

GitHub statistics as a measure of the impact of open-source bioinformatics software

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- 8 Abstract
- 9 Modern research is increasingly data-driven and reliant on bioinformatics software. Publication is a
- 10 common way of introducing new software, but not all bioinformatics tools get published. Giving
- there are competing tools, it is important not merely to find the appropriate software, but have a
- metric for judging its usefulness. Journal's impact factor has been shown to be a poor predictor of
- software popularity; consequently, focusing on publications in high-impact journals limits user's
- choices in finding useful bioinformatics tools. Free and open source software repositories on popular
- 15 code sharing platforms such as GitHub provide another venue to follow the latest bioinformatics
- trends. The open source component of GitHub allows users to bookmark and copy repositories that
- are most useful to them. This Perspective aims to demonstrate the utility of GitHub "stars,"
- 18 "watchers," and "forks" (GitHub statistics) as a measure of software impact. We compiled lists of
- impactful bioinformatics software and analyzed commonly used impact metrics and GitHub statistics
- of 50 genomics-oriented bioinformatics tools. We present examples of community-selected best
- 21 bioinformatics resources and show that GitHub statistics are distinct from the journal's impact factor
- 22 (JIF), citation counts, and alternative metrics (Altmetrics, CiteScore) in capturing the level of
- community attention. We suggest the use of GitHub statistics as an unbiased measure of the usability
- of bioinformatics software complementing the traditional impact metrics.

25 1 Introduction

- 26 It is currently undeniable that bioinformatics tools and databases represent a highly impactful part of
- 27 modern research (Wren, 2016). Many journals focus exclusively on publishing software tools and
- databases. Some of the most famous examples include application notes published in *Bioinformatics*,
- database and web-server issues published by *Nucleic Acids Research*, software articles published in
- 30 Frontiers Bioinformatics and Computational Biology, PLOS Computational Biology, BMC
- 31 *Bioinformatics*. However, given the continued growth of bioinformatics publications (Wren, 2016)
- 32 (Supplementary Figure S1), it is getting increasingly difficult to find software that will be useful in
- real-life applications. Recently, a term "software crisis" was coined to illustrate the problem of
- 34 finding useful software (Mangul et al., 2018).
- 35 Finding useful bioinformatics software is further hindered by publication lag. It often takes more than
- a year from the time of presubmission inquiry, potential resubmission and the peer-review period to

- 37 the accepted publication. Such delays inevitably diminish the potential impact of published software.
- Non-peer-reviewed preprint publishing (arXiv, biorXiv, PeerJ, AsapBio) aims to eliminate
- 39 publication lag. However, the number of preprints grows nearly ten times faster than the number of
- 40 peer-reviewed publications¹², further complicating finding useful software.
- Reviews of bioinformatics resources can help orient a scientist in the wealth of published tools and
- databases. Such reviews are typically written about bioinformatics software published in high-impact
- 43 journals while leaving preprints and unpublished software largely out of scope. Furthermore, reviews
- may be limited by the experience of the authors, as well as by a bias to review software published in
- 45 high-impact journals. Thus, while helpful in orienting a novice in the topic, reviews may overlook
- 46 useful bioinformatics resources.
- 47 Although the peer-review process helps to publish high-quality bioinformatics software, it is
- 48 unknown at the time of publication which tools and databases will be embraced by the scientific
- 49 community and which will be forgotten (Wren and Bateman, 2008). In fact, a study based on text
- 50 mining found that over 70% of published bioinformatics software resources are never reused (Duck
- et al., 2016). A recent analysis of the usability of bioinformatics software confirmed these
- observations by highlighting issues with software accessibility and installation (Mangul et al., 2018).
- Notably, a journal's impact factor, calculated as the average number of citations received in a
- calendar year by the total number of articles and reviews published in that journal in the preceding
- two years (JIF) is not a good predictor of software popularity (Seglen, 1997; Wren, 2016), making it
- hard to predict whether a bioinformatics tool or a database published in a high-impact journal will be
- 57 useful in real-life applications.

