

User Manual

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Introduction and Overview

- yes, the important theory section is currently empty; forthcoming!
- take a lot of this from the original report I made

Compiling

To compile Decisivator, uncompress the .tgz file using:

```
tar -xvzf Decisivator.tqz
```

Now, navigate to the src directory and type:

make

This will create the executable 'Decisivator'. That's it! Put this wherever in your system you prefer.

If you have troubles compiling (say, when moving from one computer to another), type:

```
make clean
```

and then continue with the make command. (If you have trouble getting this to work on your system, please complain to be at: josephwb@uidaho.edu).

Input Files

Multiple user files may be passed in to Decisivator, but this must minimally include a file containing information regarding which taxa possess data for each partition. Normally a partition will refer to a sequenced genetic fragment (e.g. DNA/RNA or amino acids), but this could refer to any kind of character that might be used for phylogenetic reconstruction (e.g. morphology). Each independent set of characters should be designated as a separate partition: for genetics this will likely involve contiguous genes; for morphology, each character should be designated as a separate partition (unless characters can be identified as *non*-independent, in which case characters should be partitioned together). This taxon-partition information will most often take the form of a Nexus-formatted data file. Regardless of data type, partition information is gleaned from CHARSET declarations. CHARSETs can either be contiguous or interval in nature:

```
CHARSET ziggy = 1-2043;
CHARSET iggy = 2044-2099\3; [e.g. for codon data]
CHARSET joey = 2100-2100; [e.g. morph. character]
```

At the moment, the formatting of these CHARSET declarations is rather Draconian. For example, there should be no spaces around the '-' and '\'. More annoyingly, Decisivator does not yet support referential CHARSET declarations. For example:

```
CHARSET nineteenseventytwo = iggy david;
```

Will cause Decisivator to die a horrible death. If you wish to use referential CHARSET declarations, first rearrange your data into contiguous format (sorry).

A second option exists for passing in taxon-partition information. This is a 'legacy' format for which support will likely not continue, although in some instances (e.g. large/sparse matrices) it may be easier to construct than a Nexus-formatted file. This second option involves a tab-delimited file. The first row should give partition names, and the first column should give taxon names. Existence of information for a taxon-partition is indicated by a '1', non-existence by a '0'. Below is a dummy example:

Gene_1	Gen	.e_2	Gen	e_3	Gene_4
Taxon_1	1	_0	1	_1	_
Taxon_2	1	1	0	1	
Taxon_3	0	1	1	1	
Taxon_4	0	0	0	1	
Taxon_5	1	1	1	0	
Taxon_6	0	0	1	0	
Taxon_7	1	0	1	0	
Taxon_8	0	1	0	0	
Taxon_9	0	0	1	1	

In addition to a required data file, several optional files are possible. Perhaps most importantly is a file containing user-tree(s). These trees must be Nexus-formatted, and may (but not necessarily) utilize a translation table. The tree file may contain multiple trees; this may involve possible trees that are of interest as hypotheses, or it may involve a distribution of trees (i.e. posterior or bootstrap distribution). In the case of a distribution of trees, conventional options (e.g. burnin and thinning) are available (see below).

The last type of file that can be passed in indicates 'weights', one file for taxa and one for partitions. These weights are used to help determine optimal targeted data acquisition (e.g. targeted sequencing). By default, all taxa and all partitions have a weight of 1.0. Weight files, if present, need only identify those taxa or partitions which receive a weight *not* equal to 1.0. Below is an example weight file for taxa:

Taxon	Weight
Taxon_3	5
Taxon_7	3
Taxon_8	0.5
Taxon 2	0.1

Note that a header line <u>is</u> expected.

Usage Options

While analysis conditions are accessed via a interactive menu prompt, several variables can be passed in to Decisivator via the command-line. To see these options, type:

./Decisivator -h

which will print out a slew of information to the screen:

Decisivator version 0.47
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Program description: Calculates phylogenetic 'decisiveness' sensu Sanderson and Steel.

