

Simulation of introgression of the *POLLED* allele into the Jersey breed via conventional breeding vs. gene editing¹

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INTRODUCTION

Dehorning, a standard cattle management practice to protect animals and humans from injury, is unpleasant, costly, and subject to public scrutiny (Stafford and Mellor, 2011). In the United States, 94% of dairy cattle producers report routine dehorning (USDA, 2009). Horns are recessively inherited; an alternative is to breed for polled (Long and Gregory, 1978).

The frequency of the *POLLED* allele is very low in U.S. dairy cattle (<0.01). Therefore, adding its economic value to the lifetime net merit index (NMS) does not effectively increase *POLLED* (Cole, 2015). Gene editing to produce high-genetic merit, polled bulls has been proposed as an efficient method to eliminate dehorning (Carlson et al., 2016).

The use of gene editing to eliminate dehorning in Holsteins was previously simulated (Mueller et al., 2018). Dehorning is also an issue in Jerseys, the second largest U.S. dairy breed. The *HORNED* allele frequency is 1.5% lower in Jerseys compared

with Holsteins (Null, 2015), and a higher proportion of polled sires are available (Spurlock et al., 2014).

This simulation tested the hypothesis that gene editing is more efficient than conventional breeding for eliminating *HORNED* from the Jersey population. The objective was to model the incorporation of *POLLED* into the U.S. Jersey population using either conventional breeding or gene editing for three polled-mating schemes and quantify changes in *HORNED* frequency, inbreeding, and rates of genetic gain.

MATERIALS AND METHODS

Simulation

Geneedit.py (Cole, 2017) simulated introgression of *POLLED* into the Jersey breed via conventional breeding or gene editing. The base population was 35,000 cows distributed over 200 herds and 350 bulls. True breeding values for NMS were determined by randomly sampling from a normal distribution, with a mean of \$0 for cows and \$300 for bulls and SD of \$200 for both. The proportion of polled bulls was set to 5.4% heterozygous (**Pp**) and 1.5% homozygous (**PP**). These bulls averaged 0.5 and 1.3 SD lower NMS, respectively, than horned bulls (NAAB, 2018). The frequency of *HORNED* was set to 0.978 (Null,

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Table 1. Parameters and results of each scenario

Mating scheme	Scenario	Gene editing	1° bull-selection criterion	Sire genotype(s)	Matings per year limit	<i>HORNED</i> allele frequency*	Inbreeding %*	NM\$ [‡]
A	A1	No	Polled	PP, Pp (pp [‡])	Yes	0.22 ^{a,b}	6.0 ^{a,b}	2,820 ^{a,b}
	A2	Yes				0.09 ^{a,b}	5.5 ^{a,b}	3,184 ^{a,b}
B	B1	No	Polled	PP (pp [‡])	Yes	0.59 ^{a,b}	5.2 ^{a,b}	2,936 ^{a,b}
	B2	Yes				0.01 ^{a,b}	5.7 ^a	3,337 ^{a,b}
C	C1	No	Polled	PP	No	0.01 ^b	14.4 ^{a,b}	2,636 ^{a,b}
	C2	Yes				0.01 ^b	7.7 ^a	3,216 ^{a,b}
D	D	No	NM\$	n/a	Yes	0.99	7.2	3,446

*Average values at year 20 of simulation.

[‡]Lifetime NM\$.

[‡]If not enough polled sires available for mating scheme, horned sires were used.

^{||}In scenario D, both PP and Pp sires may have been available, but genotype was not included as a selection criterion.

^aSignificant difference ($P \leq 0.01$) between scenarios within a mating scheme.

^bSignificant difference ($P \leq 0.01$) between scenario and baseline (D).

2015). Horned status was determined by randomly selecting one allele each from the sire and dam. The population was limited to 500 bulls and 100,000 cows total over 20 yr, with overlapping generations (Cole, 2015; Mueller et al., 2018).

Mate Allocation

Four mating schemes were modeled (A–D, Table 1). To establish a baseline, NM\$ was the only sire selection criterion in scheme D. Polled sires were preferentially selected in mating schemes A–C. Both PP and Pp bulls were used in scheme A. Only PP bulls were used in schemes B and C. In schemes A and B, bulls were limited to 5,000 matings per year; if too few polled bulls were available, horned bulls were used. Matings per bull were not limited in scheme C, so no horned sires were used. All schemes used the modified Pryce scheme to allocate bulls to cows, which penalizes the parent average NM\$ for inbreeding (Pryce et al., 2012) and the economic costs of horned and carrier calves (Cole, 2015; Mueller et al., 2018). The average cost of dehorning is \$22.50 per animal (Thompson et al., 2017). To account for breeder preferences and marketing opportunities, horned and carrier calves were penalized \$40 and \$20, respectively (Cole, 2015; Mueller et al., 2018).

