Towards Reusability of Computational Experiments
Capturing and Sharing Research Objects from Knowledge Discovery Processes

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Abstract: Calls for more reproducible research by sharing code and data are released in a large number of fields from biomedical science to signal processing. At the same time, the urge to solve data analysis bottlenecks in the biomedical field generates the need for more interactive data analytics solutions. These interactive solutions are oriented towards wet lab users whereas bioinformaticians favor custom analysis tools. In this position paper we elaborate on why Reproducible Research, by presenting code and data sharing as a gold standard for reproducibility misses important challenges in data analytics. We suggest new ways to design interactive tools embedding constraints of reusability with data exploration. Finally, we seek to integrate our solution with Research Objects as they are expected to bring promising advances in reusability and partial reproducibility of computational work.

1 INTRODUCTION

Over the last few years, calls from researchers defending better data and code sharing for computational experiments (CE) are propagated in high-ranked journals (McNutt, 2014; Peng, 2011). Usually grouped under Reproducible Research (RR), these invitations elevate reproducibility or replicability as a central key of the scientific method. One of the interpretations presents reproduction as an application, by independent researchers, of identical methods on identical data to obtain similar results whereas replication is similar except that different data is selected. According to RR proponents, benefits would be numerous. First, for verifying results of a published study (Peng, 2011). Second, for reusing previous work and build new knowledge. While the latter brings a constructive and enriching dimension to reproducible science, the first one is clearly oriented to alleviating scientific misconduct, particularly in Life Sciences (Laine et al., 2007).

Despite the fact that RR proponents are focused on suggesting to exchange code and data as a minimal threshold for “good science”, they do not examine the methods used or people participating in CEs. Methods are not of interest to RR as the main focus lays on getting similar results for verification. Hence, the end product of a CE is seen as a script, or package that should be made available by the authors of a paper as supplementary material.

The issue investigated in this work emerged from three phenomena: (1) the notorious increase of data generation and resource intensive analytics. Here in the biomedical domain, (2) ignorance about data generation processes and their impact in terms of modelling. For instance, the sequencing instruments and custom bioinformatics pipelines producing analytical data and how well they represent underlying biological facts and (3) non-specialists, not trained in data analytics, eager to participate in computationally intensive experiments but preferably via convenient end-user interfaces instead of custom scripts or programs (Holzinger et al., 2014).

The phenomena described above were observed during a design science research (DSR) (Hevner & Chatterjee, 2010) we conducted in the domain of biomedical genetics. Our research was focused on designing an interactive data mining tool for biologists to identify interesting outliers in RNA-Seq count tables. Ultimately, the goal is to seek how to facilitate access and how to reuse scripts and packages for bioinformaticians and biologists at the same time. After one design cycle of a technical artifact and its evaluation by three focus groups gathering biologists and bioinformaticians (n=15) we collected evidence against some practices proposed by RR and suggest potentially fruitful improvements.
Indeed, reproducibility of CEs should not be reduced to code and data sharing as it does not cover fundamental characteristics of modern data analysis in biology. We state that web resources and their support for multiple representations that satisfy the interest of both types of users involved will have a positive impact on reproducibility by facilitating reusability first.

2 BACKGROUND

Two aspects of knowledge creation and sharing are presented. Together, they clarify what issues emerge from code and data sharing when all stakeholders involved in a CE are not considered. We make use of a standard knowledge cycle, the Integrated Knowledge Cycle (IKC) (Dalkir, 2005) to emphasize the issues of codification implied by Reproducible Research. In knowledge management, codification aims at making implicit knowledge (from an individual) available as an object that is separated from the individual (Hislop, 2005). This can also be seen as the goal of RR which distributes experiments as packages.

![Integrated Knowledge Cycle with three stages](image)

The IKC is illustrated in Figure 1. We focus our discussion on the knowledge capture and creation and knowledge sharing and dissemination phases. The last phase acquisition is not discussed here as we believe it to be the role of academia or industry in general.

2.1 HCI-KDD

We start with Human-computer Interaction (HCI) which is the “study of the way in which computer technology influences human work and activities.” (Dix, 2009). Knowledge discovery from databases (KDD) is defined by Fayyad as “the nontrivial process of identifying valid, novel, potentially useful, and ultimately understandable patterns in data” (Fayyad et al., 1996).

