ADENINE — A Data Exploration plpeline

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Abstract

In this paper we introduce Adenine, a Python framework for data exploration. The main goal of Adenine, is twofold: helping researchers and data scientists achieving a first and quick overview on the main structures underlying their data and choosing the most suitable unsupervised learning pipeline for the problem at hand. This software tool encompasses state of the art techniques for: missing values imputing, preprocessing, dimensionality reduction and clustering tasks. Adenine exploits both process and thread level parallelism and it is capable of generating nice and clean publication-ready figures along with a quantitative description of the algorithms performance. Adenine is released under FreeBSD license and it can be downloaded from http://slipguru.github.io/adenine/.

Keywords: Data exploration, unsupervised learning, RNA-Seq gene expression

1. Introduction

Data exploration can be an extremely helpful starting point for many data analysis tasks. Researchers and data scientists are often asked to extract meaningful information from collections of complex and possibly high-dimensional, data coming from heterogenous contexts. For instance, in biomedical scenarios, physicians are likely to be interested in answering some biological questions starting from a set of observations collected from a pool of subjects enrolled in a study. Possible investigations can be: is there any relevant stratification among subjects? or is it possible to discriminate between case and control samples from my observations?. The information needed to answer such questions may be immediate, non-trivial to extract or even completely absent.

In these situations, a preliminary data exploration step can be not only good practice, but also a mandatory starting point for further investigations. In this context, several machine learning and data mining techniques were developed over the years. Among those we focus on the four most common tasks: (i) missing values imputing, (ii) data preprocessing, (iii) dimensionality reduction and (iv) unsupervised clustering.

2. Implementation

From an implementative standpoint, Adenine is built around the concept of *pipeline*, that is a sequence of the four fundamental steps mentioned in Section 1 (see Figure 1).

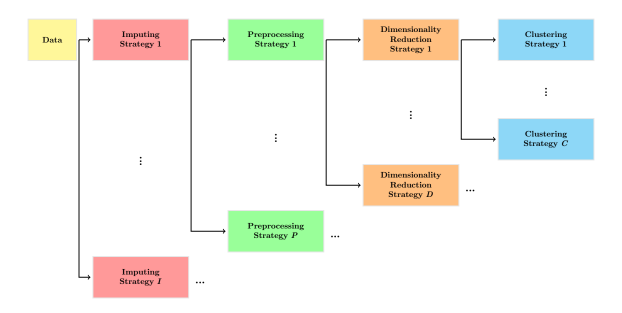


Figure 1: A schematic representation of the Adenine workflow. The list of building blocks available for each step is summarized in Table 1.

For each step, a fair number of off-the-shelf algorithms are available (see Table 2). The vast majority of their implementations is inherited, or extended, from scikit-learn (Pedregosa et al., 2011).

In order to perform exploratory analysis of large datasets, Adenine can take advantage of different parallel computing paradigms. For instance, its pipelines are designed to be independent from each other, therefore they all run in parallel as separate Python processes on different cores. Moreover, since Adenine makes large use of numpy and scipy, it automatically benefits from their bindings with optimized linear algebra libraries (such as OpenBLAS¹, or Intel[®] MKL).

3. Experiments and results

To assess the quality of the obtained results, we tested Adenine on a set of synthetic and real dataset.

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Table 1: Pipelines building blocks and relative references (which are not reported when the definition is given in Section 2).

Step	Algorithms	Ref.
Imputing	Mean	
1 0	Median KNN	(Troyanskaya et al., 2001)
Preprocessing	Recentering Standardize Normalize MinMax	(110) and any a co car, 2001)
Dimensionality reduction	Principal component Analysis (PCA) Incremental PCA Randomized PCA Kernel PCA Isomap Locally linear embedding Spectral embedding Multidimensional scaling t-Distributed Stochastic Neighbor Embedding (t-SNE)	(Jolliffe, 2002) (Ross et al., 2008) (Halko et al., 2011) (Schölkopf et al., 1997) (Tenenbaum et al., 2000) (Roweis and Saul, 2000) (Ng et al., 2002) (Borg and Groenen, 2005) (Van der Maaten and Hinton, 2008)
Clustering	K-means Affinity propagation Mean Shift Spectral Hierarchical	(Bishop, 2006) (Frey and Dueck, 2007) (Comaniciu and Meer, 2002) (Shi and Malik, 2000) (Friedman et al., 2001)

4. Conclusions

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^{1.} http://www.openblas.net/

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