

Implementation of advanced Optimum Contribution Selection in small-scale breeding schemes: prospects and challenges in Vorderwald cattle

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Vorderwald cattle are a regional cattle breed from the Black Forest in south western Germany. In recent decades, commercial breeds have been introgressed to upgrade the breed in performance traits. On one hand, native genetic diversity of the breed should be conserved. On the other hand, moderate rates of genetic gain are needed to satisfy breeders to keep the breed. These goals are antagonistic, since the native proportion of the gene pool is negatively correlated to performance traits and the carriers of introgressed alleles are less related to the population. Thus, a standard Optimum Contribution Selection (OCS) approach would lead to reinforced selection on migrant contributions (MC). Our objective was the development of strategies for practical implementation of an OCS approach to manage the MC and native genetic diversity of regional breeds. Additionally, we examined the organisational efforts and the financial impacts on the breeding scheme of Vorderwald cattle. We chose the advanced Optimum Contribution Selection (aOCS) to manage the breed in stochastic simulations based on real pedigree data. In addition to standard OCS approaches, aOCS facilitates the management of the MC and the rate of inbreeding at native alleles. We examined two aOCS strategies. Both strategies maximised genetic gain, while strategy (I) conserved the MC in the breeding population and strategy (II) reduced the MC at a predefined annual rate. These two approaches were combined with one of three flows of replacement of sires (FoR strategies). Additionally, we compared breeding costs to clarify about the financial impact of implementing aOCS in a young sire breeding scheme. According to our results, conserving the MC in the population led to significantly ($P < 0.01$) higher genetic gain (1.16 ± 0.13 points/year) than reducing the MC (0.88 ± 0.10 points/year). In simulation scenarios that conserved the MC, the final value of MC was $57.6\% \pm 0.004$, while being constraint to 58.2% . However, reducing the MC is only partially feasible based on pedigree data. Additionally, this study proves that the classical rate of inbreeding can be managed by constraining only the rate of inbreeding at native alleles within the aOCS approach. The financial comparison of the different breeding schemes proved the feasibility of implementing aOCS in Vorderwald cattle. Implementing the modelled breeding scheme would reduce costs by 1.1% compared with the actual scheme. Reduced costs were underpinned by additional genetic gain in superior simulation scenarios compared to expected genetic gain in reality (+4.85%).

Keywords: migrant contribution, native contribution, native kinship, breeding costs, regional breed

Implications

Vorderwald cattle are a regional cattle breed that has been crossbred with commercial breeds for upgrading in milk and meat performance. This led to decreasing originality and native genetic diversity of the breed. Advanced Optimum Contribution Selection facilitates the management of those parameters and simultaneously maximises the genetic gain. However, the implementation of advanced Optimum

Contribution Selection is impeded by practical limitations. Thus, we developed solutions to practically implement advanced Optimum Contribution Selection in small-scale breeding schemes of regional breeds. Eventually, this could contribute to the preservation of the agrobiodiversity.

Introduction

Vorderwald cattle are a livestock breed from the Black Forest in south western Germany. Despite the moderately low performance of the breed, it is well adapted to the local landscape, has a cultural value and contributes to the agricultural

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biodiversity. Over the past 50 years, the breeding program focused on performance traits and ignored the value of maintaining the genetic originality of the breed. This led to increased introgression from commercial breeds for upgrading (Hartwig *et al.*, 2014). Introgressed alleles replaced native gene variants and lowered the probability that alleles at a locus are identical by descent. Thus, genetic gain (ΔG) was achieved at a low rate of inbreeding (Δf) (Kohl *et al.*, 2019) but at the expense of decreasing genetic originality. The loss of genetic diversity can be measured in two different ways using: (I) the effective population size (N_e), which is the number of individuals in an idealised population under random mating that would cause the same amount of decrease in genetic diversity as the population under study and (II) the native effective population size ($N_{e(nat)}$), which is the corresponding parameter that quantifies how fast the genetic diversity at native alleles is decreasing (Wellmann *et al.*, 2012). Vorderwald cattle actually have an N_e of 102 but an $N_{e(nat)}$ of only 34. This means that genetic diversity is high and the breed is vital in the long term, but only little of its genetic diversity is explained by native gene variants (Hartwig *et al.*, 2014). Huge amounts of irreplaceable genetic resources have passed genetic bottlenecks. Genetic bottlenecks for the genetic diversity at native alleles are caused by several factors, which include (I) direct displacement of native alleles by introgressed alleles and (II) directional selection for performance traits within the mixture of native and introgressed alleles. Today, performance traits are drastically improved in commercial breeds, and consumer demands on animal welfare are rapidly growing. This leads to changes in livestock breeding with traits like robustness, fitness and fertility being increasingly focused. Selection for ΔG led to effective population sizes as low as 52 in commercial breeds like German Holstein (Koenig and Simianer, 2006). Thus, alleles with large effects on new traits have already passed genetic bottlenecks in commercial breeds (Woolliams *et al.*, 2015). This will impede the selection for new traits. The additional genetic diversity that can be found in local breeds like Vorderwald cattle is mandatory to safeguard future changes in livestock breeding. As commercial breeds like Red Holstein and Montbéliard have already been introgressed to Vorderwald cattle (Hartwig *et al.*, 2014), there is a need for the management of introgressed genetic material. Optimum Contribution Selection (OCS) has been the standard to maximise ΔG and simultaneously manage Δf (Meuwissen, 1997). However, in breeds with historical introgression for upgrading reasons, OCS would lead to reinforced selection on introgressed alleles, since these are rare in the population and positively correlated to ΔG (Wellmann *et al.*, 2012). Displacement breeding would be the consequence. This is why Wellmann *et al.* (2012) invented an advanced Optimum Contribution Selection (aOCS) method, which facilitates the simultaneous management of ΔG , Δf , the rate of inbreeding at native alleles ($\Delta f_{(nat)}$) (Wellmann *et al.*, 2012) and the amount of introgressed genetic material, that is, migrant contributions (MC). The MC is the portion of alleles descending from other

breeds. The MC calculated from pedigree data is the expected percentage of introgressed alleles carried by an individual. On a population level, the MC already exceeds 60% in Vorderwald cattle. Therefore, starting the management of MCs is imperative. This study continues the investigation of Kohl *et al.* (2019). In that preceding article, a deeper insight in the underlying simulation process with explanations of the iterative phenotyping and breeding value estimation of the population under study is given. We examined opportunities to manage the MC with aOCS in Vorderwald cattle until marker data are available. In Germany, most regional breeds cannot afford the costs of genotyping. Additionally, the populations are frequently too small to build proper reference populations. Breeds with related gene pools could extend the reference populations with additional genotype data. However, genotyping and development of methodologies is outstanding. Hence, the genomic management of regional breeds is not widespread. Many regional breeds like Vorderwalder, Hinterwalder, German Angler and German Gelbvieh have to deal with historic migration for upgrading in performance traits (Bennewitz and Meuwissen, 2005). Thus, a pedigree-based aOCS approach is of great interest to manage the native genetic diversity and originality of those breeds till marker data are affordable and methodologies are developed. The objectives of this study were (I) the development of strategies for practical implementation of an aOCS approach to manage the MC and $\Delta f_{(nat)}$ in breeding schemes of regional cattle breeds and (II) to examine the financial effects of implementing aOCS in connection with the modelled breeding scheme of Kohl *et al.* (2019) in the Vorderwald cattle breed. On one hand, conservation or reduction of the MC is needed. On the other hand, moderate rates of ΔG are mandatory to satisfy breeders to keep the breed. This is why we tested two different aOCS strategies: (I) keeping the MC at its current value while maximising ΔG at a given $\Delta f_{(nat)}$ and (II) gradual displacement of the MC by native genetic material with an annual rate of -0.35% while maximising ΔG at a given $\Delta f_{(nat)}$. Introduction of an aOCS approach to an existing breeding scheme requires modifications. Especially in regional breeds like Vorderwald cattle, where natural service is an important reproduction technique (50%). This is why we used ZPLAN+ (Täubert *et al.*, 2010) to examine the financial effects of the modelled breeding scheme compared with the actual scheme of Vorderwald cattle.

