SDA v 0.10.x

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1 Before you start

Scan Domain Architecture (SDA) is an script intended to help you with the analysis of protein functional domain architecture and evolution.

2 Feedback

Any comments, suggestions, bugs or feedback please contact to Ramon Flores, ramon.flores.r@outlook.com.

3 Requirements

SDA requires:

- Perl v5.18.2 or later version.
- Perl SVG module for graphics.
- The Pfam-A.hmm database¹.
- The HMMER² suite for hmm profiles. SDA was developed and tested under HMMER v3.1b1, but current version v3.1b2 should work as well.
- For the GUI, you'll need Java 7 or later.

SDA makes some searches inside the PFAM database to annotate the input sequences. Here, we use the PFAM version 29 (latest, 30). After installing the HMMER suite please format your database for hmmscan searches.³

¹ftp://ftp.ebi.ac.uk/pub/databases/Pfam/releases/Pfam29.0/

²http://hmmer.org/

³To format the Pfam-A.hmm file, execute hmmpress *Pfam-A.hmm* in the terminal. This will create four binary files .h3{fimp}

4 File format and headers

To search a list of proteins with PFAM annotation in order to study their functional domain architecture you will need: (1) An input file (or PFAM list file) of the sequence(s) you want to find and must be amino acid sequences. This program doesn't work with nucleic acid sequences; (2) if you want to compare your sequence PFAM annotation to other proteins, an annotation list in the **trinotate** format can be used.

It's very important that your annotation file has the following requirements: the header for the Pfam column must begin with *pfam* (capitals or small letter, doesn't matter). If the file is a genome file, the header of the genes must be or contain *gene_id*, and for a transcriptome file must be or contain *transcript*; absence of some of these headers will result in a suddenly termination of this tool.

SDA allows you to search in an annotation file like Trinotate transcriptome results, for proteins with similar architectures or new combinations of the domains of PFAM domains of your interest.

SDA works with annotation files from genomes or transcriptomes as long as they have the required format.

5 Usage

SDA creates up to three files depending on the results found. Please, verify next section for details.

The way you have to use SDA is as follows:

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./SDA -OPTION <PARAMETER> ...
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The user's interest sequence might be a FASTA transcriptome sequence in a file, or a PFAM architecture in a file or as a list.

The PFAM list may have two forms, (1) as a file with one architecture per line and each PFAM joined by a comma without spaces; each PFAM can only have the letters PF and five digits. (2) As a parameter list, in which case this input is interpreted just as one architecture; you can list up to three PFAMS this way, each one joined by a comma and without spaces.

The all possible options and their description are the next:

-a|--acc < VALUE> Defines the accuracy for each PFAM in the hmmscan summary table. It must be value between 0 and 1 (default, 0.85).

-c cpu $<\!C\!PU\!s\!>$	Defines the number of CPUs used to create the HMM profile with ${\tt hmmscan}.$
$\verb -e eval < VALUE >$	E-value threshold for hmmscan search and result filtering (default, 1e-10). Must be positive with format input as an integer or in exponential notation.
-f fasta < <i>FILE</i> >	The path for the FASTA file with a mino acid sequence(s).
-g	Use it if your annotation file is a gene file. The expected annotation file input is a transcriptome.
-h help	Displays this help menu.
-1 list < <i>FILE</i> >	Use it if your input is one or more PFAMs architecture(s) in a file.
-o output < <i>PATH</i> >	Changes the output path directory (default, parent working directory).
-p pfam <file></file>	Compulsory. Indicate the location of your Pfam-A.hmm file.
-t trin < <i>FILE</i> >	Tab-delimited file with PFAM annotation for the genome/transcriptome file in Trinotate format. $$
-v version	Shows the SDA version.

6 Output Files

SDA generates up to three different files in directory called SDA.Results in the output directory (\sim / as default).

If you feed SDA with a FASTA file, inside the results directory, SDA will create another directory with the name of your FASTA file and inside, you'll find the results. That means you can work with different FASTA files and you'll find your results in each different directory.

If you feed SDA with a PFAM architectures file, the result will be saved inside the *SDA.Results* directory if there is at least one coincidence or similarity between the PFAMs input and the annotation file.

The output files and their content is described above:

- .table Contains the hmmscan results for the hmmscan fasta file. Please, check the hmmscan documentation for further details.
- .pdf This file contains a graphical representation of your result for each sequence in the FASTA file. It draws the domains found for the sequence(s) and it's PFAM name.
- .out It's created if SDA finds at least one coincidence or similarity between each FASTA sequence (or PFAM input) and the annotation file. Contains the gene_id or transcript, location, e-value, and description of the match. At the end of this file you'll find a summary with all the PFAM architectures found, similarity score and it's relative frequency.

