khmer Documentation

Release 1.4.1+0.g1e762a3.dirty

2010-2014 Michael R. Crusoe, Greg Edvenson, Jordan Fish, Adina

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Contact khmer-project@idyll.org

GitHub https://github.com/ged-lab/khmer

Chat https://gitter.im/ged-lab/khmer

License BSD

khmer is a library and suite of command line tools for working with DNA sequence. It is primarily aimed at short-read sequencing data such as that produced by the Illumina platform. khmer takes a k-mer-centric approach to sequence analysis, hence the name.

Installing and running khmer

How to get help

There are two mailing lists dedicated to khmer, an announcements-only list and a discussion list. To search their archives and sign-up for them, please visit the following URLs:

- · Discussion: http://lists.idyll.org/listinfo/khmer
- Announcements: http://lists.idyll.org/listinfo/khmer-announce

The archives for the khmer list are available at: http://lists.idyll.org/pipermail/khmer/

khmer development has largely been supported by AFRI Competitive Grant no. 2010-65205-20361 from the USDA NIFA, and is now funded by the National Human Genome Research Institute of the National Institutes of Health under Award Number R01HG007513 through May 2016, both to C. Titus Brown.

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Introduction to khmer

1.1 Introduction

khmer is a library and toolkit for doing k-mer-based dataset analysis and transformations. Our focus in developing it has been on scaling assembly of metagenomes and mRNA.

khmer can be used for a number of transformations, include inexact transformations (abundance filtering and error trimming) and exact transformations (graph-size filtering, to throw away disconnected reads; and partitioning, to split reads into disjoint sets). Of these, only partitioning is not constant memory. In all cases, the memory required for assembly with Velvet or another de Bruijn graph assembler will be more than the memory required to use our software. Our software will not increase the memory required for Velvet, either, although we may not be able to *decrease* the memory required for assembly for every data set.

Most of khmer relies on an underlying probabilistic data structure known as a Bloom filter (also see Count-Min Sketch and These Are Not The k-mers You're Looking For), which is essentially a set of hash tables, each of different size, with no collision detection. These hash tables are used to store the presence of specific k-mers and/or their count. The lack of collision detection means that the Bloom filter may report a k-mer as being "present" when it is not, in fact, in the data set; however, it will never incorrectly report a k-mer as being absent when it *is* present. This one-sided error makes the Bloom filter very useful for certain kinds of operations.

khmer is also independent of K, and currently works for K <= 32. We will be integrating code for up to K=64 soon.

khmer is implemented in C++ with a Python wrapper, which is what all of the scripts use.

Some important documentation for khmer is provided on the Web sites for khmer-protocols and khmer-recipes. khmer-protocols provides detailed protocols for using khmer to analyze either a transcriptome or a metagenome; khmer-recipes provides individual recipes for using khmer in a variety of sequence-oriented tasks such as extracting reads by coverage, estimating a genome or metagenome size from unassembled reads, and error-trimming reads via streaming k-mer abundance.

1.2 Using khmer

khmer comes "out of the box" with a number of scripts that make it immediately useful for a few different operations, including:

- normalizing read coverage ("digital normalization")
- dividing reads into disjoint sets that do not connect ("partitioning")
- eliminating reads that will not be used by a de Bruijn graph assembler;
- removing reads with low- or high-abundance k-mers;

- trimming reads of certain kinds of sequencing errors;
- counting k-mers and estimating data set coverage based on k-mer counts;
- running Velvet and calculating assembly statistics;
- optimizing assemblies on various parameters;
- converting FASTA to FASTQ;

and a few other random functions.

1.3 Practical considerations

The most important thing to think about when using khmer is whether or not the transformation or filter you're applying is appropriate for the data you're trying to assemble. Two of the most powerful operations available in khmer, graph-size filtering and graph partitioning, only make sense for assembly datasets with many theoretically unconnected components. This is typical of metagenomic data sets.

The second most important consideration is memory usage. The effectiveness of all of the Bloom filter-based functions (which is everything interesting in khmer!) depends critically on having enough memory to do a good job. See Choosing table sizes for khmer for more information.

1.4 Copyright and license

Portions of khmer are Copyright California Institute of Technology, where the exact counting code was first developed; the remainder is Copyright Michigan State University. The code is freely available for use and re-use under the BSD License.

Contributors and Acknowledgements

khmer is a product of the GED lab at Michigan State University,

http://ged.msu.edu/

_

C. Titus Brown <ctb@msu.edu> wrote the initial ktable and hashtable implementations, as well as hashbits and counting_hash.

Jason Pell implemented many of the C++ k-mer filtering functions.

Qingpeng contributed code to do unique k-mer counting.

Adina Howe, Rosangela Canino-Koning, and Arend Hintze contributed significantly to discussions of approaches and algorithms; Adina wrote a number of scripts.

Jared T. Simpson (University of Cambridge, Sanger Institute) contributed paired-end support for digital normalization.

Eric McDonald thoroughly revised many aspects of the code base, made much of the codebase thread safe, and otherwise improved performance dramatically.

Michael R. Crusoe is the new maintainer of khmer.

MRC 2014-05-07



Citations

3.1 Software Citation

If you use the khmer software, you must cite:

Crusoe et al., The khmer software package: enabling efficient sequence analysis. 2014. http://dx.doi.org/10.6084/m9.figshare.979190

```
@article{khmer2014,
    author = "Crusoe, Michael and Edvenson, Greg and Fish, Jordan and Howe,
Adina and McDonald, Eric and Nahum, Joshua and Nanlohy, Kaben and
Ortiz-Zuazaga, Humberto and Pell, Jason and Simpson, Jared and Scott, Camille
and Srinivasan, Ramakrishnan Rajaram and Zhang, Qingpeng and Brown, C. Titus",
    title = "The khmer software package: enabling efficient sequence
analysis",
    year = "2014",
    month = "04",
    publisher = "Figshare",
    url = "http://dx.doi.org/10.6084/m9.figshare.979190"
}
```

If you use any of our published scientific methods, you should *also* cite the relevant paper(s), as directed below. Additionally some scripts use the SeqAn library for read parsing: the full citation for that library is also included below.

To see a quick summary of papers for a given script just run it without using any command line arguments.

3.2 Graph partitioning and/or compressible graph representation

The load-graph.py, partition-graph.py, find-knots.py, load-graph.py, and partition-graph.py scripts are part of the compressible graph representation and partitioning algorithms described in:

Pell J, Hintze A, Canino-Koning R, Howe A, Tiedje JM, Brown CT. Scaling metagenome sequence assembly with probabilistic de Bruijn graphs Proc Natl Acad Sci U S A. 2012 Aug 14;109(33):13272-7. http://dx.doi.org/10.1073/pnas.1121464109. PMID: 22847406

```
@article{Pell2012,
    author = "Pell, Jason and Hintze, Arend and Canino-Koning, Rosangela and
Howe, Adina and Tiedje, James M. and Brown, C. Titus",
    title = "Scaling metagenome sequence assembly with probabilistic de Bruijn
graphs",
```

```
volume = "109",
   number = "33",
   pages = "13272-13277",
   year = "2012",
   doi = "10.1073/pnas.1121464109",
    abstract = "Deep sequencing has enabled the investigation of a wide range of
environmental microbial ecosystems, but the high memory requirements for de
novo assembly of short-read shotgun sequencing data from these complex
populations are an increasingly large practical barrier. Here we introduce a
memory-efficient graph representation with which we can analyze the k-mer
connectivity of metagenomic samples. The graph representation is based on a
probabilistic data structure, a Bloom filter, that allows us to efficiently
store assembly graphs in as little as 4 bits per k-mer, albeit inexactly. We
show that this data structure accurately represents DNA assembly graphs in low
memory. We apply this data structure to the problem of partitioning assembly
graphs into components as a prelude to assembly, and show that this reduces the
overall memory requirements for de novo assembly of metagenomes. On one soil
metagenome assembly, this approach achieves a nearly 40-fold decrease in the
maximum memory requirements for assembly. This probabilistic graph
representation is a significant theoretical advance in storing assembly graphs
and also yields immediate leverage on metagenomic assembly.",
   URL = "http://www.pnas.org/content/109/33/13272.abstract",
   eprint = "http://www.pnas.org/content/109/33/13272.full.pdf+html",
    journal = "Proceedings of the National Academy of Sciences"
```

3.3 Digital normalization

The normalize-by-median.py and count-median.py scripts are part of the digital normalization algorithm, described in:

A Reference-Free Algorithm for Computational Normalization of Shotgun Sequencing Data Brown CT, Howe AC, Zhang Q, Pyrkosz AB, Brom TH arXiv:1203.4802 [q-bio.GN] http://arxiv.org/abs/1203.4802

```
@unpublished{diginorm,
    author = "C. Titus Brown and Adina Howe and Qingpeng Zhang and Alexis B.
Pyrkosz and Timothy H. Brom",
    title = "A Reference-Free Algorithm for Computational Normalization of
Shotgun Sequencing Data",
    year = "2012",
    eprint = "arXiv:1203.4802",
    url = "http://arxiv.org/abs/1203.4802",
}
```

3.4 K-mer counting

The abundance-dist.py, filter-abund.py, and load-into-counting.py scripts implement the probabilistic k-mer counting described in:

These Are Not the K-mers You Are Looking For: Efficient Online K-mer Counting Using a Probabilistic Data Structure Zhang Q, Pell J, Canino-Koning R, Howe AC, Brown CT. http://dx.doi.org/10.1371/journal.pone.0101271

```
@article{khmer-counting,
   author = "Zhang, Qingpeng AND Pell, Jason AND Canino-Koning, Rosangela
AND Howe, Adina Chuang AND Brown, C. Titus",
    journal = "PLoS ONE",
    publisher = "Public Library of Science",
    title = "These Are Not the K-mers You Are Looking For: Efficient Online
K-mer Counting Using a Probabilistic Data Structure",
   year = "2014",
   month = "07",
   volume = "9",
   url = "http://dx.doi.org/10.1371%2Fjournal.pone.0101271",
   pages = "e101271",
   abstract = "K-mer abundance analysis is widely used for many purposes in
nucleotide sequence analysis, including data preprocessing for de novo
assembly, repeat detection, and sequencing coverage estimation. We present the
khmer software package for fast and memory efficient <italic>online</italic>
counting of k-mers in sequencing data sets. Unlike previous methods based on
data structures such as hash tables, suffix arrays, and trie structures, khmer
relies entirely on a simple probabilistic data structure, a Count-Min Sketch.
The Count-Min Sketch permits online updating and retrieval of k-mer counts in
memory which is necessary to support online k-mer analysis algorithms. On
sparse data sets this data structure is considerably more memory efficient than
any exact data structure. In exchange, the use of a Count-Min Sketch introduces
a systematic overcount for k-mers; moreover, only the counts, and not the
k-mers, are stored. Here we analyze the speed, the memory usage, and the
miscount rate of khmer for generating k-mer frequency distributions and
retrieving k-mer counts for individual k-mers. We also compare the performance
of khmer to several other k-mer counting packages, including Tallymer,
Jellyfish, BFCounter, DSK, KMC, Turtle and KAnalyze. Finally, we examine the
effectiveness of profiling sequencing error, k-mer abundance trimming, and
digital normalization of reads in the context of high khmer false positive
rates. khmer is implemented in C++ wrapped in a Python interface, offers a
tested and robust API, and is freely available under the BSD license at
github.com/ged-lab/khmer.",
   number = "7",
   doi = "10.1371/journal.pone.0101271"
```

3.5 FASTA and FASTQ reading

Several scripts use the SeqAn library for FASTQ and FASTA reading as described in:

SeqAn An efficient, generic C++ library for sequence analysis Döring A, Weese D, Rausch T, Reinert K. http://dx.doi.org/10.1186/1471-2105-9-11

```
@Article{SeqAn,
  AUTHOR = {Doring, Andreas and Weese, David and Rausch, Tobias and Reinert,
      Knut},
  TITLE = {SeqAn An efficient, generic C++ library for sequence analysis},
  JOURNAL = {BMC Bioinformatics},
  VOLUME = {9},
  YEAR = {2008},
  NUMBER = {1},
  PAGES = {11},
  URL = {http://www.biomedcentral.com/1471-2105/9/11},
  DOI = {10.1186/1471-2105-9-11},
```

PubMedID = $\{18184432\}$, $ISSN = \{1471 - 2105\},\,$ ABSTRACT = {BACKGROUND: The use of novel algorithmic techniques is pivotal to many important problems in life science. For example the sequencing of the human genome [1] would not have been possible without advanced assembly algorithms. However, owing to the high speed of technological progress and the urgent need for bioinformatics tools, there is a widening gap between state-of-the-art algorithmic techniques and the actual algorithmic components of tools that are in widespread use. RESULTS: To remedy this trend we propose the use of SeqAn, a library of efficient data types and algorithms for sequence analysis in computational biology. SeqAn comprises implementations of existing, practical state-of-the-art algorithmic components to provide a sound basis for algorithm testing and development. In this paper we describe the design and content of SeqAn and demonstrate its use by giving two examples. In the first example we show an application of SeqAn as an experimental platform by comparing different exact string matching algorithms. The second example is a simple version of the wellknown MUMmer tool rewritten in SeqAn. Results indicate that our implementation is very efficient and versatile to use. CONCLUSION: We anticipate that SeqAn greatly simplifies the rapid development of new bioinformatics tools by providing a collection of readily usable, welldesigned algorithmic components which are fundamental for the field of sequence analysis. This leverages not only the implementation of new algorithms, but also enables a sound analysis and comparison of existing algorithms. },

10 Chapter 3. Citations

Release notes

Contents:

4.1 khmer v1.0 release notes

582 changed files with 40,527 additions and 31,772 deletions.

The team has been hard at work since v0.8 to refine the codebase into a stable product.

https://khmer.readthedocs.org/en/latest/

With the 1.0 release we are making a commitment to using Semantic Versioning[0]: the version number will reflect the impact of the changes between releases. New major versions will likely require you to change how you use the project. Minor versions indicate new functionality that doesn't impact the existing. Patch versions indicate backwards-compatible fixes. Right now we are limiting this promise to the command-line interface. A future release will introduce a stable and mature Python API to the khmer project and at that time we will extend the version system to include that API.

