SOFTWARE TOOL ARTICLE

Falco: high-speed FastQC emulation for quality control of sequencing data [version 1; peer review: 1 approved, 1 approved with reservations]

Guilherme de Sena Brandine, Andrew D. Smith

Department of Quantitative and Computational Biology, University of Southern California, Los Angeles, California, 90089, USA

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Abstract
Quality control is an essential first step in sequencing data analysis, and software tools for quality control are deeply entrenched in standard pipelines at most sequencing centers. Although the associated computations are straightforward, in many settings the total computing effort required for quality control is appreciable and warrants optimization. We present falco, an emulation of the popular FastQC tool that runs on average three times faster while generating equivalent results. Compared to FastQC, falco also provides greater scalability for datasets with longer reads and more flexible visualization of HTML reports.

Keywords
FastQC, high-throughput sequencing, quality control

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| report            |   |   |

1. R. Henrik Nilsson, University of Gothenburg, Gothenburg, Sweden
2. Weihong Qi, Functional Genomics Center Zurich, Zürich, Switzerland

Any reports and responses or comments on the article can be found at the end of the article.
**Introduction**

High-throughput sequencing is routinely used to profile copy number variations in cancers, assemble genomes of microbial organisms, quantify gene expression, identify cell populations from single-cell transcriptomes in a variety of tissues, and track epigenetic changes in developing organisms and diseases, among numerous other applications. New sequencing protocols are constantly being introduced, and as the cost of sequencing per base decreases, sequencing data is growing in abundance, dataset size and read length.

When high-throughput sequencing data is generated it often undergoes common upstream analysis steps involving quality control (QC), adapter trimming, filtering contaminants and low-quality reads, and mapping reads to a reference genome or transcriptome. Excluding sequence assembly applications, read mapping should be the most computationally expensive step early in analysis pipelines. In comparison, the time and computation required for QC should be negligible. However, the efficiency of mapping algorithms has improving substantially over the last decade, while software for QC has received far less attention. As a consequence, the computation required for QC is appreciable, and can no longer be ignored when considering the total cost of sequencing.

The most commonly used tool for quality control of sequencing data is FastQC, which, since 2011, has incorporated a wide range of QC checks covering multiple use cases. FastQC has been cited over 3,000 times, with citations increasing steadily since its introduction. Its analysis reports have become the standard for several QC tools, and automated analysis pipelines often rely on its evaluation as a safety criteria to proceed with downstream steps or, alternatively, to filter, trim or ultimately discard the data.

FastQC is implemented in a modular design, where multiple independent analysis procedures are run sequentially after an input record is read. This design allows new modules to be incorporated easily, but it implies that each analysis module is applied independently to each read, so the time required to process each read is the sum of the processing times for each module. If multiple modules use similar measurements, such as nucleotide content or average sequence quality, the same measurement will be calculated multiple times, causing the total analysis run time to increase.

Several QC software tools have been introduced since FastQC, many focusing on speed improvements, more flexible module visualization, incorporation of paired-end reads and filtering sequences that failed QC checks. Despite proposing different alternatives to calculate and present QC results, the modules available in these tools are largely similar to FastQC’s (Table 1).

At the same time, FastQC’s analysis results are already part of many standard initial analysis pipelines. If a new QC software tool were to be incorporated in these pipelines, it is desirable that its results, and its output formats, remain consistent with those of FastQC.

| Table 1. Comparison of analysis modules provided by fastp and HTQC, two commonly used QC software tools. |
| FastQC module | fastp | HTQC |
|----------------|-------|------|
| Per base sequence quality | No    | Yes  |
| Per base N content | Yes   | Yes  |
| Per tile sequence quality | No    | Yes  |
| Per sequence quality scores | No    | Yes  |
| Per sequence GC content | No    | No   |
| Sequence length distribution | No    | Yes  |
| Sequence duplication levels | Yes   | No   |
| Overrepresented sequences | Yes   | No   |
| Adapter content | No    | No   |
| Kmer content | Yes   | No   |

To improve the speed of quality control while retaining the behaviour of FastQC, we developed FastQC Alternative Code (falco), an emulation of FastQC’s current analysis modules. We show that falco generates the same results as FastQC across a wide variety of datasets of different read lengths, sizes, file formats and library preparation protocols at significantly shorter running times. We also present example datasets from the public domain where FastQC fails to generate reports even when run on high-performance computing hardware, demonstrating that falco expands the range of possible cases in which these quality control metrics can be applied.

**Methods**

**Implementation choices**

We designed falco to faithfully emulate FastQC’s calculations, results and text reports. Our goal was to minimize the effort required to replace FastQC with falco in the context of larger automated analysis pipelines. We use the same set of command line arguments, configuration file names and formats. We also produce the same plain text format output, and the same report structure as FastQC, allowing users to take advantage of improved speed without adjusting to different program behaviors.

There are major differences between the implementations of falco and FastQC. While FastQC’s code emphasizes modularity in a way that allows for additional types of QC information to be added easily and uniformly, falco’s design centralizes the function to read sequences from the input file and collects the minimum data necessary to subsequently create all modules after file processing. To ensure consistency with FastQC, we wrote each module’s source code based on FastQC’s implementation, adapting the portions that relate to sequence processing and maintaining the postprocessing functions that define how the collected data is used to generate summaries and reports.

