Exploiting the knowledge about model species and humans to revisit the biology of livestock animals

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Exploiting the knowledge about model species and humans to revisit the biology of livestock animals.

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Based on a large panel of approaches from ‘-omics‘ technologies to integrative modelling, prediction and multicriteria evaluation tools, the research carried by UMR1213 Herbivores aims to improve the understanding of biological mechanisms that underlie phenotypes. Especially on-going studies try to make the link between expression of the genome and the phenotypic traits linked to production (e.g. development and quality of tissues and behaviours), efficiency, and adaptation to nutritional, management and climatic disturbances through functional genomics of animal tissues, fluids and behaviours. Due to the incompleteness of annotation of the genome of livestock animal and thus of relevant information, alternative strategies are to use model species (in vivo and in vitro approaches) or to mine genome-wide sets of data from international databases (in silico approach) thank to online and interactive workflows and databases that we are developing. The knowledge gained from the studies in non-ruminant and ruminant species will foster our understanding of biological mechanisms and minimize unnecessary redundancy in research efforts.