The first project year is characterised by start of the generation of large scale and high-throughput data in Pillar 1 and the work packages (WPs) therein:

- 12 rat strains have been sequenced with more than 20X coverage (WP 1.1 Genome Sequencing).
- 362 mature miRNAs have been quantified and revealed novel dandidate miRNAs for the rat (WP 1.2 transcriptome).
- ChIP protocols have been set up and single base resolution methylation profiles of the left ventricle of the Brown Norway (BN-Lx/Cub) and spontaneously hypertensive rat (SHR/Ola) were generated (WP 1.3 epigenetics).
- A large-scale quantiative analysis of two liver proteomes (one BN-Lx and one SHR rat) and one heart proteome (left ventrical from BN-Lx rat) to a depth of 6,808 unique proteins was derived through in vitro super SILAC rat mix (WP 1.4 proteomics).
- MESA, statistical tools and analytical systems for metabolic patterns have been developed (WP 1.5 metabonomics).

Ontologies and the rat interime tool enable data integration and standards for data submission, description and analysis have been developed (WP 1.6 – data infrastructure). Mapping of different kinds of QTLs results in a transcriptional network that is published (Heinig et al (2010) A trans-acting locus regulates an anti-viral expression network and type 1 diabetes risk. Nature 467 (7314):460-464) (WP 2.1 – regulatory networks).

Rat genome modification technology has advanced through EC cells (Meek et. al, Efficient Gene Targeting by Homologous Recombination in Rat Embryonic Stem Cells, Plos ONE 5(12): e14225); ZNF technology has knocked out the first gene (WP 2.2 – functional validation).

Phenotypic and funcational imputation (WP 3.1 - comparative analysis) and tissue collection for translation of the results (WP 3.3 - validation in human tissues) has started. Orthologous regions of interest identified are accessible via Ensembl and RGD (WP 3.2 - integration with hGWAS).

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