```
In [1]: import os
    os.environ["OMP_NUM_THREADS"] = "1"
    import flexynesis
    import torch
    torch.set_num_threads(4)
Seed set to 42
```

In [2]: # parameters cell (required to pass arguments to the notebook) (see View HPO ITER = 5 # number of HPO iterations for final modeling run

Modeling Breast Cancer Subtypes

Here, we demonstrate the capabilities of flexynesis on a multi-omic dataset of Breast Cancer samples from the METABRIC consortium. The data was downloaded from Cbioportal and randomly split into train (70% of the samples) and test (30% of the samples) data folders. The data files were processed to follow the same nomenclature.

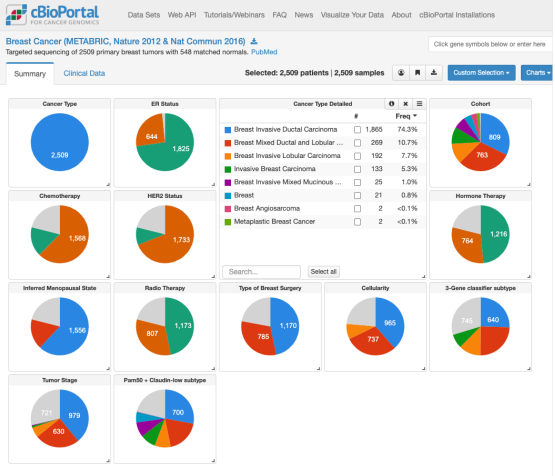
- gex.csv contains "gene expression" data
- cna.csv contains "copy number alteration" data
- mut.csv contains "mutation" data, which is a binary matrix of genes versus samples.
- clin.csv contains "clinical/sample metatada", which is a table of clinical parameters such as age, gender, therapy, subtypes.

Data Download

The data can be downloaded as follows:

```
In [3]: if not os.path.exists("brca_metabric_processed"):
    !wget -0 brca_metabric.tgz "https://bimsbstatic.mdc-berlin.de/akalin/")
```

```
--2025-03-10 12:13:54-- https://bimsbstatic.mdc-berlin.de/akalin/buyar/fl
exynesis-benchmark-datasets/brca metabric processed.tgz
Resolving bimsbstatic.mdc-berlin.de (bimsbstatic.mdc-berlin.de)... 141.80.
181.47, 141.80.181.46
connected. to bimsbstatic.mdc-berlin.de (bimsbstatic.mdc-berlin.de)|141.8
0.181.47|:443...
HTTP request sent, awaiting response... 200 OK
Length: 407225158 (388M) [application/octet-stream]
Saving to: 'brca metabric.tgz'
                     100%[========] 388.36M
brca metabric.tgz
                                                           202MB/s
                                                                       in 1.9
2025-03-10 12:13:56 (202 MB/s) - 'brca_metabric.tgz' saved [407225158/4072
25158]
brca_metabric_processed/
brca_metabric_processed/test/
brca_metabric_processed/test/gex.csv
brca_metabric_processed/test/mut.csv
brca_metabric_processed/test/clin.csv
brca_metabric_processed/test/cna.csv
brca_metabric_processed/9606.protein.aliases.v12.0.txt.gz
brca metabric processed/9606.protein.links.v12.0.txt.gz
brca_metabric_processed/train/
brca_metabric_processed/train/gex.csv
brca_metabric_processed/train/mut.csv
brca_metabric_processed/train/clin.csv
brca metabric processed/train/cna.csv
  cBioPortal
                  Data Sets Web API Tutorials/Webinars FAQ News Visualize Your Data About cBioPortal Installations
```



Let's check the number of samples and number of features in the corresponding files

under train and test folders:

Importing Multiomics Data Into Flexynesis

Procedure

We use the flexynesis.DataImporter class to import multiomics data from the data folders. Data importing includes:

- 1. Validation of the data folders
- 2. Reading data matrices
- 3. Data processing, which includes:
 - Cleaning up the data matrices to:
 - remove uninformative features (e.g. features with near-zero-variation)
 - remove samples with too many NA values
 - remove features with too many NA values and impute NA values for the rest with few NA values
- 4. Feature selection **only on training data** for each omics layer separately:
 - Features are sorted by Laplacian score
 - Features that make it in the top percentile
 - Highly redundant features are further removed (for a pair of highly correlated features, keep the one with the higher laplacian score).
- 5. Harmonize the training data with the test data.
 - Subset the test data features to those that are kept for training data
- 6. Normalize the datasets
 - Normalize training data (standard scaling) and apply the same scaling factors to the test data.
- 7. (Optional): Log transform the final matrices.
- 8. Distinguish numerical and categorical variables in the "clin.csv" file. For categorical variables, create a numerical encoding of the labels for training data. Use the same encoders to map the test samples to the same numerical encodings.