Material and methods

The aim of the stochastic simulation was to reflect reality and predict future developments of the breed by using aOCS under practical conditions. Since sufficient marker data are not yet available, and will not be in the foreseeable future, the simulations rely on pedigree data. A detailed description of the simulation protocol and the breeding scheme can be found in Kohl *et al.* (2019). The simulation process was iterated 21 times, whereby each iteration equalled 1 year.

This reflects four generations with overlapping generations, on basis of a generation interval of 5.41 (Hartwig *et al.*, 2013). This timeframe was chosen, because we assume that marker data should be available afterwards. Further opportunities of aOCS by utilising marker data have already been examined by Wang *et al.* (2017).

Base population

The raw data set, provided by the State Agency for Spatial Information and Rural Development Baden-Württemberg, consisted of 354 451 individuals with information on sex, breed, date of birth and estimated breeding values (EBVs) for the total merit index (TMI). Vorderwald cattle are a dual-purpose breed. The actual TMI is composed of performance traits in 44%, 44% and 12% for milk yield, fitness and meat production, respectively. Thus, we assumed a breed-specific TMI based on an individual's own performance, equivalent to a simulated trait with $h^2 = 0.25$ for our simulations (Kohl *et al.*, 2019). In Germany, EBVs are standardised with mean 100 and a genetic SD of 12. The iterative estimation of breeding values within the simulation process was modelled to reflect this variation and enable changes in EBVs due to increasing accuracies (r) over time. We processed the raw data set for individuals born before 2012 with at least three equivalent complete generations (MacCluer *et al.*, 1983). Pruning the raw data set for these individuals resulted in a pedigree of 89 911 individuals born between 1938 and 2012. Individuals born before 1970 and with unknown pedigree were defined as native founders and had an MC of 0%. Individuals from foreign breeds and individuals with unknown pedigree born after 1970 were defined as migrant founders with an MC of 100%. This assumption is based on the results of Hartwig *et al.* (2014) who found that severe introgression of foreign breeds to Vorderwald cattle started in 1970. After defining native and migrant founders, the MC was calculated for all individuals in the pedigree. Birth cohorts 2004 to 2012 consisted of 3 372 individuals on average. Therefore, the simulation process was designed to create 3 372 progeny per year, assuming that the population size remains constant.

Selection candidates

Each year of the simulation started with sampling of deceased individuals (Kohl *et al.*, 2019). Selection candidates were labelled as purebred in the herd book of Vorderwald cattle. As aOCS was applied for overlapping generations, we defined concrete birth cohorts which spanned 1 year. Birth cohort B_t consisted of living individuals born in year t . Available proven sires for broad deployment were aged between 3 and 15 years, so these belonged to birth cohorts B_{t-3} to B_{t-15} . Old sires were replaced by young sires, which completed 10 matings for progeny testing in B_{t-1} . As a result, those had accuracies of EBVs of ≈ 0.5 (Kohl *et al.*, 2019) by the time of getting available for broad deployment in B_{t-3} . The flow of replacement of sires (FoR strategy) has a significant impact on ΔG when aOCS is implemented (Kohl *et al.*, 2019). This is why we tested three different strategies: 10, 20

or 30 young sires for annual restock (FoR10, FoR20 and FoR30 strategies, respectively). Living dams belonging to birth cohorts B_{t-1} to B_{t-9} were available for breeding.

Advanced Optimum Contribution Selection strategies

The result of OCS approaches for overlapping generations is a vector \mathbf{c}_t with the desired genetic contributions c_{it} of each individual i to the next birth cohort $t+1$. The genetic contribution of each individual i cannot be negative, that is, $c_{it} \geq 0$. This is a general constraint for all OCS approaches. As a second constraint, the total genetic contributions of each sex equal 0.5, since the genes of diploid species originate 50% from sires and 50% from dams, that is, $\mathbf{c}_t' \mathbf{s}_t = 0.5$ and $\mathbf{c}_t' \mathbf{d}_t = 0.5$, where \mathbf{s}_t and \mathbf{d}_t are the indicative vectors of sex (0/1). Due to the limited number of progeny per cow and year, female contributions were forced to be equal, that is, $c_{ti} = c_{tj}$ for all females i, j . Advanced OCS was only applied to the bull path. Since we face a breeding scheme with a substantial amount of natural service ($\approx 50\%$), the maximum genetic contribution per male candidate was forced to not exceed 0.05, that is, $c_{ti} \leq 0.05$ for all males i . As a result, aOCS selected at least 10 sires per year ($\mathbf{c}_t' \mathbf{s}_t / 10 = 0.05$) to satisfy the population under natural service. According to the absolute number of descendants per birth cohort, a single sire could service a maximum of 337 females per year ($3372/10 \sim 337$). Maximisation of ΔG can be achieved by maximising $\mathbf{c}_t' \mathbf{EBV}$, where \mathbf{EBV} is a vector of EBVs for TMI of the selection candidates. The maximisation of $\mathbf{c}_t' \mathbf{EBV}$ is done under constraints. Effective population sizes of 50 to 100 have been suggested in the literature to keep a breed vital in the long term (Meuwissen, 2009). Thus, we chose an $N_{e(nat)}$ of 100 to be on the safe side. As an artefact of introgression, the classical kinship (*classKin*) is smaller than the native kinship (*natKin*). Restricting the *natKin* by an upper bound will automatically restrict the *classKin* and will manage both parameters simultaneously (Kohl *et al.*, 2019). We calculated the desired value for the annual increase in native kinship as

$$\Delta f_{nat} = \frac{1}{2 \times N_{e(nat)} \times L} = 0.092,$$

where L is the generation interval of 5.41. The first constraint of the aOCS procedure poses as an upper bound for *natKin* in the population at time $t+1$:

$$ub.natKin_{t+1} = \emptyset natKin_{2012} + (t - 2011) \times \Delta f_{nat},$$

where $\emptyset natKin_{2012}$ is the average native kinship of the population in 2012. The second constraint poses as an upper bound for the MC in the population at time $t+1$:

$$ub.MC_{t+1} = MC_{2012} - (t - 2011) \times \Delta MC,$$

where MC_{2012} is the average MC of 58.2% of the base population in 2012 and ΔMC is the desired annual rate of decrease in MC. Our aim was to examine the possibilities

of reducing the MC and the arising impact on ΔG . We tested two different *aOCS* strategies. Both strategies maximised ΔG . The MC was constrained in the first strategy as $\Delta MC = 0\%$ to conserve the MC (*conserve-MC* strategy), and in the second strategy as $\Delta MC = -0.35\%$ to reduce the MC annually (*reduce-MC* strategy). ΔMC of -0.35% was identified as the maximum value, for which most simulation scenarios solved the optimisation problem within the chosen time frame of four generations. Both *aOCS* strategies were combined with all *For* strategies, so six different scenarios were examined in total. Each scenario was replicated five times. The presented results are means averaged over replicates.

Genetic gain

For a better interpretation of the results, we calculated ΔG as the average annual improvement in the mean EBV of the birth cohorts between year t_1 and t_2 as:

$$\Delta G_{t_1, t_2} = \frac{\bar{\text{EBV}}_{t_2} - \bar{\text{EBV}}_{t_1}}{t_2 - t_1},$$

where $\bar{\text{EBV}}_t$ is the average EBV of birth cohort t , averaged over five replicates. To verify average value differences, we performed a two-factorial ANOVA and a Tukey honestly significant difference (HSD) test. The response variable was $\Delta G_{2012, 2033}$. The *For* strategy and the *aOCS* strategy were used as factor variables. The development of ΔG in the real population was included for comparison. The average ΔG of real data between 2005 and 2015 was 1.18 points/year. Hence, we assumed that the ΔG will evolve linearly with 1.18 points/year in reality.

