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POODLE 1.0 - To the core of your data

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Abstract

Motivation: Many life science laboratories are still using Excel files to organize their data. This leads to a huge workload for maintenance as well as a inconvenient access and update routine. POODLE provides an easy-to-use and powerful interface to improve the cloning work of a lab.

Results: POODLE is a Java-based web interface, which allows intuitive access, update and manipulation of data for cloning.

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Supplementary information: Supplementary data and the code are available at https://github.com/derjedi/Bioinformatics/Assigments/tree/master/poodle_project online.

1 Introduction

The POODLE project is a project which deals with really daily duty in a bio laboratory. Due to the data managment aspect of sequences and the task providing information fast, the project is clearly settled in the computer science field. The background data, which was obtained, was focussed on cloning for protein function studies or structure elucidation. Therefore we will focus on this application of the POODLE.

1.1 A real world pipeline

The cloning procedure is a standard strategy, which is used in nearly every life science lab. One possible application is in protein structure-function relationship studies. These studies use proteins, with a known sequence and start to modify the sequence with mutationsin order to observe changes in the protein structure or functional bahaviour. The general pipeline of such an experiment could be, introducing the mutation via cloning, isolating the mutated protein and visualisation of differences via several methods. The visualisation for example could be an NMR interaction study or via X-Ray-Crystallography or any other essay. POODLE trys to manage the first step of this pipeline, which is not only in this pieline the basic step.

1.2 cloning as lab strategy

Cloning is the key to a huge ammount of variety in the resulting protein constructs. Via cloning, it is possible to remove a function, add a function to a protein or influence the structure. Today, there are many different cloning methods available, like the classical restriction approach, the site directed mutagenisis method, restriction free cloning or the Crisp-Cas system. All methods have a specific task and are chosen in various situation. For this paper we want to stick to the site directed mutagenisis, which was initially developed by stratagene as QuickChange site-directed mutagensis kit. This method allow to introduce small directed mutation into a protein on DNA level. these mutations can be used to perform point mutations or switch single or multiple amino acids.



Fig. 1. Schema of QC

1.3 cloning methods

Classsic Cloning, Quickchange, Restriction free cloning

1.4 The resulting data

 $primer\ cloning Vectors\ protein Constructs$

2 Approach

As already mentioned above many life-science laboratories use Excel files for storing and working with data. But Excel files have a few drawbacks, which make working with them inconvenient. One of the biggest problems occurs, if more than one member of the laboratory tries to access the file at the same time. Inconsistent files can be the result of a simultaneous access. Second, searching in a Excel file is not as easy as it could be. Furthermore, licenses for Excel are expensive and switching to a free software solution saves a lot of money. The reason for using Excel rather than a database system is that most life-scientist are not familiar with database systems and prefer the GUI of Excel to work with. We are trying to solve that problem by combining the functionality of a database system with an intuitive and easy-to-learn handling.

3 Methods

POODLE is build as a two-layer software. The first layer consists of the database and routines to automatically build and update that database. The

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second layer consists of the web service that is used to access, update and manipulate data. This is the front end layer and provides access to all functions for the user, such as accessing the database, manipulating data and using the in-build Blast functionality.

3.1 Database layer

SQLite¹ is the database system running in the background. We chose this software for several reasons. First, it is free of charge. Second, and more important SQLite is a small and fast database system written in C. Since, the whole database is stored just in one file SQLite has a incredibly good performance combined with a really small among of required space. And there is the possibility to create a mobile version of POODLE without changing the whole database system, which may be interesting in the future. Besides, SQLite guaranties that transactions have all ACID (Atomicity, Consistency, Isolation and Durability) properties even if a system crash or power failure occurs. So we have a robust storage and access of data as well as a lot of functionality provided by the database system. In addition to the SQLite database POODLE's software creates a BLAST database for the provided data, too.

The build and update process is implemented in Python. POODLE combines a SQL database with several local Blast databases and this routine ensures that the SQLite database and BLAST databases are always in the same state. It also makes the database a little bit safer, since changes can be reviewed before inserting them into the database. Furthermore, the user does not need detailed knowledge about SQL to create the database, which may be helpful for many laboratories.

