349	2,143	44	4
GGP Bovine 150K ³	139,480	97	32	364	37	13,322	8,162	1,383	506
GGP Bovine 7K ³	7,083	0	0	0	0	3	376	0	16
Total		1,484	3,186	15,567	3,817	727,278	150,313	17,295	95,404

¹Illumina Inc., San Diego, CA.²Affymetrix, Santa Clara, CA.³Gene Seek, Lincoln, NE.⁴Zoetis, Florham Park, NJ.

is provided in Table 1, and Table 2 includes details about the cow and bull genotypes included in the study. Definitions of the SUP haplotypes used in this analysis are provided in Supplemental Table S1 (<http://dx.doi.org/10.3168/jds.2015-10777>).

Whereas the causal variants are now known for several recessive disorders, and those variants now are present on some of the SNP genotyping chips, many of the recessive tests are still based on haplotypes rather than individual SNP (Table 3). Most of the genotypes in the national dairy database do not include the SNP tests, so haplotype tests must be used until enough data are available to accurately impute carrier status for older genotypes. There also are cases (e.g., HH0) where SNP tests are used, but those test results are not transmitted from the genotyping laboratories to the Council on Dairy Cattle Breeding (Bowie, MD) due to intellectual property agreements. Results from custom assays, such as Sequenom MassARRAY (Agena Bioscience, San Diego, CA) genotypes, used by many research groups, are not part of the national dairy database, either.

Assignment of HCD Carrier Status

Two variants are unique to the SNP haplotype for HCD in the population, one that includes the harmful variant and one that does not. Both haplotypes share identical SNP genotypes but have different origins, which is similar to variants associated with arachnemia (Drögemüller et al., 2010) and zinc deficiency-like syndrome (Jung et al., 2014). The haplotype carrying the harmful variant is transmitted by the bull Maughlin Storm (HOCAN000005457798) and his descendants, whereas the nonharmful haplotype is transmitted by Willowholme Mark Anthony (HOCAN000000334489) and his descendants. The nonlethal haplotype actually originated with the bull Fairlea Royal Mark (HOCAN000000299855; Kipp et al., 2015), but that animal is not genotyped in the United States and pedigrees instead are traced back to Anthony. A list of carriers and putatively affected animals was constructed based on genotypes, and pedigrees are then traced back to determine the origin of the recessive haplotypes. Heterozygotes whose pedigrees trace back to Storm, but not Anthony, are coded as 1, indicating that they are known carriers of the harmful allele. Similarly, homozygotes with pedigree paths to Storm but not Anthony are coded as 2, indicating that they are homozygous for the harmful version of the haplotype. Heterozygotes and homozygotes with both Storm and Anthony in their pedigrees are coded as 3 and 4, respectively, indicating that they may be carriers of the harmful haplotype. Animals with incomplete pedigrees are

coded as 3 or 4 when their pedigrees do not trace far enough back to determine which haplotype version of the recessive haplotype they carry. Only animals known to carry 0 or 1 copies of the harmful allele (coded 0 or 1 in the haplotype analysis) were included in the analysis of genetic and phenotypic effects. Phenotypes of homozygous recessive animals were not analyzed for HCD effects because they die before any trait other than stillbirth can be measured.

Estimation of Phenotypic and Genetic Effects

The following fixed effects model was fit to phenotypic and genetic values for all traits using PROC GLM in SAS 9.4 for Linux (SAS Institute Inc., Cary, NC):

$$y_{ijk} = \mu + \text{sire}_j + \text{haplotype}_k + e_{ijk},$$

where y_{ijk} is the phenotypic value or DGV for any of the 8 traits evaluated for cow i , μ is the overall mean, sire_j is the fixed effect of the cow's sire, haplotype_k is the fixed effect of the haplotype being considered (coded as 0 or 1 copies of the minor allele), and e_{ijk} is the random residual error term. Sire effects were absorbed, and haplotype effects were tested for differences using a t test. Phenotypes were preadjusted for nongenetic factors by subtracting management group, parity-sex, and herd-by-sire effects from each observation. The DGV were calculated as the sum of individual SNP effects from the August 2015 genomic evaluation (Wiggans et al., 2011) plus breed- and trait-specific intercepts (Cole and Null, 2013). A within-trait Bonferroni adjustment was used to correct for multiple comparisons, and significance was declared when $P < 0.0028$.

Genetic Merit of Carrier Versus Noncarrier Bulls

Culling decisions are made based on a combination of predicted genetic merit and recessive carrier status. It is, therefore, desirable to know if carrier bulls are of consistently higher or lower genetic merit than noncarriers. The average genomic PTA for lifetime net merit (NMS\$; VanRaden and Cole, 2014) was calculated for the bulls of each breed with semen currently available for purchase (status codes of A, for active AI bulls; F, for foreign-sampled bulls with semen available in the United States; G, for genomically tested bulls at least 12 mo old; and L, for limited-use bulls). These were then compared based on carrier status to determine if carriers or noncarriers have a systematic advantage over other bulls. Averages were tested for differences using a z -test within breed.

