Imports

```
In [1]:
        import os
        import sys
        sys.path.append('/home/rcendre/classification')
        import itertools
        import pandas
        import webbrowser
        from pathlib import Path
        import matplotlib.pyplot as plt
        from misvm import SIL, MISVM
        from numpy import array, logspace
        from scipy.stats import randint,uniform
        from sklearn.decomposition import PCA
        from sklearn.ensemble import ExtraTreesClassifier
        from sklearn.manifold import TSNE
        from sklearn.pipeline import Pipeline
        from sklearn.preprocessing import LabelEncoder, MinMaxScaler, Rob
        ustScaler, StandardScaler
        from sklearn.ensemble import GradientBoostingClassifier
        from sklearn.svm import LinearSVC
        from sklearn.tree import DecisionTreeClassifier
        from sklearn.neural network import MLPClassifier
        from sklearn.metrics import davies bouldin score
        from toolbox.classification.common import Data, Folds, IO, Tools
        from toolbox.classification.parameters import Dermatology, Settin
        from toolbox.models.models import CustomMIL, MultimodalClassifier
        from toolbox.models.builtin import Applications
        from toolbox.IO import dermatology
        from sklearn.metrics import f1 score
        from toolbox.transforms.common import PredictorTransform, Flatten
        Transform, LinearTransform
        from toolbox.transforms.labels import OrderedEncoder
        from toolbox.transforms.images import DistributionImageTransform,
        DWTImageTransform, FourierImageTransform, HaralickImageTransform,
        SpatialImageTransform
        from toolbox.views.common import Views, ViewsTools
        from toolbox.views.images import ImagesViews
        import warnings
        warnings.filterwarnings('ignore')
```

Using TensorFlow backend.

Parameters

```
In [2]: # Advanced parameters
    data_type='Full'
    prefix = 'Cumulative_Sigmoid'
    validation = 10
    settings = Settings.get_default_dermatology()
```

Photography

```
In [3]: p inputs = IO.load('Photography.pickle')
In [4]: p inputs['Extractor'] = p inputs['ResNetAvg']
        # SVM Linear
        model = Pipeline([('scale', MinMaxScaler()), ('clf', LinearSVC(cl
        ass weight='balanced'))])
        model_params = {'clf_C': logspace(-2, 3, 6).tolist()}
In [5]: low folds = Tools.generate folds(([1, 2, 3, 4, 5], [6, 7, 8],
        [9]), validation)
        Tools.evaluate(p inputs, {'datum': 'Extractor', 'label encode': '
        LesionEncode'}, model, 'Prediction', folds=low folds, distributio
        n=model params, calibrate='sigmoid')
        Evaluation achieved!
In [6]: from IPython.display import HTML
        from IPython.display import display
        diagnosis encoder = OrderedEncoder().fit(['Benign', 'Malignant'])
        # ROC Curve
        ViewsTools.plot size((8,8))
        name = f'Prediction'
        # Label
        display(HTML(ViewsTools.dataframe_renderer([Views.report(p_input
        s, {'label_encode': 'LesionEncode', 'eval': name}, diagnosis_enco
        der)],
        title=[f'Photography'])))
```

Photography

	precision	recall	f1-score	support
Benign	0.56±0.35	0.12±0.27	0.19±0.24	86.00±2.01
Malignant	0.63±0.12	0.94±0.11	0.75±0.06	137.00±2.00
accuracy	0.62±0.10	0.62±0.10	0.62±0.10	0.62±0.10
macro avg	0.59±0.21	0.53±0.09	0.47±0.14	223.00±0.46
weighted avg	0.60±0.21	0.62±0.10	0.54±0.15	223.00±0.46