Usage

- Here, we import both train/test datasets from the data folder we downloaded and unpacked before.
- We choose which omic layers to import
- We choose whether we want to concatenate the data matrices (early integration) or not (intermediate integration) before running them through the neural networks.
- We want to apply feature selection and keep only top 10% of the features. In the end, we want to keep at least 1000 features per omics layer.
- We apply a variance threshold (for simplicity of demonstration, we want to keep a small number of most variable features). Setting this to 80, will remove 80% of the features with lowest variation from each modality.

```
[INFO] ========= Importing Data =========
[INFO] Validating data folders...
[INFO] ------ Reading Data ------
[INFO] Importing ./brca metabric processed/train/clin.csv...
[INFO] Importing ./brca metabric processed/train/cna.csv...
[INFO] Importing ./brca metabric processed/train/gex.csv...
[INFO] ------ Reading Data -----
[INFO] Importing ./brca metabric processed/test/clin.csv...
[INFO] Importing ./brca_metabric_processed/test/cna.csv...
[INFO] Importing ./brca metabric processed/test/gex.csv...
[INFO] ------ Checking for problems with the input data
[INFO] Data structure is valid with no errors or warnings.
[INFO] ------ Processing Data (train) ------
[INFO] ------ Cleaning Up Data -----
[INFO] working on layer: gex
[INFO] Imputing NA values to median of features, affected # of cells in th
e matrix 7 # of rows: 5
[INFO] Number of NA values: 0
[INFO] DataFrame gex - Removed 16482 features.
[INFO] working on layer: cna
[INFO] Imputing NA values to median of features, affected # of cells in th
e matrix 108 # of rows: 87
[INFO] Number of NA values: 0
[INFO] DataFrame cna - Removed 18033 features.
[INFO] DataFrame gex - Removed 3 samples (0.23%).
[INFO] DataFrame cna - Removed 3 samples (0.23%).
[INFO] Implementing feature selection using laplacian score for layer: gex
with 4121 features and 1303 samples
Calculating Laplacian scores: 100%| 4121/4121 [00:01<00:00, 214
6.53it/s
Filtering redundant features: 100%| 412/412 [00:00<00:00, 906
2.11it/sl
[INFO] Implementing feature selection using laplacian score for layer: cna
with 4509 features and 1303 samples
Calculating Laplacian scores: 100%| 4509/4509 [00:02<00:00, 216
4.22it/sl
Filtering redundant features: 100%| 450/450 [00:00<00:00, 41952
3.63it/sl
```

```
[INFO] ------ Processing Data (test) ------
[INFO] ------ Cleaning Up Data ------
[INFO] working on layer: gex
[INFO] Number of NA values: 0
[INFO] DataFrame gex - Removed 16482 features.
[INFO] working on layer: cna
[INFO] Imputing NA values to median of features, affected # of cells in th
e matrix 63 # of rows: 51
[INFO] Number of NA values: 0
[INFO] DataFrame cna - Removed 18033 features.
[INFO] DataFrame gex - Removed 2 samples (0.36%).
[INFO] DataFrame cna - Removed 2 samples (0.36%).
[INFO] ------ Harmonizing Data Sets -------
[INFO] ------ Finished Harmonizing -------
[INFO] ------ Normalizing Data -----
[INFO] ------ Normalizing Data -----
[INFO] Training Data Stats: {'feature_count in: cna': 450, 'feature_count
in: gex': 408, 'sample_count': 1303}
[INFO] Test Data Stats: {'feature_count in: cna': 450, 'feature_count in:
gex': 408, 'sample_count': 558}
[INFO] Merging Feature Logs...
[INFO] Data import successful.
```

