FAST: Fast Analysis of Sequences Toolbox

Travis J. Lawrence ¹, Dana L. Carper ¹, Kyle T. Kauffman ², Katherine C.H. Amrine ^{1,3}, Raymond S. Lee ⁴, Claudia J. Canales ⁴ and David H. Ardell ^{1,2*}

- ¹Quantitative and Systems Biology, University of California, Merced, CA, USA
- ² Molecular and Cell Biology Unit, University of California, Merced, CA, USA
- ³Dept. of Viticulture and Enology, University of California, Davis, CA, USA

Correspondence*:

David H. Ardell

Molecular and Cell Biology, School of Natural Sciences, University of California, Merced, 5200 North Lake Road, Merced, CA, 95343, USA, dardell@ucmerced.edu

ABSTRACT

3 FAST (Fast Analysis of Sequences Toolbox) provides simple command-line tools for rapid prototyping of powerful and reproducible bioinformatic workflows on flat-file biological sequence 4 databases. Modeled after the GNU (GNU's Not UNIX) Textutils such as grep, cut, and tr, 6 FAST tools such as fasgrep, fascut, and fastr are designed to be serially composed in UNIX-style pipelines. Unlike UNIX tools, FAST processes data per-sequence-record rather 7 than per-line. This enables efficient inline processing of biological sequence data in a concise, 9 expressive and time-tested idiom with no programming experience required. Bringing all the advantages of open-source software consistent with scientific ideals, FAST has a shallow 10 learning curve through consistency of interfaces across utilities and a high return on investment 11 of mastery through conformity with conventions of GNU utilities, Matlab, Perl, BioPerl and R. 12 Portability, ease of installation, and security of FAST are inherently derived from its BioPerl 13 and Perl foundations. The default data exchange format in FAST is MultiFastA (specifically, 14 a restriction of BioPerl FastA format), while Sanger and Illumina 1.8+ FASTQ formatted files are also compatible. Functionality highlights include fully automated numerical and taxonomic selection and sorting of sequence records, annotation-based selection of sites from alignments, 17 selection and transformation with regular expressions, molecular biological statistics including 18 composition and codon usage, and molecular population genetic statistics. FAST promotes 19 reproducibility in bioinformatic workflows by reducing manual interventions and increasing 20 documentability of data processing and facilitates the interactive investigation and control of 21 data by its users. FAST brings the power of Perl and BioPerl to bioinformatic users at the command-line without requiring previous programming skills and experience. 23

24 Keywords: Unix philosophy, MultiFASTA, pipeline, bioinformatic workflow, open source, BioPerl, regular expression, NCBI Taxonomy

1 INTRODUCTION

- 25 The field of molecular biology has changed significantly with the advent of Next Generation Sequencing
- 26 (NGS) technology. It is now commonplace to analyze gigabases of data per experiment. Commonly,
- 27 bioinformatics programs developed for visualization and basic sequence manipulation use a monolithic

⁴School of Engineering, University of California, Merced, CA, USA

application with a Graphical User Interface (GUI) (**Smith et al.**, 1994; **Rampp et al.**, 2006; **?**), usually without any facility to record and document user actions for later reproduction of a bioinformatic workflow.

Early calls for reproducible research and code publication in bioinformatics and biostatistics were motivated in part to combat fraudulent research in biomedicine (?). Open-source software in science facilitates crowd-sourced verification, correction and extension of code-based analysis (?), and reuse of software and data to make discoveries using public data (?). The absence of adequate documentation for analysis of high-dimensional data can obscure errors in processing with profound potential ethical and financial consequences (????). In one analysis, only 2 of eighteen published microarray gene-expression analyses were completely reproducible in part because of proprietary closed-source software (?). Analytical errors are a major source of retractions in the scientific literature (?). Peer-review of scientific software is generally not yet required for publication but seen as necessary for integrity of the scientific enterprise across fields and requiring more computational training for all scientists (??). Publication and documentation of bioinformatics workflows simplifies post-publication validation of research findings and may help reduce the temptation to fish for significance and encourage more objectivity in data analysis and interpretation (?)

