KLAST plugin
for CLC Bio softwares

User manual – release 4.1

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1. Introduction

KLAST is a fast, accurate and NGS scalable bank-to-bank sequence similarity search tool providing significant accelerations of seeds-based heuristic comparison methods, such as the Blast suite of algorithms.

Relying on unique software architecture, KLAST takes full advantage of recent multi-core personal computers without requiring any additional hardware devices.

KLAST being designed for bank-to-bank sequence comparisons, it is not appropriate to use on a « per-sequence » basis. In other words, compare your query sequences in batch mode using set of sequences instead of running several KLAST jobs, each of them using a single query sequence.

2. Algorithm

KLAST high-performance sequence similarity search tool relies on a new optimized and combined implementation of the Parallel Local Sequence Alignment Search Tool (PLAST) [1] and the Ordered Index Seed (ORIS) [2] algorithms developed by Inria's Genscale research team [3].

KLAST aims at comparing two large sets of sequences. Query and subject can be both proteins or nucleotides for which you'll use KLASTp or KLASTn to compare them, respectively. Considering proteins versus translated nucleotides, you'll use one of KLASTx, tKLASTx or tKLASTn. KLASTp, KLASTx, tKLASTn and tKLASTx algorithms are actually using the same algorithm, relying on KLASTp principles. KLASTn uses a slightly different approach. The following sections describes KLASTp and KLASTn.

KLASTp aims to compare two large sets of sequences. It implements a three-step seed-based algorithm: (1) indexing, (2) ungap extension and (3) gap extension. Compared to BLASTp algorithm, KLASTp differs essentially during step 1. Steps 2 and 3 are similar between the two algorithms.

1. Both sets are first indexed with subset-seeds of W characters [4]. The index is a table of K entries (K is the number of different seeds). Each entry is associated with a list L of P positions corresponding to all occurrences of the seeds in the sequence set.

2. For each seed, P1 x P2 ungap extensions are performed. P1 (resp. P2) represents the number of elements associated to the list L1 (resp. L2) for the selected seed. Ungap extensions are computed on subsequences made of 2L+W characters (size of the seed + L neighboring residues before and after the seed). Ungap alignments with a score greater than a predefined threshold value are selected for the 3rd step.

3. As the previous step can generate many ungap alignments belonging to the same gap alignment, a checking on the final set of alignments is done before launching the gap process. Dynamic programming algorithm is used to compute final gapped sequence alignments.

Moreover, KLASTp is optimized for multicore processor architectures. The algorithm is natively parallelized following a coarse-grained approach scaling well with the number of cores. Furthermore, gap and ungap extension computation are vectorized using the SSE instruction set of processors.

KLASTn compares two large sets of DNA sequences. Similarly to KLASTp, two indexes including neighboring information are built, allowing the first steps of the search to be fastened. Furthermore, KLASTn includes a new search engines based on the concept of Ordered Seed Indexing [2]. This algorithm suppresses numerous redundant computations during the extension step of alignment construction.

As KLASTp, KLASTn is optimized for today multicore microprocessor architectures, and scale well with the number of cores.

It is worth noting that KLAST uses a Karlin-Altschul statistical model, as BLAST does. As a consequence, every score, e-value, etc. computed by KLAST has the same meaning as BLAST-computed values.
3. KLAST plugin overview

The KLAST plugin comes with two major tools:

1. KLAST itself to enable the comparison of queries and reference databanks

2. KLAST Databank Manager to prepare reference databanks from various public and personal sequence files

The plugin also provides a convenient way to introduce in the results some biological classification data, such as Enzyme, Gene Ontology, Interpro, Pfam and NCBI Taxonomy.
4. Setting up a KLAST job

1. From the CLC Workbench’s Toolbox, click twice on the « KLAST on local computer » item to start the KLAST job wizard:

2. Select your query sequence from your data set:

3. Select the target (or reference) databank:

Notice: since KLAST directly uses FASTA formatted sequence file, you can provide your target databank in such a format. KLAST Databank Manager being presented on section 7.), we simply use a FASTA file for this first job.
4. Setup the KLAST search parameters.

