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Laura Morel, Peter W Harrison, Guillaume Devailly

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
VizFaDa: Visualisations of FAANG data

Laura Morel¹, Peter W. Harrison², Guillaume Devailly¹

1: GenPhySE, Université de Toulouse, INRAE, ENVT, 31326, Castanet Tolosan, France

2: EMBL-EBI, Hinxton, United Kingdom

Abstract

- Functional Annotation of Animal Genomes (FAANG) is an open consortia gathering genome regulation data in farmed animals: 
- Raw genomic data and rich sample metadata are freely available through a data portal hosted at EMBL-EBI: data.faang.org
- We aim to provide data visualisations of FAANG datasets (RNA-seq, ChIP-seq, DNA methylation), embedded to the FAANG data portal.
- Visualisations will provide an overview of the available data to foster data re-use.

➤ Data processing **nextflow**

Nextflow pipelines based on nf-core:

- A Nextflow module do get metadata & download data
- Fast RNA-seq quantification using Salmon pseudo-aligner
 - Pearson correlation on $\log_{10}(\text{TPM} + 1)$
- ChIP-seq processing using bwa and macs2
 - Jaccard index from bedtools

➤ Notable by-products

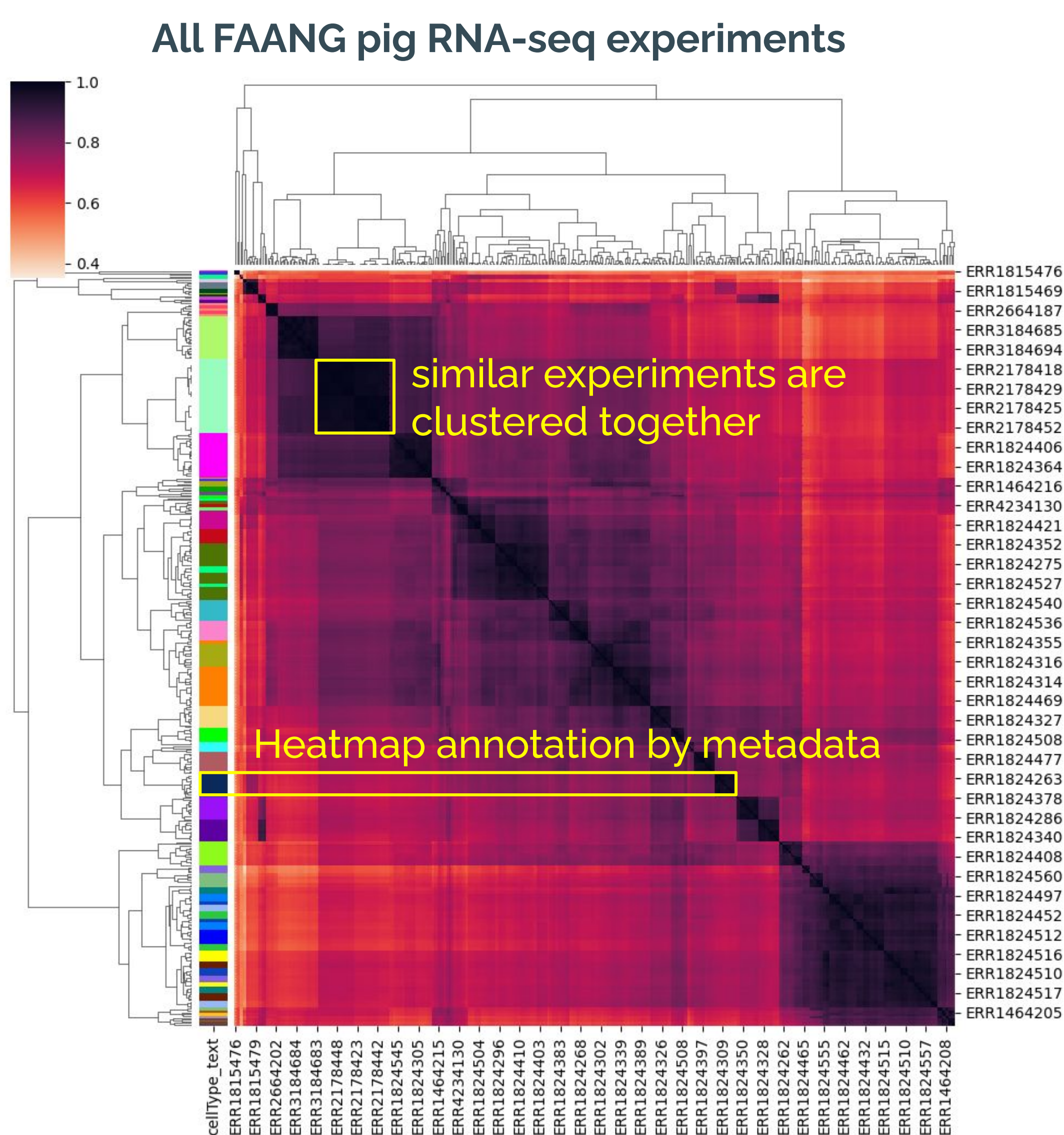
Uniformly processed outputs on FAANG data:

- FastQC reports
- Gene and transcript expression matrices (count & TPM)
- ChIP-seq peaks and bigwig tracks

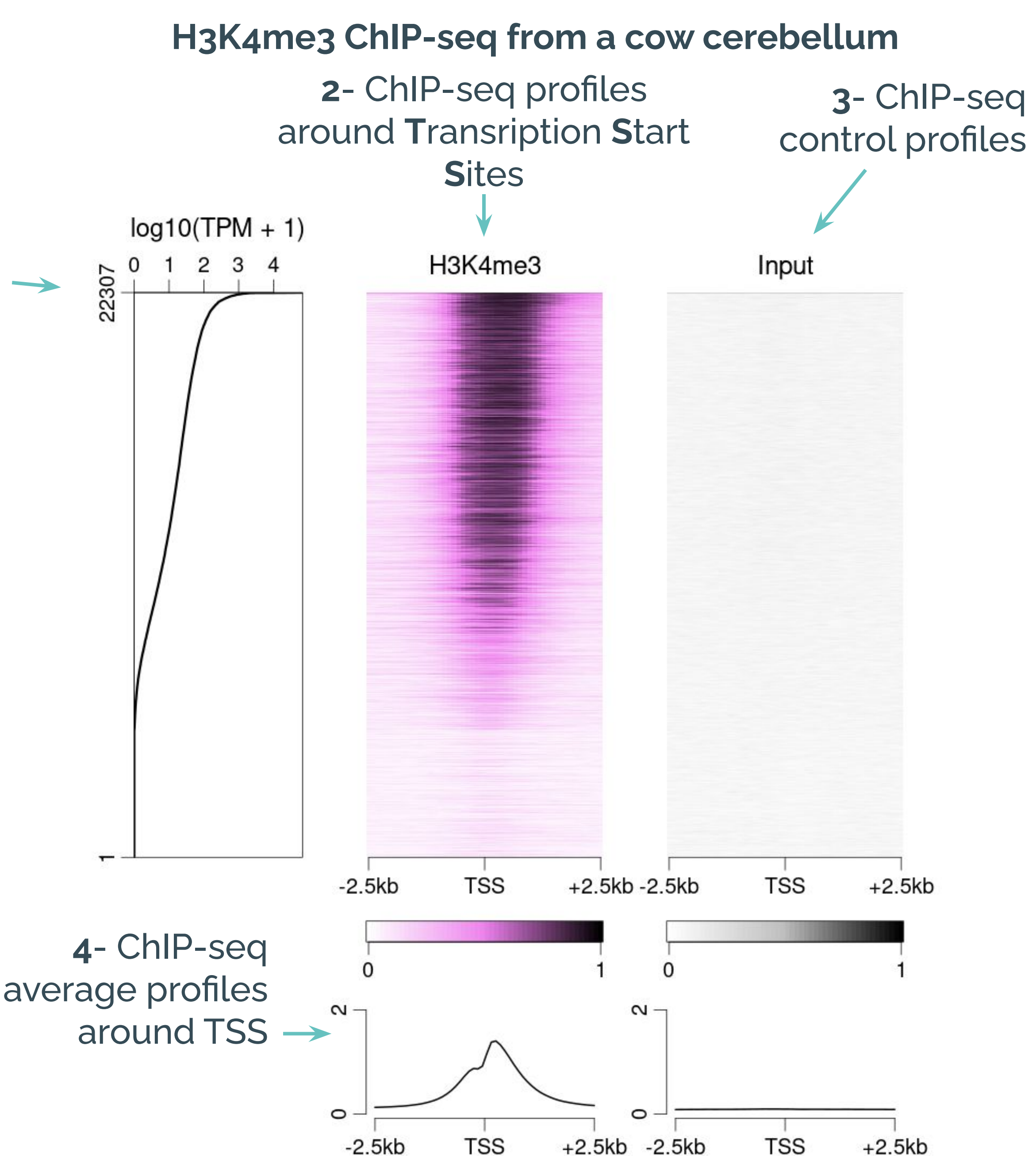
➤ Sample-to-sample correlation heatmaps

By species and by experiment types (RNA-seq, ChIP-seq, DNA methylation)

- overview of available data
- visualising the relationships between samples
- outlier identification

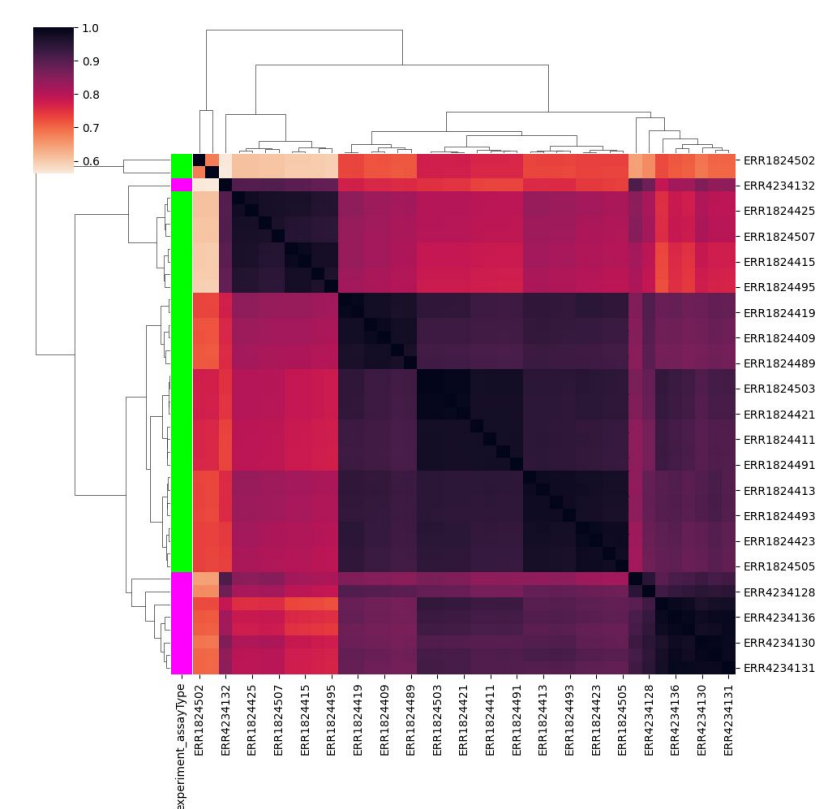


1- The 22,307 annotated *Bos taurus* genes are sorted according to their expression level from a matching FAANG RNA-seq experiment



4- ChIP-seq average profiles around TSS

Heatmaps are subsetable by metadata fields



- Soon: upload your own data
- RNA-seq: gene expression table (count, TPM)
 - ChIP-seq: peak bed file

➤ VizFaDa timeline

2020-07	2021-01	2021-07	2022-01	2022-07
VizFaDa starts	All RNA-seq processed	Web-app (alpha)	All ChIP-seq processed	Web-app (beta)
			All DNA methylation processed	Web app (RC)

- Maintenance:
- new FAANG data
 - using new reference genomes and annotations

➤ Web applications

Soon: a dedicated web-app for visualisations

Soon: integrations to the data.faang.org portal

➤ Fundings

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