Table 3. Availability of SNP tests, the type of test used, the number of genotypes that included haplotype and SNP tests, and SNP genotype frequencies for recessive disorders from the August 2015 evaluation

Haplotype ¹	SNP test available	Type of test used	Number of haplotype tests	Number of SNP tests ²	Genotype count			
					AA	Aa	aa	aa (LD) ³
AH1	Yes	Haplotype	4,670	0 ⁴	3,603	1,063	4	1
BH1	No	Haplotype	19,384	—	16,328	3,042	14	7
BH2	No	Haplotype	19,384	—	16,381	2,986	17	11
BHD	Yes	Haplotype	19,384	3,223	18,774	609	1	1
BHM	Yes	Haplotype	19,384	3,223	18,456	927	1	0
BHW	Yes ⁵	Haplotype	19,384	2,596	19,056	327	1	0
HCD ⁶	No ⁷	Haplotype	877,591	—	820,953	56,363	275	188
HH0	Yes	Haplotype	877,591	0 ⁸	838,546	39,035	10	10
HH1	Yes	SNP	877,591	237,132	847,936	29,637	18	18
HH2	No	Haplotype	877,591	—	855,677	21,892	22	22
HH3	Yes	Haplotype	877,591	2,506	826,523	51,052	16	12
HH4	Yes	Haplotype	877,591	2,506	872,504	5,085	2	2
HH5	No	Haplotype	877,591	—	841,186	36,397	8	4
HHB	Yes	SNP	877,591	217,188	873,019	4,568	4	3
HHC	Yes	SNP	877,591	18,369	855,740	21,810	41	39
HHM	Yes	SNP	877,591	9,019	876,652	939	0	0
JH1	Yes	SNP	112,699	45,337	87,808	24,855	36	32
JH2	No	Haplotype	112,699	—	109,810	2,853	36	36

¹The recessives analyzed include the fertility haplotypes in each breed; haplotypes for spinal muscular atrophy (BHM), spinal dysmyelination (BHD), and Weaver (BHW) in Brown Swiss; and haplotypes for bovine leukocyte adhesion deficiency (HHB), brachyspina (HH0), complex vertebral malformation (HHC), and mulefoot (syndactyly; HHM) in Holsteins (http://aipl.arsusda.gov/reference/recessive_haplotypes_ARR-G3.html).

²A blank value indicates that the functional mutation is not currently known.

³The number of homozygous calls from low-density (<30K SNP) genotypes.

⁴No genotypes from chips including the AH1 causal SNP were received before the August 2015 genetic evaluation run.

⁵Several SNP associated with Weavers are included on some SNP genotyping chips, including the recent candidate variant reported in the *PNPLA8* gene.

⁶The heterozygote counts for HCD include code 3 and 4 animals.

⁷The causal variant for HCD has recently been identified, but test results from SNP genotyping chips are not yet available.

⁸The SNP genotype for HH0 (brachyspina) is not transmitted from the genotyping laboratories to the Council on Dairy Cattle Breeding.

Economic Impact of Recessives

The economic impact on producers of the recessives carried by each breed was calculated as the product of the total national herd size (9 million), the proportion of the population represented by each breed based on the number of records from cows calving in 2014 included in the national genetic evaluations (CDCB, 2016), and the sum of squared carrier frequencies multiplied by the economic impact of the loss for each recessive. An average value of \$200 was used for pregnancy loss [A. De Vries (University of Florida, Gainesville), personal communication] and a value of \$342 was used for lost calves (assumed dead at 21 d), which is the average calf value assigned to stillbirths in the lifetime net merit calculations (VanRaden and Cole, 2014) plus \$2 per day for rearing costs for 21 d. Pregnancy losses are coded E (early pregnancy) in Table 1, and calf losses are coded B (death at or near birth) or W (calf death weeks after birth), and costs were calculated using the latest time of loss for a recessive with effects over multiple time periods. All haplotypes within a breed were

assumed independent although, for example, HH2 and HH4 both are located on chromosome 1.

RESULTS AND DISCUSSION

Not all haplotypes are fatal before or near birth, and small numbers of homozygous genotypes were observed for all recessives studied (Table 3). A substantial number of homozygotes (275) for CD were identified, 188 of those using lower-density SNP genotypes. VanRaden and Null (2015) reported that females in the US homozygous for CD had neither breeding nor lactation records, strongly suggesting that they exited the herd as calves. The number of homozygous genotypes for other lethal haplotypes, such as HH0 and JH2, were very small, and most homozygotes are likely to be imputation errors based on low-density genotypes. However, homozygous genotypes also can be the result of imperfect linkage disequilibrium between haplotypes and causal mutations or incomplete penetrance.

The phenotypic and genetic effects of the recessive haplotypes on phenotypic and genetic values of 8 traits

from heterozygous daughters of heterozygous bulls are shown in Tables 4 and 5, respectively. *P*-values were adjusted on a within-trait basis to account for multiple comparisons. A significant haplotype effect does not necessarily indicate a causal relationship, and the association of the genotype with trait differences may be due to physical linkage of the recessive with the true causal variant. For example, Hedrick (2013) demonstrated that changes in the frequencies of coat color alleles in mice selected for weight gain were properly attributable to physical linkage of the recessive with the true causal variant, not pleiotropy. It should be possible to test such a hypothesis in the US dairy cattle population once more generations are available to test for linkage decay over time.

Cholesterol Deficiency

In most cases studied previously, it was straightforward to determine carrier status using haplotype tests. This has not proven to be the case with cholesterol deficiency, making it an excellent example of how to use inconclusive genotypes to identify probable carriers. Cholesterol deficiency was identified recently because farmers in Germany reported problems with calves that suffered from loss of appetite, decreases in BW, and diarrhea that were not responsive to veterinary intervention (Kipp et al., 2015). The haplotype has a higher frequency (2.5%, based on all known and suspected heterozygotes) than many other recessives that also result in calf deaths (Figure 1), such as CVM (1.37%) and BLAD (0.25%). Therefore, it is important that carriers and affected animals be identified as quickly as possible. We identified 56,641 Holsteins as carriers of HCD in the August 2015 genomic evaluation. Of those animals, 30,928 (54.6%) were heterozygous for the harmful haplotype (code 1) and parental origin could be determined, 275 (0.48%) were homozygous for the harmful haplotype (code 2), 25,077 (44.2%) were heterozygous for the recessive haplotype but parental origin could not be determined (code 3), and 358 (0.63%) were homozygous for the recessive haplotype but parental origin could not be determined (code 4). Approximately half of the putative carriers and affected animals may carry the normal form of the recessive haplotype, which means that the economic impact of HCD is further increased because some unaffected animals will be culled. A laboratory test now is available for the causal variant in the *APOB* gene (Charlier, 2016; Menzi et al., 2016; Schütz et al., 2016), and if the exact test is used rather than the haplotype test fewer noncarriers will be culled.