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2 Limitations of alternative metrics to measure the impact of bioinformatics software

- Alternative metrics have been proposed to alleviate the shortcomings of JIF or the lack of it in
- preprint publishing. CiteScore, a metric developed by Scopus includes more document types and
- 61 citation sources, and uses the 3-year time window to calculate the ratio of citations over the total
- number of citable items, has been proposed as a consistent alternative to JIF (Silva and Memon,
- 63 2017). Article-level metrics, or Altmetrics, is currently the most widely used alternative to measure
- the impact of scholarly material, including preprints (Priem et al., 2010; Shema et al., 2014). In
- addition to academic citations, this metric aggregates mentions in social media networks, such as
- Twitter, online discussions and recommendations. Although in principle Altmetrics can be applied to
- any research output that has a digital object identifier (DOI), including datasets, code and software
- 68 (Piwowar, 2013), its use for measuring the impact of bioinformatics software is less common.
- 69 Furthermore, Altmetrics may still be biased by high impact factor (hence, greater exposure and
- discussion) (Adie, 2013), and overlook the practical usability of software. The usefulness of these
- alternative metrics on measuring the impact of bioinformatics software remains unknown.

3 Community-guided selection of bioinformatics resources

- An increasing number of bioinformaticians choose to develop their tools on popular code sharing
- web services, such as GitHub (Wilson et al., 2017). Besides code-sharing services, GitHub combines
- a version control system (Bryan, 2017) with features found in popular social network sites such as

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 $^{^{1}\ \}underline{https://www.crossref.org/blog/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-higher-than-journal-articles/preprints-growth-rate-ten-times-growth-r$

² http://www.prepubmed.org/monthly_stats/?Subject=Bioinformatics

- Facebook and Twitter (Lima et al., 2014). Users may try the tools and bookmark the most practically
- useful ones by "starring," "watching," and/or "forking" them. "Starring" a repository is similar to
- bookmarking it as a favorite, while "watching" is a more advanced feature allowing a user to receive
- 79 all, or selected, updates about a repository. "Forking" further advances user's involvement by
- 80 creating a copy of a forked repository under the user's account, allowing him/her to offer code
- 81 enhancements by creating pull requests. GitHub creates a natural ecosystem for software
- 82 development where the amount of community attention to a repository is directly proportional to its
- popularity (Hu et al., 2016). We expect the number of stars, watchers, and forks ("GitHub statistics")
- 84 to reflect some evidence of the practical utility of the software and suggest they should be used to
- 85 inform selection of the most useful resources.

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4 Lists of community-selected software as reviews of practical utility

- 87 Although using GitHub statistics as a guide for selecting the most popular software, including
- bioinformatics tools, has been suggested³ (Hu et al., 2016; Russell et al., 2018), it does not alleviate
- 89 the problem of finding the right field-specific resources among a large number of bioinformatics
- 90 repositories⁴. The abundance of GitHub repositories gave rise to field-specific collections of the most
- 91 useful resources (tools, databases, papers, books, and videos), frequently referred to as "awesome"
- 92 lists (Table 1, Supplementary Table S1). They are assembled by inspired individuals who empirically
- 93 try them and bookmarks the most valuable repositories (Marlow et al., 2013). These collections of
- 94 links and notes are themselves published on GitHub and starred by the community. The collections
- 95 may themselves be assembled into field-specific "awesome" lists of lists (Supplementary Table S2).
- 96 Being analogous to bookmarks freely accessible on the web, they do not require any programming
- 97 skills to be used. These collections may be compared with field-specific reviews peer-reviewed by
- 98 the community and may be used to quickly prioritize practically useful resources.

5 Community attention as a distinct and universal measure of software impact

- 100 To better understand the relationship between community attention-based and traditional impact
- metrics, we compared GitHub statistics, JIF, CiteScore, Altmetrics, citation count, and software age
- of 50 popular genomics-oriented bioinformatics tools published in peer-review journals, developed
- on GitHub, and starred 50 times or more (Supplementary Table S3, Methods⁵). Principal component
- analysis (PCA, Figure 1) and correlation analysis (Supplementary Figure S2) showed the expected
- analysis (1 C/1, 1 igure 1) and correlation marysis (Supplementary 1 igure 32) showed the expected
- 105 correlation between similarly calculated JIF and CiteScore (Pearson Correlation Coefficient, PCC =
- 106 0.73). The software age and citation counts were also correlated (PCC = 0.60) as would be expected
- for older software having more chance of being cited. However, neither the software age nor citation
- 108 counts were correlated with JIF (PCC = -0.23/-0.02, respectively), suggesting that citations of
- bioinformatics software have minimal effect on JIF. Furthermore, the correlation between JIF and
- Altmetrics was relatively modest (PCC = 0.49), suggesting that Altmetrics captures a different level
- of impact. The poor correlation among traditional impact metrics complicates their use for measuring
- the software impact.

