BioProv - A provenance library for bioinformatics workflows

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Summary

In an era where it can be argued that all biology is computational biology (Markowetz, 2017), properly managing computational analyses and data is crucial to evaluate the findings of \textit{in silico} experiments. A major aspect of best practices in scientific computing is managing the provenance of data analysis workflows (Irving et al., 2021; Pasquier et al., 2017; Wilson et al., 2017). The World Wide Web Consortium (W3C) Provenance Working Group defines provenance as “a record that describes the people, institutions, entities, and activities involved in producing, influencing, or delivering a piece of data or a thing” (Moreau & Groth, 2013).

Therefore, for bioinformatics workflows (BWFs), where there are usually numerous steps in data processing, capturing and storing provenance rapidly becomes a challenge. This provenance data should not only be comprehensible to humans, but structured and queryable; this is to support reproducibility in present and future research in bioinformatics and many other fields of scientific research (Kanwal et al., 2017; Pasquier et al., 2017). A proposed standard for interoperability of provenance data is the W3C-PROV data model, specifically designed to share provenance data across the web and among diverse applications and systems. Adding provenance to BWFs can be costly to both developers, responsible for storing information about these workflows, and researchers designing and analyzing workflow results. While some workflow systems already provide provenance capturing, modelling bioinformatics data in compliance with W3C-PROV is not automatic, requiring great effort from workflow developers and researchers. We introduce BioProv as a library that aims to facilitate the creation of W3C-PROV compliant documents for BWFs, automatically capturing the provenance of workflow steps between different users and computing environments.

W3C-PROV

The W3C-PROV recommendation is endorsed by the World Wide Web Consortium (W3C), the leading global community for web standards. It divides provenance data into three separate views (Figure 1): the data flow view, comprised of entities, that are any physical, digital or conceptual thing; the process flow view, that focuses on activities, that are processes that happen over time and act upon or with entities, either by consuming, processing, using, or generating them; and the responsibility view, that concerns the assignment of agents that are responsible for entities, activities, or other agents. These three elements of provenance have a set of seven relations between them, that can be further described by relation patterns. For a full introduction to the W3C-PROV standard, we recommend Moreau & Groth (2013).

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Statement of need

BioProv is a Python library for generating provenance documents of bioinformatics workflows. The challenge of provenance capture in the field of bioinformatics has been characterized and is standing for more than a decade (Stevens et al., 2007). Presently, there are many freely available tools for managing provenance through workflow systems (Afgan et al., 2018; Di Tommaso et al., 2017; Hull et al., 2006; Köster & Rahmann, 2012; Oliveira et al., 2012; Vivian et al., 2017) and provenance systems for capturing and storing provenance data from workflow scripts (Khan et al., 2019; Silva et al., 2018). Several studies have been able to implement solutions that model BWFs and adequately capture and store provenance data (K. Ocaña et al., 2014; K. A. C. S. Ocaña et al., 2015; Paula et al., 2013). However, to the best of our knowledge, there is not yet any software library that specializes in capturing the provenance of BWFs. In the case of workflow management systems, they provide execution reports such as execution trace or graph, but these documents are not W3C-PROV compliant and/or are not serializable, or the collection of domain-specific information must be manually designed by the user with an ad hoc approach. Domain-specific data are particularly relevant in BWFs, as they can be used to help researchers make decisions and steer workflow parameters during runtime (Costa et al., 2013). They refer to metadata that are characteristic of biological data formats, e.g. the distribution of the length of sequences in a nucleotide sequence file, or the number of nodes in a phylogenetic tree file. Implementing a system to capture these data can be very costly to both users and developers of BWFs, as most provenance capture software are generic and do not support, for example, parsing of biological data formats. This may imply the need to either manually develop specific parsing solutions for the files involved or to create database schemas that support domain-specific data. BioProv attempts to fill this gap, by providing features that support the specificities of bioinformatics applications and allow the automatic capture of provenance data and generation of documents in a W3C-PROV compliant format.
Features and data modelling