Hypocholesterolemia is characteristic of CD (Kipp et al., 2015), but the relationship of blood cholesterol

levels with milk fat production is not known. Carriers of HCD had significantly higher protein yields (2.36 ± 0.75 kg, $P = 0.0016$) than noncarriers. It is not clear how HCD might affect protein yield, and this probably represents a physical linkage of the recessive with the true causal variant effect rather than pleiotropy because Maughlin Storm (HOCAN000005457798), the bull in which the HCD mutation originated, had 2 very good copies of chromosome 11 that were transmitted to his daughters and granddaughters along with HCD. However, HCD carriers had significantly higher genetic merit for fat, protein, SCS, PL, DPR, CCR, and HCR than noncarriers. It is important to note, however, that the magnitude of the effects is small (<1 kg of fat and protein and <1% of DPR, CCR, and HCR) and may reflect increased power due to a large sample size rather than biologically important differences.

Effects of Other Recessive Haplotypes

The phenotypic effects of the minor allele for each recessive on the 8 traits included in the analysis are shown in Table 4, and the genetic effects in Table 5. The number of observations (*N*) is smaller in Table 5 than Table 4 because many cows had more than one lactation record for production and fertility traits, increasing the number of records used, but only one genetic evaluation for each trait. Of the 136 phenotype tests conducted, 4 were significant at the 0.0028 level (Bonferroni-adjusted within trait) or higher, including 2 with fat yield, 1 with protein yield, and 1 with CCR. A total of 35 of 136 DGV effects were different from 0, including 2 for milk, 3 for fat, 6 for protein, 3 for SCS, 5 for PL, 4 for DPR, 5 for CCR, and 7 for HCR. The majority of the significant genetic effects (29 of 35) were observed for HO haplotypes, and probably reflect greater statistical power due to large sample sizes rather than differences of great biological importance. The following discussion groups results by breed.

Ayrshire. We found no significant phenotypic or genetic effects on fertility, although Cooper et al. (2014) reported a significant effect on sire conception rate. Although not reflected by the fertility traits included in this analysis, Venhoranta et al. (2014) also reported that AH1 is associated with increased juvenile mortality. The lack of significant effects is probably due to the very limited number of animals (15 carrier bulls with a total of 32 genotyped daughters) available for analysis. Whereas some effects appear to be large, such as the 763.22 ± 349.59 kg effect on milk yield, the standard errors also are large, reflecting the limited information available.

Brown Swiss. VanRaden et al. (2011b) reported a negative effect of BH1 on sire conception rate, whereas

Table 4. Effect of the minor allele for recessive haplotypes on phenotypes preadjusted for nongenetic factors from heterozygous daughters of heterozygous bulls

Breed ¹	Haplo ²	Statistic ³	Trait ⁴							
			Milk (kg)	Fat (kg)	Protein (kg)	SCS	PL (mo)	DPR (%)	CCR (%)	HCR (%)
AY	AH1	N	67	67	67	58	16	37	37	37
		Effect	763.22	-25.27	-6.57	-0.463	7.65	7.18	22.63	-4.12
		SE	349.59	15.80	12.34	0.300	10.99	11.72	20.13	13.85
BS	BH1	N	628	628	628	600	173	427	427	427
		Effect	-160.09	2.99	0.83	-0.003	3.89	4.18	5.07	3.69
		SE	157.78	5.99	4.05	0.089	2.50	4.67	8.53	3.33
	BH2	N	848	848	848	813	286	575	575	575
		Effect	-57.83	-3.91	4.22	-0.023	-3.30	-1.69	-4.47	0.89
		SE	111.48	5.12	3.48	0.077	1.74	4.36	6.62	3.03
	BHD	N	208	208	208	195	61	149	149	149
		Effect	413.21	20.12	6.08	0.627	-3.81	-4.53	4.83	4.42
		SE	271.32	9.80	7.20	0.145	4.24	5.80	12.22	6.44
	BHM	N	255	255	255	234	48	175	175	175
		Effect	725.84	-25.49	-24.02	0.300	0.15	11.60	-18.70	-9.15
		SE	273.49	11.04	8.24	4.76	0.13	5.41	12.94	3.87
BHW	N	24	24	24	11	1	11	11	11	
	Effect	NE ⁵	NE	NE	NE	NE	NE	NE	NE	
	SE	NE	NE	NE	NE	NE	NE	NE	NE	
HO	HCD	N	14,909	14,909	14,909	14,583	3,560	10,694	10,694	10,694
		Effect	-9.37	1.69	2.36*	0.64	0.64	1.95	2.97	-0.43
		SE	28.47	1.12	0.75	0.49	0.49	0.97	1.30	0.78
	HH0	N	12,836	12,836	12,836	12,557	3,905	10,360	10,360	10,360
		Effect	-26.46	5.92*	0.36	-0.003	0.07	1.10	3.38	0.95
		SE	28.22	1.16	0.80	0.019	0.48	0.86	1.14	0.79
	HH1	N	7,420	7,420	7,420	7,305	2,601	6,025	6,025	6,025
		Effect	-87.51	2.61	-0.01	-0.005	-0.66	-2.09	-3.39	0.77
		SE	31.27	1.51	1.07	0.025	0.59	1.18	1.43	1.05
	HH2	N	10,370	10,370	10,370	10,126	3,152	8,273	8,273	8,273
		Effect	-33.00	2.71	0.71	-0.016	0.04	-0.82	-1.70	-1.12
		SE	32.49	1.29	0.86	0.021	0.52	1.11	1.36	0.91
	HH3	N	19,131	19,131	19,131	18,731	6,033	15,965	15,965	15,965
		Effect	9.96	0.95	-1.36	-0.033	0.13	0.04	0.31	-0.03
		SE	23.45	0.91	0.64	0.016	0.37	0.80	0.94	0.66
	HH4	N	1,521	1,521	1,521	1,481	456	1,224	1,224	1,224
		Effect	-101.16	-3.26	-2.78	-0.020	1.97	6.32	-1.37	3.29
		SE	98.37	3.13	2.25	0.058	1.33	3.04	3.53	2.27
	HH5	N	14,473	14,473	14,473	14,225	3,835	11,232	11,232	11,232
		Effect	5.71	1.97	0.62	-0.029	0.21	-0.66	-1.81	-0.60
		SE	25.44	1.06	0.72	0.019	0.47	0.97	1.20	0.76
	HHB	N	306	306	306	286	52	204	204	204
		Effect	139.87	-11.55	-4.84	0.101	1.71	4.29	5.93	3.31
		SE	353.82	10.57	7.27	0.168	4.71	4.84	10.38	5.54
	HHC	N	4,393	4,393	4,393	4,207	794	3,097	3,097	3,097
		Effect	89.69	-6.91*	-2.77	-0.016	-0.15	0.33	-2.49	-1.05
		SE	47.64	2.23	1.47	0.034	1.26	1.21	2.32	1.17
	HHM	N	112	112	112	111	23	91	91	91
		Effect	-0.18	5.18	-4.84	-0.178	-8.25	-1.22	-6.76	3.12
		SE	0.24	16.76	10.18	0.241	7.85	6.95	10.18	8.68
JE	JH1	N	21,152	21,152	21,152	20,833	5,791	17,348	17,348	17,348
		Effect	6.43	-0.99	1.43	0.020	0.34	-0.52	-0.86	0.19
		SE	14.86	0.76	0.53	0.014	0.39	0.62	0.76	0.49
	JH2	N	2,446	2,446	2,446	2,364	529	1,865	1,865	1,865
		Effect	-55.59	1.30	-0.91	-0.058	-0.77	-2.41	-7.17*	-0.98
		SE	62.39	2.57	1.80	0.043	1.57	1.99	2.33	1.66