• dataset.dat contains the data matrices

```
In [7]: train_dataset.dat
Out[7]: {'cna': tensor([[ 0.2908, 0.2509, 0.2583, ..., 0.2730, 0.2473, 0.2
        537],
                 [0.2908, 0.2509, 0.2583, \ldots, 0.2730, 0.2473, 0.2537],
                 [0.2908, 0.2509, 0.2583, \ldots, 0.2730, 0.2473, 0.2537],
                 [0.2908, 0.2509, 0.2583, \ldots, 0.2730, 0.2473, 0.2537],
                 [-1.0024, -1.0362, -1.0212, \ldots, -1.0112, -1.0365, -1.0374],
                 [-1.0024, -1.0362, -1.0212, \ldots, -1.0112, -1.0365, -1.0374]]),
         'gex': tensor([[ 0.4363, 0.6053, 0.7949, ..., 0.8402, 1.2532, 1.1
        986],
                 [0.5032, 0.6295, 0.6994, \ldots, 0.3461, 0.0405, 0.1700],
                 [1.0687, 0.8074, 1.1943, \ldots, 0.1788, -0.5907, -0.2283],
                 [0.7849, 0.6369, 1.0743, \ldots, 0.2282, -0.1475, 0.3063],
                 [0.8006, 0.2211, 1.2079, \dots, -0.2034, -0.6651, -0.6278],
                 [-3.1475, -3.2875, -1.6456, \ldots, 0.5189, 0.5569, 0.8683]])
In [8]: train_dataset.dat['gex'].shape, train_dataset.dat['cna'].shape
Out[8]: (torch.Size([1303, 408]), torch.Size([1303, 450]))
```

• dataset.ann contains the sample annotation data (from clin.csv), where the keys are variable names and values are tensors.

```
In [9]: train dataset.ann
Out[9]: {'LYMPH NODES EXAMINED POSITIVE': tensor([0, 1, 5, ..., 2, 0, 0]),
          'NPI': tensor([3.0800, 4.0400, 6.1200, ..., 4.0700, 3.0240, 4.0160],
                dtype=torch.float64),
          'AGE AT DIAGNOSIS': tensor([70.9100, 48.1100, 58.2700, ..., 70.1000, 6
        9.1900, 60.5900],
                dtype=torch.float64),
          'OS MONTHS': tensor([220.3000, 138.1000, 90.8000, ..., 128.4000, 152.
        3000, 108.0667],
                dtype=torch.float64),
          'RFS_MONTHS': tensor([217.4000, 136.2800, 53.1600, ..., 126.7100, 150
         .3000, 102.7600],
                dtype=torch.float64),
          'CELLULARITY': tensor([0., 1., 2., ..., 0., nan, nan], dtype=torch.flo
          'CHEMOTHERAPY': tensor([0., 1., 0., ..., 0., 0., 0.], dtype=torch.floa
        t64),
          'COHORT': tensor([4., 0., 3., ..., 2., 0., 1.], dtype=torch.float64),
         'ER_IHC': tensor([1., 1., 1., ..., 1., 0.], dtype=torch.float64),
          'HER2 SNP6': tensor([2., 2., 2., ..., 2., 2.], dtype=torch.float6
        4),
         'HORMONE_THERAPY': tensor([1., 1., 1., ..., 1., 0.], dtype=torch.f
        loat64),
          'INFERRED MENOPAUSAL STATE': tensor([0., 1., 0., ..., 0., 0., 0.], dty
        pe=torch.float64),
          'SEX': tensor([0., 0., 0., ..., 0., 0., 0.], dtype=torch.float64),
          'INTCLUST': tensor([3., 3., 9., ..., 3., 4., 5.], dtype=torch.float6
          'OS_STATUS': tensor([0., 0., 1., ..., 1., 0., 1.], dtype=torch.float6
          'CLAUDIN_SUBTYPE': tensor([2., 2., 3., ..., 3., 5., 6.], dtype=torch.f
        loat64),
          'THREEGENE': tensor([1., 1., 0., ..., 0., 1., nan], dtype=torch.float6
        4),
          'VITAL STATUS': tensor([2., 2., 0., ..., 1., 2., 0.], dtype=torch.floa
         'LATERALITY': tensor([0., 1., 1., ..., 1., 0., 0.], dtype=torch.float6
          'RADIO THERAPY': tensor([0., 1., 0., ..., 1., 0., 0.], dtype=torch.flo
        at64),
          'HISTOLOGICAL SUBTYPE': tensor([0., 0., 0., ..., 0., 0., 0.], dtype=to
        rch.float64),
         'BREAST_SURGERY': tensor([1., 1., 1., ..., 0., 0., 0.], dtype=torch.fl
        oat64),
          'RFS STATUS': tensor([0., 0., 1., ..., 0., 0., 1.], dtype=torch.float6
        4)}
          • A mapping of the sample labels for categorical variables can be found in
```

