wt petase model creation

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1 Creating a competative supervised model for wild type PETase activity at low pH

We show in a seperate paper TODO that supervised models should be used for predicting PETase activity at unique conditions (eg. low pH) once some assay labeled data is available, and this outperforms HMMs. Here we create models that: 1. Take in embeddings as input, explore over: Aligned OHE, ES|M2, SaProt, MSATransformer 2. Use linear vs non-linear models: Linear regression, Random Forest

Hyperperameter optimization is conducted over the models for each input type.

Save the final model, which can be loaded like any other sklearn model if AIDE is installed.

eg. model=joblib.load('model.pkl')

```
[]: import os
     import pandas as pd
     import numpy as np
     from sklearn.model selection import RandomizedSearchCV, KFold, cross validate
     from sklearn.feature_selection import VarianceThreshold
     from sklearn.linear_model import ElasticNet, LinearRegression
     from sklearn.neural_network import MLPRegressor
     from sklearn.preprocessing import StandardScaler
     from sklearn.decomposition import PCA
     from sklearn.pipeline import Pipeline
     from sklearn.metrics import roc_auc_score
     from sklearn.ensemble import RandomForestRegressor
     from scipy.stats import loguniform, spearmanr
     import seaborn as sns
     import matplotlib.pyplot as plt
     sns.set style("whitegrid")
     sns.set context("talk")
     import aide_predict as ap
     from aide_predict.utils.data_structures.structures import StructureMapper
```

1.1 1. Load and prepare data

We need to get: 1. The sequences and labels 2. Assign their structures (for SaProt Embedding) 3. Get and MSA of known PETases (for baseline HMMscore and MSA transformer)

```
[]: RAW_DATA_DIR = os.path.join('.', 'data', 'p740')
```

1.1.1 1.1 Label data

```
[]: df = pd.read_csv(os.path.join(RAW_DATA_DIR, 'label_data.csv'), index_col=0).

sample(frac=1, random_state=42)
```

```
[]: df = df.dropna(subset=['activity_at_5.5_40_cryPow'])
# drop rows with non canonical AAs - these were not predicted properly by AF
X = ap.ProteinSequences.from_dict(df['sequence'].to_dict())
has_non_canonical = [x.has_non_canonical for x in X]
df = df[~np.array(has_non_canonical)]
```

```
[]: X = ap.ProteinSequences.from_dict(df['sequence'].to_dict())
y = df['activity_at_5.5_40_cryPow'].values
```

```
[]: sns.kdeplot(y, bw_adjust=0.5) plt.xlabel('Activity at 5.5 pH, 40°C')
```

1.1.2 1.2 Structures

```
[]: # get the structures - Needed for SaProt embedding
mapper = StructureMapper(os.path.join(RAW_DATA_DIR, 'structures'))
mapper.assign_structures(X)
```

1.1.3 1.3 Homolog MSA (for MSA transformer)

Compute weights so that MSA transformer can sample it properly.

```
[]: msa = ap.ProteinSequences.from_fasta(os.path.join(RAW_DATA_DIR, 'D1-Scraped-513.

→mfa'))

msa.aligned
```

```
[]: msa.width
```

1.2 2. Define scoring functions

```
[]: # 5 fold cv
cv_obj = KFold(n_splits=5, shuffle=True, random_state=42)
```

```
[]: # metrics to measure
# marks magnitude of error eg R2, also score AUROC to see if the model can
classify active or not
scoring = {
```

```
'spearman': lambda est, X, y: spearmanr(y, est.predict(X))[0],
'roc_auc': lambda est, X, y: roc_auc_score(y > 0.001, est.predict(X))
}
```

```
[]: def construct_pipeline(embedder, model, pca: bool=False):
         if not pca:
             return Pipeline([
                 ('embedder', embedder),
                 ('var', VarianceThreshold()),
                 ('scaler', StandardScaler()),
                 ('model', model)
             1)
         else:
             return Pipeline([
                 ('embedder', embedder),
                 ('var', VarianceThreshold()),
                 ('scaler', StandardScaler()),
                 ('pca', PCA(n_components=0.95)),
                 ('model', model)
             ])
     def evaluate_pipeline_with_hyperopt(embedder, model_info, n_iter=10):
         """Run hyperparameter optimization on a pipeline with a given embedder and \sqcup
      ⊶model
         embedder: Embedder object eg ap.BaseProteinModel
         model_info: dict with keys:
              'model': sklearn model object
             'param_distributions': dict of hyperparameter distributions for_
      \neg RandomizedSearchCV
         pipeline = construct_pipeline(embedder, model_info['model'])
         random_search = RandomizedSearchCV(
             pipeline,
             param_distributions=model_info['param_dist'],
             n_iter=n_iter,
             cv=cv_obj,
             scoring=scoring,
             refit='spearman',
             n_jobs=1)
         random_search.fit(X, y)
         best_params = random_search.best_params_
         cv_scores = cross_validate(pipeline.set_params(**best_params), X, y,_

¬cv=cv_obj, scoring=scoring)
         return best_params, cv_scores
```