Overview

BioProv represents the provenance elements of a BWF into a class called Project, that represents the execution of a bioinformatics workflow (a sequence of programs) using a particular dataset. An instance of Project is composed by related samples, files and programs that are represented by corresponding classes from the BioProv library. The Project class has specific methods that allow for the user to specify the relationships between objects, such as a file, a biological sample or a computer program. Projects also carry information about agents, i.e. users and computing environments used to execute programs. In the context of BioProv, a “Project” is distinct from a “Workflow” in the sense that a Project refers to a particular set of samples and files and associated programs, while a Workflow refers to a sequence of programs that can be run on a set of adequate input files. A user can therefore use the same Workflow for multiple Projects. Because they are serializable in JSON and tabular formats, BioProv objects can be stored and shared across computing environments, and can be exported as W3C-PROV compliant documents, allowing better integration with web-based systems. The library can be used interactively, in an environment such as Jupyter (Ragan-Kelley et al., 2014), or from the application’s command line interface (CLI). The CLI component of BioProv allows users to quickly launch custom workflows from the command line using the bioprov <workflow_name> command.

Figure 2: Architecture of a BioProv application.

BioProv is built on top of the BioPython (Cock et al., 2009) library, which has the capacity to parse common bioinformatics file formats, and supports several file formats for both sequence and alignment data, allowing the user to easily extract domain data without having to write any parsers. Here we present some of the core features of BioProv, but for a more complete
introduction, we recommend the package's tutorials in Jupyter Notebook format, that can also be launched via Binder, and the documentation page. As example data, we provide five small bacterial genomes and a BLAST database that is a subset of MEGARES (Lakin et al., 2017). These two datasets can be used to demonstrate the installation and to illustrate some of the core features of BioProv.

Classes

BioProv implements several classes in order to represent provenance data extracted from BWfs. Its object-oriented design allows for users to benefit from the flexibility of working with extensible Python objects, that are familiar to frameworks such as the libraries in the scientific Python stack: NumPy (Harris et al., 2020), SciPy (Virtanen et al., 2020), Matplotlib (Hunter, 2007), and others. The five main classes are:

- **Project**: The higher-level structure that represents core project information, like samples, files, and programs.
- **Sample**: Describes biological samples. Contains collections of files and programs, and can group any sample attributes, such as collection date, collection site, type of sample (soil, water, tissue, etc.).
- **File**: Describes computer files that may be associated with a Sample or Project.
- **Program**: Describes programs that process and create files. Instances of Program are associated to instances of the Parameter and Run classes (the latter representing a program execution).
- **Environment**: Describes an environment that was used to run a program, including environment variables and library versions.

A Project instance is the top-level object in the BioProv library. It is composed by Samples, Files, and Programs. A Sample instance represents any biological sample, its attributes, and it is composed by Files and Programs that are associated with that particular sample. Files and Programs are associated with the Project when they contain or process information from multiple samples. In the provenance data model, instances of the Project, Sample, and File classes are represented as entities, and instances of Program are represented as activities. Instances of Environment are represented as agents that act on behalf of the current user.

BioProv detects the current user and environment variables and stores them alongside the Project; each Program, when executed, is automatically associated with the current computing environment (Figure 3). This way, BioProv can represent which execution is associated with each user and environment, allowing for traceable collaborative work.

These five classes constitute the basis of a BioProv project. The library captures provenance data through instances of these classes. For Samples, it stores the sample's attributes, a collection of files, and a collection of programs. For both Files and Programs, relevant information is automatically captured, such as the start and end time of each program execution and file size of each file. Files containing biological sequences that are supported by BioPython can be parsed with the SeqFile class. This class inherits from File and can extract metadata about the file contents, such as number of sequences, number of base pairs, GC content (if it is a nucleotide file), and other metrics. This feature allows users to extract domain data for their provenance documents by using parsers available in BioPython.

Programs in BioProv can be created manually or loaded as a preset. BioProv offers a few preset programs for common bioinformatics tasks, such as sequence alignment search, multiple sequence alignment, gene prediction and quantification of gene expression. Running a program with BioProv instead of directly from the command-line automatically captures provenance information for that execution. Some of the included programs are:
• BLAST+: sequence alignment search (Camacho et al., 2009)
• Diamond: sequence alignment search (Buchfink et al., 2014)
• MAFFT: multiple sequence alignment (Katoh et al., 2005)
• Muscle: multiple sequence alignment (Edgar, 2004)
• Prodigal: prokaryotic gene prediction (Hyatt et al., 2010)

Figure 3: Diagram showing relationships between classes in the BioProv library. Each class is related to a PROV element, depicted above the class name. The color scheme is the same as in Figure 1 (entities in yellow, activities in blue, agents in orange).

