- grepq: A Rust application that quickly filters
- 2 FASTQ files by matching sequences to a set of

regular expressions

- ⁴ Nicholas D. Crosbie, Melbourne Veterinary School, University of Melbourne,
- 5 Parkville, Victoria, Australia
- 6 ORCID: 0000-0002-0319-4248
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Abstract

- п Regular expressions (regex) (Kleene 1951) have been an im-
- portant tool for finding patterns in biological codes for decades
- (Hodgman 2000 and citations therein), and unlike fuzzy-finding
- approaches, do not result in approximate matches. The perfor-
- mance of regular expressions can be slow, however, especially
- when searching for matching patterns in large files. *grepq* is
- a Rust application that quickly filters FASTQ files by matching
- sequences to a set of regular expressions. grepq is designed
- with a focus on performance and scalability, is easy to install
- and easy to use, enabling users to quickly filter large FASTQ

files, and to update the order in which patterns are matched against sequences through an in-built *tune* command. *grepq* is open-source and available on *GitHub* and *Crates.io*.

Statement of need

- The ability to quickly filter FASTQ files by matching sequences to a set of regular expressions is an important task in bioinformatics, especially when working with large datasets. The importance and challenge of this task will only grow as sequencing technologies continue to advance and produce ever larger datasets (Katz et al. 2022). The uses cases of grepq are diverse, and include preprocessing of FASTQ files before downstream analysis, quality 31 control of sequencing data, and filtering out unwanted sequences. Where decisions need be made quickly, such as in a clinical set-33 tings (Bachurin et al. 2024), biosecurity (Valdivia-Granda 2012), and wastewater-based epidemiology in support of public health measures (Choi et al. 2018; Sims and Kasprzyk-Hordern 2020; Xylogiannopoulos 2021; Merrett et al. 2024), the ability to quickly filter FASTQ files by matching sequences to a set of regular ex-38 pressions is attractive as it circumvents the need for more timeconsuming bioinformatic workflows.
- Regular expressions are a powerful tool for matching sequences,
- ₄₂ but they can be slow and inefficient when working with large

datasets. Furthermore, general purpose tools like grep (Free Software Foundation 2023) and ripgrep (A. Gallant 2025) are not optimized for the specific task of filtering FASTQ files, and ocassionaly yield false positives as they scan the entire FASTQ record, including the sequence quality field. Tools such awk (Aho, Kernighan, and Weinberger 1988) and gawk (Free Software Foundation 2024) can be used to filter FASTQ files without yielding false positives, but they are significantly slower than grepq and can require the development of more complex scripts to achieve the same result.

Implementation

- grepg is implemented in Rust, a systems programming language known for its safety features, which help prevent common pro-
- gramming errors such as null pointer dereferences and buffer over-
- flows. These features make Rust an ideal choice for implementing
- a tool like *grepq*, which needs to be fast, efficient, and reliable.
- Furthermore, *grepq* obtains its performance and reliability, in part,
- by using the seq io (Schlegel and Seyboldt 2025) and regex (Gal-
- lant et al. 2025b) libraries. The seq_io library is a well-tested
- library for parsing FASTQ files, designed to be fast and efficient,
- and which includes a module for parallel processing of FASTQ
- records through multi-threading. The regex library is designed

- to work with regular expressions and sets of regular expressions,
- and is known to be one of the fastest regular expression libraries
- currently available (Gallant et al. 2025a). The regex library sup-
- ports Perl-like regular expressions without look-around or backref-
- erences (documented at https://docs.rs/regex/1.*/regex/#syntax).
- Further performance gains were obtained by:

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- use of the RegexSet struct from the regex library to match multiple regular expressions against a sequence in a single pass, rather than matching each regular expression individually (the RegexSet is created and compiled once before entering any loop that processes the FASTQ records, avoiding the overhead of recompiling the regular expressions for each record)
- multi-threading to process the records within an input FASTQ
 file in parallel through use of multiple CPU cores
- use of the zlib-ng backend to the flate2 library to read and
 write gzip-compressed FASTQ files, which is faster than the
 default miniz_oxide backend
 - use of an optimised global memory allocator (the *mimalloc* library (Mutiple, n.d.)) to reduce memory fragmentation and improve memory allocation and deallocation performance
 - buffer reuse to reduce the number of memory allocations and deallocations