**Operation**

Compilation of falco requires a GNU GCC compiler version 5.0.0 (July 16, 2015; full support for the C++11
standard) or greater. Once installed, falco can be run on uncompressed files (FASTQ and SAM) without any additional dependencies. In order to process files in gzip compressed FASTQ and BAM formats, falco must be compiled with the ZLib and HTSLib libraries. The full documentation on how to compile, install dependencies and run the program is available in the README file in the falco repository.

Use cases
Like FastQC, falco can be applied to any sequencing data file (i.e. a file of sequenced reads) in the accepted formats. The only required command line argument is the path to the input file. Also like FastQC, a wide range of options can be provided if users only require a given subset of its analysis modules or outputs. The letters and symbols used for command line arguments were chosen to maintain consistency with FastQC’s options. As mentioned above, this choice is to facilitate integration with larger pipelines that already employ FastQC and depend on its behaviours.

Falco can be run on a FASTQ format file named example.fastq with the following simple command:

$ falco example.fastq

This will generate three files:

1. fastqc_data.txt: The complete numerical values generated in each module’s individual analysis.
2. fastqc_report.html: A visual page display of the text report’s data and plots generated in modules.
3. summary.txt: A short summary indicating whether the input file passed or failed each module, and whether any warnings were raised.

Default configuration files are contained in a Configuration directory that is included with the program, but falco also allows users to manually define the thresholds for statistics to be considered a pass, warning or fail, the list of adapters to search for in reads and the list of contaminants to check overrepresented sequences by using configuration files in the same format used by FastQC.

Results
Falco matches FastQC’s output
We compared the output of falco to its FastQC counterpart using 11 datasets (Table 2). The tests consist of Illumina files originating from a range of different library preparation protocols for DNA, RNA and epigenetic experiments, as well as reads from the nanopore technology. For simplicity, Illumina paired-end datasets were only tested on their first read.

FASTQ files available in the Sequencing Read Archive (SRA) were downloaded using the fastq-dump command from the SRA toolkit. We used the following flags when running fastq-dump: --skip-technical, -readids, -read-filter pass, -dumpbase, -split-3 and -clip. One dataset was downloaded from the Whole Human Genome Sequencing Project.

We directly compared the text summary for each output of falco to FastQC’s output summary files, obtaining the same outputs (pass/warn/fail) for all tested criteria in all datasets.

To assess if falco’s output is consistent with FastQC’s format, we used the fastqc R package version 0.1.2 and MultiQC version 1.7. Both tools can successfully parse the text reports generated by falco for the tested files. Differences in the text reports generated by falco and MultiQC’s output summary files, obtaining the same outputs (pass/warn/fail) for all tested criteria in all datasets.

Falco is faster than popular QC tools
Some alternative software tools exist for quality control of sequencing data, and users may opt for them due to their efficiency in cases where not all FastQC analysis modules are necessary. Among these, fastp has gained popularity for its speed and versatile set of options for trimming. HTQC has demonstrated superior runtime to FastQC even when generating FASTQ format output files corrected by trimming adapters and filtering (which requires both input and output). HTQC is another tool that was developed with the intent to both improve speed performance and incorporate trimming functions after quality control. The two programs were used as benchmarks to compare with falco’s performance.

| Table 2. Datasets used for comparison with FastQC’s output and run time speed benchmarking between QC tools. |
|---|---|---|---|---|---|---|
| test | accession | reference | file size (FASTQ) | reads | length (bp) | protocol |
| 1 | SRR10124060 | unpublished | 7.3GB | 25,172,255 | 130 | RNA-Seq |
| 2 | SRR10143153 | unpublished | 11.0GB | 15,949,900 | 150 | miRNA-Seq |
| 3 | SRR3897196 | 21 | 4.2GB | 15,570,348 | 100 | BS-Seq |
| 4 | SRR6264732 | 22 | 1.6GB | 18,807,797 | 150 | ChIP-Seq |
| 5 | SRR1853178 | 23 | 130.0GB | 510,210,716 | 60 | Drop-Seq |
| 6 | SRR6267347 | 24 | 20.0GB | 305,434,830 | 100 | 10x genomics |
| 7 | SRR6954584 | 5 | 56.0GB | 152,853,013 | 150 | Microwell-Seq |
| 8 | SRR691268 | 25 | 46.0GB | 192,904,649 | 50 | ATAC-Seq |
| 9 | SRR878357 | unpublished | 38.0MB | 3,284 | 64,000 | Nanopore |
| 10 | wgs-FAB49164 | 17 | 8.4GB | 746,333 | 180,000 | Nanopore |
| 11 | SRR6059706 | unpublished | 1.4GB | 892,313 | 150,000 | Nanopore |
Although most fastp modules are both calculated and displayed equivalently to FastQC, one major difference between these tools is how overrepresented sequences are estimated. fastp counts the sequences at every $P$ reads (which users may specify), whereas FastQC stores the first 100,000 reads encountered for the first time, and subsequently checks if the following sequences match any of the stored candidates. This choice of implementation causes fastp’s runtime to greatly differ when over-representation is enabled. Conversely, FastQC’s runtime does not seem to be affected by disabling the overrepresented sequences module. For a comprehensive comparison between programs, we have measured the run times for our test datasets both with and without the overrepresented sequences module enabled. Programs were compared both in compressed (gzipped FASTQ) and uncompressed (plain FASTQ) file formats.