Notice: you have to choose at least the comparison method (klastp, klastn, etc.). KLAST’s parameters are detailed in section 8.

5. Choose whether or not to apply a processing tasks. Tasks being details on section 9. and 10., we let this panel empty in this example:
6. Choose the way you would like to see the results; here, we want to get the graphical display (*Result handling = Open*) and ask to see interactive Klast job processing (*Log handling/Make log is checked*):

![Image of KLAST user interface with selected options]

Click on [Finish] to start the job.

7. Monitor data processing during the KLAST job execution:

![Image of KLAST job execution monitoring]

8. **KLAST Results Viewer** is displayed when the comparison job is done:

![Image of KLAST results viewer]
5. Analysing results

The KLAST Results Viewer is divided into two parts:

- **a.** The above table is the best hit table: each row displays a query and its best hit, i.e. the hit with the highest score.
- **b.** The below table is the Hit List Viewer: it displays the whole set of hits reported for a single query.

The Hit List Viewer can be replaced by the Graphic Viewer, as follows:

On this view, each hit is represented as a yellow arrow mapped on the query sequence. You can click on an arrow to obtain hit details in the table presented just below the graphic view, and *vice versa.*
Table column content can be changed as needed, using the column selector located at the top right corner of the tables:

Choosing columns for the Best Hit Table

Choosing columns for the Hit List Viewer

6. Exporting results

Commands to export table content are accessible from the pulldown menu available at the top right corner of the tables (see above pictures).
7. KLAST Databank Manager

Introduction

The KLAST plugin comes with a Databank Manager (KDMS) that enables the automated installation of public and personal sequence databanks on your computer in order to run locally KLAST search jobs and sequence data retrieval tasks.

When databanks are installed locally, you can run search jobs in a very effective way and without connecting any remote servers. Regarding public data sources (e.g. databanks available from NCBI, EBI, etc.), KDMS is capable of achieving the following tasks in a fully automated way:

1. downloading data from remote servers
2. uncompressing and unarchiving data files
3. converting standard sequence data sources (Genbank, EMBL, Genpept, Uniprot, Silva and BOLD) to FASTA files
4. converting FASTA files into KLAST databanks
5. indexing standard biological classifications: Enzyme, Gene Ontology, NCBI Taxonomy, Pfam and Interpro

Steps 3 and 4 usually result in the creation of annotated KLAST databanks. Indeed, during this processing KDMS collects, if available, any term IDs from the following biological classification: Gene Ontology, Enzyme, Pfam, Interpro and NCBI Taxonomy. This information is introduced within the header of each sequence reported in the FASTA file created during step 3. Then, as a result of step 5, the KLAST databank contains information that can, in turn, be used by the KLAST plugin to produce KLAST results containing biological classification data. More on this on section 10.

It is worth noting that during the installation of a databank, you cannot run a KLAST job, and vice versa.

The databank repository

The very first step you have to check before any use of KDMS is the place where the databanks will be installed on your computer. By default, KDMS uses the directory called «KLAST_databanks» located within your home directory. If you prefer to use another location, simply use the command [Change Repository] located on the bottom right side of the KDMS main frame:

Please be advised to use a large disk space since databanks may require several hundred gigabytes of storage to be installed.

It is worth noting that KDMS can handle several repositories. Simply use the [Change Repository] command to switch between your various databank repositories.

Databanks are always installed within the active repository, i.e. the storage location displayed within the Change Repository dialogue box (see above picture).
**Overview of the Databank Manager interface**

Start the KLAST Databank Manager (KDMS) as follows: from the CLC Workbench's Toolbox, click twice on the «KLAST Databank Manager» item.

