



University of Glasgow | School of  
Computing Science

Honours Individual Project Dissertation

# PEP2PATH v2

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## Abstract

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# 1 | Introduction

One particular problem of interest for biologists is the identification of new biological products with novel properties. Until relatively recently, this process was done manually by trained experts. But with the recent advent of the field of bioinformatics, we can now bring immense computational resources to bear on problems like this, saving precious expert time.

Pep2Path (Medema et al. (2014)) is a tool that aims to accelerate drug identification by matching mass-spectrometry outputs to identified biosynthetic gene clusters and in the original paper the authors demonstrate robust benchmarking results. However, despite the utility of Pep2Path, it is implemented as a few monolithic scripts, with little room to revise elements of it in accordance with new developments and research without a complete rewrite. In this paper we present a small suite of software tools implementing a platform for functionality like Pep2Path, and the software process by which this software was designed and created.

## 1.1 Aims

The aims of the project are to:

- Implement a set of software tools that can achieve similar functionality to the original Pep2Path, subject to correctness testing.
- Achieve a degree of flexibility in the implementation such that this software can be responsively updated to changes in its environment.
- Follow good software practises in the implementation such as loose coupling such that we can achieve software reuse, a cornerstone of software development.

## 1.2 Motivation

Various biological compounds, are formed from assembly-chains of smaller compounds, such as peptides, formed from amino acids. If we look at this problem as a computer scientist might, the first thing to come to mind in identifying useful drugs within this space might be an exhaustive search of all combinations of molecule. (Of course, for other molecules questions of topology and structure come into question...) This quite quickly runs into problems, as with the addition of each component to a chain of biological molecules, the chemical space grows exponentially. For example, if we consider the relatively limited case of the twenty proteinogenic amino acids and a peptide of length 4, there are  $20^4$  possible combinations... Around 160,000! We would then have to identify which, if any, of these have useful properties.

Without some heuristic (for example being able to infer properties from chemical composition), an exhaustive search of the chemical space is impossible. Instead we identify biological products produced in the wild, by living organisms – so-called ‘natural products’. From an evolutionary perspective, we expect that if organisms are expending resources on producing a particular biological product, then it confers a survival advantage. For example, a strain of bacteria might produce an antibiotic that kills another competitor species of bacteria – which we can then use ourselves to kill a potentially deadly strain of bacteria. Natural products produced in this way to

fulfil an auxiliary function not directly related to the growth or reproduction of the organism are known as 'secondary metabolites'.

To identify potentially interesting natural products we could, for example, analyse chemical structure for similarity to other interesting compounds, or by observing similar behaviour across several species which produce a particular product. However, we also want to match this natural product to the gene cluster – a Biosynthetic Gene Cluster, henceforth BGCs – which encodes it. One way of achieving this is to cluster together species that share a BGC and produce the natural product in question – this is a good indication that the pattern is meaningful and the BGC produces the natural product.

Mass spectrometry allows us to identify the chemical composition of compounds – by breaking up molecules and then measuring the mass lost, we can infer the *potential* composition of a compound by the masses of its components using a mass-translation table. Mass spectrometry readings, or amino acid sequences, are one of the key inputs to Pep2Path, which provides two algorithms, for two different natural product classes: Non-Ribosomal Peptides (henceforth NRPs) and Ribosomally-synthesized and Post-translationally-modified Peptides (henceforth RiPPs).

The other key input to each of the two Pep2Path algorithms is the gene sequence we are attempting to match our compound to. As their name suggests, RiPPs are synthesised by ribosomes, the cell organelle responsible for translating gene sequences to proteins (peptides made up of the 20 proteinogenic amino acids) and then modified post-translation by enzymes. As a result, they are directly encoded directly in an organism's gene sequence and we can retrieve their composition directly, as Pep2Path does by obtaining their 'six translation frames'. NRPs, by contrast, are instead synthesised by Non-Ribosomal Peptide Synthetases (henceforth NRPSes) which are independent of the ribosome and can introduce non-proteinogenic amino acids. These consist of several NRPS modules which form an assembly-line of amino acids, and are controlled by three domains – A (Adenylation), T (Thiolation) and C (Condensation). Of particular interest to us is the Adenylation domain, which controls which substrate is added to the chain of amino acids. These are not directly encoded into the gene and are therefore harder to predict, so we rely on an external tool – antiSMASH (Blin et al. (2017)) – for this purpose.

