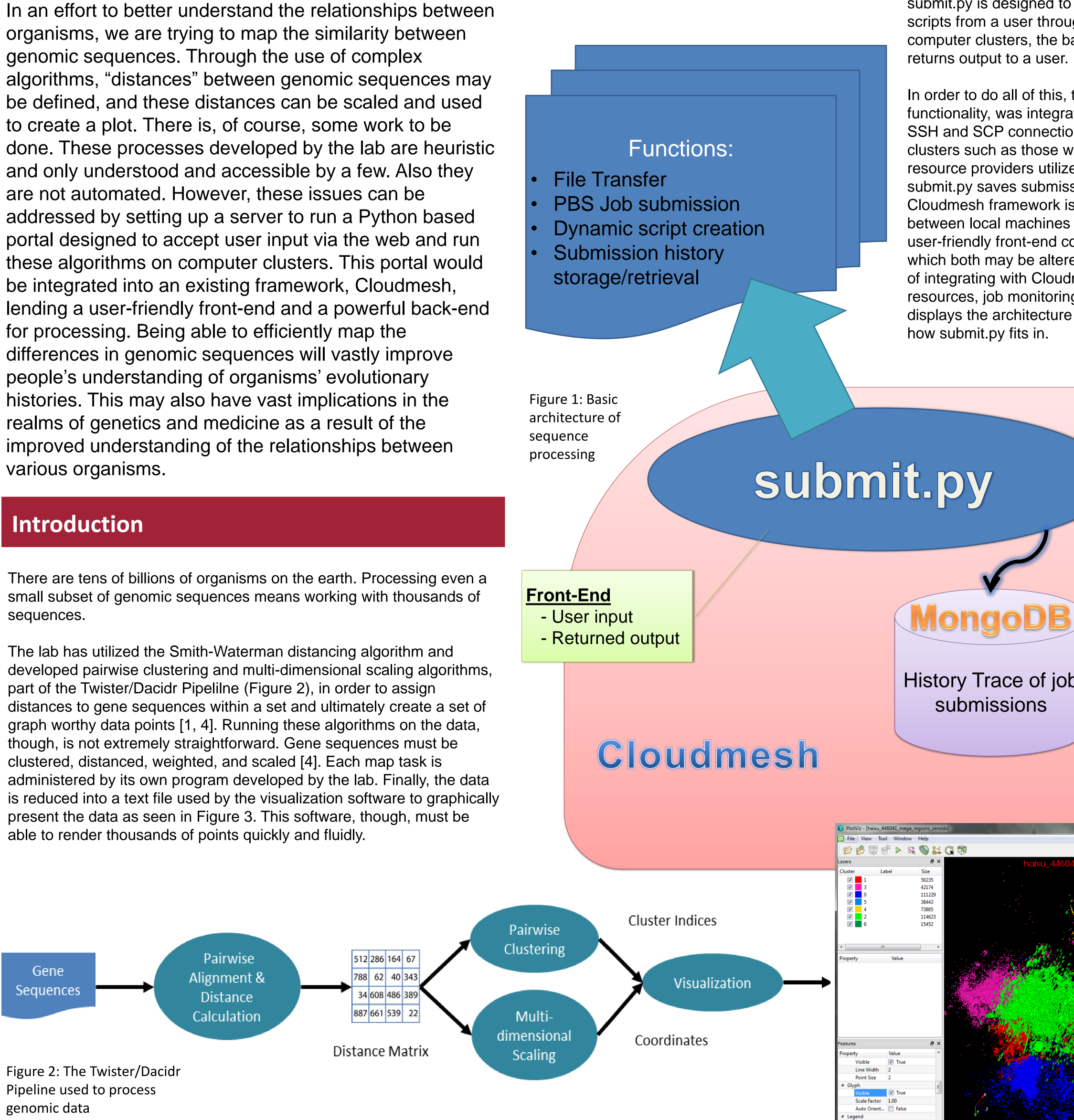
Genomic Sequence Analysis Automation

Lorander Saggu, Saliya Ekanayake, Yang Ruan, Gregor von Laszewski, Geoffrey Fox

Indiana University

Abstract



Submission Tool

submit.py is designed to accept Portable Batch System or PBS scripts from a user through the front-end and run them on computer clusters, the back-end. The tool transfers files as well as returns output to a user.