Dermoscopy

```
In [7]: d inputs = I0.load('Dermoscopy.pickle')
         d inputs['Extractor'] = d inputs['ResNetAvg']
In [8]: dp inputs = IO.load('Photography.pickle')
         dp inputs['Extractor'] = dp inputs['ResNetAvg']
         d inputs = pandas.concat([dp inputs, d inputs], axis=0)
In [9]: all image = [True] * len(d inputs.index)
         single_image = d_inputs['Modality'] == 'Dermoscopy'
         Data.build_bags(d_inputs, single_image, 'ID_Lesion', all_image, '
         ID Lesion', 'Extractor')
         d inputs = d_inputs[single_image].reset_index()
         Tools.transform(d inputs, {'datum': 'Extractor'}, FlattenTransfor
         m(), 'Flat')
In [10]: low_folds = Tools.generate_folds(([1, 2, 3, 4, 5], [6, 7, 8],
         [9]), validation)
         # SVM Linear
         model = Pipeline([('scale', MinMaxScaler()), ('clf', LinearSVC(cl
         ass weight='balanced'))])
         model params = {'clf C': logspace(-2, 3, 6).tolist()}
         Tools.evaluate(d_inputs, {'datum': 'Flat', 'label_encode': 'Lesio
         nEncode'}, model, 'Prediction', folds=low folds, distribution=mod
         el params, calibrate='sigmoid')
```

Evaluation achieved!

```
In [11]: from IPython.display import HTML
from IPython.display import display

diagnosis_encoder = OrderedEncoder().fit(['Benign', 'Malignant'])

# ROC Curve
ViewsTools.plot_size((8,8))

name = f'Prediction'
# Label
display(HTML(ViewsTools.dataframe_renderer([Views.report(d_input s, {'label_encode': 'LesionEncode', 'eval': name}, diagnosis_encoder)],

title=[f'Dermoscopy'])))
```

Dermoscopy

	precision	recall	f1-score	support
Benign	0.53±0.38	0.20±0.21	0.29±0.22	86.00±2.01
Malignant	0.64±0.10	0.89±0.13	0.74±0.07	137.00±2.00
accuracy	0.62±0.08	0.62±0.08	0.62±0.08	0.62±0.08
macro avg	0.58±0.21	0.54±0.06	0.52±0.12	223.00±0.46
weighted avg	0.60±0.19	0.62±0.08	0.57±0.12	223.00±0.46

Microscopy

```
In [12]: m_inputs = I0.load('Microscopy.pickle')
```

Merge

```
In [15]: Tools.transform(m inputs, {'datum': extractor}, LinearTransform(m
         ethod='average'), 'Avg')
         Tools.transform(m inputs, {'datum': extractor}, LinearTransform(m
         ethod='max'), 'Max')
         Tools.transform(m inputs, {'datum': extractor}, LinearTransform(m
         ethod=4), 'Norm4')
         Tools.transform(m inputs, {'datum': extractor}, LinearTransform(m
         ethod=6), 'Norm6')
         Tools.transform(m inputs, {'datum': extractor}, LinearTransform(m
         ethod=8), 'Norm8')
```

Predicting on microscopy

```
In [16]: mp inputs = IO.load('Photography.pickle')
         mp inputs['Extractor'] = mp inputs['ResNetAvg']
In [17]: | md inputs = IO.load('Dermoscopy.pickle')
         md inputs['Extractor'] = md inputs['ResNetAvg']
In [18]: m inputs['Extractor'] = m inputs['Avg']
         m inputs = pandas.concat([mp inputs, md inputs, m inputs], axis=
         0)
In [19]: all image = [True] * len(m inputs.index)
         single image = m inputs['Modality'] == 'Microscopy'
         Data.build bags(m inputs, single image, 'ID Lesion', all image, '
         ID_Lesion', 'Extractor')
         m inputs = m inputs[single image].reset index()
         Tools.transform(m inputs, {'datum': 'Extractor'}, FlattenTransfor
         m(), 'Flat')
In [20]: | # SVM Linear
         model = Pipeline([('scale', MinMaxScaler()), ('clf', LinearSVC(cl
         ass weight='balanced'))])
         model params = {'clf C': logspace(-2, 3, 6).tolist()}
In [21]: low folds = Tools.generate folds(([1, 2, 3, 4, 5], [6, 7, 8],
         [9]), validation)
         Tools.evaluate(m inputs, {'datum': 'Flat', 'label encode': 'Lesio
         nEncode'}, model, 'Prediction', folds=low folds, distribution=mod
         el params, calibrate='sigmoid')
         Evaluation achieved!
```

```
In [22]: from IPython.display import HTML
    from IPython.display import display

diagnosis_encoder = OrderedEncoder().fit(['Benign', 'Malignant'])

# ROC Curve
ViewsTools.plot_size((8,8))

name = f'Prediction'
# Label
display(HTML(ViewsTools.dataframe_renderer([Views.report(m_input s, {'label_encode': 'LesionEncode', 'eval': name}, diagnosis_encoder)],

title=[f'Microscopie'])))
```