To compile, type the following in a unix prompt:

make

To run, type:

```
./Decisivator [-d data_file] [-m taxon-gene_matrix] [-t
tree_file] [-b burnin] [-n thinning]
    [-w taxon weights] [-l locus weights]
```

where:

'data_file' is a simple 'vanilla' Nexus file containing sequences and defined CHARSETs.

- PLEASE NOTE! Only simple CHARSETs are currently supported.
- e.g. contiguous (X-Y) or interval (e.g. codon: X-Y\3) data are fine.
- CHARSET referencing is NOT allowed at present (but will be!)
- 'taxon-gene_matrix' is a table listing taxa (rows) and genes (columns). This is a legacy format.
- '1' indicates cell has been sequenced, while '0' indicates it has not.

First row should give locus names. First column should give taxon names.

'tree_file' contains user tree(s) in Nexus format to evaluate decisiveness upon.

'burnin' is the number of trees to ignore. Only makes sense with a distribution of trees.

'thinning' is the interval between sampling trees (i.e. where every nth tree sample will be retained).

Only makes sense with a distribution of trees.

'taxon_weights' is a two-column (taxon, weight; with headers) file listing weights for taxa based on some arbitrary accessibility criterion.

'locus_weights' is the analogous two-column (locus, weight; with headers) file for locus weights.

NOTE: The taxon and locus weight files need not be complete. All weights are 1.0 by default.

Enter only weights which should be changed from the default.

For a description of the various input files, see the 'Input Files' section above.

If a distribution of trees is being passed in, it might not be of interest to analyze *every* tree, but instead a sampling of this distribution. Decisivator allows both 'burnin' (where the first N trees are discarded) and 'thinning' (where only every N tree is retained).

Program Menu Options

Once in the program itself, several options are available:

Programs options:

[A]dd virtual taxon-character(s)

[M]erge taxa (i.e create a chimeric taxon)

[E]xclude taxa

[D]elete partition(s)

[P]rint current matrix to the screen

[C]alculate partial decisiveness using random trees

[T]est of complete decisiveness

[I]nvestiagte decisiveness on a provided user-tree

[S]ummarize current status

[W]rite current matrix

[R]evert to original matrix

[Q]uit

The Add virtual taxon-character(s) option allows a user to investigate how hypothetical data collection influences matrix decisiveness.

The Merge taxa option allows one to create a chimeric taxon. This can be useful for closely related taxa which, alone, have limited data, but combined have more partition-specific data. This will be useful for deep relationships (where the variance in tip values is much less than the variance between clades), but should not be done for 'shallow' trees.

The Exclude taxa option includes many sub-options for removing taxa according to some criterion dealing with missing data. The sub-options are:

Exclude:

```
Taxa by [I]ndex
Taxa by [N]ame
Taxa missing [E]xactly N partitions
Taxa missing N or [M]ore partitions
Taxa possessing [O]nly N partitions
Taxa possessing N or [F]ewer partitions
Taxa possessing data for only a [S]pecific partition
Taxa exhibiting minimal [P]artition overlapping
[B]ack to main menu
```

Most of these options are self-explanatory. 'Index' refers to the position of the taxon in the alignment (e.g. the 13th taxon). 'Partition overlap' is an index that quantifies the number of taxa with which a particular taxon shares partition information. Rationale: taxa with high 'overlap' will satisfy more taxon quartets, and so contribute more to decisiveness. On the other hand, taxa with low 'overlap' feature in few quartets, and so are less pivotal to decisiveness. In the best scenario, taxon X shares (across all partitions) a partition with all N-1 taxa. This *could* be completely decisive, but not necessarily. Take the following dummy matrix (partition names excluded for clarity):

