PRIDE Peptidome: A hub for peptide evidences from public proteomics datasets for ENSEMBL and Uniprot



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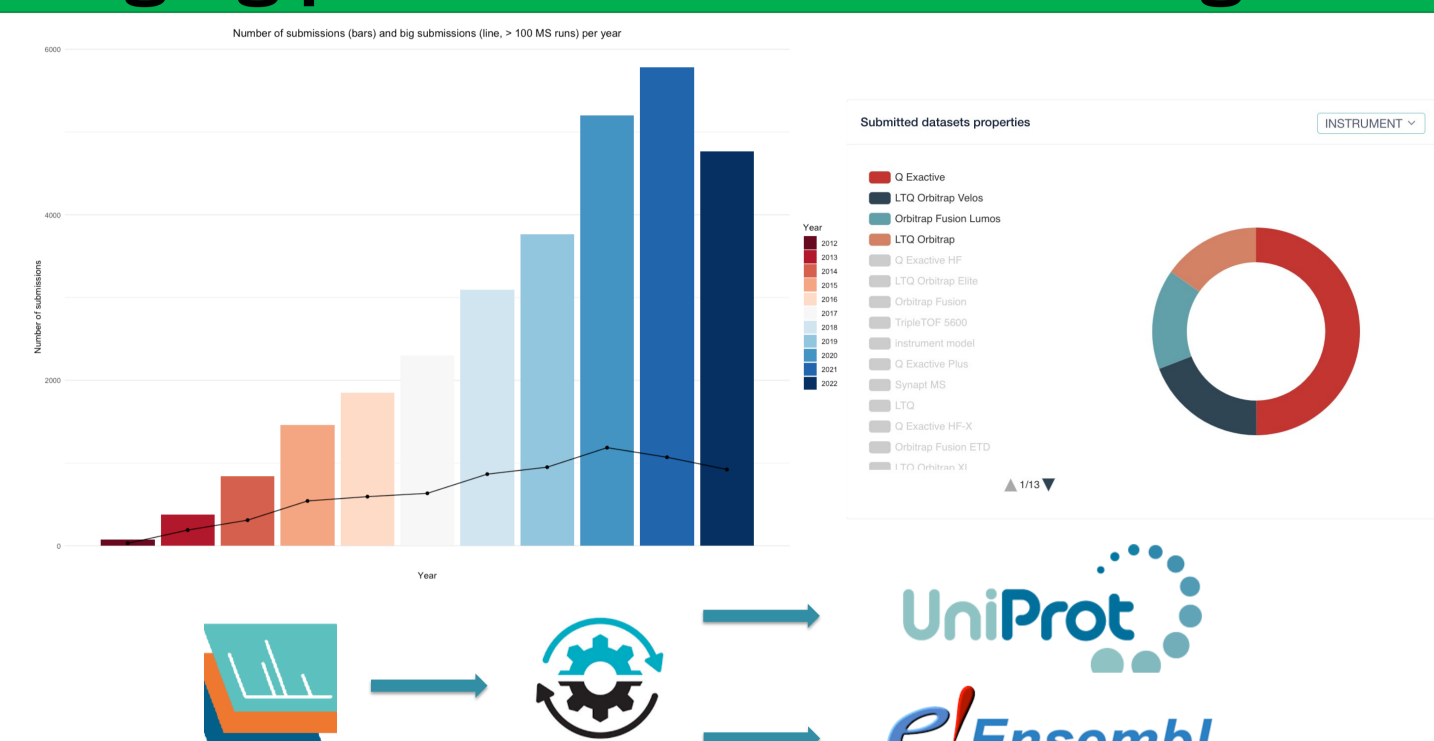
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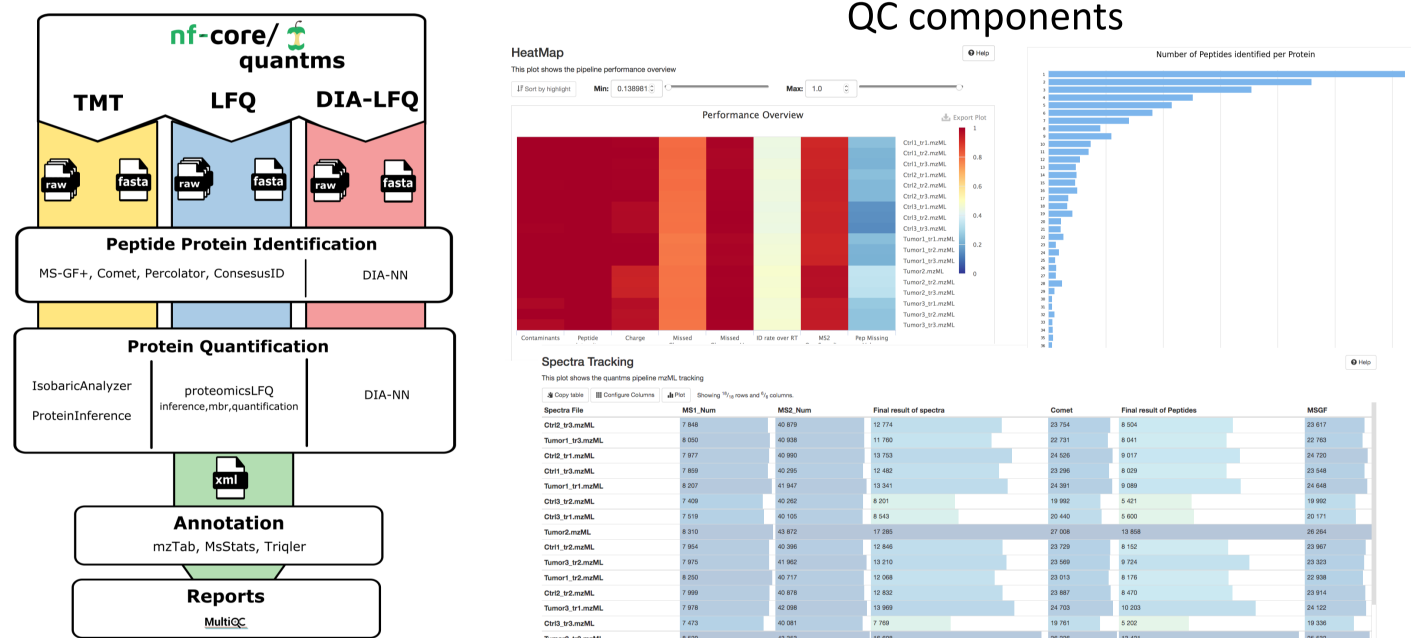
https://www.ebi.ac.uk/pride/. https://quantms.readthedocs.io/

