

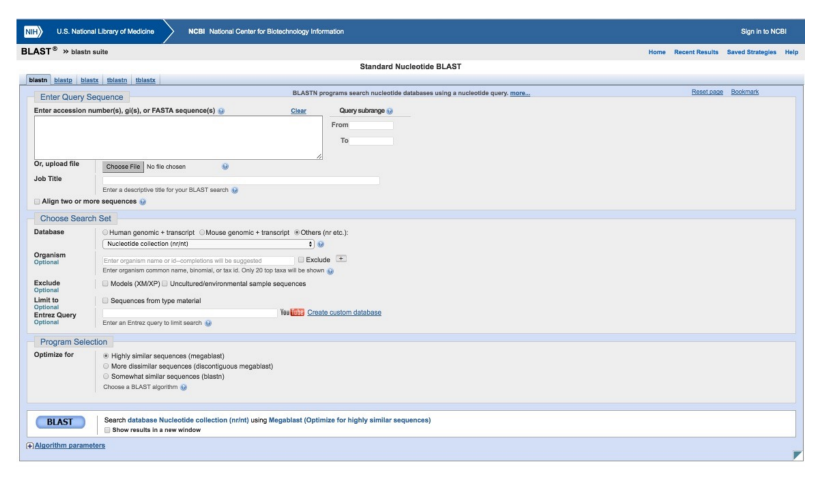


Common Themes

- Gene and functional pathway presence/absence
- Identification, alignment of orthologs
- Structural analysis and comparison



Basic Local Alignment Search Tool (BLAST)





The Core BLAST Programs

Program	Query Sequences	Database Sequences	Alignment
blastn	nucleotide	nucleotide	nucleotide
blastp	amino acid	amino acid	amino acid
blastx	nucleotide	amino acid	amino acid
tblastn	amino acid	nucleotide	amino acid
tblastx	nucleotide	nucleotide	amino acid



Downstream Applications

- Identify orthologs
 - Reciprocal BLAST
 - OrthoMCL
- Annotation and Pathway Analysis
 - e.g., BLAST2GO
- Many others



Advantages of Running BLAST Locally

- Customize databases with unpublished sequences
- Run thousands of simultaneous searches
- Integrate into custom scripts and pipelines



Parsing BLAST Output

```
BLASTN 2.2.30+
Database: bacterial-tRNAs.fas
          24,746 sequences; 1,923,058 total letters
Query= 1
Length=74
                                                                Score
Sequences producing significant alignments:
                                                               (Bits)
                                                                      Value
 Microcystis_aeruginosa_NIES_843_chr.trna42-AspGTC (409004-40893...
                                                                 111
                                                                       2e-26
 Anabaena variabilis ATCC 29413 chr.trna22-AspGTC (6126075-61260...
                                                                 111
                                                                       2e-26
> Microcystis_aeruginosa_NIES_843_chr.trna42-AspGTC (409004-408931)
Asp (GTC) 74 bp Sc: 75.49
Length=74
Score = 111 bits (122), Expect = 2e-26
Identities = 69/74 (93%), Gaps = 0/74 (0%)
Strand=Plus/Plus
Query 1
          GGGATTGTAGTTCAATTGGTTAGAGCACCGCCCTGTCAAGGCGGAAGCTACGGGTTCGAG
          GGGATTGTAGTTCAATTGGTTAGAGCACCGCCCTGTCACGGCGGAAGTTGCGGGTTCGAG
Sbjct 1
Query 61 TCCCGTCAGTCCCG 74
Sbict 61 CCCCGTCAATCCCG 74
```



Organization of BLAST Output

RESULTS – One results block for each sequence in the original query file

- HITS Within each results block, one hit block for each sequence in the database that produced a significant alignment
 - HSPs Within each hit block, one high-scoringpair block for each significant local alignment to that hit



Parsing BLAST Output



http://bioperl.org/howtos/SearchIO HOWTO.html



Parsing BLAST Output – BioPerl Results Information

Object	Method	Example	Description
Result	algorithm	BLASTX	algorithm string
Result	algorithm_version	2.2.4 [Aug-26- 2002]	algorithm version
Result	query_name	gi	20521485
Result	query_accession	AP004641.2	query accession
Result	query_length	3059	query length
Result	query_description	Oryza sativa 977CE9AF checksum.	query description
Result	database_name	test.fa	database name
Result	database_letters	1291	number of residues in database
Result	database_entries	5	number of database entries
Result	available_statistics	effectivespaceused dbletters	statistics used
Result	available_parameters	gapext matrix allowgaps gapopen	parameters used
Result	num_hits	1	number of hits
Result	hits		List of all Bio::Search::Hit::GenericHit objects for this Result
Result	rewind		Reset the internal iterator that dictates where next_hit() is pointing, useful for re-iterating through the list of hits