 A mapping of the sample labels for categorical variables can be found in dataset.label_mappings

```
In [10]: train_dataset.label_mappings
```

```
Out[10]: {'CELLULARITY': {0: 'High', 1: 'Low', 2: 'Moderate', 3: nan},
           'CHEMOTHERAPY': {0: 'NO', 1: 'YES'},
           'COHORT': {0: 'cohort1',
            1: 'cohort2',
            2: 'cohort3',
            3: 'cohort4'
           4: 'cohort5'},
           'ER_IHC': {0: 'Negative', 1: 'Positve', 2: nan},
           'HER2 SNP6': {0: 'GAIN', 1: 'LOSS', 2: 'NEUTRAL', 3: 'UNDEF'},
           'HORMONE THERAPY': {0: 'NO', 1: 'YES'},
           'INFERRED_MENOPAUSAL_STATE': {0: 'Post', 1: 'Pre'},
           'SEX': {0: 'Female'},
           'INTCLUST': {0: '1',
            1: '10',
            2: '2',
            3: '3',
            4: '4ER+',
            5: '4ER-',
            6: '5',
            7: '6',
            8: '7',
            9: '8'
           10: '9'},
           'OS_STATUS': {0: '0:LIVING', 1: '1:DECEASED'},
           'CLAUDIN SUBTYPE': {0: 'Basal',
            1: 'Her2',
            2: 'LumA',
            3: 'LumB',
            4: 'NC',
            5: 'Normal',
            6: 'claudin-low'},
           'THREEGENE': {0: 'ER+/HER2- High Prolif',
            1: 'ER+/HER2- Low Prolif',
            2: 'ER-/HER2-',
            3: 'HER2+',
            4: nan},
           'VITAL_STATUS': {0: 'Died of Disease',
           1: 'Died of Other Causes',
            2: 'Living',
            3: nan},
           'LATERALITY': {0: 'Left', 1: 'Right', 2: nan},
           'RADIO THERAPY': {0: 'NO', 1: 'YES'},
           'HISTOLOGICAL SUBTYPE': {0: 'Ductal/NST',
            1: 'Lobular',
            2: 'Medullary',
            3: 'Metaplastic',
            4: 'Mixed',
            5: 'Mucinous',
            6: 'Other',
            7: 'Tubular/ cribriform',
           'BREAST SURGERY': {0: 'BREAST CONSERVING', 1: 'MASTECTOMY', 2: nan},
           'RFS STATUS': {0: '0:Not Recurred', 1: '1:Recurred', 2: nan}}
```

 As the data matrices are stored as tensors, the row and column names cannot be stored as tensors. These are stored in the same dataset object as: dataset.samples and dataset.features

```
In [11]: train dataset.samples[1:10], train dataset.features
Out[11]: (['MB-0172',
            'MB-7140'
            'MB-0279',
            'MB-4719',
            'MB-6167',
            'MB-0381',
            'MB-5562',
            'MB-2752',
            'MB-0180'],
           {'cna': Index(['DAP3', 'FCRLA', 'TOP1P1', 'LAMC1', 'TDRKH', 'MST01', 'M
          STO2P',
                   'YY1AP1', 'EFNA1', 'DPM3',
                   'XPR1', 'SOAT1', 'SELENBP1', 'PI4KB', 'SELP', 'RFX5', 'AXDND1',
                   'KIAA1614', 'TRMT1L', 'FM09P'],
                  dtype='object', length=450),
            'gex': Index(['FOXA1', 'MLPH', 'ESR1', 'GATA3', 'SPDEF', 'TBC1D9', 'FO
          XC1', 'C1S',
                   'XBP1', 'CA12',
                   'N4BP2', 'TNFSF14', 'LEP', 'INIP', 'RPL7L1', 'MBD4', 'HCG2P7',
          'ZNF430',
                   'KIAA1791', 'IL10'],
                  dtype='object', length=408)})
```