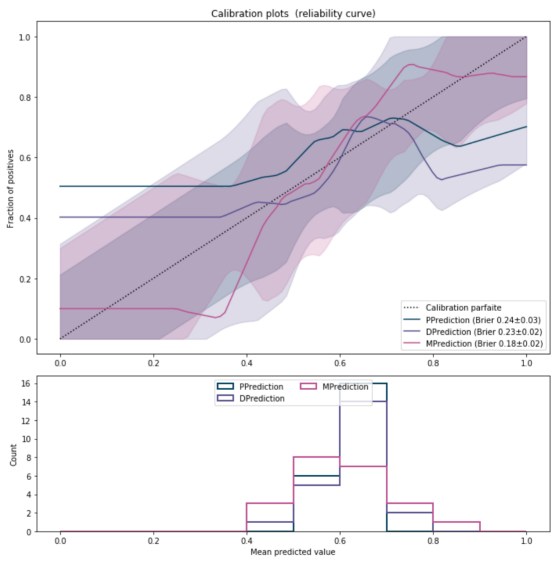
Microscopie

	precision	recall	f1-score	support
Benign	0.76±0.18	0.52±0.24	0.62±0.18	86.00±2.01
Malignant	0.75±0.11	0.90±0.09	0.82±0.06	137.00±2.00
accuracy	0.75±0.08	0.75±0.08	0.75±0.08	0.75±0.08
macro avg	0.76±0.08	0.71±0.10	0.72±0.11	223.00±0.46
weighted avg	0.75±0.06	0.75±0.08	0.74±0.09	223.00±0.46

```
In [23]: # Calibration
```

```
In [24]: p_inputs['PPrediction_Prediction'] = p_inputs['Prediction_Predict
ion']
    p_inputs['Prediction_Probability'] = p_inputs['Prediction_Probab
ility']
    p_inputs['DPrediction_Prediction'] = d_inputs['Prediction_Predict
ion']
    p_inputs['DPrediction_Probability'] = d_inputs['Prediction_Probab
ility']
    p_inputs['MPrediction_Prediction'] = m_inputs['Prediction_Predict
ion']
    p_inputs['MPrediction_Probability'] = m_inputs['Prediction_Probab
ility']

figure = Views.reliability_curve(p_inputs, 'LesionEncode', ['PPrediction', 'DPrediction', 'MPrediction'])
figure.savefig(f'{prefix}.svg')
figure.show()
```



Fusion

```
In [25]: inputs = pandas.concat([p_inputs, d_inputs, m_inputs], axis=0)
inputs = inputs.reset_index(drop=True)
# Save
I0.save(inputs, f'Low_{prefix}.pickle')
```