⁴ <u>https://www.researchgate.net/post/Is_there_too_many_bioinformatics_tools2__</u> # Community attention as a distinct and universal measure of software impact

³ https://gitstar-ranking.com/

⁵ Methods at https://github.com/mdozmorov/bioinformatics-impact (will be made public post-publication)

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- Being a measure of attention of open-source software development community, GitHub statistics are
- expected to capture the practical usability of software that may be missed by traditional impact
- metrics. Indeed, GitHub statistics (counts of "stars," "watches," and "forks") were highly correlated
- with each other (average PCC = 0.92) but were distinct from other metrics. Neither JIF nor
- 117 Altmetrics correlated with GitHub statistics (average PCC = -0.09/0.14, respectively), highlighting
- differences between community attention-based and traditional impact metrics. Interestingly, GitHub
- statistics and citation counts showed modest correlation (average PCC = 0.66), suggesting that
- practically useful software cited more frequently. However, the software age correlated with GitHub
- statistics to a much lesser extent (average PCC = 0.32), suggesting that the age of the software does
- not necessarily indicate its usefulness. We suggest that GitHub statistics should be used as an
- objective addition to JIF and other traditional impact metrics in measuring the practical utility of
- bioinformatics software.

125 6 Author contributions

MD envisioned the project, collected and analyzed the data, and wrote the manuscript.

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130 **8** Conflict of interest

The author declares that he has no conflict of interest.

132 **9 References**

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- 168 **10** Figures
- Figure 1. Principal component analysis of bioinformatics impact measures, colored by metric
- 170 **type.**

171 Table 1. Popular collections of bioinformatics resources, accessed on 2018-11-30

Name	Description	URL	Stars	Watchers	Forks		
General bioinformatics collections							
deeplearning- biology	A list of deep learning implementations in biology	https://github.com/hussius /deeplearning-biology	775	148	198		
deep-review	A collaboratively written review paper on deep learning genomics and precision medicine	https://github.com/greenel ab/deep-review	742	120	188		
Awesome- Bioinformatics	A curated list of awesome Bioinformatics libraries and software	https://github.com/daniele cook/Awesome- Bioinformatics	583	80	158		

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awesome	Awesome resources on Bioinformatics data science machine learning programming language Python Golang R Perl and miscellaneous stuff	https://github.com/shenwei356/awesome	304	21	115
genomicspapers	The Leek group guide to genomics papers	https://github.com/jtleek/genomicspapers	299	54	134
biotools	A list of useful bioinformatics resources	https://github.com/jdidion/ biotools	205	24	60
getting-started- with-genomics- tools-and- resources	Unix R and python tools for genomics	https://github.com/crazyh ottommy/getting-started- with-genomics-tools-and- resources	157	27	69
Field-specific bioinformatics collections					
awesome- single-cell	List of software packages for single- cell data analysis including RNA-seq ATAC-seq etc	https://github.com/seanda vi/awesome-single-cell	712	154	303
RNA-seq- analysis	RNAseq analysis notes from Ming Tang	https://github.com/crazyh ottommy/RNA-seq- analysis	260	44	104
ChIP-seq- analysis	ChIP-seq analysis notes from Ming Tang	https://github.com/crazyh ottommy/ChIP-seq- analysis	252	41	136
awesome- cancer-variant- databases	A community- maintained repository of cancer clinical knowledge bases and databases focused on cancer variants	https://github.com/seanda vi/awesome-cancer- variant-databases	109	23	25
awesome-10x-genomics	List of tools and resources related to the 10x Genomics GEMCode/Chromium system	https://github.com/johand ahlberg/awesome-10x- genomics	63	8	12
DNA-seq- analysis	DNA sequencing analysis notes from Ming Tang	https://github.com/crazyh ottommy/DNA-seq- analysis	53	7	34
awesome- microbes	List of computational resources for analyzing microbial	https://github.com/stevets a/awesome-microbes	33	5	16

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	sequencing data				
DNA- methylation-	DNA methylation analysis notes from	https://github.com/crazyhottommy/DNA-	25	4	22
analysis	Ming Tang	methylation-analysis			