The first aspect is that an end-user should be able to analyze data by using steps from the knowledge discovery process but interactively. This combination between KDD and human-computer interaction was theorized by Holzinger (2013). Tailored to the biomedical field, the process emphasizes that an end-user needs powerful visualization tools as much as data management and analytics capabilities. Holzinger also stresses the fact that reproducibility should be investigated further as it represents a major problem with data intensive experiments (Holzinger, 2013).

The steps of the HCI-KDD are integration, pre-processing and data mining. Integration is the activity of merging structured or unstructured data sets. Pre-processing applies normalization or transformation techniques to make the data sets suitable for data analysis. Data mining is the design and application of algorithms to identify patterns, associations or outliers.

2.2 Reproducible Research

The second aspect is the need for better reproducibility of experiments which are conducted with computers. Here we integrate notions belonging to two approaches to reuse context and computational material.

On the one hand, based on literate programming (Knuth, 1984), dynamic documents (Pérez and Granger, 2007) and compendiums (Gentleman and Lang, 2007) constrain design choice to add human and machine readable context to executable code. Compendiums aggregate dynamic documents. Dynamic documents are executable files that contain code with descriptive information. They are currently available with authoring packages in R (Knitr, Sweave) or Python (IPython notebooks, Jupyter).

On the other hand, an ontology based approach for dissemination of reusable components is assured by semantically enriched objects aggregating resources about the context of an experiment and its material. These are called Research Objects (RO) (Bechhofer et al., 2013).

3 DISCUSSION

As we noticed, the fact that one end-user deals with each step is, at least, a very optimistic view on data analytics. The HCI-KDD process implemented in our prototype was discussed among participants (see
section 3.1). The questions were oriented to the flow of analysis and presence or absence of components (e.g. charts, packages, result tables, context…) in the interface. Additionally, a survey was answered by 11 respondents (n=11) about how they are dealing with data and Reproducible Research.

3.1 Focus Groups Result

Inside our three focus groups we divide participants according to their main interests, i.e. bioinformaticians and biologists.

For the first type of participants, bioinformaticians, a friendly user interface is rejected. Scripts are preferred for analyzing data. Regarding methods applied, a participant indicated that a method is sometimes selected because “it works” and is not a matter of “hidden” assumptions. By assumption we refer to prior knowledge of the state of the world embedded in packages or statistical models. Not being aware of them makes a package acting as a “black-box” with unknown consequences on the rest of the processing.

For the second type of participants, biologists, they estimated the presence of such methods as appropriate. The indications given on the website (package name, version, reference paper, running environment and online documentation) are sufficient if kept up-to-date. The web interface offered the possibility to apply different methods on the same data set. This was judged as beneficial because the influence of a choice could be assessed by the user interactively. In that case, another concern raised by bioinformaticians is about the interpretation of results by users that would not be trained in statistics.

Regarding reproducibility, the lab part of an experiment has strong influences on the rest of the pipeline and it is perceived as challenging to integrate in the tool. Efforts for improving reproducibility are welcome but full reproducibility is impossible, as indicated by participants in the third focus group.

3.2 Code and Data for Verification

It is the view of Peng (2011) that executable code and data form a gold standard of reproducible research. We argue that these elements are not of interest for each important type of stakeholder involved in a computational experiment. We may admit though that what the author tries to achieve is a minimal level of reproducibility for verification purposes. The idea is that a reviewer would carefully inspect code shared with a paper, e.g. as an R package on Bioconductor. With that package, the entire computational workflow is runnable and shows figures that are identical to their online or printed counterpart.

But as even noticed by Peng (2011), papers validating previous work are rarely acclaimed by publishers which expect “new” knowledge to be submitted. This may be an explanation while results from our survey showed a poor interest in full replication. On a scale from 1 (never) to 5 (always), the need for full replication has a Mode of 2 (Median=2). Partial replication did slightly better with a Mode of 3 (Median=3).

