

Meta-analysis of public transcriptomics data to understand and phenotype bovine body composition

Jeanne Bazile, Muriel Bonnet, Florence Jaffrezic, Denis Laloë

▶ To cite this version:

Jeanne Bazile, Muriel Bonnet, Florence Jaffrezic, Denis Laloë. Meta-analysis of public transcriptomics data to understand and phenotype bovine body composition. 10. International Symposium on the Nutrition of Herbivores (ISNH10), Sep 2018, Clermont-Ferrand, France. hal-02734215

HAL Id: hal-02734215 https://hal.inrae.fr/hal-02734215v1

Submitted on 2 Jun2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Meta-analysis of public transcriptomics data to understand and phenotype bovine body composition

Jeanne Bazile¹, Florence Jaffrezic², Denis Laloe², Muriel Bonnet¹

¹Université Clermont Auvergne, INRA, VetAgro Sup, UMR Herbivores, 63122 Saint-Genès-Champanelle, France, ²INRA, UMR1313 Génétique Animale et Biologie Intégrative, Jouy-en-Josas, France

E-mail: jeanne.bazile@inra.fr

Take home message From a compilation of public genomics data, *HOXA5* was highlighted as a candidate driver or biomarker of bovine muscle mass accretion with age, in two bovine breeds, and for either steers or bulls.

Introduction Producing ruminant with adequate muscular and adipose tissues masses, *i.e.* lean-to-fat ratio, is an economic challenge for the bovine sector. The lean-to-fat ratio contribute to the animal adaptability, food efficiency and meat / carcass qualities. Despite numerous genomics studies that assayed how rearing factors affect the lean-to-fat ratio (Ceciliani *et al.*, 2018) there is no consensus on molecular biomarkers of the lean-to-fat ratio, perhaps due to the difficulty to compare large datasets produced with different quantitative protocols. One of the rare attempt to compile large datasets had merged micro-arrays data from a same quantitative protocol (Baron *et al.*, 2011). Our challenge is to compile public data and to implement statistical tools such as meta-analysis methods, to identify genes that could be robust biomarkers of the lean-to-fat ratio.

Materials & methods For datasets comparisons, we used R software, first to merge identifiers over datasets and to perform descriptive statistics with the ade4 package. We then used the metaMA package to look for differentially abundant genes in each study and whatever the studies (named Merged Data), by performing Benjamini-Hochberg corrections at a significant level of 5%. This pipeline was bench tested with 4 datasets from 2 transcriptomics studies of bovine *Longissimus* muscle (Qin *et al.*, 2011; Moisá *et al.*, 2013). These studies used different microarrays (UIUC Bos taurus 13.2K 70-mer condensed oligoarray and Affymetrix Bovine Genome Array), and cattle differing by the breed (Chinese Red Steppe vs Angus X Simmental), diet (compensatory growth plane of nutrition vs standard diet) and sex (steers vs bulls).

Results & discussion Only 855 GenBank accessions over thousands were shared by the 4 datasets. One gene (Homeobox protein Hox-A5, *HOXA5*), was overexpressed during muscle growth in each study and in the Merged Data (Figure 1), highlighting *HOXA5* as a potential marker of muscle accretion whatever the breed, diet and castration. The embryonic factor *HOXA5* has been found at the adulthood involved in fat depot in adipose tissue (Gesta *et al.*, 2006), but was never related to post-natal muscle biology. In dataset C, D and Merged Data, 179 genes were differentially abundant and may be drivers of muscle (including marbling) accretion with age. Of these, genes involved in *SMAD*, *TGF*- β or *activin* signalling already known to modulate muscle hypertrophy underscored the meta-analysis method.

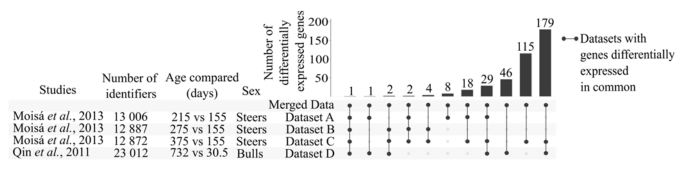


Figure 1 Descriptive data and number of differentially abundant genes in datasets used to bench test the meta-analysis.

Conclusion We provide methods to compile transcriptomics data that should foster the discovery of keys pathways or biomarkers of the lean-to-fat ratio, such as *HOXA5*. The next step is to increase the number of gene under compilation by a reannotation of transcriptomics data on the current bovine genome.

Acknowledgements The regional council of Auvergne-Rhône-Alpes and the Inra-Phase Division for funding J. Bazile.

References

Baron D, Dubois E, Bihouée A, Teusan R, Steenman M, Jourdon P, Magot A, Péréon Y, Veitia R, Savagner F, Ramstein G and Houlgatte R 2011. BMC Genomics 12, 113.

Ceciliani F, Lecchi C, Bazile J and Bonnet M 2018. In Proteomics in Domestic Animals: from Farm to Systems Biology, pp. 233–254. Springer International Publishing.

Gesta S, Blüher M, Yamamoto Y, Norris AW, Berndt J, Kralisch S, Boucher J, Lewis C and Kahn CR 2006. Proceedings of the National Academy of Sciences of the United States of America 103, 6676–6681.

Moisá SJ, Shike DW, Graugnard DE, Rodriguez-Zas SL, Everts RE, Lewin HA, Faulkner DB, Berger LL and Loor JJ 2013. Bioinformatics and Biology Insights 7, 253–270.

Qin L, Zhang G, Cao Y, Zhang J, Zhao Y and Zhao Z 2011. Animal and Veterinary Advances 10, 428-436.