Taking these two sets of inputs, NRP2Path matches the potential chemical compositions generated by mass spectrometry analysis to BGC predictions using antiSMASH, whereas RiPP2Path matches these potential chemical compositions to the six-frame translation extracted from the genetic sequence data of the RiPP-producing BGCs. Pep2Path can then automatically score these two sets of data against one another, filling a key part in an automated drug-identification pipeline, where previously investigation of natural products would be based on hand-identification.

However, the original Pep2Path is written almost entirely using singular scripts, one for each algorithm. As a result, it breaches some important software design principles, by mixing concerns such as parsing input and generating sequence tags, and at times relies on data structures passed around the entire script. This design makes Pep2Path difficult to update, but there are a number of reasons why it might be advantageous to do so. For one, the scoring method the authors use is based on antiSMASH 2.0 (Blin et al. (2013)) the current version at the time. Since the release of Pep2Path, there have been antiSMASH versions up to 4. (Blin et al. (2017)) While the original Pep2Path scoring method is backwards-compatible, antiSMASH version 4.0 offers the SANDPUMA ensemble algorithm (Chevrette et al. (2017)) which collates the Stachelhaus Code and SVM predictors that antiSMASH 2.0 uses along with several others to offer a more accurate BGC prediction.

Furthermore, we may want to investigate product classes other than NRPS and RiPP, the classes of biological product on which Pep2Path operates – there are many others, including polyketides and alkaloids. (For an example as to why this might be useful, the pain-medication morphine is an alkaloid extracted from the opium poppy.) It would also be useful to integrate different algorithms for identifying biological products from sequence data (such as the RiPPQuest (Mohimani et al.

(2014)) method for lanthipeptides – a subtype of RiPP), decouple the mass spectrometry process from the Pep2Path process or otherwise update the scoring mechanism with new developments. We also expect more generally that having a more transparent and modular software design might lead to easier discovery of bugs or integration with other codebases.

## 2 | Background

### 2.1 Mass Spectrometry

Mass spectrometry is a common technique in chemistry for the identification of the chemical composition of a molecule by ionising it, causing it to break into fragments. This gives a series of ordered readings of mass and intensity – we can infer the gaps between mass peaks are due to the fragmentation of a molecule, and can then translate the mass being broken off into chemical composition using knowledge of molecular weights stored in a dedicated mass translation table. Traditionally, this was a problem hand-solved by chemists, but has long been a target for computational processes due to the ease of translation of these numerical processes.

Once mass shifts have been translated to potential compounds, these can be joined together, end-to-end into longer sequences of compounds – ‘sequence tags’. Then, computational resources can be used to easily mine a group of mass spectrometry readings for sequence tags. However, there will inevitably be noise in mass spectrometry readings, whether from measurement error or the breaking off of small fragments that don’t represent a significant part of the molecule. For this reason, two variables are introduced: an *intensity threshold* and a *mass tolerance*. We can firstly cut out all low-quality readings by cutting out all readings below a certain intensity; secondly, we can account for slight discrepancies in mass measurements by measuring mass shifts within some interval rather than taking the exact values from the mass table. Good values for these variables depend on the dataset in question, and a good general pair of values is currently unknown.

### 2.2 BGC Substrate Detection

#### 2.2.1 NRPs and antiSMASH

antiSMASH (antibiotics and Secondary Metabolite Analysis Shell) (Blin et al. (2017)) is a widely-used piece of bioinformatics software used for the automated labelling of BGCs and geonome-mining of secondary metabolites from raw sequence geonome sequence data extracted from bacteria, fungi or plants. It is usable both as a standalone program on MacOS or Linux and as a web-server and has gone through several revisions and is, at the time of writing, currently up to version 4.0. Therefore, it aggregates many of the latest developments in natural product research and offers many features for the handling of such sequence data and the annotation of its BGCs, outputting data in the standard GenBank format.

For the purposes of this paper, we are interested in antiSMASH’s ability to predict the substrate specificity of an NRPS’ adenylation domains, that is, give us a prediction for the chemical makeup of a particular NRP given the sequence data of the organism that produces it. The current version of antiSMASH uses the SANDPUMA ensemble algorithm to do this, which aggregates the results of several other predictive algorithms, including those in previous versions of antiSMASH (maintaining backwards-compatibility), to produce significantly better results. However SANDPUMA and antiSMASH 4.0 are relatively recent compared to Pep2Path, and instead we are interested in two key predictors of antiSMASH 2.0: the Stachelhaus Code, a



set of rules for comparing different adenylation domains created from empirical observation (Stachelhaus et al. (1999)) and a machine learning Support Vector Machine (SVM) based method.

