Where differences resemble: Sequence-feature analysis in curated databases of intrinsically disordered proteins

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Supplementary Material

**Figure S1** – Merge strategy for overlapping annotation regions. Overlapping annotations in the origin datasets are converted to a series of adjacent annotations in MobiDB 3.0. In this analysis, adjacent annotations are considered as a single annotation.

**Figure S2** – Number of overlapping residues. Each cell represents the ID regions overlap between a database pair. Unique residues are those annotated uniquely by a single database. The count is by row, e.g. DIBS annotates 3,325 residues.
Figure S3 – Fraction of overlapping residues. As in Figure S2 but numbers represent percentages calculated over the database size, e.g. DIBS uniquely annotates 26.04% of its ID residues.

Figure S4 – Taxonomic diversity significance. Significance is calculated with the Chi-square test and represent diversity of the taxonomic distribution, P-value < 0.05 represents significantly different distributions. The Chi-square is directional, but the test is consistent in both directions.
**Figure S5** – Heat map of statistical significance of differences in region lengths. The P-value represents the statistical significance of the difference between database length distribution and it is calculated with a T-test.

**Figure S6** – Hierarchically clustered heat map of amino acid absolute frequencies. Clustering is based on Euclidean distance between frequency vectors. Each value represents amino acid frequencies of each database.
**Figure S7** – Correlation matrix of absolute amino acid frequencies. Correlation coefficients (in red) and correlation p-values (in blue) of absolute amino acid.

**Figure S8** – Correlation matrix of fold increase amino acid frequencies. Correlation coefficients (in red) and correlation p-values (in blue) of fold increase compared to TrEMBL reference frequencies.
Figure S9 – Heat map of statistical significance of differences in conformational propensity. Significance is calculated with the Chi-square test and represent diversity in the distribution of the five different Pappu’s classes. P-value < 0.05 represents significantly different distributions. The Chi-square is directional, but the test is consistent in both directions.

Figure S10 – Heat map of statistical significance of differences in region low complexity content. The P-value represents the statistical significance of the difference between region low complexity content distribution and it is calculate with a T-test.