Open source bioimage informatics for cell biology

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Significant technical advances in imaging, molecular biology and genomics have fueled a revolution in cell biology, in that the molecular and structural processes of the cell are now visualized and measured routinely. Driving much of this recent development has been the advent of computational tools for the acquisition, visualization, analysis and dissemination of these datasets. These tools collectively make up a new subfield of computational biology called bioimage informatics, which is facilitated by open source approaches. We discuss why open source tools for image informatics in cell biology are needed, some of the key general attributes of what make an open source imaging application successful, and point to opportunities for further operability that should greatly accelerate future cell biology discovery.

Bioimage Informatics as a discovery tool in cell biology

Imaging is used as a tool for discovery throughout basic life science, and biomedical and clinical research. In these domains, advances in light and electron microscopy have transformed biological discovery, enabling visualization of mechanism and dynamics across scales of nanometers to millimeters and picoseconds to many days. Fluorescent protein (FP)-tagged fusions can be used as reporters of biomolecular interactions in cultured living cells [1], and the same reporter can reveal the localization and growth of a tumor in a living animal [2,3]. In short, the last 20 years have provided us with a wealth of sophisticated biological reporters and image data acquisition tools for biomedical research. Many of these imaging and instrumentation developments have been driven by partnerships between academic laboratories that invent and prototype new technology and commercial entities that develop and market them as commercial products. This development and delivery pipeline of commercial imaging instrumentation and software has been quite successful, having delivered the laser scanning confocal [4,5], spinning disc confocal [6,7], wide-field deconvolution [8,9] and multiphoton microscopes [10] that are engines of discovery in cell and developmental biology.

All of these methodologies produce complex, multidimensional data sets that must be transformed into reduced representations that scientists can manipulate, analyze, share with colleagues, and ultimately understand. Despite the diversity of applications of imaging in biology, there are common unifying challenges such as displaying a multi-gigabyte time-lapse movie on a laptop screen, or identifying, tracking, and measuring the objects in that movie and presenting the resulting measurements in a graph that reveals the mechanisms that drive their movements. These requirements have spawned the new field of bioimage informatics [11], which aims to deliver tools for data visualization, management, storage, and analysis. While still a relatively young field, bioimage informatics has already had a major impact in cell biology particularly in the area of quantitative cell imaging where advanced feature recognition, segmentation, annotation and data mining approaches are used regularly [12–20].

Almost all commercially provided image acquisition systems include software tools that provide sophisticated image visualization and analysis functions for the images recorded by the instrument they control. However, in recent years, many non-commercial projects have appeared, almost always based in research laboratories that require functionality not available in commercial products. Here, we discuss the application of bioimage informatics in cell biology and focus specifically on the development of open source solutions for bioimage informatics that have emerged over the last few years.

What are the informatics challenges in quantitative cell biology imaging?

Given the rapid development in image acquisition systems in the last 20 years, it is worth considering why a corresponding rapid development of informatics tools has occurred only recently. Certainly, one of the barriers to providing universal tools for bioimage informatics is the diversity of data structures and experimental applications that produce imaging data. In optical microscopy alone there are a substantial number of different types of imaging modalities and, indeed, a method like fluorescence microscopy encapsulates a huge and rapidly growing field of image acquisition approaches [21]. Informatics tools that support this range of methods must be capable of capturing the raw data (the individual pixels) and the metadata.
around the acquisition methodology itself, including
instrument settings, exposure details etc. This diversity
of data structures makes delivering common informatics
solutions difficult, and this complexity is multiplied by the
large number of commercial imaging systems that use
individually specified, and often proprietary, file formats
for data storage. Our current estimates are that there are
approximately 80 proprietary file formats for optical micro-
scopy alone (and not including other common imaging
techniques) that must be supported by any bioimage infor-
matics tool that aims to provide a generalizable solution.
In short, the lack of standardized access to data makes the
generation of informatics tools quite difficult.

A deeper challenge resides in each individual laboratory
that uses imaging as part of its experimental repertoire.
The sheer size of the raw data sets and the rate of pro-
duction mean that individual researchers can easily gen-
erate many tens of gigabytes of data per day. This means
that large laboratories or departmental imaging facilities
generate many hundreds of gigabytes to terabytes per
week, and are now enterprise-level data production facili-
ties. However, the expertise for developing enterprise soft-
ware tools or even simply running the hardware necessary
for this scale of data management and analysis rarely
exists in individual laboratories. In short, the sophisticated
systems and development expertise that are used to deliver
genomics databases and applications are required in indi-
vidual imaging laboratories and facilities. The delivery of
tools that provide access to a broad range of data types,
manage and analyze large sets of data, and help run the
systems that store and process this data is the challenge
that bioimage informatics seeks to address.