- use of byte slices to avoid the overhead of converting to and
 from string types
- in-lining of performance-critical functions
- use of the *write_all* I/O operation that ensures the data is written in one go, rather than writing data in smaller chunks

Feature set

- ⁹⁴ *grepq* has the following features:
- support for presence and absence (inverted) matching of a
 set of regular expressions
- IUPAC ambiguity code support (N, R, Y, etc.)
- gzip support (reading and writing)
- JSON support for pattern file input and *tune* command output,
 allowing named regular expression sets and named regular
 expressions (pattern files can also be in plain text)
- the ability to set predicates to filter FASTQ records on the
 header field (= record ID line) using a regular expression, minimum sequence length, and minimum average quality score
 (supports Phred+33 and Phred+64)
- the ability to output matched sequences to one of four formats
 (including FASTQ and FASTA)
 - the ability to tune the pattern file with the tune command: this command will output a plain text or JSON file with the patterns

sorted by their frequency of occurrence in the input FASTQ file or gzip-compressed FASTQ file (or a user-specified number of FASTQ records). This can be useful for optimizing the pattern file for performance, for example, by removing patterns that are rarely matched

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 the ability to count and summarise the total number of records and the number of matching records (or records that don't match in the case of inverted matching) in the input FASTQ file

Other than when the *tune* command is run, a FASTQ record is deemed to match (and hence provided in the output) when any of the regular expressions in the pattern file match the sequence field of the FASTQ record. An example (abridged) output of the *tune* command (when given with the **–json-matches** flag) is shown below:

```
"regexCount": 298,
    "regexName": "Primer contig 06aR",
    "regexString": "CCCCG[CT]C[AT]ATT[CT]"
},
{
    "regexCount": 1143,
    "regexName": "Primer contig 03",
    "regexString": "GG[AG][ACGT]GGC[ACGT]GCAG"
}
],
    "regexSetName": "conserved 16S rRNA regions"
}
```

When the count option (-c) is given with the tune command, grepq will count the number of FASTQ records containing a sequence 126 that is matched, for each matching regular expression in the pat-127 tern file. If, however, there are multiple occurrences of a given 128 regular expression within a FASTQ record sequence field, grepq 129 will count this as one match. When the count option (-c) is not given with the tune command, grepq provides the total number of 131 matching FASTQ records for the set of regular expressions in the 132 pattern file. 133

Colorized output for matching regular expressions is not implemented to maximise speed and minimise code complexity, but can be achieved by piping the output to *grep* or *ripgrep* for testing purposes.

Performance

- The performance of *grepq* was compared to that of *fqgrep*, *seqkit grep*, *ripgrep*, *grep*, *awk*, and *gawk* using the benchmarking tool *hyperfine*. The test conditions and results are shown in **Table 1**, **Table 2** and **Table 3**.
- Table 1: Clock times and speedup of various tools for filtering FASTQ records against a set of regular expressions. Test FASTQ file: SRX26365298.fastq (not gzip-compressed) was 874MB in size, and contained 869,034 records.

tool	clock time (s)		speedup		
	mean	S.D.	× grep	× ripgrep	× awk
grepq	0.19	0.0021	1814.71	18.74	870.79
fqgrep	0.34	0.01	1010.24	10.43	484.77
ripgrep	3.56	0.01	96.85	1.00	46.48
seqkit grep	122.05	0.90	2.83	0.03	1.36
grep	344.79	1.24	1.00	0.01	0.48
awk	165.45	1.59	2.08	0.02	1.00
gawk	287.66	1.68	1.20	0.01	0.58

grepq v1.3.6, fqgrep v.1.02, ripgrep v14.1.1, seqkit grep v.2.9.0, grep 2.6.0-FreeBSD, awk v. 20200816, and gawk v.5.3.1. fqgrep and seqkit grep were run with default settings, ripgrep was run with -colors 'match:none' -no-line-number, and grep was run with -color=never. The tools were configured to output matching records in FASTQ format. The clock times, given in

seconds, are the mean of 10 runs, and S.D. is the standard deviation of the clock times, also given in seconds.

