Abstract—Protein crystallography produces most of the protein structures used in structure-based drug design. The process of protein structure determination is computationally intensive and error-prone because many software packages are involved. Here, we attempt to support the reproducibility of this computational work by using Jupyter notebooks to document the decisions made, the code, and selected output. We have made libraries of code templates to ease running the crystallography packages in Jupyter notebooks when editing them with JupyterLab or Colab. Our combined use of GitHub, snippet libraries, Jupyter notebooks, JupyterLab, and Colab will help modernize the computing done by structural biologists.

Index Terms—literate programming, reproducible research, scientific rigor, electronic notebooks, JupyterLab, Jupyter notebooks, computational structural biology, computational crystallography, biomolecular crystallography, protein crystallography, biomolecular structure, biomedical research, protein*drug interactions, RNA*drug interactions, molecular graphics, molecular visualization, scientific communication, molecular artwork, computational molecular biophysics

Introduction

Structural biologists study the molecular structures of proteins and nucleic acids to understand how they function in biology and medicine. The underlying premise of the field is that molecular function follows molecular form. More precise aliases for these scientists include molecular structural biologists, structural biochemists, and molecular biophysicists. Some of the methods used to determine the near-atomic resolution molecular structures include molecular modeling, X-ray crystallography, nuclear magnetic resonance (NMR), and cryo electron microscopy (cryo-EM). These scientists often use the molecular structures of these large biomolecules to design small-molecule drugs for improved therapies. As a result, structural biology plays a vital role in drug discovery and development, and many structural biologists work in the pharmaceutical industry. Those in academia in the United States generally have their work funded by the National Institutes of Health, the National Science Foundation, the Department of Defense, the Department of Energy, or one of several disease oriented medical foundations.

Structural biology is at the intersection of biochemistry, molecular biology, molecular biophysics, and computer science. Structural biologists have diverse backgrounds and varying levels of experience with computer programming ranging from minimal to very advanced. Several decades ago, the barriers to entry into the field included expertise with running command-line-driven programs and the ability to write programs to meet data analysis needs not met by existing software packages. However, these barriers have been lowered over the past two decades by the widespread availability of GUI-driven software that is often free for academics (e.g., CCP4 [Winn11], Phenix [Lieb19], CNS [Brun98], ATAS [Mana21], BioXTAS [Hopk17], CCPEM [Burn17]). As a result, biologists, who often have little formal training in computing, have become the largest component of the field.

Computing is involved in the six or more steps from structural data acquisition to publication. Several alternate software packages are often available for each step. Different combinations of these alternatives lead to a combinatorial explosion of possible workflows. In some situations, workers have set up software pipelines for some of the steps. However, these pipelines are difficult to transfer or have trouble with the challenging samples that cannot yet be handled without human intervention. The current heterogenous computing environment makes the computational work vulnerable to errors in the tracking of input and output files. Storing the code and outputs for some of the steps in Jupyter notebooks would be one way to reduce this vulnerability [Kluy16], [Gran21].

To ease crystal structure determination in Jupyter, we made libraries of code templates for crucial programs. We formatted the libraries for two extensions of JupyterLab that provide access to code snippets. One extension (jupyterlab-snippets) displays the snippets in nested pull-down menus [jLsnip]. The other extension (elyra-code-snippet-extension) uses a search box to locate the desired snippet [ELSN] (also see the blog post by Luciano Resende [Rese20]). The user can easily add new code snippets to both systems.

We also ported the libraries to Google Colaboratory or Google Colab or just Colab [Carn18], [Cola21]. Colab is an integrated development environment (IDE) for running Jupyter notebooks on the Google Cloud Platform (GPC). Colab was designed to promote the adaptation of deep learning software to new problems and facilitate collaborative computing. Colab is a free service that provides a temporary instance of a Linux operating system with access to one K80 GPU through a Jupyter notebook. Access to
TPUs is also available. The access is terminate after long periods of inactivity or a 12-hour time limit, whichever is reached first. The time limit can be extended with a small subscription fee.

