

Plugin Based Microbiome Analysis (PLUMA , formerly MIAMI) Version 2.0.0 - User Guide

Trevor Cickovski and Giri Narasimhan
School of Computing and Information Sciences
Florida International University
tcickovs@fiu.edu, giri@fiu.edu

<http://biorg.cis.fiu.edu/PluMA/>

Other Contributors from Florida International University:

Vanessa Aguiar-Pulido
Michael Campos
Mitch Fernandez
Wenrui Huang
Shamsed Mahmoud
Jingan Qu
Juan Daniel Riveros
Victoria Suarez-Ulloa
Camilo Valdez
Joseph R. Quinn

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Abstract

We present PLUMA, a lightweight and flexible package for constructing general software pipelines. PLUMA is designed to be infinitely extensible, allowing researchers to select a specific set of dynamically loaded plugins as sequential stages of their pipelines. These plugins can be implemented by either themselves or other users, in their language of choice.

We begin by introducing the key features of PLUMA, and follow with a discussion of how to download and install the latest version, compile, and run the software. We also include information on setting up configuration files that specify desired plugins for a pipeline, and how to extend PLUMA with new plugins in various programming languages. Finally, we conclude with a full pipeline example and a brief discussion of our envisioned future of PLUMA.

We distribute PLUMA under the MIT License, copyrighted by Florida International University.

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Chapter 1

Introduction

Software pipelines are applicable to any field of study and involve control flow execution as a series of *stages*, with an output of a given stage serving as an input to the next. As an example, Figure 1.1 shows a common metagenomics analysis pipeline, with an initial set of DNA sequences passing through a denoising stage that improves read quality, followed by clustering through some similarity or compositional metric, and finally labelling sequence clusters with the closest matching taxonomic unit [2].

Many pipelines have been developed by independent teams as standalone tools, although stages of individual pipelines could potentially be reused amongst one another. Particularly in the field of metagenomics analysis, there are many software pipelines that perform similar tasks and may even share stages that are still constructed independently, because there is no standardized framework for developing, testing and particularly *integrating* these stages. PLUMA is designed to address this need by providing a lightweight and generic back end that can execute these stages as dynamically loaded *plugins*, specified by a user through a configuration file or GUI.

We show the conceptual design of PLUMA in Figure 1.2, which follows a Problem-Solving Environment (PSE, [13]) using three conceptual tiers [8]: a compiled machine layer, a middle scripted layer, and an upper user layer. The user layer consists of *plugins* [3, 14] which can be developed by various users in the PLUMA community in multiple languages, and subsequently integrated to form pipelines. A user currently performs this integration through a configuration file, though we plan on adding a graphical interface later.

Plugins can be developed standalone using either the *scripted* or *computational* (compiled) interfaces to PLUMA. The scripted interface currently supports Python, Perl and R, and the compiled interface supports C++ for the CPU and CUDA for the GPU. We also provide a *PluginGenerator* (PluGen) module which can produce plugins that wrap existing tools on a user's machine (we will later show examples with Mothur [26] and Cytoscape [10]). Scripted plugins can also refer to each other, and we have a future goal of creating wrappers to the computational layer using SWIG [4]. We also will be expanding our set of supported languages, with Java as our next target.



Figure 1.1: An example metagenomics analysis pipeline. Each stage gets executed sequentially, with the output of a specific stage serving as input to a later stage of the pipeline.

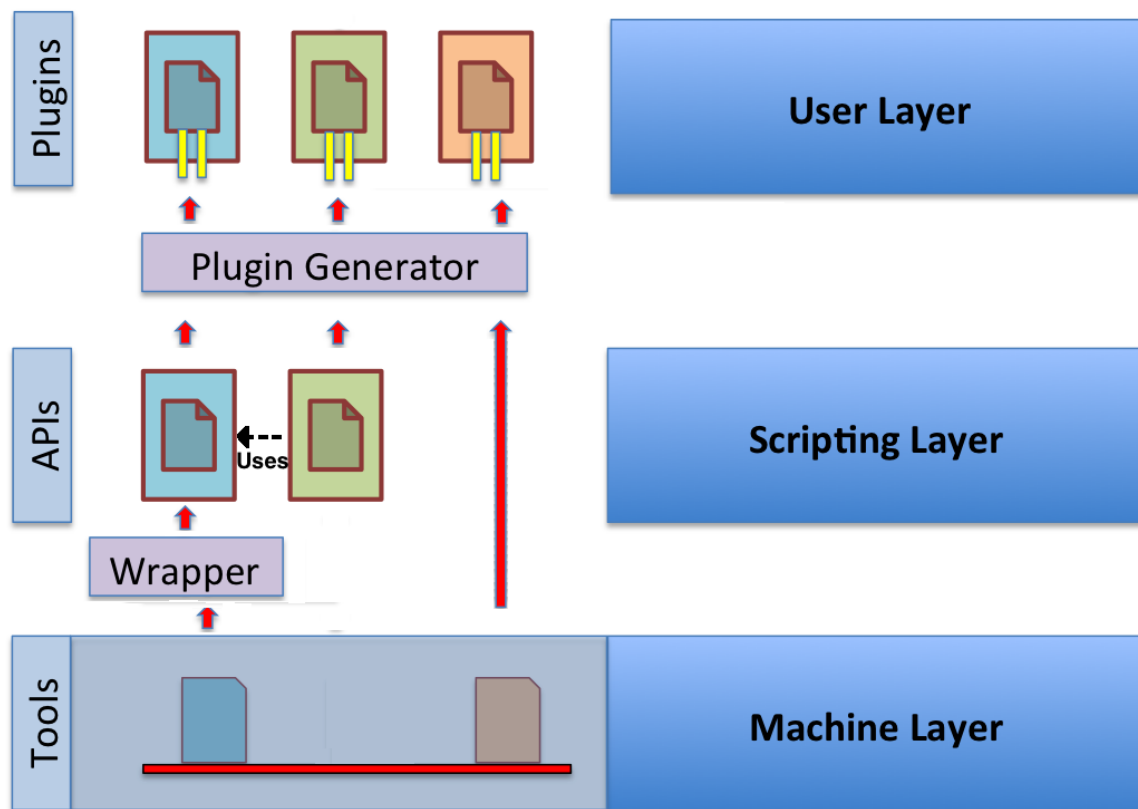


Figure 1.2: Conceptual design of PLUMA. Users interact with the software through the user layer, where they can assemble pipelines using plugin extensions in a variety of languages. Python, Perl and R plugins interface to the scripted layer of PLUMA, and compiled plugins in C++ or CUDA to the computational layer. From [5].

Chapter 2

Availability and Installation of PLUMA

2.1 How to Download PLUMA

PLUMA is hosted by the Bioinformatics Research Group (BioRG) at Florida International University at <http://biorg.cis.fiu.edu/pluma>, with source code now available through GitHub:

<https://github.com/FIUBioRG/PluMA>.