The PRIDE database – Public data reuse: Bringing proteomics data to biologists

- EMBL-EBI's PRIDE database is the largest data resource of MS proteomics data worldwide.
- During 2021 alone, 490 datasets were submitted per month on average to PRIDE.
- Reuse of public proteomics datasets is increasing dramatically, with multiple applications.
- Rather than in the creation of new bioinformatic resources, for sustainability reasons, our focus is to disseminate and integrate PRIDE proteomics data into added-value EMBL-EBI resources, namely UniProt (PTM data, protein sequences), Expression Atlas (protein expression information, the focus of this poster), Ensembl (proteogenomics data) and MGnify (metaproteomics and metagenomics data).
- The goal is to make proteomics data more accessible to life science scientists, especially those non-expert in proteomics.



Internal reanalysis of dataset using quantms (DDA-LFQ, DDA-TMT, DIA)



Major Data-related features:

- Use multiple search engines in combination (msgf+, comet), boost the number of peptides by using percolator.
 Perform multiple protein quantification methods: LFQ-intensity based, spectral counting, TMT intensity, DIA-LFQ
 - Perform multiple protein quantification methods: LFQ-intensity based, spectral counting, TMT intensity, DIA-LFQ intensity based.
 - False localization rate analysis for PTMs including Phosphorylation and acetylation.
 Based on HUPO-PSI standard file formats including mzTab, mzML, SDRF.