Users can create their own presets with either the Program class or the PresetProgram class, that inherits from Program and possesses additional methods for batch execution. To manually create programs (that are not presets), the user should create the program as it is called from the command line, and add Parameters to it. A Parameter is another BioProv class that represents specific parameters associated with a program. Presets contain parameters specific to that program. Parameters can be added to the command string that will be evaluated on the system’s shell, by means of Python’s subprocess module. For a more complete walkthrough of how to build programs and add them to the workflow, please refer to the tutorials. Additionally, workflow presets can be created. Workflow presets are a sequence of programs which are to be executed on a project’s files. These presets can then be run using the library’s CLI, as their command-line arguments and parser are constructed automatically.

Workflow execution with BioProv

To set up a workflow with BioProv to capture provenance data, users must either write a Python script and replace usual program calls with BioProv code, or launch custom or preexisting preset workflows from the CLI. Again, the tutorials and documentation page are the best

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resources on how to do this. Once this is done, data must be imported as a BioProv project so the workflow can be executed. In the following section, we provide a brief demonstration of how to import data and run programs.

**Importing data**

There are a few ways to import and export data with BioProv. If a project has not been previously imported, the most convenient way to import it is by generating a table containing one sample per row, and columns with the path to each file associated with that sample. Columns that are not files will be processed as sample attributes. For example, assume the following table:

| sample-id | assembly     | report     | source  |
|-----------|--------------|------------|---------|
| sample_1  | contigs_1.fasta | report_1.txt | seawater |
| sample_2  | contigs_2.fasta | report_2.txt | soil     |

The `sample-id` column is our index, i.e., each sample is identified by it. The `assembly` column contains the path to the genome assembly of each sample (therefore, a “sequence file”). The `report` column points to a plain text file containing the assembly report (therefore, a “file”). The other columns will be parsed as sample attributes. This can be easily done with the `read_csv()` function:

In [1]: import bioprov as bp

In [2]: project = bp.read_csv("myTable.csv",
   file_cols="report",
   sequencefile_cols="assembly",
   tag="myProject",
   import_data=True)

The table from which the data was sourced is automatically added as a Project file:

In [3]: project.files
Out[3]: {'project_csv': /home/user/myProject/myTable.csv}

And Samples are created with associated files and attributes:

In [4]: project["sample_1"]
Out[4]: Sample sample_1 with 2 file(s).

In [5]: project["sample_1"].files
Out[5]:
{ 'report': /home/user/myProject/report_1.txt,
  'assembly': /home/user/myProject/contigs_1.fasta}

In [6]: project["sample_1"].attributes
Out[6]: { 'source': 'seawater'}

Sequence metadata is extracted from sequence files, as set by the `import_data=True` parameter:
To run a program using BioProv, it must be added to the project or to a sample. We can use the preset prodigal program to illustrate this. Prodigal runs a gene prediction algorithm for prokaryotic genomes, and creates three output files:

Now that the project is loaded, the user can add new files, samples and programs. Programs can run and execution provenance is captured (such as stdout and stderr, start and end time, and files involved).

To export the project, there are a few options. The user can either:
- export the project in a tabular format such as the one presented in myTable.csv. This omits project information (about programs, for example), but preserves information of samples and files. This action is done with the Project.to_csv() method;
- export the project as JSON, the preferred option, as BioProv can deserialize this format back into a Project object with all related information; This is done with the Project.to_json() method;
- store the project in BioProv's database. BioProv has a builtin document-oriented database, as explained in the following section.

To query an existing project, it must be loaded with the load_project() function, and it can then be queried with Python syntax. The Project's .samples, .files and .programs

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attributes are Python dictionaries, and can be manipulated as so. A Sample instance will also contain .files and .programs, and additionally a .attributes dictionary with additional information about the sample (e.g. the source column in our data table). When iterating over a Project object, we can loop over each sample:

```
In [4]: for sample in project:
   ....:    # loop over samples
   ....:    source = sample.attributes['source']
   ....:    print(sample.name, 'is a', source, 'sample.')
   ....:    # make conditional statements
   ....:    if 'prodigal' in sample.program.keys():
   ....:       print(sample.name,
   ....:               'contains an instance of the Prodigal program.')
```

sample_1 is a seawater sample.
sample_2 is a soil sample.
sample_1 contains an instance of the Prodigal program.

