Overview

It was noted in Chapter 11 that breeding program design can be pre-determined and implemented through sets of rules, or it can emerge as a consequence of decisions made at the level of individual matings. This latter approach is the tactical approach, with decisions made tactically in the face of prevailing animals and other resources.

Tactical implementation of breeding programs provides a practical means to integrate technical, logistical and cost issues facing animal breeders. Moreover, tactical implementation benefits from opportunistic use of prevailing animals and other resources, resulting in better outcomes.

In any breeding operation, there is an almost infinite range of actions that can be made, involving decisions on issues such as animal selection, semen collection and purchase, and mate allocations. Each set of actions is predicted to have a given utility to the breeder - based on factors such as genetic gains, risk, costs and constraints satisfied. The tactical approach described in this chapter works by searching across all these possible routes ahead, and finding the one that is predicted to best suit the breeder's needs. This has only recently become possible because of development of efficient computing algorithms that mimic evolutionary processes to find the best solution.

Introduction

The animal breeder must juggle many issues when s/he makes decisions resulting in implementation of the breeding program, including concerns about breeding objectives, genetic gains, crossbreeding, inbreeding, logistical constraints, and various types of operational cost.

One approach to solving these problems is to follow sets of rules recommended by geneticists and other practitioners, as outlined in Chapter 11. However, such rules are derived from generalised theories and concepts - and these are usually not well integrated with each other. For example, theories and rules about selection, crossbreeding and inbreeding have been developed largely in isolation from each other, such that it is difficult to mix them in real applications, and we are likely to miss the best overall strategy.

Mate Selection is an approach that can be used both to integrate all the key issues facing animal breeders, and to implement the program tactically. A simple example involving selection, crossbreeding and running costs is given in Chapter 11. Mate selection incorporates decisions on animal selection and mate allocation. Because the best
animals to select depends on pattern of mate allocation, and vice versa, we can best
make these decisions simultaneously as mate selection - we just decide what mating
pairs and groups to make.

Moreover, there can be added advantage in making decisions tactically, rather than
following a pre-set strategy. This is because a tactical approach will make use of
knowledge of the full range of actual animals available for breeding at the time of
decision making, as well as other factors such as availability of mating paddocks,
current costs of specified semen, current quarantine restrictions on animal migration,
current or projected market prices, etc. Tactical implementation of breeding programs
gives the power to capitalise on prevailing opportunities - opportunities that would often
be missed when adhering to a set of rules.

The Mate Selection Index (MSI)

The MSI quantifies the value to the breeder of matings made. It is in fact equivalent to
the objective function of Kinghorn and Shepherd (1994), described in Chapter 11.

In some cases, the consequences of a particular mating might be simple and
quantifiable. For example, if the predicted merit of progeny from a mating is, say,
310Kg yearling weight, or +$12 in breeding objective units, then either of these figures
constitutes an MSI for that mating. This can be done because the value of a mating in
such a scheme is independent from what other matings might be made. However, in
most progressive programs this is not the case - the value of a mating depends on what
other matings are actually going to be made. For example, the value of a mating using a
‘new blood’ imported sire to help reduce inbreeding depends on how many other
matings will be made using sires from the same outside source.

This means that for most applications the MSI cannot be specified at the level of
individual matings - we can only calculate an overall MSI that characterises the
combined value of all matings in the mating set. Examples of such an MSI are given by
Kinghorn (1998), Shepherd and Kinghorn (1998) and Kinghorn et al. (1999), and a
further example will be given later in this paper.

Implementation of mate selection

The mate selection approach to breeding is driven by specifying desired outcomes. An
outline of the approach is shown in figure 1. For each mating set tested, the component
outcomes evaluated constitute the overall Mate Selection Index (MSI). Each
component must be evaluated on the same scale, typically the scale of the breeding
objective in units of, for example, dollars profit per breeding cow per year. The MSI
can be set to an arbitrarily low and uncompetitive value for mating sets that break a
constraint - for example mating sets that imply migration against a hard quarantine
barrier, or greater use of liquid funds than a limit specified by the breeder or group.
Figure 1. An outline for implementation of a mate selection index. The set of matings shown is an hypothetical test mating set. The matings specified imply the need for collection of semen, etc., as shown. The mating set is evaluated for all components in the MSI. An efficient algorithm for finding the best mating set is required.