The interface of KDMS contains two panels. On the left side, you can see the panel Databanks Sources that let you choose which databank to install. On the right side, you can see the panel Installed Databanks which displays the list of databanks installed on your system; actually, that list is divided into three sub-lists: Klast Databanks, Sequence Annotations and Biological Classifications.

The Databank Manager provides two major ways to install databanks: from FTP servers (public institutes or in-house servers) and from your personal data files. In both cases, you will use the left panel of KDMS to start the installation job.

KDMS is capable of handling major sequence databank file formats: Genbank, Refseq, Embl, Genpept, Swissprot, TrEmbl, Fasta, Silva and BOLD. In addition to sequence data files, KDMS can install major biological classifications: NCBI Taxonomy, Gene Ontology, Enzyme Commission, Interpro and Pfam domains. Plain text as well as compressed (gzip) and archived (tar) files are accepted by KDMS.
Installing public databanks

Within the Databanks Sources panel, select the tab called Public Databanks. There, you can see a list of pre-configured databank descriptors. Each descriptor contains the relevant information enabling the software to install a particular databank: location of the remote server and the list of files to retrieve. You will see later on how to edit and add a new databank descriptor.

Usually, a databank descriptor relating to sequence databanks (Genbank, Swissprot, etc.) can be used to deploy locally two types of banks: a Klast databank to be used to run sequence comparison jobs and a sequence annotation bank (also called index) to be used for sequence data retrieval tasks.

Not only a databank descriptor provides the material to install a databank, it can also be used to get some information before the installation: availability of the remote server, number of files to download, size of the data to download and an estimation of the databank size on your disk storage after installation.

If you want to install a particular databank, or if you want to get its information without installing it, simply select the corresponding descriptor in the list (it is possible to select several descriptors), and click on the button called [Install]. KDMS will ask if you want to see the databank information before the installation: answer [Yes]. After a short period of time, during which KDMS queries the remote server, a dialogue box will appear to display the databank information. Click on the [Cancel] button to close the dialogue box if you do not want to start the install. On the other hand, if you click on the [Install] button, KDMS will provide you with the Installation Scheduler dialogue box. Use it to set up when you want to install the databank and click on the [Ok] button.

It is worth noting that KDMS has to be up and running when a databank installation has been scheduled. Otherwise, the task will not be executed. When an installation task is running, you can monitor it using the bottom part of the Databanks Sources panel. The Process sub-panel displays several progress bars that give you a simplified overview of what is going on. Use the Logs sub-panel to get a more detailed view of the installation processing.

You have two options if you want to install a databank not listed in the Public Databanks panel:

- option 1: create the appropriate databank descriptor
- option 2: install the databank using the Personal Databank panel

Using option 1 requires that sequence data files are available from FTP servers, and that you know which files to retrieve from these servers. Within KDMS, you create a new databank descriptor from an existing one: select an existing descriptor that is similar to the new databank you want to install, then click on the [Create] button. Finally, follow the instructions of the software. More information is available in the KDMS guide available at http://www.korilog.com/klast/kdms_user_guide.pdf.

Using option 2 requires you have the sequence data files already on your local computer. See section « Install personal databanks » for more information.

Installing public databanks: biological classifications

One of the major features of the KLAST plugin is its capability to introduce biological classification data within the KLAST results. To enable this feature, you have to install two types of databanks:

1. the biological classifications managed by KDMS; they are listed by the end of the Public databanks panel: Enzyme, GeneOntology_terms, InterPro_terms, NCBI_Taxonomy and Pfam_terms
2. a reference sequence databank that is annotated with such classifications; a very well known example is Uniprot_Swissprot

So, you have to install the above mentioned classifications and annotated reference sequence databanks before any attempt to produce KLAST results containing biological classification data. See section 10. for more information.
Installing personal databanks

Within the Databanks Sources panel, select the tab called Personal Databanks. There, you can see a panel that will enable you to provide various sequence files that will be converted to KLAST databanks.

