Making Request to Database

To make a request to a local BLAST database do the following:

- If you're using BLAST open Tools BLAST BLAST Search.
- If you're using BLAST+ open Open Tools BLAST BLAST+ Search.

If there is a sequence opened you can also initiate the request to a local BLAST database from the Sequence View.

- If you're using *BLAST* select the *Analyze Query with BLAST* item in the context menu or in the *Actions* main menu.
- If you're using BLAST+ select the Analyze Query with BLAST+ item in the context menu or in the Actions main menu.

The Request to local BLAST database dialog will appear:

8 Request to Local BLAST Database	<u>ହ</u> <u>x</u>
General options Advanced options Extension	n options
Select search:	blastn Search for short, nearly exact matches
Expectation value	10.000000 ি 🔲 Megablast
Culling limit:	20 🗟 💿 Both strands 🔘 Direct 💿 Complement
Database path:	Select
Base name for BLAST DB files:	a database file
✓ Save annotation(s) to	
Existing table	• E
Create new table	
Use auto-annotations table	
 Annotation parameters 	
Group name <auto></auto>	*
	Number of CPUs being used 8
	Restore to default Search Cancel Help

The following general options are available:

Select search - here you should select the tool you would like to use. If the query sequence is a nucleotide sequence then blastn, blastx and tblastx it ems are available. For a protein sequence the items are blastp and tblastn.

Expectation value - this option specifies the statistical significance threshold for reporting matches against database sequences. Lower expect thresholds are more stringent, leading to fewer chance matches being reported.

Culling limit - the maximum number of hits that will be shown (not equal to number of annotations). The maximum available number is 5000.

Search for short, nearly exact matches - automatically adjusts the word size and other parameters to improve results for short queries.

Megablast - select this option to compare query with closely related sequences. It works best if the target percent identity is 95% or more, but it is very fast.

Database path - path to the database files.

Base name for BLAST DB files - base name for the BLAST database files.

You can see the description of the annotation saving parameters here.

The following advanced parameters are available:

% Request to Local BLAST Database	ନ <u>୍</u>
General options Advanced options Extension options	
Word size 7 🗘 Gap costs 2 2	•
Match scores 1 -3	•
Filters	Masks
Low complexity filter	Mask for lookup table only
Human repeats filter	Mask lower case letters
	Restore to default Search Cancel Help

Word size - the size of the subsequence parameter for the initiated search.

Gap costs - costs to create and extend a gap in an alignment. Increasing the Gap costs will result in alignments which decrease the number of Gaps introduced.

Match scores - reward and penalty for matching and mismatching bases.

Filters - filters for regions of low compositional complexity and repeat elements of the human's genome.

Masks for lookup table only — this option masks only for purposes of constructing the lookup table used by BLAST so that no hits are found based upon low-complexity sequence or repeats (if repeat filter is checked).

Mask lower case letters — with this option selected you can cut and paste a FASTA sequence in upper case characters and denote areas you would like filtered with lower case.

The view of the Advanced options tab depends on the selected search. For the blastn search it looks like on the picture above. When the blastn search is selected in the general options, the view of the Advanced options tab is the following:

ℜ Request to Local BLAST Data	itabase	? X
General options Advance	ed options Extension options	
Word size	3 Gap costs 11 1	•
Threshold	12.00	
Matrix	BLOSUM62 Service plain	
	D or d: default (equivalent to 2)	•
Filters	Masks	
Low complexity filter	Mask for lookup table only Mask lower case letters	
Human repeats filter	Mask lower case letters	
	Restore to defaultSearchCancel	Help

As you can see there is no Match scores option, but there are Threshold, Matrix, Composition-based statistics and Service options.

Threshold - threshold for extending hits.

Matrix — key element in evaluating the quality of a pair-wise sequence alignment is the "substitution matrix", which assigns a score for aligning any possible pair of residues.

Service — blastp service which needs to be performed: plain, psi or phi.

Composition-based statistics - composition-based statistics.

When the tblastx search is selected in the general options, the view of the Advanced options tab is the following:

Request to Local BLAST Database	ି ୪
General options Advanced options Extension options	
Word size 3	
Threshold 13.00	
Matrix BLOSUM62 Service plain	▼
Filters	Masks
Low complexity filter	Mask for lookup table only
Human repeats filter	Mask lower case letters
	Restore to default Search Cancel Help

The following extension options are available:

% Request to Local BLAST Database		? X
General options Advanced options Extension options		
X dropoff value (in bits)		
For gapped alignment	30	
For ungapped extensions	20.00	
For final gapped alignment	100	
Multiple Hits Window Size	0	
Perform gapped alignment		
	Destans to default	Uala
	Restore to default Search Cancel	Help

For gapped alignment - X dropoff value (in bits) for gapped alignment.

For ungapped alignment - X dropoff value (in bits) for ungapped alignment.

For final gapped alignment - X dropoff value (in bits) for final gapped alignment.

Multiple hits window size - multiple hits window size.

Perform gapped alignment - performs gapped alignment.