The computing challenge is to find the mating set that gives the best MSI. For this purpose, an evolutionary algorithm was developed (Kinghorn 1998), based on Differential Evolution (Price and Storn 1997). The mate selection driver shown in Table 1 was developed to conduct the search across all legal mating sets. The underlined values in Table 1 drive the three matings noted, and these are the values to be optimised. “No. of uses” (second column for males, second row for females) is the number of matings for which each animal should be used, and this in turn drives selection, including extent of use of each animal. An animal is culled if this is set to zero. “Ranking criterion” is simply a real number. It is ranked to give the column “Rank”. This in turn drives the mate allocation. The first ranked male mating is the single mating from male 3. He is thus allocated to the first available female mating (the one nearest to the left) - the one mating from female 1. The second ranked male mating is the first mating from male 1. He is thus allocated to the second available female mating (the one second nearest to the left) - the one mating from female 1. The third ranked male mating is the second mating from male 1. He is thus allocated to the third available female mating - the one mating from female 4.

Table 1. This table illustrates the components to be optimised for mate selection - they are underlined.
An example mate selection index (optional reading)

The following example MSI pays attention to genetic gain, long-term inbreeding, short-term inbreeding, crossbreeding effects, running costs and logistical constraints. This section is included for completeness. It adds little in concept to what is shown in figure 1, and so this section can be skipped by those not wanting to know more about the nuts and bolts of an MSI.

For any given mate selection set (list of matings to be made):

\[
MSI = \frac{x'G}{2M} + \frac{x'Ax}{4M^2} + \phi F + \chi C - \cos t \quad \text{when no logistical constraint is broken, or}
\]

\[
MSI = \text{a very low value when a logistical constrain is broken. This low value is sufficiently low to ensure that the mating set is not the solution of the mate selection algorithm.}
\]

- \(M\) is the total number of matings to be made. This is typically the number of breeding females, unless MOET or some other form of reproductive boosting is to be an option, whence some breeding females will effectively be mated more than once.

- \(x\) is a vector of number of matings to be made for each candidate, over both sexes. These numbers are the same as ‘number of uses’ in table 1. Thus, for each sex of candidate, the elements of \(x\) are restricted to sum to the total number of matings to be made, giving a total sum of \(2M\) for the elements of \(x\). Meuwissen (1997) and Grundy et al. (1999) treat elements of \(x\) as proportional contributions, with \(x\) restricted to sum to \(\frac{1}{2}\) for each sex of candidates. However, using number of matings as elements of \(x\) is useful for practical application of selection and mate allocation. The difference is handled by dividing by \(2M\) for each instance of \(x\) in the MSI. Restrictions on the maximum value of each element of \(x\) are made as described later. Vector \(x\) could also be extended to accommodate predicted future contributions from existing juveniles and adults, following Meuwissen and Sonesson (1998) or Grundy et al. (1999).
• $G$ is a vector of selection index values for candidates based on multi-trait EBV's, typically in dollar units.

• $\frac{x'G}{2M}$ is the weighted mean EBV of selected parents - it is in fact an estimate of the mean genetic value of progeny arising from the mating set.

• $\lambda$ is a weighting factor on mean coancestry for selected parents (see next item). $\lambda$ is typically negative, to discourage low effective population sizes. Meuwissen (1997) calculates $\lambda$ to give a constrained value of $x'Ax$. However, different values of $\lambda$ can be chosen, effectively giving different index weights on genetic gain (1) and long-term inbreeding ($\lambda$), to give a range of results for these two factors, as shown in figure 3.

• $A$ is the numerator relationship matrix for candidates.

• $\frac{x'Ax}{4M^2}$ is the weighted mean coancestry of selected parents. This reflects long-term inbreeding, reliability of predicted selection response, and risk of poor response achieved. Just as the numerator relationship between two animals is twice the inbreeding predicted in their progeny, this value is equivalent to twice the rate of inbreeding, $2\Delta F$.

• $\phi$ is a weighting factor on predicted progeny mean inbreeding coefficient. A small value for $\phi$ is often sufficient to have a notable effect to reduce progeny inbreeding. This can also be true even when there are competing mate allocation issues in the MSI. Higher values of $\phi$ will affect which animals are selected, as well as mate allocation (Kinghorn et al., 1999).