Files used to assess falco’s output comparison to FastQC (Table 2) were also used for speed benchmarking. Tests were executed in an Intel Xeon CPU E5-2640 v3 2.60GHz processor with a CentOS Linux 7 operating system. All file I/O was done using local disk to reduce variability in execution runtime. Programs were instructed to run using a single thread.

Real run times for falco, fastp and FastQC on uncompressed FASTQ format, with the overrepresented sequences module on and off. Asterisks (*) indicate tests in which tools did not run to completion.

| test | falco overrep off | fastp overrep off | FastQC overrep off | falco overrep on | fastp overrep on | FastQC overrep on | HTQC |
|------|------------------|------------------|-------------------|------------------|------------------|-------------------|------|
| 1    | 1m19s            | 2m19s            | 3m49s             | 1m25s            | 6m23s            | 3m50s             |      |
| 2    | 45s              | 1m31s            | 2m21s             | 51s              | 5m23s            | 2m24s             |      |
| 3    | 33s              | 1m10s            | 1m35s             | 35s              | 2m26s            | 1m36s             |      |
| 4    | 1m01s            | 2m06s            | 3m01s             | 1m03s            | 3m50s            | 3m00s             |      |
| 5    | 16m05s           | 42m40s           | 44m57s            | 18m17s           | 53m09s           | 44m59s            |      |
| 6    | 12m26s           | 23m18s           | 26m39s            | 12m29s           | 47m32s           | 26m38s            |      |
| 7    | 8m40s            | 17m34s           | 22m31s            | 8m34s            | 44m41s           | 22m31s            |      |
| 8    | 7m08s            | 14m37s           | 16m06s            | 6m31s            | 18m19s           | 16m11s            |      |
| 9    | 2s               | 1s               | *                 | 1s               | 27s              | *                 |      |
| 10   | 2m23s            | 2m32s            | *                 | 2m34s            | 5m22s            | *                 |      |
| 11   | 22s              | 31s              | *                 | 23s              | 1m14s            | *                 |      |

Real run times for falco, fastp and FastQC on gzip compressed FASTQ format.

| test | falco overrep off | fastp overrep off | FastQC overrep off | falco overrep on | fastp overrep on | FastQC overrep on |
|------|------------------|------------------|-------------------|------------------|------------------|-------------------|
| 1    | 1m19s            | 2m19s            | 3m49s             | 1m25s            | 6m23s            | 3m50s             |
| 2    | 45s              | 1m31s            | 2m21s             | 51s              | 5m23s            | 2m24s             |
| 3    | 33s              | 1m10s            | 1m35s             | 35s              | 2m26s            | 1m36s             |
| 4    | 1m01s            | 2m06s            | 3m01s             | 1m03s            | 3m50s            | 3m00s             |
| 5    | 16m05s           | 42m40s           | 44m57s            | 18m17s           | 53m09s           | 44m59s            |
| 6    | 12m26s           | 23m18s           | 26m39s            | 12m29s           | 47m32s           | 26m38s            |
| 7    | 8m40s            | 17m34s           | 22m31s            | 8m34s            | 44m41s           | 22m31s            |
| 8    | 7m08s            | 14m37s           | 16m06s            | 6m31s            | 18m19s           | 16m11s            |
| 9    | 2s               | 1s               | *                 | 1s               | 27s              | *                 |
| 10   | 2m23s            | 2m32s            | *                 | 2m34s            | 5m22s            | *                 |
| 11   | 22s              | 31s              | *                 | 23s              | 1m14s            | *                 |
that ran to completion demonstrate that falco’s analysis times are also significantly smaller in comparison.

**Falco scales for longer Nanopore reads**

Nanopore sequencing is gaining popularity in genome assembly applications and as a low-cost protocol to quantify short reads. Nanopore sequencers can generate reads of up to millions of bases, and assessing quality metrics for these datasets is fundamental to test for potential problems in quality or bias in specific regions of such long reads. While FastQC is capable of making summaries for protocols such as 454\(^\text{27}\) PacBio\(^\text{28}\), which generate sequences with around 10,000 bases per read, we have observed that it does not run to completion when given files with larger reads of over 100,000 bases. Files for which FastQC’s analysis does not finish are marked with an asterisk in Table 3 and Table 4. Falco successfully completes its analysis on these datasets, demonstrating that it can equally be used as a QC tool for longer reads.

**Falco allows dynamic visualization of results**

Despite FastQC’s clarity in its HTML reports, graphs are displayed as static images and have limited visualization flexibility, such as tile heatmaps not displaying raw deviations from average Phred scores in base positions or raw values in line plots not being visible. We have opted to display falco’s analysis results using the Plotly JavaScript library\(^\text{29}\), which allows interactive changes of axis labels, hovering on data points to visualize raw values and screenshots from specific position on the plot (Figure 1). This choice of presentation provides greater options to explore and interpret QC results.