PLUMA users must agree to MIT software license regulations, part of the Open Source Initiative (OSI). We provide a copy of this license at the end of this User Guide. Finally, we make available a plugin pool of dynamically loadable plugin libraries implemented in various languages at:

<http://biorg.cs.fiu.edu/pluma/plugins.html>

which contains links to repositories of various PLUMA plugins developed by external users. Pipelines can be assembled using combinations of these plugin extensions.

2.2 Compiling PLUMA

PLUMA uses the SCons (<http://www.scons.org>) open source software construction tool to compile its back end. Please download and install SCons before compiling PLUMA. Once SCons is installed, the steps to compile PLUMA are:

1. Change to the main `pluma` directory.
2. Run the command `scons /`. If you run only `scons` and pass no flags, PLUMA will assume you would like compatibility with plugin extensions for all supported languages (currently C++, CUDA, Perl, Python, and R).

If you do not need plugins in some of these languages and/or do not wish to install their facilities, you can turn off their compilation by setting the flags to off. A list of flags that can be set during compilation can be viewed by running `scons --help`.

2.2.1 Dependencies

Compiling with R support additionally requires installation of the `RInside` package, which can be done using the standard R installation process using the command `install.packages`. This will automatically install another package `Rcpp`, which you will also need.

Perl and Python also assume the user has `PThreads` installed, though these now come standard on most *nix systems.

2.2.2 Environment Variables

Additionally, it is possible that your language tool installation locations are non-standard and you may need to command line options so that `scons` will include and link them properly. We have outlined each of these below:

Flag	Variable	Description	Default
Prefix	<code>--prefix</code>	The directory prefix for the PluMA installation	None
R	<code>--r-include-dir</code>	The include directory of the system's R installation.	None
R	<code>--r-lib-dir</code>	The lib director of the system's R installation.	None
CUDA	<code>--with-cuda</code>	Enable compilation with CUDA plugins	false
CUDA	<code>--gpu-architecture</code>	Specify the name of the class of NVIDIA 'virtual' GPU architecture for CUDA input file compilation	sm35
Python	<code>--without-python</code>	Disable compilation of Python support	false
Perl	<code>--without-perl</code>	Disable Perl plugin compilation	false
R	<code>--without-r</code>	Disable R plugin compilation	false
Rust	<code>--with-rust</code>	Enable experimental support for Rust plugins	false

Note: CUDA will work as long as the NVIDIA compiler `nvcc` is in the system `PATH`.

2.2.3 Compiling Plugins

Running `scons` will also automatically compile plugins inside the `plugins/` folder of the PLUMA source tree, which comes included (empty) within a PLUMA installation. We recommend installing any desired plugins in this folder; however other plugin directories can also be included in the compilation by specifying them in the environment variable `PLUMA_PLUGIN_PATH`. As an example, setting `PLUMA_PLUGIN_PATH` in this way:

```
export PLUMA_PLUGIN_PATH=/usr/local/bin/plugins:/home/johndoe/myplugins
```

Would automatically include the folders: `/home/johndoe/myplugins`, `/usr/local/bin/plugins`, and `plugins/` in the `scons` compilation. At the time of this release, the supported compiled languages for PLUMA is C++ only. To include CUDA plugins, pass the flag `texttt-with-cuda`.

After everything has compiled successfully, the user may run PLUMA from its root directory, which is described in the next chapter.

Chapter 3

Getting Started

We now introduce the commands needed to run PLUMA, including command line formats on *nix machines and configuration file formats. We will also demonstrate some useful features of PLUMA such as log and restart files. In the next section, we will show some real example configuration files to help further illustrate concepts. Although only *nix machines (UNIX, Linux and Mac) are currently supported, we are currently developing a Windows release for PLUMA .

3.1 Command Line

We conveniently name the application file `pluma`, which can be typed at a UNIX prompt followed by a configuration file and, optionally, a restart point:

```
./pluma (configuration file) (optional restart point)
```

The configuration file can be specified using an absolute or relative path. The restart point is a plugin name from this same configuration file, which if specified causes PLUMA to start from its stage, rather than the first plugin in the configuration file (the default).

As an example, see the three PLUMA executions below and their meanings. These executions assume that PLUMA has been installed in the location `/home/johndoe` and that the file `/home/johndoe/examples/myconfig.txt` specifies three plugins *Stage1*, *Stage2* and *Stage3*, in this order. Note that *Stage1*, *Stage2* and *Stage3* immediately become valid possible restart points:

<code>./pluma /home/johndoe/examples/myconfig.txt</code>	Run using the configuration file <code>/home/johndoe/examples/myconfig.txt</code>
<code>./pluma examples/myconfig.txt</code>	Same
<code>./pluma examples/myconfig.txt Stage1</code>	Same
<code>./pluma examples/myconfig.txt Stage2</code>	Same, but start with second stage
<code>./pluma examples/myconfig.txt Stage3</code>	Same, but only run last stage

Restart points work conveniently with *log files*, which PLUMA automatically outputs on every pipeline execution into a folder labelled with the date and time. These include each plugin that executed and particularly for a pipeline with many stages, can be a convenient way to determine a point of failure or an interesting intermediate result [25]. The user can subsequently start their pipeline using this intermediate stage as their restart point, rather than rerunning all stages before it (some may be expensive) just to reproduce the same outputs.

3.1.1 Plugin Location(s)

Dynamic loading of PLUMA plugins takes a similar approach to compiling PLUMA plugins. As before, the default location that PLUMA will assume for plugins is the `plugins/` folder in the PLUMA root directory. PLUMA will also search for plugins in any directory within the environment variable `PLUMA_PLUGIN_PATH`. The order is also the same for compilation, though becomes more significant here if two plugins have the same name. Once again, if the variable was set using:

```
export PLUMA_PLUGIN_PATH=/usr/local/bin/plugins:/home/johndoe/myplugins
```

PLUMA will first search the default `plugins/` directory, followed by `/usr/local/bin/plugins`, followed by `/home/johndoe/myplugins`. *If two plugins have the same name, PLUMA will use the second plugin.* Therefore `PLUMA_PLUGIN_PATH` can be used to define default versions of individual plugins, which can be overridden by local changes from a user.

3.1.2 Other Options

PLUMA also provides command-line options that can provide a user specific information about their software and plugins, as specified in the table below.

<code>./pluma help</code>	Help with PLUMA command line
<code>./pluma</code> (with no arguments)	Same
<code>./pluma version</code>	Version number of PLUMA that is running
<code>./pluma plugins</code>	List of all currently installed plugins (will reference <code>PLUMA_PLUGIN_PATH</code> as well)

3.2 Configuration File

We now show an example PLUMA configuration file, which runs a sample eight-stage metagenomics analysis pipeline that includes some downstream analysis. In the next section, we will illustrate how to build PLUMA plugins in various languages by building some of these plugins:

Metagenomics analysis pipeline

Prefix pipelines/Mouse

Plugin Mothur **inputfile** input.mothur **outputfile** none

Plugin CountTableProcessing **inputfile** mouse.trim.abund.pick.an.unique_list **outputfile** abund.csv

Plugin CSVNormalize **inputfile** abund.csv **outputfile** abund.norm.csv

Plugin Spearman **inputfile** abund.norm.csv **outputfile** spearman.csv

Plugin CSVPad **inputfile** spearman.csv **outputfile** network.csv

Plugin GPUATria **inputfile** network.csv **outputfile** ATria.noa

*# Plugin ATria **inputfile** network.csv **outputfile** ATria.noa*

Plugin CSV2GML **inputfile** network.csv **outputfile** network.gml

Plugin Cytoscape **inputfile** network.visualization.txt **outputfile** none

This configuration file uses all three PLUMA configuration file components: comments, prefixes and plugins. We now discuss each of these.