Rates of inbreeding

We calculated Δf and Δf_{nat} for a time interval spanning from year t_1 to t_2 as

$$\Delta f_{t_1, t_2} = \frac{\bar{\text{classKin}}_{t_2} - \bar{\text{classKin}}_{t_1}}{t_2 - t_1}$$

and

$$\Delta f_{\text{nat}(t_1, t_2)} = \frac{\bar{\text{natKin}}_{t_2} - \bar{\text{natKin}}_{t_1}}{t_2 - t_1},$$

whereby the average was taken over five replicates.

Migrant contributions

We calculated ΔMC as the average annual change of the mean MC of the birth cohorts for the time interval from year t_1 to t_2 as:

$$\Delta MC_{t_1, t_2} = \frac{\bar{\text{MC}}_{t_2} - \bar{\text{MC}}_{t_1}}{t_2 - t_1},$$

where $\bar{\text{MC}}_t$ is the average MC of a birth cohort t , averaged over five replicates.

Number of deployed sires

Since natural service is an important reproduction technique in Vorderwald cattle, the number of annually contributing sires influences the feasibility of a future breeding scheme. This is why we calculated the average number of annually contributing sires:

$$\bar{\text{Sires}}_t = \frac{\Sigma \text{Sires}_t}{5},$$

where ΣSires_t is the sum of sires that contributed to birth cohort t , averaged over five replicates.

Breeding costs

We used the software *ZPLAN+* (Täubert *et al.*, 2010) to calculate breeding costs of the young sire breeding scheme (Kohl *et al.*, 2019) in connection with a *For23* strategy and 10 matings for progeny tests per young bull. The calculated costs of the young sire breeding scheme were compared with the costs of the actual scheme of Vorderwald cattle (Kohl *et al.*, 2019). Reasons for choosing a *For23* strategy rather than a *For10*, *For20* or *For30* strategy will be given in the discussion (see the "Transfer to a practical breeding scheme" section). *ZPLAN+* utilises a deterministic evaluation of breeding schemes. It is based on selection index theory (Hazel and Lush, 1942) and geneflow (Hill, 1974) in populations with overlapping generations. The results of *ZPLAN+* provide population parameters, discounted costs and discounted returns of a breeding scheme. However, the results of *ZPLAN+* were exclusively used to estimate breeding costs per animal and year, because population parameters and discounted returns are based on selection index theory rather than *aOCS*-based simulation results. Alternative software to evaluate various breeding schemes are *SelAction* (Rutten *et al.*, 2002) and *ADAM* (Pedersen *et al.*, 2009). The output of *SelAction* includes the response to selection measured in economic units (Rutten *et al.*, 2002). The software *ADAM* uses stochastic simulations to evaluate selective breeding schemes. However, *SelAction* and *ADAM* do not facilitate a monetary evaluation of the costs of different schemes. Hence, we chose *ZPLAN+* for our study. The costs of a breeding scheme are composed of fix and variable costs. The fix costs of the actual breeding scheme and the young sire scheme were assumed to be equal. The variable costs included: milk recording (33€/year/cow), selection of bull calves (10€/calve), performance tests of bull calves (50€/bull), keeping of waiting bulls (100€/bull/year), evaluation of daughters (50€/daughter), production of straws (1€/straw) and storage of straws (0.03€/straw) (Priv. Doz. Dr. Pera Herold, personal communication, 10 May 2018). The sum of all breeding costs is covered by each individual of a breeding scheme. Thus, the total breeding costs were divided by the number of individuals (Vereinigte Informationssysteme Tierhaltung w.V., 2011).

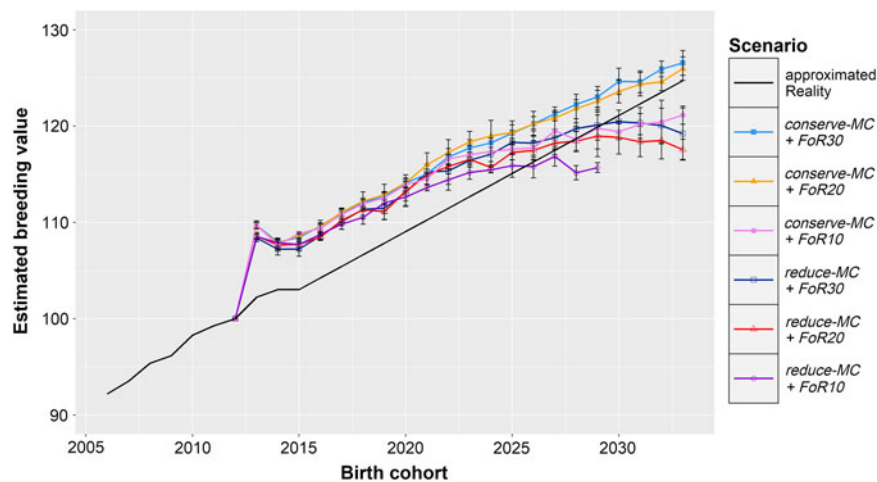


Figure 1 (colour online) Development of genetic gain of birth cohorts in different simulation scenarios and extrapolated reality for Vorderwald cattle – Development of mean estimated breeding values for the total merit index of birth cohorts in simulation scenarios and reality. Development in the real population was calculated based on the real data between 2005 and 2015. We assumed that the genetic gain will evolve linearly with an annual rate of 1.18 points/year. In simulation scenarios, we examined varying flows of replacement of sires with 10, 20 or 30 on an annual basis (*FoR10*, *FoR20* and *FoR30*, respectively) in combination with two different *aOCS* strategies. The first one conserved the mean MC in subsequent birth cohorts (*conserve-MC*). The second one reduced MC with an annual rate of -0.35% (*reduce-MC*). Results of simulation scenarios were averaged over five replicates ($\pm SD$). *aOCS* = advanced Optimum Contribution Selection; MC = migrant contribution.

Results

Genetic gain

Figure 1 shows the development of ΔG in the simulation scenarios and the calculated ΔG in reality. All simulations show a strong increase of the EBVs in the first birth cohort ($\Delta G_{2012,2013} = +9.1 \pm 0.7$), which was followed by slightly lower EBVs in the following two birth cohorts ($\Delta G_{2013,2015} = -0.51 \pm 0.32$). After 2015, the EBVs increased linearly in scenarios *conserve-MC + FoR10*, *conserve-MC + FoR20* and *conserve-MC + FoR30* with $\Delta G_{2012,2033}$ of 1.01 ± 0.05 , 1.24 ± 0.06 and 1.27 ± 0.06 , respectively. In scenarios *reduce-MC + FoR20* and *reduce-MC + FoR30*, $\Delta G_{2015,2029}$ (0.80 ± 0.14 and 0.92 ± 0.10 , respectively) was significantly higher ($P = 9 \times 10^{-10}$) than the annual increase in the following birth cohorts ($\Delta G_{2029,2033} = -0.36 \pm 0.36$ and -0.22 ± 0.32 , respectively). As a result, $\Delta G_{2012,2033}$ was 0.84 ± 0.05 in *reduce-MC + FoR20* and 0.92 ± 0.13 in *reduce-MC + FoR30*. The optimisation problem could not be solved in 2029 for scenario *reduce-MC + FoR10* because no solution existed. However, $\Delta G_{2012,2029}$ was 0.91 ± 0.05 . According to the Tukey HSD test, $\Delta G_{2012,2033}$ was significantly affected by the *aOCS* strategy and by the *FoR* strategy ($P = 2 \times 10^{-11}$ and 2×10^{-6} , respectively). Conserving the MC realised higher $\Delta G_{2012,2033}$ (1.16 ± 0.13) than reducing the MC (0.88 ± 0.1). The *FoR10* strategy produced lower $\Delta G_{2012,2033}$ (1.0 ± 0.05) than the *FoR20* and *FoR30* strategy (1.04 ± 0.22 and 1.09 ± 0.21 , respectively). The lower variation of $\Delta G_{2012,2033}$ in *FoR10* scenarios is reasoned in the missing results of scenario *reduce-MC + FoR10* in 2033. However, we performed an additional Tukey HSD test for $\Delta G_{2012,2029}$ that proved present results. Additionally, *conserve-MC + FoR20* and *conserve-MC + FoR30* produced a higher $\Delta G_{2012,2033}$ than calculated for reality. Other scenarios created less (Figure 1 and Table 1).