Major cloud features:

- Can be executed in LSF, AWS, Google Cloud, Kubernetes, local PC, HPC. Based on BioContainers.
- Open-source, open access.

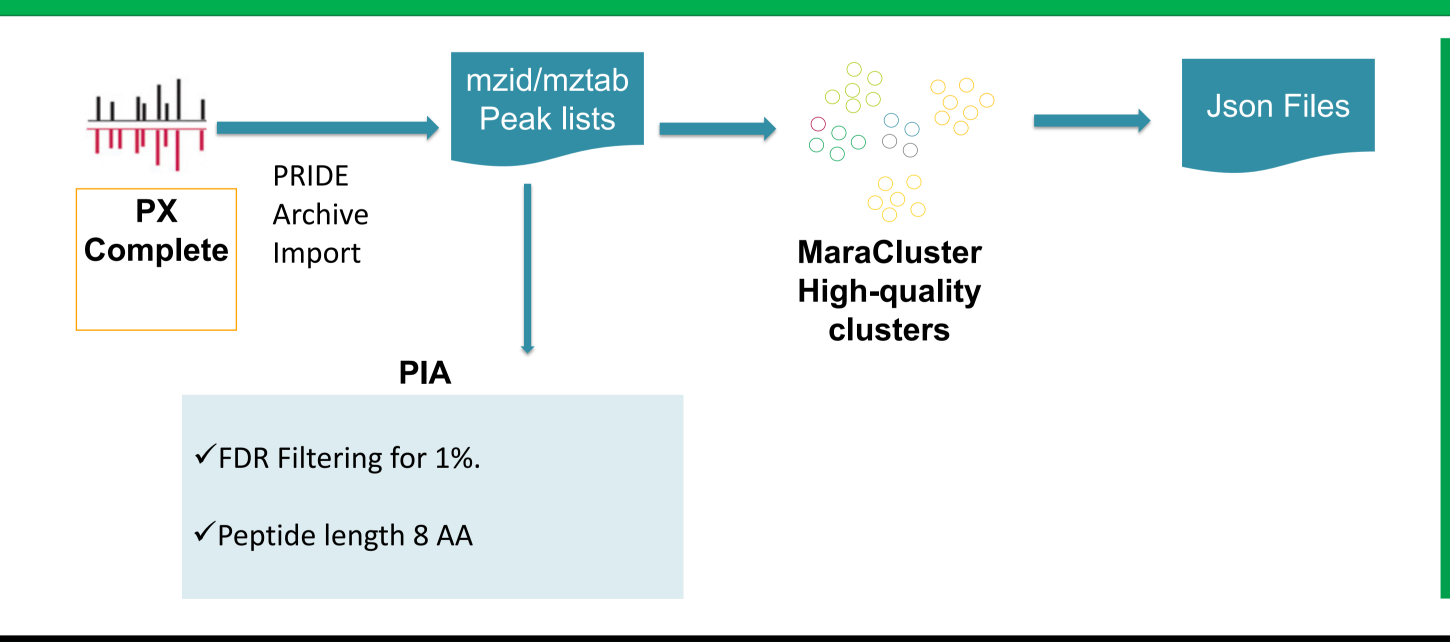
 Include features for acres.
- Include features for compute resource allocation, automatic fail-tolerance and resume workflows.

References:

- 1. Umer, Husen M., et al. Bioinformatics 38.5 (2022): 1470-1472.
- 2. Bai, Mingze, et al. chemRxiv (2022). 10.26434/chemrxiv-2022-x18d5