¹AY = Ayrshire, BS = Brown Swiss, HO = Holstein, and JE = Jersey.²The recessives analyzed include the fertility haplotypes in each breed; haplotypes for spinal muscular atrophy (BHM), spinal dysmyelination (BHD), and Weaver (BHW) in Brown Swiss; and haplotypes for bovine leukocyte adhesion deficiency (HHB), brachyspina (HH0), complex vertebral malformation (HHC), and mulefoot (syndactyly; HHM) in Holsteins (http://aipl.arsusda.gov/reference/recessive_haplotypes_ARR-G3.html).³n = the number of observations used in the analysis.⁴PL = single-trait productive life; DPR = daughter pregnancy rate; HCR = heifer conception rate; and CCR = cow conception rate.⁵NE = not estimable.*Significance at the 5% level following within-trait Bonferroni adjustment ($P < 0.0028$).

Table 5. Effect of the minor allele for recessive haplotypes on direct genomic values of heterozygous daughters of heterozygous bulls

Breed ¹	Haplo ²	Statistic	Trait ³								
			Milk (kg)	Fat (kg)	Protein (kg)	SCS	PL (mo)	DPR (%)	CCR (%)	HCR (%)	
AY	AH1	N	32	32	32	32	32	32	32	32	32
		Effect	-7.92	-1.06	0.42	-0.016	0.07	0.75	0.24	0.37	
		SE	53.67	1.93	1.53	0.033	0.49	0.38	0.30	0.33	
BS	BH1	N	300	300	300	300	300	300	300	300	
		Effect	2.17	-0.18	-0.42	0.007	-0.23	-0.11	-0.04	0.12	
		SE	18.19	0.78	0.52	0.014	0.23	0.14	0.13	0.13	
	BH2	N	508	508	508	508	508	508	508	508	
		Effect	17.39	-0.001	0.42	0.001	-0.09	-0.01	0.10	-0.05	
		SE	13.43	0.522	0.38	0.010	0.16	0.10	0.10	0.10	
	BHD	N	97	97	97	97	97	97	97	97	
		Effect	-44.86	-1.53	-0.82	0.026	-0.06	0.17	-0.06	0.18	
		SE	29.70	1.03	0.84	0.024	0.41	0.24	0.24	0.21	
	BHM	N	77	77	77	77	77	77	77	77	
		Effect	-70.11	-1.47	-1.70	-0.002	-0.10	-0.38	-0.10	-0.17	
		SE	41.90	1.36	1.32	0.026	0.43	0.30	0.26	0.27	
BHW	N	8	8	8	8	8	8	8	8		
	Effect	-135.62	-3.17	-3.63	-0.040	1.60	0.40	2.10	0.30		
	SE	66.70	14.15	10.62	0.213	3.93	1.66	0.89	1.06		
HO	HCD	N	11,332	11,332	11,332	11,332	11,332	11,332	11,332	11,332	
		Effect	-4.26	0.91*	0.72*	-0.022*	0.30*	0.17*	0.35*	0.11*	
		SE	4.02	0.15	0.09	0.002	0.03	0.02	0.03	0.02	
	HH0	N	9,150	9,150	9,150	9,150	9,150	9,150	9,150	9,150	
		Effect	5.30	0.70*	-0.14	-0.0002	-0.01	-0.07	0.08	-0.07*	
		SE	4.30	0.16	0.10	0.0026	0.03	0.03	0.03	0.02	
	HH1	N	5,749	5,749	5,749	5,749	5,749	5,749	5,749	5,749	
		Effect	-21.94*	-0.46	-0.54*	0.020*	-0.41*	-0.22*	-0.30*	-0.026*	
		SE	5.31	0.20	0.12	0.003	0.04	0.03	0.04	0.03	
	HH2	N	8,312	8,312	8,312	8,312	8,312	8,312	8,312	8,312	
		Effect	12.07	0.44	0.32*	-0.001	0.18*	0.04	0.14*	0.15*	
		SE	4.59	0.17	0.10	0.003	0.03	0.03	0.03	0.02	
	HH3	N	17,869	17,869	17,869	17,869	17,869	17,869	17,869	17,869	
		Effect	-20.76*	0.19	-0.53*	-0.014	0.05	-0.07*	-0.14*	-0.27*	
		SE	2.97	0.11	0.07	0.002	0.02	0.02	0.02	0.02	
	HH4	N	1,218	1,218	1,218	1,218	1,218	1,218	1,218	1,218	
		Effect	-11.26	-0.85	-0.52	-0.013	0.09	0.06	0.10	0.04	
		SE	11.10	0.42	0.26	0.008	0.09	0.08	0.08	0.06	
	HH5	N	12,587	12,587	12,587	12,587	12,587	12,587	12,587	12,587	
		Effect	-5.48	-0.05	-0.30*	-0.012	-0.01	0.002	-0.04	-0.11*	
		SE	3.62	0.13	0.08	0.002	0.03	0.024	0.03	0.02	
	HHB	N	131	131	131	131	131	131	131	131	
		Effect	-48.17	-0.07	-0.07	0.09	0.04	0.06	0.08	-0.15	
		SE	41.02	1.62	0.33	0.96	0.03	0.28	0.31	0.23	
	HHC	N	1,762	1,762	1,762	1,762	1,762	1,762	1,762	1,762	
		Effect	-11.38	-0.49	0.16	0.003	0.14	0.01	-0.02	-0.02	
		SE	10.03	0.36	0.08	0.006	0.23	0.06	0.07	0.05	
	HHM	N	36	36	36	36	36	36	36	36	
		Effect	104.61	1.59	1.88	0.161*	-1.56*	-0.73	-0.41	0.02	
		SE	73.78	2.82	1.38	0.044	0.46	0.47	0.54	0.40	
JE	JH1	N	13,300	13,300	13,300	13,300	13,300	13,300	13,300	13,300	
		Effect	5.39	-0.48*	0.26*	0.002	-0.19*	-0.18*	-0.27*	-0.23*	
		SE	2.69	0.10	0.07	0.001	0.02	0.02	0.02	0.01	
	JH2	N	1,279	1,279	1,279	1,279	1,279	1,279	1,279	1,279	
		Effect	10.06	0.02	-0.02	-0.001	-0.10	-0.12	-0.09	-0.11	
		SE	9.44	0.35	0.25	0.005	0.08	0.07	0.07	0.05	

¹AY = Ayrshire, BS = Brown Swiss, HO = Holstein, and JE = Jersey.

²The recessives analyzed include the fertility haplotypes in each breed; haplotypes for spinal muscular atrophy (BHM), spinal dysmyelination (BHD), and Weaver (BHW) in Brown Swiss; and haplotypes for bovine leukocyte adhesion deficiency (HHB), brachyspina (HH0), complex vertebral malformation (HHC), and mulefoot (syndactyly; HHM) in Holsteins (http://aipl.arsusda.gov/reference/recessive_haplotypes_ARR-G3.html).

³PL = single-trait productive life; DPR = daughter pregnancy rate; HCR = heifer conception rate; and CCR = cow conception rate.

*Significance at the 5% level following within-trait Bonferroni adjustment ($P < 0.0028$).