To reduce human error and increase reproducibility in the management and documentable processing of biological data, it is desirable to create self-documenting automated bioinformatic workflows — scripts and other literate programming that encode scientific workflows for machine processing of biological data. Web-based open-source workflow suites such as Galaxy (?), Taverna (?) and BioExtract (?) facilitate this, but an even more rapid and time-tested development platform for rapid prototyping of reproducible workflows is the UNIX command-line, specifically UNIX pipelines.

The FAST utilities are modeled after the standard Unix toolkit(**Peek**, 2001) and follow the Unix philosophy to "do one thing and do it well" (**Stutz**, 2000).

Command-line utilities for bioinformatics such as the EMBOSS package (**Rice et al.**, 2000) or the scripts that come with BioPerl (**Stajich et al.**, 2002) typically offer suites of tools with simple, well-defined functions that lend themselves to scripting, but are not necessarily designed according to the UNIX toolbox philosophy specifically to interoperate through composition.

They are written in Perl using BioPerl packages (**Stajich et al.**, 2002). This makes FAST utilities easy to adopt if you are familiar with the Unix toolbox and allows fast sequence analysis even on large datasets. Extensive documentation has been developed for each utility along with useful error messages following the recommendations of (**Seemann**, 2013) to increase usability. Lastly, FAST is open source, which makes it available to anyone free of cost. This is in line with the call to make science more assessable, open, and reproducible by other scientists and the public (**Groves and Godlee**, 2012).

2 DESIGN

An overview of Version 1.0 of the FAST project is shown in Figure 1. Descriptions of the function of utilities along with GNU Textutil analogs are given in Table 1.

FAST is split into three categories selection, transformation, and annotation and analysis. The selection category contains utilities designed to select sequences and sites from alignments based on several different criteria. For example fasgrep selects sequences by matching a regular expression to the ID, description, or sequence. The transformation utilities are used to modify the ID, description, sequence, or order of sequences using several criteria. For example, fastaxsort sorts sequences within a multifasta file based on NCBI taxonomy (**Benson et al.**, 2009; **Sayers et al.**, 2009). The annotation and analysis category contains utilities to calculate sequence composition, codon usage, sequence length, and basic population genetic statistics. Additionally these utilities can also append the results of the analysis to

Table 1. FAST 1.0 utilities

Tool	Function	Textutil analog	Default field processed
fasgrep fasfilter fastax fashead fastail fascut fasuniq	regex selection of records numerical selection of records taxonomic selection of records order-based selection of records order-based selection of records index-based selection and reordering of data record reduction by content and order	grep head tail cut uniq	identifiers identifiers descriptions sequences sequences
alncut gbfalncut fassort fastaxsort	selection of sites by content selection of sites by features numerical or text sorting of records taxonomic sorting of records	sort	sequences sequences identifiers identifiers
faspaste fastr fassub faslen fascomp fascodon fasxl fasrc fasconvert	merging of records character transformations on records regex substitutions on records annotate sequence lengths annotate monomeric compositions annotate codon usage annotate biological translations annotate reverse complements convert format of records	paste tr	sequences identifiers identifiers descriptions descriptions descriptions descriptions
gbfgrep gbf2fas alnpi faswc	select feature neighborhoods by context emit sequences by regex matching on features molecular population genetic statistics tally sequences and characters	grep grep wc	features features sequences

- the sequence description, which then can be used as selection and sorting criteria by the utilities in the selection category.
- Learnability of the FAST tools is helped by making interface components such as specific options, consistent with the standard UNIX tools and across the FAST suite. Learning one FAST tool generally helps the user anticipate how to use others. In addition, specification of numerical ranges, regular expressions and other useful parameters follows standard Perl and UNIX conventions, all with the intent of making the tools fast and easy to learn.

