Mr. Clean: a Tool for Tracking and Comparing the Lineage of Scientific Visualization Code

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Abstract—Visualization is a key step in scientific analysis and understanding in many fields. Scientific studies often require development of software that produces visualizations. However, as a study proceeds, the software evolves, and both developers and expert users have to periodically ascertain how code modifications affect visualization output and hence the results of the study. To our knowledge, no current visualization framework enables tracking and comparison of the lineage of scientific visualizations. We describe an approach for comparing and maintaining the code for scientific analysis and modeling through interactive comparison of visualization output. We have realized this approach in a tool called Mr. Clean. This tool provides a framework for combining different visualization tools, interaction devices, and display middleware for visual comparisons on high-resolution displays. Mr. Clean also provides user-configurable interactions supported by many devices. We provide use cases and a requirement analysis for our approach, and we describe the design and implementation of Mr. Clean. Source code is available at: https://github.com/UniversityofTromso/mrclean.

I. INTRODUCTION

Software for processing and managing large amounts of data is vital to the advancement of knowledge in many scientific disciplines. Such data-intensive science is often referred to as the fourth paradigm of scientific research where theory, experiment, and simulation are unified for data exploration [1]. One of the most important tools for data exploration is visualization [2], [3].

Scientific visualizations are often the end result of complex statistical analysis. They are often problem-specific, and to provide novel insights, it is often necessary to develop novel analysis methods. Scientists face the challenges not only of understanding and interpreting one visualization, but also of comparing the visualization with other visualizations resulting from alternative statistical methods or models. During the development of new statistical models, code changes may affect the model’s output. It is important to understand how a given change in code relates to a change in output. Due to the complexity of these models and their dependence on the input data, such understanding is typically obtained by comparing the visualizations produced by the different models.

No existing code revision tool enables model developers to interactively compare the differences between visualizations resulting from statistical and mathematical models. Instead such comparison is typically done with ad hoc implementation of multi-graph visualizations either in scripting languages such as R, or by combining images in, e.g., PDF documents. This severely limits research capabilities by forcing scientists to perform boring, time-consuming manual studies to understand how changes to code affect analysis results. An interactive approach for tracking and comparing the lineage of scientific visualizations will allow scientists to better evaluate, understand, and improve the analytical performance of their methods.

We propose an approach for maintaining and comparing scientific data analysis and modeling code through interactive comparison of the code’s visual outputs. We have implemented our approach in a system called Mr. Clean, which we have deployed for comparative analysis of genomics data cleaning and image processing methods. In the next section we describe two case studies that we use to provide a requirement analysis for our approach. We then outline this approach, and the design and implementation of the Mr. Clean system.

II. CASE STUDIES AND REQUIREMENTS ANALYSIS

We built Mr. Clean to solve problems we and our collaborators met when developing genomics data cleaning and image processing software. We present these two tasks as case studies and use them to formulate requirements for an approach for tracking and comparing the lineage of scientific visualization code.

A. Microarray data cleaning

Our genomics collaborators have developed several data cleaning methods for the Norwegian Women and Cancer (NOWAC) postgenome biobank [4]. Data cleaning is an important first step in genomics data analysis that may severely bias the statistical analysis at later stages. For microarray data analysis, this includes selection of a normalization method and
methods for identification and removal of outliers. A human expert uses these methods to identify and visualize outliers. Oldham et al. [5] describe one such outlier removal process in detail. The statistical methods are often implemented in a framework for statistical computing such as R, which provides a big ecosystem of packages for implementing different methods and parsers for genomics data.

The expert uses visualizations to determine which outliers should be removed and which should be kept. An expert typically wants to remove outliers resulting from instrumentation or experimental error and keep outliers resulting from natural biological variation. The expert removes unwanted outliers from the input data and repeats the process as many times as necessary. Although we focus on microarray data, the approach is similar to data cleaning for other data types [6]. There is no one-size-fits-all data cleaning solution, so the analyst must typically explore several statistical methods to find the best approach for a particular dataset. Such a comparative approach is often a collaboration between developers, statisticians, and domain experts.

B. KEGG pathway image processing

We developed a system, Amdex [7], that extracts metadata such as reactions and entities from KEGG pathway images [8] using computer vision algorithms from the OpenCV library [9].

Computer vision problems are notoriously difficult to tackle. Results can vary greatly when different algorithms are used in different orders, but also due to the input images having slightly different features. The common approach is to use an iterative process to figure out which algorithms, parameters, and order of algorithms to apply. To compare two approaches, the developer typically inspects the resulting images. Although we focus on pattern recognition on KEGG images, development of computer vision software in other domains faces similar challenges.

C. Requirements

Each of the above use cases produces many images. For an expert to make meaningful comparisons, she must organize and layout the images in a way that makes it easy to understand the relationships between the images and their association with the model code that produced them. The images can be grouped and sorted based on their underlying approach, dataset, iteration, or statistical method. By showing multiple integrated images simultaneously, users can compare how different data cleaning approaches perform. However, there is no predefined hierarchy among the groups, so the user must explore several groupings. For example grouping by method and ordering by iteration, or grouping by iteration and ordering by method. It may also be necessary to compare visualizations produced by latter-stage analysis tools to truly understand the effects of the data cleaning on the final analysis results.