Parsing BLAST Output – BioPerl Hit Information

Object	Method	Example	Description	
Hit	name	gb	443893	
Hit	length	331	Length of the Hit sequence	
Hit	accession	443893	accession (usually when this is a Genbank formatted id this will be an accession number - the part after the gb or emb)	
Hit	description	LaForas sequence	hit description	
Hit	algorithm	BLASTX	algorithm	
Hit	raw_score	92	hit raw score	
Hit	significance	2e-022	hit significance	
Hit	bits	92.0	hit bits	
Hit	hsps		List of all Bio::Search::HSP::GenericHSP objects for this Hit	
Hit	num_hsps	1	number of HSPs in hit	
Hit	locus	124775	locus name	
Hit	accession_number	443893	accession number	
Hit	rewind		Resets the internal counter for next_hsp() so that the iterator will begin at the beginning of the list	



Parsing BLAST Output – BioPerl HSP Information

Object	Method	Example	Description
HSP	algorithm	BLASTX	algorithm
HSP	evalue	2e-022	e-value
HSP	expect	2e-022	alias for evalue()
HSP	frac_identical	0.884615384615385	fraction identical
HSP	frac_conserved	0.923076923076923	fraction conserved (conservative and identical replacements aka "fraction similar")
HSP	gaps	2	number of gaps
HSP	query_string	DMGRCSSG	query string from alignment
HSP	hit_string	DIVQNSS	hit string from alignment
HSPt	homology_string	D+ + SSGCN	string from alignment
HSP	length('total')	52	length of HSP (including gaps)
HSP	length('hit')	50	length of hit participating in alignment minus gaps
HSP	length('query')t	156	length of query participating in alignment minus gaps
HSPt	hsp_length	52	Length of the HSP (including gaps) alias for length('total')
HSPt	frame	0	\$hsp->query->frame,\$hsp->hit->frame
HSP	num_conserved	48	number of conserved (conservative replacements, aka "similar") residues
HSP	num_identical	46	number of identical residues
HSPt	rank	1	rank of HSP
HSP	seq_inds('query','identical')	(966,971,972,973,974,975)	identical positions as array
HSP	<pre>seq_inds('query','conserved-not- identical')</pre>	(967,969)	conserved, but not identical positions as array
HSP	seq_inds('query','conserved')	(966,967,969,971,973,974,975,	conserved or identical positions as array
HSP	seq_inds('hit','identical')	(197,202,203,204,205,)	identical positions as array
HSP	<pre>seq_inds('hit','conserved-not- identical')</pre>	(198,200)	conserved not identical positions as array
HSP	seq_inds('hit','conserved',1)	(197,202-246)	conserved or identical positions as array, with runs of consecutive numbers compressed



Parsing BLAST Output – BioPerl HSP Information (Cont.)

HSPt	score	227	score
HSP	bits	92.0	score in bits
HSP	range('query')	(2896,3051)	start and end as array
HSP	range('hit')	(197,246)	start and end as array
HSP	percent_identity	88.4615384615385	% identical
HSP	strand('hit')	1	strand of the hit
HSP	strand('query')	1	strand of the query
HSP	start('query')	2896	start position from alignment
HSP	end('query')	3051	end position from alignment
HSP	start('hit')	197	start position from alignment
HSP	end('hit')	246	end position from alignment
HSP	matches('hit')	(46,48)	number of identical and conserved as array
HSP	matches('query')	(46,48)	number of identical and conserved as array
HSP	get_aln	sequence alignment	Bio::SimpleAlign object
HSPt	hsp_group	Not available in this report	Group field from WU-BLAST reports run with -topcomboN or -topcomboE specified
HSP	links	Not available in this report	Links field from WU-BLAST reports run with -links showing consistent HSP linking



Exercise

https://dbsloan.github.io/TS2022/exercises/local_blast.html

- Make BLAST databases
- Run local BLAST searches
- Parse BLAST output with BioPerl
- Make dot-plot comparing two bacterial genomes in R using parsed BLAST output