• $F$ is predicted progeny mean inbreeding coefficient for the mating set under consideration.

• $\chi$ is a weighting factor on predicted progeny mean crossbreeding value $C$. A sensible value for $\chi$ is 1 - this is the implied weight on the genetic gain component $\frac{x'G}{2M}$, and both these components have direct effects on progeny merit, making them of equal importance if merit of later descendants does not feature in the objective.

• $C$ is predicted progeny crossbreeding value - the value predicted using information on breed genotype alone. This is typically predicted using a dominance model of heterosis, incorporating direct and maternal components of both additive and dominance effects (Chapter 4). Use of $\chi C$ aims just one generation ahead. A more involved approach is required in order to aim further ahead (Shepherd and Kinghorn, 1999), making investment matings (eg. to generate first cross females) as well as realisation matings (eg. terminal sire by first cross female). If $\chi C$ is included in the MSI then EBVs in $G$ should be net of breed genotype effects, to avoid double counting of these effects.
cost is the cost of the mating policy implied by x. This can include costs of AI and MOET. It can be calculated to discourage solutions that, for example, nominate allocation of just a few females to a natural mating male, as well as giving both genetic and economic consideration to use of reproductive manipulation. Figure 2 gives a simple example for females. In one mode of operation, the price of reproductive techniques used to drive figure 2 can be decreased until reproductive technology starts to feature in the best mating set, and this illustrates a break-even price for use of that technology. cost can include other components such as seedstock purchase prices and transport costs, expressed in the same units as the dollar EBV’s in G (see Chapter 11 for a simple example).

Other MSI components not in this example include penalties on variation in progeny trait expression, attention to connection between flocks and optimising QTL expression in progeny.

![Graph](image)

Figure 2. An illustration of one way to formulate costs for female matings. Calculated costs are summed over all candidates, of both sexes, to contribute to the component cost in the MSI.

Logistical constraints are simply applied by examining each contending mating set and applying an MSI value of low value, or an overriding penalty, if any constraint is broken. A related strategy is to apply a moderate penalty - this means that matings sets that break a constraint but are otherwise of high merit can contribute to finding the best solution. However, the penalty must be applied in such a manner that the final solution contains no broken constraints. Here are some example constraints:

- Nominated maximum number of matings for each candidate. This might be, for example, 40 matings for males that cannot have semen taken from them, 1000 for males that can have semen taken, 1 for females that cannot enter a MOET program and 8 for females that can. The figure for dead males might be the number of semen doses available. Minimum numbers can also been set, where a
minimum number of semen doses per animal must be purchased. Of course zero is an accepted value in such cases.

- Migration constraints include not permitting young rams to migrate from flock of birth, and restricting older natural mating rams to be used in just one flock alone. Quarantine barriers can also set set in a simple manner.

- Any factor in the MSI can be included as a constraint instead of an index component. For example, long-term inbreeding can be included as a constraint by using a simplified MSI, (MSI = $\frac{x'G}{2M} + \phi F + \chi C - \cos t$) and penalising any mating set for which $\frac{x'Ax}{4M^2} \equiv 2\Delta F$ exceeds a predetermined value. For example, to constrain $\Delta F$ to 0.02 per generation, this value should be set at $\frac{1}{2}$ times 0.02 equals 0.01.

To calculate optimal values for MSI index weights $\chi$, $\phi$ and $\lambda$ would be a complex undertaking. However, these can be manipulated to give a desire outcome. An example of this is shown in figure 3, where $\lambda$ is varied in order to give a frontier of outcomes for genetic gain and long-term inbreeding. (see also Dynamic control of desired outcomes, below).

![Figure 3](image-url)

**Figure 3.** Plot of predicted progeny merit (EBV Index) against mean predicted long-term inbreeding per generation, for 13 alternative mate selection sets generated by using 13 values of $\lambda$. The chosen solution is circled.
Application of the mate selection approach

The mate selection approach outlined here has been implemented as Total Genetic Resource Management (TGRM, trademarked to LAMBPLAN). Information to implement TGRM includes parameters that describe conditions and desires, and data on animals.