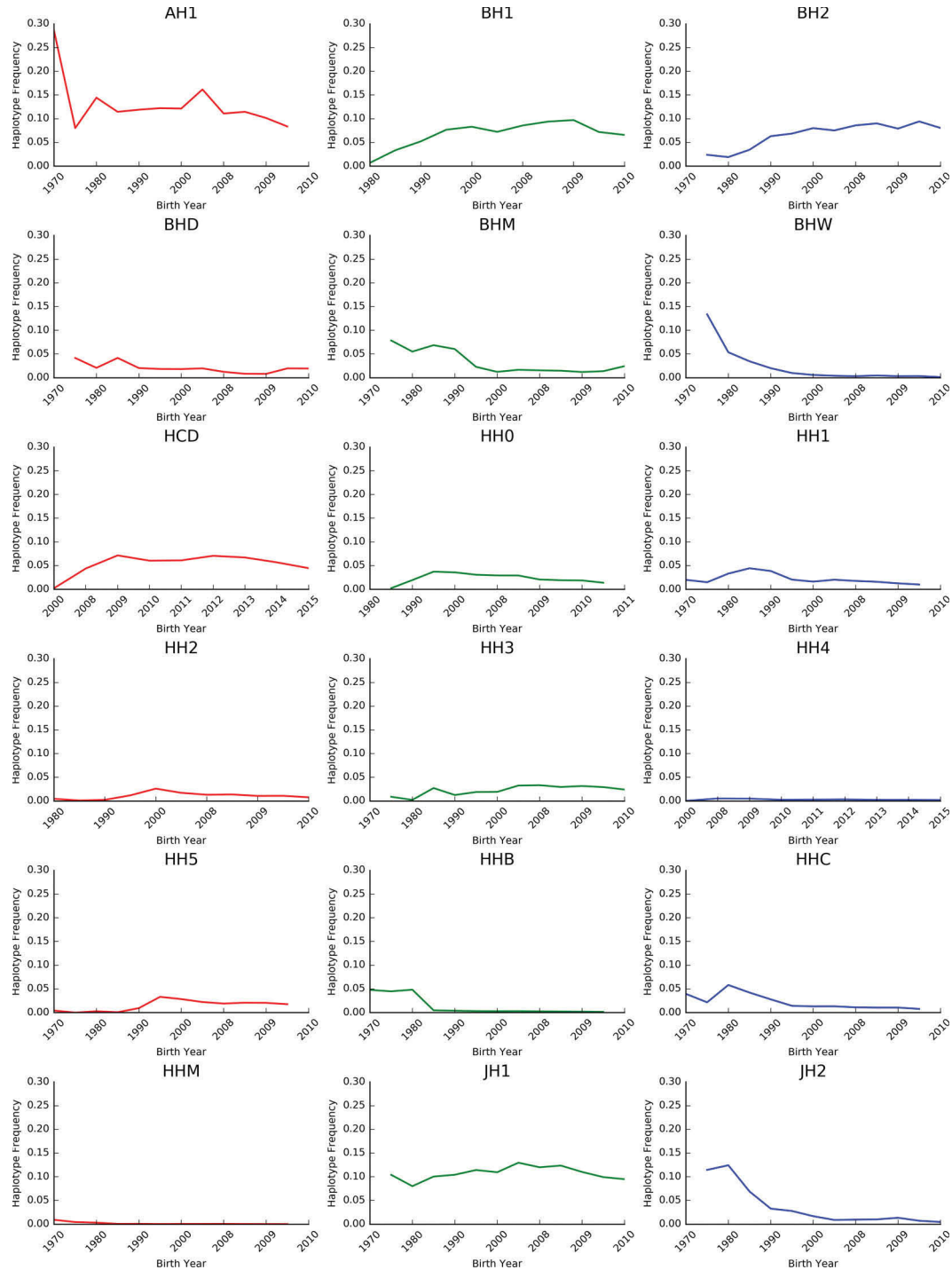


Figure 1. Changes in the frequency of recessive haplotypes. The recessives analyzed include the fertility haplotypes in each breed; haplotypes for spinal dysmyelination (BHD), spinal muscular atrophy (BHM), and Weaver (BHW) in Brown Swiss; and haplotypes for bovine leukocyte adhesion deficiency (HHB), brachyspina (HH0), complex vertebral malformation (HHC), and mulefoot (syndactyly; HHM) in Holsteins (http://aipl.arsusda.gov/reference/recessive_haplotypes_ARR-G3.html). Color version available online.

the current results show positive but nonsignificant phenotypic effects of BH1 on female fertility. No genetic effects were significant, and estimates were undesirable for DPR and CCR but favorable for HCR. The BH2

haplotype had no significant associations with traits included in our study, although Schwarzenbacher et al. (2012) reported significant phenotypic effects on still-birth and calf survival for at-risk matings. Neither the

Table 6. Differences in average lifetime net merit (\$) between bulls that are carriers of at least one recessive disorder and those that are free of known recessives by breed

Breed ¹	Carriers			Noncarriers			Difference	P-value
	N	Mean	SD	N	Mean	SD		
AY	16	286.43	194.67	53	272.41	191.69	13.02	0.407
BS	30	223.10	218.39	59	284.19	160.36	-61.09	0.087
HO	550	394.08	216.77	1,765	479.49	213.89	-85.41	<0.001
JE	99	340.51	149.00	378	338.82	153.99	1.69	0.460

¹AY = Ayrshire, BS = Brown Swiss, HO = Holstein, and JE = Jersey.

spinal dysmyelination (BHD) nor the spinal muscular atrophy (BHM) haplotype were associated with phenotypic or genetic differences. No phenotypic effects were estimable for the Weaver haplotype, possibly due to the extremely small sample size ($n = 24$), and no genetic effect differed from 0. As was the case with Ayrshire, the lack of significant effects may reflect the limited information available.

Holstein. Brachyspina (HH0) carriers had higher phenotypic and genetic fat yields of 5.92 ± 1.16 ($P < 0.0001$) and 0.70 ± 0.16 kg ($P < 0.0001$), respectively. Direct genomic values for HCR were also lower for HH0 carriers ($-0.07 \pm 0.02\%$). There were no phenotypic differences between HH1 carriers and noncarriers, but carriers had significantly lower DGV for milk and protein yields, PL, and fertility, as well as higher (unfavorable) DGV for SCS. In contrast, HH2 carriers had higher genetic merit for protein yield, PL, CR, and HCR than noncarriers. Holstein haplotype 3 carriers had lower milk and protein yields and DPR, CCR, and HCR than noncarriers. No significant results were observed for HH4, which had the fewest observations (1,218) of the Holstein fertility haplotypes. The HH5 carriers had lower DGV for both protein yield and DPR. Whereas HHB carriers did not differ from noncarriers, HHC carriers had lower phenotypes for fat yield (-6.91 ± 2.23 kg; $P = 0.0020$) than noncarriers. Mulefoot (HHM) carriers had higher SCS (0.161 ± 0.044 ; $P = 0.0009$) and lower PL (-1.56 ± 0.46 mo; $P = 0.0018$) DGV than noncarriers. It was expected that carriers of HH1 to HH5 would have lower fertility than noncarriers because those haplotypes were identified based, in part, on their effects on fertility. This was the case for all significant CCR, HCR, and DPR effects with the exception of HH2 carriers, which had higher DGV for CCR and DPR.

Jersey. The JH1 haplotype was not associated with any phenotypic effects, but we found significant differences among DGV for fat, protein, PL, DPR, CCR, and HCR. All fertility effects were undesirable, which is consistent with the reduction in sire conception rate associated with JH1 described by Sonstegard et al. (2013). The JH2 haplotype was associated with sig-

nificantly lower phenotypic CCR ($-7.17 \pm 2.33\%$, $P = 0.0022$), which is consistent with the significant effect on sire conception rate reported by VanRaden et al. (2014).