Migrant contributions

The development of the average MC of the birth cohorts is shown in Figure 2. In the *conserve-MC* scenarios, $\Delta MC_{2012,2013}$ was $-4.33\% \pm 0.25$. In the *reduce-MC* scenarios, $\Delta MC_{2012,2013}$ was $-6.20\% \pm 0.25$. This decline in MC was followed by an increase till 2015 in all scenarios ($\Delta MC_{2013 \text{ to } 2015} = +0.62\% \pm 0.27$). After 2015, the MC evolved linearly and eventually reached $57.6\% \pm 0.004$ and $49.3\% \pm 0.002$ for *conserve-MC* and *reduce-MC* scenarios, respectively. As a result, the corresponding $\Delta MC_{2012,2033}$ were $-0.18\% \pm 0.02$ and $-0.58\% \pm 0.01$. Overall, the *reduce-MC* scenarios reduced the MC by $8.9\% \pm 0.002$.

Classical kinship and kinship at native alleles

The realised Δf , Δf_{nat} and the upper bound on Δf_{nat} are visualised in Figure 3. Across all simulated populations, the Δf_{nat} was 0.091 ± 0.0006 per year, which was only slightly below the constraint setting of Δf_{nat} (0.092 per year). Δf followed similar trends with 0.091 ± 0.007 across all scenarios, although it was not constrained by the *aOCS* procedures. As a result, the realised $N_{e(nat)}$ and N_e were 101.4 ± 0.4 and 102.1 ± 8.7 across all simulated populations, respectively.

Selected sires

The mean number of sires that contributed to the birth cohorts is visualised in Figure 4. Different bar graphs are shown for the different *FoR* strategies. In all scenarios increasing numbers of sires contributed to the birth cohorts at the beginning of the simulation with $\bar{\theta}Sires_{2012} = 14.6 \pm 2.0$, $\bar{\theta}Sires_{2015} = 21.4 \pm 4.3$ and $\bar{\theta}Sires_{2020} = 36.5 \pm 7.2$. Subsequently, the number of contributing sires was rather constant with $\bar{\theta}Sires_{2020 \text{ to } 2033} = 46.2 \pm 3.6$ across *conserve-MC* scenarios and 29.7 ± 6.0 across *reduce-MC* scenarios.

Table 1 Different simulation scenarios are explained by a combination of implemented aOCS strategy and FoR strategy of Vorderwald cattle

Scenarios	FoR ¹	aOCS ²	n	$\Delta G_{2012 \text{ to } 2033}$ ³	Δf_{nat}	$N_{e(nat)}$	Δf	N_e	MC ₂₀₃₃
Reality	$\approx 2 \text{ to } 3^4$	TS	1	1.18 ⁵	NA ⁶	NA	NA	NA	NA
Conserve-MC + FoR10	10	Conserve-MC	5	1.01 ± 0.05 ^{AX}	0.092 ± 0.0004	100.8	0.097 ± 0.002	94.6	57.8 ± 0.18
Conserve-MC + FoR20	20	Conserve-MC	5	1.24 ± 0.06 ^{BX}	0.091 ± 0.0006	101.3	0.099 ± 0.004	93.2	57.8 ± 0.28
Conserve-MC + FoR30	30	Conserve-MC	5	1.27 ± 0.06 ^{BX}	0.091 ± 0.0005	101.5	0.093 ± 0.004	99.3	57.3 ± 0.51
Reduce-MC + FoR10	10	Reduce-MC	5	NA/0.91 ± 0.05 ^{7AY}	0.092 ± 0.0007	101.0	0.090 ± 0.002	103.1	NA/ 50.7 ± 0.29 ⁷
Reduce-MC + FoR20	20	Reduce-MC	5	0.84 ± 0.05 ^{BY}	0.091 ± 0.0007	101.8	0.083 ± 0.002	111.8	49.4 ± 0.10
Reduce-MC + FoR30	30	Reduce-MC	5	0.92 ± 0.13 ^{BY}	0.091 ± 0.0004	101.9	0.081 ± 0.001	114.0	49.2 ± 0.23

Scenarios = different scenarios are explained by a combination of FoR strategy and aOCS strategy; aOCS = advanced Optimum Contribution Selection; FoR = annual flow of replacement of sires; n = replicates per scenario; ΔG = genetic gain; Δf_{nat} = rate of native Inbreeding for overlapping generations per year. Restricted to 0.092; $N_{e(nat)}$ = native effective population size; Δf = rate of Inbreeding for overlapping generations per year. Not restricted by aOCS; N_e = effective population size; TS = truncation selection; MC = migrant contribution; MC₂₀₃₄ = Average migrant contribution of birth cohort 2033 as final value.

¹Three different FoR strategies were examined with 10, 20 or 30 young sires for restock per year.

²Two different aOCS strategies were examined. Either conserving or reducing MCs in the next birth cohort with an annual rate of 0.0% or -0.35%, respectively.

³Genetic gain was defined as improvement in mean estimated breeding values for the total merit index among birth cohorts B_{2012} to B_{2033} .

⁴According to personal communication (Dr Franz Maus, 22 February 2018).

⁵Genetic gain in reality was calculated based on the real data between 2005 and 2015.

⁶NA = Not available.

⁷Reduce-MC + FoR10 was the only simulation scenario for which the aOCS optimisation problem could not be solved in 2029. Thus, NAs relate to 2033. The given figure relates to 2029.

^{A,B,X,Y}Different superscripts label significantly different values at $P < 0.01$ in terms of FoR strategies (A v. B) or aOCS strategies (X v. Y).

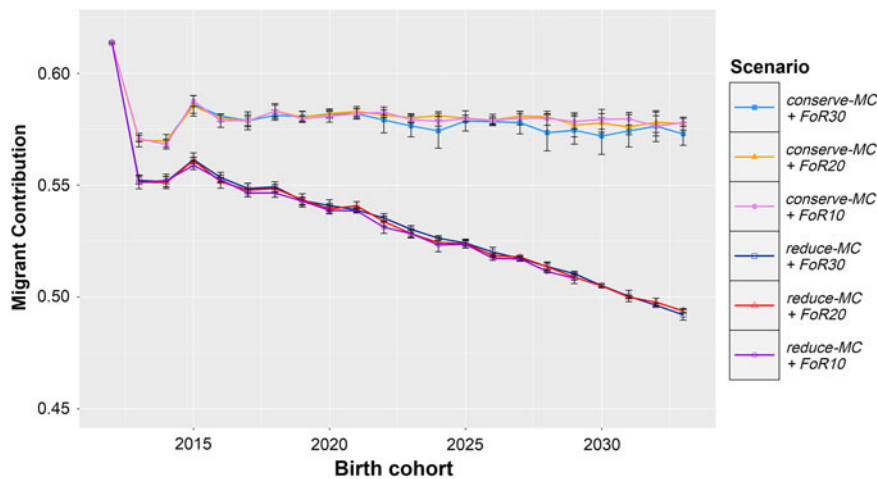


Figure 2 (colour online) Development of MCs of birth cohorts in different simulation scenarios for Vorderwald cattle – Development of mean MC of birth cohorts in simulation scenarios. We examined varying flows of replacement of sires with 10, 20 or 30 on an annual basis (FoR10, FoR20 and FoR30, respectively) in combination with two different aOCS strategies. The first one conserved the mean MC in subsequent birth cohorts (conserve-MC). The second one reduced MC with an annual rate of -0.35% (reduce-MC). Results of simulation scenarios were averaged over five replicates ($\pm 5D$). MC = migrant contribution; aOCS = advanced Optimum Contribution Selection.

