## **HyPnOBrain**

your homology based HPO neural network predictor

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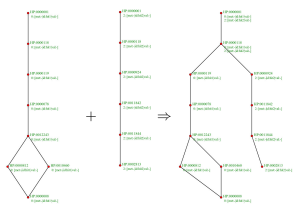
# Homology based function prediction

#### General approach:

- Search for annotated similar sequences with blast and hhblits (hits)
- ▶ Build subgraph of HPO containing the found annotations
- ► Calculate confidence for every annotation from some distance measure to the hits

## Preparations

- Prepare databases for annotated sequences
- ▶ Represent HPO Graph in predictor
- Merge trees corresponding to hits

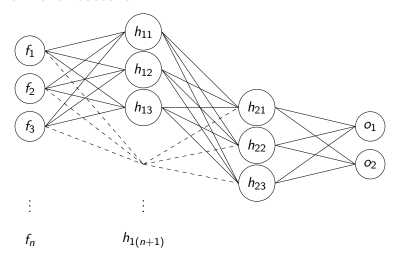


#### **Features**

- use neural network to calculate confidence per node
- each node is assigned 12 features derived from the merged tree

```
number of hits
min. E-value
avg. E-value
product of E-values
longest hit
...
```

### Network architecture



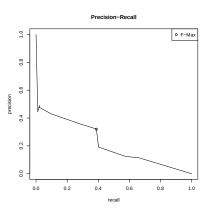
- fully connected net with two hidden layers
- two output nodes trained for the two possible predictions
- difference of predictions as confidence



### Validation

- ► Inspection of the dataset shows: Most sequences have pairwise similarity < 80%
  - $\Rightarrow$  reduce set at 80%-level to remove highly similar clusters
- Crossvalidate over reduced set but allow non-reduced trainingset for similarity search during testing
- Calculate precision and recall per test sequence and average over all sequences

### Results



- $\blacktriangleright$  F-measure 0.35  $\pm$  0.03 (at confidence level 0.34)
- ightharpoonup Precision 0.32  $\pm$  0.04 (at same confidence level)
- ▶ Recall  $0.39 \pm 0.10$  (at same confidence level)