3 IMPLEMENTATION DETAILS AND BENCHMARKING

All FAST utilities can process files or input on what is called the "standard input" stream. They all by default out to the "standard output" stream which may be connected to the "standard input" of another utility by a UNIX "pipe." It is this latter facility that eases serial processing of data. Since most FAST utilities process sequences inline, they should mostly linear runtime complexity in number of sequences. Two exceptions to inline processing in FAST utilities concern fassort and fastail which both require some paging of data into temporrary files. However, some preliminary benchmarking suggests that fassort runtimes also scale linearly in sequence number (fig 2). Benchmarking was performed using the Benchmark v1.15 perl module on a MacBook Pro ??Ghz Intel i7, 8 Gb of RAM. The average CPU time of 100 iterations to complete the analysis of six datasets consisting of 25,000, 250,000, or 1,000,000 100bp sequences (fig 2A) and 10,000, 100,000, or 1,000,000 1,000bp sequences (fig 2B) were used to estimate performance.

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- FAST is compatible with the zero-based indexing if the sequence identifier is thought as the zeroth
- 91 field of the identifier line. This field must exist in Data selection in FAST is one-based as is conventional
- 92 BioPerl coordinates and bioinformatics generally.
- 93 FAST supports automated logging to ease the reproducibility of workflows. The BioPerl backend of
- 94 FAST was version 1.6.901 downloaded in January, 2012. BioPerl dependencies of FAST scripts were
- 95 analyzed with the Cava packager (http://www.cavapackager.com). To fix some problems with
- 96 I/O, the SeqIO components were updated to version 1.6.923 on June 4, 2014 and the Root components
- 97 were updated on July 10, 2014 (github commit 50f87e9a4d). Some customizations of the BioPerl code-
- 98 base were introduced, primarily to enable the translation of sequences containing gaps.

4 INSTALLATION AND USAGE EXAMPLES

4.1 INSTALLATION AND DEPENDENCIES

- 99 FAST requires a working Perl installation and is distributed through the Comprehensive Perl Archive
- 100 Network (CPAN). In a manual install, after download, installation follows standard Perl install procedure:
- 101 perl Makefile.PL; make; make test; (sudo) make install. A small footprint of
- 102 BioPerl dependencies has been packaged together in the FAST namespace. Other CPAN dependencies
- 103 can be detected and installed by the cpan package manager. A fully automated install may on many
- 104 systems be initiated by executing perl -MCPAN -e 'install FAST'.

4.2 SELECTING SEQUENCES BY ENCODED MOTIFS

- An advantage of the annotation approach in FAST is the ability to select and sort sequences by attributes
- 106 computed and annotated into data by utilities upstream in the pipeline. For example, to select protein-
- 107 coding genes from a file cds. fas whose translations contain the N-glycosylation amino acid motif (?),
- 108 one could execute:
- 109 fasxl -a cds.fas | fasgrep -t xl0 "N[^P][ST][^P]" | fascut -f 1..-2
- The first command in the pipeline translates each sequence and appends the translation to the description
- with the tag "xl0" (indicating translation in the zeroth reading frame). The second command in the pipeline
- does a regex match using the specified motif pattern on the value of a "name:value" pair in the description
- 113 with tag "xl0", hence processing the annotations produced by fasxl. The regex argument to fasgrep is
- 114 quoted to protect the argument from interpretation by the shell. The last command in the pipeline removes
- the last field in the description, restoring records as they were before they were annotated by fasxl.

4.3 SORTING SEQUENCES BY THIRD CODON POSITION COMPOSITION

- 116 Another example illustrates the powerful expression of ranges in fascut. An optional "by" parameter in
- 117 ranges allows increments or decrements in steps larger than one. To extract the third codon position bases
- 118 from a gene, annotate their compositions, and sort sequences by their third-position adenosine contents,
- 119 do:
- 120 fascut 1:-1:3 cds.fas | fascomp | fassort -nt comp_A

5 CONCLUDING REMARKS

- 121 Planned additions in future versions of FAST include fastand and alneand for sampling,
- 122 permutations and bootstrapping of sequences and sites, respectively.