https://quantms.readthedocs.io/

From complete submissions to peptide evidences



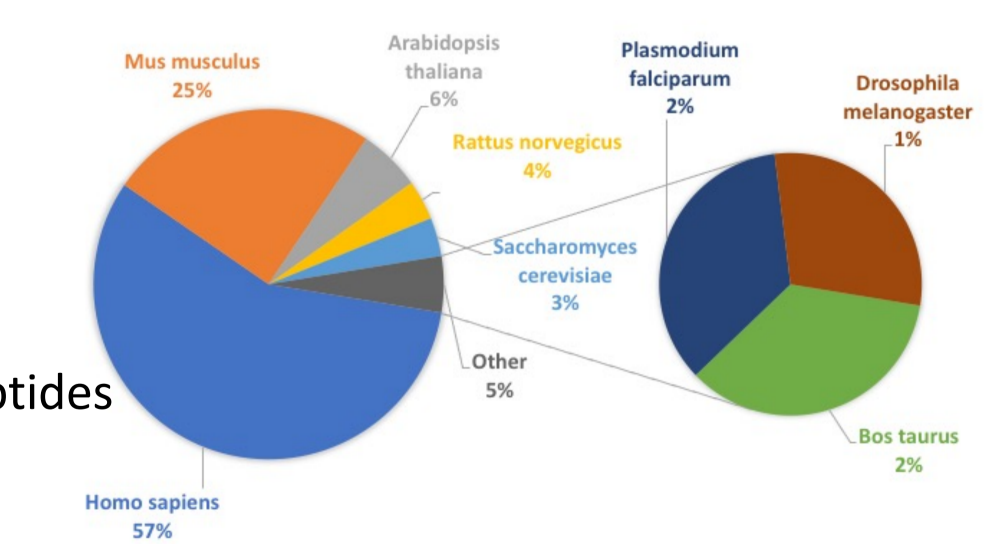
√1294 Projects

✓ 217′261 Proteins

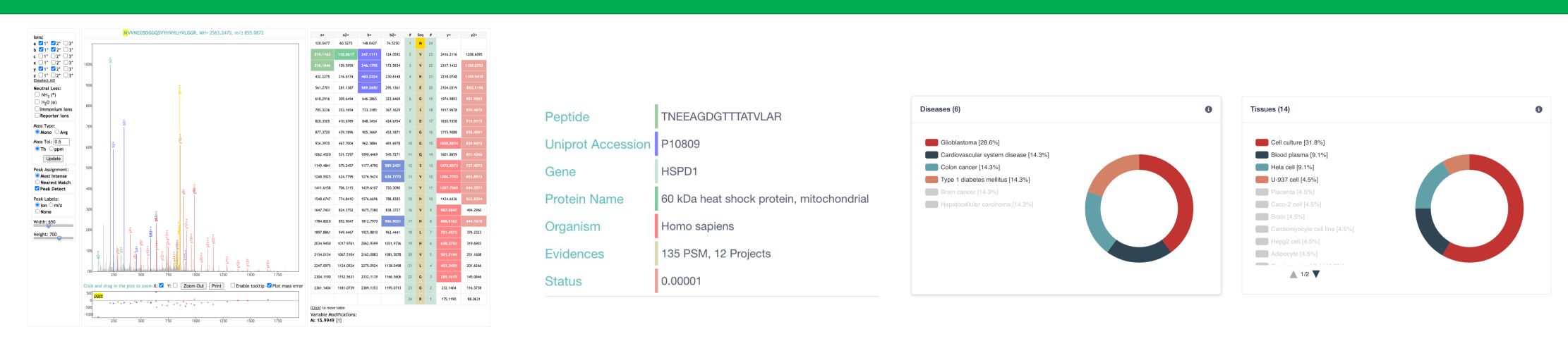
✓ 2′300′389 Peptides

✓ 1'922'855 Unique Peptides

√150′658′201 PSMs



Peptidome and spectra archive



- ✓ Search by peptides, protein accessions, names, gene names.
- ✓ Peptide + protein visualization including names, number of psms and projects.
- ✓ ENSEMBL and Uniprot links to the corresponding spectra that validate a specific sequence feature (e.g., PTM or variant)

References:

- 1. Yasset Perez-Riverol, et. al. Nucleic Acids Research (2022), Pages D543–D552,
- 2. Deutsch, E. W., Perez-Riverol, et. Al. (2021). Nature methods, 18(7), 768-770.

https://www.ebi.ac.uk/pride/peptidome