Why are open source approaches essential?
A critical development in the field of bioimage informatics
has been the introduction of many open source projects in
the last few years [11,22–30]. These projects range from
being open source distributions where the code is available
but new development is not specifically encouraged, to open
development projects that are community-driven projects
that actively encourage the help and participation of pro-
jects for the support and addition of new features. Therefore,
before we proceed, it is worth considering what constitutes
open source and open development efforts and why they are
valuable or even necessary for bioimage informatics.

Open source software is a well-established movement
with strong paradigms in many very successful projects such
as Linux (http://www.linuxfoundation.org/), Java (http://
java.sun.com/), MySQL (http://www.mysql.com/products/
database/), and Apache (http://www.apache.org/). A funda-
mental tenet of open source software projects is that the
copyright holder (usually the software developer or his/her
employer) determines the software license, which defines
how the software is distributed and what end-users may do
with the software. For open source software, the original
source code is made available under the terms of this license.
An open source license usually allows end-users to use the
software for any purpose, make changes to the software
source code or link their own software to it and, if they desire,
distribute those ‘derivative works. However, the software
license also defines under what terms and license derivative
works may be distributed. Table 1 gives some examples of
commonly used open source software licenses and summar-
izes their terms. For any users or developers, these details
are important and must be understood given the great
implications for development and deployment.

The ability to see and make changes to the work of
another developer is a critical component of open source
software. The attractive aspect of this approach for science
is that users and developers can directly see, evaluate, and
use another’s work (really, their intellectual property) and,
if necessary, build upon it. This is a key and often over-
looked part of open source software. Successful open source
software development projects are dynamic, evolving
enterprises allowing input, feedback, and often contribu-
tions from their community.

This evolving, adaptable aspect makes open source soft-
ware particularly useful for scientific discovery and, more
specifically, for the rapidly evolving and diverse set of
imaging applications used in biological research. Commer-
cial and closed source applications have certainly sup-
ported many significant advances in imaging. However,
an essential part of bioimaging data analysis is the ability
to easily try new methodology and approaches or even to
combine existing ones to generate a derivative result based
on the combination of two approaches. Open source
approaches make this possible. As such, there is a natural
fit between open source software and the process of scien-
tific discovery. In addition, a consequence of the growth
of the open source community is a de facto establishment of
standardized documentation methods (http://java.sun.-
com/j2se/javadoc/) and software specifications (http://java.
sun.com/products/ejb/docs.html). These specifications
ensure that developers can understand and use each
other’s code and, most importantly, that two independent
software packages can use a specified, common interface.
This software ‘interoperability’, enforced by the community
either formally or informally, is a general hallmark of open
source software, and perhaps one of its most underappre-
ciated strengths. Because standardization is so well estab-
lished in the open source community, open source software
has a critical role in providing the specifications and tools
for common file formats or common interfaces that enable
two otherwise incompatible packages to communicate
their input and output data to one another. This type of
interoperability is critical to support the rapidly evolving
needs of bioimage informatics. For all these reasons, many
of the recent developments in bioimage informatics are
based on an open source foundation.

Recently, a subclass of open source project know as ‘open
development’ has been defined (http://www.oss-watch-a-
c.uk/resources/odm.xml). Open development projects take
the open source concepts and add a significant role for the
community in the development process. In truth, com-
unity interaction and feedback was a component of most
initial open source projects, but as open source projects
have expanded, not all have included efforts to engage and
respond to their user community. Community interaction
and support is expensive, it takes precious developer time
and often requires the use of forums, mailing lists,
and other resources to manage the interactions with
the project’s community. However, open source, and open
development approaches in particular, have proven to be particularly attractive for funding agencies supporting biomedical research. They provide a way to measure the success of the project, by providing measures of uptake and participation. As the community grows around an open development project, it provides a measure of protection for the research investment and sustainability of the software past the duration of the initial award. Many agencies are now requiring that applicants have a software sharing plan in their grant application and, if an open source approach is not possible, justify this decision. In our opinion, the value for the developers, the community and the funding investment will be maximized if open development models are also followed.

Open source tools for data acquisition, visualization, analysis, and dissemination

It is beyond the scope of this article to provide a comprehensive review of all available open source tools in image informatics and features and applications. Many other papers [20,27–36] have reviewed particular applications in depth. Instead, we list representative ones (Table 2) to help illustrate the current impact and future potential of open source tools in biological imaging.