Breeding costs

We give a brief overview of costs of the young sire scheme in combination with the FoR23 strategy. Costs are compared to the actual scheme to give a better understanding of the financial opportunities. According to the results of ZPLAN+, costs sum up to 18.44€/breeding animal/year and 18.23€/breeding animal/year for the actual breeding scheme and the young sire scheme, respectively. Thus, costs could be decreased by 1.1% by implementing aOCS in connection with the young sire scheme.

Discussion

Comparison of scenarios

The first two years belong to Stage 1 of the simulation process of Kohl *et al.* (2019) in which no new male

selection candidates became available. The EBVs increased strongly in all scenarios in the first birth cohort ($\Delta G_{2012,2013} = +9.1 \pm 0.7$; Figure 1), while the MC decreased ($\Delta MC_{2012,2013} = -4.33\% \pm 0.25$ and $-6.20\% \pm 0.25$; Figure 2). This reflects the huge short-term optimisation opportunities of aOCS in breeding schemes that formerly used truncation selection (Kohl *et al.*, 2019). Probably, the MC decreased because the most influential sires had by chance an MC that was slightly below average. In the following years, the EBVs of the birth cohorts were slightly smaller ($\Delta G_{2013,2015} = -0.51 \pm 0.32$) because keeping the desired level of the native kinship in the population required the use of less superior sires. These sires were chosen from the same pool as in 2012 because until 2015 no new male

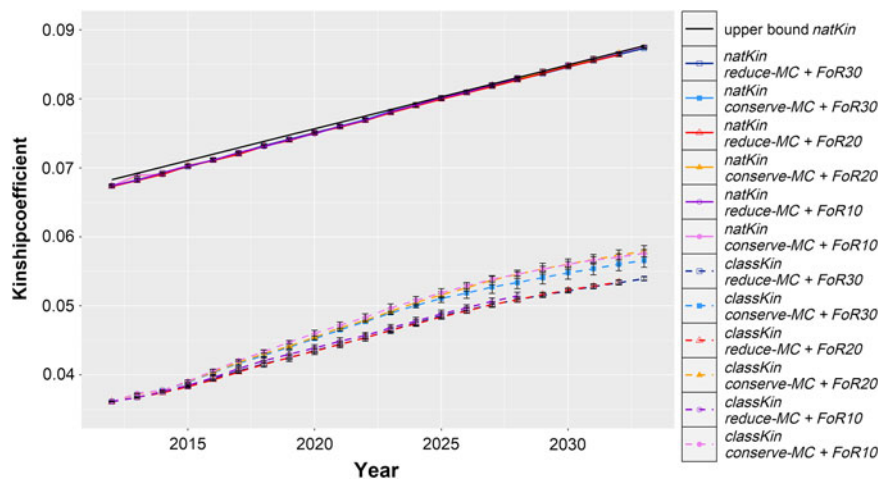


Figure 3 (colour online) Development of average classical and native kinship coefficients of evolving populations in different simulation scenarios for Vorderwald cattle – We examined varying flows of replacement of sires with 10, 20 or 30 on an annual basis (*FoR10*, *FoR20* and *FoR30*, respectively) in combination with two different *aOCS* strategies. The first one conserved the mean MC in subsequent birth cohorts (*conserve-MC*). The second one reduced the MC with an annual rate of -0.35% (*reduce-MC*). Both *aOCS* strategies restricted the average kinship at native alleles (*natKin*) of the population by an upper bound (black and solid), meanwhile the classical kinship (*classKin*) was not managed. The graphs are subdivided for different simulation scenarios (colour) and both kinship coefficients (solid and dashed). Results of simulation scenarios were averaged over five replicates ($\pm 5D$). *aOCS* = advanced Optimum Contribution Selection; MC = migrant contribution.

candidates were introduced. The number of contributing sires indeed increased from 2012 to 2015 ($\bar{\sigma}Sires_{2012} = 14.6 \pm 2.0$, whereas $\bar{\sigma}Sires_{2015} = 21.4 \pm 4.3$; Figure 4). The MCs increased slightly between 2013 and 2015 but were kept below the constraint setting ($\Delta MC_{2013,2015} = +0.62\% \pm 0.27$ in all scenarios). In 2015, first young sires got available for broad deployment, so the introduction of new proven sires finished the impediment of selection. Those were less related to the population and had on average higher EBVs compared to their older companions (results not shown). As expected, the EBVs increased linearly and the MCs were conserved in scenario *conserve-MC* from birth cohort 2015 onwards. The numbers of contributing sires further increased until ~ 2020 across all scenarios ($\bar{\sigma}Sires_{2015} = 21.4 \pm 4.3$ towards $\bar{\sigma}Sires_{2020} = 36.5 \pm 7.2$). Over the course of the simulation process, $\Delta MC_{2012,2033}$ were -0.58 ± 0.01 and $-0.18\% \pm 0.02$ for *reduce-MC* and *conserve-MC*, respectively, while the constraint setting for ΔMC were -0.35% and 0.0% , respectively. Thus, all scenarios reduced the MC at a higher rate than desired. This was an artefact of the random decrease of the MC in 2013.

All scenarios that selected for reduced MC reached a selection plateau for the EBVs around 2023, which was after the second generation was born. The reason is that pedigree data were used for estimating the MC. When pedigree data are used, then the MC of an individual is always the average of the MCs of its parents. Continued averaging of these values in newborn individuals narrows the range of the MCs in the population, and the mean of the MCs converges to a value which is above the minimum MC that was present in a selection candidate in 2012. Hence, the number of bulls whose MC surpasses the threshold value for MC is strongly decreasing after a few years. This reduces the number of males that can be used for breeding, which in turn reduces

strongly the achievable selection intensity on the EBV for TMI. After some years of management, the OCS problem can no longer be solved. The optimisation problem could not be solved for scenario *reduce-MC + FoR10* in 2029 because not enough sires with low MC existed that had a sufficiently low native kinship with the population. The situation was slightly better in scenarios *FoR20* and *FoR30* because sires could be selected from a larger pool. The number of sires to select from is 130, 260 and 390 in *FoR10*, *FoR20* and *FoR30* scenarios, respectively. This shows clearly that pedigree data cannot be used longer than two generations to reduce the MC of a population. We assumed a breed-specific heritability for the TMI of $h^2 = 0.25$ (Kohl *et al.*, 2019) to run the stochastic simulations. However, Gandini *et al.* (2014) reported that the ΔG will increase as heritabilities of traits increase when an OCS approach is implemented. Thus, the ΔG that will be realised by implementing *aOCS* in reality might deviate from the observed results. However, the utilised $h^2 = 0.25$ is realistic with respect to the composition of the actual TMI of Vorderwald cattle.