4.1.1 New items of note:

CITATION: Each script now outputs information on how to cite it. There is a new paper to describes the project overall: MR Crusoe et al., 2014. doi: 10.6084/m9.figshare.979190

The documentation for the scripts has undergone an overhaul. The scripts now output extensive notes and the formal documentation website is generated from the scripts themselves and will never be out of sync.

https://khmer.readthedocs.org/en/latest/scripts.html

4.1.2 Notable bugs fixed/issues closed:

git clone of the khmer repo reqs > 0.5 GiB #223 @mr-c new khmer/file module #357 @RamRS Floating point exception in count-overlap.py #282 @qingpeng add documentation for sample-reads-randomly #192 @mr-c only build zlib and bzip2 when needed #168 @mr-c

4.1.3 Minor updates

khmer tools should output intelligent error messages when fed empty files #135 @RamRS set IParser::ParserState::ParserState::fill_id to zero at initialization #356 @mr-c demote nose & sphinx to extra dependencies. #351 @mr-c CID 1054792 (Medium) Uninitialized scalar field (UNINIT_CTOR) #179 @mr-c CID

1077117 (Medium): Division or modulo by zero (DIVIDE_BY_ZERO) #182 @mr-c if -savehash is specified then don't continue if there is not enough free disk space #245 @RamRS finish fixing implicit downcasts #330 @mr-c Clean up compile warnings in subset.cc #172 @mr-c all scripts need to output their version #236 @mr-c environmental variables need documenting #303 @mr-c C++ code should be consistently formatted #261 @mr-c Clean up ancillary files #146 @mr-c squash option not implemented in abundance-dist-single.py #271 @RamRS Add documentation on how to tie into a particular tagged version #29 @mr-c pip install -e fails with compile error #352 @mr-c remove the unused KTable object #337 @luizirber zlib 1.2.3 -> zlib 1.2.8 #336 @mr-c CID 1173035: Uninitialized scalar field (UNINIT_CTOR) #311 @mr-c CID 1153101: Resource leak in object (CTOR_DTOR_LEAK) #309 @mr-c remove khmer::read_parsers::IParser::ParserState::thread_id #323 @mr-c several modifications about count-overlap.py script #324 @qingpeng fixed runscript to handle SystemExit #332 @ctb CID 1063852: Uninitialized scalar field (UNINIT_CTOR) #313 @mr-c [infrastructure] update to new Doxyfile format, make version number autoupdate #315 @mr-c Removed an extraneous using namespace khmer; in kmer.hh, #276 @fishjord Minimum and recommended python version #94 @mr-c KmerCount class appears to be unused #302 @mr-c If loadhash is specified in e.g. normalize-by-median, don't complain about default hashsize parameters #117 @RamRS

4.1.4 Known Issues

All of these are pre-existing.

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

Some FASTQ files confuse our parser when running with more than one thread. For example, while using load-into-counting.py. If you experience this then add "-threads=1" to your command line. This issue is being tracked in https://github.com/ged-lab/khmer/issues/249

If your k-mer table (hashfile) gets truncated, perhaps from a full filesystem, then our tools currently will get stuck. This is being tracked in https://github.com/ged-lab/khmer/issues/247 and https://github.com/ged-lab/khmer/issues/96 and https://github.com/ged-lab/khmer/issues/246

Paired-end reads from Casava 1.8 currently require renaming for use in normalize-by-median and abund-filter when used in paired mode. The integration of a fix for this is being tracked in https://github.com/ged-lab/khmer/issues/23

annotate-partitions.py only outputs FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

A user reported that abundance-dist-single.py fails with small files and many threads. This issue is being tracked in https://github.com/ged-lab/khmer/issues/75

4.1.5 Contributors

@camillescott, @mr-c, @ctb, @luizirber, @RamRS, @qingpeng [0] http://semver.org/

4.2 khmer v1.0.1 release notes

This is bugfix release. Note: the installation instructions have been slightly simplified.

https://khmer.readthedocs.org/en/v1.0.1/

4.2.1 New items of note:

This release successfully installs and passes its unit tests on Debian 6.0 "Squeeze", Debian 7.0 "Wheezy", Fedora 19, OS X 7 "Lion", OS X 8 "Mountain Lion", Red Hat Enterprise Linux 6, Scientific Linux 6, Ubuntu 10.04 LTS, and Ubuntu 12.04 LTS. Thanks to the UW-Madison Build and Test Lab for their testing infrastructure.

4.2.2 Notable bugs fixed/issues closed:

fixed thread hanging issue #406 @ctb Explicit python2 invocation #404 @mr-c MANIFEST.in,setup.py: fix to correct zlib packaging #365 @mr-c fixed check_space_for_hashtable to use args.n_tables #382 @ctb Bug fix: make-initial-stoptags.py error on missing .ht input file, actual input file is .pt #391 @mr-c

4.2.3 Minor updates

include calc-best-assembly.py in v1.0.1 #409 @ctb updated normalize-by-median documentation for loadtable #378 @ctb updated diginorm for new FP rate info; corrected spelling error #398 @ctb Add spellcheck to code review checklist. #397 @ctb

4.2.4 Known Issues

All of these are pre-existing.

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

Some FASTQ files confuse our parser when running with more than one thread. For example, while using load-into-counting.py. If you experience this then add "-threads=1" to your command line. This issue is being tracked in https://github.com/ged-lab/khmer/issues/249

If your k-mer table (hashfile) gets truncated, perhaps from a full filesystem, then our tools currently will get stuck. This is being tracked in https://github.com/ged-lab/khmer/issues/247 and https://github.com/ged-lab/khmer/issues/246

Paired-end reads from Casava 1.8 currently require renaming for use in normalize-by-median and abund-filter when used in paired mode. The integration of a fix for this is being tracked in https://github.com/ged-lab/khmer/issues/23

annotate-partitions.py only outputs FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

A user reported that abundance-dist-single.py fails with small files and many threads. This issue is being tracked in https://github.com/ged-lab/khmer/issues/75

4.2.5 Contributors

@mr-c, @ctb, @luizirber, @RamRS, @ctSkennerton

4.3 khmer v1.1 release notes

This is v1.1, a minor version release; this version adds several new scripts.

Docs at: https://khmer.readthedocs.org/en/v1.1/

Release notes w/links: https://github.com/ged-lab/khmer/releases/tag/v1.1

4.3.1 New items of note:

- removed unnecessary files from PyPI package; distribution is now under 2 MB (#419) @mr-c
- tests are now distributed with package and can be run after 'pip install' (#451) @mr-c
- complain properly on file read failures (#333) @ctb
- Sequence loading scripts will now report total numbers of k-mers if given –report_total_kmers (#491/#429)
 @mr-c
- added metagenome protocol to acceptance testing (#472) @SherineAwad @ctb

4.3.2 Notable bugs fixed/issues closed:

- removed sandbox/load-into-hashbits.py (superseded by scripts/load-graph.py -no-tagset) (#480, @wrightmhw)
- promoted extract-long-sequences.py to scripts (#461, @wrightmhw)
- promoted fastq-to-fasta.py to scripts (#436, @wrightmhw)
- remove incorrect filesystem space check from abundance-dist.py (#452, @chuckpr)
- when counting hash writes fail, produce error message (#411, @znruss)
- removed a number of memory leaks found by Coverity and valgrind (#451, @mr-c)
- updated reservoir sampling to produce multiple subsamples with -S (#197, @ctb)
- fixed pip2, python2 issues (#428 and #485, @accaldwell @mr-c)
- removed untested/unused code and scripts (#438, @mr-c)

4.3.3 Known issues:

All of these are pre-existing.

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

Some FASTQ files confuse our parser when running with more than one thread. For example, while using load-into-counting.py. If you experience this then add "-threads=1" to your command line. This issue is being tracked in https://github.com/ged-lab/khmer/issues/249

If your k-mer table is truncated on write, an error may not be reported; this is being tracked in https://github.com/ged-lab/khmer/issues/443. However, khmer will now (correctly) fail when trying to read a truncated file (See #333).

Paired-end reads from Casava 1.8 currently require renaming for use in normalize-by-median and abund-filter when used in paired mode. The integration of a fix for this is being tracked in https://github.com/ged-lab/khmer/issues/23

Some scripts only output FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

A user reported that abundance-dist-single.py fails with small files and many threads. This issue is being tracked in https://github.com/ged-lab/khmer/issues/75

4.3.4 Contributors

@mr-c, @ctb, @camillescott, @wrightmhw, @chuckpr, @luizirber, @accaldwell, @znruss

4.4 khmer v1.2 release notes

This is the v1.2 release of khmer: minor new features and bug fixes. The start of this release cycle coincided with the Mozilla Science Lab Global Sprint 2014. We honor and thank the 19 new contributors (including four Michigan State University undergraduates) who volunteered their time to contribute!

Docs at: https://khmer.readthedocs.org/en/v1.2/

4.4.1 New items of note:

@mr-c and @ctb are proud to announce khmer's code of conduct http://khmer.readthedocs.org/en/v1.2/dev/CODE_OF_CONDUCT.html
#664 All scripts list which files have been created during their execution #477 @bocajnotnef All scripts now only
output status messages to STDERR instead of STDOUT #626 @b-wyss docs/ a fairly major re-organization and
brand new developer docs @ctb @mr-c load-into-counting.py: --summary-info: machine readable summary in
JSON or TSV format #649 @kdmurray91 scripts/extract-partitions.py: added documentation for .dist columns #516
@chuckpr Makefile: a new target make install-dependencies is useful for developers #539 @mr-c Sandbox
scripts have been cleaned up, or removed (see the sandbox/README.rst for details) #589 @ctb

4.4.2 Notable bugs fixed/issues closed:

do-partition.py's excessive spawning of threads fixed. #637 @camillescott Fixed unique k-mer count reporting in load-graph, load-into-counting, and normalize-by-median. #562 @mr-c Clarified and test the requirement for a 64-bit operating system #529 @Echelon9 Removed some of the broken multi-threading options #511 @majoras-masque Fix table.get("wrong_length_string") gives core dump #585 @Echelon9 filter-abund lists parameters that it doesn't use #524 @jstapleton Reduction of memory required to run the test suite #542 @leogargu BibTeX included in CITATIONS #541 @HLWiencko

4.4.3 Additional fixes/features

delete ScoringMatrix::assign as it is unused #502 @RodPic Root all of our C++ exceptions to a common base exception #508 @iglpdc deleted KhmerError #503 @drlabratory normalize-by-median reporting output after main loop exits, in case it hadn't been triggered #586 @ctb Many issues discovered by cppcheck cleaned up #506 @brtaylor92 Developers have a new Makefile target to autofix formatting: make format #612 @brtaylor92 normalize-by-median.py test coverage increased #361 @SherineAwad Several unused functions were removed #599 @brtaylor92 Developer docs now link to the stdc++ docs as appropriate #629 @mr-c Added tests for non-sequential access to input files #644 @bocajnotnef Removed khmer/theading_args.py #653 @bocajnotnef Improved test for maximum k value #658 @pgarland ReadParser no longer crashes if n_threads = 0 #86 @jiarong

4.4.4 Known issues:

All of these are pre-existing.

Multithreaded reading will drop reads. This major issue has been present for several khmer releases and was only found via a much larger test case that we had been previously using. Credit to @camillescott. Workaround: disable threading. The next release will fix this and the other FAST[AQ] parsing issues. https://github.com/ged-lab/khmer/issues/681

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

Some FASTQ files confuse our parser when running with more than one thread. For example, while using load-into-counting.py. If you experience this then add "-threads=1" to your command line. This issue is being tracked in https://github.com/ged-lab/khmer/issues/249

If your k-mer table is truncated on write, an error may not be reported; this is being tracked in https://github.com/ged-lab/khmer/issues/443. However, khmer will now (correctly) fail when trying to read a truncated file (See #333).

Paired-end reads from Casava 1.8 currently require renaming for use in normalize-by-median and abund-filter when used in paired mode. The integration of a fix for this is being tracked in https://github.com/ged-lab/khmer/issues/23

Some scripts only output FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

A user reported that abundance-dist-single.py fails with small files and many threads. This issue is being tracked in https://github.com/ged-lab/khmer/issues/75

4.4.5 Contributors

@mr-c, @ctb, *@bocajnotnef, *@Echelon9, *@jlippi, *@kdmurray91, @qingpeng, *@leogargu, *@jiarong, *@brtaylor92, *@iglpdc, @camillescott, *@HLWiencko, *@cowguru2000, *@drlabratory, *@jstapleton, *@bwyss, *@jgluck, @fishjord, *@SherineAwad, *@pgarland, *@majoras-masque, @chuckpr, *@RodPic, @luizirber, *@jrherr

* Denotes new contributor

4.5 khmer v1.3 release notes

This is the v1.3 release of khmer featuring a new FAST[AQ] parser from the SeqAn project.

Docs at: https://khmer.readthedocs.org/en/v1.3/

4.5.1 New items of note:

Fixes the two multithreaded reading of sequence files issues: FASTQ parsing and the recently found read dropping issue. Several khmer scripts now support reading from non-seekable plain and gziped FAST[AQ] files (a.k.a pipe or streaming support). @mr-c #642

4.5.2 Notable bugs fixed/issues closed:

restore threading to load-graph.py #699 @mr-c

4.5.3 Additional fixes/features

increase filter_abund.py coverage #568 @wrightmhw Provide scripts/ testing coverage for check_space_for_hashtable #386 #678 #718 @b-wyss Use absolute URI in CODE_OF_CONDUCT #684 @jsspencer give SeqAn credit #712 @mr-c Added testing to make sure all sandbox scripts are import-able and execfile-able. #709 @ctb reduce memory requirements to run tests #701 @ctb Two minor bug fixes to sandbox scripts #706 @ctb Upgrade of trim-low-abund for better, more profitable streaming. #601 @ctb Add –force or –expert or –ignore flag to all khmer scripts that do sanity checking #399 #647 @jessicamizzi Add XDECREF for returned read tuple in ReadParser.read_pair_iterator() #693 @mr-c @camillescott

4.5.4 Known issues:

All of these are pre-existing.

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in #266

If your k-mer table is truncated on write, an error may not be reported; this is being tracked in https://github.com/ged-lab/khmer/issues/443. However, khmer will now (correctly) fail when trying to read a truncated file (See #333).

Paired-end reads from Casava 1.8 currently require renaming for use in normalize-by-median and abund-filter when used in paired mode. The integration of a fix for this is being tracked in #23

Some scripts only output FASTA even if given a FASTQ file. This issue is being tracked in #46

A user reported that abundance-dist-single.py fails with small files and many threads. This issue is being tracked in #75

4.5.5 Contributors

@mr-c, @ctb, @camillescott, @b-wyss, @wrightmhw, @jsspencer

4.6 khmer v1.4 release notes

This is the v1.4 release of khmer featuring the results of our March and April (PyCon) coding sprints and the 16 new contributors; the use of the new v0.8 release of screed (the library we use for pure Python reading of nucleotide sequence files); and the addition of @luizirber's HyperLogLog counter for quick cardinality estimation.

