Benchmark comparison of protein sequence preprocessing effect on learning task for Pfam family classification

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Sunday 12th June, 2022

Inspiration and problem overiew

- Data preprocessing for size reduction without loosing crucial information.
- Growing amount of biological data.
- Full Swissprot database: >250 MB
- Full Uniprot database: around 10 GB
- Combining data from other databases would increase those numbers even more.
- Analysis becomes much harder and time-consuming

Inspiration and problem overiew

- Data preprocessing for size reduction without loosing crucial information.
- 0.25% sequences manually annotated in Uniprot
- Automated data annotating
- Need for better models
- More complicated models take longer to train

Biovec

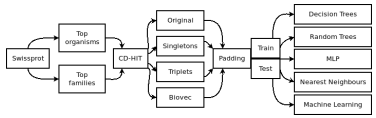
- Biovec [1][2] a multidisciplinary model combining knowledge from biology, linguistics and statistics.
- Analysis of k-mers in bigger sequential context.
- Calculating vectors of probability of k-mers in the given context.
- Using preprocessed metadata for training.

Original Sequence $(1)\overrightarrow{M}^{(2)}\overrightarrow{A}^{(3)}\overrightarrow{F}SAEDVLKEYDRRRRMEAL..$ **Splittings**

- 1) MAF, SAE, DVL, KEY, DRR, RRM, .. 2) AFS, AED, VLK, EYD, RRR, RME, .. 3) FSA, EDV, LKE, YDR, RRR, MEA, ..

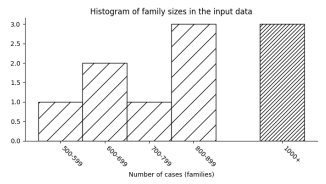
Compared methods of preprocessing

- Data source: Swissprot manually reviewed part of the Uniprot[3] database
- Filtering by most frequent organisms and families
- CD-HIT[4] removing very simmilar protein sequences
- Padding
- Splitting to stratified train and test pools
- Shuffling
- Original
- Singletons dtype int8
- Triplets dtype int16
- Biovec

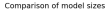


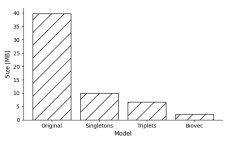
Tested models

- Decision trees
- Random trees
- MLP
- Nearest neighbours
- Machine Learnig (Simple Dense model)
- Grid Search for optimal parameters
- Cross-validation

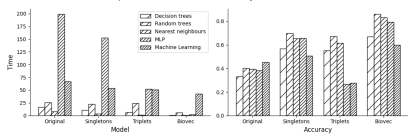


Results





Comparison of time and accuracy between test



Summary

- Treating protein data as string objects might have a negative impact on the training process.
- Simple conversion to numerical data improves both accuracy and runtime.
- Further numerical categorization for triplets while speeding up the learning process is loosing some of the information leading to decreased accuracy.
- Neighbourhood of aminoacids can be analysed using more sophisticated biovec model.
- Biovec yields both better results and up to 100 times faster training process (MLP) compared to the original model.
- Biovec model data weights over 20 times less than the original one.

References

- Article Source: Continuous Distributed Representation of Biological Sequences for Deep Proteomics and Genomics
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- Li W, Godzik A. Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences. Bioinformatics. 2006 Jul 1;22(13):1658-9. doi: 10.1093/bioinformatics/btl158. Epub 2006 May 26. PMID: 16731699.