Colab comes with some deep learning software and the Python scientific computing stack including SciPy[SciP20]. Colab spares the user of the maintenance of the hardware and the operating system software. Colab can also serve as a test platform for software on Linux when one’s primary computing environment is a Mac or Windows. Colab also eases collaborative work and provides a uniform computing environment for classes and workshops. The use of Colab requires that the user have a Google Drive account for storing software, Jupyter notebooks, and data files.

The user can install additional Python and other packages including structural biology software, provided the user has the required software licenses. This software installation step requires extra time at the start of the Colab session because most structure biology software packages have numerous dependencies. To accelerate this setup step, we provide notes and code snippets for the installation of this software in the Jupyter notebook that serves as the carrier of the snippet libraries on Colab. The user can install the required software in several minutes. Although the software is deleted automatically from Google Cloud at the end of session, the software can be stored on the user’s Google Drive for faster reinstallation.

Methods

We created snippet libraries for each structural biology package to support structural biology computations in Jupyter and Colab. Any particular workflow is unlikely to require all of the libraries. For example, a beginner’s workflow is unlikely to use CCTBX, a library of Python wrapped C++ routines for building molecular structure determination software. Likewise, a cryo-EM workflow will not need XDS, a package for processing X-ray diffraction images. We created a GitHub site for each library to ease the downloading of only those libraries that interest users (Table ??). This modularization of the project should ease the correction and augmentation of individual libraries as the extensions, and structural biology software packages evolve. We only provided libraries for JupyterLab because the Jupyter Project plans to phase out support for the Jupyter Notebook software. Among the several alternative extensions for code snippets in JupyterLab, we choose jupyterlab-snippets [JLsnip] and Elyra [Elyra] because these two extensions are actively maintained and have different features. We also support a snippet library for Jupyter notebooks on Google Colab as described below because Colab provides access to GPUs, which can accelerate some of the larger computational tasks.

The jupyterlab-snippets extension

The jupyterlab-snippets extension adds a snippet menu to the JupyterLab menu bar. The user accesses the snippets through a cascading pulldown menu. Each snippet resides in a separate plain text file without any formatting. This feature dramatically eases adding new snippets by users and eases keeping the snippets under version control. The snippets are stored in the Jupyter data directory (which is found by entering jupyter --path; it is in ~/Library/Jupyter/snippets on Mac OS). Each snippet library is stored in a separate subfolder, which appears on the menu bar as a part of a cascading pulldown menu (Figure 1).

We clustered snippets into categories. Each category has a cascading submenu. Clicking on a snippet name in the submenu triggers its insertion into the current cell in the notebook. The nested menu hierarchy serves well the user who is familiar with the content of the snippet libraries.

Like most other snippet extensions for Jupyter Notebook and JupyterLab, the jupyterlab-snippets extension does not support tab stops or tab triggers. These are common features of snippet libraries for most text editors and IDEs that accelerate the editing of parameter values in snippets. The tab stops are particularly valuable because they direct the user to sites that may need changes in their parameter values, and they guide the user to all of the site to ensure that none are overlooked. The overlapping of parameter values that require changing can be a major source of bugs. The tab triggers are also often mirrored, so a change at one instance of the same parameter will be propagate automatically to other identical instances of the parameter. To compensate for the lack of tab triggers, we include a second copy of the code in the same snippet but in a comment and with the tab triggers marked with curly braces and numbers (Figure 2). The user uses the code in the comment to direct their editing of the active code. The user can delete the commented out comment when they have finished editing. Separate versions of the libraries were made with commented out code. These versions are distinguished by having "plus" appended to their names.

The elyra-code-snippet extension

A menu icon labeled with <KeyPress> provides access to snippets in the elyra-code-snippet-extension system. After the icon is clicked, the snippets appear in the left margin of the JupyterLab GUI. Snippets from all libraries appear in alphabetical order. The user can scroll through the list of snippets. Hovering the mouse cursor over the snippet’s name triggers the display of a description of the snippet.