Genetic Merit of Carrier Versus Noncarrier Bulls

Bulls that were carriers of at least one recessive had higher average NM\$ (VanRaden and Cole, 2014) than noncarriers in the Ayrshire and Jersey breeds, but the differences were not significant (Table 6). Brown Swiss and Holstein carriers had genomic PTA for NM\$ that were \$61.09 ($P = 0.087$) and \$85.41 ($P < 0.001$) lower than noncarriers. The differences between Brown Swiss and Holstein bulls may reflect variation in sample size or random differences between birth year cohorts. These results suggest that the use of only noncarrier bulls in breeding programs would not affect the overall genetic trend in the population, but would require using substantially fewer bulls. This may be undesirable because it could result in higher rates of inbreeding.

Economic Impact of Recessives

Total annual losses across all 4 breeds were estimated as \$10,743,308. The majority of losses (\$7,500,265) were attributable to Holstein, with smaller losses for Ayrshire (\$109,238), Brown Swiss (\$233,414), and Jersey (\$2,900,390). The costliest recessives in each breed were AH1, BH2, JH1, and HH0. In Holsteins, HCD (\$1,696,555) and HH3 (\$1,381,452) had somewhat lower values than HH0, the former because of a lower carrier frequency and the latter due to losses occurring earlier in pregnancy. Average losses for animals of each breed were \$5.77, \$3.65, \$0.94, and \$2.96 in Ayrshire, Brown Swiss, Holstein, and Jersey, respectively, which represent the economic impact of genetic load as it affects fertility and perinatal mortality (Supplemental Table S2; <http://dx.doi.org/10.3168/jds.2015-10777>). However, these results underestimate the total effect of recessives in Holstein, Brown Swiss, and Jersey because there are large populations of those breeds in other countries that use semen from US bulls. The \$342

value used for calf deaths likely is an underestimate of actual losses because it assumes that matings are made at random, rather than considering breeding values and other selection criteria, and does not include labor and veterinary costs associated with treatment of sick calves.

Management of Carriers in the Population

Each human genome contains approximately 100 loss-of-function mutations, including about 20 genes that are completely inactivated (MacArthur et al., 2012), and the total is probably similar for cattle. In the current study, no bull or cow carried copies of more than 5 recessive haplotypes, and most animals carried no copies of any harmful haplotypes (Figure 2). The birth year, number of sons and grandsons in AI, and number

of daughters and granddaughters for the founder bull of each recessive are shown in Table 7. Most mutations were spread throughout the population because the founder sired many daughters directly, and then the founder's sons provided many granddaughters. Popular sire effects can be amplified when a bull is heavily used in a small population. For example, Selwood Betty's Commander (AYUSA000000117936), the source of the AH1 haplotype, produced 4,747 daughters and 23,964 granddaughters in a small breed (Ayrshire, with 5,405 cows on milk recording born in 2015).

It is easy to reduce the frequency of a deleterious allele in a population under selection, but is extremely difficult to eliminate it entirely from the population (e.g., Lush, 1945). Known carriers may be removed from the population, but in practice it is more common to avoid carrier-to-carrier matings because carrier