To format a set of sequence files to be used with KLAST system, proceed as follows:

1. provide the set of file(s) to prepare. You can provide several sequence files at once, however be sure that they are all of the same sequence type (protein or nucleotide). Accepted sequence file formats are: Genbank, Refseq, Embl, Genpept, Swissprot, TrEmbl, Fasta, Silva or Bold. Plain text and gzipped are accepted (please note that gzipped files must have the .gz file extension).

2. select the sequence type and enter the name of the databank.

3. click on the [Install databank] button.

When the software terminates the installation of your databank, you will see it on the right panel of KDMS. Klast/Blast Databanks tab (prepared from Fasta files), or Sequence Annotations tab (prepared from annotated sequence files).

When you provide sequence files to KDMS, you should verify their content and especially check that sequence identifiers are unique and well formatted. Well formatted sequence IDs means that they follow the NCBI recommendations.

Here is an example of the installation of COI DNA barcoding data sets. Source files have been manually retrieved from the BOLD data server (http://www.boldsystems.org/index.php/datarelease), then KLAST Databank Manager has been setup as follows:

![Databank Sources](image)

It is worth noting that you have first to install the NCBI Taxonomy classification before installing taxonomy-based sequence data sets (Silva and BOLD).
8. KLAST parameters

Basic search parameters

KLAST parameters are provided by the KLAST Wizard, as follows:

Parameters from sections «Basic» and «Extended» are similar to BLAST ones. However, there are two minor differences. First, comparison method (or Program) is one of the following values:

<table>
<thead>
<tr>
<th>Program</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>KlastP</td>
<td>Search protein database using a protein query</td>
</tr>
<tr>
<td>KlastN</td>
<td>Search a nucleotide database using a nucleotide query</td>
</tr>
<tr>
<td>KlastX</td>
<td>Search protein database using a translated nucleotide query</td>
</tr>
<tr>
<td>tKlastN</td>
<td>Search translated nucleotide database using a protein query</td>
</tr>
<tr>
<td>tKlastX</td>
<td>Search translated nucleotide database using a translated nucleotide query</td>
</tr>
</tbody>
</table>

Then, there are two parameters to control how many hits and HSPs are reported in the results:

<table>
<thead>
<tr>
<th>Program</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max Hit/Query</td>
<td>Set the maximum number of hits for each query</td>
</tr>
<tr>
<td>Max HSP/Hit</td>
<td>Set the maximum number of HSPs for each hit</td>
</tr>
</tbody>
</table>
Optimizing search job using KLAST specific arguments

KLAST’s default configuration has been setup to provide an optimal ratio between speed and quality in order to produce results with quality similar to Blast. Even in such a configuration, you'll have great speedup factors.

However, depending on your needs you can enhance speed factors with little loss of quality in your results.

KLAST specific parameters are provided in section « KLAST algorithm configuration », and three of them are of interest:

<table>
<thead>
<tr>
<th>Program</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent of seeds to use (or seed-ratio)</td>
<td>Ratio of seeds to be used (see below). [1..100], default is 100. Decrease value to speedup algorithm with little loss of quality. Available for all methods but KlastN.</td>
</tr>
<tr>
<td>Threshold score</td>
<td>Ungapped threshold triggering a small gapped extension (see below). [25..127], default is 28 and 55 for protein-based and nucleic-based comparisons, respectively. Increase value to speedup algorithm with little loss of quality.</td>
</tr>
<tr>
<td>Maximum DB Size</td>
<td>Maximum allowed size (in bytes) for a databank. If greater, database is segmented (see below).</td>
</tr>
</tbody>
</table>

Fine tuning seed-ratio, threshold score and max-database-size may provide impressive acceleration of the KLAST comparison engine, with little loss of quality in the results. Carefully read the following sections.

