Machine Learning Engineer Nanodegree

Capstone Proposal

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Proposal

Domain Background

Peptidic natural products (PNPs) are small bioactive compounds consisting of amino acids connected via peptide bonds. A PNP may be represented as a graph with amino acids as nodes and bonds as edges. These graphs have either linear, cyclic, or more complex structure. PNPs are important for medicine since many of them are active against bacteria i.e. could be antibiotics. One of the main ways to study PNPs is through mass spectrometry. For each PNP you can get a spectrum (intensity as a function of the mass-to-charge ratio) or a few by examining it in a black box -- mass spectrometer. These spectra can further be compared against databases of previously characterized compounds using computational methods such as DEREPLICATOR (Mohimani H. et al., 2017).

Understanding which spectra correspond to which types of PNPs structure will significantly speed up the DEREPLICATOR since it will be possible to search through smaller sets (cyclic spectra only against cyclic compounds and linear only against linear). At the same time it will increase precision of the algorithm because initial DEREPLICATOR compares any spectrum with any compound and thereby can get such false positive matching as linear spectrum to nonlinear compound and nonlinear spectra to linear compound (not present in an improved algorithm). Also knowledge about the structure itself (separately from DEREPLICATOR) tells scientists some biological properties of the compound represented by its own spectrum. Cyclic PNPs are more stable and biologically more active on average so we can focus on studying of only such spectra thereby saving our resources.

Problem Statement

The problem of this Capstone project is to categorize PNPs spectra into spectra corresponding to cyclic compounds and linear. Thus the program requires spectrum of the unknown compound as input and defines type of the compound structure as output.

Datasets and Inputs

There is already a huge amount of publicly available mass spectra of natural products. It turned out to be possible to detect natural products by their mass spectra and also find new ones missing in the database using a high-throughput technology built on computational algorithms such as DEREPLICATOR.

I'm going to use this **one hundred million tandem mass spectra** in the Global Natural Products Social (GNPS) molecular networking infrastructure (Wang M. et al., 2016) to select peptide compounds and categorize them into cyclic and non-cyclic by Machine learning algorithms. The labels can be taken from molecular structures from GNPS library (trustworthy labels manually obtained by biologists) or from highly-reliable DEREPLICATOR identifications. In both cases it's **several hundred structures** (about 200 cyclic and 100 non-cyclic structures) and about a **thousand spectra** related to them (3-5 different spectra for the structure on average).

Solution Statement

It's Supervised learning task because example input-output (namely spectrum-structure) pairs exists. I will start with the simplest model so the baseline is some simple **Neural network** (most likely **CNN** to utilize a multi-dimensional data). The advantage of Neural networks approach is the possibility of non-linear models with respect to the features. I plan to try various data

representations and then do some preprocessing steps. There are two ways to work with these continuous space of input data: discretize the raw spectra or directly approximate them by functions. Of course I also will try a different models (various layers and etc.) and most Keras optimizers. The solution can be measured by common metrics such as accuracy, precision, recall and more since there is labeled data.

Benchmark Model

A good result that relates to the domain of Natural products identification would be less elapsed time and less FP at the same time obtained by **target matching DEREPLICATOR** (cyclic spectra against cyclic compounds and linear against linear) than by current DEREPLICATOR pipeline. It will mean that the model correctly classify the spectra by their structures into two groups. Thus the benchmark model is **current DEREPLICATOR** results.

Evaluation Metrics

Accuracy, precision, recall, F1 score and FP as the primary metric are a good choice for evaluation metrics that can be used to quantify the performance of both the benchmark model and the target matching DEREPLICATOR. Here FP means that DEREPLICATOR got a structure that actually doesn't match input spectrum.

Also we can simply compare results of the solution on test set from GNPS library using **FP** metric where false means that spectrum corresponds to other cyclicality than our ML algorithm got.

Project Design

I will be programming in Python 3 using pandas, Num Py, scikit-learn and mainly Keras.

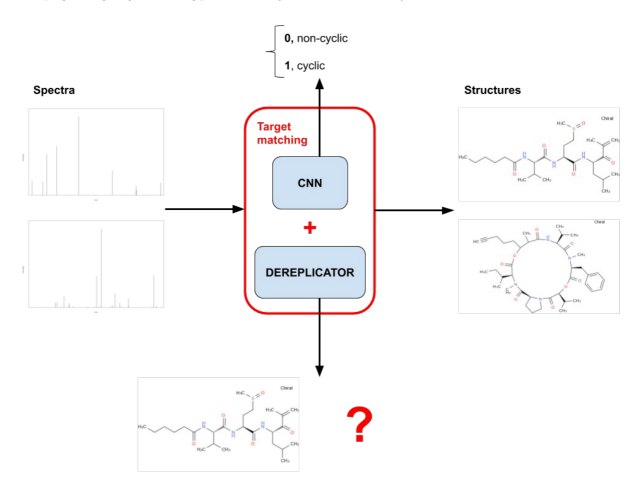


Fig. 1. Linear and cyclic spectra are passed to CNN, DEREPLICATOR, and sequentially to both of them in Target matching DEREPLICATOR. CNN outputs types of its compounds, DEREPLICATOR could identify only linear PNP, and together both structures.

The workflow for approaching a solution given the problem includes

- **Collect** the data. Choose peptide not complex compounds from GNPS Public Spectral Library and also the same highly-reliable DEREPLICATOR identifications.
- **Preprocess** the data. It's necessary to think thoroughly here about a representation of the input spectra since w hat features will consider our algorithm completely depends on it. It can be some tiny step discretization of raw spectra or spectra approximation by basis functions like RBF. Maybe it will be meaningful to use some data augmentation to increase the set of input data. After that when I understand the data I will identify what kind of preprocessing is needed: scaling, normalization and so on.
- **Split** the data into training, validation and test sets such that both linear and cyclic compounds fall into each of these sets in acceptable proportions.
- Choose, train and tune the model. The baseline is CNN using two various input data representations with a few first come to mind preprocessing steps. Get some intuitions about how these networks work on spectra data by testing them and plotting some scores, change initial model varying layers and other hyperparameters, use different optimizers and also try any more models.
- Evaluate the solution. After getting two groups of spectra by approved network run DEREPLICATOR for cyclic spectra against cyclic compounds and linear against linear separately. Compare FP and elapsed time for these results and for DEREPLICATOR on full set of spectra. Also compute FP for approved network without considering DEREPLICATOR pipeline.