Table 2: Clock times and speedup of various tools for filtering gzip-compressed
FASTQ records against a set of regular expressions. Test FASTQ file:
SRX26365298.fastq.gz was 266MB in size, and contained 869,034 records.

tool	clock t	ime (s)	speedup
1001	mean	S.D.	× grep
grepq	1.707	0.002	2.09
fqgrep	1.84	0.01	1.94
ripgrep	3.57	0.01	1.00

Test conditions and tool versions as above, but *grepq* was run with the -x option, *ripgrep* with the -z option, and *grep* with the -z option.

Table 3: Clock times and speedup of various tools for filtering FASTQ records against a set of regular expressions. Test FASTQ file: SRX22685872.fastq (not gzip-compressed) was 104GB in size, and contained 139,700,067 records.

tool	clock tir	speedup	
1001	mean	S.D.	× ripgrep
grepq	26.97	0.22	4.38
fqgrep	50.47	0.62	2.34
ripgrep	118.161	1.068	1.00

Test conditions and tool versions as described in the footnote to Table 1. Note that when *grepq*was run under the same conditions but SRX22685872.fastq was gzip-compressed, and a gzipcompressed output was generated, the clock time was 148.01 seconds, with a memory resident time for the *grepq* process of 116M as reported by the *top* command (Apple Inc. 2023c).

4 Testing

- The output of *grepq* was compared against the output of *fqgrep*, *seqkit grep*, *ripgrep*, *grep*, *awk* and *gawk*, using the *stat* command (Apple Inc. 2023b), and any difference investigated using the *diff* command (Apple Inc. 2023a). Furthermore, a custom utility, *spikeq* (Crosbie 2024b), was developed to generate synthetic FASTQ files with a known number of records and sequences with user-specified lengths that were spiked with a set of regular expressions a known number of times. This utility was used to test the performance of *grepq* and the aforementioned tools under controlled conditions.
- Finally, a bash test script (see *examples/test.sh*, available at *grepq*'s Github repository) and a simple Rust CLI application, *predate* (Crosbie 2024a), were developed and utilised to automate system testing, and to monitor for performance regressions.
- grepq has been tested on macOS 15.0.1 (Apple M1 Max) and Linux Ubuntu 20.04.6 LTS (AMD EPYC 7763 64-Core Processor). It may work on other platforms, but this has not been tested.

Availability and documentation

- grepq is open-source and available at *GitHub* (https://github.com/Rbfinch/gre pq) and *Crates.io* (https://crates.io/crates/grepq).
- Documentation and installation instructions for *grepq* are available at the same

 GitHub repository, and through the **-h** and **-help** command-line options, which

 includes a list of all available commands and options, and examples of how to

 use them. Example pattern files in plain text and JSON format are also provided,

 as well as test scripts. *grepq* is distributed under the MIT license.

Conclusion

The performance of grepq was compared to that of fqgrep, seqkit grep, ripgrep, 189 grep, awk, and gawk using the benchmarking tool hyperfine. The results show 190 that grepq is significantly faster than the other tools tested, with a speedup of 191 1814.71 times over grep, 870.79 times over awk, and 18.74 times over ripgrep. 192 The performance of grepg was also compared to that of fggrep and ripgrep 193 when filtering gzip-compressed FASTQ files, with grepq being 2.09 times faster 194 than ripgrep and 1.94 times faster than fggrep. When coupled with its exceptional runtime performance, grepg's feature set make it a powerful and flexible 196 tool for filtering large FASTQ files.

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205 Conflicts of interest

206 The author declares no conflicts of interest.

References

Aho, Alfred V., Brian W. Kernighan, and Peter J. Weinberger. 1988. *The AWK*Programming Language. https://www.cs.princeton.edu/~bwk/btl.mirror/.