3.3 Reusability and Interactivity

Regarding Research Objects, they sometimes appear to be developed as external solutions or repositories. We would lose a major group of researchers if the goal of an application is to purely manage research objects. Instead, the software application should produce resources that might be automatically aggregated in a RO. This is a transparent manner for users more interested in advanced visualization capabilities.

Therefore, we claim that Research Objects could be a hidden component of any interactive mining tool. By doing this, we encourage RO generation and usage without transforming such tools in a “reproducibility manager” for users interested in getting precious insights from their experiments. Exaggerating any requirement of RO management for these stakeholders will most probably result in a rejection of the entire application. This could be achieved by automatically extracting information from earlier processing stages and intermediate data sets in the analysis flow.

3.4 Resources and Representations

An interesting proposal in compendium design was the notion of transformer. We present it in this work as the creation of a representation (or view) from a single resource. A resource is an object of interest whereas a representation is a usable form of a resource which corresponds to the consumer’s interest. We designate by consumers both human and machine readers or interpreters.

In the RO world, it implies to work on ontologies and machine readable standards. For biologists, it means that a chart resource has to render a dynamic representation. We can imagine that after exchanging a RO, we find a data object resource and a chart resource. A chart shows the content of a data object as, for instance, a scatterplot. We expect an end-user to be willing to select parts of this scatterplot, zoom-
in or display labels. We also expect that this chart resource is identical to what was generated by a team of researchers which created this RO.

As we show in the next section, open source technologies for visualization “as a resource” exist and are under heavy development. They are able to create Json or html/JS serialization of a chart resource while providing enough interactivity for end-users.

4 SOLUTION

The evaluation of our prototype yielded limitations of both HCI-KDD and current practices defended by Reproducible Research. Hence, we suggest an improved knowledge discovery process embedding the HCI-KDD in an extended process named Reproducible Research Oriented Knowledge Discovery in Databases (RRO-KDD).

When conducting a DSR, four stages appear at each design cycle (Hevner and Chatterjee, 2010). The problem specification resulted from a literature review and meetings with experts in biomedical genetics. The other steps found in design science research are Intervention, Evaluation and Reflection.

Each of them are described in the next subsections.

4.1 Specification

The problem addressed in this work encompasses reproducibility and visualization for researchers in biology who are collaborating with bioinformaticians. As explained in the background section, computational experiments are not only conducted on the bioinformatics side of data analysis. Hence, an application enabling self-service data analytics for biologists has additional constraints. Self-service is understood as letting users perform analytics tasks without advanced knowledge of programming or statistical modelization.

4.2 Intervention

As technical outcome of the DSR we conducted, a prototype was developed and deployed in a research lab for structural genomics at the University Medical Centre Utrecht (UMCU) in the Netherlands.

The prototype started from the HCI-KDD process by implementing interactive visualization capabilities together with methods to pre-process and mine data sets. Pre-processing consisted in normalization and transformation of table of counts generated by RNA-Seq technologies and tools. A table of counts has samples of patients in columns and a list of genes as rows (60 000 in the files used).

Via the web interface, users start with these tables
in a virtual experiment (gathering data and contextual information). Then a possibility is offered to normalize or transform data sets by calling packages from Bioconductor. Normalization is an important pre-processing task to make samples comparable due to the presence of (technical) biases in the raw data.

4.3 Evaluation

Exploratory focus groups with biologists and bioinformaticians provided input for conducting additional iterations, similar to an agile approach. From requirements and discussions with specialists a set of functionalities for KDD and visualization were implemented. The facet of RR was imposed as it was not a primary requirement from the field experts. Hence, design choices for RR were inspired by previously described literature about compendiums and ROs.

Next, three confirmatory focus groups invited bioinformaticians and biologists to discuss about the prototype and judge the applicability of the KDD steps implemented. We addressed results obtained from the focus groups in section 3. These results are further processed in section 4.4. We present a design proposition which is an outcome of the evaluation of the prototype. Furthermore, our design proposition covers architectural choices which are mainly grounded in the web architecture.

4.4 Reflection

The lessons learned from our DSR are described in the RRO-KDD process. We processed the input of three confirmatory focus groups with 15 participants. We described the results earlier and elaborate on their processing further in the next section.

