



INTEGRATED BIOINFORMATICS PROJECT

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# CRISPR Exposed

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# 1 Introduction

Clustered, Regularly Interspaced, Short Palindromic Repeats (CRISPR) and CRISPR-associated proteins (Cas proteins) form a complex bacterial/archaeal immune response system that mitigates foreign DNA activity. The identification of these systems first occurred in 1987 in *Escherichia coli* [1], followed by other species of bacteria and archae in the 90's [2]. Their function however was not elucidated until 2007 [3]. CRISPR-Cas systems work by capturing short signatures of invading DNA elements and inserting them into the genomic material in regions known as CRISPR arrays.

These arrays consist of the captured elements, known as spacers, which are separated by similarly sized, conserved DNA sequences known as repeats. The arrays are transcribed and processed into CRISPR RNA (crRNA). crRNA, together with Cas proteins, form an active complex that patrols the cell. If an invader DNA element with a similar signature is encountered by the complex, it will be degraded and its activity will be prevented [4].

CRISPR-Cas systems can be categorized into multiple types. Each type having a different set of features and modes of action that makes it somewhat unique [5].

Previously [6], our team used computational CRISPR detection software like CRT [7] to detect spacers from all “reference” bacterial and archaeal genomes on NCBI (about 5000 genomes in total in May 2015). A subsequent step had been to blast those detected elements against the NCBI nucleotide database. This revealed the origin of about 7% of spacers, outlining the problem of the so-called *biological dark matter*. The identified spacers were also searched against the genomes of the bacteria they originated from. Spacers showing up outside of the CRISPR region(s) were classified as *hits*. Hits putatively showed that there was a possibility for given phages to integrate into bacterial genome despite the presence of spacers against them.

The main objective of this project was to make a large database of CRISPR elements from all the bacterial/archaeal genomes available in the NCBI genome database. Hence, removing the limitations we had before when we focused only on “reference genomes”. The number of genomes considered is thus substantially higher, about 54000 at the time of this project (December 2015). A database of these elements was made accessible via a web application. The associated website exposes the CRISPR details of the strains, along with a number of services, such as blasting against spacer elements, CRISPR array detection, and elements of data visualization.

## 2 Methods and results

### 2.1 Classification of CRISPR-Cas systems

We attempted to classify CRISPR-Cas systems based on methods proposed by Makarova *et al.*. The dataset from their studies (supplementary document 7) was first processed into a clean subset (removed of “partial” subtypes and ambiguous classification) to search for possible classifiers. Loci were classified as “partial” if they contained neither the full complement of effector module genes nor *cas9* or *cpf1*. The classification criteria used was the polythetic classification mentioned in their studies [8].

Two approaches were attempted to tackle the problem, random forest and Naive Bayes. Random forest showed unpromising results as the total factor level exceeded computational limit. On the other hand, Naive Bayes model performed well with the clean dataset with

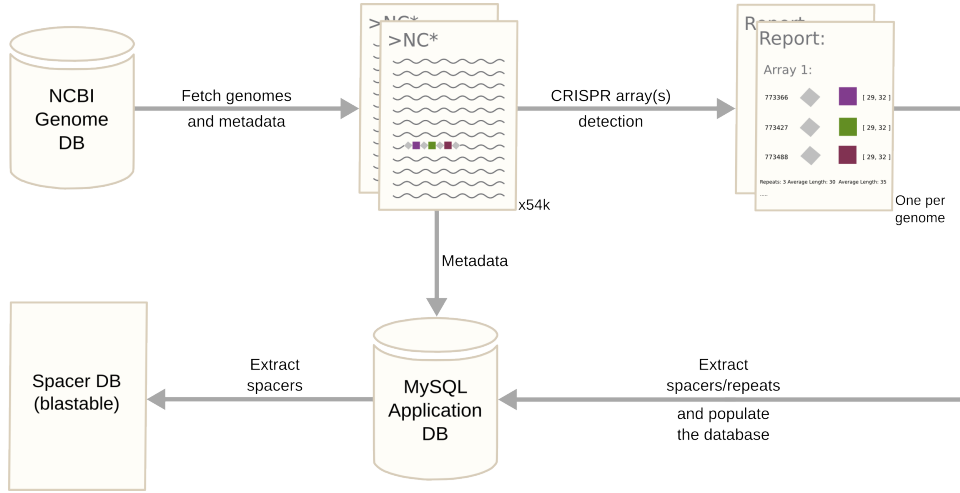


Figure 1: Data processing pipeline

an overall accuracy of 0.98, high sensitivity and specificity to all subtypes except lower sensitivity to CAS-I-A (0.85) and CAS-III-C (0.93) in the validation set.