Cumulative

```
In [26]: inputs = IO.load(f'Low {prefix}.pickle')
In [27]: all image = [True] * len(inputs.index)
         single image = inputs['Modality'] == 'Photography'
         Data.build bags(inputs, single image, 'ID Lesion', all image, 'ID
         _Lesion', f'Prediction_{Tools.PROBABILITY}')
         inputs = inputs[single image].reset index(drop=True)
         high folds = Tools.generate folds(([1], [2]), validation)
In [28]:
         modality = MultimodalClassifier(method='modality', metric=f1 scor
         modality rev = MultimodalClassifier(method='modality', metric=f1
         score. ordered=False)
         classe = MultimodalClassifier(method='modality class', metric=f1
         score)
         classe rev = MultimodalClassifier(method='modality class', metric
         =f1 score, ordered=False)
         modality ones = MultimodalClassifier(method='modality', metric=f1
         score, from zero=False)
         modality rev ones = MultimodalClassifier(method='modality', metri
         c=f1 score, ordered=False, from zero=False)
         classe ones = MultimodalClassifier(method='modality class', metri
         c=f1 score, from zero=False)
         classe rev ones = MultimodalClassifier(method='modality class', m
         etric=f1 score, ordered=False, from zero=False)
```

```
In [29]: | Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, modality, 'simple increase
         increase')
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, modality rev, 'simple incre
         ase decrease')
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, classe, 'double increase in
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, classe rev, 'double increas
         e decrease')
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, modality ones, 'simple decr
         ease increase')
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, modality rev ones, 'simple
         decrease decrease')
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label_encode': 'LesionEncode'}, classe ones, 'double decrea
         se increase')
         Tools.evaluate(inputs, {'datum': f'Prediction {Tools.PROBABILIT
         Y}', 'label encode': 'LesionEncode'}, classe rev ones, 'double de
         crease decrease')
         IO.save(inputs, f'High_{prefix}.pickle')
```

Evaluation achieved!

Score

```
In [30]: from IPython.display import HTML
         from IPython.display import display
         inputs = I0.load(f'High {prefix}.pickle')
         diagnosis encoder = OrderedEncoder().fit(['Benign', 'Malignant'])
         # ROC Curve
         ViewsTools.plot size((8,8))
         name = f'simple increase increase'
         display(HTML(ViewsTools.dataframe renderer([Views.report(inputs,
         {'label encode': 'LesionEncode', 'eval': name}, diagnosis_encode
         r) 1.
         title=[f'Test - {name}'])))
         figure = Views.steps visualization(inputs, {'label encode': 'Lesi
         onEncode', 'eval': name})
         figure.savefig(f'{prefix} {name}.svg')
         figure.show()
         name = f'simple increase decrease'
         display(HTML(ViewsTools.dataframe_renderer([Views.report(inputs,
         {'label_encode': 'LesionEncode', 'eval': name}, diagnosis encode
         r)],
         title=[f'Test - {name}'])))
         figure = Views.steps visualization(inputs, {'label encode': 'Lesi
         onEncode', 'eval': name})
         figure.savefig(f'{prefix} {name}.svg')
         figure.show()
         name = f'double increase increase'
         display(HTML(ViewsTools.dataframe renderer([Views.report(inputs,
         {'label encode': 'LesionEncode', 'eval': name}, diagnosis encode
         r) 1.
         title=[f'Test - {name}'])))
         figure = Views.steps visualization(inputs, {'label encode': 'Lesi
         onEncode', 'eval': name})
         figure.savefig(f'{prefix} {name}.svg')
         figure.show()
         name = f'double increase decrease'
         display(HTML(ViewsTools.dataframe_renderer([Views.report(inputs,
         {'label encode': 'LesionEncode', 'eval': name}, diagnosis encode
         r)],
         title=[f'Test - {name}'])))
         figure = Views.steps visualization(inputs, {'label encode': 'Lesi
         onEncode', 'eval': name})
         figure.savefig(f'{prefix} {name}.svg')
         figure.show()
         name = f'simple decrease increase'
         display(HTML(ViewsTools.dataframe_renderer([Views.report(inputs,
         {'label encode': 'LesionEncode', 'eval': name}, diagnosis encode
         r)],
         title=[f'Test - {name}'])))
```

```
figure = Views.steps visualization(inputs, {'label encode': 'Lesi
onEncode', 'eval': name})
figure.savefig(f'{prefix} {name}.svg')
figure.show()
name = f'simple decrease decrease'
display(HTML(ViewsTools.dataframe renderer([Views.report(inputs,
{'label encode': 'LesionEncode', 'eval': name}, diagnosis encode
r)],
title=[f'Test - {name}'])))
figure = Views.steps visualization(inputs, {'label encode': 'Lesi
onEncode', 'eval': name})
figure.savefig(f'{prefix} {name}.svg')
figure.show()
name = f'double decrease increase'
display(HTML(ViewsTools.dataframe renderer([Views.report(inputs,
{'label encode': 'LesionEncode', 'eval': name}, diagnosis_encode
r)],
title=[f'Test - {name}'])))
figure = Views.steps visualization(inputs, {'label encode': 'Lesi
onEncode', 'eval': name})
figure.savefig(f'{prefix} {name}.svg')
figure.show()
name = f'double decrease decrease'
display(HTML(ViewsTools.dataframe renderer([Views.report(inputs,
{'label_encode': 'LesionEncode', 'eval': name}, diagnosis_encode
r)],
title=[f'Test - {name}'])))
figure = Views.steps visualization(inputs, {'label encode': 'Lesi
onEncode', 'eval': name})
figure.savefig(f'{prefix} {name}.svg')
figure.show()
```

