MLViz: How to guide

MLViz is a suite of interactive visualisation tools to assist the exploratory analysis of high-dimensional datasets. It is based on the Bokeh and sci-kit learn libraries and is developed to integrate directly with Jupyter notebooks so it can be used in with exisiting exploratory analysis workflows.

This document introduces the tools (as of V0.1) and has some hints on how to use the tools.

1 The Univariate feature selection tool

The feature selection tool computes univariate statistical tests of each feature and allows the user to select a feature subset based on these tests. When there are a large number of features it is often unfeasible to use all the features to build a machine learning model or perform exploratory analysis of and this tool can help the users prioritise the features to investigate further with the other tools in the MLviz library.

A list of the most important features (as determined by the univariate statistical tests) can be extracted as a list (see fig. 2) so they can be directly fed into different tools in the MLViz library.

2 The HDViz tool

HDViz is a tool which supports the exploratory analysis of high-dimensional datasets. It allows the user to generate a 2-dimensional representation of their dataset and extract clusters of datapoints (instances which are 'close' in the high dimensional space). The extracted clusters of datapoints can then be investigated using other visualisation tools to glean insights from the data set.

2.1 Bokeh tools

The HDViz tool is configured with standard Bokeh tools to enable the user to navigate the data, the available tools are outlined in tab. 1. They allow the user to pan the data, zoom and select different instances of the data set.

¹Andrea Dal Pozzolo, Olivier Caelen, Reid A. Johnson and Gianluca Bontempi. Calibrating Probability with Undersampling for Unbalanced Classification. In Symposium on Computational Intelligence and Data Mining (CIDM), IEEE, 2015.

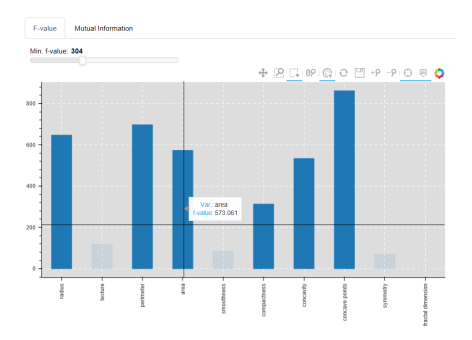


Figure 1: Overview of the UVFS tool. The horizontal axis displays each feature and the vertical axis the calculated statistical test of that feature (F-value or mutual information, shown by the highlighted tab in the top left corner). The user has the ability to select (dark shading) or de-select (light shading) using the Bokeh 'Tap tool' (Top right tool bar). A user can also threshold the features by selecting the 'minimum' value of the statistical test (using the slider, top left) to de-select features of potential low-importance.

```
tool.get_selected_features(metric='f-value')
Getting features selected by f-value
```

Figure 2: Extracting selected features from the UVFS tool. This method can be used to extract the names of the features selected in the UVFS tool (i.e., any features which are shaded in dark blue in fig. 1).

2.2 Using the tool

The tool is initialised by calling the HDViz class, passing it the appropriate arguments (i.e., as a minimum, the training set as a pandas dataframe), this command is shown in fig. 4. It is important to assign this class to a variable so the instance attributes of HDViz can be accessed later by the user.

2.2.1 Generating the low dimensional representation

Once the plot has loaded the user will be presented with an empty figure panel with various buttons for interaction (see previous section). To get a 2-dimensional representation

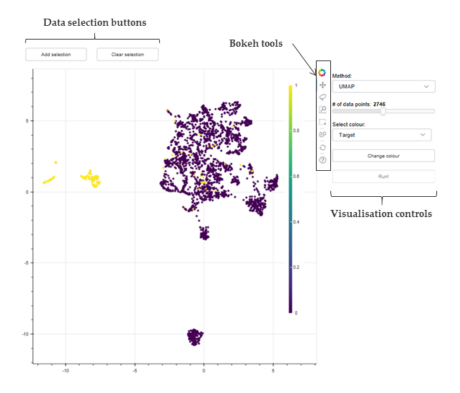


Figure 3: Overview of the HDViz tool. Graphic showing the 2-dimensional representation of the credit card data set¹ generated using the HDViz tool. Buttons on the right hand side are used to control the plot visualisations and parameters for performing the dimensionalty reduction and buttons along the top row are used in extracting 'brushed' data.