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Fig. 1: Cascading pull-down menu for the Jupyter categories of the jupyterlabpymolpysnips library.
multiple libraries are stored together in the code-snippets folder. Unlike the JupyterLab snippets extension, the snippets from directory url{~/Library/Jupyter/metadata/code-snippets} on the url{JUPYTER_DATA/metadata/code-snippets}. This is the opening of a window with a warning. The snippet into a notebook with an active Python kernel will trigger a sanity check. For example, an attempt to insert a C++ snippet into a notebook with an active Python kernel will trigger the display of the snippet in a textbox.

Each snippet is stored in a separate JSON file. Each JSON file has the snippet code plus several rows of metadata, including a list of tags and the programming language of the snippet. The latter has the snippet code plus several rows of metadata, including a list of tags and the programming language of the snippet. The latter provides a sanity check. For example, an attempt to insert a C++ snippet into a notebook with an active Python kernel will trigger the opening of a window with a warning.

All of the snippets reside in the folder url{JUPYTER_DATA/metadata/code-snippets}. This is the directory url{~/Library/Jupyter/metadata/code-snippets} on the Mac. There are no subfolders for individual snippet libraries, unlike the jupyterlab-snippets extension. The snippets from multiple libraries are stored together in the code-snippets folder.

Fig. 2: Comparison of active code in the bottom code block and the commented out code above the active code from a code snippet. The commented lines of code serve as guides for editing because they have curly braces marking sites to be considered for editing. The commented lines of code compensate for the absence of tab stops.

Alternatively, the user can enter a search term in the search box at the top of the menu to reduce the list of snippets. The search terms can be part of a snippet name or a tag stored with each snippet.

A tag icon displays all of the available tags in the snippets as separate icons. The user can select tags to be used to choose snippets by clicking on the icons.

Each snippet is displayed with several icons (Figure 3). A triangular toggle can trigger the display of the snippet in a textbox. A pencil icon enables the editing of the code. Other icons enable copying the code to the clipboard, inserting code into the current cell in the notebook, and deleting the snippet.

A plus sign in the upper-right hand corner opens a GUI for the creation of a new snippet. The GUI occupies a new tab in the window that houses the Jupyter notebooks. The GUI has a text box for each kind of metadata: name, description, tags, language, and the snippet code. There is a save button at the bottom to add the new snippet to the current library.

Each snippet is stored in a separate JSON file. Each JSON file has the snippet code plus several rows of metadata, including a list of tags and the programming language of the snippet. The latter provides a sanity check. For example, an attempt to insert a C++ snippet into a notebook with an active Python kernel will trigger the opening of a window with a warning.

The tag system can be used to select all snippets from one library. The tag system serves well the user who is not familiar with the content of the installed libraries. The user can download the snippets from GitHub as zip file and then uncompress this file and move the snippet files to the final destination.

Colab snippet library

The Colab snippet system resembles the Elyra snippet system in that the snippets appear in a menu to the left of the notebook and that search terms in a search box retrieve snippets. However, the Colab system differs from the Elyra system ins that the snippets are stored in one or more Jupyter notebooks. The user’s Google Drive stores the notebook of snippets. The user enters the url for the notebook in a the Tools --> Settings --> Site --> Custom Snippet Notebook URL. Multiple URLs for multiple notebooks can be entered at one time. The user logs out of Colab and upon logging in again to install the snippets. The user will see the newly added snippets in the left margin after opening the snippet menu by clicking on the </> icon.

Each snippet had a markdown cell followed by a code cell. The markdown cell contained the name of the snippet, a description of what the snippet does, and the structural biology software. These features are searched in the search box to narrow the list of snippets to inspect for selection.

The first snippet in each notebook provided the steps for installing the software on Colab. The markdown cell listed these installation steps. Then a series of code snippets contained the code for carrying out the steps. This installation snippet was the only one in a notebook that contained more then one code snippet.