7 Similarity score

The similarity of the PFAMs for the .out file depends on the PFAM cardinality of each sequence in the FASTA file or PFAM list, and the number of matches. This way, the similarity scores is defined as

$$S_{SDA} = \frac{n}{\max\{l_1, l_2\}} \tag{1}$$

where n is the amount of domains shared by the both of sequences, and l_1, l_2 are the total amount of domains found for each sequence.

8 Working with SDA

There are many ways to use SDA, we shown some ways to use it.

8.1 Identifying domains architectures with SDA

The first use of SDA is to identify domains in a sequence of amino acids. The simplest way to do this is typing

$$\ \ ./SDA -p < Pfam-A path> -f < fastaFile path>$$

SDA shown the results of processing the file and at the end, it will indicate you where you will find the results.

SDA will create the files .table and .pdf for results (figure 1).

Additionaly, you can indicate the number of CPUs used to create the hmmscan profiles, the e-value treshold, and the minimum accuracy of the results adding

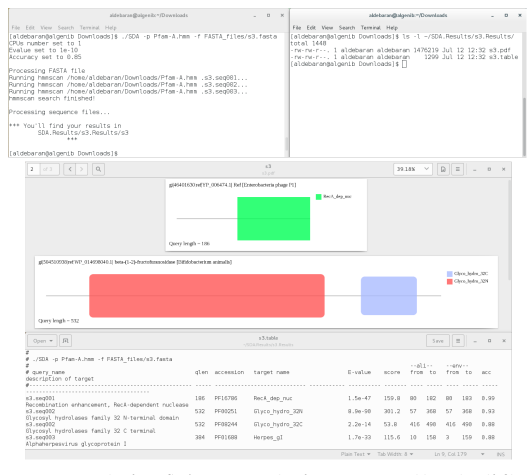


Figure 1: Results from SDA. Up, example of run. Down, .table and .pdf files results.

$$-c$$
 $< CPUs > -a < accuracy > -e < e-value >$

8.2 SDA with a FASTA file and an annotation file

For searching domain architectures in a trinotate annotation file, you must add the option -t with the path to the annotation file, and the option -g if the annotation is a genome file (default is a transcriptome). Remember that the column names in the annotation file must have a format (section 4).

This way, the command will be

And again, you can specify the number of processors and accuracy of results. The example run and results are shown in figure 2.

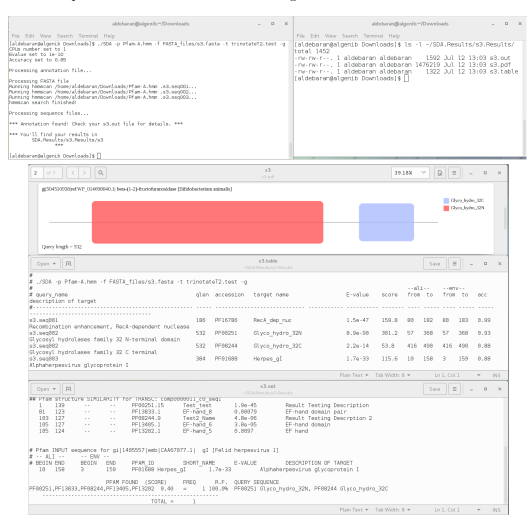


Figure 2: Results from SDA with annotation file. Up, example of run. Down, .table, .pdf and .out files results.

The .out file contains a table with information of the query domain and the domain(s) found. At the end, there is a resume with the domain architectures found.

9 Using the GUI

There is a GUI for the users that dislike using shell, it is call SDA.run and was created with Java.

To execute it just open terminal in the folder where you have the files SDA.run and SDA (both of them must be in the same location). The GUI is easy to use, you just have to select the files in your computer and click the button **start** to get results (figure 3).

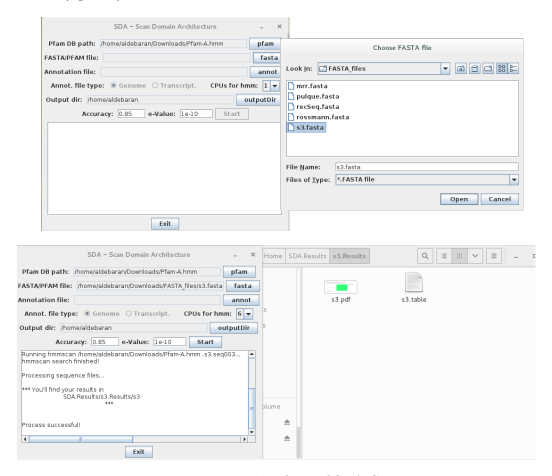


Figure 3: Example of use of SDA GUI.

When the program finishes, SDA automatically will open the file with the results.