Deciding for superior strategies

According to the results of the Tukey HSD tests, ΔG was affected significantly by *aOCS* strategies ($P = 2 \times 10^{-11}$) as well as *FoR* strategies ($P = 2 \times 10^{-6}$). The *conserve-MC* scenarios resulted in a higher ΔG than the *reduce-MC* scenarios. The total selection intensity was the same in both scenarios, because the population sizes and the increases in the native kinship were equal (Figure 3). However, in the *conserve-MC* scenarios, selection was only on EBVs for TMI, whereas in the *reduce-MC* scenarios, selection was on both the EBVs and the MCs. The constraint for the MC became difficult to satisfy after a few years of selection because pedigree data were used to estimate the MC. Consequently, more and more of

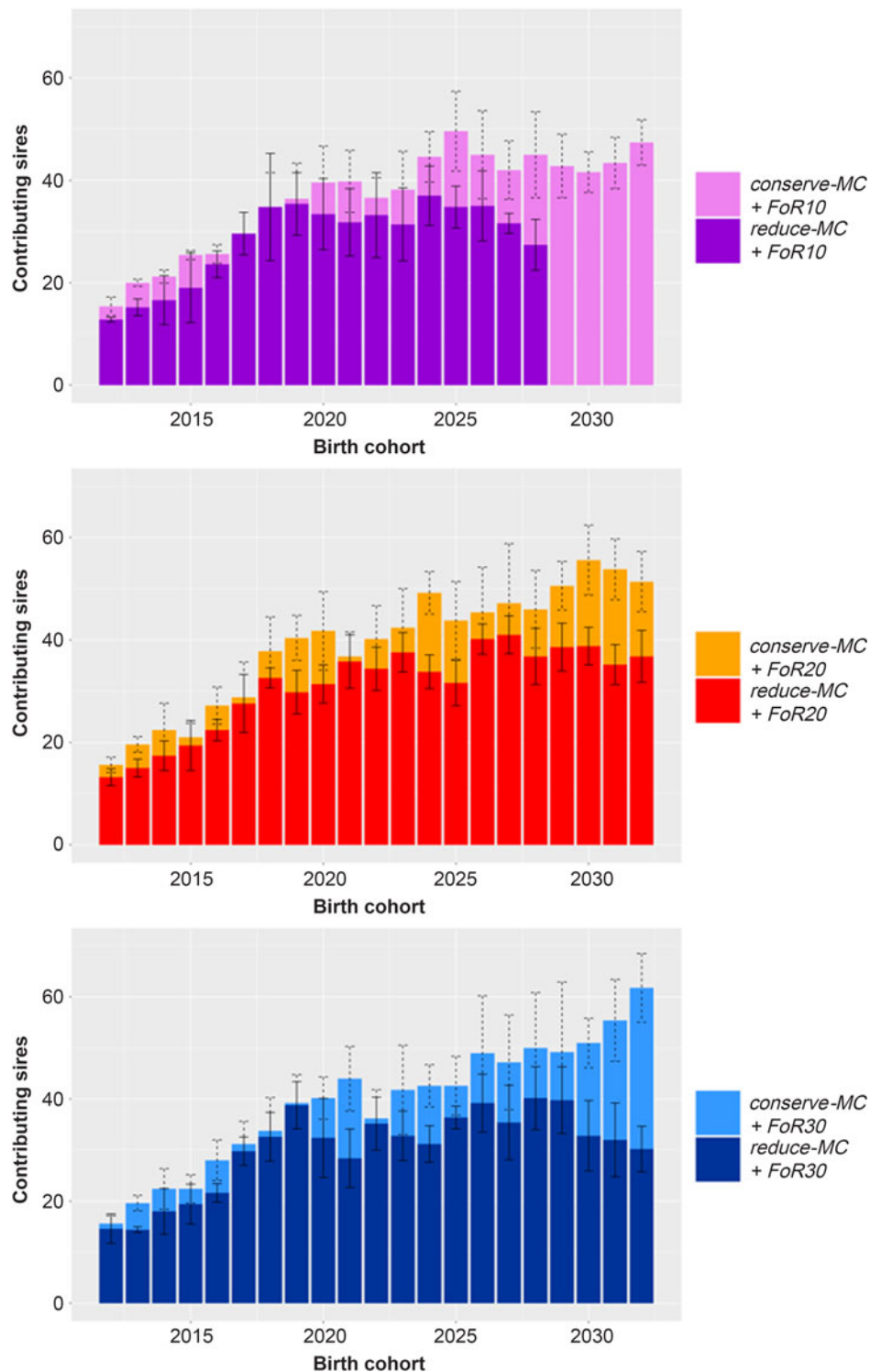


Figure 4 (colour online) Bar graph of annually contributing sires in different simulation scenarios for Vorderwald cattle – We examined varying flows of replacement of sires with 10, 20 or 30 on an annual basis (*FoR10*, *FoR20* and *FoR30*, respectively) in combination with two different *aOCS* strategies. The first one conserved the mean MC in subsequent birth cohorts (*conserve-MC*). The second one reduced MC with an annual rate of -0.35% (*reduce-MC*). The bar graphs are subdivided for *FoR* strategies. Bars visualise the average of annually contributing sires averaged over five replicates ($\pm SD$). *aOCS* = advanced Optimum Contribution Selection; MC = migrant contribution.

the selection intensity was allocated to reduce the MC and less of it was allocated to achieve genetic gain. Consequently, genetic gain was lower in the *reduce-MC* scenarios than in *conserve-MC* scenarios and eventually

approached zero. Subsidies for breeders for conserving the breed instead of keeping commercial breeds are not sufficient to cover the expenses (Kohl and Herold, 2017). Thus, further economical disadvantages through a lower ΔG must be

avoided. In a breeding program that reduces the MC, pedigree data must be replaced by genomic data after no more than two generations of selection because otherwise, the EBVs would soon reach a selection plateau. This reduced genetic gain could lead to a growing probability of extinction for the Vorderwald breed (Meuwissen, 2009). Conclusively, we recommend to implement the *conserve-MC* strategy until genotype data are available to reduce the MC. Genomic data enhance the aOCS approach by estimation of MC based on shared haplotype segments (runs of homozygosity) (Wang *et al.*, 2017). As soon as genotypes are accessible, further studies are needed to examine possibilities of reducing the MC in connection with an adjusted breeding scheme.

The *FoR20* and *FoR30* strategy resulted in a significantly higher ΔG than the *FoR10* strategy ($P = 2 \times 10^{-6}$). The highest ΔG was achieved by the *conserve-MC + FoR30* strategy but was not significantly different from the genetic gain of the *conserve-MC + FoR20* strategy. Since selection as well as husbandry and progeny testing of young sires is cost-intensive, we recommend to implement the *FoR20* strategy and conserve the MC.

Transfer to a practical breeding scheme

The actual breeding scheme of Vorderwald cattle selected 42 bull calves annually to pass a performance test on station. Subsequently, 2 to 3 of them were selected to replace sires for artificial insemination after passing a progeny test (Kohl *et al.*, 2019). According to our results, the *FoR20* strategy is sufficient to maximise ΔG when aOCS is used. However, the simulation is based on the assumption that semen will be taken from all young sires. This is unrealistic because some of them will not be available at the time of production. Based on the results of Wathes *et al.* (2008), we assume that 10% of bull calves might pass away between selection (within 1st month) and progeny testing (≈ 12 th month). Consequently, we recommend to select 22 to 23 bull calves for a performance test on station. Subsequently, all surviving young bulls should pass a progeny test and produce straws, irrespective of their performance. Thus, at least 20 young sires will be available for annual restock of proven sires. Conclusively, the superior *FoR20* strategy of simulations is transferred to a practical breeding scheme by implementing a *FoR23* strategy in reality. This is why we compared costs of the actual breeding scheme to a young sire scheme in connection with a *FoR23* strategy in *ZPLAN+*.

