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Combining dynamic epidemiological and genetic models to assess the impact of genetic selection strategies on the spread of classical scrapie within a sheep flock

Ram PrP

genotype

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Introduction

Classical scrapie is a transmissible spongiform encephalopathy that affects small ruminants. Due to potential health risks, it is submitted to eradication measures. It is characterised in sheep by a genetic factor based on the PrP major susceptibility gene. Therefore, control strategies partly rely on selective breeding. Scrapie transmission mechanisms are still incompletely understood. The long infectious and undetectable incubation period makes direct data analyses difficult, hence the interest of a modelling approach.

Aim: study the **impact of selective breeding strategies** on the disease spread within a flock, using a dynamic epidemiological model coupled to the optimal outputs of a genetic model through the breeding function.







FIG 1. a- Genetic model. Population description with 7 categories (rectangles), 7 matings paths (thick lines), selection steps (dotted lines) and selection proportion Wsac (sex s=1: males, 2 : females; age a=1,...,as; category c=1: not elite, 2: elite).

Modelling

The genetic deterministic dynamic model (FIG 1.a) [1] represents breeding programs including:

- •overlapping generations,
- different selection steps,
- different strategies for males and females,

b-Epidemiological model. Flock of susceptible (S) and infected (I) renewal ewes, structured by genotype (g), age class (a), and infection stage (i); demographic (purple) and epidemiological (green) processes.

The realistic SI-based epidemiological model (FIG 1.b) [2] takes into account demographic and epidemiological processes, including: seasonality in breeding and transmission,

- stochastic mating,
- genetic and age-dependent susceptibilities,
- long and variable incubation periods, etc.

•non-random mating. It maximises scrapie resistance, with a constraint on loss of genetic progress in milk production.

Results

The model was applied to 3 French dairy flocks corresponding to breeds exhibiting contrasting PrP genotype frequencies: Lacaune (FIG 2.A), MTN (FIG 2.B), MTR (FIG 2.C). It was simulated for 9 scenarios combining:

- •3 scrapie introduction scenarios,
- 3 breeding strategies.

The robustness of the results was tested by a sensitivity analysis.

► Parameters with the greatest impact on the incidence levels and patterns:

- (i) the transmission scaling factor;
- (ii) the flock;
- (iii) the age and genetic susceptibilities.

Conclusion

It was calibrated using French outbreak data.



Breeding strategies have a strong impact on the disease outbreak and are therefore an efficient tool to control scrapie. However, their efficacy in terms of disease eradication strongly depends on the scrapie introduction scenarios.

emphasise hence These results the contribution of our **original approach**, which combines genetic and epidemiological models in a realistic framework.

[1] A. D. Costard, Z. G. Vitezica, C. R. Moreno, and J.-M. Elsen. A dynamic deterministic model to optimize a multiple-trait selection scheme. J. Anim. Sci. 87(3): 885-894, 2009.

[2] S. Touzeau et al. Modelling the spread of scrapie in a sheep flock: evidence for increased transmission during lambing seasons. Arch. Virol. 151(4):735-751, 2006.

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