Optimizing KLAST: sample recipes

In order to tune KLAST correctly, we always invite the users to try the software with sample data sets. When you need to compare large set of sequences, always start your work by comparing a small subset of your data. This way, you can check the parameters, the results and the speed of the software.

As an example, if you have to compare 300,000 sequences against NCBI nt, start your work by comparing 300 query sequences against NCBI nt using default KLASTn parameters. Then, fine tune it (see below the use of seed-ratio, max-database-size and threshold score) and check the results. As soon as your parameters are fine, go ahead with 3,000 and/or 10,000 query sequences, and check results and speed. If everything is fine, then run the full comparison.

Optimizing KLAST at runtime: using seed-ratio

When using KLAST for protein-based sequence comparisons, the algorithm can be speedup using the seed-ratio parameter. As stated earlier, KLASTp algorithm relies on a finite table of seeds; there are about 6,200 seeds for BLOSUM50 and BLOSUM62 matrices, whatever the input sequence databanks (for more information, see Reference [3]). During the comparison, KLAST orders seeds by occurrences, starting to process seeds producing the highest number of hits. So, it is possible to ask KLAST to use either the entire set of seeds to achieve a comparison, or a subset. This fine-tuning KLAST feature is achieved using the seed-ratio parameter, ranging from 1% to 100%. The highest seed-ratio you use, the highest sensitivity you get... the lowest seed-ratio you use, the highest speed you get with little loss in quality, as illustrated on this example:
Reducing number of seeds to use during a comparison still provides high results quality while dramatically reducing search time.

The seed-ratio parameter is available for KLASTp, KLASTx, tKLASTx and tKLASTn.

Optimizing KLAST at runtime: using threshold score

A second way to fine tune KLAST, and speedup the search, consists in using the 'threshold score' parameter. During a search, KLAST computes a score for each ungapped sequence alignment matching a query and a hit. As soon as this score is above the threshold, that alignment is retained for further processing. By default, this 'threshold score' is set to a small value (38 for protein comparisons, 55 for nucleotide comparisons) to let KLAST be as sensitive as possible. However, if you suspect that your query sequences may be closely related to the reference databank, you could increase the 'threshold score': KLAST can still produce high-quality results, but with an additional speedup.

As an example, when comparing 900 reads (500 nucleotides on average) against Silva SSU databank (740,000 sequences) on a 8 cores Intel-Xeon based computer, search time was 73 hours using TScore=55, but only 8 minutes using TScore=127; results were the same in terms of quality, i.e. we got the same best hit for each query in both results.

The TScore parameter is available for KLASTn, KLASTp, KLASTx, tKLASTx and tKLASTn. Its value is in the range 25..127 (default is 28 and 55 for protein-based and nucleic-based comparisons, respectively).

Optimizing KLAST at runtime: using max-database-size

Another way to fine tune KLAST, and again speedup the search, consists in using the max-database-size parameter. It sets the amount of bytes to reserve in RAM in order to load databank pages into memory. Indeed, during the comparison of query vs. subject databanks, KLAST automatically paginates databanks if they do not fit entirely into RAM. For that purpose, KLAST relies on the max-database-size parameter; when setting up that parameter, compare the amount of RAM you have in your computer with 'max-database-size x 8 x 2' (each databank index requires '8 x max-database-size' bytes, and you have two databanks). For instance, when using KLAST on a 32 Gb computer, increasing max-database-size from 5M (default value) to 100M may produce an additional speedup of 4x.

We advise you to setup the max-database-size value to enable the full load of the query databank into memory. For instance, if your query file sizes 12 Mb, then set the max-database-size parameter to 15000000 (15 Mb). Also, we do not recommend to set max-database-size to value exceeding 100000000 (1Gb) ; in such a case, if your query file is very big, let KLAST paginates the query, or run several KLAST jobs, each of them processing one partition of your query file.