Being able to query BioProv objects with Python syntax allows users familiar with the language to create custom queries and functions, and integrate BioProv into their existing Python applications. Additionally, it is possible to use a tool such as the ObjectPath library, that implements a query language for semi-structured data, and is compatible with BioProv’s JSON outputs.

**Command line application and workflows**

To use the CLI, after installing, type bioprov in the system’s Shell:

```
~/$ bioprov
usage: bioprov [-h]
       [-v | -l]
       {genome_annotation,blastn,kaiju} ...
```

BioProv command-line application. Choose a command to begin.

optional arguments:
- `-h`, `--help` show this help message and exit
- `--show_config` Show location of config file.
- `--show_provstore` Show location of ProvStore credentials file.
- `--create_provstore` Create ProvStore credentials file.
- `--show_db` Show location of database file.
- `--clear_db` Clears all records in database.
- `--version` Show BioProv version
- `--list` List Projects in the BioProv database.

workflows:
- `{genome_annotation,blastn,kaiju}`

The bioprov --show_db and bioprov --list commands, for example, can be used to show the location of BioProv’s database and list all projects it contains. The workflows are preset sequences of programs that can be run directly from the CLI. They are implemented...
with BioProv's Workflow class and handle the creation of the BioProv project while running the desired workflow. Users can write their own workflow and the command-line parser will be automatically generated based on the parameters set by the user. For more information, please refer to BioProv's workflows subpackage, where preset workflows are stored, or run the bioprov <workflow_name> command in the CLI for help about a particular workflow. Presently, BioProv workflows are limited, serving mainly as a reference for users to write their own workflows; one of the main future goals of the library is to support a range of common BWFs. We actively encourage users to contribute their workflows by following the contributing guidelines or request one by opening an issue in the repository.

Provenance documents

To create W3C-PROV documents, BioProv utilizes the PROV library (Dong, 2020), and models its native objects onto the three PROV elements: entities, agents, and activities (Moreau & Groth (2013) provide a detailed description of PROV concepts). To do this, a BioProvDocument object is created from project. A W3C-PROV compatible document is then created, with associated relationships between PROV elements for each object of the project. BioProv defines one “bundle” for the project, and one for each sample and user. Bundles are documents nested within the top-level BioProvDocument, used to describe individual entities (for the Sample bundles) or agents (for the user bundles). Computing environments are also regarded as agents, that act on behalf of users through activities (that correspond to BioProv's Program instances). By leveraging the PROV library, the resulting document can be exported in a number of ways, such as graphical format and PROV-N (a human-readable provenance format).

The following code generates the Figure 4 and a PROV-N record.

In [5]: prov = bp.BioProvDocument(project)
In [6]: prov.write_provn()
In [7]: prov.dot.write_pdf("myProject.pdf")

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Figure 4: Provenance graph created by BioProv with the PROV and PyDot libraries. This graph represents a Project containing two samples associated with bacterial genomes. The prodigal program uses the assembly file as input to create the proteins, genes, and scores files. The square shapes grouping sets of icons correspond to each bundle in the project.

Lastly, **BioProvDocuments** can be uploaded to ProvStore, a web service for storage and visualization of W3C-PROV documents. The credentials to the ProvStore API are set with the bioprov --create_provstore command, and a document can be uploaded with the upload_to_provstore() method:

In [8]: prov.upload_to_provstore()

This feature enables the immediate publication of provenance documents that have a uniform resource identifier (URI) through their ProvStore address. Future developments for further supporting the W3C-PROV standard in the field of bioinformatics should focus on integrating more web services, such as NCBI Entrez and Wikidata, for continuous improvement of the reproducibility of BWFs.

**Conclusion**

We present a software library that allows users to create and run bioinformatics workflows while capturing provenance data during runtime, that can then be exported as W3C-PROV compliant documents. Provenance data can be imported and exported in widely-used and web-compatible formats, and this provenance is enriched with domain-specific data without much effort from the user. BioProv is written entirely in Python, benefitting from numerous existing libraries that complement it, and its objects can be extended and customized to suit users’ needs. This results in a simple, lightweight yet powerful library that can make Python workflows provenance-aware, setting BioProv as a valuable addition to the ecosystem of bioinformatics workflow tools.

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