Documentation is at https://khmer.readthedocs.org/en/v1.4/

4.6.1 New items of note:

Casava 1.8 read naming is now fully supported and in general the scripts no longer mangle read names. Side benefits: split-paired-reads.py will no longer drop reads with 'bad' names; count-median.py can generate output in CSV format. #759 #818 @ctb #873 @ahaerpfer

Most scripts now support a "broken" interleaved paired-read format for FASTA/ FASTQ nucleotide sequence files. 'trim-low-abund.py "has been promoted from the sandbox as well (with streaming support). #759 @ctb #963 @sguermond #933 @standage

The script to transform an interleaved paired-read nucleotide sequence file into two files now allows one to name the output files which can be useful in combination with named pipes for streaming processing #762 @ctb

Streaming everywhere: thanks to screed v0.8 we now support streaming of almost all inputs and outputs. #830 @aditi9783 #812 @mr-c #917 @bocajnotnef #882 @standage

Need a quick way to count total number of unique k-mers in very low memory? the unique-kmers.py script in the sandbox uses a HyperLogLog counter to quickly (and with little memory) provide an estimate with a controllable error rate. #257 #738 #895 #902 @luizirber

normalize-by-median.py can now process both a paired interleaved sequence file and a file of unpaired reads in the same invocation thus removing the need to write the counting table to disk as required in the workaround. #957 @susinmotion

4.6.2 Notable bugs fixed/issues closed:

Paired-end reads from Casava 1.8 no longer require renaming for use in normalize-by-median.py and abund-filter.py when used in paired mode #818 @ctb

Python version support clarified. We do not (yet) support Python 3.x #741 @mr-c

If a single output file mode is chosen for normalize-by-median.py we now default to overwriting the output. Appending the output is available by using the append redirection operator from the shell. #843 @drtamermansour

Scripts that consume sequence data using C++ will now properly throw an error on truncated files. #897 @kdmurray91 And while writing to disk we properly check for errors #856 #962 @mr-c

abundance-dist-single.py no longer fails with small files and many threads. #900 @mr-c

4.6.3 Additional fixes/features

Of interest to users:

Many documentation updates #753 @PamelaM, #782 @bocajnotnef, #845 @alameldin, #804 @ctb, #870 @SchwarzEM, #953 #942 @safay, #929,@davelin1, #687 #912 #926 @mr-c

Installation instructions for Conda, Arch Linux, and Mac Ports have been added #723 @reedacartwright #952 @elmbeech #930 @ahaerpfer

The example script for the STAMPS database has been fixed to run correctly #781 @drtamermansour

split-paired-reads.py: added -o option to allow specification of an output directory #752 @bede

Fixed a string formatting and a boundry error in sample-reads-randomly.py #773 @qingpeng #995 @ctb

CSV output added to abundance-dist.py, abundance-dist-single.py, and count-overlap.py, and readstats.py #831 #854 #855 @drtamermansour #959 @anotherthomas

TSV/JSON output of load-into-counting.py enhanced with the total number of reads processed #996 @kd-murray91 Output files are now also checked to be writable *before* loading the input files #672 @pgarland @bocajnotnef

interleave-reads.py now prints the output filename nicely #827 @kdmurray91

Cleaned up error for input file not existing #772 @jessicamizzi #851 @ctb

Fixed error in find-knots.py #860 @TheOneHyer

The help text for load-into-counting.py for the --no-bigcounts/-b flag has been clarified #857 @kd-murray91

@lexnederbragt confirmed an old bug has been fixed with his test for whitespace in sequence identifiers interacting with the extract-partitions.py script #979

Now safe to copy-and-paste from the user documentation as the smart quotes have been turned off. #967 @ahaerpfer

The script make-coverage.py has been restored to the sandbox. #920 @SherineAwad

normalize-by-median.py will warn if two of the input files have the same name #932 @elmbeech

Of interest to developers:

Switched away from using --user install for developers #740 @mr-c @drtamermansour & #883 @standage

Developers can now see a summary of important Makefile targets via make help #783 @standage

The unused khmer.load_pe module has been removed #828 @kdmurray91

Versioneer bug due to new screed release was squashed #835 @mr-c

A Python 2.6 and 2.7.2 specific bug was worked around #869 @kdmurray91 @ctb

Added functions hash_find_all_tags_list and hash_get_tags_and_positions to CountingHash objects #749 #765 @ctb

The make diff-cover and ChangeLog formatting requirements have been added to checklist #766 @mr-c

A useful message is now presented if large tables fail to allocate enough memory #704 @mr-c

A checklist for developers adding new CPython types was added #727 @mr-c

The sandbox graduation checklist has been updated to include streaming support #951 @sguermond

Specific policies for sandbox/ and scripts/ content, and a process for adding new command line scripts into scripts/ have been added to the developer documentation #799 @ctb

Sandbox scripts update: corrected #! Python invocation #815 @Echelon9, executable bits, copyright headers, no underscores in filenames #823 #826 #850 @alameldin several scripts deleted, docs + requirements updated #852 @ctb

Avoid running big-memory tests on OS X #819 @ctb

Unused callback code was removed #698 @mr-c

The CPython code was updated to use the new checklist and follow additional best practices #785 #842 @luizirber

Added a read-only view of the raw counting tables #671 @camillescott #869 @kdmurray91

Added a Python method for quickly getting the number of underlying tables in a counting or presence table #879 #880 @kdmurray91

The C++ library can now be built separately for the brave and curious developer #788 @kdmurray91

The ReadParser object now keeps track of the number of reads processed #877 @kdmurray91

Documentation is now reproducible #886 @mr-c

Python future proofing: specify floor division #863 @mr-c

Miscellaneous spelling fixes; thanks codespell! #867 @mr-c

Debian package list update #984 @mr-c

khmer.kfile.check_file_status() has been renamed to check_input_files() #941 @proteasome filter-abund.py now uses it to check the input counting table #931 @safay

normalize-by-median.py was refactored to not pass the ArgParse object around #965 @susinmotion

Developer communication has been clarified #969 @sguermond

Tests using the 'fail_okay=true' parameter to runscript have been updated to confirm the correct error occurred. 3 faulty tests were fixed and the docs were clarified #968 #971 @susinmotion

FASTA test added for extract-long-sequences.py #901 @jessicamizzi

'added silly test for empty file warning' #557 @wltrimbl @bocajnotnef

A couple tests were made more resilient and some extra error checking added in CPython land #889 @mr-c

Copyright added to pull request checklist #940 @sguermond

khmer_exceptions are now based on std::strings which plugs a memory leak #938 @anotherthomas

Python docstrings were made PEP257 compliant #936 @ahaerpfer

Some C++ comments were converted to be Doxygen compliant #950 @josiahseaman

The counting and presence table warning logic was refactored and centralized #944 @susinmotion

The release checklist was updated to better run the post-install tests #911 @mr-c

The unused method find_all_tags_truncate_on_abundance was removed from the CPython API #924 @anotherthomas

OS X warnings quieted #887 @mr-c

4.6.4 Known issues:

All of these are pre-existing.

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

Some scripts only output FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

4.6.5 Contributors

```
@ctb, @kdmurray91, @mr-c, @drtamermansour, @luizirber, @standage, @bocajnotnef, *@susinmotion, @jessicamizzi, *@elmbeech, *@anotherthomas, *@sguermond, *@ahaerpfer, *@alameldin, *@TheOneHyer, *@aditi9783, *@proteasome, *@bede, *@davelin1, @Echelon9, *@reedacartwright, @qingpeng, *@SchwarzEM, *@scottsievert, @PamelaM, @SherineAwad, *@josiahseaman, *@lexnederbragt,
```

4.6.6 Issue reporters

@moorepants, @teshomem, @macmanes, @lexnederbragt, @r-gaia-cs, @magentashades

4.7 khmer v1.4.1 release notes

This is the v1.4.1 release of khmer. Due to the upcoming Python 3 compatibility in both khmer and Screed we need to modify the dependency between khmer and the Screed library to be only the existing version 0.8, and not some future version.

If you have khmer 1.4 installed then there is no benefit to upgrading; this point release is to keep pip install khmer still working when we release the next version of Screed with Python 3 support. The next version of khmer, v2.0, will also have Python 3 support.

Documentation is at https://khmer.readthedocs.org/en/v1.4.1/ (no changes from v1.4)

4.7.1 Known issues:

All of these are pre-existing.

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

Some scripts only output FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

^{*} Indicates new contributors

The khmer user documentation

Contents:

5.1 Installing and running khmer

You'll need a 64-bit operating system, Python 2.7.x and internet access.

The khmer project currently works with Python 2.6 but we target Python 2.7.x.

5.1.1 Build requirements

OS X

1. From a terminal download the virtualenv package and create a virtual environment with it:

```
curl -O https://pypi.python.org/packages/source/v/virtualenv/virtualenv-1.11.6.tar.gz
tar xzf virtualenv*
cd virtualenv-*; python2.7 virtualenv.py ../khmerEnv; cd ..
source khmerEnv/bin/activate
```

Linux

- 1. Install the python development environment, virtualenv, pip, gcc, and g++.
 - On recent Debian and Ubuntu this can be done with:

```
sudo apt-get install python2.7-dev python-virtualenv python-pip gcc \
    g++
```

• For RHEL6:

```
sudo yum install -y python-devel python-pip git gcc gcc-c++ make sudo pip install virtualenv
```

2. Create a virtualenv and activate it:

```
cd a/writable/directory/
python2.7 -m virtualenv khmerEnv
source khmerEnv/bin/activate
```

Linux users without root access can try the OS X instructions above.

5.1.2 Installing khmer inside the virtualenv

1. Use pip to download, build, and install khmer and its dependencies:

```
pip2 install khmer
```

- 2. The scripts are now in the env/bin directory and ready for your use. You can directly use them by name, see khmer's command-line interface.
- 3. When returning to khmer after installing it you will need to reactivate the virtualenv first:

```
source khmerEnv/bin/activate
```

Run the tests

After installing you can run the embedded test suite:

```
nosetests khmer --attr '!known_failing'
```

If the nosetests binary isn't installed then:

```
pip2 install khmer[tests]
nosetests khmer --attr '!known_failing'
```

5.2 A few examples

See the 'examples' subdirectory for complete examples.

5.2.1 STAMPS data set

The 'stamps' data set is a fake metagenome-like data set containing two species, mixed at a 10:1 ratio. The source genomes are in 'data/stamps-genomes.fa'. The reads file is in 'data/stamps-reads.fa.gz', and consists of 100-base reads with a 1% error rate.

The example shows how to construct k-mer abundance histograms, as well as the effect of digital normalization and partitioning on the k-mer abundance distribution.

See the script for running everything and the IPython Notebook.

For an overall discussion and some slides to explain what's going on, visit the Web site for a 2013 HMP metagenome assembly webinar that Titus Brown gave.

5.3 An assembly handbook for khmer - rough draft

```
date 2012-11-2
```

An increasing number of people are asking about using our assembly approaches for things that we haven't yet written (or posted) papers about. Moreover, our assembly strategies themselves are also under constant evolution as we do more research and find ever-wider applicability of our approaches.

Note, this is an exact copy of Titus' blog post, here – go check the bottom of that for comments.

5.3.1 Authors

This handbook distills the cumulative expertise of Adina Howe, Titus Brown, Erich Schwarz, Jason Pell, Camille Scott, Elijah Lowe, Kanchan Pavangadkar, Likit Preeyanon, and others.

5.3.2 Introduction

khmer is a general framework for low-memory k-mer counting, filtering, and advanced trickery.

The latest source is always available here.

khmer is really focused on short read data, and, more specifically, Illumina, because that's where we have a too-much-data problem. However, a lot of the prescriptions below can be adapted to longer read technologies such as 454 and Ion Torrent without much effort.

Don't try to use our k-mer approaches with PacBio – the error rate is too high.

There are currently two papers available on khmer: the partitioning paper and the digital normalization paper.

There are many blog posts about this stuff on Titus Brown's blog. We will try to link them in where appropriate.

5.3.3 Asking for help

There's some documentation here:

https://khmer.readthedocs.org/en/latest/

There's also a khmer mailing list at lists.idyll.org that you can use to get help with khmer. To sign up, just go to the khmer lists page and subscribe.

5.3.4 Preparing your sequences

Do all the quality filtering, trimming, etc. that you think you should do.

Most of the khmer tools currently work "out of the box" on interleaved paired-end data. Ask on the list if you're not sure.

All of our scripts will take in .fq or .fastq files as FASTQ, and all other files as FASTA. gzip files are always accepted. Let us know if not; that's a bug!

Most scripts *output* FASTA, and some mangle headers. Sorry. We're working on outputting FASTQ for FASTQ input, and removing any header mangling.

5.3.5 Picking k-mer table sizes and k parameters

For k-mer table sizes, read Choosing table sizes for khmer

For k-mer sizes, we recommend k=20 for digital normalization and k=32 for partitioning; then assemble with a variety of k parameters.

5.3.6 Genome assembly, including MDA samples and highly polymorphic genomes

1. Apply digital normalization as follows.

Broadly, normalize each insert library separately, in the following way:

For high-coverage libraries ($> \sim 50x$), do three-pass digital normalization: run normalize-by-median to C=20 and then run filter-abund with C=1. Now split out the remaining paired-end/interleaved and single-end reads using strip-and-split-for-assembly, and normalize-by-median the paired-end and single-end files to C=5 (in that order).

For low-coverage libraries (< 50x) do single-pass digital normalization: run normalize-by-median to C=10.

- 2. Extract any remaining paired-end reads and lump remaining orphan reads into singletons using strip-and-split-for-assembly
- 3. Then assemble as normal, with appropriate insert size specs etc. for the paired end reads.

You can read about this process in the digital normalization paper.

5.3.7 mRNAseq assembly

1. Apply single-pass digital normalization.

Run normalize-by-median to C=20.

- 2. Extract any remaining paired-end reads and lump remaining orphan reads into singletons using strip-and-split-for-assembly
- 3. Then assemble as normal, with appropriate insert size specs etc. for the paired end reads.

You can read about this process in the digital normalization paper.

5.3.8 Metagenome assembly

1. Apply single-pass digital normalization.

Run normalize-by-median to C=20 (we've also found C=10 works fine).

- 2. Run filter-below-abund with C=50 (if you diginormed to C=10) or C=100 (if you diginormed to C=20);
 - 3. Partition reads with load-graph, etc. etc.
- 4. Assemble groups as normal, extracting paired-end reads and lumping remaining orphan reads into singletons using strip-and-split-for-assembly.

(We actually use Velvet at this point, but there should be no harm in using a metagenome assembler such as MetaVelvet or MetaIDBA or SOAPdenovo.)

Read more about this in the partitioning paper. We have some upcoming papers on partitioning and metagenome assembly, too; we'll link those in when we can.

5.3.9 Metatranscriptome assembly

(Not tested by us!)

1. Apply single-pass digital normalization.

Run normalize-by-median to C=20.

- 2. Extract any remaining paired-end reads and lump remaining orphan reads into singletons using strip-and-split-for-assembly
- 3. Then assemble with a genome or metagenome assembler, *not* an mRNAseq assembler. Use appropriate insert size specs etc. for the paired end reads.

5.3.10 Preprocessing Illumina for other applications

(Not tested by us!)

Others have told us that you can apply digital normalization to Illumina data prior to using Illumina for RNA scaffolding or error correcting PacBio reads.

Our suggestion for this, based on no evidence whatsoever, is to diginorm the Illumina data to C=20.

5.3.11 Quantifying mRNAseq or metagenomes assembled with digital normalization

For now, khmer only deals with assembly! So: assemble. Then, go back to your original, unnormalized reads, and map those to your assembly with e.g. bowtie. Then count as you normally would:).

5.3.12 Philosophy of digital normalization

The basic philosophy of digital normalization is "load your most valuable reads first." Diginorm gets rid of redundancy iteratively, so you are more likely to retain the first reads fed in; this means you should load in paired end reads, or longer reads, first.

5.3.13 Iterative and independent normalization

You can use --loadtable and --savetable to do iterative normalizations on multiple files in multiple steps. For example, break

```
normalize-by-median.py [ ... ] file1.fa file2.fa file3.fa
```

into multiple steps like so:

```
normalize-by-median.py [ ... ] --savetable file1.ct file1.fa
normalize-by-median.py [ ... ] --loadtable file1.ct --savetable file2.ct file2.fa
normalize-by-median.py [ ... ] --loadtable file2.ct --savetable file3.ct file3.fa
```

The results should be identical!