The max-database-size parameter is available for KLASTn, KLASTp, KLASTx, tKLASTx and tKLASTn.
9. Filtering KLAST results

Introduction

A data filter, or simply a filter, is made of \( n \geq 1 \) rules. In turn, each rule is defined with a data accessor, an operator and a value. Several rules are combined with boolean operators ‘or’ & ‘and’.

Data accessors

A data accessor enables the access to an information contained in a KLAST result. They are very straightforward, as they correspond to standard sequence comparison tool results:

<table>
<thead>
<tr>
<th>Accessor name</th>
<th>Accessor name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hit Accession</td>
<td>HSP query from</td>
</tr>
<tr>
<td>Hit rank</td>
<td>HSP query to</td>
</tr>
<tr>
<td>Number of HSPs</td>
<td>HSP query frame</td>
</tr>
<tr>
<td>Hit definition</td>
<td>HSP query gaps</td>
</tr>
<tr>
<td>Hit identifier</td>
<td>HSP hit from</td>
</tr>
<tr>
<td>Hit Length</td>
<td>HSP hit to</td>
</tr>
<tr>
<td>HSP rank</td>
<td>HSP hit frame</td>
</tr>
<tr>
<td>Query coverage</td>
<td>HSP hit gaps</td>
</tr>
<tr>
<td>Hit coverage</td>
<td>HSP % of positives</td>
</tr>
<tr>
<td>HSP bit score</td>
<td>HSP % of gaps</td>
</tr>
<tr>
<td>HSP score</td>
<td>HSP % of identities</td>
</tr>
<tr>
<td>HSP E-Value</td>
<td>HSP alignment length</td>
</tr>
</tbody>
</table>

Data operators

This table lists the available operators you can use to compare values:

<table>
<thead>
<tr>
<th>Operator</th>
<th>Description</th>
<th>Accepted values</th>
</tr>
</thead>
<tbody>
<tr>
<td>==</td>
<td>Equal to</td>
<td>Numeric, string</td>
</tr>
<tr>
<td>!=</td>
<td>Not equal to</td>
<td>Numeric, string</td>
</tr>
<tr>
<td>&lt;</td>
<td>Less than</td>
<td>Numeric</td>
</tr>
<tr>
<td>&gt;</td>
<td>Greater than</td>
<td>Numeric</td>
</tr>
<tr>
<td>&lt;=</td>
<td>Less than or equal to</td>
<td>Numeric</td>
</tr>
<tr>
<td>&gt;=</td>
<td>Greater than or equal to</td>
<td>Numeric</td>
</tr>
<tr>
<td>::</td>
<td>Match</td>
<td>String vs. RegExp</td>
</tr>
<tr>
<td>!:</td>
<td>Do not match</td>
<td>String vs. RegExp</td>
</tr>
<tr>
<td>[]</td>
<td>Contain in the range</td>
<td>Numeric</td>
</tr>
<tr>
<td>][]</td>
<td>Not contain in the range</td>
<td>Numeric</td>
</tr>
</tbody>
</table>

All operators but [] and ][] are equality and relational operators; they are used to compare a left and a right operand.

Operators :: and !: are used for pattern matching to figure out whether or not the right operand is contained within the left operand; left operand is a string, but right operand is a regular expression.

By default, all string-based operators are ‘case sensitive’. However, the Filter Editor (see below) enables you to modify this behavior.
Data types

Data handled by Klast are quite simple: integer, real, string (of characters) and date.

Real numbers can be represented using scientific notation; e.g. 0.001 and 1e-3 are equivalent.

Dates are represented using the following format: YYYYMMDD, where YYYY, MM and DD are the year, month and day, respectively. MM is in the range [01-12] and DD is in the range [01-31]. Example: 20050512 stands for May 12th, 2005.

String can be a literal string, or it can be a regular expression (see below).

Creating a data filter

A filter is created with the Filter Editor during the setup of a KLAST job (step 4). On this panel, simply click on the [Apply Filter] check box to display the editor, then setup your rules as needed:

Each row corresponds to a single rule. You can add/remove rules using the two command [+]/[x] located on the right side of each rule.