A search box at the top of the list of snippets is used to recover a snippet (Figure 5). The user enters a snippet name in the search box to display the snippet and its documentation. The user hits the ‘Install’ button to install the snippet’s code at the current position in the notebook. Unlike the Elyra snippets which insert a whole snippet into one code block, a Colab snippet can have multiple code blocks that are inserted into the notebook at the current position of the mouse cursor. One snippet can have different types of code blocks. For example, the snippet in Figure 5 has a three blocks of Python code, two blocks of shell commands, and two blocks of bash cell magics with multiple lines of bash commands.

The list snippet for a library will print in a table below the current cell a list of the snippets in the library and a brief description. This table is stored in a pandas DataFrame that can be searched with the pandas search function. This table can also be searched for key terms with the search function in the notebook. The code block and output can be hidden by clicking on the three blue dots on the left margin of the cell.

Notebooks on Colab open very quickly, but the user must reinstall their software on each login. We ease this annoying task by supplying the complete chain of installation steps. For example, the installation of the molecular graphics program PyMOL requires seven code blocks of different types. Some involve the use of curl, and others use the conda package management system. We include all steps in one snippet, which is uniquely possible with the snippet system for Colab (Figure 5). The user only has to select one snippet and then run each code block in succession.

The use of Colab requires that the user has a Google account and a Google Drive. Many structural biologists already have both.
Fig. 3: The GUI from the elyra-code-snippet extension for accessing code snippets is shown on the left. A preview of the hbonddash snippet is shown in the lower left. A Jupyter notebook with the inserted the hbonddash snippet is shown on the right.

Notebooks with sample workflows
We created a library of Jupyter Notebooks with sample workflows. This library of notebooks is only representative and not exhaustive because the combinatorial explosion of possible workflows makes covering all workflows impractical. These notebooks can serve as templates for the creation of new notebooks and are available on our GitHub site [MLGH].

Availability of the snippet libraries
We have shared these libraries on GitHub [MLGH]. Each library is also archived in zenodo.

Results
We describe here a set of libraries of code templates to support computational crystallography in Jupyter Notebooks on the cloud and on local computers. The libraries and notebooks can be loaded on and run on Google Colab where the user can share the notebook with collaborators or gain access to GPUs and TPUs. The user uploads the libraries and notebook to their Google Drive account and accesses the notebook from Colab. The storage of the libraries and notebooks on Google Drive persists between logins to Google Colab, but the crystallographic software must be reinstalled on each use of Colab. These libraries are installed only once; however, the crystallographic software must be reinstalled upon each login. We describe below installation scripts in the form of snippets that can be quickly run at the top of a Notebook to minimize the effort required to re-install the software. Another limitation of the Colab snippet system is that snippets from all libraries are stored in one pool and have to be accessed by either scrolling through a long list or by entering the snippet name in a search box. We addressed this limitation with a snippet for each library that prints a list of the available snippets with a short description. This list can span more than the length of a paper, but it can be collapsed to hide it or can be deleted when no longer needed. After the snippet is pulled out of the list by the search box, more detailed documentation about the snippet is displayed. Next, we describe the content of each library with representative output in the Colab notebook.

Structure determination and refinement workflows with Phenix
A team of professional software developers based at the Berkeley-Lawrence National Laboratory (BLNL) develops the Phenix software to refine protein crystal structures determined from X-ray diffraction data [Adam02]. The project includes several collaborators located around the world who develop auxiliary components of the package. Phenix uses Python to interface with the Computational Crystallography Tool Box (CCTBX), which is written in C++ for speed [Gros02]. CCTBX is also wrapped in Python and can be imported at as module. While Python eases the use of CCTBX, mastery of CCTBX requires at least an intermediate level of Python programming skills. On the other hand, Phenix is easy to use via the command line or a GUI and has become of the most popular software packages for biological crystallography.