![Figure 1. Sample HTML report for test 8 (accession SRR891268). Layout and plots are based on FastQC’s HTML report.](image-url)
results while maintaining the visualization standards set by FastQC.

Conclusions
Falco\(^1\) is a faster alternative to calculate the wide range of QC metrics generated by FastQC. It is entirely based on emulating the analysis modules FastQC provides while running faster than popular QC tools and generating dynamic visual summaries of analysis results. Both falco’s text and HTML outputs provide the same information generated by FastQC’s report, so tools that parse these files for custom visualization and downstream analysis can seamlessly incorporate falco into their pipeline.

Data availability
Datasets used to compare Falco and FastQC are shown in Table 2. Guidance for how to accept accession wgs-FAB49164 is available from the Benchmark directory of the falco GitHub page.

Software availability
Source code for falco available at: https://github.com/smithlabcode/falco.

The scripts used to download files and reproduce the benchmarking steps described are also available in the same repository within the “benchmark” directory.

Archived source code at time of publication: http://doi.org/10.5281/zenodo.3520933\(^2\).

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Weihong Qi
Functional Genomics Center Zurich, Zürich, Switzerland

The authors developed falco, an emulation of the popular FastQC tool, which is faster and can handle very long Nanopore reads. It is a useful development, especially for core facilities and research labs that produce high volumes of sequencing data regularly, where generating read QC reports in a timely fashion is indeed helpful. I only have a few questions and one minor comment:

**Main questions:**
1. The implementation session could be expanded with more details. From my understanding, the major improvement was identified duplicated analysis in FastQC analysis modules, and implemented a single analysis workflow that was sufficient to generate the same modularized results. But it is not clear to me which changes make falco to handle long ONT reads successfully, while the original FastQC failed.

2. The original FastQC is portable (Unix, Mac and Windows). It also has a GUI version for less experienced users. These features are not important for experienced users and automated workflows where analyzing large amounts of data in a short time is the focus. But they can be important for other type of end users. The authors should at least point out these differences.

3. In results, run times of multiple QC tools analyzing different datasets were compared, how about the RAM usages?

**Minor comment:**
The sentence “While FastQC is capable of making summaries for protocols such as 454 PacBio, which generate sequences with around 10,000 bases per read” should be updated to "While FastQC is capable of making summaries for protocols such as 454 PacBio, which generate sequences with around 10,000-20,000 bases per read".

**Is the rationale for developing the new software tool clearly explained?**
Yes

Is the description of the software tool technically sound?
Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Genome informatics.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 09 Jan 2021

Guilherme de Sena Brandine, University of Southern California, Los Angeles, USA

The reviewer has raised some important questions about points not addressed by the manuscript. We provide our responses below, and highlight changes made to the manuscript to address the reviewer’s comments.

The authors developed falco, an emulation of the popular FastQC tool, which is faster and can handle very long Nanopore reads. It is a useful development, especially for core facilities and research labs that produce high volumes of sequencing data regularly, where generating read QC reports in a timely fashion is indeed helpful. I only have a few questions and one minor comment:

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We really appreciate the reviewer highlighting the missing details regarding FastQC’s
behavior. Upon trying to address this comment, we have further explored the FastQC code to understand why it failed for long reads, and we have learned that the perl script that wraps the FastQC call imposes a maximum memory limit of 250 MB per thread, which we were not aware of at the time we wrote the manuscript, and was prohibitive for the long read samples we have selected. Changing this configuration internally allowed us to run FastQC in the samples that we were previously unable to, thus allowing us to report a more comprehensive comparison of both time and memory for every test we have gathered. For this reason, we have removed the section named “Falco scales for larger nanopore reads”, instead replacing it with a paragraph at the end of the subsection named “Falco is faster than popular QC tools”, where the memory usage of each mapper is discussed. We have also filled Tables 3 and 4 with the results of running FastQC in the three long-read samples in our tests under the same hardware settings used for other tests.

2) The original FastQC is portable (Unix, Mac and Windows). It also has a GUI version for less experienced users. These features are not important for experienced users and automated workflows where analyzing large amounts of data in a short time is the focus. But they can be important for other type of end users. The authors should at least point out these differences.

We thank the reviewer for this observation. We have made modifications to the first paragraph of the “implementation choices” subsection under “methods” to clarify that falco was designed for UNIX systems and does not include a graphical user interface.

3) In results, run times of multiple QC tools analyzing different datasets were compared, how about the RAM usages?

We appreciate the observation about RAM comparison. We have added two paragraphs in the manuscript that address memory requirement in more detail. A “system requirement” subsection under “methods” was added to emphasize that falco requires about 100 MB for short read samples and under 1 GB for samples with read lengths of at most 1 million. As stated in question (1), we also reported the RAM usage for the software tools compared in the tests, both for short-read and long-read tests in the section “Falco is faster than popular QC tools”.