Gene Editing

Gene editing was modeled as an added step to the Kasinathan (2015) production system, which uses advanced reproductive technologies and somatic cell nuclear transfer cloning. In gene-editing schemes, designated with a number 2 in the “scenario” column of Table 1, the live bulls were sorted on NM\$ in descending order, and the top

1% of Pp and pp bulls were cloned and edited. All edited bulls were assumed to be PP. Schemes designated with a number 1 in the “scenario” column of Table 1, and the baseline scenario (D), used only conventional breeding.

Analysis

Ten replicates of each scenario were compared using the Student *t*-test. *P* values of ≤ 0.01 were considered significantly different.

RESULTS AND DISCUSSION

HORNED Allele Frequency

Baseline results (D) support findings by Cole (2015) and Mueller et al., (2018) that including a horned economic penalty (\$40) in the selection index is not effective at decreasing the frequency of *HORNED* (Figure 1A).

Similar to Holstein, gene editing decreased *HORNED* frequency as fast, or faster, than conventional breeding in each scheme. Scenarios B1 and B2 exhibited the largest difference (0.58) in *HORNED* frequency within a mating scheme after 20 yr. Only a small number ($n = 8$) of PP sires were available in the base population. Since there was also a mating limit, several horned sires were used in B1. Therefore, *HORNED* frequency did not decrease as quickly as in other scenarios. In contrast, using gene editing in B2 rapidly increased the number of high NM\$ PP sires, so fewer horned sires were used each year. As a result, *HORNED* frequency in B2 decreased significantly faster ($P \leq 0.01$) than B1. Since only PP sires were used in scheme C, both conventional breeding