Test - simple_increase_increase

	precision	recall	f1-score	support
Benign	0.53±0.36	0.09±0.22	0.16±0.21	86.00±2.01
Malignant	0.62±0.11	0.95±0.11	0.75±0.06	137.00±2.00
accuracy	0.62±0.09	0.62±0.09	0.62±0.09	0.62±0.09
macro avg	0.58±0.20	0.52±0.07	0.46±0.12	223.00±0.46
weighted avg	0.59±0.20	0.62±0.09	0.52±0.13	223.00±0.46

Test - simple_increase_decrease

	precision	recall	f1-score	support
Benign	0.50±0.35	0.08±0.20	0.14±0.19	86.00±2.01
Malignant	0.62±0.11	0.95±0.11	0.75±0.06	137.00±2.00
accuracy	0.61±0.08	0.61±0.08	0.61±0.08	0.61±0.08
macro avg	0.56±0.19	0.52±0.05	0.45±0.10	223.00±0.46
weighted avg	0.57±0.19	0.61±0.08	0.52±0.12	223.00±0.46

Test - double_increase_increase

	precision	recall	f1-score	support
Benign	0.27±0.09	0.03±0.18	0.06±0.12	86.00±2.01
Malignant	0.61±0.10	0.94±0.12	0.74±0.06	137.00±2.00
accuracy	0.59±0.08	0.59±0.08	0.59±0.08	0.59±0.08
macro avg	0.44±0.09	0.49±0.03	0.40±0.06	223.00±0.46
weighted avg	0.48±0.14	0.59±0.08	0.48±0.11	223.00±0.46

Test - double_increase_decrease

	precision	recall	f1-score	support
Benign	0.27±0.09	0.03±0.18	0.06±0.12	86.00±2.01
Malignant	0.61±0.10	0.94±0.12	0.74±0.06	137.00±2.00
accuracy	0.59±0.08	0.59±0.08	0.59±0.08	0.59±0.08
macro avg	0.44±0.09	0.49±0.03	0.40±0.06	223.00±0.46
weighted avg	0.48±0.14	0.59±0.08	0.48±0.11	223.00±0.46

Test - simple_decrease_increase

	precision	recall	f1-score	support
Benign	0.69±0.28	0.52±0.25	0.60±0.22	86.00±2.01
Malignant	0.74±0.11	0.85±0.10	0.79±0.05	137.00±2.00
accuracy	0.73±0.07	0.73±0.07	0.73±0.07	0.73±0.07
macro avg	0.72±0.15	0.69±0.10	0.69±0.12	223.00±0.46
weighted avg	0.72±0.11	0.73±0.07	0.72±0.09	223.00±0.46