Based on the case studies, we believe a tool for tracking and comparing the lineage of scientific visualizations should satisfy the following requirements:

1) User-defined visualization scripts. It is not realistic to change the many different tools used to generate visualizations, nor to provide libraries for the many different frameworks used to generate visualizations.

2) Interactive grouping. To compare the lineage of visualizations, it is necessary to interactively group these based on methods, iterations, versions, etc. It is difficult to automatically create the best grouping and ordering of the visualizations, so the tool should provide an interactive user interface to change these.

3) Multiple visualizations. There are often a large number of visualizations. Displaying multiple visualizations simultaneously may make the comparison easier.

4) Automated provenance management. The tool should automate provenance management so that the user can experiment with alternative methods and parameters, and revert to the best methods and parameters.

5) Collaboration. The approaches are often compared in a collaborative setting. The tools should enable multiple people to view and interact with the visualizations.

III. Mr. Clean

We have implemented the Mr. Clean tool to fulfill the above requirements. Our approach is as follows:

1) Domain experts supply visualization scripts that produce image files (requirement 1) to a directory tree managed by Mr. Clean. The path in this tree is user-defined and represents the metadata of the image files.

2) Mr. Clean records new images and changes to the scripts to track the visualizations provenance (requirement 4). We use a revision control system to manage data provenance for both the modeling code and the output visualizations.

3) Mr. Clean reads the visualization files to populate an internal data structure with files and their corresponding metadata. It uses this metadata to group and sort the images. The images are then displayed spatially organized according to group and sort order. The user interacts with the groups and sorted images to explore data by, e.g., regrouping the images or rearranging the spatial display of the images (requirement 2). We use a flexible user input event system that enables the use of the interaction devices most suitable for a given display platform, and the adaptation of these to application-specific interaction patterns.

4) We use large high-resolution displays that show many images simultaneously and enable collaborative image comparisons (requirements 3 and 5). Mr. Clean is deployed and in use on the Tromsø display wall [10].

A. Architecture

Mr. Clean is a component-based distributed system (Figure 1). The Core component is responsible for coordinating the other components, and for providing these components with communication interfaces. It holds the state of the whole system, and it processes and propagates events received from
A core feature of Mr. Clean is its flexible interface for different input systems. This interface accepts input events and maps these to operations that manipulate the visuals’ data structure and their layout on the display. We currently use the gesture system described by [13], which we have extended to support different devices.

Our current implementation allows the users to associate a particular gesture with a sorting criterion for the visuals. The sorting criterion is a user-defined list of parameters that must
match the directory structure from Section III-B. This allows
the users to cycle through different visual arrangements and
compare the different results of the scripts executions (Figure
2), satisfying requirement 2.

F. Informal Evaluation

We are currently evaluating Mr. Clean with our domain-
expert collaborators. In this paper we report our initial expe-
riences developing and using Mr. Clean.

We found it easy to modify the data cleaning scripts to
output files with the necessary metadata. The modifications
had the added benefit of better structuring the code, and
making it easier to keep track of code changes.

The high resolution and large size of the display wall
enables multiple users to compare images by walking along
the wall. In addition, the gesture interface allows the users to
interact with the images while standing in front of the display
wall without having to walk back to a workstation. We believe
that a more refined and intuitive gesture interface can improve
and possibly speed up the data cleaning process.

IV. RELATED WORK

Display walls are often used for scientific visualization [14].
Hibbs et. al. [15] demonstrate how integrated views enable
novel biological discoveries. WindowScape [16] is a window
manager that uses flexible implicit grouping of windows, but it relies on mouse and keyboard for interaction. It is
common to use different interaction approaches depending on
the interaction space [14]. Mr. Clean is, to our knowledge,
the first display wall system that provides a specialized dis-
play and interaction approach for interacting with scientific
visualization code lineages.

Systems for lineage management of scientific code and
data include Galaxy [11], which is popular in Bioinformatics.
However, the data lineage is usually managed manually. DEVis
[17] is an example of a visualization tool that focuses on the
evolution of non-code artifacts in a software development
setting, namely technical documentation. VisTrails [18] is a
system for work-flow and data provenance management with
support for data exploration, visualization and simulations.

Recent advances in human–computer interfaces include per-
ceptual input systems [19], and ubiquitous gesture recognition
systems [20].

V. CONCLUSION

Mr. Clean is a tool for maintaining and comparing scientific
data analysis and modeling code through interactive compar-
isons of their produced output.

Our approach allows users to understand how changes to
their scientific code affect analytical results. By automati-
cally organizing the visualization output of different versions,
datasets, iterations, and methods, the user can overcome the
limitations of ad hoc comparisons.

We believe that our approach is general enough to be
applicable to other fields of scientific visualization code than
only those of our two case studies. Understanding the effects
of output data from changes to scientific data analysis and
modeling code is vital for the development of robust analysis
and modeling software. Mr. Clean enables such understanding.

Although Mr. Clean is already useful we plan to improve
it. The first improvement planned is to provide a code diff
by selecting two or more images with a gesture. The diffs
provide the user information about the cause of the changes
in the images by looking at the changes in the visualization
code.

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