If you want to independently normalize multiple files for speed reasons, go ahead. Just remember to do a combined normalization at the end. For example, instead of

```
normalize-by-median.py [ ... ] file1.fa file2.fa file3.fa
```

you could do

```
normalize-by-median.py [ ... ] file1.fa
normalize-by-median.py [ ... ] file2.fa
normalize-by-median.py [ ... ] file3.fa
```

and then do a final

```
normalize-by-median.py [ ... ] file1.fa.keep file2.fa.keep file3.fa.keep
```

The results will not be identical, but should not differ significantly. The multipass approach will take more total time but may end up being faster walltime because you can execute the independent normalizations on multiple computers.

For a cleverer approach that we will someday implement, read the Beachcomber's Dilemma.

5.3.14 Validating and comparing assemblies

More here soon:).

5.4 khmer's command-line interface

The simplest way to use khmer's functionality is through the command line scripts, located in the scripts/ directory of the khmer distribution. Below is our documentation for these scripts. Note that all scripts can be given -h which will print out a list of arguments taken by that script.

Many scripts take -x and -x parameters, which drive khmer's memory usage. These parameters depend on details of your data set; for more information on how to choose them, see Choosing table sizes for khmer.

You can also override the default values of --ksize/-k, --n_tables/-N, and --min-tablesize/-x with the environment variables *KHMER KSIZE*, *KHMER N TABLES*, and *KHMER MIN TABLESIZE* respectively.

- 1. k-mer counting and abundance filtering
- 2. Partitioning
- 3. Digital normalization
- 4. Read handling: interleaving, splitting, etc.

Note: Almost all scripts take in either FASTA and FASTQ format, and output the same. Some scripts may only recognize FASTQ if the file ending is '.fq' or '.fastq', at least for now.

Files ending with '.gz' will be treated as gzipped files, and files ending with '.bz2' will be treated as bzip2'd files.

5.4.1 k-mer counting and abundance filtering

load-into-counting.py

Build a k-mer counting table from the given sequences.

usage: load-into-counting.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-threads THREADS] [-b] [-summary-info FORMAT] [-report-total-kmers] [-f] output_countingtable_filename input_sequence_filename [input_sequence_filename ...]

output_countingtable_filename

The name of the file to write the k-mer counting table to.

input_sequence_filename

The names of one or more FAST[AQ] input sequence files.

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

```
--ksize \langle int \rangle, -k \langle int \rangle
```

k-mer size to use

--n_tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--threads <int>, -T <int>

Number of simultaneous threads to execute

-b, --no-bigcount

The default behaviour is to count past 255 using bigcount. This flag turns bigcount off, limiting counts to 255.

--summary-info <format>, -s <format>

What format should the machine readable run summary be in? (json or tsv, disabled by default)

--report-total-kmers, -t

Prints the total number of k-mers to stderr

-f, --force

Overwrite output file if it exists

Note: with -b the output will be the exact size of the k-mer counting table and this script will use a constant amount of memory. In exchange k-mer counts will stop at 255. The memory usage of this script with -b will be about 1.15x the product of the -x and -N numbers.

Example:

```
load-into-counting.py -k 20 -x 5e7 out.ct data/100k-filtered.fa
```

Multiple threads can be used to accelerate the process, if you have extra cores to spare.

Example:

load-into-counting.py -k 20 -x 5e7 -T 4 out.ct data/100k-filtered.fa

abundance-dist.py

Calculate abundance distribution of the k-mers in the sequence file using a pre-made k-mer counting table.

usage: abundance-dist.py [-h] [-z] [-s] [-csv] [-version] [-f] input_counting_table_filename input_sequence_filename output_histogram_filename

input_counting_table_filename

The name of the input k-mer counting table file.

input_sequence_filename

The name of the input FAST[AQ] sequence file.

output_histogram_filename

The columns are: (1) k-mer abundance, (2) k-mer count, (3) cumulative count, (4) fraction of total distinct k-mers.

-h, --help

show this help message and exit

-z, --no-zero

Do not output 0-count bins

-s, --squash

Overwrite existing output_histogram_filename

--csv

Use the CSV format for the histogram. Includes column headers.

--version

show program's version number and exit

-f, --force

Continue even if specified input files do not exist or are empty.

abundance-dist-single.py

Calculate the abundance distribution of k-mers from a single sequence file.

usage: abundance-dist-single.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-threads THREADS] [-z] [-b] [-s] [-csv] [-savetable filename] [-report-total-kmers] [-f] input_sequence_filename output_histogram_filename

input_sequence_filename

The name of the input FAST[AQ] sequence file.

output_histogram_filename

The name of the output histogram file. The columns are: (1) k-mer abundance, (2) k-mer count, (3) cumulative count, (4) fraction of total distinct k-mers.

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n_tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--threads <int>, -T <int>

Number of simultaneous threads to execute

-z, --no-zero

Do not output 0-count bins

-b, --no-bigcount

Do not count k-mers past 255

-s, --squash

Overwrite output file if it exists

--csv

Use the CSV format for the histogram. Includes column headers.

--savetable <filename>

Save the k-mer counting table to the specified filename.

--report-total-kmers, -t

Prints the total number of k-mers to stderr

-f, --force

Overwrite output file if it exists

Note that with -b this script is constant memory; in exchange, k-mer counts will stop at 255. The memory usage of this script with -b will be about 1.15x the product of the -x and -N numbers.

To count k-mers in multiple files use load_into_counting.py and abundance_dist.py.

filter-abund.py

Trim sequences at a minimum k-mer abundance.

usage: filter-abund.py [-h] [-threads THREADS] [-cutoff CUTOFF] [-variable-coverage] [-normalize-to NORMAL-IZE_TO] [-o optional_output_filename] [-version] [-f] input_counting_table_filename input_sequence_filename [input_sequence_filename ...]

input_counting_table_filename

The input k-mer counting table filename

input_sequence_filename

Input FAST[AQ] sequence filename

-h, --help

show this help message and exit

```
--threads <int>, -T <int>
```

Number of simultaneous threads to execute

```
--cutoff <int>, -C <int>
```

Trim at k-mers below this abundance.

--variable-coverage, -V

Only trim low-abundance k-mers from sequences that have high coverage.

```
--normalize-to <int>, -Z <int>
```

Base the variable-coverage cutoff on this median k-mer abundance.

```
-o <optional_output_filename>, --out <optional_output_filename>
```

Output the trimmed sequences into a single file with the given filename instead of creating a new file for each input file.

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

Trimmed sequences will be placed in \${input_sequence_filename}.abundfilt for each input sequence file. If the input sequences are from RNAseq or metagenome sequencing then --variable-coverage should be used.

Example:

```
load-into-counting.py -k 20 -x 5e7 table.ct data/100k-filtered.fa filter-abund.py -C 2 table.ct data/100k-filtered.fa
```

filter-abund-single.py

Trims sequences at a minimum k-mer abundance (in memory version).

usage: filter-abund-single.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-threads THREADS] [-cutoff CUTOFF] [-savetable filename] [-report-total-kmers] [-f] input_sequence_filename

input_sequence_filename

FAST[AQ] sequence file to trim

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--threads <int>, -T <int>

Number of simultaneous threads to execute

--cutoff <int>, -C <int>

Trim at k-mers below this abundance.

--savetable <filename>

If present, the name of the file to save the k-mer counting table to

--report-total-kmers, -t

Prints the total number of k-mers to stderr

-f, --force

Overwrite output file if it exists

Trimmed sequences will be placed in \${input_sequence_filename}.abundfilt.

This script is constant memory.

To trim reads based on k-mer abundance across multiple files, use **load-into-counting.py** and **filter-abund.py**.

Example:

```
filter-abund-single.py -k 20 -x 5e7 -C 2 data/100k-filtered.fa
```

trim-low-abund.py

Trim low-abundance k-mers using a streaming algorithm.

usage: trim-low-abund.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-cutoff CUTOFF] [-normalize-to NORMALIZE_TO] [-o filename] [-variable-coverage] [-l filename] [-s filename] [-force] [-ignore-pairs] [-tempdir TEMPDIR] input_filenames [input_filenames ...]

input_filenames

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

```
--ksize <int>, -k <int>
k-mer size to use

--n_tables <int>, -N <int>
number of k-mer counting tables to use

--min-tablesize <float>, -x <float>
lower bound on tablesize to use

--cutoff <int>, -C <int>
remove k-mers below this abundance

--normalize-to <int>, -Z <int>
base cutoff on this median k-mer abundance
```

-o <filename>, --out <filename>
 only output a single file with the specified filename; use a single dash "-" to specify that output should go to
 STDOUT (the terminal)

--variable-coverage, -V

Only trim low-abundance k-mers from sequences that have high coverage.

- -1 <filename>, --loadtable <filename> load a precomputed k-mer table from disk
- -s <filename>, --savetable <filename>
 save the k-mer counting table to disk after allreads are loaded.
- --force
- --ignore-pairs

```
--tempdir <str>, -T <str>
```

The output is one file for each input file, <input file>.abundtrim, placed in the current directory. This output contains the input sequences trimmed at low-abundance k-mers.

The -V/--variable-coverage parameter will, if specified, prevent elimination of low-abundance reads by only trimming low-abundance k-mers from high-abundance reads; use this for non-genomic data sets that may have variable coverage.

Note that the output reads will not necessarily be in the same order as the reads in the input files; if this is an important consideration, use load-into-counting.py and filter-abund.py. However, read pairs will be kept together, in "broken-paired" format; you can use extract-paired-reads.py to extract read pairs and orphans.

Example:

```
trim-low-abund.py -x 5e7 -k 20 -C 2 data/100k-filtered.fa
```

count-median.py

Count k-mers summary stats for sequences

usage: count-median.py [-h] [-version] [-f] [-csv] input_counting_table_filename input_sequence_filename out-put_summary_filename

input_counting_table_filename

input k-mer count table filename

input sequence filename

input FAST[AQ] sequence filename

output_summary_filename

output summary filename

-h, --help

show this help message and exit

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

--csv

Use the CSV format for the histogram. Includes column headers.

Count the median/avg k-mer abundance for each sequence in the input file, based on the k-mer counts in the given k-mer counting table. Can be used to estimate expression levels (mRNAseq) or coverage (genomic/metagenomic).

The output file contains sequence id, median, average, stddev, and seq length; fields are separated by spaces. For khmer 1.x count-median.py will split sequence names at the first space which means that some sequence formats (e.g. paired FASTQ in Casava 1.8 format) will yield uninformative names. Use --csv to fix this behavior.

Example:

```
count-median.py counts.ct tests/test-data/test-reads.fq.gz medians.txt
```

NOTE: All 'N's in the input sequences are converted to 'G's.

count-overlap.py

Count the overlap k-mers which are the k-mers appearing in two sequence datasets.

usage: count-overlap.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-csv] [-f] input_presence_table_filename input_sequence_filename output_report_filename

input_presence_table_filename

input k-mer presence table filename

$\verb"input_sequence_filename"$

input sequence filename

output_report_filename

output report filename

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n_tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--csv

Use the CSV format for the curve output in \${output_report_filename}.curve, including column headers.

-f, --force

Overwrite output file if it exists

An additional report will be written to \${output_report_filename}.curve containing the increase of overlap k-mers as the number of sequences in the second database increases.

5.4.2 Partitioning

do-partition.py

Load, partition, and annotate FAST[AQ] sequences

usage: do-partition.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-threads THREADS] [-subset-size SUBSET_SIZE] [-no-big-traverse] [-keep-subsets] [-f] graphbase input_sequence_filename [input_sequence_filename ...]

graphbase

base name for output files

input_sequence_filename

input FAST[AQ] sequence filenames

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--threads <int>, -T <int>

Number of simultaneous threads to execute

--subset-size <float>, -s <float>

Set subset size (usually 1e5-1e6 is good)

--no-big-traverse

Truncate graph joins at big traversals

--keep-subsets

Keep individual subsets (default: False)

-f, --force

Overwrite output file if it exists

Load in a set of sequences, partition them, merge the partitions, and annotate the original sequences files with the partition information.

This script combines the functionality of load-graph.py, partition-graph.py, merge-partitions.py, and annotate-partitions.py into one script. This is convenient but should probably not be used for large data sets, because do-partition.py doesn't provide save/resume functionality.

load-graph.py

Load sequences into the compressible graph format plus optional tagset.

usage: load-graph.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-threads THREADS] [-no-build-tagset] [-report-total-kmers] [-write-fp-rate] [-f] output_presence_table_filename input_sequence_filename [input_sequence_filename ...]

output_presence_table_filename

output k-mer presence table filename.

input_sequence_filename

input FAST[AQ] sequence filename

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--threads <int>, -T <int>

Number of simultaneous threads to execute

--no-build-tagset, -n

Do NOT construct tagset while loading sequences

--report-total-kmers, -t

Prints the total number of k-mers to stderr

--write-fp-rate, -w

Write false positive rate into .info file

-f, --force

Overwrite output file if it exists

See **extract-partitions.py** for a complete workflow.

partition-graph.py

Partition a sequence graph based upon waypoint connectivity

usage: partition-graph.py [-h] [-stoptags filename] [-subset-size SUBSET_SIZE] [-no-big-traverse] [-version] [-f] [-threads THREADS] basename

basename

basename of the input k-mer presence table + tagset files

-h, --help

show this help message and exit

--stoptags <filename>, -S <filename>

Use stoptags in this file during partitioning

--subset-size <float>, -s <float>

Set subset size (usually 1e5-1e6 is good)

--no-big-traverse

Truncate graph joins at big traversals

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

--threads <int>, -T <int>

Number of simultaneous threads to execute

The resulting partition maps are saved as '\${basename}.subset.#.pmap' files.

See 'Artifact removal' to understand the stoptags argument.

merge-partition.py

Merge partition map '.pmap' files.

usage: merge-partition.py [-h] [-ksize KSIZE] [-keep-subsets] [-version] [-f] graphbase

graphbase

basename for input and output files

-h, --help

show this help message and exit

--ksize <int>, -k <int>

k-mer size (default: 32)

--keep-subsets

Keep individual subsets (default: False)

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

Take the \${graphbase}.subset.#.pmap files and merge them all into a single \${graphbase}.pmap.merged file for annotate-partitions.py to use.

annotate-partitions.py

Annotate sequences with partition IDs.

usage: annotate-partitions.py [-h] [-ksize KSIZE] [-version] [-f] graphbase input_sequence_filename [input_sequence_filename ...]

graphbase

basename for input and output files

input sequence filename

input FAST[AQ] sequences to annotate.

-h, --help

show this help message and exit

```
--ksize <int>, -k <int>
```

k-mer size (default: 32)

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

Load in a partitionmap (generally produced by partition-graph.py or merge-partitions.py) and annotate the sequences in the given files with their partition IDs. Use **extract-partitions.py** to extract sequences into separate group files.

Example (results will be in random-20-a.fa.part):

```
load-graph.py -k 20 example tests/test-data/random-20-a.fa partition-graph.py example merge-partitions.py -k 20 example annotate-partitions.py -k 20 example tests/test-data/random-20-a.fa
```

extract-partitions.py

Separate sequences that are annotated with partitions into grouped files.

usage: extract-partitions.py [-h] [-max-size MAX_SIZE] [-min-partition-size MIN_PART_SIZE] [-no-output-groups] [-output-unassigned] [-version] [-f] output_filename_prefix input_partition_filename [in-put_partition_filename ...]

```
output_filename_prefix
```

```
input_partition_filename
```

-h, --help

show this help message and exit

```
--max-size <int>, -X <int>
```

Max group size (n sequences)

--min-partition-size <int>, -m <int>

Minimum partition size worth keeping

--no-output-groups, -n

Do not actually output groups files.