AVAILABILITY

- FAST is available through the Comprehensive Perl Archive Network (CPAN) at http://search.
- 124 cpan.org/~dhard/FAST

DISCLOSURE/CONFLICT-OF-INTEREST STATEMENT

- The authors declare that the research was conducted in the absence of any commercial or financial 125
- relationships that could be construed as a potential conflict of interest. 126

AUTHOR CONTRIBUTIONS

- D.H.A. conceived, designed, and wrote much of FAST. T.J.L. contributed major code factorizations and 127
- reorganization and fastail. K.T.K. contributed code including faspaste, and fashead. R.S.L. 128
- contributed an analysis of code dependencies for the FAST installer. All authors, especially D.L.C. 129
- and C.J.C., contributed documentation, testing, and code fixes. K.C.H.A. and D.H.A. wrote the FAST 130
- Cookbook. D.H.A. wrote the paper with major contributions from D.L.C. and T.J.L. All authors made 131
- minor contributions to the manuscript, reviewed the final version of the manuscript and agree to be 132
- accountable for its contents. 133

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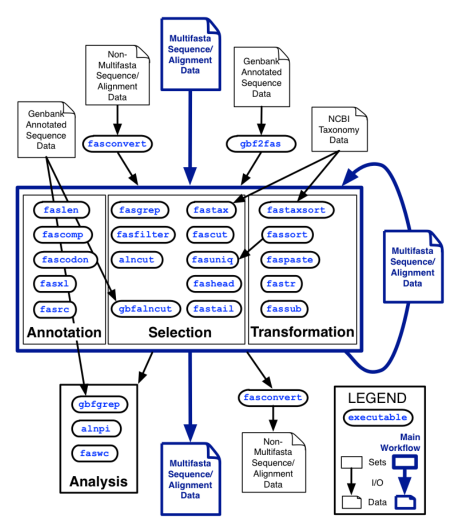


Figure 1. FAST version 1.0 with data and workflow dependencies indicated.

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FIGURES

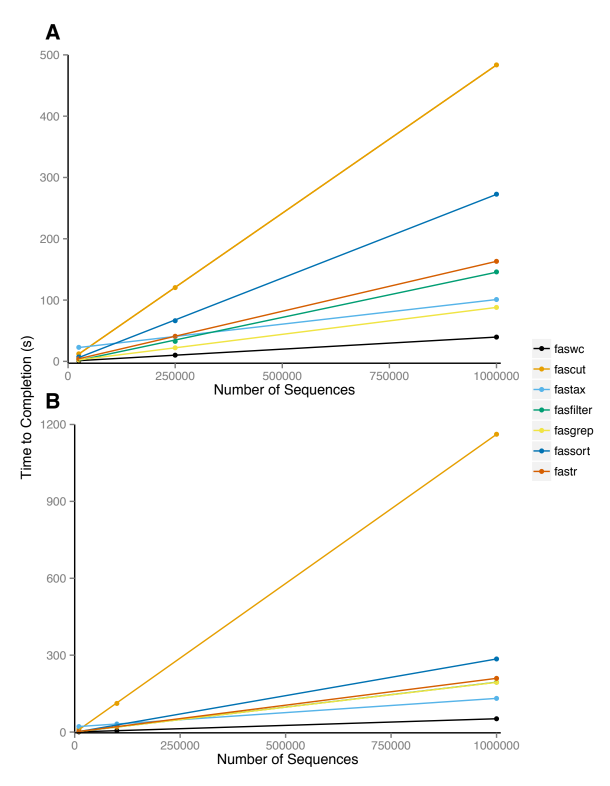


Figure 2. Average processor time of 100 repetitions required to complete analysis using indicated utility. Utilities were run on six datasets consisting of (a) 25000, 250000, and 1000000 100bp sequences and (b) 10000, 100000, and 1000000 1000bp sequences.

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