Machine learning is frequently used in bioinformatics to extrapolate underlying trends from the vast quantities of data often involved, learning from a provided dataset useful understandings of programmer-selected features by performing some optimisation task. SVMs in particular are a classifier (that is, assign inputs to one of a set of discrete classes, in this case a substrate prediction) and attempt to draw a decision boundary separating those classes so as to minimise the distance between a certain number of the training datapoints plotted in a hyperplane. The SVM implementation for antiSMASH is provided by NRPSPredictor2, (Röttig et al. (2011)) a standalone piece of software which has since been integrated into the antiSMASH pipeline. One of the things that makes the original Pep2Path results robust is that it was tested on datasets NRPSPredictor2 had *not* been trained on, avoiding the introduction of bias and showing the generalisability of the Pep2Path method.

### 2.2.2 RiPPs, their six-frame translation and RiPPQuest

For RiPPs, the translation process is more direct than with NRPs, and does not require an external platform like antiSMASH to make substrate predictions. This process can be done via the 'six-frame translation'. DNA strands are made of long strings of four bases (Alanine, Glycine, Thymine and Cytosine) which can easily be represented and processed in computer systems in large numbers, and which bind in A-T and G-C pairs across the two strands. These bases, in groups of three known as codons, reliably encode amino acids in ways we can extract. However, when looking at a genome, we do not know where the first codon begins - it could begin at any of three positions. Then, the encoding could be done by either strand - we only store one strand, but we can retrieve its reverse complement by converting the base to its corresponding base and reversing the strand - for a total of six different encodings. This is the six-frame translation method we use for RiPPs.

Among RiPPs, there are many subtypes. One particular subclass of interest is the lanthipeptide, which is specifically targeted by the bioinformatics software RiPPQuest. (Mohimani et al. (2014)) The RiPPQuest method centres on the prediction of the 'LANC-like' domain in the genome, which is important for the biosynthesis of lanthipeptides in particular. It then centres its translation window around the LANC-like domain and begins searching using the six translation frames. As a result, it provides more accurate results for lanthipeptides in particular. We do not implement the RiPPQuest method here, and instead implement a more general method with looser assumptions in order to target all classes of RiPP, but we highlight this method as a possibility for future extension.