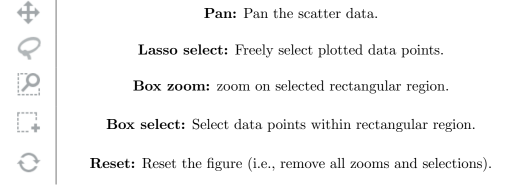


Table 1: The different Bokeh tools included with the HDViz tool. The left column displays the graphic representing the tool and the right column displays its name (bold) and a short description of its function.

Figure 4: Initialisation of the HDViz plot in a Jupyter notebook. When calling the HDviz class it is important to assign it to a variable (in this case 'HD_plot'). It takes one argument as default (a pandas dataframe of the training data) but can also take the target and the location ('url' kwarg) of the notebook for output.

of the data set the user must:

- Select the dimensionality reduction method to use (PCA, t-SNE or UMAP).
- Select the number of (randomly selected) data points to embed (by default, equal to half the training set).
- Click the 'Run!' button to begin the calculation.

The dimensionality reduction algorithms can take a long time to run and it is important to note the computational complexity for each algorithm (see appendix). A good default is the UMAP algorithm as it scales linearly with the number of training instances and generally provides fairly robust embeddings. We recommended trying an embedding with a sample of the training set (e.g., 2500 data points) before running the full data set.

2.2.2 Exploring and extracting data

Once the algorithm has run successfully the 'Run!' button is disabled and the figure should now look like the one displayed fig. 3. The user can now select to colour the data points by different features or the target (using the 'Visualisation controls' located to the right hand side of the plot). Additionally, at this stage the Bokeh tools can be used to select different clusters of data points and then add them (ready for extraction) using the 'Add selection' button, located on the top row. This will append the brushes to a list so all user brushes can be accessed in the jupyter notebook in which the plot is embedded. The 'Clear selections' button removes all of the brushes previously added with the 'Add selection' button.

Once a sufficient number of brushes have been extracted (5 is a reasonable maximum number, or visualisation is inhibited) the data can be extracted by calling the 'get_user_brushes()' method of the HDViz instance (shown in fig. 5). The returned data (stored in a pandas dataframe) also have a new column 'cluster_number' which shows in which user brushing the datapoint was part of (i.e., their first, second and so on brush). This data can now be fed into other tools in the MLViz pipeline to enable

3 Draughtsman plot

The draughtsman plot (a.k.a a pairplot) can be used in syenergy with the HDviz tool to investigate the clusters observed in the HDViz plot in the original feature space. It allows

```
X, y = HD_plot.get_brushed_data()
```

Figure 5: Method for extracting the brushed data from the HDViz plot. This graphic shows the method a user must call to extract the brushed data, it returns the brushed data and the targets of these brushed instances.

users to investigate correlations and class seperability in the original feature space.

3.1 Using the tool

The tool is initialised by calling the DraughtPlot class, passing it the appropriate arguments (i.e., as a minimum, the training set and target values or the output of the HDViz get_brushed_data method), this command is shown in fig. 7. The user can select which features to display in the plot using the 'features' kwarg, if none are provided the first five features in the training set are plotted.

Once initialised, users can control the colour of the data points (corresponding to the values of a selected feature), zoom in and brush data points (highlighting the same instances in different panels) on the scatter plots. When brushing, the computed values (top-right off diagonal panels) are updated to reflect the metrics for the selected data points only.

