What Is the Price of Open-Source Software?

The notion that all scientific software should be open-source and free has been actively promoted in recent years, mostly from the top down via mandates from funding agencies but occasionally from the bottom up, as exemplified by a recent Viewpoint in this journal. A commonly articulated rationale is that the results of scientific research funded by government grants should be free for society and that the scientific community benefits from free access. The purpose of this Viewpoint is to examine the consequences of these opinions.

What Is Scientific Software? Modern computational chemistry software is an extremely complex product based on advanced scientific ideas (models and theories) and sophisticated algorithms that transform these ideas from equations into useful tools. The development of practical software that can be used by nonexperts to solve contemporary research problems requires considerable technical effort to produce and maintain robust, efficient, and validated code. Unlike the development of, for example, a smart-phone app, where the code base is small and a relatively large community can easily write extensions and add-ons, production of scientific software involves the curation of millions of lines of source code. The complexity of this code demands long-term user and developer support to maintain its integrity and performance while keeping up with new computer architectures, fixing bugs, and adding features. Recognizing the importance of these ideas, various funding agencies in the U.S. have made “sustainable software” a key priority in the distribution of research support. Sustainability is a critical goal, but one that can be realized in various ways.

Good Software Is Important to Science. Computational chemistry software is an essential scientific instrument that facilitates discovery and innovation far beyond the laboratories in which it is created, an achievement that was recognized by the 1998 and 2013 Nobel Prizes in Chemistry. Focusing on quantum chemistry software in this Viewpoint, we note that today any chemist can (with very little training) use numerous quantum chemistry programs as teaching and research tools that aid in the design and interpretation of experiments.

A software package should be more than just a tool for end users, however; it should also be a platform to develop and test new models and algorithms. Maintaining a code base requires extensive validation, and given the complexity of modern computational methods, even testing of “pilot code” or a “proof-of-principle” implementation requires access to basic software infrastructure, for example, an integrals library, a self-consistent field procedure, efficient I/O and memory management, tools for manipulating tensors, and so forth. Modularity is a laudable goal, but in reality, “interoperability” often comes at the expense of performance. In high-performance codes, the aforementioned components are tightly interwoven, to the extent that expert help is often required to modify key components or to develop nonstandard interfaces to them. As such, the ability to innovate along either applied or theoretical lines depends crucially on the quality of the software and the availability of documentation and expert support.

As examples, consider two widely used electronic structure programs, QCHEM and MOLPRO. These codes consist of ~5.5 and ~2.5 million lines of source code, respectively, written in multiple languages and each in continuous development over several decades. QCHEM incorporates scientific advances reported in more than 300 peer-reviewed scientific publications, whereas methods implemented for the first time in MOLPRO have led to 20 high-impact papers that have each been cited over 300 times. Neither code is static: more than 70 scientists are actively contributing to MOLPRO, and the QCHEM developer base numbers more than 100. Such agile innovation comes at a price, however. Significant effort is required to keep the code robust, efficient, and sound and to provide the documentation that ensures the usability of new methods and the extensibility of older ones.

Software from academia is often developed with an emphasis on ideas rather than implementation, fed by the need for timely peer-reviewed journal publications that provide ongoing grant support and future jobs for graduate students. To bring new ideas to the production level, with software that is accessible (and useful for) the broader scientific community, contributions from expert programmers are required. These technical tasks usually cannot—and generally should not—be conducted by graduate students or postdocs, who should instead be focused on science and innovation. To this end, QCHEM employs four scientific programmers. Other quantum chemistry codes (e.g., MOLPRO, TURBOMOLE, JAGUAR, MOLCAS, PQS, and ONETEP) face the same challenges and adopt similar models to ensure sustainability.

It is important to distinguish these academically led software ventures from purely commercial endeavors. The large majority of the code in a package like QCHEM is funded by the government, either through grants to academic groups or, in some cases, through technology grants to the company itself. The role of the company programmers is to enable sustainability through bug fixes, user support, release management, and the addition of features that academic developers either cannot or will not add themselves. Programmers employed by the company place emphasis on functionality, robustness, and performance, more so than scientific innovation. They are directly addressing the “reproducibility problem.” Sales revenue cannot support the entire development cost of an academic code, but it contributes critically to its sustainability. The cost that the customer pays for a code like QCHEM reflects this funding model: it is vastly lower than the development cost, particularly for academic customers but also for industry. It primarily reflects the sustainability cost.

Software Is Not Data. In his Viewpoint, Gezelter argues that both software and data should be open, yet it is important not to conflate the two. Software is not data, and simply because it is feasible to put software on the Internet does not imply that it should be posted. Software is a product that contains an intellectual component (models and algorithms) but owes its

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There Is No Free Software

The absence of a commercial platform. Software is not di
clear in the case of commercial software, where license fees are
new laser system may enable the pursuit of new science,
when purchasing a computer or other lab equipment. Just like a
computing technologies, information storage media, computer
commonly conducted in an industrial setting. This is a common
software projects by granting academic groups access to the
consideration is whether the software is a product, not a scientific
Consider an analogy from the field of photovoltaics. Scientific findings concerning the mechanistic details of charge
generation and exciton propagation in a given material are results that merit discussion in the peer-reviewed literature.
However, creating a new solar panel based on this research requires significant additional engineering effort, which is most
commonly conducted in an industrial setting. This is a common
impact on the manufacturing process. In this model, users buy the software that o
software is a product, not a scientific finding, more akin to, say, an
NMR spectrometer—a sophisticated instrument—than to the
spectra produced by that instrument.

