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Cebo

▶ To cite this version:

Marine Gelé, Sofia Meurisse, Philippe Trossat, Aurore Oudotte, Didier Boichard, et al.. An equation based on MIR spectra to explore the genetic determinism of spontaneous lipolysis in dairy cows. 2022 ICAR/Interbull Conference, May 2022, Montréal, Canada. pp.132-136. hal-03796270

HAL Id: hal-03796270 https://hal.inrae.fr/hal-03796270v1

Submitted on 4 Oct 2022

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An equation based on MIR spectra to explore the genetic determinism of spontaneous lipolysis in dairy cows

*M. Gelé*¹, *S. Meurisse*¹, *P. Trossat*², *A. Oudotte*², *D. Boichard*³, *L. Bernard*⁴, *C. Hurtaud*⁵, *A. Barbat*³ and *C. Cebo*³

¹Institut de l'Elevage, Maison Nationale des Eleveurs, 149 rue de Bercy, 75595, Paris cedex 12, France (<u>marine.gele@idele.fr</u>)

²Actalia-Cécalait, Rue de Versailles, 39800 Poligny, France

³Université Paris-Saclay, INRAE, AgroParisTech, GABI, 78350 Jouy-en-Josas, France ⁴INRAE, Université Clermont Auvergne, VetAgroSup, UMR Herbivores, 63122, Saint-Genès-Champanelle, France

⁵PEGASE, INRAE, Institut Agro Rennes-Angers, 35590, Saint-Gilles, France

Abstract (400 words maximum)

Mid infrared (MIR) spectra have been used since the late 2000s to phenotype new traits at large scale through prediction equations. It has been used in France since 2012 to quantify free fatty acids on herd milks in the framework of quality-based milk payment system. In addition to the loss of income, a high lipolysis level leads to a degradation of organoleptic (rancid taste) and technological (processing inability) properties of milk. Monitoring lipolysis is required to maintain the quality of milk and dairy products.

Lipolysis depends on many factors: genetics, farming practices, milking equipment, transformation process. Regarding genetics, Vanbergue highlighted in 2017 the existence of a genetic susceptibility to spontaneous lipolysis. The present study aims to investigate this issue further, by studying the genetic determinism of spontaneous lipolysis in a larger number of cows. This work is carried out within the framework of the LIPOMEC project which aims to better understand the molecular mechanisms controlling the degradation of milk fat.

Studying the genetic determinism of lipolysis requires the phenotyping of a large number of cows. As the equation initially developed on herd milk was not fully appropriate for individual milks (range, precision), a new prediction equation was estimated from the MIR spectra of individual milks. For this purpose, 432 milk samples were collected in 4 experimental farms in 2018 (approximately 40 cows per farm sampled 2 or 3 times a year) to maximize the variability of breeds (Holstein, Normande, Montbéliarde) and diets. A joint analysis of lipolysis according ISO/TS 22113 standard (BDI method) and by MIR spectrometry was carried out on each sample. Lipolysis measured by BDI method averaged 0.53 mmol/100 g fat (sd=0.41 mmol/100 g fat). The equation was developed by Partial Least Square regression after LOG transformation. Its coefficient of determination R² reached 0.72, with a residual standard deviation S_{y,x} of 0.19 mmol/100 g of fat. The equation was then applied to obtain phenotypes on more than 300,000 MIR milk spectra from Holstein, Normande and Montbéliarde breeds. Genetic parameters were estimated using a repeatability animal model. Heritability and repeatability estimates were 0.21 and 0.40 in both Normande and Hostein breeds but higher (0.31 and 0.46) in Montbéliarde breed.

This work opens the opportunity to new uses of MIR spectra to improve the control of lipolysis in farm, by a closer management of the herd, or even by selection.

The LIPOMEC project was funded by APIS-GENE.

Keywords: lipolysis, dairy cattle, mid infrared spectroscopy, genetic