 We can get a summary of sample metadata using print_summary_stats. For categorical variables, we can the sample counts per label and for numerical variables, we get mean/median statistics.

```
In [12]: flexynesis.print_summary_stats(train_dataset)
```

```
Summary for variable: LYMPH NODES EXAMINED POSITIVE
Numerical Variable Summary: Median = 0.0, Mean = 1.9286262471220261
Summary for variable: NPI
Numerical Variable Summary: Median = 4.04, Mean = 4.017291158864159
Summary for variable: AGE AT DIAGNOSIS
Numerical Variable Summary: Median = 61.79, Mean = 61.30643898695319
Summary for variable: OS MONTHS
Numerical Variable Summary: Median = 114.4666667, Mean = 125.0357380406669
Summary for variable: RFS_MONTHS
Numerical Variable Summary: Median = 100.63, Mean = 109.94034535686878
Summary for variable: CELLULARITY
Categorical Variable Summary:
  Label: High, Count: 656
  Label: Low, Count: 136
  Label: Moderate, Count: 484
  Label: nan, Count: 27
Summary for variable: CHEMOTHERAPY
Categorical Variable Summary:
  Label: NO, Count: 1044
  Label: YES, Count: 259
Summary for variable: COHORT
Categorical Variable Summary:
  Label: cohort1, Count: 308
  Label: cohort2, Count: 196
  Label: cohort3, Count: 521
  Label: cohort4, Count: 159
  Label: cohort5, Count: 119
_ _ _ _ _ _
Summary for variable: ER_IHC
Categorical Variable Summary:
  Label: Negative, Count: 289
  Label: Positve, Count: 994
  Label: nan, Count: 20
Summary for variable: HER2 SNP6
Categorical Variable Summary:
  Label: GAIN, Count: 279
  Label: LOSS, Count: 67
  Label: NEUTRAL, Count: 955
  Label: UNDEF, Count: 2
Summary for variable: HORMONE THERAPY
Categorical Variable Summary:
  Label: NO, Count: 508
  Label: YES, Count: 795
Summary for variable: INFERRED MENOPAUSAL STATE
Categorical Variable Summary:
  Label: Post, Count: 1034
  Label: Pre, Count: 269
Summary for variable: SEX
Categorical Variable Summary:
```

```
Label: Female, Count: 1303
Summary for variable: INTCLUST
Categorical Variable Summary:
  Label: 1, Count: 93
  Label: 10, Count: 151
 Label: 2, Count: 50
  Label: 3, Count: 195
 Label: 4ER+, Count: 154
 Label: 4ER-, Count: 50
 Label: 5, Count: 127
 Label: 6, Count: 60
 Label: 7, Count: 129
 Label: 8, Count: 197
  Label: 9, Count: 97
Summary for variable: OS STATUS
Categorical Variable Summary:
  Label: 0:LIVING, Count: 539
  Label: 1:DECEASED, Count: 764
Summary for variable: CLAUDIN_SUBTYPE
Categorical Variable Summary:
  Label: Basal, Count: 145
  Label: Her2, Count: 152
  Label: LumA, Count: 468
 Label: LumB, Count: 328
  Label: NC, Count: 5
 Label: Normal, Count: 90
  Label: claudin-low, Count: 115
Summary for variable: THREEGENE
Categorical Variable Summary:
  Label: ER+/HER2- High Prolif, Count: 414
  Label: ER+/HER2- Low Prolif, Count: 425
 Label: ER-/HER2-, Count: 197
  Label: HER2+, Count: 128
  Label: nan, Count: 139
Summary for variable: VITAL_STATUS
Categorical Variable Summary:
  Label: Died of Disease, Count: 432
  Label: Died of Other Causes, Count: 331
 Label: Living, Count: 539
  Label: nan, Count: 1
Summary for variable: LATERALITY
Categorical Variable Summary:
  Label: Left, Count: 649
  Label: Right, Count: 588
 Label: nan, Count: 66
Summary for variable: RADIO THERAPY
Categorical Variable Summary:
  Label: NO, Count: 522
 Label: YES, Count: 781
Summary for variable: HISTOLOGICAL_SUBTYPE
Categorical Variable Summary:
  Label: Ductal/NST, Count: 999
```

```
Label: Lobular, Count: 98
 Label: Medullary, Count: 18
 Label: Metaplastic, Count: 1
 Label: Mixed, Count: 135
 Label: Mucinous, Count: 15
 Label: Other, Count: 12
 Label: Tubular/ cribriform, Count: 15
  Label: nan, Count: 10
Summary for variable: BREAST_SURGERY
Categorical Variable Summary:
  Label: BREAST CONSERVING, Count: 524
 Label: MASTECTOMY, Count: 763
 Label: nan, Count: 16
Summary for variable: RFS_STATUS
Categorical Variable Summary:
  Label: 0:Not Recurred, Count: 763
 Label: 1:Recurred, Count: 539
 Label: nan, Count: 1
```