Furthermore, we highly recommend to accelerate the implementation of selection of elite dams. This would facilitate directed mating with sires of sires. Selection of dams should be optimised by a modified aOCS approach that additionally optimises female contributions. Consequently, bull calves could be selected not only in the broad population but also among descendants of elite matings (Kohl *et al.*, 2019). This would result in three benefits: (I) Further enhanced population parameters (Gandini *et al.*, 2014). (II) Unavailability of bull calves at the time of selection (sale and culling) could be avoided. (III) Simplified selection of bull calves, since dams are known. Currently, 3 160 dams are

serviced naturally (Kohl *et al.*, 2019). This is a further obstacle to overcome. In the actual breeding scheme, 80 sires were available for that task. According to our results, aOCS deployed 39.9 ± 1.3 sires in *conserve-MC + FoR20*. This shows that fewer sires will be available for natural mating. To meet this practical constraint, we recommend to keep young sires alive after production of straws till birth cohort B_{t-4} . Thus, those would be available for broad deployment of ≈ 2 years, i.e. parts of B_{t-2} (i.e., after progeny testing), B_{t-3} and B_{t-4} . Hence, 40 to 60 young sires (replacement rate and mortality rate) will be available for natural service. The biggest challenge of implementing optimum contributions in the naturally serviced dam population is the organisation of stock bull availability in single herds. Kohl *et al.* (2019) proposed a stock-bull-to-herd rotation program. On one hand, the accuracy of breeding value estimation could be improved through enhanced estimation of herd effects. On the other hand, this will cause organisational challenges, since sharing stock bulls will compromise the hygienic status of herds. Several approaches are conceivable to overcome that problem. The first option might be to put stock bulls under quarantine as those arrive at each herd. However, this approach has some drawbacks: Reasonable quarantine periods of mammals should last for a minimum of 30 days (Miller, 1996). At best, one stock bull could serve 11 herds per year, by complying this quarantine period. However, as aOCS is introduced to a naturally serviced dam population, there is a need for several stock bulls per herd. Thus, practicability of this approach depends on the number of herds using natural service. As a second approach, we thought of a farm cooperative solution. Thus, consistent groups of farms would share stock bulls, minimising hygienic risks of single herds since their hygienic status should converge over time. However, quarantine periods would still be necessary as stock bulls are supplied from one farm cooperative to the other. Additionally, the necessity of introducing optimum contributions in farm cooperatives by usage of several stock bulls is still a problem. Small-scale breeders might have space problems related to the need of bull paddocks to keep unused bulls till these are delivered. Furthermore, it is unclear which farms should set up the quarantine facilities and who will bear the costs. As a result, we propose two approaches, where one of these should be chosen depending on the possibilities of the breeding organisation: (I) A bull leasing system with the breeding organisation as lessor. Since genomic breeding value estimation has conquered the livestock breeding environment, most breeding organisations still have huge properties where waiting bulls have formerly been kept. Those could be used to keep stock bulls to be leased by the breeders. Thus, the hygienic status of stock bulls would be equal and could be managed professionally. (II) Epidemic units, where higher scale breeders should set up quarantine facilities to keep several stock bulls and organise stock bull delivery to connected smaller scale breeders. In both approaches, hygienic concepts have to be developed by the breeding organisation in connection with the breeders themselves. Additionally, the increasing aggressiveness of

eldering stock bulls is a well-known problem among practitioners. This is the main reason of restricting the usage of stock bulls to 4 years of age. This restriction is flexible as it is subjective and might be breed-specific. Additionally, the increasing aggressiveness of eldery stock bulls could be a new trait to be implemented in the breeding objective of Vorderwald cattle. Further studies are needed to examine the possibilities for that. Schreiner (2018) has shown that higher costs of required measures on farm distract breeders to sign conservation breeding contracts. Additionally, the payout structures of subsidies for conserving genetic agrobiodiversity at Germany are too inflexible. Breeding organisations are legal entities. Thus, those are disqualified from subsidies by the 'GAK-Förderzahlungen' (Bundesministerium für Ernährung und Landwirtschaft, 2016). The subsidies for breeders are exclusively connected to keeping a breeding animal and participation in a corresponding breeding scheme. This problem has been outspoken several times, but political actions are still missing. Required measures, that is, setting up quarantine facilities or a bull leasing system, to effectively conserve endangered livestock breeds should be subsidised separately. In reality, breeders are discouraged to invest in required measures without having a financial benefit. Thus, applicability of the modelled breeding scheme is questionable till the funding of required measures is clarified. Additionally, Schreiner (2018) mentioned that monitored pairing discourages breeders to sign conservation breeding contracts. However, the underlying aOCS approach optimises exclusively the contributions of sires to the following birth cohort. The selection in the dam path and the pairing remains a core competence of breeders. However, the pairing decisions of breeders will be compromised by the estimated optimum contributions of sires. As soon as the estimated contribution of a sire reaches the optimum, the breeding organisation should stop supplying the corresponding semen and stock bulls within the chosen time frame. In addition to the scientific work of this study, we had participative meetings about the aOCS methodology and necessary adjustments to the breeding scheme to sensitise the breeders. Marsoner *et al.* (2018) investigated the socio-ecological and cultural value of indigenous breeds. The authors reported about the maintenance of cultural landscapes, the contribution to cultural heritage and identity and the promotion of tourism through locally adopted breeds. The Vorderwald cattle breed contributes to those benefits in the region of the Schwarzwald in Germany. Thus, the lost profits of keeping Vorderwald cattle, instead of a commercial breed, were tolerated among breeders so far. Additionally, the breeders of Vorderwald cattle obtain subsidies for conserving the native genetic background of an endangered breed, although the MC of Vorderwald cattle already exceeds 60% with an $N_{e(nat)}$ of 34. Eventually, the breeders might accept the compromises on breeding decisions. The lost profits have already been accepted in the past, and the effective management of the MC is imperative to not lose the eligibility for future subsidies.

Breeding costs

ZPLAN+ was used to compare costs of the actual breeding scheme and the young sire scheme in connection with a *FoR23* strategy and 10 matings for progeny testing of each young sire. Since ZPLAN+ is based on selection index theory, population parameters were omitted. The actual breeding scheme of Vorderwald cattle accepts accuracies of EBVs of young sires depending on 25 to 30 daughters (Hartwig *et al.*, 2013). Actually, 250 to 300 matings are required for progeny tests of 2 to 3 young sires annually. According to Kohl *et al.* (2019), 14 to 15 matings per young sire are sufficient to maximise ΔG in the broad population when aOCS is used. Therefore, progeny tests of 23 young sires would result in 322 to 345 matings in total. Thus, the number of matings for progeny testing, and subsequent costs, increases by 15% to 30%. However, the *FoR23* strategy will reduce performance tests of bull calves on station (23 towards 42). The arising savings of reduced performance tests will overcompensate the expenses of increased progeny testing. As a result, implementing aOCS with the recommended superior strategies will reduce costs by 1.1%, according to ZPLAN+. Finally, a comparison of expected ΔG underpins financial benefits. The rate of ΔG of *conserve-MC + FoR20* was 4.85% higher compared to extrapolated ΔG in reality. Unfortunately, accurate values for monetary assessment of ΔG are not available for Vorderwald cattle. However, financial benefits will further expand.

Behind practical aspects

Behind practical aspects, data flow, work flow and accountable actors of the breeding scheme have to be defined. In most cases, the breeding organisation is responsible for registration of breeding animals in the stud book. At Baden-Württemberg in Germany, breeding value estimation is a state-owned task. Therefore, the breeding organisation of Vorderwald cattle supplies studbook data to governmental authorities. Following the breeding value estimation, aOCS should be conducted. Two options are conceivable here. On one hand, aOCS could be implemented at the governmental authority that drives the breeding value estimation. In this way, the breeding organisation has to supply additional data that identify the living dam population and available sires. The backflow of data would contain EBVs, optimum contributions of breeding animals and selection proposals for elite animals calculated by aOCS (dams of sires and sires of sires). As an opportunity of this option, all calculation and estimation tasks are consolidated. A drawback is the more complicated data flow and the need for communication paths, as results of the aOCS approach might seem questionable sometimes. On the other hand, governmental authorities could supply merely the EBVs to the breeding organisation (as done so far). In this way, aOCS would be conducted at the breeding organisation itself. As an opportunity of this approach, data division would remain the same and questions about results of the aOCS approach

could be scrutinised within the breeding organisation itself. Normally, all of the mentioned tasks are consolidated within the breeding organisation. Thus, the data flow, work flow and appointment of accountable actors are an internal issue of the breeding organisation.