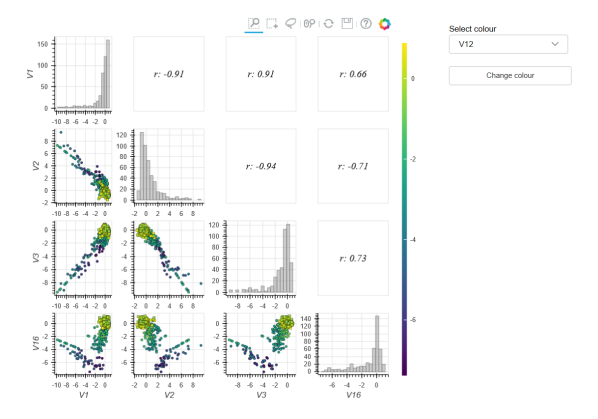


Figure 6: The Draughtsman plot tool with example data. Users can select which features to plot when calling the function and data points can be coloured by a user specified dimension. The upper off-diagonal panels shows the pearsons correlation coefficient for the two relevant features and the lower off-diagonals show a scatter plot of the training instances for the two relevant features. Panels on the diagonal display a histogram training instances for the feature on the horizonal axis.

DMPlot = DraughtPlot(X, y, features = columns[7:])

Figure 7: Initiating the Draughtsman plot. X and y can either be the brushed data returned by the HDViz get_brushed_data method or the raw feature matrix and target values. The features kwarg argument is a list of features to include in the list. We recommend that the number of features is kept to approximately five or less. If a binary target (y) is provided instances with a target equal to 1 are plotted as squares and instances with a target of 0 are plotted as circles.

A Dimensionality reduction techniques

The HDviz tool relies upon using dimensionality reduction techniques, consistent with the sklearn API, to produce a 2-dimensional embedding of the high-dimensional dataset. There are three different methods currently implemented, all of which have different advantages and disadvantages.

A.1 PCA

Principle component analysis (PCA) is, arguably, the most popular dimensionality reduction technique. It works by projecting the high-dimensional data into a lower dimensional hyper plane, returning a smaller feature set whose features are a linear combination of the old features. Typically it is used to make a large feature set more managable, by lowering its dimensionality while preserving the maximum amount of variance (achieved by projecting the data into a lower dimensional hyperplane which lies closest to the data). It is not, however, that effective at embedding high-dimensional data sets into two dimensions as the information loss in discarding dimensions in the way is generally too large. It is included in MLViz as a benchmark algorithm and for rapid testing but is not generally recommend for wide-scale use.

- Scalability: Exact solution: Slow, $O(m \times n^2) + n^3$. However, Randomized PCA scales as $O(m \times d^2) + d^3$ (where d is the number of dimensions to embed into and n the number of features). Sklearn uses Randomized PCA automatically for large problems.
- **Performance:** Generally performs poorly when d is much less than n. Not practically used to visualise high-dimensional data.

A.2 t-SNE

t-SNE is currently the go to method for embedding high dimensional data into two dimensions, it excels in the image domain (c.f., MNIST). It generates a similarity measure between datapoints in high-dimensional space and then distributes the datpoints in the lower dimensional space where another similarity measure is as close as possible to the high-dimensional one.

- Scalability: Full: $O(m \times n^2)$. But sklearn, as default, uses an approximation which scales $O(m \log(n)n)$.
- **Performance:** Known to perform well on image data sets.

t-SNE resources:

- L.J.P. van der Maaten and G.E. Hinton. Visualizing High-Dimensional Data Using t-SNE. Journal of Machine Learning Research 9(Nov):2579-2605, 2008
- 2. Wattenberg, et al., "How to Use t-SNE Effectively", Distill, 2016. http://doi.org/10.23915/distill.0000

A.3 UMAP

UMAP inherts from sklearn classes and is therefore consistent with its API but it is not currently part of the sklearn package and is still under development. It should be considered as an experimental feature.

UMAP is a less mature technique than t-SNE but has the potential to overtake it as the household technique for visualising high dimensional data sets. It scales much better than t-SNE, there is a direct mapping from the high-dimensional to low dimensional space and works effectively when embedding into an arbitray number of dimensions (where as t-SNE does not perform well when m > 3).

• Scalability: Very fast: O(kn)

• Performance: Known to perform well on image datasets.

UMAP resources:

1. McInnes et al., (2018). UMAP: Uniform Manifold Approximation and Projection. Journal of Open Source Software, 3(29), 861. jhttps://doi.org/10.21105/joss.00861j.