It is worth noting that rules are combined with boolean operators 'or' and 'and' using the 'any' and 'all' items, respectively, displayed in the dropdown list located on top of the rules (see above picture).

The filter will be applied on the KLAST results automatically during the job execution. Then, the KLAST engine will only report hits validated by the filter rules.

Filtered data can be easily identified on the viewer, as illustrated on this figure:
Notice: it is always a good idea to setup and try a filter on a reduced data set in order to validate the rules with regard to the results. Then, go ahead and apply your filter on larger data sets.

To help you identifying what was the filter used during a particular KLAST job, simply use the job history panel:

Managing filters

The Filter Editor provides two commands, [Save Filter] and [Open Filter], to enable you to save and reopen a filter, respectively. This way, you can store and reuse your filters without writing them from scratch.
Using regular expressions

Basic concepts

A regular expression, often called a pattern, is an expression that describes a set of strings. They are usually used to give a concise description of a set, without having to list all elements. Alternatively, a pattern can be used to identify strings containing at least one of the possible strings defined by the pattern. In the field of bioinformatics, there is a well-known example of the usage of regular expressions: the Prosite patterns are nothing else than such expressions.

By the way of operators :: and !:, ngKLAST’s filtering system gives you the opportunity to use regular expressions. In the context of filtering KLAST data, these expressions are quite useful to locate a string within another. As an example, if you plan to locate in a KLAST result the database sequences for which the description field contains a particular word, a regular expression will help you, since it allows you to locate a misspelt word. Remember that operators == and != do an exact comparison of two strings: in no way they can be used to locate a particular string within another.

Using a regular expression in a filter

Using a regular expression in a filter's rule is quite easy: as soon as you use one of the operators :: or !:, the right operand (which is the value you specify for the comparison) is considered by ngKLAST as a regular expression.

Regular expression syntax

Very basic expressions

The simplest regular expression is a simple string: A, ATG, kinase, etc.

Alternation

A vertical bar separates alternatives. For example, gray|grey, can match gray or grey.

Grouping

Parentheses are used to define the scope and precedence of the operators. For example, gr(a|e)y, means gr follows by a or e, follows by y. It is equivalent to gray|grey and can also match gray or grey.

Case sensitivity

By default, regular expressions are case-sensitive. As a consequence the pattern atg cannot recognize the string ATG. Using the special construct (?i) allows to search for case-insensitive patterns. The (?i) modifier affects all characters to the right and in the same group, if any. For example in the pattern a(?i)t,g, only t is allowed to be case-insensitive, whereas in (?i)(atg) all characters are allowed to be case-insensitive. The pattern (?i)(atg) can match atg, aTg, ATg, ATG, etc.

Quantification

A quantifier after a character or group specifies how often consecutive occurrences of that preceding expression are allowed to occur. The available quantifiers are ?, *, and +.

The question mark indicates there is 0 or 1 of the previous expression. For example, colou?r matches both color and colour.

The asterisk indicates there are 0, 1 or any number of the previous expression. For example, "go*gle" matches ggle, gogle, google, gooogle, etc.
The plus sign indicates that there is at least 1 occurrence of the previous expression. For example, "go+gle" matches gogle, google, gooogle, etc. (but not ggle).

**Extended quantification**

The expression \{x,y\}, where x and y are numbers, can be used to define more precise quantification. For example \(T\{x\}\) means \(T\) exactly \(x\) times. \(T\{x,\}\) means \(T\) at least \(x\) times. \((ATG)\{x,y\}\) means \(ATG\) at least \(x\) but not more than \(y\) times.

**Character classes**

Character classes are defined using [ and ] to match a single character that is contained within the brackets. For example, \[abc\], matches a, b or c at any given position of a string. Expression \[^abc\] matches any character except a, b, or c (negation). Expression \[a-zA-Z\] matches a through z or A through Z, inclusive (range).