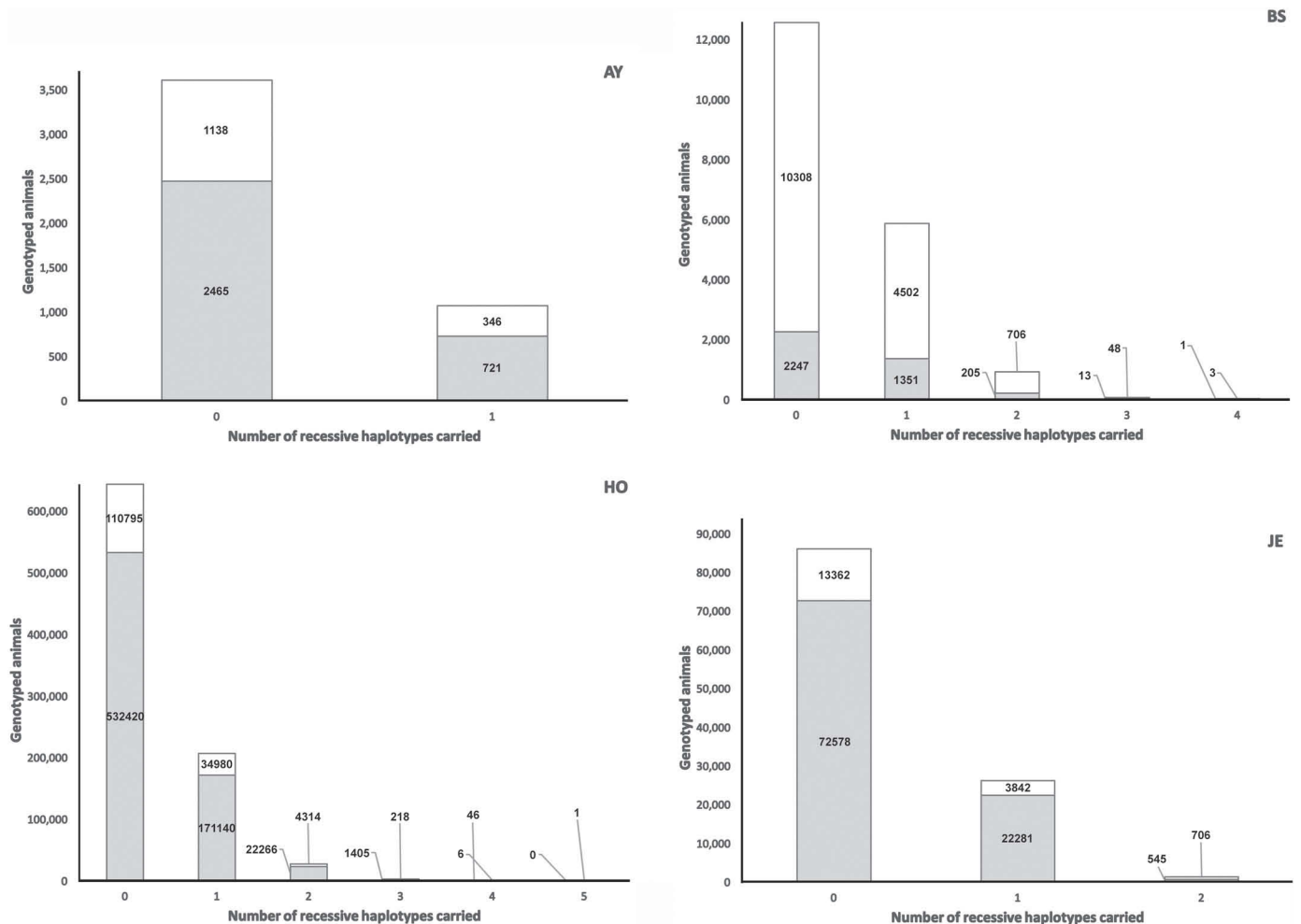


Figure 2. Number of copies of recessive haplotypes carried by genotyped bulls (white bars) and cows (gray bars) of the Ayrshire (AY), Brown Swiss (BS), Holstein (HO), and Jersey (JE) breeds in the US national dairy database. Counts of bulls (upper number) and cows (lower number) are shown for bars that are too small to appear in the figure.

Table 7. The earliest genotyped ancestor bulls for each recessive haplotype, and the number of daughters, granddaughters, sons, and grandsons in each animal's pedigree in the US national dairy database

Breed	Haplo ¹	Earliest genotyped ancestor name	Earliest genotyped ancestor identification	Birth year	Daughters	Grand-daughters	Sons	Grand-sons
AY	AH1	Selwood Betty's Commander	AYUSA000000117936	1953	4,747	23,964	30	125
BS	BH1	West Lawn Stretch Improver	BSUSA000000163153	1972	4,598	7,107	13	49
	BH2	Rancho Rustic My Design	BSUSA000000144488	1963	532	1,570	2	16
HO	BHD	White Cloud Jason's Elegant	BSUSA000000148551	1966	2,194	20,810	33	80
	BHM	Meadow View Destiny	BSUSA000000118619	1953	806	6,586	7	36
	BHW	Autumn Sun	BSDEU000803611398	1951	662	3,565	6	8
	HCD	Maughlin Storm ²	HOCAN0000005457798	1991	14,466	143,829	112	1,080
	HH0	Sweet-Haven Tradition	HOUSA000001682485	1974	17,342	227,064	573	1,446
	HH1	Pawnee Farm Arlinda Chief	HOUSA000001427381	1962	16,367	528,383	472	3,941
	HH2	Willowholme Mark Anthony	HOCAN000000334489	1975	1,312	2,258	9	20
	HH3	Gray View Skyliner	HOUSA000001244845	1954	8,653	33,000	28	97
	HH4	Besne Buck ³	HOFRA004486041658	1986	6	1,354	4	85
	HH5	Thornlea Texal Supreme	HOCAN000000264804	1957	486	7,744	11	8
JE	HHB	Osborndale Ivanhoe	HOUSA000001189870	1952	10,194	267,158	137	934
	HHC	Ideal Fury Reflector	HOUSA000001381027	1959	3,794	57,792	47	186
	HHM	Gar-Bar-Dale Burke Kate	HOUSA000001410387	1961	5,508	11,848	14	6
	JH1	Observer Chocolate Soldier	JEUSA000000596832	1962	1,458	58,841	47	251
	JH2	S.S. Quicksilver ff Fallneva	JEUSA000000593883	1960	3,754	58,047	52	213