3.2.1 Comments

PLUMA implements comments by ignoring all characters on a line that follow a pound (#) sign. Comments are useful for writing well-documented configuration files, or also to quickly swap out a plugin and test a different one. For example in the case above we included a plugin for the *Ablatio Triadum* (ATria, [6]) algorithm, but then implemented a more efficient version on the GPU. To test for accuracy and performance improvement, we could simply comment out the CPU version of ATria and add the GPU version.

3.2.2 Prefix

The **Prefix** keyword specifies a relative or absolute path for input and output data files. This saves having to retype this path for multiple input and output files when specifying them for plugins. PLUMA will automatically use the last value of **Prefix** if more than one is specified, and the configuration file is read sequentially (so a user can specify one prefix for part of the configuration file and a different one for the rest). A user can also specify no **Prefix** at all, but then must include paths when specifying input and output files for plugins (or just have them reside in their current working directory).

3.2.3 Plugins

The core component of the PLUMA configuration file is a sequential collection of unique plugin identifiers (one for each stage of the pipeline), their input files, and their output files:

Plugin	<i>plugin1</i>	inputfile	<i>inputfilename1</i>	outputfile	<i>outputfilename1</i>
Plugin	<i>plugin2</i>	inputfile	<i>inputfilename2</i>	outputfile	<i>outputfilename2</i>
Plugin	<i>plugin3</i>	inputfile	<i>inputfilename3</i>	outputfile	<i>outputfilename3</i>
	.				
	.				
	.				

Plugins (*plugin1*, *plugin2*, *plugin3*, etc.) must be installed within the `PLUMA_PLUGIN_PATH` or the `PLUMA plugins/` folder. Output files are written by their respective plugins, and input files are read. If a particular plugin does not read and/or write a file, specify *none* as the value of *inputfile* and/or *outputfile*, respectively. A plugin's input files must exist when it executes, but not necessarily at the start of the pipeline – since very often with pipelines an input file will be an output file from a prior plugin, and probably most often the one immediately before. We had one exception in our above example, with both GPUATria (Stage 6) and CSV2GML (Stage 7) taking the same file `network.csv` as input, which was output by CSVPad (Stage 5).

3.2.4 PLUMA Plugin Pool

Existing PLUMA plugins developed by the PLUMA community can be downloaded from the PLUMA plugin pool at:

<http://biorg.cs.fiu.edu/pluma/plugins>

Figure 3.1 shows the current PLUMA plugin pool, which as of this release has 66 plugins. The plugin pool specifies each plugin along with a link to its source code repository, the source language, and a short

PLUMA: Plugin-Based Microbiome Analysis

New Release: PLUMA 1.1 Now Available

Key Features: *Kitties, Pipelines within Pipelines, Customizable Testing, Alpha GUI, 225+ Plugins*

Plugin Pool (415)

These plugins have been tested and are compatible with the latest version of PLUMA. You can download just the ones you wish to use. The easiest way to install them is to place them in the `plugins/` directory of the PLUMA source tree, however you can install them elsewhere by setting the environment variable `PLUMA_PLUGIN_PATH`.

Have a new plugin that you would like to add to the pool? Please send your source code and a description to [Trevor Cickovski](#). All plugins in the pool will be added as test cases for future releases of PLUMA, giving you automatic forward compatibility!

File Converters	Stats/Visualization	Transformations	Dissimilarity	Correlation	Centrality	Clustering	Time Series	External Tools	Miscellaneous
File Converters (53)					Stats/Visualization (79)				
Name	Short Description	Language	Name	Short Description	Language				
AbundCSV2NOA	Convert CSV file of abundances to NOA for Cytoscape	Python	Abundance	Average abundance of an entity in a sample set.	Python				
BIOM2CSV	Convert BIOM file to CSV	Python	AlphaDiversity	Alpha-diversity using Qiime (Caporaso et al, 2010)	C++				
ClusterCSV2NOA	Convert CSV File Of Clusters to NOA	Python	AlphaStats	Alpha-diversity statistics	Python				
Copy	Make a copy of a file	Python	AssemblyStats	Full genome assembly statistics (Generated by PluGen)	C++				
CountTableProcessing	Converts Mothur Counts To Abundance CSV	R	Atlas	Plot the abundance of a taxon over samples, colored by category	R				
CSVAppend	Append a new row or column to a CSV file	Python	ANOSIM	Analysis of Similarities (ANOSIM) (Clarke, 1993)	python				
CSVChop	Return a range of rows from a CSV file	Python	Bar	Bar graph of OTU abundances	R				
CSV2EDA	Converter from CSV to Cytoscape EDA file format	Python							
CSV2GML	CSV To GML Converter	Python							

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[PLUMA Plugin Pool](#)

[PLUMA Pipeline Pool](#)

PLUMA Add-Ons

[GUI](#)

[PipeViz](#)

[Upstream-Downstream](#)

[Windows](#)

Figure 3.1: PLUMA plugin pool. Available at <http://biorg.cs.fiu.edu/pluma/plugins>.

description of its functionality. Many of these plugins are also file format converters, which can be seamlessly integrated between two pipeline stages for input/output file compatibility. To assemble a PLUMA pipeline a user can install the plugins they need from the pool inside the `PLUMA plugins` directory or any directory within their `PLUMA_PLUGIN_PATH`. A user can also develop their own PLUMA plugins and run them alongside other plugins in the pool, which we describe next. If you develop a PLUMA plugin and would like to add your repository to the pool for other users in the community, please contact Trevor Cickovski at tcickovs@fiu.edu and provide a link to your repository and a description of your plugin.

3.2.5 PLUMA Pipelines

We have also provided an empty `pipelines/` folder in the PLUMA root directory, to install any PLUMA pipelines that have been made publicly available. For example, our sample pipeline that we reference in this userguide can be found at: <https://github.com/movingpictures83/Mouse>. This pipeline contains scripts that will automatically clone the appropriate PLUMA plugins into the `plugins` directory.

Chapter 4

Extending PLUMA

With plugin-based analysis the ultimate goal is infinite extensibility and flexibility, therefore we now illustrate this important property of PLUMA through some sample plugin extensions in various languages. We first show how to use our tool `PluGen` to automatically generate a plugin for an existing software package, followed by a discussion on how to build a plugin in each programming language supported by PLUMA, and finally an example of how to take components of an existing plugin and use them to build a new one. All plugins mentioned in this section are currently part of the PLUMA plugin pool.

