Hermes Documentation (add references!)

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1 Introduction

This documentation aims to outline the basic dependencies, installation, and usage of the Hermes RNA software suite, as well as provide extended commentary on the flags and options available to code. Particular attention is payed to the invocation style of mashup programs (using multi_param) to dispatch flags to the appropriate submodules. To get started as quickly as possible, see the quickstart one-liner in 2.1. Should you still have questions, you can reach the main author of the source code at evansenter@gmail.com.

2 Installation

2.1 Quick Start

From the root directory of Hermes, execute the following command:

```
cd build && cmake .. && make
```

If you encounter errors in configuring or compiling the software, we recommend checking out section 2.4 on common troubleshooting solutions.

2.2 Dependencies

```
cmake (\geq 2.6-patch 4, tested through 3.0.0) http://www.cmake.org/
```

CMake is used as the build system for Hermes.

GNU99 compiler:

Most C compilers should support the -std=gnu99 flag, which is required for GNU library extensions, particularly unistd.h

C++11 compiler:

C++11 support (included in $g++ \ge 4.7$) is necessary for proper struct initialization in FFTbor2D.

OpenMP support http://openmp.org/wp/

OpenMP support comes by default in most modern compilers, and is required for loop-optimization in FFTbor2D.

LAPACK ($\geq 3.4.2$) http://www.netlib.org/lapack/

Various LAPACK routines are used in RNAmfptto compute the inverse, or pseudoinverse, of a transition probability matrix.

GSL (≥ 1.15 , tested through 1.16) http://www.gnu.org/software/gsl/

GSL is required to compute the eigendecomposition of a (possibly) non-symmetric transition rate matrix for RNAeq.

FFTW3 ($\geq 3.3.4$) http://www.fftw.org/

FFTW3 functions are used to compute the inverse discrete Fourier transform in FFTbor2D.

libRNA.a ($\geq 2.0.7$, tested through 2.1.7) http://www.tbi.univie.ac.at/RNA/

Various ViennaRNA functions and data structures are leveraged for homogenous energy model support, as well as fold_par, pf_fold_par, and subopt_par. We additionally make use of ViennaRNA functions to compute the necessary polynomial size for FFTbor2D, determined in the following fashion. Parameters K (resp. L) are defined to be the sum of the number of base pairs in reference structure A (resp. reference structure B) plus the number of base pairs in the maximum matching (Nussinov) structure which contains no base pair of A (resp. B).

2.3 Compilation

cd build && cmake ...

First, ensure that your system has the dependencies outlined above (the presence of CMake can be verified with cmake --version). While not required, is widely considered best practice to perform an out-of-source build, where the compilation of the code happens in a separate directory from the location of the source itself. To this end, hermesprovides an empty build directory that can be used for compiling the code. From hermes/build execute cmake with the path to hermes/CMakeLists.txt(..) provided as argument.

make

Compiles the code, and generates binary executables for FFTbor2D, RNAmfpt, RNAeq, FFTmfpt, FFTeq, and RateEq. Additionally generates both static and shared libraries for FFTbor2D, RNAmfpt, and RNAeq. The output directory for binaries is hermes/bin and output directory for libraries is hermes/lib.

make install (optional)

Installs the executables built with make to \$DESTDIR/bin and copies libraries / archives to \$DESTDIR/lib (on *nix systems, \$DESTDIR defaults to /usr/local).

2.4 Troubleshooting

While we have done the utmost to try and ensure that CMake is able to infer locations of third-party libraries and add compiler-appropriate flags in an automated fashion, due to the diversity of build environments possible, it is possible that you will need to specify additional command-line flags to cmake when generating the Makefiles in order to successfully build Hermes. The following are five useful flags for CMake, and a brief explanation of when they may need to be employed:

• The default compiler I'd like to use for C code is installed in a non-standard location, or not the globally default C compiler.

CMAKE_C_COMPILER

i.e. cmake -DCMAKE_C_COMPILER=/path/to/c/compiler ...

This variable sets the path to the compiler to use for configuration and subsequent compiliation via make. This is the compiler that will be used by CMake to test for the presence of various flags, i.e. -03 and -Wall.

• The default compiler I'd like to use for C++ code is installed in a non-standard location, or not the globally default C++ compiler.

CMAKE_CXX_COMPILER

i.e. cmake -DCMAKE_CXX_COMPILER=/path/to/cxx/compiler ...

Same as above.

• I'm getting a compile-time error indicating undefined symbols for _get_iindx, _maximumMatchingConstraint or something similar.

CMAKE_LIBRARY_PATH

i.e. cmake -DCMAKE_LIBRARY_PATH=/dir/for/libRNA-2.0.7+/ ...

These are libRNA.a symbols specific to the 2.0+ release of ViennaRNA. In all cases identified thus far, this error means that the version of libRNA.a found by CMake is not out of date, and can be resolved by explicitly providing the library path to a 2.0+ ViennaRNA static library using the CMAKE_LIBRARY_PATH flag.

• Libraries required by Hermes are not installed in a location visible by the linker (in LD_LIBRARY_PATH), and CMake is unable to validate their existence.

CMAKE_LIBRARY_PATH

i.e. cmake -DCMAKE_LIBRARY_PATH="/more/libraries;/even/more/libraries" ...