Apple Inc. 2023a. *The Diff Command*.

```
——. 2023b. The Stat Command.
    ——. 2023c. The Top Command.
    Bachurin, Stanislav S, Mikhail V Yurushkin, Ilya A Slynko, Mikhail E Kletskii,
213
       Oleg N Burov, and Dmitriy P Berezovskiy. 2024. "Structural Peculiarities
214
       of Tandem Repeats and Their Clinical Significance." Biochemical and Bio-
215
       physical Research Communications 692: 149349.
216
    Choi, Phil M, Ben J Tscharke, Erica Donner, Jake W O'Brien, Sharon C Grant,
217
       Sarit L Kaserzon, Rachel Mackie, et al. 2018. "Wastewater-Based Epidemi-
218
       ology Biomarkers: Past, Present and Future." TrAC Trends in Analytical
219
       Chemistry 105: 453-69.
220
    Crosbie, Nicholas D. 2024a. "predate: Catch bugs and performance regres-
       sions through automated system testing." https://github.com/Rbfinch/preda
222
       te.
223
             2024b. "spikeq: Generates synthetic FASTQ records free of se-
224
       quences defined by regex patterns, or containing spiked sequences based
225
       on regex patterns." https://github.com/Rbfinch/spikeg.
226
    Free Software Foundation. 2023. GNU Grep 3.11. Free Software Foundation.
227
       https://www.gnu.org/software/grep/manual/grep.html.
228
       —. 2024. GAWK: Effective AWK Programming: A User's Guide for GNU
229
       Awk, for the 5.3.1. Free Software Foundation. https://www.gnu.org/softwa
230
       re/gawk/manual/gawk.html.
231
    Gallant et al. 2025a. "rebar." https://github.com/BurntSushi/rebar.
232
    ——— et al. 2025b. "regex." https://github.com/rust-lang/regex.
233
    Gallant, Andrew. 2025. "Ripgrep: Recursively Search the Current Directory for
       Lines Matching a Pattern." https://github.com/BurntSushi/ripgrep.
235
    Hodgman, T. Charles. 2000. "A Historical Perspective on Gene/Protein Func-
```

tional Assignment." Bioinformatics 16 (1): 10-15.

- ²³⁸ Katz, Kenneth, Oleg Shutov, Richard Lapoint, Michael Kimelman, J Rodney
- Brister, and Christopher O'Sullivan. 2022. "The Sequence Read Archive:
- A Decade More of Explosive Growth." Nucleic Acids Research 50 (D1):
- D387-90.
- Kleene, SC. 1951. "Representationof Events in Nerve Nets and Finite Automata." *CE Shannon and J. McCarthy*.
- Merrett, James E, Monica Nolan, Leon Hartman, Nijoy John, Brianna Flynn,
- Louise Baker, Christelle Schang, et al. 2024. "Highly Sensitive Wastewater
- Surveillance of SARS-CoV-2 Variants by Targeted Next-Generation Ampli-
- con Sequencing Provides Early Warning of Incursion in Victoria, Australia."
- Applied and Environmental Microbiology 90 (8): e01497–23.
- Mutiple. n.d. "Mimalloc: A Rust Wrapper over Microsoft's MiMalloc Memory

 Allocator."
- Schlegel, Markus, and Adrian Seyboldt. 2025. "seq_io: FASTA and FASTQ
 parsing and writing in Rust." https://github.com/markschl/seq_io.
- Sims, Natalie, and Barbara Kasprzyk-Hordern. 2020. "Future Perspectives of Wastewater-Based Epidemiology: Monitoring Infectious Disease Spread and Resistance to the Community Level." *Environment International* 139:
- ²⁵⁶ 105689.
- Valdivia-Granda, Willy A. 2012. "Biodefense Oriented Genomic-Based
 Pathogen Classification Systems: Challenges and Opportunities." *Journal* of Bioterrorism & Biodefense 3 (1): 1000113.
- Xylogiannopoulos, Konstantinos F. 2021. "Pattern Detection in Multiple
 Genome Sequences with Applications: The Case of All SARS-CoV-2
 Complete Variants." bioRxiv, 2021–04.