--output-unassigned, -U

Output unassigned sequences, too

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

Example (results will be in example.group0000.fa):

```
load-graph.py -k 20 example tests/test-data/random-20-a.fa
partition-graph.py example
merge-partitions.py -k 20 example
annotate-partitions.py -k 20 example tests/test-data/random-20-a.fa
extract-partitions.py example random-20-a.fa.part

(extract-partitions.py will produce a partition size distribution
in <br/>base>.dist. The columns are: (1) number of reads, (2) count
of partitions with n reads, (3) cumulative sum of partitions,
(4) cumulative sum of reads.)
```

Artifact removal

The following scripts are specialized scripts for finding and removing highly-connected k-mers (HCKs). See Partitioning large data sets (50m+ reads).

make-initial-stoptags.py

Find an initial set of highly connected k-mers.

usage: make-initial-stoptags.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-subset-size SUBSET_SIZE] [-stoptags filename] [-f] graphbase

graphbase

basename for input and output filenames

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n_tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

--subset-size <float>, -s <float>

Set subset size (default 1e4 is prob ok)

--stoptags <filename>, -S <filename>

Use stoptags in this file during partitioning

-f, --force

Overwrite output file if it exists

Loads a k-mer presence table/tagset pair created by load-graph.py, and does a small set of traversals from graph waypoints; on these traversals, looks for k-mers that are repeatedly traversed in high-density regions of the graph, i.e. are highly connected. Outputs those k-mers as an initial set of stoptags, which can be fed into partition-graph.py, find-knots.py, and filter-stoptags.py.

The k-mer counting table size options parameters are for a k-mer counting table to keep track of repeatedly-traversed k-mers. The subset size option specifies the number of waypoints from which to traverse; for highly connected data sets, the default (1000) is probably ok.

find-knots.py

Find all highly connected k-mers.

usage: find-knots.py [-h] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-version] graphbase

graphbase

Basename for the input and output files.

-h, --help

show this help message and exit

--n_tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on the size of the k-mer counting table(s)

--version

show program's version number and exit

Load an k-mer presence table/tagset pair created by load-graph, and a set of pmap files created by partition-graph. Go through each pmap file, select the largest partition in each, and do the same kind of traversal as in **make-initial-stoptags.py** from each of the waypoints in that partition; this should identify all of the HCKs in that partition. These HCKs are output to <graphbase>.stoptags after each pmap file.

Parameter choice is reasonably important. See the pipeline in Partitioning large data sets (50m+ reads) for an example run.

This script is not very scalable and may blow up memory and die horribly. You should be able to use the intermediate stoptags to restart the process, and if you eliminate the already-processed pmap files, you can continue where you left off.

filter-stoptags.py

Trim sequences at stoptags.

usage: filter-stoptags.py [-h] [-ksize KSIZE] [-version] [-f] input_stoptags_filename input_sequence_filename [input_sequence_filename ...]

```
input_stoptags_filename
```

input sequence filename

-h, --help

show this help message and exit

--ksize <int>, -k <int>
 k-mer size

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

Load stoptags in from the given .stoptags file and use them to trim or remove the sequences in <file1-N>. Trimmed sequences will be placed in <fileN>.stopfilt.

5.4.3 Digital normalization

normalize-by-median.py

Do digital normalization (remove mostly redundant sequences)

usage: normalize-by-median.py [-h] [-version] [-q] [-ksize KSIZE] [-n_tables N_TABLES] [-min-tablesize MIN_TABLESIZE] [-C CUTOFF] [-p] [-u unpaired_reads_filename] [-s filename] [-R filename] [-f] [-save-on-failure] [-d DUMP_FREQUENCY] [-o filename] [-report-total-kmers] [-force] [-l filename] in-put_sequence_filename [input_sequence_filename ...]

input_sequence_filename

Input FAST[AQ] sequence filename.

-h, --help

show this help message and exit

--version

show program's version number and exit

-q, --quiet

--ksize <int>, -k <int>

k-mer size to use

--n_tables <int>, -N <int>

number of k-mer counting tables to use

--min-tablesize <float>, -x <float>

lower bound on tablesize to use

-C <int>, --cutoff <int>

-p, --paired

- -u <unpaired_reads_filename>, --unpaired-reads <unpaired_reads_filename>
 with paired data only, include an unpaired file
- -s <filename>, --savetable <filename>
 save the k-mer counting table to disk after allreads are loaded.
- -R <filename>, --report <filename>

-f, --fault-tolerant

continue on next file if read errors are encountered

--save-on-failure

Save k-mer counting table when an error occurs

-d <int>, --dump-frequency <int>
dump k-mer counting table every d files

-o <filename>, --out <filename>

only output a single file with the specified filename; use a single dash "-" to specify that output should go to STDOUT (the terminal)

--report-total-kmers, -t

Prints the total number of k-mers post-normalization to stderr

--force

Overwrite output file if it exists

-1 <filename>, --loadtable <filename>
load a precomputed k-mer table from disk

Discard sequences based on whether or not their median k-mer abundance lies above a specified cutoff. Kept sequences will be placed in <fileN>.keep.

Paired end reads will be considered together if -p is set. If either read will be kept, then both will be kept. This should result in keeping (or discarding) each sequencing fragment. This helps with retention of repeats, especially. With coption: -ul-unpaired-reads, unpaired reads from the specified file will be read after the paired data is read.

With -s/-savetable, the k-mer counting table will be saved to the specified file after all sequences have been processed. With -d, the k-mer counting table will be saved every d files for multifile runs; if -s is set, the specified name will be used, and if not, the name *backup.ct* will be used. -l/-loadtable will load the specified k-mer counting table before processing the specified files. Note that these tables are are in the same format as those produced by **load-into-counting.py** and consumed by **abundance-dist.py**.

-f/-fault-tolerant will force the program to continue upon encountering a formatting error in a sequence file; the k-mer counting table up to that point will be dumped, and processing will continue on the next file.

To append reads to an output file (rather than overwriting it), send output to STDOUT with *-out* - and use UNIX file redirection syntax (>>) to append to the file.

Example:

```
normalize-by-median.py -k 17 tests/test-data/test-abund-read-2.fa
```

Example:

```
normalize-by-median.py -p -k 17 tests/test-data/test-abund-read-paired.fa
```

Example:

```
normalize-by-median.py -p -k 17 -o - tests/test-data/paired.fq >> appended-output.fq
```

Example:

```
normalize-by-median.py -k 17 -f tests/test-data/test-error-reads.fq tests/test-data/test-fastq-reads
```

Example:

```
normalize-by-median.py -k 17 -d 2 -s test.ct tests/test-data/test-abund-read-2.fa tests/test-data/test
```

5.4.4 Read handling: interleaving, splitting, etc.

extract-long-sequences.py

Extract FASTQ or FASTA sequences longer than specified length (default: 200 bp).

usage: extract-long-sequences.py [-h] [-o OUTPUT] [-l LENGTH] input_filenames [input_filenames ...]

input filenames

Input FAST[AQ] sequence filename.

-h, --help

show this help message and exit

-o, --output

The name of the output sequence file.

-l <int>, --length <int>

The minimum length of the sequence file.

extract-paired-reads.py

Take a mixture of reads and split into pairs and orphans.

usage: extract-paired-reads.py [-h] [-version] [-f] infile

infile

-h, --help

show this help message and exit

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

The output is two files, <input file>.pe and <input file>.se, placed in the current directory. The .pe file contains interleaved and properly paired sequences, while the .se file contains orphan sequences.

Many assemblers (e.g. Velvet) require that you give them either perfectly interleaved files, or files containing only single reads. This script takes files that were originally interleaved but where reads may have been orphaned via error filtering, application of abundance filtering, digital normalization in non-paired mode, or partitioning.

Example:

extract-paired-reads.py tests/test-data/paired.fq

fastq-to-fasta.py

Converts FASTQ format (.fq) files to FASTA format (.fa).

usage: fastq-to-fasta.py [-h] [-o filename] [-n] input_sequence

input_sequence

The name of the input FASTQ sequence file.

-h, --help

show this help message and exit

-o <filename>, --output <filename>

The name of the output FASTA sequence file.

-n, --n_keep

Option to drop reads containing 'N's in input_sequence file.

interleave-reads.py

Produce interleaved files from R1/R2 paired files

usage: interleave-reads.py [-h] [-o filename] [-version] [-f] infiles [infiles ...]

infiles

-h, --help

show this help message and exit

```
-o <filename>, --output <filename>
```

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

The output is an interleaved set of reads, with each read in <R1> paired with a read in <R2>. By default, the output goes to stdout unless -o/--output is specified.

As a "bonus", this file ensures that if read names are not already formatted properly, they are reformatted consistently, such that they look like the pre-1.8 Casava format (@name/1, @name/2).

Example:

```
interleave-reads.py tests/test-data/paired.fq.1 tests/test-data/paired.fq.2 -o paired.fq
```

readstats.py

Display summary statistics for one or more FASTA/FASTQ files.

usage: readstats.py [-h] [-o filename] [-csv] filenames [filenames ...]

filenames

```
-h, --help show this help message and exit
```

```
-o <filename>, --output <filename>
   output file for statistics; defaults to stdout.
```

--csv

Use the CSV format for the statistics, including column headers.

Report number of bases, number of sequences, and average sequence length for one or more FASTA/FASTQ files; and report aggregate statistics at end.

With $-\phi$:options:'-output', the output will be saved to the specified file.

Example:

```
readstats.py tests/test-data/test-abund-read-2.fa
```

sample-reads-randomly.py

Uniformly subsample sequences from a collection of files

usage: sample-reads-randomly.py [-h] [-N NUM_READS] [-M MAX_READS] [-S NUM_SAMPLES] [-R RAN-DOM_SEED] [-force_single] [-o output_file] [-version] [-f] filenames [filenames ...]

filenames

```
-h, --help
    show this help message and exit
-N <int>, --num_reads <int>
-M <int>, --max_reads <int>
-S <int>, --samples <int>
```

-R <int>, --random-seed <int>

--force single

Ignore read pair information if present

```
-o <output_file>, --output <output_file>
```

--version

show program's version number and exit

-f, --force

Overwrite output file if it exits

Take a list of files containing sequences, and subsample 100,000 sequences $(-N/--num_reads)$ uniformly, using reservoir sampling. Stop after first 100m sequences $(-M/--max_reads)$. By default take one subsample, but take -S/--samples samples if specified.

The output is placed in -o/--output <file> (for a single sample) or in <file>.subset.0 to <file>.subset.S-1 (for more than one sample).

This script uses the reservoir sampling algorithm.

split-paired-reads.py

Split interleaved reads into two files, left and right.

usage: split-paired-reads.py [-h] [-o output_directory] [-1 output_first] [-2 output_second] [-p] [-version] [-f] infile

infile

-h, --help

show this help message and exit

- -o <output_directory>, --output-dir <output_directory>
 Output split reads to specified directory. Creates directory if necessary
- -1 <output_first>, --output-first <output_first>
 Output "left" reads to this file
- -2 <output_second>, --output-second <output_second>
 Output "right" reads to this file

-p, --force-paired

Require that reads be interleaved

--version

show program's version number and exit

-f, --force

Overwrite output file if it exists

Some programs want paired-end read input in the One True Format, which is interleaved; other programs want input in the Insanely Bad Format, with left- and right- reads separated. This reformats the former to the latter.

The directory into which the left- and right- reads are output may be specified using -o/--output-dir. This directory will be created if it does not already exist.

Alternatively, you can specify the filenames directly with -1/--output-first and -2/--output-second, which will override the -o/--output-dir setting on a file-specific basis.

-p/--force-paired will require the input file to be properly interleaved; by default, this is not required.

Example:

```
split-paired-reads.py tests/test-data/paired.fq
```

Example:

```
split-paired-reads.py -o ~/reads-go-here tests/test-data/paired.fq
```

Example:

```
split-paired-reads.py -1 reads.1 -2 reads.2 tests/test-data/paired.fq
```

5.5 Blog posts and additional documentation

5.5.1 Hashtable and filtering

The basic inexact-matching approach used by the hashtable code is described in this blog post:

http://ivory.idyll.org/blog/jul-10/kmer-filtering

A test data set (soil metagenomics, 88m reads, 10gb) is here:

http://angus.ged.msu.edu.s3.amazonaws.com/88m-reads.fa.gz

5.5.2 Illumina read abundance profiles

khmer can be used to look at systematic variations in k-mer statistics across Illumina reads; see, for example, this blog post:

http://ivory.idyll.org/blog/jul-10/illumina-read-phenomenology

The fasta-to-abundance-hist and abundance-hist-by-position scripts can be used to generate the k-mer abundance profile data, after loading all the k-mer counts into a .ct file:

```
# first, load all the k-mer counts:
load-into-counting.py -k 20 -x 1e7 25k.ct data/25k.fq.gz
# then, build the '.freq' file that contains all of the counts by position
python sandbox/fasta-to-abundance-hist.py 25k.ct data/25k.fq.gz
# sum across positions.
python sandbox/abundance-hist-by-position.py data/25k.fq.gz.freq > out.dist
```

The hashtable method 'dump_kmers_by_abundance' can be used to dump high abundance k-mers, but we don't have a script handy to do that yet.

You can assess high/low abundance k-mer distributions with the hi-lo-abundance-by-position script:

```
load-into-counting.py -k 20 25k.ct data/25k.fq.gz
python sandbox/hi-lo-abundance-by-position.py 25k.ct data/25k.fq.gz
```

This will produce two output files, <filename>.pos.abund=1 and <filename>.pos.abund=255.

5.6 Choosing table sizes for khmer

If you look at the documentation for the scripts (khmer's command-line interface) you'll see two mysterious parameters --N and -x, or, more verbosely, $-n_tables$ and --tablesize. What are these, and how do you specify them?

5.6.1 The really short version

There is no way (except for experience, rules of thumb, and intuition) to know what these parameters should be up front. So, make the product of these two parameters be the size of your available memory:

-N 4 -x 4e9

for a machine with 16 GB of free memory, for example. Also see the rules of thumb, below.

5.6.2 The short version

These parameters specify the maximum memory usage of the primary data structure in khmer, which is basically N big hash tables of size x. The **product** of the number of hash tables and the size of the hash tables specifies the total amount of memory used.

This table is used to track k-mers. If it is too small, khmer will fail in various ways (and should complain), but there is no harm in making it too large. So, **the absolute safest thing to do is to specify as much memory as is available**. Most scripts will inform you of the total memory usage, and (at the end) will complain if it's too small.

For normalize-by-median, khmer uses one byte per hash entry, so: if you had 16 GB of available RAM, you should specify something like $-\mathbb{N}$ 4 $-\mathbb{x}$ 4e9, which multiplies out to about 16 GB.

For the graph partitioning stuff, khmer uses only 1 bit per k-mer, so you can multiple your available memory by 8: for 16 GB of RAM, you could use

-N 4 -x 32e9

which multiplies out to 128 Gbits of RAM, or 16 Gbytes.

Life is a bit more complicated than this, however, because some scripts – load-into-counting and load-graph – keep ancillary information that will consume memory beyond this table data structure. So if you run out of memory, decrease the table size.

Also see the rules of thumb, below.

5.6.3 The real full version

khmer's scripts, at their heart, represents k-mers in a very memory efficient way by taking advantage of two data structures, Bloom filters and Count-Min Sketches, that are both *probabilistic* and *constant memory*. The "probabilistic" part means that there are false positives: the less memory you use, the more likely it is that khmer will think that k-mers are present when they are not, in fact, present.