In the case when libraries required by Hermes are not visible to CMake, or the global library is an out-of-date version, it is possible to provide hints to the build tool for additional directories to search. Directories specified by the CMAKE_LIBRARY_PATH flag will be prepended onto the linker search path, and thus override global matches (handling the case where default libraries aren't sufficiently up to date). When desiring to provide multiple locations to search for libraries, CMake uses the semicolon (;) character as a separator and the entire string should be quoted to escape the shell environment.

• Headers required by Hermes are not installed in a location visible by the compiler (in CPATH or a derivative), and as a result I'm seeing undefined reference to errors.

CMAKE_INCLUDE_PATH

i.e. cmake -DCMAKE_INCLUDE_PATH="/more/includes;/even/more/includes" ...

It is generally likely that the CMAKE_LIBRARY_PATH and CMAKE_INCLUDE_PATH will both be necessary, when either one is required. This flag operates in a fashion identical to CMAKE_LIBRARY_PATH described above, and uses the same syntax. Alternatively a user can update their CPATH environment variable, but this may have unpredictable results when headers are found, but out of date.

• I don't have permissions to make install to the default location (generally /usr/local for *nix) on my system.

CMAKE_INSTALL_PREFIX

i.e. cmake -DCMAKE_INSTALL_PREFIX=/make/install/path/prefix ..

This variable sets the destination directory of the make install command. Binaries will be placed in the bin subdirectory and libraries will be placed in the lib subdirectory. This is analogous to ./configure --prefix=/make/install/path/prefix in Autotools and can also be achieved by setting the DESTDIR environment variable.

3 Software Organization

3.1 General Principles

FFTeq, located in hermes/mashup/population_from_fftbor2d uses functions from libfftbor_static.a (derived from FFTbor2D) to compute the energy landscape and functions from librnaeq_static.a (derived from RNAeq) to estimate population occupancy for s, A, B without requiring an investigator to copy pieces of individual packages to achieve their goals, instead using the static libraries automatically produced for all hermes/src software, shared headers made available in hermes/h and libmulti_param.a (from hermes/src/multi_param) to dispatch command-line arguments to the appropriate underlying function. The result? The entirety of FFTeqis 67 lines of C++ code, and uses native binary data structures the entire way through; there is no command-line funneling of FFTbor2Dinto RNAeq.

3.2 multi_param Overview

All of FFTbor2D, RNAmfpt, and RNAeqpresent a wide selection of command-line arguments to meet the diverse demands of end users. When we moved on to developing mashup software between these three programs, there were a number of requirements that we came up with to make both implementing and using these programs as easy as possible. We decided that a) the developer should not have to reimplement command-line parsing for mashups b) all existing flags for underlying libraries (i.e. FFTbor2D, RNAmfpt, RNAeq) would be supported, and c) flags would be namespaced to have deterministic targets.

To achieve these goals, the library multi_paramwas developed. This library simply takes a collection of command line arguments and re-dispatches them to the appropriate underlying library. Given a set of command-line flags such as --all-v --fftbor2d-i GGGAAACCC --fftbor2d-j '.....' --fftbor2d-k '(((...)))' --mfpt-x --mfpt-h, the code ensures that FFTbor2Dis passed -v -i GGGAAACCC -j '.....' -k '(((...)))' as options in argv and RNAmfptis passed -v -x -h as options in argv.

3.3 multi_param Details

An example from hermes/mashup/mfpt_from_fftbor2d/mfpt_from_fftbor2d.cpp:

```
PARAM_CONTAINER* params;
FFTBOR2D_PARAMS fftbor2d_params;

/* ...omitted for clarity... */
char* subparams[] = { "fftbor2d", "mfpt" };
params = split_args(argc, argv, subparams, 2);

fftbor2d_params = init_fftbor2d_params();
parse_fftbor2d_args(fftbor2d_params, params[0].argc, params[0].argv);
```

The way this code works is by first declaring which packages can be used by the mashups, out of fftbor2d, mfpt, or population. In the snippet above the selection is saved in the variable subparams. All of the FFTbor2D, RNAmfpt, and RNAeqlibraries have init_*_params functions available, which return an object with the default parameters for that package. They also all have parse_*_args functions, which take three arguments, 1) a pointer to the parameters object 2) argc, and 3) argv.

split_args takes the prefixed command-line arguments (see the example in 3.2) and bins them by their prefix, with the special --all prefix being supplied to all declared subparams. The prefixes are then removed and the grouped arguments are returned from the function in a PARAM_CONTAINER array, with the same order as the subparams were passed to the function. It is then trivial to call the parse_*_args functions with the corresponding subarrays from PARAM_CONTAINER to get a final parameters object.

Leveraging the example from 3.2 a final time, the variable params would look as follows after invoking split_args:

```
[
    { argv: ["-v", "-i", "GGGAAACCC", "-j", "......", "-k", "(((...)))"], argc: 7 },
    { argv: ["-v", "-x", "-h"], argc: 3 }
]
```

4 Core Programs

4.1 FFTbor2D

4.1.1 Applications

TEST

4.1.2 Example Usage

TEST2

4.1.3 Options

TEST3

- 4.2 RNAmfpt
- 4.3 RNAeq
- 5 Mashups
- 5.1 FFTmfpt
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- 5.3 RateEq
- 6 References