Training flexynesis models

We create a tuner object by specifying:

- 1. dataset: the training dataset (as we constructed above)
- 2. model_class: which model architecture to use: a) DirectPred: a fully connected network (standard multilayer perceptron) with supervisor heads (one MLP for each target variable) b) Supervised Variational Autoencoder: A variational autoencoder (MMD-loss) with supervisor heads (one MLP for each target variable) c) MultiTripletNetwork: A network structured in triplets to enable contrastive learning (using triplet loss) and additiona supervisor heads (one MLP for each target variable)
- 3. target_variables : A comma separated list of target variables (specify the column headers from the clin.csv).
 - One MLP per each target variable will be created.
 - The target variables may contain NA values
- 4. config name: which hyperparameter search space configuration to use.
- 5. n_iter: How many hyperparameter search steps to implement.
- This example runs 1 hyperparameter search step using DirectPred architecture and a hyperparameter configuration space defined for "DirectPred" with a supervisor head for "CLAUDIN_SUBTYPE" variable:

 $tuner = flexynesis. Hyperparameter Tuning (dataset = train_dataset, model_class = flexynesis. DirectPred, target_variables = ["CLAUDIN_SUBTYPE"], config_name = "DirectPred", n_iter=1)$

 We use perform_tuning function to run the hyperparameter optimisation procedure. At the end of the parameter optimisation, best model will be selected and returned.

model, best_params = tuner.perform_tuning()

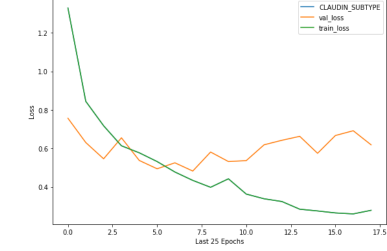
Early Stopping

Training a model longer than needed causes the model to overfit, yield worse validation performance, and also it takes a longer time to train the models, considering if we have to run a long hyperparameter optimisation routine, not just for 1 step, but say more than 100 steps.

It is possible to set early stopping criteria in flexynesis, which is basically a simple callback that is handled by Pytorch Lightning. This is regulated using the early_stop_patience. When set to e.g. 10, the training will stop if the validation loss has not been improved in the last 10 epochs.

One can also visualize the training setting plot_losses to True. This will print the loss values training/validation splits and also the individual loss values for each target variable. In this case, the total loss value for the training equals the loss value of the single variable we chose.