¹The recessives analyzed include the fertility haplotypes in each breed; haplotypes for spinal muscular atrophy (BHM), spinal dysmyelination (BHD), and Weaver (BHW) in Brown Swiss; and haplotypes for bovine leukocyte adhesion deficiency (HHB), brachyspina (HH0), complex vertebral malformation (HHC), and mulefoot (syndactyly; HHM) in Holsteins (http://aipl.arsusda.gov/reference/recessive_haplotypes_ARR-G3.html).

²Maughlin Storm is the earliest genotyped carrier of the HCD mutation but may not be the founder. The original mutation may have occurred in a nongenotyped female ancestor.

³Besne Buck is the earliest genotyped carrier of the HH4 mutation but may not be the founder. The original mutation may have occurred in a nongenotyped female ancestor.

bulls may have high genetic merit for economically important traits. The results in Table 6 suggest that the complete avoidance of carrier bulls could reduce the frequencies of the recessives studies without major effects on genetic trend, but this would require coordinated action by cattle breeders that seems unlikely. Segelke et al. (2016) recently suggested that selection of cows on an index including haplotypes of interest and bulls on breeding values can be used to balance selection for (or against) specific alleles with genetic gain, and Cole (2015) has demonstrated a strategy for mate allocation that can accommodate many recessives simultaneously. Dairy farmers are unlikely to completely avoid the use of carriers, thus the inclusion of recessives in selection programs is needed to ensure that harmful allele frequencies remain low.

CONCLUSIONS

Effects of the recessive haplotypes on other traits studied were generally small even when significant. Almost \$11 million of economic losses due to reduced fertility and perinatal calf death could be avoided by

selecting mate pairs that will not produce affected embryos. Carrier animals may continue to be selected if the merit of their favorable alleles exceeds the loss from their recessive alleles, but carrier bulls can be generally avoided without reducing the average genetic merit of the sires available for mating.

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