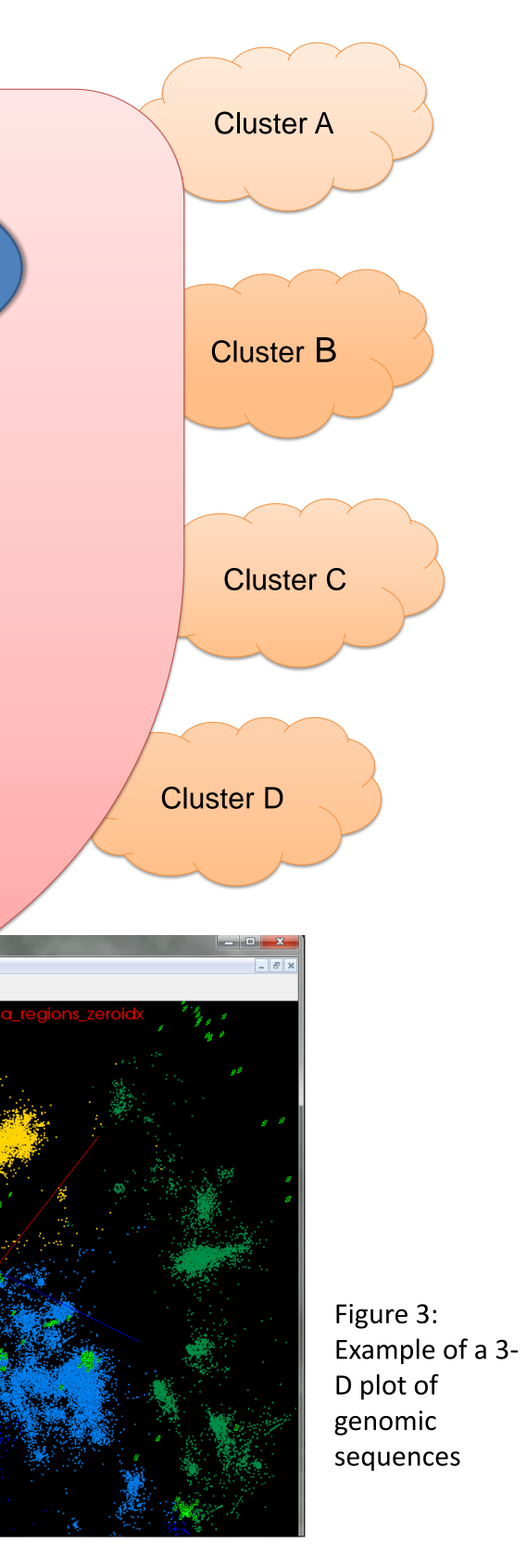
In order to do all of this, the submit.py program, with command line functionality, was integrated with the Cloudmesh framework. Via SSH and SCP connection, submit.py may access computer clusters such as those within FutureGrid, one of the computer grid resource providers utilized by Cloudmesh. Also, within Cloudmesh, submit.py saves submission information into a history trace. The Cloudmesh framework is designed to allow efficient communication between local machines and computer clusters [3]. It possesses a user-friendly front-end command line and graphical user interface which both may be altered for specific purposes. Also, the benefits of integrating with Cloudmesh include access to multiple cluster resources, job monitoring, and web based interfacing [3]. Figure 2 displays the architecture of the genomic sequence processing and how submit.py fits in.

History Trace of job submissions

False

Visible





Conclusion

The results of processing genomic sequence data and creating 3dimensional visualizations have huge implications within the fields of biology, medicine, and genetics. Being able to quickly and reliably process genomic sequences into friendly visualizations may allow biologists to recognize unnoticed relationships, geneticists to understand and explain evolutionary changes, and medical experts to infer the effectiveness of potential treatments. By working to streamline the job submission process, analysis of genomic data will become more accessible to these groups: biologists, doctors, geneticists, and other interested parties.

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Primary Contact

Gregor von Laszewski, Indiana University, laszewski@gmail.com