(latent_dim=105, hidden_dim_factor=0.25503043695984917, lr=0.00362561763457623, supervisor_hidden_dim=22, epochs=500, batch_size=32)



| 0/? [00:00<?, ?it/s]

Validation: |

Validate metric	DataLoader 0			
CLAUDIN_SUBTYPE	0.6191389560699463			
val_loss	0.6191389560699463			

```
Tuning Progress: 100% | | 1/1 [00:16<00:00, 16.16s/it, Iteration=1, Best Loss=0.619] [INFO] current best val loss: 0.6191389560699463; best params: {'latent_dim': 105, 'hidden_dim_factor': 0.25503043695984917, 'lr': 0.003625617634576 23, 'supervisor_hidden_dim': 22, 'epochs': 500, 'batch_size': 32} since 0 hpo iterations
```

• One can also provide own parameter optimisation spaces via a yaml file as input:

tuner = flexynesis.HyperparameterTuning(dataset = train_dataset, model_class =
flexynesis.DirectPred, target_variables = ["CLAUDIN_SUBTYPE"], config_name = "DirectPred",
n_iter=1, plot_losses=True, config_path='./conf.yaml') model, best_params =
tuner.perform_tuning()

 We can also provide multiple target variables as input. This will create multiple MLP heads (one per variable) and the network will be trained to learn to predict both variables.

tuner = flexynesis.HyperparameterTuning(dataset = train_dataset, model_class =
flexynesis.DirectPred, target_variables = ["CLAUDIN_SUBTYPE, "CHEMOTHERAPY"], config_name
= "DirectPred", n_iter=1, plot_losses=True, early_stop_patience=10) model, best_params =
tuner.perform_tuning()

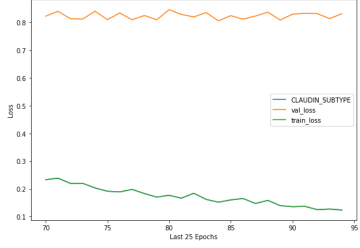
• We can mix numerical and categorical variables. The relevant network structure and evaluation procedures will be applied depending on the type of variable

tuner = flexynesis.HyperparameterTuning(dataset = train_dataset, model_class =
flexynesis.DirectPred, target_variables = ["CLAUDIN_SUBTYPE, "CHEMOTHERAPY",
"LYMPH_NODES_EXAMINED_POSITIVE"], config_name = "DirectPred", n_iter=1,
plot losses=True, early stop patience=10) model, best params = tuner.perform tuning()

Longer Training

In reality, hyperparameter optimisation should run for multiple steps so that the parameter search space is large enough to find a good set. However, for demonstration purposes, we only run it for 5 steps here.

HPO Step=5 out of 5 (latent_dim=107, hidden_dim_factor=0.29138413075201125, lr=0.00015679933916723006, supervisor_hidden_dim=24, epochs=500, batch_size=64)



Validation: |

| 0/? [00:00<?, ?it/s]

Validate metric	DataLoader 0			
CLAUDIN_SUBTYPE val_loss	0.8315162658691406 0.8315162658691406			

Tuning Progress: 100%| 5/5 [02:26<00:00, 29.22s/it, Iteration=5, Best Loss=0.627]

[INFO] current best val loss: 0.6270639300346375; best params: {'latent_di m': 82, 'hidden_dim_factor': 0.21393512381599933, 'lr': 0.0016409286730647 93, 'supervisor_hidden_dim': 12, 'epochs': 500, 'batch_size': 128} since 1 hpo iterations

In [15]: model

Out[15]: DirectPred(

```
(log vars): ParameterDict( (CLAUDIN SUBTYPE): Parameter containing:
          [torch.FloatTensor of size 1])
            (encoders): ModuleList(
              (0): MLP(
                (layer 1): Linear(in features=450, out features=96, bias=True)
                (layer out): Linear(in features=96, out features=82, bias=True)
                (relu): ReLU()
                (dropout): Dropout(p=0.1, inplace=False)
                (batchnorm): BatchNorm1d(96, eps=1e-05, momentum=0.1, affine=True,
          track_running_stats=True)
              )
              (1): MLP(
                (layer 1): Linear(in features=408, out features=87, bias=True)
                (layer out): Linear(in features=87, out features=82, bias=True)
                (relu): ReLU()
                (dropout): Dropout(p=0.1, inplace=False)
                (batchnorm): BatchNorm1d(87, eps=1e-05, momentum=0.1, affine=True,
          track running stats=True)
              )
            )
            (MLPs): ModuleDict(
              (CLAUDIN SUBTYPE): MLP(
                (layer 1): Linear(in features=164, out features=12, bias=True)
                (layer out): Linear(in features=12, out features=7, bias=True)
                (relu): ReLU()
                (dropout): Dropout(p=0.1, inplace=False)
                (batchnorm): BatchNorm1d(12, eps=1e-05, momentum=0.1, affine=True,
          track running stats=True)
              )
            )
          )
In [16]: best_params
Out[16]: {'latent_dim': 82,
           'hidden dim factor': 0.21393512381599933,
           'lr': 0.001640928673064793,
           'supervisor hidden dim': 12,
           'epochs': 42,
           'batch size': 128}
```