1.3 3. Baseline model: HMM

```
[]: hmm = ap.HMMWrapper()
hmm.fit(msa)

baseline_scores = {
     k: v(hmm, X, y) for k, v in scoring.items()
}
print('Baseline scores:', baseline_scores)
```

1.4 4. Supervised learning: Define embedders, models, and hyperparameter space

```
[]: embedders = {
        'ESM2': ap.ESM2Embedding(
           metadata_folder='esm2_embeddings',_
     'SaProt': ap.SaProtEmbedding(metadata_folder='saprot_embeddings',_

device='mps', pool='mean'),
        'MSATransformer': ap.MSATransformerEmbedding(
           metadata_folder='msa_embeddings', device='mps', pool='mean',
           n_msa_seqs=32
       ),
        'AlignedOneHot': ap.OneHotAlignedEmbedding(
           metadata_folder='onehot_embeddings')
    # fit the models that have fixed fitting over folds
    embedders['ESM2'].fit([])
    embedders['SaProt'].fit([])
    embedders['MSATransformer'].fit(msa)
```

```
[ ]: models = {
         'ElasticNet': {
             'model': ElasticNet(),
             'param_dist': {
                 'model__alpha': loguniform(1e-5, 1e2),
                 'model__l1_ratio': np.linspace(0, 1, 11)
             }
         },
         'RandomForest': {
             'model': RandomForestRegressor(n_estimators=10),
             'param_dist': {
                 'model__max_depth': [None, 10, 100],
                 'model__min_samples_split': [2, 5, 10],
                 'model__min_samples_leaf': [1, 5, 10]
             }
         }
```

```
}
```

5. Train and evaluate models with hyperparameter optimization

```
[]: import joblib
     if not os.path.exists('search_results.pkl'):
         results = {}
     else:
         results = joblib.load('search_results.pkl')
[]: for embedder_name, embedder in embedders.items():
         for model name, model info in models.items():
             if f'{embedder_name}_{model_name}' in results:
                 print(f"Skipping {embedder name} with {model name}...")
                 continue
             else:
                 print(f"Evaluating {embedder_name} with {model_name}...")
             best_params, scores = evaluate_pipeline_with_hyperopt(embedder,_
      →model_info, n_iter=200)
             results[f'{embedder_name}_{model_name}'] = {
                 'embedder': embedder_name,
                 'model': model name,
                 'best_params': best_params,
                 'spearman': scores['test spearman'],
                 'roc_auc': scores['test_roc_auc']
             joblib.dump(results, 'search_results.pkl')
[]: # convert to long
     df_list = []
     for item in results.values():
         for i in range(5): # Assuming 5 values for each metric
             df_list.append({
                 'embedder': item['embedder'],
                 'model': item['model'],
                 'spearman': item['spearman'][i],
                 'roc_auc': item['roc_auc'][i]
             })
     df = pd.DataFrame(df_list)
     # Melt the DataFrame to create a column for the metric type
     df_melted = pd.melt(df, id_vars=['embedder', 'model'], var_name='metric',__
      →value name='value')
```

```
[]: plt.figure(figsize=(16, 10))
     # Create the faceted plot
     g = sns.catplot(
         data=df_melted,
         kind="bar",
         x="model",
         y="value",
         hue="metric",
         col="embedder",
         height=4,
         aspect=1.0,
         palette="Set2",
         col_wrap=4,
         ci="sd",
         legend=True # We'll add the legend manually
     # remove legend from seaborn
     g._legend.remove()
     # add baselines of the same color as the bars
     for ax in g.axes.flat:
         for i, metric in enumerate(['spearman', 'roc_auc']):
             ax.axhline(baseline_scores[metric], color=sns.color_palette("Set2")[i],_
      ⇒linestyle='--', label=f'Baseline {metric}')
         ax.set_ylim(0, 1)
     plt.legend(loc='upper center', bbox_to_anchor=(-1.1, -0.28), ncol=4)
     # Customize the plot
     g.set_axis_labels("Model", "Value")
     g.set_titles("{col_name}")
     # Display the plot
     plt.savefig('p740_model_comparison.png', bbox_inches='tight', dpi=300)
    1.6 6. Train final model and save
```

```
best_row = df.groupby(['embedder', 'model']).mean().sort_values('roc_auc',__
ascending=False).iloc[0]
best_row

[]: best_pipeline = construct_pipeline(embedders[best_row.name[0]], models[best_row.
aname[1]]['model'])

[]: best_pipeline.fit(X, y)

[]: joblib.dump(best_pipeline, 'p740_best_pipeline.pkl')
```