Test - simple_decrease_decrease

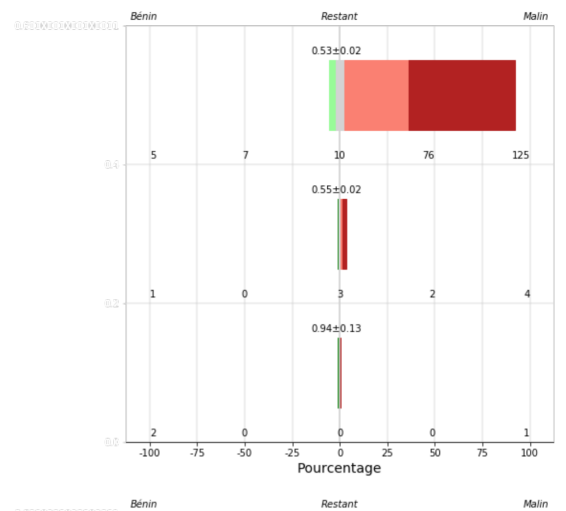
	precision	recall	f1-score	support
Benign	0.70±0.29	0.50±0.26	0.59±0.21	86.00±2.01
Malignant	0.73±0.11	0.87±0.10	0.80±0.05	137.00±2.00
accuracy	0.73±0.06	0.73±0.06	0.73±0.06	0.73±0.06
macro avg	0.72±0.14	0.68±0.09	0.69±0.11	223.00±0.46
weighted avg	0.72±0.11	0.73±0.06	0.71±0.08	223.00±0.46

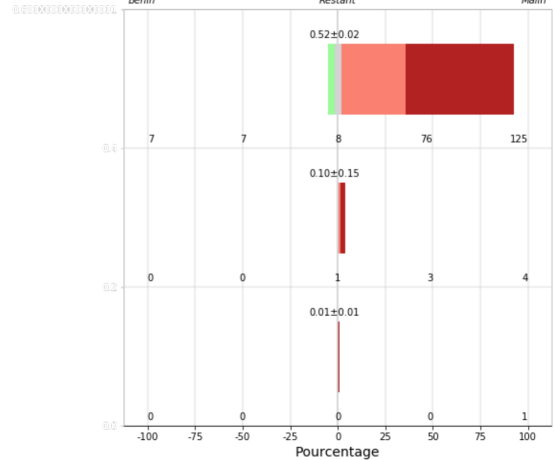
Test - double_decrease_increase

	precision	recall	f1-score	support
Benign	0.73±0.40	0.41±0.32	0.52±0.32	86.00±2.01
Malignant	0.71±0.13	0.91±0.12	0.79±0.08	137.00±2.00
accuracy	0.71±0.11	0.71±0.11	0.71±0.11	0.71±0.11
macro avg	0.72±0.23	0.66±0.14	0.66±0.19	223.00±0.46
weighted avg	0.72±0.21	0.71±0.11	0.69±0.16	223.00±0.46

Test - double_decrease_decrease

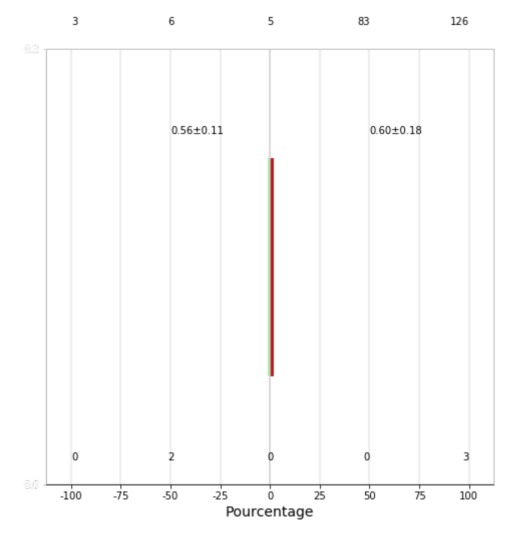
	precision	recall	f1-score	support
Benign	0.71±0.40	0.40±0.32	0.51±0.32	86.00±2.01
Malignant	0.70±0.13	0.90±0.13	0.79±0.08	137.00±2.00
accuracy	0.70±0.12	0.70±0.12	0.70±0.12	0.70±0.12
macro avg	0.71±0.23	0.65±0.14	0.65±0.18	223.00±0.46
weighted avg	0.70±0.21	0.70±0.12	0.68±0.16	223.00±0.46





Bénin Restant Malin

0.62±0.04 0.51±0.08



Bénin Restant Malin

0.62±0.04 0.50±0.08