Α	1	1	0	0	1
В	0	1	0	0	0
С	0	0	0	0	1
D	1	0	0	0	0
E	0	0	0	0	1
F	0	1	1	0	0
G	0	0	1	0	0
H	0	0	1	0	0
I	0	1	1	0	0
J	0	0	1	0	0
K	1	0	0	0	0
L	0	0	0	0	1
M	0	0	0	1	0
N	0	1	0	0	1
0	1	0	0	0	0

```
Ρ
        1
                  0
                          0
                                    0
                                            0
                  0
                          0
                                    0
Q
        1
                                             0
        0
                  0
                          1
                                    1
                                             0
R
S
        0
                          0
                                    1
                  0
                                             0
Т
        0
                  0
                          0
                                    1
                                             0
        0
U
                  0
                          1
                                    1
                                             0
V
        1
                  0
                          0
                                    1
                                             0
        0
                          0
                                    1
W
                  0
                                             0
        0
                  0
                          0
                                    1
                                             0
X
        0
                  1
                          0
                                    0
                                             0
Υ
                                    0
\mathbf{Z}
        1
                  0
                          0
                                             0
```

Given that we are dealing with 26 taxa, a particular taxon can potentially share data with at most 25 other taxa. If we walk through the matrix, we come up with the following overlap counts:

```
Overlap
           Shares a gene with:
Α
     15
           B, C, D, E, F, I, K, L, N, O, P, Q, V, Y, Z
В
     5
           A, F, I, N, Y
С
     4
           A, E, L, N
D
     7
           A, K, O, P, Q, V, Z
Ε
     4
           A, C, L, N
F
     10
           A, B, G, H, I, J, N, R, U, Y
G
     6
           F, H, I, J, R, U
Η
     6
           F, G, I, J, R, U
Ι
     8
           F, G, H, J, N, R, U, Y
J
     6
           F, G, H, I, R, U
K
     7
           A, D, O, P, Q, V, Z
L
     4
           A, C, E, N
М
     8
           R, S, T, U, V, W, X, Y
Ν
     8
           A, B, C, E, F, I, L,
0
     7
           A, D, K, P, Q, V,
Ρ
     7
           A, D, K, O, Q, V,
     7
           A, D, K, O, P, V,
Q
                              \mathbf{z}
           F, G, H, I, J, M, S, T, U, V, W, X
R
     12
S
     7
           M, R, T, U, V, W,
                              X
Т
     7
           M, R, S, U, V, W, X
           F, G, H, I, J, M, R, S, T, V, W, X
U
     12
V
     10
           A, D, K, R, S, T, U, W, X, Z
W
     7
           M, R, S, T, U, V, X
X
     7
           M, R, S, T, U, V, W
     5
Υ
           A, B, F, I, N
\mathbf{Z}
     7
           A, D, K, O, P, Q, V
```

As an example, let us focus on taxon A. Taxon A only has data for partitions 1, 2, and 5, so only consider these partitions for this taxon. Taxon A only shares partition information with taxa B, C, D, E, F, I, K, L, N, O, P, Q, V, Y, Z, but no others. Taxon A thus has an 'overlap' of 15 (i.e. do not count sharing with taxon N twice). This is the highest overlap,

so we definitely want to keep it. Taxa C, E, and L have an overlap of only 4. Throwing out these taxa will leave a more overly-decisive matrix. We can therefore use minimal overlap as a criterion for throwing out taxa. Decisivator supports two exclusion criteria based on 'overlap':

Exclude [M]inimally overlapping taxa (i.e. just the worst)
or implement [T]hreshold

That it, exclude the taxon (or taxa, in the case of a tie) that exhibits the worst overlap, or exclude *all* taxa below a particular threshold (chosen by the user). Use of the threshold criterion tends to yield matrices with 1) more taxa, 2) lower coverage, but 3) higher decisiveness than naïve exclusion practices (such as excluding all taxa with information for only a single partition).

The Delete partition(s) option probably is not useful, as deleting a partition cannot possibly increase decisiveness, although it can be informative as to the relative influence on decisiveness of different partitions. The option is mainly used for the situation where more CHARSETs exist in a file than are to be analyzed (say, overlapping partitions or CHARSETs grouped according to some criterion).

The Print current matrix to the screen option is meant only to double-check that matrix changes are recorded correctly.