Digital normalization (normalize-by-median and filter-abund) uses the Count-Min Sketch data structure.

Graph partitioning (load-graph etc.) uses the Bloom filter data structure.

The practical ramifications of this are pretty cool. For example, your digital normalization is guaranteed not to increase in memory utilization, and graph partitioning is estimated to be 10-20x more memory efficient than any other de Bruijn graph representation. And hash tables (which is what Bloom filters and Count-Min Sketches use) are really fast and efficient. Moreover, the optimal memory size for these primary data structures is dependent on the number of k-mers, but not explicitly on the size of k itself, which is very unusual.

In exchange for this memory efficiency, however, you gain a certain type of parameter complexity. Unlike your more typical k-mer package (like the Velvet assembler, or Jellyfish or Meryl or Tallymer), you are either guaranteed not to run out of memory (for digital normalization) or much less likely to do so (for partitioning).

The biggest problem with khmer is that there is a minimum hash number and size that you need to specify for a given number of k-mers, and you cannot confidently predict what it is before actually loading in the data. This, by the way, is also true for de Bruijn graph assemblers and all the other k-mer-based software – the final memory usage depends

on the total number of k-mers, which in turn depends on the true size of your underlying genomic variation (e.g. genome or transcriptome size), the number of errors, and the k-mer size you choose (the k parameter) [see Conway & Bromage, 2011]. **The number of reads or the size of your data set is only somewhat correlated with the total number of k-mers.** Trimming protocols, sequencing depth, and polymorphism rates are all important factors that affect k-mer count.

The bad news is that we don't have good ways to estimate total k-mer count a priori, although we can give you some rules of thumb, below. In fact, counting the total number of distinct k-mers is a somewhat annoying challenge. Frankly, we recommend *just guessing* instead of trying to be all scientific about it.

The good news is that you can never give khmer too much memory! k-mer counting and set membership simply gets more and more accurate as you feed it more memory. (Although there may be performance hits from memory I/O, e.g. see the NUMA architecture.) The other good news is that khmer can measure the false positive rate and detect dangerously low memory conditions. For partitioning, we actually *know* what a too-high false positive rate is – our k-mer percolation paper lays out the math. For digital normalization, we assume that a false positive rate of 10% is bad. In both cases the data-loading scripts will exit with an error-code.

Rules of thumb

Just use -N 4, always, and vary the -x parameter.

For digital normalization, we recommend:

- -x 2e9 for any amount of sequencing for a single microbial genome, MDA-amplified or single colony.
- -x 4e9 for up to a billion mRNAseq reads from any organism. Past that, increase it.
- -x 8e9 for most eukaryotic genome samples.
- -x 8e9 will also handle most "simple" metagenomic samples (HMP on down)
- For metagenomic samples that are more complex, such as soil or marine, start as high as possible. For example, we are using -x 64e9 for ~300 Gbp of soil reads.

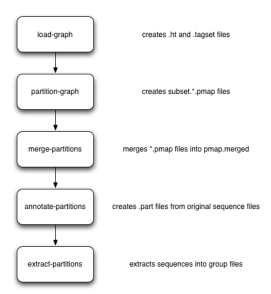
For partitioning of complex metagenome samples, we recommend starting as high as you can – something like half your system memory. So if you have 256 GB of RAM, use -N 4 -x 256e9 which will use 4 x 256 / 8 = 128 GB of RAM for the basic graph storage, leaving other memory for the ancillary data structures.

5.7 Partitioning large data sets (50m+ reads)

"Partitioning" is what khmer calls the process of separating reads that do not connect to each other into different logical bins. The goal of partitioning is to apply divide & conquer to the process of metagenomic assembly.

5.7.1 Basic partitioning

The basic workflow for partitioning is in the figure below:

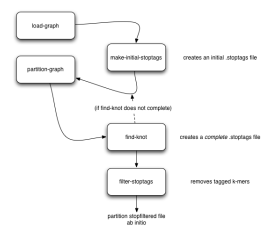


Briefly, you load everything into khmer's probabilistic graph representation; exhaustively explore the graph to find all disconnected sequences; merge the results of the (parallelized) graph exploration; annotate sequences with their partition; and then extract the different partitions into files grouped by partition size. These groups can then be assembled individually.

5.7.2 Artifact removal

As part of our partitioning research, we discovered that large Illumina data sets tend to contain a single large, connected component. This connected component seems to stem from sequencing artifacts that causes knots in the assembly graph. We have developed tools to forcibly remove the knot at the heart of the graph.

Here's the workflow:



5.7.3 Running on an example data set

Here is a set of commands for running both basic partitioning and artifact removal on a small soil metagenomics data set that we've made available for this purpose.

The data set is about 1.1G and you can download it from here:

https://s3.amazonaws.com/public.ged.msu.edu/khmer/iowa-corn-50m.fa.gz

```
cd /path/to/data
# the next command will create a '50m.ct' and a '50m.tagset',
# representing the de Bruijn graph
load-graph.py -k 32 -N 4 -x 16e9 50m iowa-corn-50m.fa.gz
# this will then partition that graph. should take a while.
# update threads to something higher if you have more cores.
# this creates a bunch of files, 50m.subset.*.pmap
partition-graph.py --threads 4 -s 1e5 50m
# now, merge the pmap files into one big pmap file, 50m.pmap.merged
merge-partitions.py 50m
# next, annotate the original sequences with their partition numbers.
# this will create iowa-corn-50m.fa.gz.part
annotate-partitions.py 50m iowa-corn-50m.fa.gz
# now, extract the partitions in groups into 'iowa-corn-50m.groupNNNN.fa'
extract-partitions.py iowa-corn-50m iowa-corn-50m.fa.gz.part
# at this point, you can assemble the group files individually. Note,
# however, that the last one them is quite big? this is because it's
# the lump! yay!
# if you want to break up the lump, go through the partitioning bit
# on the group file, but this time with a twist:
mv iowa-corn-50m.group0005.fa corn-50m.lump.fa
# create graph,
load-graph.py -x 8e9 lump corn-50m.lump.fa
# create an initial set of stoptags to help in knot-traversal; otherwise,
# partitioning and knot-traversal (which is systematic) is really expensive.
make-initial-stoptags.py lump
# now partition the graph, using the stoptags file
partition-graph.py --stoptags lump.stoptags lump
\# use the partitioned subsets to find the k-mers that nucleate the lump
find-knots.py -x 2e8 -N 4 lump
\# remove those k-mers from the fasta files
filter-stoptags.py *.stoptags corn-50m.lump.fa
# now, reload the filtered data set in and partition again.
# NOTE: 'load-graph.py' uses the file extension to determine
# if the file is formatted as FASTA or FASTQ. The default is
# fasta, therefore if your files are fastq formatted you need
# to append 'fastq' to the name so that 'load-graph.py'
# will parse the file correctly
load-graph.py -x 8e9 lumpfilt corn-50m.lump.fa.stopfilt
partition-graph.py -T 4 lumpfilt
merge-partitions.py lumpfilt
annotate-partitions.py lumpfilt corn-50m.lump.fa.stopfilt
extract-partitions.py corn-50m-lump corn-50m.lump.fa.stopfilt.part
# and voila, after all that, you should now have your de-knotted lump in
```

```
# corn-50m-lump.group*.fa. The *.group????.fa files can now be
# assembled individually by your favorite assembler.
```

5.7.4 Post-partitioning assembly

The 'extract-partitions' script takes reads belonging to each partition and aggregates them into 'group' files; each group file contains at least one entire partition (and generally a lot more). Note, you can control the number of reads in each file (equiv, the size of these files) with some of the arguments that 'extract-partitions' takes.

Now that you have these files... what do you do with them? The short answer is: assemble them! Each of these group files contains reads that do not connect to reads in other files, so the files can be assembled individually (which is the whole point of partitioning).

If you're using Velvet, checkout the sandbox/velvet-assemble.sh script, which you can run like this:

```
bash /path/to/khmer/sandbox/velvet-assemble.sh <groupfile> <k>
```

This script does three things:

- first, it breaks the reads up into paired reads and single reads, and puts them in separate files (.pe and .se);
- second, it strips off the partition information from the reads, which confuses Velvet;
- and third, it runs velveth and velvetg to actually assemble.

You can implement your own approach, of course, but this is an example of what we do ourselves.

5.8 Known Issues

Some users have reported that normalize-by-median.py will utilize more memory than it was configured for. This is being investigated in https://github.com/ged-lab/khmer/issues/266

If your k-mer table is truncated on write, an error may not be reported; this is being tracked in https://github.com/ged-lab/khmer/issues/443. However, khmer will now (correctly) fail when trying to read a truncated file (See #333).

Paired-end reads from Casava 1.8 currently require renaming for use in normalize-by-median and abund-filter when used in paired mode. The integration of a fix for this is being tracked in https://github.com/ged-lab/khmer/issues/23

Some scripts only output FASTA even if given a FASTQ file. This issue is being tracked in https://github.com/ged-lab/khmer/issues/46

5.9 Deploying the khmer project tools on Galaxy

This document is for people interested in deploying the khmer tools on the Galaxy platform.

We are developing the support for running all the khmer scripts in Galaxy.

5.9.1 Install the tools & tool description

In the administrative interface select "Search and browse tool sheds" under the heading "Tool sheds". Click on "Galaxy test tool shed" and search for khmer. Click on the "khmer" button and choose "Preview and install". Click the "Install to Galaxy" button at the top. At the bottom of the next page click the "Install" button.

5.8. Known Issues 49

5.9.2 Single Output Usage

For one or more files into a single file:

- #. Choose 'Normalize By Median' from the 'khmer protocols' section of the 'Tools' menu.
- #. Compatible files already uploaded to your Galaxy instance should be listed. If not then you may need to set their datatype manually.
- #. After selecting the input files specify if they are paired-interleaved or not.
- #. Specify the sample type or show the advanced parameters to set the tablesize yourself. Consult Choosing table sizes for khmer for assistance.

5.10 An incomplete bibliography of papers using khmer

5.10.1 Digital normalization

Multiple Single-Cell Genomes Provide Insight into Functions of Uncultured Deltaproteobacteria in the Human Oral Cavity. Campbell et al., PLoS One, 2013, doi:10.1371/journal.pone.0059361. [paper link]

Insights into archaeal evolution and symbiosis from the genomes of a nanoarchaeon and its inferred crenarchaeal host from Obsidian Pool, Yellowstone National Park. Podar et al., Biology Direct, 2013 doi:10.1186/1745-6150-8-9. [paper link]

5.11 How to get help

First, be sure that you:

- 1. Read the documentation (this site)
- 2. Google search for the error output and/or keywords related to your problem. Here you can search results from the mailing list, where others may have discussed solutions to the same issue.

5.11.1 Mailing List

The primary way to get help is through the khmer discussion list: http://lists.idyll.org/listinfo/khmer

5.11.2 Asking a question

- 1. Include your:
 - OS version (Mac OS X or Linux): uname -mrs
 - Python version: python --version
 - and khmer version: pip freeze | grep khmer
- 2. Precisely describe what you are trying to do. Reread it from the perspective of someone else trying to reproduce your task.
- 3. Copy-and-paste the exact command that is causing the problem. Include the steps you performed leading up to the issue.
- 4. Include the complete error message; if it is large, include a link to a file.

5.11.3 GitHub

You are also welcome to report an issue you are having using GitHub:: https://github.com/ged-lab/khmer/issues/new

The khmer developer documentation

This section of the documentation is for people who are contributing (or would like to contribute to) the khmer project codebase, either by contributing code or by helping improve the documentation.

Please note that this project is released with a Contributor Code of Conduct. By participating in the development of this project you agree to abide by its terms.

Contents:

6.1 Contributor Code of Conduct

As contributors and maintainers of this project, we pledge to respect all people who contribute through reporting issues, posting feature requests, updating documentation, submitting pull requests or patches, and other activities.

We are committed to making participation in this project a harassment-free experience for everyone, regardless of level of experience, gender, gender identity and expression, sexual orientation, disability, personal appearance, body size, race, age, or religion.

Examples of unacceptable behavior by participants include the use of sexual language or imagery, derogatory comments or personal attacks, trolling, public or private harassment, insults, or other unprofessional conduct.

Project maintainers have the right and responsibility to remove, edit, or reject comments, commits, code, wiki edits, issues, and other contributions that are not aligned to this Code of Conduct. Project maintainers or contributors who do not follow the Code of Conduct may be removed from the project team.

Instances of abusive, harassing, or otherwise unacceptable behavior may be reported by emailing khmer-project@idyll.org which only goes to C. Titus Brown and Michael R. Crusoe. To report an issue involving either of them please email Judi Brown Clarke, Ph.D. the Diversity Director at the BEACON Center for the Study of Evolution in Action, an NSF Center for Science and Technology.

This Code of Conduct is adapted from the Contributor Covenant, version 1.0.0, available at http://contributor-covenant.org/version/1/0/0/

6.2 Getting started with khmer development

This document is for people who would like to contribute to khmer. It walks first-time contributors through making their own copy of khmer, building it, and submitting changes for review and merge into the master copy of khmer.

Start by making your own copy of khmer and setting yourself up for development; then, build khmer and run the tests; and finally, claim an issue and start developing!

If you're unfamiliar with git and branching in particular, check out the git-scm book.

We've provided a quick guide to the khmer code base here: A quick guide to the khmer codebase.

6.2.1 One-time Preparation

1. Install the dependencies.

OS X users

- (a) Install Xcode from the Mac App Store (requires root).
- (b) Register as an Apple Developer.
- (c) Install the Xcode command-line tools: Xcode -> preferences -> Downloads -> Command Line Tools (requires root).

Linux users

(a) Install the python development environment, virtualenv, pip, gcc, and g++.

On recent Debian and Ubuntu this can be done with:

```
sudo apt-get install python2.7-dev python-virtualenv python-pip gcc \
g++ git astyle gcovr cppcheck
```

For RHEL6:

```
sudo yum install -y python-devel python-pip git gcc gcc-c++ make sudo pip install virtualenv
```

For Arch Linux:

```
sudo pacman -S python2 python2-pip python2-virtualenv gcc make
```

2. Get a GitHub account.

(We use GitHub to manage khmer contributions.)

3. Fork github.com/ged-lab/khmer.

Visit that page, and then click on the 'fork' button (upper right).

(This makes a copy of the khmer source code in your own GitHub account.)

4. Clone your copy of khmer to your local development environment.

Your clone URL should look something like this:

```
https://github.com/empty-titus/khmer.git
```

and the UNIX shell command should be:

```
git clone https://github.com/empty-titus/khmer.git
```

(This makes a local copy of khmer on your development machine.)

5. Add a git reference to the khmer ged-lab repository:

```
cd khmer
git remote add ged https://github.com/ged-lab/khmer.git
cd ../
```

(This makes it easy for you to pull down the latest changes in the main repository.)

6. Create a virtual Python environment within which to work with virtualenv:

```
python2.7 -m virtualenv env
```

This gives you a place to install packages necessary for running khmer.