The Phenix project greatly eased the incorporation of simulated annealing into crystal structure refinement by hiding the
Simulated annealing involves molecular dynamics (MD) simulation at high temperatures to move parts of a molecular model out of local energy minima and into conformations that fit the experimental data better. Twenty minutes of applying simulated annealing to an early model that still has numerous errors can significantly improve the model while saving the user a day or more of the tedious manual rebuilding of the molecular model. The PDB file does not have sufficient information about chemical bonding for MD simulations. The molecular dynamics software that carries out the simulated annealing requires two parameter files and the coordinate file. The preparation and debugging of the parameter files manually takes many hours, but Phenix automates this task.

More recently, Phenix has been extended to refine crystal structures with neutron diffraction data and for structure determination and refinement with cryo-EM data [Lieb19]. The addition of support for cryo-EM help address the recent need for the ability to fit atomic models to cryo-EM maps that have recently become available at near atomic resolution because of the dramatic improvements in detector technology. Users can interact with Phenix via a GUI interface or the command line, as mentioned before, but users can also use PHIL, domain-specific language scripting language for more precise parameter settings for Phenix. In addition, users can use the phenix.python interpreter. Unfortunately, the phenix.python interpreter is still limited to Python2, whereas CCTBX has been available for Python3 for over a year.
Fig. 5: Code snippet for installing PyMOL on Colab. The <> icon opens a menu on the left side that lists all of the snippets. The search term ‘pymol’ was used to reduce the list of candidate snippets. The highlighted snippets name ‘Install PyMOL is new Colab notebook’. Selecting this snippets opens the snippet below. The snippet description is displayed followed by the seven blocks of code. The description includes the seven steps for installing the molecular graphics programs. Clicking with the mouse cursor on ‘INSERT’ in blue inserts the code into the cells on the notebook on the right.

Jupyter Lab and its extensions are also best run with Python3. The most practical approach to using Phenix in Jupyter Lab is to invoke Phenix by utilizing the shell rather than using Python. For example, the command shown below invokes statistical analysis of the B-factors in a Protein Data Bank (PDB) file by using one line of code in the shell. The PDB file uses a legacy, fixed-format file for storing the atomic coordinates and B-factors of crystal structures. The B-factors are a measure of the atomic motion, statistical disorder, or both in individual atoms in a protein structure. The PDB file format was defined and popularized by the Protein Data Bank, a repository for atomic coordinates and structural data that has over 170,000 entries from around the world. The PDB was started in 1972 and unified with the branches in Japan and Europe in 2003 as the wwPDB [Berm03]. The wwPDB continues to play a central role in promoting the principles of open science and reproducible research in structural biology.

Since 2019, the wwPDB requires the PDBx/mmCIF format for new depositions [Adam19]. Many structural biology software packages now have the ability to read files in the PDBx/mmCIF format.

!phenix.b_factor_statistics 1lw9.pdb

The output form this command is printed below the cell that invokes the command. Some of the output is shown below.

Starting phenix.b_factor_statistics
on Wed Jun 2 04:49:01 2021 by blaine

Processing files:

Found model, /Users/blaine/pdbFiles/1lw9.pdb

Processing PHIL parameters:

No PHIL parameters found

Final processed PHIL parameters:

    data_manager {
        model {
            file = "~/Users/blaine/pdbFiles/1lw9.pdb"
        }
    }
    default_model = "~/Users/blaine/pdbFiles/1lw9.pdb"

Starting job

Validating inputs

    min    max    mean   <Bi,j>   iso   aniso
Overall:  6.04 100.00  24.07    N/A  1542    0
Protein:  6.04 100.00  23.12    N/A  1328    0
Water:   9.98  55.93  30.47    N/A  203     0
Other:   14.11  54.47  21.10    N/A   11     0
Chain A:  6.04  100.00  24.07    N/A  1542    0

Histogram:

    Values   Number of atoms
   6.04 - 15.44    309
  15.44 - 24.83    858
  24.83 - 34.23    187
  34.23 - 43.62     78
  43.62 - 53.02     32
  53.02 - 62.42     16
  62.42 - 71.81      8
  71.81 - 81.21     46
  81.21 - 90.60     10
  90.60 - 100.00    46

Job complete

usr+sys time: 1.92 seconds
wall clock time: 2.93 seconds
There are several dozen commands that can be run via the shell and return useful output that can be captured in one Jupyter Notebook rather than in dozens of log files. The output can be copied and pasted into a new cell and then reformatted in markdown as a table or the copied output be used as input data to make a plot with matplotlib. While these are basic data science tasks, they are intimidating to new users of Jupyter and some of the details are easy for more experienced users to forget. To overcome this problem, we supply snippets that demonstrate how to transform the output and that can be used as templates by the users.

These commands are becoming harder to find as the online documentation has been migrating to serving only the GUI interface. The bash script files that run the Phenix commands can be found on Mac OSX by running the following command:

```bash
!ls /Applications/phenix-*/build/bin/phenix.*
```

These shell scripts invoke Python scripts that capture the command line arguments and pass them to the Phenix Python interpreter. This Python script files can be found on Mac OSX by running the following command:

```bash
!ls /Applications/phenix-1.19.2-4158/modules/phenix/phenix/command_line/*.py.
```

### Molecular graphics with PyMOL

The end result of the crystal structure refinement in Phenix is a set of atomic coordinates. They can be displayed in one of the many available molecular graphics programs like PyMOL [PyMO21]. If PyMOL is available in the current Python environment, PyMOL’s Python API can be accessed by importing the `cmd` class. In addition, it is useful to import the `Image` class from IPython to be able to upload images written to disk by PyMOL.

```python
from pymol import cmd
from IPython.display import Image
```

After installing PyMOL in Colab as outlines in Figure 5 and the PyMOL snippet library, the `T4L` snippet was inserted into a Colab notebook and executed. The snippet includes the IPython command that was used to upload the image into the Notebook as shown in Figure 6.

![Fig. 6: The code of the T4L snippet inserted into a code block in Colab.](image)

There are several other methods of importing images including using Markdown or HTML code.

### Discussion

Amazon introduced the first cloud computing service in 2006; there are now over 200 services. These services have the advantage of providing access to computer hardware and software. These services can lower barriers for those labs that have limited access to computer hardware or that have trouble with installing software. Many of these services supply disk spaces and access to CPUs, GPUs, and sometimes TPUs. Access to basic services is often free with additional services, computing power, and disk space being available for a modest fee. In principle, consumer computers could be used as an interface for doing all manner of crystallographic computing on the cloud.

### Why Colab?

Colab was developed internally and first released for public use in 2018. Numerous research papers in the physical and medical sciences have been published that used Colab. Google Colab provides fast and easy access for users with a Google account and Google drive, so many workers in crystallography already have the prerequisites. Many readers are also familiar with Jupyter Notebooks (10 million shared on GitHub as of early 2021). Jupyter Notebooks can be loaded onto Google Drive and then opened in Colab. Colab is a specialized IDE for editing Jupyter Notebooks. the Colab interfaces has more features that the easy-to-use nteract IDE, but fewer features than JupyterLab. Colab provides almost instant loading of specific Jupyter notebooks but at the cost of needing reinstall the software used in a notebook upon logging in again or after a 12-hour session. The first point lower the barrier to resuming work while the second point can be addressed by including the code for installing the required software at the head of the notebook.

Microsoft has stopped supporting its Azure Notebook and has asked users to migrate to several alternative approaches. One approach is to use of Visual Studio Code (VSC) rather to JupyterLab to edit and run Jupyter notebooks locally and on Microsoft’s cloud service. VSC is an advanced text editor that has stronger support for code snippets because it supports the use of tab triggers and tab stops, two important features that are missing from Colab, JupyterLab, and the Classic Jupyter Notebook. However, VSC is so feature-rich that it can be overwhelming for some beginning users. To support a wider group of users, we developed the libraries for Google Colab. We plan to develop libraries for editing Jupyter Notebooks in VSC.