Prediction and Model Evaluation

We can use the best model (chosen based on the hyperparameter optimisation procedure) to make predictions on the test dataset

```
In [17]: y_pred_dict = model.predict(test_dataset)
In [18]: y_pred_dict
```

The predictions are class labels for both variables. Now, we can run
 evaluate_wrapper to evaluate all predictions. The wrapper goes through each
 variable and figures out which type of evaluation to apply to the corresponding
 variable (whether to report metrics relevant to regression tasks or classification
 tasks)

```
In [19]: metrics_df = flexynesis.evaluate_wrapper(method = 'DirectPred', y_pred_di
metrics_df
```

Out[19]:	: method		var	variable_type	metric	value	
	0	DirectPred	CLAUDIN_SUBTYPE	categorical	balanced_acc	0.591356	
	1 DirectPred		CLAUDIN_SUBTYPE	categorical	f1_score	0.724952	
	2	DirectPred	CLAUDIN_SUBTYPE	categorical	kappa	0.645426	
	3	DirectPred	CLAUDIN_SUBTYPE	categorical	average_auroc	0.924110	
	4	DirectPred	CLAUDIN SUBTYPE	categorical	average aupr	0.792086	

Extracting the sample embeddings

All models trained within flexynesis comes with a transform method, which extracts the sample embeddings that are generated by the encoding networks (whether it is an MLP or a variational autoencoder). The embeddings reflect a merged representation of multiple omic layers.

```
In [20]: ds = test_dataset
E = model.transform(ds)
In [21]: E.head()
```

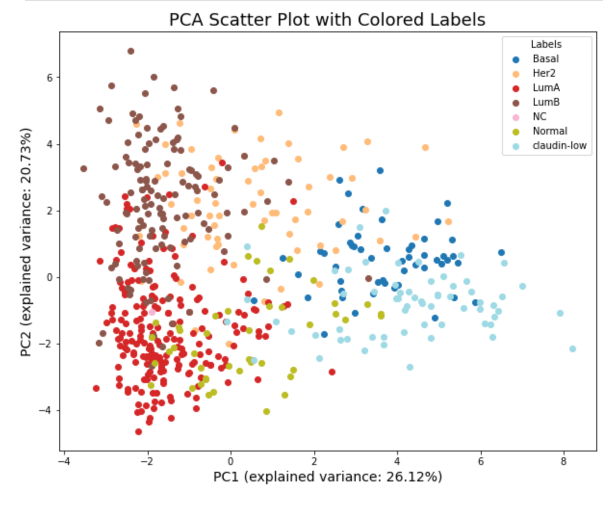
Out[21]:		E0	E1	E2	E3	E4	E 5	E6	
	MB-7253	0.001160	0.154013	0.075810	-0.052389	0.225307	0.437584	0.349962	0.
	MB-0601	-0.021648	-0.016646	0.044106	-0.000734	-0.134784	0.165325	-0.018294	0.
	MB-2929	0.001160	0.154013	0.075810	-0.052389	0.225307	0.437584	0.349962	0.
	MB-4329	-0.021648	-0.016646	0.044106	-0.000734	-0.134784	0.165325	-0.018294	0.
	MB-0599	0.001160	0.154013	0.075810	-0.052389	0.225307	0.437584	0.349962	0.

5 rows × 164 columns

Visualizing the sample embeddings

Let's visualize the embeddings in a reduced space and color by the target variables.

```
In [22]: f = 'CLAUDIN_SUBTYPE'
labels = [ds.label_mappings[f][x] for x in ds.ann[f].numpy()] #map the sa
In [23]: flexynesis.plot_dim_reduced(E, labels, color_type = 'categorical', method
```



We can also use UMAP visualisation

```
In [24]: flexynesis.plot_dim_reduced(E, labels, color_type = 'categorical', method
```