The task proved to be non-trivial and was eventually discontinued due to time constraint. Challenges encountered in the process include ambiguous classification, and a need for a pipeline to detect and annotate cas genes, and genomes bias etc.

## 2.2 Data Processing Pipeline

The first step of the data processing pipeline was to gather all possible bacterial and archaeal genomes available from the NCBI genome database. This task was accomplished using a custom python script which fetched from the RefSeq ftp site [9] the 54000 prokaryote genomes available (all assembly levels). In practice that meant downloading the genome files (\*.fna). Alongside each genome file, the corresponding annotation files (\*.gff) were also downloaded. Finally the corresponding metadata available on NCBI for each genome was captured into JSON files. All the files, representing about 300 Gb worth of data, were stored in a “one genome, one folder” fashion on our server.

The second step was to detect all CRISPR arrays (i.e spacers and repeats) in each genome. For this, the CRT software [7] was used. In order to treat the the multi-FASTA files correctly (e.g for genomes containing plasmid sequences, or multiple contigs) the CRT software was wrapped in a python script and then applied to the fetched genomes to create the CRT report files containing detected CRISPR array(s) in each strain. Next, a custom parser was implemented to extract the spacers and repeats from each CRT report and structure them into JSON files.

These CRISPR elements and the metadata fetched previously were used to populate the relational MySQL database. All spacers available were extracted from the MySQL database and consolidated into a multi-FASTA file and a blastable database was made using the *formatdb* tool from NCBI BLAST.

## 2.3 Software Architecture

The approach taken to expose the database and let the user interact with the data was to develop a web application. We used Django to accomplish that. Django is a free and open-

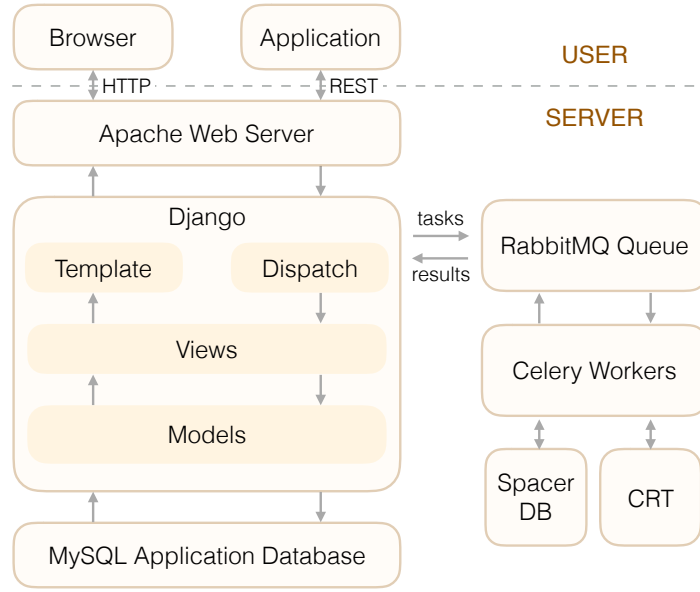


Figure 2: System Architecture

source, Model-Template-View (MTV) web application framework written in python. It encourages rapid development and clean, pragmatic design. The web application is running on a server using an Apache web server.

A key element of the web application is the REST API which allows the users to programmatically query the MySQL database in order to obtain the data for their downstream analysis. This was done using the django REST framework, a flexible toolkit for building Web APIs.

As an additional service, users have the option to blast their own sequences against our database of spacers elements. They can also submit their bacterial or archaeal sequences for CRISPR array(s) detection using the CRT software installed on the server. In order to efficiently handle user requests for BLAST and CRT services, the requests are queued before they are run by the backend, allowing for more control over the tasks and their resources usage. It also provides the option to run them asynchronously. This element was developed using Celery as a task queue and RabbitMQ as a message broker.