4.1 Converting an Existing Tool to a Plugin

It is very possible for a user to have installed an existing software package that they would like to convert to a PLUMA plugin to run alongside other stages. A perfect example of this is our `Mothur` plugin from our above example, which references the metagenomics software package `Mothur` [26]. In this case we are using `Mothur` to take a sample of raw gut microbiome sequences from a mouse and produce a set of abundances for each microbial taxon in the sample. We now use `PluGen` to produce a PLUMA plugin for `Mothur`. All plugins generated by `PluGen` will be placed inside the default PLUMA `plugins/` directory. Note this also automatically includes them in the `scons` compilation as well.

`PluGen` is also automatically compiled by `scons`. To run `PluGen`, first change to the `PluGen/` subdirectory.

4.1.1 Option 1: Single Command (No Arguments)

If your software package does not accept command line arguments or its command line arguments are predefined somehow, you can generate a PLUMA plugin as follows:

```
./plugin plugin_name command_to_run
```

Where *plugin_name* will be the name of the new PLUMA plugin (be sure it is unique) and *command_to_run* is the command to run the software (with any predefined command line arguments). Although this was not applicable for our `Mothur` plugin since `Mothur` requires an input file, since its executable name is `mothur` we could have produced a plugin to run `Mothur` with no input parameters by running `./plugin Mothur mothur`. Most software tools, like `Mothur`, will require some type of input (and possibly output) parameters, so we now outline how to generate plugins with these possibilities.

4.1.2 Option 2: Input and Output File Arguments

If a plugin only requires an input and output file, then its structure can be mapped directly to that of a PLUMA plugin, where in the configuration file we provide an `inputfile` and `outputfile`. This option was applicable for our `Mothur` plugin, since `Mothur` accepts a command line parameter for an input file (it also outputs files, but those are specified in the input file and not on the command line). It would then make sense to allow the input file to be the same `inputfile` specified by a user in the PLUMA configuration file, and similar for `outputfile`. To do this, `PluGen` allows these same placeholders in its command line arguments, to represent user-specified configuration file values. For example, to generate our `Mothur` plugin we used the command:

```
./PluGen Mothur mothur inputfile
```

Now in our configuration file, we specified:

```
Plugin Mothur inputfile input.mothur outputfile none
```

`Mothur` would then be executed by our new plugin using the command `mothur input.mothur`. If we later ran the same plugin but with a different value for `inputfile` in the configuration file, that value would then be substituted as the argument to `mothur`.

4.1.3 Option 3: Fully Customized (Multiple Arguments)

Other software tools may accept multiple input parameters. For example, a common method for running Cytoscape [10] is to provide on the command line a network file to visualize, and a Cytoscape (*.cys) file for visualization properties. For our pipeline, we would have executed on the command line:

```
./cytoscape -N network.gml -s myproperties.cys
```

Since there are now multiple configuration file parameters, `PluGen` will generate a plugin that accepts a single input file with multiple keyword-value pairs. Therefore in our configuration file above, we provide:

```
Plugin Cytoscape inputfile network.visualization.txt outputfile none
```

In `network.visualization.txt`, we then provide as its contents:

```
networkfile network.gml  
sessionfile myproperties.cys
```

When running `PluGen` we use the command line as a template. The keywords in the single input file then become placeholders for their corresponding values. Our command to generate the `Cytoscape` plugin was:

```
./plugin Cytoscape cytoscape -N networkfile -s sessionfile
```

As before, if we were to run the `Cytoscape` plugin with a different input file and provide different values for `networkfile` and `sessionfile`, those values would automatically be substituted into the command line parameters above. Note we also provide flags like `-N` and `-s` in addition to the name of the `cytoscape` executable, as arguments to `PluGen`.

4.2 Building a New Plugin From Scratch

We now describe how to build a new plugin using one of PLUMA's supported languages. These languages have a variety of syntax and semantics, some are object-oriented and some are not, some are compiled and others scripted, some run on the CPU and others on the GPU. As one can imagine, the process will vary with the language. However, from the perspective of running PLUMA nothing will be different. Independent of its source language, each plugin uses the same configuration file specification outlined above. The one requirement of all PLUMA plugins is three procedures (these can be empty if desirable):

1. An `input()` procedure that accepts one parameter (an input file). Generally this will read that file and initialize the plugin.
2. A `run()` procedure that accepts zero parameters, and runs the plugin.
3. An `output()` procedure that accepts one parameter (an output file). Generally this will finalize the plugin and write that file.

We now review PLUMA's supported languages, and show through examples how to write plugins in each language. Some of these plugins were in our example configuration file from the previous session, and all are currently available in the PLUMA plugin pool.

4.2.1 C++

For C++ we show the `ATria` plugin, which accepts a signed and weighted network in CSV format, computes node centrality (importance) using the *Ablatio Triadum* algorithm (ATria, [7]), and produces a list of the most central nodes in NOde Attribute (.noa) format, which Cytoscape can use to color nodes based on centrality. New plugins should be constructed in the `PLUMA_plugins` folder and another directory within your `PLUMA_PLUGIN_PATH`. A subdirectory will then uniquely identify the plugin, in which C++ source files will use that same name followed by `Plugin`. For example, assuming we used the `PLUMA_plugins` folder our directory tree structure looked like this:

```
plugins/ATria/ATriaPlugin.h
plugins/ATria/ATriaPlugin.cpp
```

The file name should also be the same name as a new C++ class that they define. This class will then inherit from a parent class `Plugin`, defined with PLUMA in the file `Plugin.h`, which should be included as shown in Program 1, which defines the header file `ATriaPlugin.h`. Note all three required methods are present, and `input` and `output` each accept a single parameter of type `string` for the input and output files, respectively. Therefore the Standard Template Library [19] `string` class should also be included. All other included headers, member procedures and variables are at the user's discretion.

We now show a template for the corresponding source file `ATriaPlugin.cpp` in Program 2. This source file will include definitions for the three required procedures, along with any other procedures defined

Program 1 Plugin header file ATria/ATriaPlugin.h.

```
#ifndef ATRIAPLUGIN_H
#define ATRIAPLUGIN_H

#include "Plugin.h"
#include <string>
// Other necessary includes...

class ATriaPlugin : public Plugin
{
public:
    // These are required
    void input(std::string file);
    void run();
    void output(std::string file);

    // Other member procedures...

private:
    float* OrigGraph;
    std::string* bacteria;
    // Other member variables...
};

#endif
```

by the user. As is typical, the corresponding header `ATriaPlugin.h` should be included, along with two PLUMA headers: `PluginManager.h` and `PluginProxy.h`. The proxy [3] and manager [9] work together to interface this new plugin to the PLUMA computational core. Note when assembling the proxy, we also specify the name (`ATria`) which PLUMA uses to reference this plugin in the configuration file. Upon this reference, code for this plugin will be dynamically loaded at runtime, keeping the software lightweight. Note that the plugin class `ATriaPlugin` should also be passed into the proxy template as shown.

