## **Annotating Proteins**

### Finding homologs

```
For Protein to Protein searches BLASTP, phmmer (HMMER), FASTA module load fasta fasta36 query database > results.FASTA fasta36 -m 8c -E 1e-3 query database > results.FASTA.tab
```

#### To Find Domains

```
See Overview lecture Domains lecture
Searching with HMMer against Pfam
See the HMMER tutorial
Searching with Interpro
```

#### Searchin Intepro on HPCC

```
Note this can be slow.
```

```
#SBATCH -p batch -N 1 -n 8
module load iprscan
CPU=4
interproscan.sh --goterms --pathways -f tsv -i PROTEINFILE.fa --cpu $CPU > SEARCH.log
The results will contain information like
Gene Ontology http://geneontology.org/
```

## Running Analyses on Biocluster

```
module load hmmer
module load db-pfam
hmmscan --domtbl domtbl_results.out $PFAM_DB/Pfam-A.hmm proteins.fa > proteins.hmmscan
hmmsearch --domtbl domtbl_results.out $HMM protein-db.fa > protein.hmmsearch
Pfam2GO - http://current.geneontology.org/ontology/external2go/pfam2go
```

# Workshop

- $1. \ \ Searching for Pfam \ domains in sets of proteins \ https://github.com/biodataprog/GEN220\_2019\_examples_See \ search\_SOD1.sh$
- 2. Parsing report files