OS X users and others may need to download virtualenv first:

```
curl -O https://pypi.python.org/packages/source/v/virtualenv/virtualenv-1.11.6.tar.gz
tar xzf virtualenv*
cd virtualenv-*; python2.7 virtualenv.py ../env; cd ..
```

Mac ports users on the OS X platform can install pip by execution from the command line:

```
sudo port install py27-pip
```

Homebrew users on the OS X platform will have pip already installed

Conda users on any platform should instead create a separate Conda environment:

```
conda create -n khmer anaconda
```

7. Activate the virtualenv and install a few packages:

```
source env/bin/activate
cd khmer
make install-dependencies
```

(This installs Sphinx and nose, packages we use for building the documentation and running the tests.)

In Conda to activate the previously created environment and install dependencies:

```
source activate khmer
cd khmer
make install-dependencies
```

8. Cppcheck installation:

Debian and Ubuntu Linux distro users can install eppcheck by executing from the command line:

```
sudo apt-get install cppcheck
```

Mac ports users on the OS X platform can install cppcheck by executing from the command line:

```
sudo port install cppcheck
```

Homebrew users on the OS X platform can install cppcheck by executing from the command line:

```
sudo brew install cppcheck
```

6.2.2 Building khmer and running the tests

1. Activate (or re-activate) the virtualenv:

```
source ../env/bin/activate
```

... or for Conda users:

```
source activate khmer
```

You can run this many times without any ill effects.

(This puts you in the development environment.)

2. Build khmer:

make

If this fails, we apologize – please go create a new issue, paste in the failure message, and we'll try to help you work through it!

(This takes the C++ source code and compiles it into something that Python can run.)

3. Run the tests:

```
make test
```

You should see lots of output, with something like:

```
Ran 360 tests in 10.403s
OK
```

at the end.

(This will run all of the Python tests in the tests/ directory.)

Congratulations! You're ready to develop!

6.2.3 Claiming an issue and starting to develop

1. Find an open issue and claim it.

Go to the list of open khmer issues and find one you like; we suggest starting with the low-hanging fruit issues).

Once you've found an issue you like, make sure that no one has been assigned to it (see "assignee", bottom right near "notifications"). Then, add a comment "I am working on this issue." You've staked your claim!

(We're trying to avoid having multiple people working on the same issue.)

2. In your local copy of the source code, update your master branch from the main khmer master branch:

```
git checkout master
git pull ged master
```

(This pulls in all of the latest changes from whatever we've been doing on ged-lab.)

3. Create a new branch and link it to your fork on GitHub:

```
git checkout -b fix/issue_number
git push -u origin fix/issue_number
```

where you replace "issue_number" with the number of the issue.

(This is the set of changes you're going to ask to be merged into khmer.)

4. Make some changes and commit them.

This will be issue dependent;).

(You should visit and read Coding guidelines and code review checklist.)

5. Periodically update your branch from the main khmer master branch:

```
git pull ged master
```

(This pulls in all of the latest changes from whatever we've been doing on ged-lab - important especially during periods of fast change or for long-running pull requests.

6. Run the tests and/or build the docs *before* pushing to GitHub:

make doc test pep8

Make sure they all pass!

7. Push your branch to your own GitHub fork:

git push origin

(This pushes all of your changes to your own fork.)

- 8. Repeat until you're ready to merge your changes into "official" khmer.
- 9. Set up a Pull Request asking to merge things into the central khmer repository.

In a Web browser, go to your GitHub fork of khmer, e.g.:

https://github.com/empty-titus/khmer

and you will see a list of "recently pushed branches" just above the source code listing. On the right side of that should be a "Compare & pull request" green button. Click on it!

Now:

- add a descriptive title ("updated tests for XXX")
- put the issue number in the comment ("fixes issue #532")

then click "Create pull request."

(This creates a new issue where we can all discuss your proposed changes; the khmer team will be automatically notified and you will receive e-mail notifications as we add comments. See GitHub flow for more info.)

- 10. Paste in the committer checklist from Coding guidelines and code review checklist and, after its pasted in, check off as many of the boxes as you can.
- 11. As you add new commits to address bugs or formatting issues, you can keep pushing your changes to the pull request by doing:

git push origin

- 12. When you are ready to have the pull request reviewed, please mention @luizirber, @camillescott, @mr-c, or @ctb with a comment 'Ready for review!'
- 13. The khmer team will now review your pull request and communicate with you through the pull request page. Please feel free to add 'ping!' and an @ in the comments if you are looking for feedback this will alert us that you are still on the line but we will automatically get notified of your pull request and any new comments, so use sparingly.

If this is still your first issue, please don't take another issue until we've merged your first one - thanks!

14. If we request changes, return to the step "Make some changes and commit them" and go from there. Any additional commits you make and push to your branch will automatically be added to the pull request (which is pretty dang cool.)

6.2.4 After your first issue is successfully merged...

You're now an experienced GitHub user! Go ahead and take some more tasks; you can broaden out beyond the low hanging fruit if you like.

Here are a few suggestions:

- If you're knowledgeable in C++ and/or Python and/or documentation and/or biology, we'd love to attract further contributions to khmer. Please visit the issues list and browse about and find something interesting looking.
- One general thing we'd like to do is increase our test coverage. You can go find test coverage information on
 our continuous integration server by clicking down to individual files; or, ask us on khmer-project@idyll.org for
 suggestions.
- Ask us! Ask khmer-project@idyll.org for suggestions on what to do next. We can suggest particularly ripe low-hanging fruit, or find some other issues that suit your interests and background.
- You can also help other people out by watching for new issues or looking at pull requests. Remember to be nice and polite!

6.3 A quick guide to testing (for khmer)

This document is for contributors new to automated testing, and explains some of the motivation and logic behind the khmer project's testing approach.

One of our most important "secret sauces" for khmer development is that we do a fair bit of testing to make sure our code works and keeps working!

- We maintain fairly complete test coverage of our code. What this means is that we have automated tests that, when run, execute most of the lines of Python and C++ code in our lib/, khmer/ and scripts/ directories. This doesn't *guarantee* things are correct, but it does mean that at least most of the code works at some basic level.
- we have other tests that we run periodically (for example, before each release) see Releasing a new version of khmer for details. These tests check that our code works on multiple systems and with other people's software.

CTB and others have written a great deal about testing, and testing in Python in particular. Here's an introductory guide CTB wrote a long time ago. You might also be interested in reading this description of the different kinds of tests.

For the more general motivation, see the Lack of Testing Death Spiral.

But... how do you do testing??

First, let's talk about specific goals for testing. What should you be aiming for tests to do? You can always add more testing code, but that might not be useful if they are redundant or over-complicated.

An overall rule is to "keep it simple" - keep things as simple as possible, testing as few things as possible in each test.

We suggest the following approach to writing tests for **new code**:

- 1. Write a test that just *runs* the new code, generally by copying existing test code to a new test and changing it. Don't do anything clever for the first test just run something straightforward, and try to use existing data.
- 2. Decide which use cases should be tested. This is necessarily code specific but our main advice is "don't be clever" write some tests to make sure that the code basically works.
- 3. Add in tests for edge cases. By this we mean look for special cases in your code if statements, fence-post bound errors, etc. and write tests that exercise those bits of code specifically.
- 4. Make sure tests that expect a function call to fail (esp. with fail_ok=True) are failing for the expected reason. Run the code from the command line and see what the behavior is. For troubleshooting tests, catch the error with try: ... except: or print err.

For adding tests to **old code**, we recommend a mix of two approaches:

- 1. use "stupidity driven testing" and write tests that recapitulate bugs before we fix those bugs.
- 2. look at test coverage (see khmer's cobertura test coverage, here) and identify lines of C++ or Python code that are not being executed by the current tests. Then write new tests targeting the new code.

Next, to add a test, you have two options: either write a new one from scratch, or copy an existing one. (We recommend the latter.)

To write a new one, you'll need to know how to write tests. For getting an idea of the syntax, read this introductory guide and the official documentation. Then find the right file in tests/*.py and add your test!

A better approach is, frankly, to go into the existing test code, find a test that does something similar to what you want to do, copy it, rename it, and then modify it to do the new test.

Finally, where do you add new tests and how do you run just your test?

Put new tests somewhere in tests/*.py. If you have trouble figuring out what file to add them to, just put them in *some* file and we'll help you figure out where to move them when we do code review.

To run one specific test rather than all of them, you can do:

```
./setup.py nosetests --tests tests/test_scripts.py:test_load_into_counting
```

Here, you're running just one test - the test function named test_load_into_counting in the file test_scripts.py.

6.4 A quick guide to the khmer codebase

This document describes the khmer project's directory layout.

The ChangeLog file lists changes to the codebase, most recent first.

The lib/ directory contains all of the C++ code.

The khmer/ directory contains the khmer package (khmer/_init__.py etc) and the C++-to-Python bridge (khmer/ khmermodule.cc).

The scripts/ and sandbox/ directory contain Python command-line scripts.

The tests/ directory contains all of the tests. Each test is a function in one of the tests/test*.py files.

6.5 Coding guidelines and code review checklist

This document is for anyone who want to contribute code to the khmer project, and describes our coding standards and code review checklist.

6.5.1 Coding standards

All plain-text files should have line widths of 80 characters or less unless that is not supported for the particular file format.

For C++, we use Todd Hoff's coding standard, and astyle -A10 / "One True Brace Style" indentation and bracing. Note: @CTB needs Emacs settings that work for this.

Vim users may want to set the ARTISTIC_STYLE_OPTIONS shell variable to "-A10 -max-code-length=80" and run `: %!astyle` to reformat. The four space indentation can be set with:

```
set expandtab
set shiftwidth=4
set softtabstop=4
```

For Python, PEP 8 is our standard. The 'pep8' and 'autopep8' Makefile targets are helpful.

Code, scripts, and documentation must have its spelling checked. Vim users can run:

```
:setlocal spell spelllang=en_us
```

Use Js and Is to navigate between misspellings and z= to suggest a correctly spelled word. zg will add a word as a good word.

GNU aspell can also be used to check the spelling in a single file:

```
aspell check --mode ccpp $filename
```

6.5.2 Code Review

Please read 11 Best Practices for Peer Code Review.

See also Code reviews: the lab meeting for code and the PyCogent coding guidelines.

6.5.3 Checklist

Copy and paste the following into a pull request comment when it is ready for review:

```
- [] Is it mergeable?
- [] Did it pass the tests?
- [] If it introduces new functionality in scripts/ is it tested?
Check for code coverage with `make clean diff-cover`
- [] Is it well formatted? Look at `make pep8`, `make diff_pylint_report`,
   `make cppcheck`, and `make doc` output. Use `make format` and manual
   fixing as needed.
- [] Did it change the command-line interface? Only additions are allowed
   without a major version increment. Changing file formats also requires a
   major version number increment.
- [] Is it documented in the ChangeLog?
   http://en.wikipedia.org/wiki/Changelog#Format
- [] Was a spellchecker run on the source code and documentation after
   changes were made?
- [] Is the Copyright year up to date?
```

Note that after you submit the comment you can check and uncheck the individual boxes on the formatted comment; no need to put x or y in the middle.

6.5.4 CPython Checklist

Here's a checklist for new CPython types with future-proofing for Python 3:

```
- [ ] the CPython object name is of the form `khmer_${OBJECTNAME}_Object`
- [ ] Named struct with `PyObject_HEAD` macro
- [ ] `static PyTypeObject khmer_${OBJECTNAME}_Type` with the following
 entries
  - [ ] `PyVarObject_HEAD_INIT(NULL, 0)` as the object init (this includes
    the `ob_size` field).
  - [ ] all fields should have their name in a comment for readability
  - [ ] The `tp_name` filed is a dotted name with both the module name and
    the name of the type within the module. Example: `khmer.ReadAligner`
  - [ ] Deallocator defined and cast to `(destructor)` in tp_dealloc
    - [ ] The object's deallocator must be
       `Py_TYPE(obj)->tp_free((PyObject*)obj);`
  - [ ] Do _not_ define a `tp_getattr`
  - [ ] BONUS: write methods to present the state of the object via
     `tp_str` & `tp_repr`
  - [ ] _Do_ pass in the array of methods in `tp_methods`
  - [ ] _Do_ define a new method in `tp_new`
- [ ] PyMethodDef arrays contain doc strings
  - [ ] Methods are cast to `PyCFunctions`s
- [ ] Type methods use their type Object in the method signature.
- [ ] Type creation method decrements the reference to self
 (`Py_DECREF(self); `) before each error-path exit (`return NULL; `)
- [ ] No factory methods. Example: `khmer_new_readaligner`
- [ ] Type object is passed to `PyType_Ready` and its return code is checked
 in `init_khmer()`
 [ ] The reference count for the type object is incremented before adding
 it to the module: `Py_INCREF(&khmer_${OBJECTNAME}_Type);`.
```

6.6 Command line scripts, scripts/, and sandbox/

Note: This document applies through khmer/oxli 2.0/3.0 (see Roadmap to v2.0, v3.0, v4.0) - we will revisit when the Python API falls under semantic versioning for oxli 4.0.

khmer has two conflicting goals: first, we want to provide a reliable piece of software to our users; and second, we want to be flexible and enable exploration of new algorithms and programs. To this end, we've split our command line scripts across two directories, scripts/ and sandbox/. The former is the staid, boring, reliable code; the latter is a place for exploration.

As a result, we are committed to high test coverage, stringent code review, and Semantic Versioning for files in scripts/, but explicitly *not* committed to this for files and functionality implemented in sandbox/. So, putting a file into scripts/ is a big deal, especially since it increases our maintenance burden for the indefinite future.

We've roughed out the following process for moving scripts into scripts/:

- Command line scripts start in sandbox/;
- Once their utility is proven (in a paper, for example), we can propose to move them into scripts/;
- There's a procedure for moving scripts from sandbox/ into scripts/.

Read on!

6.6.1 Sandbox script requirements and suggestions

All scripts in sandbox/ must:

- be importable (enforced by test_import_all in test_sandbox_scripts.py)
- be mentioned in sandbox/README.rst
- have a hash-bang line (#! /usr/bin/env python2) at the top
- be command-line executable (chmod a+x)
- have a Copyright message (see below)
- have lowercase names
- use '-' as a word separator, rather than '_' or CamelCase

All new scripts being added to sandbox/ should:

- have decent automated tests
- be used in a protocol (see khmer-protocols) or a recipe (see khmer-recipes)
- be pep8 clean and pylint clean-ish (see make pep8 and make diff pylint).

6.6.2 Command line standard options for scripts/

All scripts in scripts/ should have the following options, if they could apply:

- --version should always apply
- --help should always apply
- --force override any sanity checks that may prevent the script from running
- --loadtable and --savetable where appropriate (see khmer args.py)

6.6.3 Copyright message

Our current Copyright message is:

```
#
# This file is part of khmer, http://github.com/ged-lab/khmer/, and is
# Copyright (C) Michigan State University, 2009-2015. It is licensed under
# the three-clause BSD license; see doc/LICENSE.txt.
# Contact: khmer-project@idyll.org
#
```

The beginning year should be the first year that this file existed in the repo; the end year should be the last year a coding change was made in the file.

Upgrading a script from 'sandbox' to 'scripts'

First, everything needed (all library support code) should be already committed to khmer master after the usual review process; the relevant script(s) should be in sandbox/.

Second, an issue should be started explicitly to discuss whether the script(s) should be moved from sandbox/ into scripts/. This issue should discuss the general need for this script, outside of a particular paper pipeline. (Note that there is no imperative to move a script out of sandbox/; if we think it's useful code to have around and want to keep it functioning, we should just add in automated tests and otherwise level it up.)