4.2.2 CUDA

In addition to C++, CUDA can be incorporated into a plugin to take advantage of GPU parallelism, assuming you have an NVIDIA graphics card installed. We will demonstrate CUDA functionality by taking the C++ `ATria` plugin from the previous section and converting it to a version `GPUATria` which runs on the GPU. We have previously used this plugin to increase the speed of `ATria` by an order of magnitude [7] for large networks, allowing us to analyze a 3000-node fruit fly network in a few hours as opposed to a few days.

Since CUDA is an extension of C and the NVIDIA compiler (NVCC) now accepts C++ code, building a CUDA plugin will work similarly to building a C++ plugin. The source file must now end in `.cu` as opposed to `.cpp` so that `scons` will compile the code with `nvcc`. CUDA *kernel* functions can then be added standalone as shown in Program 3.

CUDA uses the `__global__` keyword to preface kernel declarations. These can in turn be invoked from the C++ class functions in the source file, as shown in Program 4. For our particular implementation we used kernels for the all-pairs shortest path algorithm [12], [21] and a final pass to compute centrality.

CUDA kernels will most likely be invoked from the `run` procedure, though not necessarily. The amount

Program 2 Plugin source file ATria/ATriaPlugin.cpp.

```
#include ‘‘ATriaPlugin.h’’
#include ‘‘PluginManager.h’’
#include ‘‘PluginProxy.h’’
// Other necessary includes...

void ATriaPlugin::input(std::string file) {
    // Read file , and initialize member variables...
}

void ATriaPlugin::run() {
    // Run the algorithm...
}

void ATriaPlugin::output(std::string file) {
    // Perform any final operations , and write file...
}

// Other member procedure definitions...

// Required , connects the plugin to the PluMA back end
PluginProxy<ATriaPlugin> ATriaPluginProxy
    = PluginProxy<ATriaPlugin> (‘‘ATria’’, PluginManager::getInstance());
```

Program 3 GPUATria/GPUATriaPlugin.h

```
#ifndef GPUATRIAPLUGIN_H
#define GPUATRIAPLUGIN_H

// Same includes as C++...

class GPUATriaPlugin : public Plugin
{
    public:
        // These are still required
        void input(std::string file);
        void run();
        void output(std::string file);

        // Other member procedures...

    private:
        // Member variables...
};

__global__ void _GPU_Floyd_kernel(int k, float *G, int N);
__global__ void _GPU_Pay_kernel(float* D, float* P, int N);
// Other GPU kernels...

#endif
```

Program 4 GPUATria/GPUATriaPlugin.cu

```
// Same includes as C++...

void GPUATriaPlugin::input(std::string file) {
    // Read file , and initialize member variables...
}

void GPUATriaPlugin::run() {
    // ...
    // First kernel
    _GPU_Floyd_kernel<<<dimGrid ,BLOCK_SIZE>>>(k,dG,N);
    // ...
    // Second kernel
    _GPU_Pay_kernel<<<numblocks , BLOCK_SIZE>>>(dG, dPay, (N/2));
    // ...
}

void GPUATriaPlugin::output(std::string file) {
    // Perform any final operations , and write file...
}

// Other member procedure definitions...

__global__ void _GPU_Pay_kernel(float* D, float* P, int N) {
    // GPU code...
}

__global__ void _GPU_Floyd_kernel(int k, float *G, int N){
    // GPU code...
}

// Other kernel procedure definitions...

// Still required...
PluginProxy<GPUATriaPlugin> GPUATriaPluginProxy
    = PluginProxy<GPUATriaPlugin>("GPUATria", PluginManager::getInstance());
```

of CUDA cores and threads you allocate for each kernel (specified within the <<< and >>>) can be critical to its efficiency [17]. For more information, please view the CUDA Programmer’s Guide [20].

4.2.3 Python

The remaining PLUMA-supported languages are all scripted. The first major difference when constructing a plugin in one of these languages is that the entire plugin should be encapsulated in one file. For example, in this section we will build a template for a Python plugin `PageRank` that runs Google’s PageRank [22] centrality algorithm. The filename should follow similar conventions to the compiled plugins; using the plugin name followed by `Plugin`. We thus name our file `PageRankPlugin.py`. Note that because these plugins are scripted, they are not detected in the `scons` PLUMA compilation. However, they still should be installed in either `plugins/` or a location in the `PLUMA_PLUGIN_PATH` to be recognized at execution time. In addition, we will not need to include a proxy to interface properly to the PLUMA back end, unlike C++ and CUDA.

We show our plugin template for PageRank in Program 5. Our particular implementation imports various Python libraries [15, 18, 27] to aid in the computation. Since Python is object-oriented we once again will build the plugin as a class and as with C++ and CUDA, the classname (in our case, `PageRankPlugin`) should be the same as the filename. We then declare the three required procedures as member functions of this Python class, with `input` and `output` accepting their file parameter. Since Python is dynamically typed, as is the case for most scripting languages, no specification is required for the types of these parameters.

Program 5 `PageRank/PageRankPlugin.py`

```
import numpy
import networkx
from pythonds.graphs import PriorityQueue, Graph, Vertex
# Other imports...

numdiff = 0
ALPHA=0.5
def buildNetworkXGraph(myfile):
    # Read myfile, build and return graph

# Other global variables/procedures...

class PageRankPlugin:
    def input(self, file):
        self.bacteria, self.graph = buildNetworkXGraph(file)
    def run(self):
        self.U = networkx.pagerank(self.graph, alpha=ALPHA, max_iter=100)
    def output(self, file):
        UG = []
        for key in self.U:
            UG.append((self.U[key], key))
        UG.sort()
        UG.reverse()

        # Write file ...
```

4.2.4 R

Since the remaining PLUMA-supported language, R and Perl, are not by default object-oriented (although there are importable packages that facilitate object-oriented programming), PLUMA does not use classes for plugins in either of these languages but instead assumes that the three required procedures will be implemented standalone. Program 6 shows a plugin `Spearman` that we developed for calculating Spearman correlations, following similar conventions with regard to the filename.

Program 6 `Spearman/SpearmanPlugin.R`

```
p_value <- 0.01;
libs <- c('Hmisc');
lapply(libs, require, character.only=T);
# Other global variables and libraries...

# Required
input <- function(inputfile) {
  pc <- read.csv(inputfile, header = TRUE);
}

# Required
run <- function() {
  # Some preprocessing...
  correlations <- rcorr(pc[,], type=c("spearman"));
  pc <- as.matrix(correlations$r);
  # Post-processing and p-value thresholding...
}

# Required
output <- function(outputfile) {
  write.table(pc, file=outputfile, sep=',', append=FALSE,
    row.names=unlist(cn), col.names=unlist(cn), na='');
}

# Other helper procedures...
```

We declare any global variables and import any necessary libraries at the top of the file before the three procedure definitions. This particular plugin takes input and output in CSV format and computes correlations using the `rcorr` method from the R `Hmisc` package following some preprocessing. We perform some final p-value thresholding before output.