Minor comment:
The sentence “While FastQC is capable of making summaries for protocols such as 454\protect\cite{27} PacBio\protect\cite{28}, which generate sequences with around 10,000 bases per read” should be updated to “While FastQC is capable of making summaries for protocols such as 454\protect\cite{27} PacBio\protect\cite{28}, which generate sequences with around 10,000-20,000 bases per read”.

We thank the reviewer for the observation. The section containing this sentence was removed given the shift in the focus of the manuscript to memory comparison, as addressed in questions (1) and (3).

Competing Interests: No competing interests were disclosed.
R. Henrik Nilsson
Gothenburg Global Biodiversity Centre, University of Gothenburg, Gothenburg, Sweden

The authors present a welcome addition to the flora of FastQC-style read processing packages. The fact that it is a drop-in replacement for FastQC is particularly nice.

The manuscript is a bit too short in my opinion. I miss some background information and some performance-related data. If the authors want to address a wide audience, they should probably work a bit more on the installation instructions and documentation too. The authors should probably also consider defining who their target audience is.

**Introduction:**
- As a discretionary comment: the authors sometimes, but not always, use the Oxford comma (see example below). I wonder if this is something that should be streamlined. “sequencing data is growing in abundance, dataset size and read length⁹,” vs. “...adapter trimming, filtering contaminants and low-quality reads, and mapping reads to a reference genome or transcriptome.”

- “as a safety criteria” should probably be “as a safety criterion”.

**Methods:**
- Good thinking behind the “We designed falcoⁱ³ to faithfully...” paragraph. Nobody would be helped by yet another set of new file formats. A drop-in replacement is the way to go, if you ask me. And that is indeed what the authors deliver.

- I must make the observation, though, that the name “falco” may not be available (?): https://www.falcoseed.com/ca/article/cibus-registers-new-falco-brand-79k-canola-hybrid/ (some other more or less commercial uses of “Falco” can be found on https://en.wikipedia.org/wiki/Falco). I'm not sure how North American trademark/proprietary laws operate, but I suggest that the authors consult with the lawyers of their university. Better safe than sorry, right. (I consulted with our lawyers at one point, and they had me change the name of a software package we were working on at the time.)

- Two unintended line breaks:
  - “uniformly, falco’s design centralizes”

**Results:**
Please cite the Sequence Read Archive formally; see Kodama et al. (2012).  

I like the reproducible nature of the “Results” section.  

Two superfluous line breaks:  

“gzip  
compressed files.”  

Conclusions:  
Both “Falco” and “falco” are used in the manuscript. You’d think that the name would be fixed as either “Falco” or “falco”.  

Software availability:  
Why not use the term “open source” at least once in the manuscript?  

Miscellaneous questions and observations:  
Out of curiosity: how does falco compare to Liu et al. (2019)?  

I’ve seen one software tool for quality-score-based trimming of sequences that actually reads the entire query file into memory, and then started to process it. This works less well as data files continues to grow, obviously. Is it worth pointing out that falco does not do this? What is, in fact, the maximum file size allowed by falco? Or is this dictated solely by the operating system?  

You can produce some pretty funny behavior in some other tools for sequence QC/trimming by feeding them a file with a single FastQC entry in it, speaking of nothing. Is there, then, a minimum file size or number of query sequences for falco?  

Unless I’m mistaken, there are no fastq files available on https://github.com/smithlabcode/falco. I think the authors should make one or two available, so that it will be smooth and easy for the reader to try the software out. “The scripts used to download files and reproduce the bench-marking steps described are also available in the same repository within the “benchmark” directory.” comes across as somewhat indirect to me.  

What, exactly, are the hardware and software requirements for falco? This may be worth pointing out. Between the lines I read “any computer you can install the GNU GCC compiler on” – but not all readers will probably read it this way. Instead you’ll get the question: “Does it run on Windows 10?”  

Also, how much memory is needed to run falco? And how much memory is used up by falco when it processes a large file?  

Between the lines (“Programs were instructed to run using a single thread.”), I take it that falco can use multiple threads. Is that correct? And if so, why not point it out more explicitly? And out of curiosity: when you run falco on 4 cores, do you see a 4x speedup? Or is the
bottleneck something else (disk IO?) than raw computational power? How does it scale with the number of cores, in other words?

- Suppose my dataset is 10 Gb, and that I have 15 Gb left of free disk space. Do I dare to run falco on that dataset? How large is the output compared to the input? In a “minimum” (no extra features) as well as “maximum” (all extra features) mode?

- Suppose a user finds a bug, or wants to put forward a feature request. How can the user do that? Should this be mentioned in the manuscript?

- The first question I always get from users of my software tools is: “where can I download the Windows binaries?”. Is it, actually, not a good idea to be explicit about the fact that this is a command-line tool that you compile on your own computer? Would probably save the authors some time to be upfront with this.

- The list of references comes across as somewhat untidy. Some examples follow below. The authors should probably go through all references to make sure they comply with journal specifications.

  - Journal names are sometimes abbreviated, sometimes not. Ref 2 is abbreviated, whereas ref 3 should be abbreviated “BMC Bioinform.”