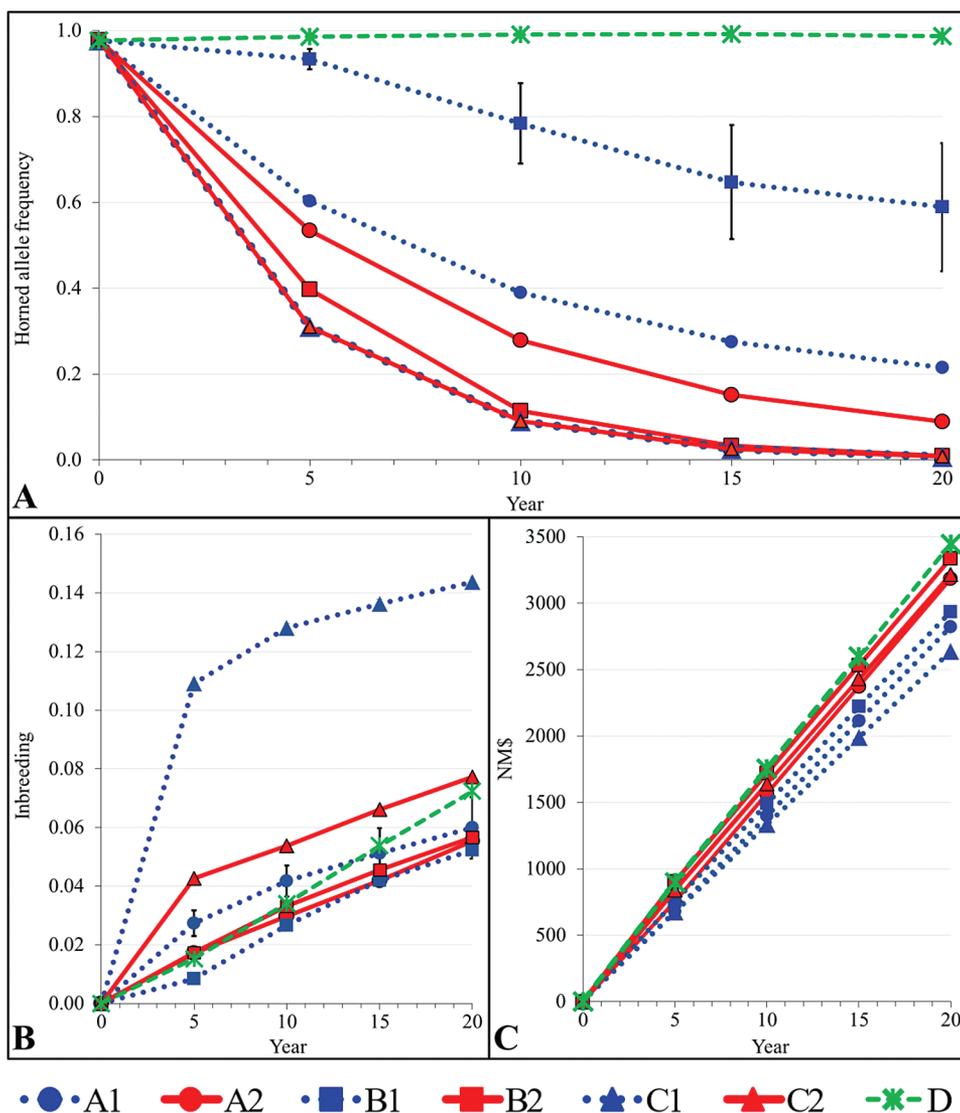


Figure 1. Effect of each mating scenario on (A) *HORNED* allele frequency, (B) inbreeding, and (C) NMS. Conventional breeding is a dashed green line, conventional breeding for polled scenarios are dotted blue lines, and gene editing for polled scenarios are solid red lines. Bars are used to represent the SEM.

and gene editing resulted in the same ($P = 0.16$) rapid change in *HORNED* frequency. Results for scheme A were intermediate to B and C.

Inbreeding

In scenario D, inbreeding reached 7% after 20 yr. Polled schemes A and B showed lower inbreeding than D (A1, A2, B1 ($P \leq 0.01$); B2 ($P = 0.28$)) (Figure 1B). In contrast, selecting for polled in Holstein resulted in higher inbreeding vs. D in all but B1 (Mueller et al., 2018). The polled-mating schemes used an additional selection criterion in the breeding objective, so herds used a wider variety of sires, resulting in lower inbreeding levels than D. However, this simulation assumed that all base population animals were initially unrelated, which is

unlikely in a production population. Since a mixture of sire genotypes were used in schemes A and B, inbreeding reached 6% regardless of the introgression method after 20 yr. When herds were forced to use only PP sires, inbreeding was significantly higher (14%, $P \leq 0.01$) than D for the conventional breeding scheme (C1) but was not significantly different (8%, $P = 0.09$) for the gene-editing scenario (C2).

Genetic Gain

Polled-mating schemes resulted in significantly slower ($P \leq 0.01$) rates of genetic gain (NMS) vs. D (Figure 1C), which is consistent with previous findings (Spurlock et al., 2014; Mueller et al., 2018). In all polled schemes, gene editing resulted in significantly greater ($P \leq 0.01$) genetic gain than conventional

breeding. The greatest difference in NMS within a mating scheme after 20 yr, \$580, was observed between C1 and C2. Although only PP sires were also used in C2, gene editing allowed for the top NMS bulls to be PP in just a few years. The greatest average rate of genetic gain per year, \$167, was achieved with gene editing in B2, when herds preferentially used PP sires, but could also use pp sires.

Consistent with our hypothesis, these results show that gene editing was more efficient at rapidly reducing the frequency of *HORNED*, while keeping inbreeding at acceptable levels and maintaining rates of genetic gain in Jersey. Scenario B2, which used both PP and pp sires in combination with gene editing, was the optimal scenario (Figure 1). This is consistent with the results observed in Holstein (Mueller et al., 2018). Scheme C models a case where consumer and market expectations force the dairy industry to eliminate dehorning immediately, thereby requiring the exclusive use of PP sires.

IMPLICATIONS

Our simulations show, given the current NMS of dairy sires, conventional breeding to decrease *HORNED* frequency will increase inbreeding and slow the rate of genetic gain (NMS). Resulting economic considerations hinder the dairy industry's ability to address this animal welfare concern through currently available approaches. Although long-term progress can be made through conventional breeding, the negative impact on inbreeding and NMS is greater than if gene editing was used. If consumers demand an immediate end to dehorning, producers may have limited time to change their practices. In this case, gene editing will be necessary to avoid long-term detrimental effects to the U.S. dairy industry. This study demonstrates how gene editing to produce high-genetic merit-polled sires could relieve dairy producers' economic concerns while also alleviating consumers' animal welfare concerns.

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