Additional information

In this study, we used

$$ub.natKin_{t+1} = \emptyset natKin_{2012} + (t - 2011) \times \Delta f_{nat},$$

as the upper bound for *natKin* in birth cohort $t+1$. An alternative would be to use

$$ub.natKin_{t+1} = \emptyset natKin_t + (1 - \emptyset natKin_t) \times \Delta f_{nat},$$

where $\emptyset natKin_t$ is the mean native kinship of selection candidates in year t . However, this led to a bias because matings for progeny tests of young sires were not optimised by aOCS, although these contribute to $\emptyset natKin$ in year $t+1$. Thus, $\emptyset natKin$ increased at another rate than expected. This bias in $ub.natKin_{t+1}$ would accumulate over the years. As a result, $N_{e(nat)}$ would vary between 78 and 93, although it should have been constrained to 100. To be on the safe side and avoid an accumulation of errors over the years, $ub.natKin_t$ should be constrained as done in this study.

Additionally, this study proves the possibility of controlling Δf by setting an upper bound only for Δf_{nat} . Values for N_e varied within 93.2 to 114.0 across all scenarios. Despite the large variation of N_e across scenarios, the variation within scenarios was rather small (Table 1, SDs of Δf). Effective population sizes of 50 to 100 have been suggested to keep a breed vital in the long term (Meuwissen, 2009). Thus, restricting only Δf_{nat} is sufficient to manage the genetic diversity of the breed.


Conclusion

The examined scenarios and the corresponding breeding costs revealed the potential to reduce the MC with pedigree data and the potential of implementing aOCS in Vorderwald cattle in connection with a young sire breeding scheme. Replacing pedigree-based estimates of the MC by genomic estimates after no more than two generations of selection on reduced MC turned out to be mandatory for a breeding program that aims at recovering the native genetic background of a breed. From a conservational perspective, it would be desirable to implement the *reduce-MC* strategy. However, subsidies for keeping the breed are not sufficient to compensate expenses. Thus, the ΔG should not decrease in comparison to the actual population parameters. Implementation of the *conserve-MC + FoR23* strategy would create sufficient ΔG , manage Δf_{nat} at the desired level, *conserve-MC* and keep costs within an acceptable range. Unfortunately, the implementation of aOCS to an existing breeding scheme presumes investments in required

measures. The funding of those measures has to be clarified before. Additionally, the freedom of breeding decisions by breeders will be restricted by the estimated optimum contributions of sires. Eventually, the funding of required measures and the required contracts with breeders might be the biggest obstacles to overcome when implementing aOCS in a practical breeding scheme.

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Declaration of interest

None.

Ethics statement

None.

Software and data repository resources

The data and the simulation code will be archived at the State Agency for Spatial Information and Rural Development Baden-Württemberg at Kornwestheim, Germany. Access might be given after a consultation with the Rinderunion Baden-Württemberg e.V. and the Breeding Value Estimation Team Baden-Württemberg.

References

- Bennewitz J and Meuwissen TH 2005. Estimation of extinction probabilities of five German cattle breeds by population viability analysis. *Journal of Dairy Science* 88, 2949–2961.
- Bundesministerium für Ernährung und Landwirtschaft 2016. Rahmenplan der Gemeinschaftsaufgabe "Verbesserung der Agrarstruktur und des Küstenschutzes" für den Zeitraum 2016–2019. Retrieved on 23 January 2017 from <https://www.bmel.de/SharedDocs/Downloads/Landwirtschaft/Foerderung/GAK-Foerderungsgrundsaeetze/2016/Foerderbereich1-A.html>.
- Gandini G, Stella A, Del Corvo M and Jansen GB 2014. Selection with inbreeding control in simulated young bull schemes for local dairy cattle breeds. *Journal of Dairy Science* 97, 1790–1798.
- Hartwig S, Wellmann R, Hamann H and Bennewitz J 2013. Pedigreeanalysen zur Beschreibung der genetischen Variabilität bei Vorderwälder, Hinterwälder und Limpurger. *Zuchtungskunde* 85, 270–88.
- Hartwig S, Wellmann R, Hamann H and Bennewitz J 2014. The contribution of migrant breeds to the genetic gain of beef traits of German Vorderwald and Hinterwald cattle. *Journal of Animal Breeding and Genetics* 131, 496–503.
- Hazel LN and Lush JL 1942. The efficiency of three methods of selection. *Journal of Heredity* 33, 393–399.
- Hill WG 1974. Prediction and evaluation of response to selection with overlapping generations. *Animal Science* 18, 117–139.

- Koenig S and Simianer H 2006. Approaches to the management of inbreeding and relationship in the German Holstein dairy cattle population. *Livestock Science* 103, 40–53.
- Kohl S and Herold P 2017. Problemanalyse zur Implementierung der Selektion nach optimierten Genbeiträgen in kleinen Populationen. *Züchtungskunde* 89, 345–358.
- Kohl S, Wellmann R and Herold P 2019. Advanced optimum contribution selection as a tool to improve regional cattle breeds: a feasibility study for Vorderwald cattle. *Animal* 1–12. doi: [10.1017/S1751731119001484](https://doi.org/10.1017/S1751731119001484).
- MacCluer JW, Boyce AJ, Dyke B, Weitkamp LR, Pfenning DW and Parsons CJ 1983. Inbreeding and pedigree structure in Standardbred horses. *Journal of Heredity* 74, 394–399.
- Marsoner T, Egarter Vigl L, Manck F, Jaritz G, Tappeiner U and Tasser E 2018. Indigenous livestock breeds as indicators for cultural ecosystem services: a spatial analysis within the Alpine Space. *Ecological Indicators* 94, 55–63.
- Meuwissen TH 1997. Maximizing the response of selection with a predefined rate of inbreeding. *Journal of Animal Science* 75, 934–940.
- Meuwissen TH 2009. Genetic management of small populations: a review. *Acta Agriculturae Scandinavica, Section A — Animal Science* 59, 71–79.
- Miller RE 1996. Quarantine protocols and preventive medicine procedures for reptiles, birds and mammals in zoos: -EN- -FR- -ES-. *Revue Scientifique et Technique de l'OIE* 15, 183–189.
- Pedersen LD, Sørensen AC, Henryon M, Ansari-Mahyari S and Berg P 2009. ADAM: a computer program to simulate selective breeding schemes for animals. *Livestock Science* 121, 343–344.
- Rutten MJM, Bijma P, Woolliams JA and van Arendonk JAM 2002. SelAction: software to predict selection response and rate of inbreeding in livestock breeding programs. *The Journal of Heredity* 93, 456–458.
- Schreiner JA 2018. Saving the breeds: German Farmers' preferences for endangered dairy breed conservation programs. *Proceedings in Food System Dynamics* 182–193.
- Täubert H, Reinhardt F and Simianer H 2010. ZPLAN+, a new software to evaluate and optimize animal breeding programs. In *Proceedings of the 9th world congress on genetics applied to livestock production, 2010, Leipzig, Germany*, p. 0950.
- Vereinigte Informationssysteme Tierhaltung w.V 2011. ZPLAN+: a software to evaluate and optimise animal breeding programs. Retrieved on 1 September 2018 from <https://service.vit.de/zplanplus/help/ZPLANOnlinedocumentation.pdf>.
- Wang Y, Segelke D, Emmerling R, Bennewitz J and Wellmann R 2017. Long-term impact of optimum contribution selection strategies on local livestock breeds with historical introgression at the example of German angler cattle. *G3: Genes, Genomes, Genetics* 7, 4009–4018.
- Wathes DC, Brickell JS, Bourne NE, Swali A and Cheng Z 2008. Factors influencing heifer survival and fertility on commercial dairy farms*. *Animal* 2, 1135–1143.
- Wellmann R, Hartwig S and Bennewitz J 2012. Optimum contribution selection for conserved populations with historic migration. *Genetics Selection Evolution* 44, 34.
- Woolliams JA, Berg P, Dagnachew BS and Meuwissen TH 2015. Genetic contributions and their optimization. *Journal of Animal Breeding and Genetics* 132, 89–99.