4.2.5 Perl

For our Perl plugin we show a simple example of a plugin `CytoViz` in Program 7 that calls Cytoscape and visualizes a network represented in the Graph Modeling Language (GML), assuming the path to Cytoscape is defined in the environment variable `CYTOSCAPE_HOME`. Following naming conventions, we implement this plugin in a file `CytoVizPlugin.pl` and include all three required procedures. Note Perl procedure parameters are implicitly stored in the array `@_`, so there is no need to specify parameters for `input` and `output`. Perl also requires global variables to be preceded with the `my` keyword, which we use for the network file. In this case our plugin will either invoke Cytoscape and visualize the network using its `-N` flag, or produce an error.

Program 7 CytoViz/CytoVizPlugin.pl

```
my $gmlfile;
# Any other global variables...

# Required
sub input {
    $gmlfile = @_[0];
    return;
}

# Required
sub run {
    $cytohome = $ENV{'CYTOSCAPE_HOME'};
    length($cytohome) != 0 or die "Please set CYTOSCAPE_HOME\n";
    @args = ($cytohome . "/cytoscape.sh", "-N", $gmlfile);
    system(@args) == 0 or die "system @args failed: $? \n";
    return;
}

# Required
sub output { return; }
```

4.3 Building a New Plugin using Existing Plugins

PLUMA plugins can also use functionality from other plugins, which can be very useful. For example, Google's PageRank can be artificially biased [28] to favor nodes with certain properties. PageRank internally involves a random walker that constantly moves between neighboring network nodes and determines centrality based on the amount of times they land on a node. We could thus tailor our centrality values to *leader* nodes of tightly connected network components by biasing PageRank to make the walker more likely to visit nodes in the same cluster (enabling cluster leaders to get hit the most). This does assume that we have already run a clustering algorithm of some kind on our network, and there are a few currently available as PLUMA plugins.

We now design our `BiasedPageRank` plugin and import the `PageRank` plugin as a Python module, as shown in Program 8. For this to work properly, the directories `plugins` and `plugins/PageRank` must be available as importable Python packages. This can be done by inserting an empty `__init__.py` file in each of these folders (the default PLUMA `plugins/` directory includes one by default). We recommend any user designing Python plugins to include this file in their specific installation directory, since this increases potential for reusing or building upon their code to produce new and creative ideas, fundamental to the purpose of PLUMA.

`BiasedPageRankPlugin` can then *inherit* from `PageRankPlugin`, as shown in the class header. We do this because the source code of our new plugin is almost exactly the same as `PageRankPlugin`, except for one small change. PageRank uses a variable `alpha` to represent the likelihood of the random walker advancing to a particular neighbor node; in this case we use two different values `alpha1` (larger) and `alpha2` (smaller), depending on whether or not the candidate neighbor is in the same cluster.

To obtain this knowledge, we also borrow functionality from a plugin `Clusterize`, which contains functions to both read a cluster file (assumed to be in CSV format mapping node names to cluster identifiers), and to determine given this data if two parameter nodes are in the same cluster. As that is also a Python plugin, we import it in a similar fashion. Our `input` method thus changes to read two CSV files (one for

the network, and one for the clusters), but since all other procedures are the same as PageRank we do not need to redefine them.

Program 8 BiasedPageRank/BiasedPageRankPlugin.py

```
import numpy
import networkx as nx

import plugins.Clusterize.ClusterizePlugin
import plugins.PageRank.PageRankPlugin

def biasedpagerank(G, clusters, alpha1=0.5, alpha2=0.35, max_iter=100,
                  tol=1.0e-8, nstart=None):

    # ...

    # Use appropriate alpha value
    for nbr in W[n]:
        if (ClusterizePlugin.inSameCluster(n, nbr, clusters)):
            x[nbr] += alpha1 * xlast[n] * W[n][nbr]['weight']
        else:
            x[nbr] += alpha2 * xlast[n] * W[n][nbr]['weight']

    # ...

#####
class BiasedPageRankPlugin(PageRankPlugin.PageRankPlugin):
    def input(self, filename):
        PageRankPlugin.input(self, filename + '.csv')
        self.clusters = ClusterizePlugin.readClusterFile(filename + '.clusters.csv')
    def run(self):
        self.U = biasedpagerank(self.graph, self.clusters, alpha1=0.5, alpha2=0.35,
                                max_iter=100)
```

Chapter 5

PLUMA Full Pipeline

We now return to our earlier example of a complete metagenomics pipeline that we used to illustrate the PLUMA configuration file:

```
# Metagenomics analysis pipeline
Prefix pipelines/Mouse
Plugin Mothur inputfile input.mothur outputfile none
Plugin CountTableProcessing inputfile mouse.trim.abund.pick.an.unique.list outputfile abund.csv
Plugin CSVNormalize inputfile abund.csv outputfile abund.norm.csv
Plugin Spearman inputfile abund.norm.csv outputfile spearman.csv
Plugin CSVPad inputfile spearman.csv outputfile network.csv
Plugin GPUATria inputfile network.csv outputfile ATria.noa
# Plugin ATria inputfile network.csv outputfile ATria.noa
Plugin CSV2GML inputfile network.csv outputfile network.gml
Plugin Cytoscape inputfile network.visualization.txt outputfile none
```

Figure 5.1 provides a conceptual description of this pipeline, which consists of plugins in all PLUMA-supported languages except Perl, although `CytoViz` could easily be substituted for `Cytoscape` above to execute our Perl example. There are also two plugins (`Mothur`) and `Cytoscape`) that were generated by `PluGen`. Some stages are simple file converters and others are more involved, but the setup is uniform in PLUMA configuration file.

We now outline the details of each pipeline stage.

5.1 Stage 1: Mothur

For the first stage, we provide a file `input.mothur` to our `Mothur` plugin. This file executes an example from the Mothur Mealybugs tutorial [16] and uses Illumina mouse genome data from MiSeq [24]. This tutorial removes noise from the reads including chimeric sequences before performing similarity-based clustering [29] and finally mapping each cluster to the closest Operational Taxonomic Unit (OTU) through lookups in the SILVA [23] database. While Mothur produces a series of output files for our pipeline we are most interested in OTUs and their abundances, which are provided in `*.shared` (Program 9) and `*.cons.taxonomy` (Program 10) files with the same prefix. Both will be necessary to run our next stage.