**Predefined character classes**

The following table gives some useful predefined character classes.

<table>
<thead>
<tr>
<th>Construct</th>
<th>Matches</th>
</tr>
</thead>
<tbody>
<tr>
<td>.</td>
<td>Any character</td>
</tr>
<tr>
<td>\d</td>
<td>A digit: [0-9]</td>
</tr>
<tr>
<td>\D</td>
<td>A non-digit: [^0-9]</td>
</tr>
<tr>
<td>\w</td>
<td>a word character: [a-zA-Z_0-9]</td>
</tr>
<tr>
<td>\W</td>
<td>a non-word character: [^\w]</td>
</tr>
</tbody>
</table>

**Special characters**

\(^\) matches the beginning of a string. \(\$\) matches the end of a string. For example, \(^[hc]\)at matches hat and cat but only at the beginning of a string, and \([hc]\)at\(\$\) matches hat and cat but only at the end of a string.
10. Adding biological classification data to KLAST results

Preparing annotated KLAST results using biological classifications

In order to introduce biological classification data within KLAST results you have to install at least:

1. the biological classifications managed by KDMS; they are listed by the end of the Public databanks panel: Enzyme, GeneOntology_terms, InterPro_terms, NCBI_Taxonomy and Pfam_terms
2. a reference sequence databank that is annotated with such classifications; a very well known example is Uniprot_Swissprot

If not already done, simply proceed as follows:

1. start the KLAST Databank Manager tool from CLC Workbench's Toolbox
2. on the left side of KDMS panel, scroll down into the list of databank descriptors to select the five descriptors targeting the biological classifications managed by KDMS:

   ![Databank Manager Screenshot]

3. click on the [Install] command
4. as soon as biological classifications are installed, it is time to install an appropriate reference databank; for instance, it can be Uniprot_Swissprot:

   ![Databank Manager Screenshot]

5. Again, click on the [Install] command to install the reference databank
Now, you can setup a KLAST job as usual. Two steps are of interest:

1. during the selection of the target databank, use a reference databank containing biological classifications data (see step 4, above):

![KLAST on local computer](image1)

2. Then, during the selection of post-processing tasks, click on «Retrieve biological classification data»:

![KLAST on local computer](image2)

During the KLAST job, and more precisely during the «results retrieval step», the KLAST annotation system will automatically introduce the classification data into the hits reported in the KLAST results.
Finally, you can display the information, as illustrated on the following screenshot:

On the first table, you can add the « Organism » column which enables a direct overview of NCBI Taxonomy data retrieved within the reference databank (Swissprot in this example).

To display the various classification terms (Enzyme, GO, etc.), you can switch to the « Graphic Viewer »: a « source » feature is associated to every hit when such data is available. In turn, classification data will be displayed within the « Feature Table » displayed at the bottom of the « Graphic Viewer ».
Exporting annotated KLAST results with biological classifications

The current release of the KLAST plugin enables an interactive view of the biological classification data on a «per query» basis. Now, if you want to get a global overview of all available data, the software allows you to export the data as a Fasta or Tabular file. The former export command will save in a file each query sequence along with the classification data. The latter export command saves in a file queries and classification data in a more compact way.

Simply use the «Best Hit table» command tool\(^1\) to access the export commands:

As you can see, two commands are available: “Export Hits” and “Export Queries”.

Both commands export data for the selected rows of the table, so you may have to use the «Select all» command (available in the same pulldown menu) before using the export commands; you can also only select relevant rows using the mouse, then export data for that selection.

\(^1\) It is the little white arrow on a blue circle, located on the top right corner of the table
11. Citing KLAST

Use the following reference to cite KLAST in your publications: « Nguyen VH, Lavenier D (2009) PLAST: parallel local alignment search tool for database comparison. BMC Bioinformatics. 10:329 ».

12. References


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