Third, assuming we reach general agreement about moving the script(s) into scripts/, start a pull request to do so, referencing the issue and containing the following checklist. The PR should start by moving the script from sandbox/into scripts/, and moving the tests out of the test_sandbox_scripts.py file.

Last but not least, intensive code review may raise more general issues that could apply to the entire code base; if contentious or needing discussion, these issues may be punted to general issues so as to not block a merge.

6.6.4 A checklist for moving a script into the scripts/ directory from sandbox/

Copy or paste this checklist into the PR, in addition to the normal development/PR checklist:

```
- [] most or all lines of code are covered by automated tests (see output of ``make diff-cover``)
- [] ``make diff_pylint`` is clean
- [] the script has been updated with a ``get_parser()`` and added to doc/user/scripts.txt
- [] argparse help text exists, with an epilog docstring, with examples and options
- [] standard command line options are implemented
- [] version and citation information is output to STDERR (`khmer_args.info(...)`)
- [] support '-' (STDIN) as an input file, if appropriate
- [] support designation of an output file (including STDOUT), if appropriate
- [] runtime diagnostic information (progress, etc.) is output to STDERR
- [] script has been removed from sandbox/README.rst
```

6.7 A guide for khmer committers

This document is for people with commit rights to github.com/ged-lab/khmer.

If you have commit privileges to the ged-lab/khmer repository, here are a few useful tips.

First, never merge something unless it's been through a review! This rule can be broken under specific conditions when doing a release; see Releasing a new version of khmer.

Second, need to force another continuous integration run? Put "test this please" in a comment. This can be used to ask our continuous integration system to run on someone else's pull request – by default, it only runs on commits from people who have write privileges to khmer, so you may need to do this if you're reviewing someone else's pull request.

Third, we ask that all contributors set up standing Pull Requests while they are working something. (This is a **requirement** if you're in the GED lab.) This lets us track what's going on. On the flip side, please do not review pull requests until they are indicated as "ready for review".

6.8 Releasing a new version of khmer

This document is for khmer release managers, and details the process for making a new release of the khmer project.

6.8.1 How to make a khmer release candidate

Michael R. Crusoe, Luiz Irber, and C. Titus Brown have all been release makers, following this checklist by MRC.

1. The below should be done in a clean checkout:

```
cd `mktemp -d`
git clone git@github.com:ged-lab/khmer.git
cd khmer
```

2. (Optional) Check for updates to versioneer:

```
pip install --upgrade versioneer
versioneer-installer

git diff

./setup.py versioneer
git diff
git commit -m -a "new version of versioneer.py"
# or
git checkout -- versioneer.py khmer/_version.py khmer/_init__.py MANIFEST.in
```

3. Review the git logs since the last release and diffs (if needed) and ensure that the ChangeLog is up to date:

```
git log --minimal --patch `git describe --tags --always --abbrev=0`..HEAD
```

- 4. Review the issue list for any new bugs that will not be fixed in this release. Add them to doc/known-issues.txt
- 5. Verify that the build is clean: http://ci.ged.msu.edu/job/khmer-master/
- 6. Submit a build to Coverity Scan if it hasn't been done recently. You can get the token from https://gitlab.msu.edu/ged-lab/ged-internal-docs/wikis/coverity-scan or https://scan.coverity.com/projects/621?tab=project_settings

```
virtualenv coverityenv
source coverityenv/bin/activate
make install-dependencies
make clean
cov_analysis_dir=~/src/coverity/cov-analysis-linux64-7.5.0/ make coverity-build
COVERITY_TOKEN=${COVERITY_TOKEN} make coverity-upload
```

7. Set your new version number and release candidate:

```
new_version=1.3
rc=rc1
```

and then tag the release candidate with the new version number prefixed by the letter 'v':

```
git tag v${new_version}-${rc}
git push --tags git@github.com:ged-lab/khmer.git
```

8. Test the release candidate. Bonus: repeat on Mac OS X:

```
mkdir ../not-khmer # if there is a subdir named 'khmer' nosetest will execute tests
# there instead of the installed khmer module's tests
pushd ../not-khmer; nosetests khmer --attr '!known_failing'; popd
# Secondly we test via pip
cd ../testenv2
source bin/activate
pip install -U setuptools==3.4.1
pip install -e git+https://github.com/ged-lab/khmer.git@v${new_version}-${rc}#egg=khmer
make install-dependencies
make dist
make test
cp dist/khmer*tar.gz ../../testenv3/
pip uninstall -y khmer; pip uninstall -y khmer; make install
cd ../.. # no subdir named khmer here, safe for nosetesting installed khmer module
normalize-by-median.py --version 2>&1 | grep ${new_version}-${rc} && \
        echo 2nd manual version check passed
nosetests khmer --attr '!known_failing'
# Is the distribution in testenv2 complete enough to build another
# functional distribution?
cd ../testenv3/
source bin/activate
pip install -U setuptools==3.4.1
pip install khmer*tar.gz
pip install nose
tar xzf khmer*tar.gz
cd khmer*
make dist
make test
pip uninstall -y khmer; pip uninstall -y khmer; make install
mkdir ../not-khmer
pushd ../not-khmer ; nosetests khmer --attr '!known_failing' ; popd
```

9. Publish the new release on the testing PyPI server. You will need to change your PyPI credentials as documented here: https://wiki.python.org/moin/TestPyPI. You may need to re-register:

```
python setup.py register --repository test
```

Now, upload the new release:

```
python setup.py sdist upload -r test
```

Test the PyPI release in a new virtualenv:

- 10. Do any final testing (BaTLab and/or acceptance tests).
- 11. Make sure any release notes are merged into doc/release-notes/.

6.8.2 How to make a final release

When you've got a thoroughly tested release candidate, cut a release like so:

1. Create the final tag and publish the new release on PyPI (requires an authorized account).:

```
cd ../../khmer
git tag v${new_version}
python setup.py register sdist upload
```

2. Delete the release candidate tag and push the tag updates to GitHub.:

```
git tag -d v${new_version}-${rc}
git push git@github.com:ged-lab/khmer.git
git push --tags git@github.com:ged-lab/khmer.git
```

- 3. Add the release on GitHub, using the tag you just pushed. Name it 'version X.Y.Z', and copy and paste in the release notes.
- 4. Make a binary wheel on OS X.:

```
virtualenv build
cd build
source bin/activate
pip install -U setuptools==3.4.1 wheel
pip install --no-clean khmer==${new_version}
cd build/khmer
./setup.py bdist_wheel upload
```

- 5. Update Read the Docs to point to the new version. Visit https://readthedocs.org/builds/khmer/ and 'Build Version: master' to pick up the new tag. Once that build has finished check the "Activate" box next to the new version at https://readthedocs.org/dashboard/khmer/versions/ under "Choose Active Versions". Finally change the default version at https://readthedocs.org/dashboard/khmer/advanced/ to the new version.
- 6. Delete any RC tags created:

```
git tag -d ${new_version}-${rc}
git push origin :refs/tags/${new_version}-${rc}
```

- 7. Tweet about the new release.
- 8. Send email including the release notes to khmer@lists.idyll.org and khmer-announce@lists.idyll.org

6.8.3 BaTLab testing

The UW-Madison Build and Test Lab provides the khmer project with a free cross-platform testing environment.

1. Connect to their head node:

```
ssh mcrusoe@submit-1.batlab.org
```

2. Move into the khmer directory and download a release from PyPI's main server or the test PyPI server:

```
cd khmer/
wget https://testpypi.python.org/packages/source/k/khmer/khmer-1.0.1-rc3.tar.gz
vim khmer-v1.0.inputs # change the 'scp_file' to point to the release
vim khmer-v1.0.run-spec # change 'project_version' at bottom
nmi_submit khmer-v1.0.run-spec
```

6.8.4 Setuptools Bootstrap

ez_setup.py is from https://bitbucket.org/pypa/setuptools/raw/bootstrap/

Before major releases it should be examined to see if there are new versions available and if the change would be useful

6.8.5 Versioning Explanation

Versioneer, from https://github.com/warner/python-versioneer, is used to determine the version number and is called by Setuptools and Sphinx. See the files versioneer.py, the top of khmer/__init__.py, khmer/_version.py, setup.py, and doc/conf.py for the implementation.

The version number is determined through several methods: see https://github.com/warner/python-versioneer#version-identifiers

```
the
                                                                                       derived
         source
                 tree
                       is
                           from
                                  a
                                      git
                                           checkout
                                                     then
                                                           the
                                                                 version
                                                                         number
                                                                                  is
                                                                                       format
     git describe --tags --dirty --always.
                                                          This
                                                                 will
                                                                       he
                                                                            in
                                                                                 the
${tagVersion}-${commits_ahead}-${revision_id}-${isDirty}.
                                                                                     Example:
v0.6.1-18-g8a9e430-dirty
```

If from an unpacked tarball then the name of the directory is queried.

Lacking either of the two git-archive will record the version number at the top of khmer/_version.py via the \$Format:%H\$ placeholders enabled by the "export-subst" entry in .gitattributes.

Non source distributions will have a customized khmer/_version.py that contains hard-coded version strings. (see build/*/khmer/_version.py after a python setup.py build for an example)

ez_setup.py bootstraps Setuptools (if needed) by downloading and installing an appropriate version

6.9 Miscellaneous implementation details

Partition IDs are "stored" in FASTA files as an integer in the last tab-separated field. Yeah, dumb, huh?

6.10 Development miscellany

6.10.1 Third-party use

We ask that third parties who build upon the codebase to do so from a versioned release. This will help them determine when bug fixes apply and generally make it easier to collaborate. If more intensive modifications happen then we request that the repository is forked, again preferably from a version tag.

6.10.2 Build framework

'make' should build everything, including tests and "development" code.

6.10.3 git and GitHub strategies

Still in the works, but read this.

Make a branch on ged-lab (preferred so others can contribute) or fork the repository and make a branch there.

Each piece or fix you are working on should have its own branch; make a pull-request to ged-lab/master to aid in code review, testing, and feedback.

If you want your code integrated then it needs to be mergable

Example pull request update using the command line:

- 1. Clone the source of the pull request (if needed) git clone git@github.com:mr-c/khmer.git
- 2. Checkout the source branch of the pull request git checkout my-pull-request
- 3. Pull in the destination of the pull request and resolve any conflicts git pull git@github.com:ged-lab/khmer.git master
- 4. Push your update to the source of the pull request git push
- 5. Jenkins will automatically attempt to build and test your pull requests.

6.10.4 Code coverage

Jenkins calculates code coverage for every build. Navigate to the results from the master node first to view the coverage information.

Code coverage should never go down and new functionality needs to be tested.

6.10.5 Pipelines

All khmer scripts used by a published recommended analysis pipeline must be included in scripts/ and meet the standards therein implied.

6.10.6 Command line scripts

Python command-line scripts should use '-' instead of '_' in the name. (Only filenames containing code for import imported should use _.)

Please follow the command-line conventions used under scripts/. This includes most especially standardization of '-x' to be hash table size, '-N' to be number of hash tables, and '-k' to always refer to the k-mer size.

Command line thoughts:

If a filename is required, typically UNIX commands don't use a flag to specify it.

Also, positional arguments typically aren't used with multiple files.

CTB's overall philosophy is that new files, with new names, should be created as the result of filtering etc.; this allows easy chaining of commands. We're thinking about how best to allow override of this, e.g.

```
filter-abund.py <ct file> <filename> [ -o <filename.keep> ]
```

All code in scripts/ must have automated tests; see tests/test_scripts.py. Otherwise it belongs in sandbox/.

When files are overwritten, they should only be opened to be overwritten after the input files have been shown to exist. That prevents stupid command like mistakes from trashing important files.

It would be nice to allow piping from one command to another where possible. But this seems complicated.

CTB: should we squash output files (overwrite them if they exist), or not? So far, leaning towards 'not', as that way no one is surprised and loses their data.

A general error should be signaled by exit code l and success by l. Linux supports exit codes from l to l where the value l means a general error. An exit code of l will get converted to l

CLI reading:

http://stackoverflow.com/questions/1183876/what-are-the-best-practices-for-implementing-a-cli-tool-in-perl

http://catb.org/esr/writings/taoup/html/ch11s06.html

http://figshare.com/articles/tutorial_pdf/643388

6.10.7 Python / C integration

The Python extension that wraps the C++ core of khmer lives in khmer/_khmermodule.CC

This wrapper code is tedious and annoying so we use a static analysis tool to check for correctness.

https://gcc-python-plugin.readthedocs.org/en/latest/cpychecker.html

Developers using Ubuntu Precise will want to install the gcc-4.6-plugin-dev package

Example usage:

```
CC="/home/mcrusoe/src/gcc-plugin-python/gcc-python-plugin/gcc-with-cpychecker --maxtrans=512" python setup.py build_ext 2>&1 | less
```

False positives abound: ignore errors about the C++ standard library. This tool is primarily useful for reference count checking, error-handling checking, and format string checking.

Errors to ignore: "Unhandled Python exception raised calling 'execute' method", "AttributeError: 'NoneType' object has no attribute 'file'"

Warnings to address:

```
khmer/_khmermodule.cc:3109:1: note: this function is too complicated for the reference-count checker to fully analyze: not all paths were analyzed
```

Adjust –maxtrans and re-run.

```
khmer/_khmermodule.cc:2191:61: warning: Mismatching type in call to
Py_BuildValue with format code "i" [enabled by default]
argument 2 ("D.68937") had type
"long long unsigned int"
but was expecting
"int"
for format code "i"
```

See below for a format string cheat sheet One also benefits by matching C type with the function signature used later. "I" for unsigned int "K" for unsigned long long a.k.a khmer::HashIntoType.

6.11 Crazy ideas

1. A JavaScript preprocessor to do things like count k-mers (HLL), and do diginorm on data as uploaded to server. Inspired by a paper that Titus reviewed for PLoS One; not yet published.

Roadmap to v2.0, v3.0, v4.0

7.1 Background

To make the khmer project easier to use and easier to build upon several fundamental changes need to happen. This document outlines our plan to do so while minimizing the impact of these changes on our existing users.

The discussion that lead to this document can be read at https://github.com/ged-lab/khmer/issues/389

7.2 Remainder of v1.x series

Start of transition to a single entrypoint named *oxli*. This will be exempt from the project's semantic versioning and will be advertised as experimental and unstable.

Migration of Python script functionality to a Python module named *oxli*. As the code moves over there will be no change to external script functionality or their command line interfaces.

7.3 v2.x series

oxli command is now under semantic versioning. Scripts are still the advertised and preferred entry point for users. Developers and workflow systems can start to trial *oxli* but need not switch until 3.0. New functionality is added to both the scripts and the *oxli* command.

7.4 v3.0 and project renaming

Project renamed to 'oxli'; all references to 'khmer' removed from the code and documentation except for a single note in the docs. All scripts dropped as their functionality has been moved to the *oxli* command. Websites that we maintain that have 'khmer' in the URL will have redirects installed.

Refinement of the Python API continues, however it is not part of the semantic versioning of the project.

7.5 v4.0

The semantic versioning now extends to the Python API.

7.6 Python API wishlist

Python 3.0 support

API for multiple container types and implementation of the same.

Cleanup of Python/C++ class hierarchy to cut down on boilerplate glue code.

Switch to new-style Python objects (see LabelHash & Hashbits)

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