### What is new

We report a set of code template libraries for doing biomolecular crystallographic computing on Colab. These template libraries only need to be installed once because they persist between

```python
from pymol import cmd
from IPython.display import Image
```

```python
!ls /Applications/phenix-*/build/bin/phenix.*
```
More software packages and even programming languages. More libraries. That is the development of notebooks that use two or more libraries supported by the snippet names used to invoke magic are under autocompletion. This system would also load the snippet’s code into the code cell. This system would also provide a converter from RasMol to PyMOL. Language converters for translating code between the leading molecular graphics programs would allow users to more easily find and use the optimal molecular graphics program for the task at hand.

We also provided snippets for PyMOL, which has 100,000 users, for use in text editors [Moo21a] and Jupyter notebooks [Moo21b]. The former support tab triggers and tab stops; the latter does not.

The libraries have to be molecular graphics program specific because molecular graphics programs have been written in a range of programming languages. The user issues the commands in either in a general programming language like Python or a domain specific language (DSL) like pml. It would cause confusion to mix snippets from multiple languages. To counter this growing tower of babel, the OpenStructure initiative was formed [Bias13].

We have also worked out how to deploy this snippet libraries in OnDemand notebooks at High-Performance Computing centers. These notebooks resemble Colab notebooks in that JupyterLab extensions cannot be installed. However, they do not have any alternate support for accessing snippets from menus in the GUI. Instead, we had to create IPython magic for each snippet that load the snippet’s code into the code cell. This system would also work on Colab and may be preferred by expert users because the snippet names used to invoke magic are under autocompletion. That is, the user enters the start of a name and IPython suggests the remainder of the name in a pop-up menu. We offer a variant library that inserts a commented out copy of the code that has been annotated with the sites that are to be edited by the user.

Opportunities for Interoperability

The set of template libraries can encourage synergistic interoperability between software packages supported by the snippet libraries. That is the development of notebooks that use two or more software packages and even programming languages. More general and well-known examples of interoperability include the Cython packages in Python that enable the running of C++ code inside Python, the reticulate package that enables the running of Python code in R, and the PyCall package in Julia that enables the running of the Python packages in Julia. The latter package is widely used to run matplotlib in Julia. Interoperability already occurs between the CCP4, clipper, and CCTBX projects and to a limited extent between CCTBX and PyMOL, but interoperability could be more widespread if the walls around the software silos were lowered. The snippet libraries provided here can prompt interoperability on Colab by their proximity on Colab.

Polyglot snippets

The unique feature of the Colab snippets is that a given snippet can contain multiple cells. The cells can be a mix of markdown (text cells) and code cells. The cells can also use a mix of programming languages invoked by different cell magics. Cell magics are an alternate method to kernels for switching between programming languages. The code for defining various cell magics are included in our snippet library. The supported compiled programming languages include C, C++, Julia, and Fortran.

The bash cell magic is built into Colab. This ability to two or more programming languages in one snippet leads to polyglot snippets. Some operations involving two or more programming languages need to be executed sequentially. These can be best grouped together in one snippet. This feature of polyglot snippets save time because the user does not have to reinvent the workflow by finding and inserting into the notebook a series of snippets.

Ubiquitous computing platform on the cloud

Colab provides the user with a ubiquitous instance of Ubuntu. Colab is accessed by opening Jupyter Notebooks stored on the users’ Google Drive account. Colab can be accessed from devices that can access the Google Drive account. The opening of the Colab instance is rapid in contrast to the Binder service where the building of a new Ubuntu instance requires a wait of many minutes. In addition, the Colab session remains active for up to 12 hours on the free plan and longer on paid plans whereas a Binder instance closes after ten minutes of inactivity. Binder is an open-source project while Colab is a closed source project. Colab maintains the Ubuntu operating system so the user does not need to spend time on software updates.

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