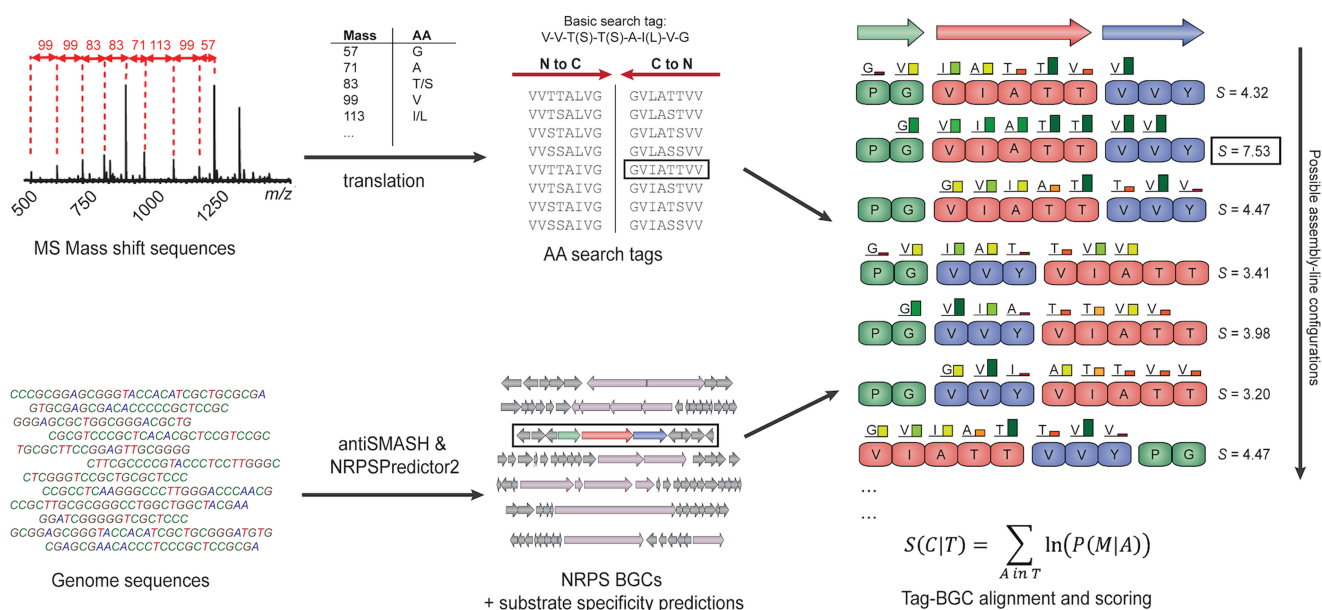
## 2.3 Pep2Path

Pep2Path (Medema et al. (2014)) provides two algorithms, NRP2Path and RiPP2Path. Both rely on the same principles of mass spectrometry. In the original Pep2Path source program, mass search tags can either be given directly or can be derived from a mass-shift sequence. They then extract relevant BGC information from a sequence, and compare potential sequence tags to potential BGCs using a different scoring function for each algorithm in order to enable finding the best match.

### 2.3.1 NRP2Path

NRP2Path first takes potential sequence tags and antiSMASH substrate specificity predictions. These sequence tags can be arranged either forwards or backwards, and within the prediction there are several different NRPS modules, which themselves can either be arranged forwards or

backwards and can be arranged in any order to make up the full BGC sequence. So in order to test all possible gene sequences, Pep2Path must generate all permutations of the cartesian products of the forwards and backwards modules (a total of  $n! \cdot 2^n$  different orderings for  $n$  modules). The sequence tag and the gene sequence are then aligned with one another and scored for their match, testing every possible alignment in order to find the best score.



**Figure 2.1:** An illustration of the NRP2Path algorithm. On the top is a mass spectrometry output being put through a mass translation table to produce a comprehensive list of potential sequence tags. On the bottom is BGC substrate specificity being extracted from raw sequence data by antiSMASH. These two sets of data are then used to compare every alignment of every sequence tag to every possible ordering of NRP blocks extracted from the BGC, using the abbreviated scoring function shown below. Figure adapted from the publicly-available Pep2Path paper and online documentation.

In order to compare a sequence tag to a BGC, NRP2Path uses its own scoring function loosely inspired by Bayes' Rule. We omit the details of its derivation here for brevity's sake, but they are available in the original paper.

$$S(C|T) = \sum_{A \in T} \ln \left( \frac{P(A) + c \cdot (I_{A,M}^\eta + x \cdot P(A))}{P(A) \cdot (1 + c \cdot (\sum_{A \in \mathbb{A}} (I_{A,M}^\eta + x)))} \right)$$

$S(C|T)$  is the score of a gene cluster given the sequence tag.  $A$  is the amino acid making up part of a tag, and  $\mathbb{A}$  is the total amino acid alphabet being used.  $c$  and  $x$  are parameters that express the degree of confidence that final probability relies on substrate specificity rather than prior probability, and a pseudo-count to correct for small sample size, with default values  $c = 1$  and  $x = 0.01$ , respectively.  $\eta$  is a regularisation term allowing for the increased penalisation of repeated mismatches.

$I_{A,M}$  is the average of two values calculated separately for the Stachelhaus Code and SVM predictions. The Code prediction is based on the closeness to the closest known NRPS module for NRPSpredictor2, whereas the SVM value is assigned between five evenly-spaced thresholds between 0 and 1 based on how closely the classes of the amino acid match (0 for no relation, 1 for exact match).

$P(A)$  is the prior probability of  $A$ , calculated like so:

$$P(A) = \frac{n(A) + \frac{k}{|\mathbb{A}|}}{\sum_{A \in \mathbb{A}} n(A) + k}$$

where  $k$  is another pseudocount with  $k=1$  and  $n(A)$  is the number of appearances within the NORINE database. (Caboche et al. (2008))

### 2.3.2 RiPP2Path

## 3 | Analysis/Requirements

### 3.1 Requirements

## 4 | Design

## 5 | Implementation

### 5.1 Language Choice

We chose to use Python for our implementation, as it is the *lingua franca* of scientific computing. It offers highly expressive language constructs and useful libraries, such as `itertools` (Python (0000)) offering many convenient ways to iterate through combinations, permutations and more, `NumPy` (NumPy (0000)) for high-speed numerical operations and `BioPython` (BioPython (0000)) for bioinformatics. All of these were used in the course of implementing our design. Additionally, a great volume of scientific software is already implemented in Python (indeed, the original `Pep2Path` is implemented using Python) and many scientists are familiar with Python already, making it easier to interface with the already extant body of work. Particularly, the client's codebase was implemented in Python, so this choice made for easier integration without the need for a language-spanning gateway. Other options like Julia (Julialang (0000)) exist, but given their lack of maturity when compared to Python we elected not to explore these options.

## 6 | Evaluation

## 7 | Conclusion



## A | Appendices

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