Parameters that describe conditions and desires

These currently include:

- Number of matings to be made, across all breeding units in the analysis
- Whether costs are to be applied
- Information on costs
- MSI weights

Plus, as required:

- Direct and maternal breed and heterosis effects
- Effects of known quantitative trait genes
- Mating contraints
- Constraints to be applied to trait expression
- Any other constraints

Data on animals

If possible, pedigree data should be extensive, including all relatives of all animals that are candidates for breeding. This helps when calculating the numerator relationships among candidates. Pedigree data are simply animal’s ID, sire’s ID and dam’s ID for each animal, with 0 or * entered for unknown parents.

Additional fields should be filled in for animals that are potential candidates for breeding. These fields must include:

- Sex of the animal. This is required in order to separate male and female candidates.
- EBV of the animal, or some other criterion of genetic merit. This is usually the multi-trait EBV calculated from a BLUP run followed by application of economic weights (see Chapter 20).
- Candidate status of the animal. This is the maximum number of matings that can be made by the animal, and reflects natural mating versus use of reproductive boosting (AI, MOET etc.). Values are typically higher for males (25 to 1000+) than for females (1 to 8+). Candidate status defines a limit, and does not mean that the animal will automatically be used for that number of matings.

Other information on each animal can be included as required, for example:
• Information on individual traits, in order to place restrictions on progeny expression of these traits, or simply to report expected outcomes in terms of these individual traits.

• Information on breed genotype of the animal, in order to accommodate crossbreeding effects.

• Information on the animal’s genetic markers for known quantitative trait loci.

Output and reporting

The mean value of key variables for the chosen mating set is reported, such as predicted genetic merit of progeny, long-term rate of inbreeding, progeny inbreeding, progeny heterosis and program costs.

The sires selected are listed together with their number of matings and distribution of these matings across flocks/herds.

The part of the report to be acted on is the mating list. This lists the male and female to be used for each mating, together with predicted merit, inbreeding etc. for progeny from each mating. This mating list constitutes decisions on all the breeding issues addressed in the mate selection run.

Dynamic control of desired outcomes

As the mate selection analysis is running it is possible to view key aspects of the currently best solution in a visual manner. This means showing predicted progeny trait merit, trait distributions, inbreeding, heterosis, costs and structural components, such as the pattern of use of sires over flocks, using real-time graphical output. The user can then change weighting factors and constraints during the analysis so that these outcomes change in desired directions. This approach will give great flexibility to learn about the potential outcomes and to optimally balance them, without having to rely on theoretical calculations about what weighting factors to be used a priori.

This approach is similar to the desired gains selection index approach, except that here the index (MSI) covers much more ground than selection alone.

Use over multiple stages

It is possible to carry out mate selection runs to make culling decisions well before joining time. In this way it is possible to undertake, for example, relatively heavy culling by castrating males, at a relatively early stage, while accommodating concerns about (lack of) relevance of early measures of merit, inbreeding, cost savings, etc.

A separate run can be made well before mating for the purpose of identifying semen, embryos and seedstock to purchase. A later run for the main mating round will benefit from knowledge of purchases made and any change in the candidate status of other animals. A further run can be made to make backup mating decisions in the light of knowledge of which females did not conceive.
Dispersal of breeding males to commercial units

A mate selection analysis can be run over both commercial and stud operations, such that it solves the problem of dispersal of bulls to commercial units, simultaneously with selection of bulls into the stud(s) (Figure 4). The competition between commercial units for bulls can be handled in a manner that optimises overall profit, in harmony with bull selection for the breeding operation, and all the components in the MSI.

As the value of prospective progeny is calculated specifically for the herds in which they will be born, the benefits will be highest where the commercial units have different breeding objectives, as in figure 4. Moreover, where crossbreeding is practiced in some commercial units, the range of terminal and maternal attributes of candidate sires can be well accommodated via their EBVs and knowledge of their breed genotype and that of their prospective commercial mates. This also holds when the commercial destinations involve different end-uses (eg. fully terminal versus ‘daughters may be bred’).

This can be done without individual information on commercial cows, by considering each cow herd, or part thereof, as a single group in the analysis - one ‘nymphomaniac cow’ for each commercial herd.

Figure 4. The fate of stud born bulls. Mate selection can be used to make decisions on dispersal of breeding males to commercial units, simultaneously with stud selection decisions.

Getting the most out of the tactical approach
The tactical approach to breeding is driven by specifying desired outcomes. Although mate selection analysis is a very powerful computing tool, the results that it gives are closely aligned to the ‘outcome instructions’ that it receives. This means that the breeder can have a high degree of control, not by specifying which animals should be selected, but by specifying desires in terms of direction of genetic change, maintenance of genetic diversity, limits in money spent, constrains to be satisfied etc.