Supporting open source software

Open source software drives further innovation by allowing the free exchange of code and algorithms. Commercial applications are largely driven by market demand for a specific function or feature, so proprietary software has to be economically viable and thus must have feature limitations, code access restrictions and design parameters focused on a particular user base. Open source software complements these commercial packages and allows for new scientific ventures where a desired feature or code addition may not be commercially viable to develop.

Any open project must be viable, it must deliver valuable products to its community, and it must be sustainable and have a strategy for long-term funding. In academic science, many projects receive grant funding to initiate their work, but it is common for software development to require more than three years to achieve a fully developed product that

### Table 2. Representative survey of open source bioimage tools

<table>
<thead>
<tr>
<th>Software tool</th>
<th>Main feature set</th>
<th>License type</th>
<th>Website</th>
</tr>
</thead>
<tbody>
<tr>
<td>ImageJ</td>
<td>Image processing</td>
<td>Public domain</td>
<td><a href="http://rsweb.nih.gov/ij/">http://rsweb.nih.gov/ij/</a></td>
</tr>
<tr>
<td>OME</td>
<td>Database and image informatics</td>
<td>GPL</td>
<td><a href="http://www.openmicroscopy.org">www.openmicroscopy.org</a></td>
</tr>
<tr>
<td>Bio-Formats</td>
<td>Metadata interchange library</td>
<td>GPL</td>
<td><a href="http://www.loi.wisc.edu/ome/formats.html">www.loi.wisc.edu/ome/formats.html</a></td>
</tr>
<tr>
<td>CellProfiler</td>
<td>Automated identification of features in cells</td>
<td>GPL</td>
<td><a href="http://www.cellprofiler.org/">www.cellprofiler.org/</a></td>
</tr>
<tr>
<td>VisBio</td>
<td>Multidimensional image analysis</td>
<td>GPL</td>
<td><a href="http://www.loi.wisc.edu/visbio">www.loi.wisc.edu/visbio</a></td>
</tr>
<tr>
<td>Bisque</td>
<td>Database system for semantic analysis</td>
<td>GPL</td>
<td><a href="http://www.bioimage.ucsb.edu/">www.bioimage.ucsb.edu/</a></td>
</tr>
<tr>
<td>PSLID</td>
<td>Subcellular localization data model</td>
<td>GPL</td>
<td>murphylab.web.cmu.edu/services/PSLID/</td>
</tr>
<tr>
<td>Micromanager</td>
<td>Microscopy control</td>
<td>BSD</td>
<td><a href="http://www.micro-manager.org/">www.micro-manager.org/</a></td>
</tr>
<tr>
<td>ITK</td>
<td>Set of extensive tools for image analysis, including segmentation and registration</td>
<td>BSD</td>
<td><a href="http://www.itk.org/">www.itk.org/</a></td>
</tr>
<tr>
<td>FARSIGHT</td>
<td>Toolkit for associative image analysis</td>
<td>BSD</td>
<td><a href="http://www.farsight-toolkit.org">www.farsight-toolkit.org</a></td>
</tr>
<tr>
<td>Osirix</td>
<td>Multidimensional medical image viewer</td>
<td>GPL</td>
<td><a href="http://www.osiris-viewer.com/">www.osiris-viewer.com/</a></td>
</tr>
<tr>
<td>BioImageXD</td>
<td>Python package that leverages ITK for image processing and volume rendering</td>
<td>GPL</td>
<td><a href="http://www.bioimagexd.net/">www.bioimagexd.net/</a></td>
</tr>
</tbody>
</table>
can be distributed and used by the community. Sustaining these efforts exclusively through grants is possible, but requires convincing demonstration of the software's utility, and must accept the reality that continued funding is subject to variations in availability of funding and the priorities of funding organizations. As they mature, open source software efforts develop a non-profit foundation (e.g., Apache Software Foundation, http://www.apache.org) or a commercial arm (e.g. http://www.kitware.com and http://glencoe-software.com) that can directly access funding from user communities through licensing and customization fees that support the targeted customer base and help finance additional code development and maintenance for the open source package. However, there are still few examples of this maturation in scientific software. An important question for the scientific community is what priority funding agencies should place on the continued funding of software development tools for its use. If continued funding is to be considered, the application and reviewing processes will need to be modified to properly capture and assess the value of these projects. In our opinion, in exchange for periodic review and consideration for sustained funding, publicly funded scientific software projects should be required to follow open development models, where engagement and support for the community is required. This can occur only if funding for support and community engagement is available, and if career development and evaluation include publication record and delivery of useful tools to and engagement with the community.