  - Article titles: should verbs and key nouns in article titles have a leading uppercase letter, as in, e.g., ref 5, or should they not, as in ref 1?

  - Should page ranges be written out in full (ref 23, “1202–1214.”) or should they be abbreviated (ref 25, “1213–8.”)?

Are the installation instructions a bit too thin? I'd say yes, at least if the intention of the authors is to address a diverse audience and not just readers with Linux-style experience. To simulate a less experienced user, I used my son's MacBook Pro and tried to install falco on it following the manual:

“Upon downloading, inflating and moving to the source directory, installation can be done through the following commands:

... $ ./configure CXXFLAGS="-O3 -Wall"
$ make all
$ make install"

So I did:

$ cd src
$ ./configure CXXFLAGS="-O3 -Wall"
-bash: ./configure: No such file or directory

And then
conda install -c bioconda falco
-bash: conda: command not found

And that was it. No further clues or assistance to be found in the instructions.

If the authors are happy with this behavior, then they should make it clear in the manuscript that falco is not for everyone, but rather only for those with significant Linux-style experience.

“Source code for falco available at: https://github.com/smithlabcode/falco.” – the trailing “.” should be removed, I’d say. The link won’t work for users who copy-and-paste it into their browser. The same thing goes for

“Archived source code at time of publication: http://doi.org/10.5281/zenodo.352093313.” where both the reference and the “.” cause problems.

References
1. Kodama Y, Shumway M, Leinonen R, International Nucleotide Sequence Database Collaboration: The Sequence Read Archive: explosive growth of sequencing data. Nucleic Acids Res. 2012; 40 (Database issue): D54-6 PubMed Abstract | Publisher Full Text
2. Liu X, Yan Z, Wu C, Yang Y, et al.: FastProNGS: fast preprocessing of next-generation sequencing reads. BMC Bioinformatics. 2019; 20 (1): 345 PubMed Abstract | Publisher Full Text

Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Partly

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Metabarcoding; molecular ecology; systematics; mycology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have
significant reservations, as outlined above.

Author Response 09 Jan 2021

Guilherme de Sena Brandine, University of Southern California, Los Angeles, USA

The reviewer has presented a thorough feedback to the description of the Falco software tool, as well as its implementation, description and documentation. We truly appreciate the very helpful comments, and provide our responses to improvement suggestions below. Comments were divided and numbered to allow us to refer to them in other places in our response if certain modifications to the manuscript pertain to multiple comments.

(1) The authors present a welcome addition to the flora of FastQC-style read processing packages. The fact that it is a drop-in replacement for FastQC is particularly nice.

The manuscript is a bit too short in my opinion. I miss some background information and some performance-related data. If the authors want to address a wide audience, they should probably work a bit more on the installation instructions and documentation too. The authors should probably also consider defining who their target audience is.

We appreciate the comments on the introduction. We have expanded on the target audience on further comments (we address these in more detail on comment 16). We also fully agree that more background information can be provided. We expanded the second paragraph of the “introduction” section to add a brief description of some common quality control tests applied to most next-generation sequencing datasets.

(2) As a discretionary comment: the authors sometimes, but not always, use the Oxford comma (see example below). I wonder if this is something that should be streamlined. “sequencing data is growing in abundance, dataset size and read length” vs. “…adapter trimming, filtering contaminants and low-quality reads, and mapping reads to a reference genome or transcriptome.”

We thank the reviewer for this observation. We have revised the manuscript and ensured that the Oxford comma is adopted across the entire manuscript.

(3) “as a safety criteria” should probably be “as a safety criterion”.

We fully agree with the reviewer. This correction was made on the manuscript.

(4) Good thinking behind the “We designed falco to faithfully...” paragraph. Nobody would be helped by yet another set of new file formats. A drop-in replacement is the way to go, if you ask me. And that is indeed what the authors deliver.

We appreciate the comments, thank you!

(5) I must make the observation, though, that the name “falco” may not be available (?): https://www.falcoseed.com/ca/article/cibus-registers-new-falco-brand-79k-canola-hybrid/ (some other more or less commercial uses of “Falco” can be found on
I'm not sure how North American trademark/proprietary laws operate, but I suggest that the authors consult with the lawyers of their university. Better safe than sorry, right. (I consulted with our lawyers at one point, and they had me change the name of a software package we were working on at the time.)

We really appreciate the reviewer raising the possible legal issue that may arise from the program name. We have researched the matter and believe that the software name should not raise legal issues, both due to the program not having any commercial or profitable goals and the name “Falco” being a common proper noun in many Latin languages.

(6) Two unintended line breaks: “uniformly, falco’s design centralizes”

We thank the reviewer for this observation. This line was removed from the manuscript.

(7) Please cite the Sequence Read Archive formally; see Kodama et al. (2012).¹

We followed the reviewer’s suggestion and added the appropriate citation when referring to the Sequence Read Archive. We also corrected the meaning of the acronym from “Sequencing Read Archive”, as it was previously written.

(8) I like the reproducible nature of the “Results” section.

We appreciate the comment, thank you!

(9) Two superfluous line breaks: “gzip compressed files.”