3.2 Web service

POODLE web service is build with Vaadin². We chose that framework because it allows a safe communication between the front end and the database layer as well as a smooth and intuitive user interface. Since a website developed with Vaadin is highly modular, it is easy to extend the web service in the future. That could be necessary, if new functionalities are required. Currently, there are three pages with different roles. The search page provides a formula that can be used to perform a simple SQL search in the local database. Therefor, different fields are displayed and can be filled by the user. A click on the search button will automatically build a SQL query and results are displayed, if there are available results. The new entry page allows to insert a new entry into the database. It also provide a Blast search against the NCBI databases to check, whether there are already available information about the new sequence or not. Last but not least, users can perform a local Blast search against their one database or a remote Blast search against the NCBI databases on the Blast page. That functionality can be found on the Blast page.

3.3 Blast functionality

Our software package provides not just a database functionality for powerful and safe data storage and an easy-to-use interface but also a Blast (Altschul *et al.*, 1990) functionality. This function is implemented by using a combination of Blast binaries provided by NCBI and the Biojava Api (Prlić *et al.*, 2012). There are two different Blast searches available in POODLE. On one hand, there is a remote Blast search on the NCBI server. And on the other hand, there is a local Blast search. The remote Blast search is not a fully functional copy of the NCBI web service but can be used to perform a quick and simple search against databases provided by NCBI, e.g. Swissprot (O'Donavan *et al.*, 2002). But we do not recommend to overuse that because too many requests may cause a blacklisting by

NCBI. This remote Blast routine is mainly for the insertion of a new entry. The idea is to check whether there are information about the new sequence. But it can also used separated from the new entry routine. The local Blast routine does everything someone would expect from a Blast search but against the local database, hence local Blast. There a several programs, e.g. blastn or blastp, available for both Blast methods.

4 Discussion

POODLE is completely open-source. So the user can adapt the code at any point of the software. To make this adaption as easy as possible for the user we chose to use widely known software inside of POODLE. Most scientist have heard of SQL as a database managment system. So it is not too venture to assume that there are people with basic SQL knowledge in most of the laboratories. And that means these laboratories do not have to use POODLE as a black box. We chose SQLite over all other SQL databases for the reasons described in the method section.

The build up process of the database is written in Python. There are two reasons for that. First, Python is an easy and powerful language which makes it possible for the user to adapt the build up process without detailed programming knowledge. And second Python2.7 is a default package of most operating systems installed on a server. We tried to make POODLE as platform independent as possible and that helps to reach our goal.

Vaadin was chosen as the running framework for several reasons. Java is a widely used programming language in the scientific world. So there is a relative high chance for a person in the laboratory, who is able to work with Java. Again we want to make adaption as easy as possible for the user. Furthermore, Vaadin needs a tomcat server. Such a server is provided free of charge by Apache³. So the user does not have to face extra charges just to be able to use POODLE. More reasons for Vaadin are discussed above.

5 Conclusion

POODLE provides a safe storage of cloning data as well as a easy-to-use interface for interacting with these data. Besides, it comes with functionalities like local and remote Blast search that can improve the workflow during cloning.

It is our goal to improve the software in the future by adding new functionalities like a chromatogram viewer and the possibility to add annotations to sequences.

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References

Altschul, S.F., Gish, W., Miller, W., Myers, E.W., Lipman, D.J. (1990) Basic local alignment search tool., *J. Mol. Biol.*, **215**, 403-410.

Prlić, A., Yates, A., Bliven, S. E., Rose, P. W., Jacobsen, J., Troshin, P. V., ... Willis, S. (2012) BioJava: an open-source framework for bioinformatics in 2012. *Bioinformatics*, 28(20), 2693-2695. http://doi.org/10.1093/bioinformatics/bts494

¹ http://www.sqlite.org/

² https://vaadin.com/home

³ http://www.apache.org/

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O'Donovan, C., Martin, M.J., Gattiker, A., Gasteiger, E., Bairoch, A., Apweiler, R. (2002) High-quality protein knowledge resource: SWISS-PROT and TrEMBL *Brief. Bioinform.* **3** (3), 275-284. doi: 10.1093/bib/3.3.275

Alberts B, Johnson A, Lewis J, et al. (2002) Molecular Biology of the Cell. 4th edition. *New York: Garland Science*. Studying Gene Expression and Function. Available from: http://www.ncbi.nlm.nih.gov/books/NBK26818/