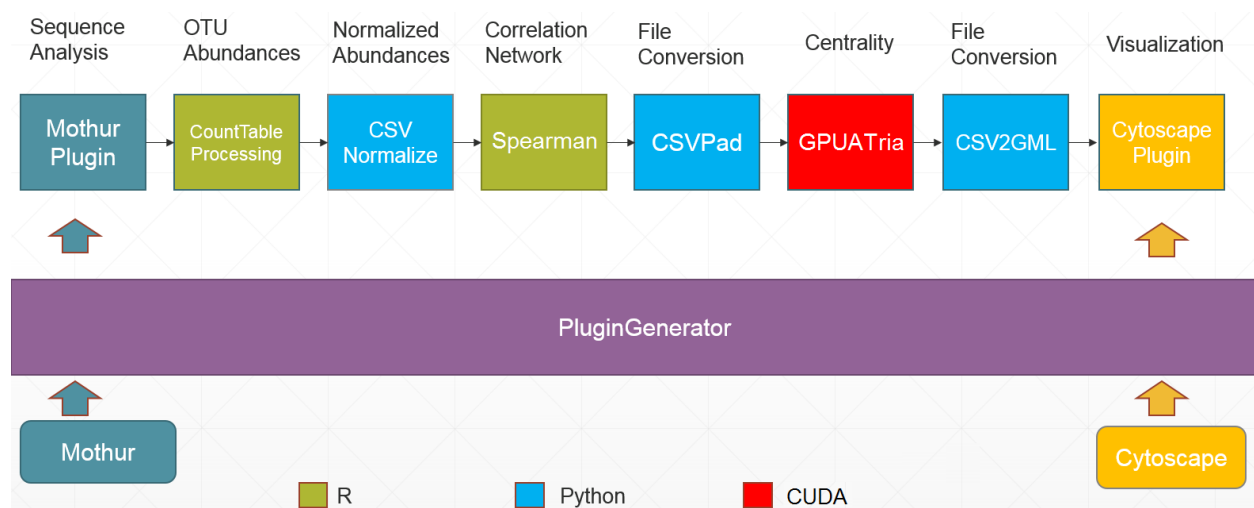


Figure 5.1: Conceptual description of a full metagenomics pipeline in PLUMA, with plugins in a variety of supported languages and two generated by `PluGen`.

Program 9 OTU abundances produced by Mothur in a `(.shared)` file).

bel	Group	numOtu	Otu001	Otu002	Otu003	Otu004	Otu005	...
0.03	F3D0	528	500	306	394	408	636	...
0.03	F3D1	528	352	311	189	64	73	...
0.03	F3D141	528	388	335	300	482	427	...
...								

Program 10 OTUs classifications produced by Mothur in a `(.cons.taxonomy)` file).

OTU	Size	Taxonomy
Otu001	12307	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
Otu002	8904	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
Otu003	7799	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
Otu004	7485	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
Otu005	7459	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
Otu006	6633	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
Otu007	6307	Bacteria(100); 'Bacteroidetes'(100); 'Bacteroidia ...
...		

5.2 Stage 2: CountTableProcessing

Now that we have the OTU abundances, we are ready to perform downstream analysis in our pipeline. However, many of our latter stages use the Comma-Separated-Value (CSV) format for data. Therefore, we include a file conversion plugin as the second stage of our pipeline. This R plugin, `CountTableProcessing`, will take the above two files from Mothur and produce an abundance matrix in CSV format. This file will include samples as rows and OTUs as columns, with entry $[i, j]$ corresponding to the abundance of OTU j in sample i . Since `CountTableProcessing` needs both Mothur output files, we provide their common prefix as input and allow the plugin to attach their respective extensions. Program 11 shows the output of this plugin.

Program 11 OTU abundance matrix produced by Stage 2 of our pipeline.

```
‘‘‘’, ‘‘Family.Porphyromonadaceae.0001’’’, ‘‘Family.Porphyromonadaceae ...
‘‘F3D0’’’, 500,306,394,408,636,356,168,164,135,19,27,110,52,104,87,78...
‘‘F3D1’’’, 352,311,189,64,73,117,127,174,83,111,53,35,115,253,288,0,7...
‘‘F3D141’’’, 388,335,300,482,427,279,205,327,120,175,112,101,8,15,28,95...
‘‘F3D142’’’, 243,258,142,152,253,192,201,81,72,43,51,82,96,6,8,45,24,5...
‘‘F3D143’’’, 189,152,172,206,309,183,116,92,63,76,69,43,37,2,7,36,26,0...
‘‘F3D144’’’, 347,243,260,316,469,282,132,36,140,255,96,84,13,7,10,100,38...
...

```

5.3 Stage 3: CSVNormalize

For our third stage, we use a Python plugin to normalize this abundance matrix, so that each sample’s OTU counts sum to 1. Normalizing guards against artifacts produced by variants such as sample quality. Measuring each OTU as a percentage of the total OTUs in the sample creates a uniform metric across all samples. Program 12 shows the normalized abundances produced by `CSVNormalize`.

Program 12 Normalized abundance matrix produced by Stage 3 of our pipeline.

```
‘‘‘’, ‘‘Family.Porphyromonadaceae.0001’’’, ‘‘Family.Porphyromonadaceae ...
‘‘F3D0’’’, 0.0803987779386,0.0492040520984,0.0633542370156,0.06560540...
‘‘F3D1’’’, 0.0754555198285,0.0666666666667,0.0405144694534,0.01371918...
‘‘F3D141’’’, 0.0833333333333,0.0719501718213,0.0644329896907,0.103522...
‘‘F3D142’’’, 0.10028889806,0.10647957078,0.0586050350805,0.0627321502...
‘‘F3D143’’’, 0.0788485607009,0.0634125990822,0.0717563621193,0.085940...
‘‘F3D144’’’, 0.100696459663,0.070516540917,0.0754497968659,0.09170052...
...

```

5.4 Stage 4: Spearman

Normalized abundances enable us to produce a *correlation matrix* where OTUs occupy both rows and columns, and entry (i, j) is the correlation value between the abundances of OTU i and OTU j over all samples. This can also be represented as a correlation *network* [], where OTUs are network nodes and edges are weighted with their correlation values (a correlation of zero would mean no edge in the network). Here

Program 14 Padded OTU correlation matrix produced by Stage 5 of the pipeline.

```
‘‘‘’, ‘‘Family . Porphyromonadaceae .0001’’’, ‘‘Family . Porphyromonadaceae .0002’’’...
‘‘Family . Porphyromonadaceae .0001’’’, 1, 0.743859648704529, 0, 0, 0, 0, 0, 0, 0...
‘‘Family . Porphyromonadaceae .0002’’’, 0.743859648704529, 1, 0, 0, 0, 0, 0, 0, 0...
‘‘Family . Porphyromonadaceae .0003’’’, 0, 0, 1, 0, 0, 0, 0, 0, 0.752631604671478...
‘‘Family . Porphyromonadaceae .0004’’’, 0, 0, 0, 1, 0.624561429023743, 0.76666...
‘‘Family . Porphyromonadaceae .0005’’’, 0, 0, 0, 0.624561429023743, 1, 0.81052...
‘‘Family . Porphyromonadaceae .0006’’’, 0, 0, 0, 0.766666650772095, 0.8105263...
...
```

We use the faster CUDA version in this pipeline as stage 6, and show its output in Program 15. The output takes the form of a NNode Attribute (NOA) file, which we will provide to Cytoscape for visualizing nodes differently based on centrality value or rank.

Program 15 NOA file of nodes, centrality values, and ranks produced by Stage 6.

Name	Centrality	Rank
Phylum . Bacteroidetes .0001	116.068	536
Kingdom . Bacteria .0012	86.5733	535
Class . Clostridia .0001	72.5422	534
Escherichia .Shigella .0001	67.3395	533
Family . Enterobacteriaceae .0002	57.8472	532
Family . Porphyromonadaceae .0057	52.0528	531
...		