There Is No Free Software. The creation of scientific software is a labor-intensive process, and its support and curation even
more so. How do we pay for these labor costs? The answer is clear in the case of commercial software, where license fees are
used to defray the costs of development and support. In this model, users buy the software that fits their research needs and
affords them the highest productivity. The decision mechanism and price-versus-deliverables choices are similar to those faced
when purchasing a computer or other lab equipment. Just like a
new laser system may enable the pursuit of new science, software that offers a competitive advantage is a sensible
investment of research funds.

Interestingly, Gezelter specifically mentions the Quantum Chemistry Program Exchange (QCPE), an early repository for
open-source software, as having contributed greatly to the growth of the field. It is therefore telling to note that QCPE was
supported initially by the Air Force Office of Scientific Research and then by the National Science Foundation before
subsequently becoming a fee-for-service operation with paid employees to do the time-consuming work of testing, documenting, and distributing the contributed programs. Gezelter acknowledges the cost of maintaining scientific software and suggests alternative models to defray these costs including selling support, consulting, or an interface, all the while making the source code available for free. These suggestions strike us as naive, something akin to giving away automobiles but charging for the mechanic who services them. Such a model creates a financial incentive to release a less-than-stellar product into the public domain, then charge to make it useful and usable. It is better to release a top-of-the-line product for a nominal fee.

Is “free” software genuinely free of charge to individual researchers? Consider software developed in the U.S. national
laboratories. These ventures are supported by full-time scientific programmers employed specifically for the task, and the
cost to support and develop these products is subtracted from the pool of research funding available to the rest of the community. The individual researcher pays for these codes, in a sense, with his rejected grant proposals in times of lean funding. In contrast to using one’s own performance metrics to guide software purchases, within this system, one has no choice in what one pays for. In other words, “free software” is not free for you; the only sense in which it is “free” is that you are freed from making a choice about how to spend your research money.

Computational chemistry software must balance the needs of two audiences: users, who gauge their productivity based on the
speed, functionality, and user-friendliness of a given program; and developers, who may be more concerned with whether the
structure “under the hood” provides an environment that fosters innovation and ease of implementation. As a quantitative example, consider that the cost of supporting a postdoctoral associate (salary plus benefits) is perhaps $4,800/month. If the use of well-supported commercial software can save 2 weeks of a postdoc’s time, then this would justify an expense of ≥$2,000 to purchase a software license. This amount exceeds the cost of an academic license for many computational chemistry programs. Given the choice between a free product and a commercial one, a scientist should make a decision based on her own needs and her own criteria for doing innovative research.

What Is “Open Source”? The term “open source” is ubiquitous but its meaning is ambiguous. Some codes are “free” but are not open, whereas others make the source code available, albeit without binary executables, so that responsibility for compilation and installation is left to the user. Insofar as the use of commercial quantum chemistry software is a mainstream of modern chemical research and teaching, there exists a broad consensus that the commercial model offers the stability and assurance of support that the community desires. Strict coding guidelines can be enforced within a model where source code access is limited to qualified developers, and this kind of stability offers one counterbalance to the “reproducibility crisis.”

Occasionally the open-source model is touted on the grounds that one can use the source code to learn about the underlying algorithms, but this hardly seems relevant if the methods and algorithms are published in the scientific literature. Source code itself rarely constitutes enjoyable reading, and using source code to learn about an algorithm is a last resort forced by poorly written scientific papers. Better peer review is a more desirable solution.

A more practical use of openly available source code is to reuse parts of it in other programs, provided that the terms of the software license allow this. Often, they do not. Some ostensibly “open” chemistry codes forbid reuse, or even redistribution. Others, such as CP2k, use the restrictive General Public License that requires any derivative built on the original code to be open-source itself. Variation in design structure from one program to the next also severely hampers transferability, even if the license terms are amenable.

Access to source code allows developers to introduce their own innovations, but this is distinct from reuse. To facilitate innovation by developers, source code needs only to be available to people who intend to build upon it. This is commonly accomplished in the framework of “closed-source” software projects by granting academic groups access to the
source code for development purposes. Given the large number of developers for many quantum chemistry codes, this developer community should not be envisaged as some small, secluded cabal but rather as a rich, diverse community of academic scientists.

Let us analyze accessibility using Q-Chem as a specific example. This is commercial software whose source code is not in the public domain because that would eliminate the product that Q-Chem, Inc., is selling. Consider, however, how many in the public domain because that would eliminate the product example. This is commercial software whose source code is not

A rigid, mindless focus on an open-source mantra is a blanket requirement that software be free and open-source, more-competitive ideas are selected over less-competitive ones, intellectual merit, productivity, and impact. In the long run, the quick, most efficient, and highest-impact way. A strictly

open-source environment may furthermore disincentivize young researchers to make new code available right away, lest their ability to publish papers be short-circuited by a more senior researcher with an army of postdocs poised to take advantage of any new code. This would contribute directly to the scenario that Gezelter wishes to avoid, namely, one where students leave behind “orphaned” code that will never be incorporated into mainstream, production-level software. Viewed in these terms, an open-source mandate degrades, rather than enhances, cyberinfrastructure.

How should the impact of software be measured? Scientific publications are a more sound metric than either the price of a product or whether its source code is available in the public domain. Software is meant to serve scientific research, in the same way that any other scientific instrument is intended. As such, the question should not be whether software is free or open source, but rather, what new science can be accomplished with it? Let us not allow political rhetoric to dictate how we are to do science. Let different ideas and different models (including open source!) compete freely and flourish, and let the community focus instead on the most important metric of all: what is good for scientific discovery.

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■ REFERENCES


(13) See, for example, the Orca license. https://orcaforum.cec.mpg.de/license.html (accessed April 2015).


(17) See, for example, the GAMESS license: http://www.msg.ameslab.gov/gamess/License_Agreement.html (accessed April 2015).