In comparing open source and commercial software products, one of the biggest differences is support for the software itself. In general, commercial software packages are supported with instructions, manuals, and direct user support, and this is a key advantage of using commercial software. The cost of such support is either included in the original purchase price or paid for by purchase of a software maintenance agreement. Covering the costs of user support is difficult for open source projects because there is no corresponding fee structure to cover such support costs and, often, the academic grants that fund open source projects cover only the innovative research components and do not support the personnel or infrastructure needed. This is gradually changing with funding agencies and scientists alike realizing the importance of producing innovative and feature-rich code but ensuring that it is well supported and maintained. There are well-established standards and tools in the open source community for support, mailing lists, user forums, screencast demos, and wiki-based user documentation, that all contribute to making software successful. Within our own Open Microscopy Environment Consortium (http://openmicroscopy.org), we use project management tools such as Subversion (http://subversion.tigris.org/) to manage our source code repository, Trac (http://trac.edgewall.org/) for all project management and issue and revision tracking, Jabber (http://www.jabber.org) for real-time communication, Hudson (https://hudson.dev.java.net/) for continuous integration, Plone for managing our web site (http://plone.org/), and PHPBB for running our user forums (http://www.phpbb.com/). In addition to these tools, we hold annual user meetings to assess progress and define roadmaps for future works. We participate as presenters or exhibitors in large meetings of the community in order to capture as much feedback as possible. These tools and activities help support and engage a very broad user and developer community and are an important part of ensuring community wide adoption, but installing, running, and maintaining these tools, as well as answering queries and moderating discussions, requires time and resources (both people and money). Many successful open source packages have shown the importance of transforming the conventional user base into an additional support mechanism where the user community interacts with the original developers and with each other for support and new code developments. Users and developers that are new to the project are often supported by the community, and not just the main development team. This transformation takes some time and investment because it results from releasing useful software and investing a moderate amount of resources in support. However, we strongly advocate that direct funding of support personnel and tools be made available for research-based open source software development. In our experience, many of our academic colleagues hesitate to release their software because of the burden of supporting use of their software, thus preventing the synergies that should occur within the scientific community.

While many of these arguments are in support of open source software for scientific research in general, there are specific advantages for biological imaging. Bioimaging is inherently interdisciplinary and covers a wide range of technical and biological applications. Given this great heterogeneity of its technology and applications, bioimaging needs the open exchange of techniques and principles for further innovation. There has always been a rich tradition of this from the physical instrumentation side of bioimaging development; many current emerging imaging technologies were first developed in other fields (e.g., adaptive optics [37] was first developed in astronomy). Open source software development builds upon this collaborative instrumentation approach to allow for further innovation by sharing specific algorithms or leveraging specific code for data acquisition, visualization or data sharing.

Partnering commercial and open source efforts
Most of the imaging systems in biological laboratories are commercially developed and provided, and thus driven by commercially licensed, closed software. These powerful tools are the workhorses of modern biological research. There are many examples of companies using open source specifications to increase the functionality and value of their products, including major vendors such as Red Hat and IBM. In addition, the expertise and know-how in commercial companies is valuable, and open source projects are often aided by commercial partners working as supporters and as active developers. We therefore strongly advocate partnerships between commercial providers and open source software projects. With the appropriate licensing models, companies can be actively involved in open source imaging software for the benefit of all. Micromanager (http://www.micro-manager.org/) and the Open Microscopy Environment [20,28,38] are two prominent examples.

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of this where commercial providers have made significant contributions and played active roles in software development. Academic and commercial partnerships are vital to the long-term success and innovation in bioimage informatics, just as this same arrangement has facilitated new innovation in imaging hardware development.

Summary
The rapid innovation in imaging technology for biomolecules, cells, and tissues requires a parallel development in software tools for managing visualizing and analyzing image data. Open source software has an important role in this development, as open code development and sharing enable rapid exchange and experimentation with new tools and ideas. As open source software tools become more sophisticated, funding mechanisms that enable laboratories to provide long-term support to a broad user and developer community must be made a priority by funding bodies. The open community is very interactive and evaluates performance based on merit; i.e. good software is used by its target audience. Thus, further funding can be tied to feedback from and uptake by the community. Our experience is that academic software should follow an open development model, even if this approach deviates from the standard models used for academic research. It is important that any funded open source program be managed efficiently and integrate previous efforts and community specifications. Finally, the community must understand that a development project does ‘develop’; it grows, matures, and ultimately, if properly run and integrated with its user community, delivers useful tools. The community’s comments and feedback during this growth is critical. This process is slow and iterative, but the paradigms are well established and can be used to deliver successful tools and ultimately new discoveries.

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