5.7 Stage 7: CSV2GML

Cytoscape will also need our network, but cannot visualize networks in CSV format. We thus include one final stage before visualization `CSV2GML` that accepts a CSV file and converts it to the Graph Modeling Language (GML) which is recognizable by Cytoscape. Program 16 shows the equivalent GML file to our CSV network in Program 14, which we will provide to Cytoscape along with the NOA output from Program 15.

5.8 Stage 8: Cytoscape

For our final stage we will use our plugin for Cytoscape generated by `PluGen`. Recall that this plugin accepts as input a plaintext file (we called this `network.visualization.txt` that includes both a properties file (.cys) and a network file (.gml). We provide our output GML file from Program 16 as the latter, and include our own properties file that colors nodes based on an attribute "Centrality". Figure 5.3 shows Cytoscape initially opening when our plugin executes, although no nodes are colored yet because we must input our NOA file from Program 15 manually through the Cytoscape menu bars. Once we input this file our node colors change based on centrality value. Figure 5.4 shows this network, with red (violet) indicating high (low) centrality. Zero centrality is indicated by white nodes.

Program 16 Our correlation network from Program 14 in GML format, produced by Stage 7 of our pipeline..

```
graph [  
  node [  
    id 0  
    label ``Family.Porphyromonadaceae.0001``  
  ]  
  node [  
    id 1  
    label ``Family.Porphyromonadaceae.0002``  
  ]  
  ...  
  edge [  
    source 0  
    target 1  
    weight 0.743859648705  
  ]  
  edge [  
    source 0  
    target 27  
    weight -0.601754367352  
  ]  
  ...  
]
```

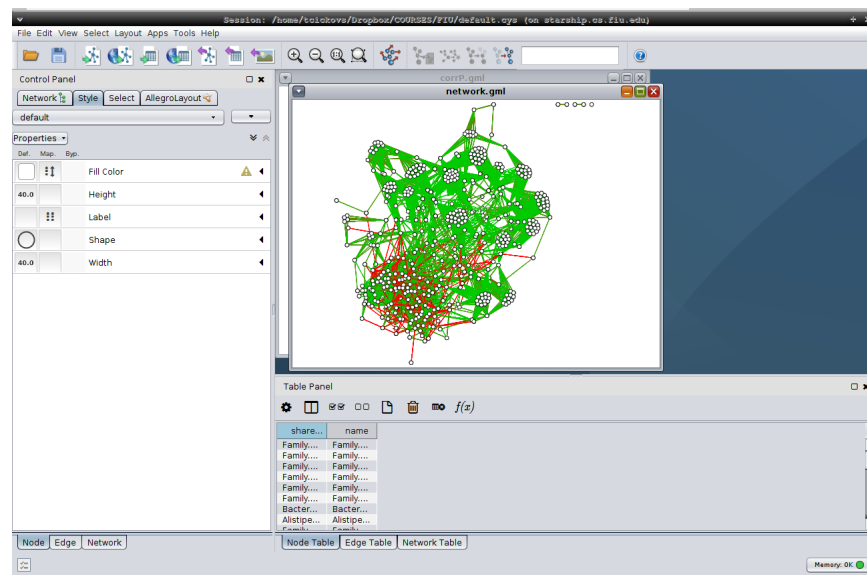


Figure 5.3: Our network visualized with Cytoscape upon executing Stage 8 of our pipeline.

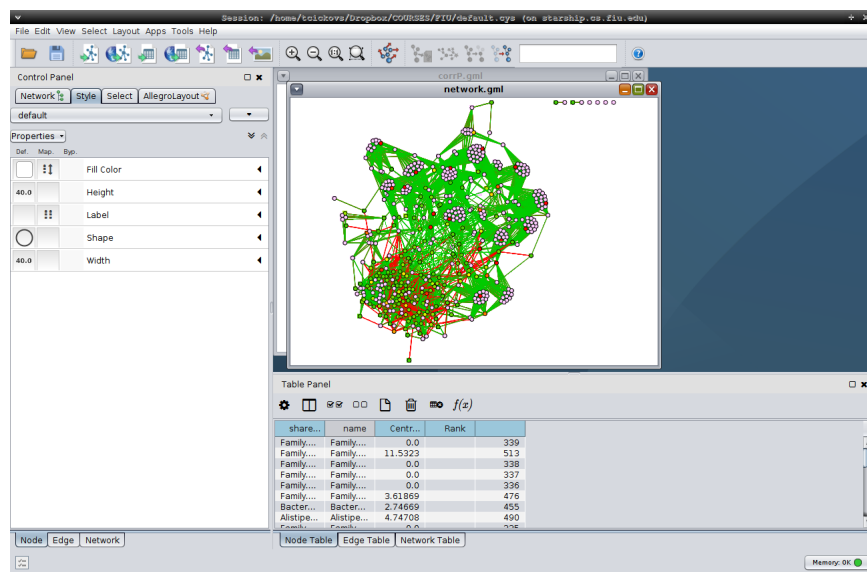


Figure 5.4: Our Cytoscape visualization after importing our NOA file from Stage 6.

Chapter 6

The Future of PLUMA

As mentioned, our vision with PLUMA is to allow users to develop new and creative analysis algorithms as plugins in their language of choice, and run them seamlessly alongside other plugins within a lightweight and flexible software framework. We will develop any future goals of PLUMA with this central purpose in mind.

The Simplified Wrapper and Generator Interface (SWIG, [4]) provides a good opportunity for PLUMA to interface plugins in scripting languages with plugins in compiled languages, or even PLUMA's computational core. SWIG can be used to wrap C++ code into functions that can be called from a scripted API. Incorporating this tool into PLUMA will allow users to develop algorithms not just alongside existing plugins, but using portions of existing plugins. We showed an example of this through our `BiasedPageRank` plugin which extended an existing `PageRank` plugin in the same language. SWIG would enable this to be done across languages, increasing reusability and further minimizing reinventing of the wheel. We will explore this soon.

In addition there are several other features that we feel will improve usability of the software itself. First, we are currently developing a Windows version of PLUMA to expand our user community. We have also prototyped a Graphical User Interface (GUI) as an alternative to the configuration file in the PLUMA User Layer, where plugins can be inserted and removed using a drag-and-drop interface. This GUI requires further testing and debugging but we hope to include it soon. Finally, we would like to PLUMA's set of supported languages to include Java, and possibly other languages that make sense given our user community. We look forward to the continued growth of the PLUMA plugin pool, and future development of its computational core.

Appendix A

PLUMA Software License

A.1 Conditions and Regulations

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A.2 Contact Information

The best contact path for licensing issues is by e-mail to lead developer Trevor Cickovski at tcickovs@fiu.edu or send correspondence to:

PLUMA Team
c/o Prof. Giri Narasimhan
Bioinformatics Research Group (BioRG)
School of Engineering and Computer Science
Florida International University
11200 SW 8th Street Miami, FL 33199 USA

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