We have removed the line break from the manuscript.

(10) Both “Falco” and “falco” are used in the manuscript. You’d think that the name would be fixed as either “Falco” or “falco”.

We thank the reviewer for this observation. The notation used in the manuscript used “Falco” at the start of sentences and “falco” everywhere else, to resemble the name of the binary program used in the command-line interface. We have modified the manuscript to use “Falco” everywhere except in the “use cases” section, where an example command-line call for the program is shown.

(11) Why not use the term “open source” at least once in the manuscript?

We thank the reviewer for this observation. To address this, we started the “Implementation choices” subsections with the following sentence: Falco is an Open Source C++ implementation of the FastQC software tool built for UNIX-based operating systems.

(12) Out of curiosity: how does falco compare to Liu et al. (2019)?

We have downloaded the software from the URL provided in the manuscript (github.com/megagenomics/fastprongs) and performed comparisons on identical hardware
to what was used in the manuscript. We tested on two datasets: Dataset 1 consisted of 76 million 150 base reads from arabidopsis (SRR12075121 in SRA), and dataset 2 contained 139 million 100 base reads from chicken (SRR5015166 in SRA). On dataset 1, Falco ran in 9:40 and FastProNGS ran in 5:16 with 3 threads (the default program configuration) and 10:42 on a single thread. On dataset 2, Falco ran in 14:30 and FastProNGS ran in 7:55 with 3 threads and 16:13 with a single thread. We noticed that FastProNGS only reports the following modules in their output: Basic statistics, adapter content, per base sequence quality, per base sequence content and sequence length distribution. We also tried to run Falco by enabling only these modules. Under these settings Falco ran in 4:35 for dataset 1 and 7:12 for dataset 2. In all tests, Falco ran with 92 MB of RAM, whereas FastProNGS used 1.26 GB. In a long-read dataset (test 12 in the manuscript), FastProNGS did not run successfully unless we configured it to only consider the first 200 bases of each read, in which case FastProNGS ran in 2 seconds, but only reported summaries for the first 200 bases of all reads. Falco ran in 12 seconds for this dataset.

A very meaningful conclusion of this comparison is the potential advantage of multithreading in QC, as evidenced by the steep decrease in processing time from FastProNGS when multithreading is enabled. We noticed, upon inspecting its source code, that this performance improvement can be explained by FastProNGS reading multiple reads in batch and allowing a new set of reads to be loaded while the previous batch of reads is processed. In contrast, Falco loads and processes each read sequentially, which reduces RAM usage but makes multithreading difficult in its current implementation. The performance of FastProNGS suggests that switching to a “batch processing” paradigm may have significant speed advantages when multiple cores are used and enough RAM is available to load reads in batch, and this is something we will incorporate in future versions of Falco, especially in order to address (18). We thank the reviewer for bringing this tool to our attention.

(13) I’ve seen one software tool for quality-score-based trimming of sequences that actually reads the entire query file into memory, and then started to process it. This works less well as data files continues to grow, obviously. Is it worth pointing out that falco does not do this? What is, in fact, the maximum file size allowed by falco? Or is this dictated solely by the operating system?

Both disk and memory requirements will depend on the length of the largest read in the dataset, as Falco processes the input FASTQ one read at a time. To address the computational requirements necessary to run Falco in more detail in the manuscript, we created an additional subsection named “system requirements” under the “Methods” section, where the computational resources (memory and disk) required to run Falco successfully are discussed. Furthermore, we have added a paragraph in the “results” section summarizing the memory and disk usage for the tests used for comparison across programs.

(14) You can produce some pretty funny behavior in some other tools for sequence QC/trimming by feeding them a file with a single FastQC entry in it, speaking of nothing. Is there, then, a minimum file size or number of query sequences for falco?

There are no minimum or maximum file sizes required by Falco. We have tested (although
not disclosed in the manuscript) that Falco successfully runs on empty files and single-read files. We thank the reviewer for having raised this issue, and have addressed that there are no constraints in file size or number of reads in the “system requirements” section stated in (13).

(15) Unless I’m mistaken, there are no fastq files available on https://github.com/smithlabcode/falco. I think the authors should make one or two available, so that it will be smooth and easy for the reader to try the software out. “The scripts used to download files and reproduce the benchmarking steps described are also available in the same repository within the “benchmark” directory.” comes across as somewhat indirect to me.

We thank the reviewer for this observation. While we cannot provide the full FASTQ files used to perform our comparisons in the GitHub repository, we do agree that the documentation of our tests can be made simpler for users who wish to test the program. We made modifications in our repository to simplify both testing in an example file and testing in the FASTQ files used in the manuscript for comparison. Specifically, we added (1) direct links to the SRA files under the “benchmark” directory and (2) an “example.fq” file, consisting of a FASTQ file of 1000 reads, which is used as input for the example commands provided in the README.

(16) What, exactly, are the hardware and software requirements for falco? This may be worth pointing out. Between the lines I read “any computer you can install the GNU GCC compiler on” – but not all readers will probably read it this way. Instead, you’ll get the question: “Does it run on Windows 10?”.