We used free and open-source softwares to achieve our goal. We also chose to publish our code on github, under a public license. Special attention and care were given to reproducibility. As such, the code was well documented. Moreover, the environment was also documented using an “infrastructure as code” approach with the help of tools like Ansible. This approach eased the developement phase, and provides enthusiast users key elements when trying to reproduce everything from scratch on their own infrastructure.

**Repository address:** <https://github.com/Milt0n/CRISPR-Exposed>

**Website address:** <http://www.crispex.net/>

## 2.4 Data Visualization

### 2.4.1 method

Four plots were created for data visualization using D3.js libraries. A row chart and two histograms were created using dc.js, stable version 1.7.5 [10] while a parallel coordinates visualization was plotted using parallel-coordinates version 0.7.0 [11]. The charts were plotted using datasets in csv format queried from the MySQL application database.

The first dataset was used to plot a row chart which shows the *log10* number of genomes available in the database and 2 histograms that shows the frequency distribution of repeats' and spacers' length detected by CRT. Since these three charts were plotted using the same data set, the information is cross-linked. Users can interactively filter the information in these charts (it will not affect the display in parallel coordinates).

The second data set which only consists of genomes with complete assembly level was used to plot parallel coordinates which shows the groups of prokaryotes, size of genomes and number of CRISPR arrays detected per genome. Each line represents a strain, which are coloured by their kingdom (i.e. bacteria in purple, archaea in orange). Axis can be reordered and can be filtered by range.

### 2.4.2 results

The row chart shows the *log10* number of genome sequences used to populate CRISPR Exposed for each group of prokaryotes. The purpose of showing this chart is to inform users that the content of the database is highly biased towards some species and this should be taken into account when interpreting the distribution of repeats' and spacers' length. About half of the total of the sequences originate from firmicutes, proteobacteria and actinobacteria. This chart could be further improved into scale-stack row chart in the future to reduce the cognitive load of interpreting the results.

The histograms of repeats' and spacers' length mainly serve the purpose of quality control of CRISPR elements detected by CRT. Length of direct repeats should not deviate too much from length of spacers. The dataset used to plot the histograms was preprocessed to remove false positive that is not bound by the constraint of parameter settings in CRT. For instance, the maximum length of spacers for CRISPR detection was set to 48 nucleotides long but there were about 3.6% (28663 out of 790903) of total number of spacers detected by CRT is longer than the length specified. These entries were stripped from the data set to avoid misleading information.

Finally, parallel coordinates shows that most of the complete assembly genomes available in the database have sizes around 2-7 million bp long and contain 1 to 5 CRISPR arrays. No notable correlation between number of CRISPR arrays and genome size was observed.

## 3 Discussion

The application and the database attached to it is novel in a couple of ways. To our knowledge, it is the first time such an extensive identification of CRISPR elements is conducted. About 54,000 genomes were used in this analysis. Other projects such as CRISPRdb [12], and CRISPI [13] only focus on the subset of genomes that have complete level assembly (4,000 genomes for CRISPRdb and 1,200 for CRISPI). As such, the spacers database constructed is the largest presently available online, consisting of 800,000 spacer sequences. We hope it can be useful to researchers, and can envision a couple of uses of our dataset:

- Comparative analysis of CRISPR arrays at species, class, or phylum level
- Distribution of the origin of spacers in given bacteriophages, or plasmids
- Phage host prediction

Future work could include:

- CRISPR-Cas systems identification and classification
- PAM sequences information
- Detection of anti-CRISPR genes

Building upon those features, improvements in data visualization would be welcome in order to bring further insights to users. Additional information that could be interesting to be visualized include:

- Cas genes detected: it is possible that CRISPR arrays detected is not associated with any cas genes in near proximity. Such arrays may be degenerated or have other functions [14]
- Information about secondary structures of repeats: Kunin *et al.* clustered putative secondary structures of crRNA based on sequence similarity and ability to form stable secondary structures [15]. Combining this information from the cas genes and the repeats may provide insights on function and regulation of CRISPR-Cas systems.
- Protospacer-adjacent-motifs (PAM) sequences: the selection of protospacers from invading DNA was shown to be determined by recognition of PAMs which are a couple of nucleotides long and may vary between different CRISPR-Cas systems [16, 17]

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