The Calculate partial decisiveness using random trees option is the meat-and-potatoes function of Decisivator. Ideally it would be preferable to know whether a particular matrix is decisive *for all possible trees*. However, in practice this might not be possible if taxon sampling is extensive enough, as there are N choose 4 quartet checks that must be made (see Introduction and Overview above). The way to get around this is to instead analyze a *large number of random trees*. In many ways, this test is much more useful than testing for complete decisiveness (see below), as this particular metric tells one *how decisive* (or, if you are a pessimist, *how indecisive*) a particular matrix is.

Decisivator offers two flavours of partial decisiveness:

```
Calculate partial decisiveness:
[T]ree-wise
[B]ranch-wise
```

These are very different takes on phylogenetic decisiveness. The first option gives the estimated proportion of trees for which the given matrix is decisive. This analysis proceeds very quickly, as only a single satisfaction of a given edge need be found; subsequent satisfactions of the edge are not considered. Furthermore, as it only take a single *unsatisfied* edge to make a given tree indecisive; Decisivator will bail on the current tree and move on to the next one. The second option gives the estimates proportion of decisive *branches* across random tree for which the given matrix is decisive. This analysis is considerably slower, as for each edge all possible relevant taxon quartets

are investigated for the purpose of determining, edge-by-edge, the proportion of taxon quartets that satisfy the edge. These decisiveness scores can inform targeted sampling, and can be correlated with nodal support values, etc. These two decisiveness numbers can vary, as it only takes a single indecisive branch to make an entire tree indecisive.

The Test of complete decisiveness option checks to see if the current matrix is decisive for *all possible trees*. As mentioned above, this can take a large (potentially prohibitive) amount of time. The outcome of this test is rather limited: either a matrix is decisive or it is not (and a matrix that is 99.9999999% partially decisive is simply *not* completely decisive). This option will thus be less useful than the partial decisiveness analogue above, although it may be of use to users with nearly completely decisive matrices hoping to make it wholly decisive.

Beyond the test for partial decisiveness, the Investigate decisiveness on a provided user-tree option may be the most useful. Here, a user may pass in a tree or distribution of trees for decisiveness analysis. Tree distributions may undergo burnin and thinning via command-line options (see above). Note that all such tree distributions will necessarily contain decisive trees (or they wouldn't have been inferred in the first place!). Much more useful is comparing the branch-wise decisiveness of tree representing distinct phylogenetic hypotheses. Decisivator calculates these numbers, and can attach them to the tree(s) for viewing in a tree visualization program (e.g. FigTree ¹). The results are also printed out to a bipartition table similar to those from PAUP*² or MrBayes³:

Bipartition Decisiveness Scores:

12345678	Freq	Poss	용
**	8	8	100
* . *	8	8	100
**.*.*	12	12	100
****	8	8	100
*****	5	5	100

These numbers may be useful in determining, say, whether a particular branch is really more highly supported or simply that more data are present with respect to that branch. Trees from these analyses can also be printed out to file with decisiveness annotations.

Summarize current status simply prints summary information to the screen (the number of 'reference taxa', the number of trees in memory, matrix coverage, and results of the various decisiveness tests).

The Write current matrix option allows the user to export a modified matrix in either Nexus or Phylip format. Obviously 'virtual taxon-characters' cannot be included; this function is mainly use for exporting matrices where taxa (or, possibly, partitions) have been excluded, so that matrix phylogenetic-sensitivity can be explored.

Finally, the Revert to original matrix option is self-explanatory: go back to the original taxon-partition matrix and original weights, and reset all decisiveness scores.

References

- 1. FigTree v.1.0 (Available from the author (http://evolve.zoo.ox.ac.uk/software.html?id=figtree), 2006).
- 2. PAUP*. Phylogenetic Analysis Using Parsimony (*and Other Methods) v.4 (beta 10). (Sinauer Associates, Sunderland, Massachusetts, 2003).
- 3. Ronquist, F. & Huelsenbeck, J. P. MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* **19**, 1572-1574 (2003).