We agree that constraints should be disclosed in more detail, and that the limited support for usage of Falco on Windows should be more explicit. We have rephrased the first paragraph in the “implementation choices”. The last sentences disclose more explicitly that Falco, by design, is a UNIX-centric program made to be run on a command line and that, unlike FastQC, it cannot be run in a graphical user interface.

(17) Also, how much memory is needed to run falco? And how much memory is used up by falco when it processes a large file?

Falco requires under 1 GB of memory for any short or long read file generated by the current sequencing technologies. More memory will be required when technologies expand read lengths to the order of millions or billions of bases per read. We have added a discussion of disk and memory requirements under the “systems requirement” section, and also discussed the memory usage of the programs compared in the manuscript under the section “Falco is faster than popular QC tools”.

(18) Between the lines (“Programs were instructed to run using a single thread.”), I take it that falco can use multiple threads. Is that correct? And if so, why not point it out more explicitly? And out of curiosity: when you run falco on 4 cores, do you see a 4x speedup? Or is the bottleneck something else (disk IO?) than raw computational power? How does it scale with the number of cores, in other words?
Falco, like FastQC currently does not use multiple threads to process a single file, and no significant speed difference was observed when running fastp with multiple threads, which is why we focused our comparison on single-thread across the software tools. Despite QC computations being fast relative to IO, our comparison with FastProNGS described in (12) suggests that multithreading can lead to speed improvements if reading and processing are done in parallel, and we certainly plan on exploring this paradigm in the next release of Falco. We have also rephrased the third paragraph of the subsection “Falco is faster than popular QC tools” to say “Both fastp and fastqc were instructed to run on a single thread” to avoid ambiguities regarding Falco’s multithread option.

(19) Suppose my dataset is 10 Gb, and that I have 15 Gb left of free disk space. Do I dare to run falco on that dataset? How large is the output compared to the input? In a “minimum” (no extra features) as well as “maximum” (all extra features) mode?

We have addressed the disk requirement on the “system requirements” section disclosed in (13) and (14), specifically adding the sentence “The total disk space required to store the three output files generated by Falco is under 1 MB”. Like FastQC, Falco’s output is a set of reports whose size scales with the maximum read length of the input but are never under 1 MB in total. We fully agree that the fact that disk space is not crucial to run Falco should be made more explicit.

(20) Suppose a user finds a bug, or wants to put forward a feature request. How can the user do that? Should this be mentioned in the manuscript?

We thank the reviewer for this observation. All issues and bug reports can be done through our GitHub page, the same one provided for the source code. To make this clearer for users, we have added a sentence at the “software availability” section, stating that errors, installation problems and bugs can be reported in the “Issues” section in the same URL provided to download the source code.

(21) The first question I always get from users of my software tools is: “where can I download the Windows binaries?”. Is it, actually, not a good idea to be explicit about the fact that this is a command-line tool that you compile on your own computer? Would probably save the authors some time to be upfront with this.

We fully agree with the reviewer, and have added the statements of a more specific target audience as discussed in (16).

(22) The list of references comes across as somewhat untidy. Some examples follow below. The authors should probably go through all references to make sure they comply with journal specifications.

Journal names are sometimes abbreviated, sometimes not. Ref 2 is abbreviated, whereas ref 3 should be abbreviated “BMC Bioinform.”

Article titles: should verbs and key nouns in article titles have a leading uppercase letter, as in, e.g., ref 5, or should they not, as in ref 1?

Should page ranges be written out in full (ref 23, “1202–1214.”) or should they be abbreviated (ref 25, “1213–8.”)?
We really appreciate the keen observations on the reference standards. We have reviewed our citations and standardized journals to their non-abbreviated names, uppercase letters only in leading words, and pages written in full.

(26) Are the installation instructions a bit too thin? I’d say yes, at least if the intention of the authors is to address a diverse audience and not just readers with Linux-style experience. To simulate a less experienced user, I used my son’s MacBook Pro and tried to install falco on it following the manual:

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And then

conda install -c bioconda falco
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And that was it. No further clues or assistance to be found in the instructions.

If the authors are happy with this behavior, then they should make it clear in the manuscript that falco is not for everyone, but rather only for those with significant Linux-style experience.

We really appreciate the reviewer bringing up this observation about our documentation. We agree that the wording of “moving to the source directory” was misleading and may cause users to try to run the commands on the “src” directory. We have updated our README with clearer command line instructions that show the user how to clone the repository or download a release file, as well as which directory to move to in order to run the commands. We are striving to make the documentation as clear and simple as possible, and truly appreciate these suggestions on how these can be improved.

(27) “Source code for falco available at: https://github.com/smithlabcode/falco.” – the trailing “,” should be removed, I’d say. The link won't work for users who copy-and-paste it into their browser. The same thing goes for

“Archived source code at time of publication: http://doi.org/10.5281/zenodo.352093313.” where both the reference and the “,” cause problems.
We appreciate this observation and the potential problems punctuation near links may cause. We have ensured that the trailing dots and citations are clearly separated from the links and that they will not cause problems when copying URLs directly from the manuscript.

**Competing Interests:** No competing interests were disclosed.

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