4.5 RRO-KDD Process

In Figure 2, the RRO-KDD process is modeled with its related “deliverables” in a so-called process-deliverable diagram (PDD) (Weerd and Brinkkemper, 2008). Here, the elements of the HCI-KDD process are integrated with contextual and technological outputs. These outputs are directed to reusability of previous experiment code, data and methods. Below, we shortly describe the steps and deliverables:

1) Understand is an activity where sufficient description of the data sets are provided. For instance, information about instruments, sequencing platforms, sample preparation. It builds a container for an experiment which is denoted by virtual experiment. Virtual experiments are uniquely identified aggregation of resources and group data sets together with context and methods.

2) Integrate, pre-process and data mining are the steps elaborated by the HCI-KDD process. Visualization is an activity that occurs in parallel to KDD and enables to get insight of what happens at each step. For instance, it helps the users to judge the impact of pre-processing methods on the data set. Activity Integrate results in data objects, and Pre-process will normalize or transform these integrated data sets into analytical data which are more easily interpretable than raw data, e.g. from sequencing instruments. Finally, data mining results find useful patterns from data, according to Fayyad’s definition (Fayyad et al., 1996). Visualization is here a subpart of the whole HCI field of research as it was not extensively investigated in this work.

3) Visualization has a deliverable called insight, which informs researchers on patterns, scores or relations in their data on an interactive manner. Interactive plots were rendered with bokeh, a python library for creating browser compatible visualizations.

4) Access presents previous, interactively created components of an experiment (like charts and new data objects) as REST resources that might be accessed without the user interface via RESTAPIs.

5) These resources, aggregated in a virtual experiment can be semantically enriched for reuse as ROs. This is made possible because each component is uniquely identified and accessible via a programming interface. As an example, a mining task created by a biologist is reusable via a RO with its unique identifier.

The code of the prototype is hosted on GitHub under MIT license and is available here: https://github.com/armell/RNASEqTool.

5 CONCLUSION

Results suggest that reproducibility cannot be reduced to data and code sharing and that the field of biomedical genetics suffers from a lack of software solutions that are both satisfactory for bioinformaticians and biologists who are mutually engaged in CEs. There are overlapping data analytics practices but also serious apprehensions from bioinformaticians to offer such a type of application to biologists if they exceed data visualization.

Despite these concerns, we found that there is gap to fill both in terms of data analytics and reuse of previous work.
As we have seen, biologists were more inclined to ask for more visualization capabilities whereas bioinformaticians expect a solution where scripting or custom data processing is allowed. Unique identifiers of resources and platform-independent information exchange via REST enables this. Nevertheless, HCI alone for biologists is not satisfactory as they want to query data and compare the impact of different methods. These comparisons require pre-processing and mining.

Reusability of data, workflows or parts of experiments seems to be more interesting for the two types of end-users which evaluated the artifact than reproducibility.

6 FUTURE WORK

The suggested RRO-KDD is still in a design proposition phase that needs to be evaluated in other settings and the interest in sharing Research Objects must be assessed. For this assessment, the mining tools have to be upgraded and provide more realistic possibilities to exchange and reuse virtual experiments and their components.

In addition, extending the RRO-KDD to distributed systems will have similar problems encountered in previous studies and known as workflow decay. This issue still holds in the RRO-KDD context which is built around web services and URLs that may be inactive after some time. Permanent Identifiers may moderate accessibility issues but not the support of data objects or remote implementations of analysis packages.

Recommendations to face these issues are an integration with virtual environments or containers (e.g., Docker), dynamic documents and proper data management solutions. More research on integrating virtual containers for reusability of computational experiments for bioinformaticians and biologists is needed. Dynamic documents generated by the tool could also play a role for bioinformaticians to understand what decisions were taken by biologists processing data via a user-friendly interface.

These investigations should be made by effectively combining HCI and KDD as suggested by Holzinger. But the multiplicity of actors, analysis tools and techniques remains a great challenge first for reusability then for reproducibility.

Hence, reproducibility arguments in literature should be replaced by better designs for reusability in IT solutions, at least for enhancing collaboration between bioinformatics and biologists. Reusability is broader than reproducibility as it enables repurposing of previous work and, in essence, reproducibility.

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REFERENCES