Using the tactical approach is like driving a good car in a competitive race. We have control of the steering wheel, accelerator and brakes, and we can drive in a manner that is fast, yet safe, economical and in the proper direction. We no longer need to have our head under the bonnet, monitoring every piston beat, and missing opportunities to overtake or avoid crashes. To make the most of mate selection, we should let it monitor the piston beats, and give it good head to find the best way ahead. There is plenty of opportunity to do test laps of the circuit before committing to a decision - if it does something we do not like, we need to adjust the way we steer it, rather than getting out and pushing it round the track. Here are some examples of how we can give mate selection room to maneuver:

- Pre-culling of animals should be restricted to ‘definite culls’. The mate selection approach will only use competitive animals, but benefits from a bigger pool of candidates.
- It is worth considering the numerical scoring of important visually classed traits. This will permit the use of information from relatives to make faster progress in these traits and monitor their genetic change. It also gives more opportunity to make corrective matings.
- Consider a wide range of outside sires. These can help increase gains, lower inbreeding levels, and provide connections to outside seedstock sources that will result in better gains in the longer term.
- Include all key costs. These can include costs of semen, transport, quarantine holding and even fencing for natural mating paddocks. Limits on finances available can also be set.
- Make flock size variable. By factoring in the cost of maintaining breeding females, flock size can be an outcome of the analysis. This can provide a way to give controlled reduction of flock size through periods of drought or financial hardship, with parallel accommodation of concerns about genetic gains, inbreeding, etc.
- Select sires for commercial units as well as breeding units. This is likely to work well in large enterprises in which the breeding objectives differ between commercial units. This means that the fate of stud males can be use in the stud, use in any one of several production units, or culled.
- There is potential for constraining outcomes. For example, it could be declared that all progeny should be expected to be below a given fat thickness or micron diameter. This is most relevant to breeding operations in which the value of product is high.
• Drive outcomes using a production model. Mate selection could usefully be driven by a dynamic production model, with each mating set evaluated on profit from the optimal production and processing pathway(s) for prospective progeny, as described below.

From TGRM to TRM - Total Resource Management

Mate selection as implemented in TGRM could usefully be driven by a dynamic production model, rather than static breeding objectives. This means that breeding decisions (including dispersal of young bulls to commercial units) could be based on the optimal production and processing pathway(s) for prospective progeny, as suggested in figure 5. The result would account for eg. animal merit, variance in merit, prevailing feed and market conditions, and options for multiple pathways to multiple product endpoints.

Figure 5. Tactical breeding program design could be extended to the full production system. “Total Genetic Resource Management” becomes “Total Resource Management”.

Extension to give *ad hoc* tactical optimisation of the production systems themselves would constitute TRM - Total Resource Management. Properly implemented, TRM would accommodate factors such as:

• Opportunities to draft into separate management groups at different ages and weight ranges, constrained by paddocks, labour etc.
• Levels of feeding, stocking rates, and management within groups.
• Time and space scheduling of limited facilities such as feedlot spaces.
• Predicted optimal pathways from any one point in the chain, through to different product end-points.

• Targeted outcomes to satisfy product objectives, cost constraints, prevailing pricing systems and contracts undertaken.

It is evident that TRM would be of most benefit in vertically integrated enterprises, or cooperating groups with vertical alliances or contracts. This is because of the integration of decision making along part or all of the chain from breeding through to processing.

However, TRM is likely to be much more challenging than TGRM. There is one key critical control point (CCP) for TGRM - mate selection at joining time. Other CCP’s for TGRM involve culling, castration, and semen purchase phases, but these are all based on provisional mate selections.

On the other hand, CCP’s in TRM could include mate selection, drafting at various ages, stocking rates, fertiliser and feed use, timing of facilities use, etc. Moreover, whereas our model of genetic effects and gene transmission in TGRM is relatively straightforward (everybody has a dad and a mum), the bio-economic models underpinning TRM will be more complex and varied.

Notwithstanding this, the tactical decision making approach inherent in TRM would make a useful backbone to place the fruits of scientific and practical research, and make them, by definition, immediately applicable for practitioners in the animal industry chains.

References


