SPIMAP: Species Informed Maximum A Posteriori Gene Tree Reconstruction

Documentation for the SPIMAP software package April 8, 2011

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citation:

Matthew D. Rasmussen and Manolis Kellis. *A Bayesian Approach for Fast and Accurate Gene Tree Reconstruction*. Molecular Biology and Evolution. 2010. doi: 10.1093/molbev/msq189

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```
>KLLA0C08239g
ATGAGTCTCAAACGTGTAGTTGTCACTGGTCTTGGGGCCTACACGCCCCTTGGTTCTACAGTTTCAAAGTCTTGGGCAGG
TTTGCTT—GCTGCTAAGCAATCACTAATACCCTTAGATGCTTTCTACAACAGAGAA—GACTTTGCAAAAAGTGAAAA
AGTTGGTCCCACTAGATACAGCAGTGAGTAGGTTACAT—
>ADL072C
—ATGCATCCCCGAGTGGTCGTGACCGGCATTGGGTGCTATACTCCTCTGGGGCCCGTCGCTAGCCCAGTCTTGGAAGGA
GCTGTTG—CGAGGGACGACGCCTTGTCAGGCTGCAAGATCTGGCAGAGTACCAGGGCCGATTACAAACCACTGTCGA
GGCTTATATCCGGTGATCTTCGAGTCGGGAAAGTTGGATTTGAG—
>kwal_5828
—ATGACTTCCAGAGTCGTGTTACTGGGCTTGGTGCTATCACTCCACTTGGGAGGACTGTTTCCGAGTCATGGAGGC
TTTATTG—GCAGGCAAGTCCGGAATTCGTCCCATTCCCGATCTTCCC
—AATGCTAAAAGCTACGAAG
GACACTGTCCTGCATCTGTTGCCGTTGCAGACATTCCTGATTTC—GATCCA
```

Figure 1: **Example *.align file.** Three gene DNA sequences are given, each with 240 sites.

1 File formats

1.1 Sequence alignment format (*.align)

SPIMAP uses the FASTA file format (http://en.wikipedia.org/wiki/FASTA_format) for sequences alignments. The file extension is not important and many different extensions are in common use (*.fa, *.mfa, *.fasta, *.align).

Each line starting with ">" is used to indicate a gene name (Figure 1). The entire line after the ">" is used as the gene name. The aligned sequence of is then given on the following lines. It may be wrapped to any number of columns (no not wrapped at all). Gaps in the alignment are represented with the "-" character.

At this time (version 1.1), SPIMAP can only use DNA sequences. The sequence can be in both upper case and lower case (SPIMAP ignores case) and degeneracy codes can be used ("NnRrYyWwSsKkMmBbD-dHhVv"), however at this time SPIMAP treats all degeneracies as completely missing data ("N").

1.2 Species tree file format (*.stree)

Species trees should be specified using the Newick file format. See http://en.wikipedia.org/wiki/Newick_format for details. Beyond the newick format, SPIMAP has only a few additional requirements. First, the species names given in the species tree should match those given in the SMAP file (Section 1.3). Second, the branch lengths of the species tree should be expressed in units of time (Figure 2). Any unit of time can be used (e.g. millions of years, generations, relative units, etc). The only requirement is that the duplication and loss rates are also expressed in compatible units. Therefore, if branch lengths are in millions of years, the duplication rate (specified by spimap's "-D" option) should be in units of duplications/gene/million years.

Naming ancestral nodes. SPIMAP also supports naming ancestral nodes in the species tree using the newick format. For example, the parental node of human and chimp can be named primate using the following syntax:

```
(((((((scer:7.061760
       spar:7.061760
      ) n7:4.999680,
      smik:12.061440
    )n6:5.970600.
    sbay:18.032040
    ) n5:52.682400,
   cgla:70.714260
  )n4:7.220700,
  scas:77.934960
 )n3:23.181480,
   agos:78.553260
    klac:78.553260
  )n9:10 434960
  kwal:88.988220
 )n8:12.128400
)n2:78.883560.
   calb:41.275620
  ctro:41.275980
)n12:29.632860,
    cpar:52.323120
   lelo:52.323120
  )n13:18.585720
  )n11:31.149540.
   cgui:75.615840
   dhan:75.615840
  )n15:14.006880,
  clus:89.622720
 )n14:12.435660
)n10:77.941620
```

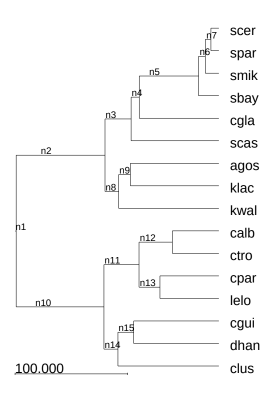


Figure 2: **Example *.stree file and corresponding tree.** This file (left) specifies the species tree (right) using the newick file format. Branch lengths should be expressed in units of time (e.g. millions of years). Ancestral nodes can also optionally be named (the names "n1", "n2", etc are used in this example).

```
((human:5,chimp:5) primate:70,mouse:75) mammal;
```

If ancestral nodes are named, they will be used in the output of the reconciliation mapping (Section 1.4).

1.3 Gene to species name mapping file format (*.smap)

SPIMAP uses a special file format (*.smap) to specify which genes belong to which species. Each line contains two tab-delimited fields:

- 1. pattern matching a gene ID
- 2. species ID

Only 3 types of gene ID patterns are supported. The pattern can either be an exact matching string, a prefix (denoted "text*"), or a suffix (denoted "*text"). The "*" is the only special wildcard character.

The species ID should be the same as the ones used in the species tree files. All patterns and IDs are case-sensitive.

When mapping a gene name to a species name all exact matches are processed first. If no exact match is found, the patterns are then processed in the same order as they appear in the file until a match is found.

```
A*
        agos
orf19*
        calb
CDUG *
        cdub
CAGL*
        cgla
IPF *
        cgla
CGUG *
        cgui
sbay_*
        sbay
scas *
        scas
smik *
        smik
spar_*
        spar
SCP*
        spom
YALI*
        ylip
Y*
        scer
Q*
        scer
```

Figure 3: **Example *.smap file.** This file specifies how to map gene names to their corresponding species. The first column indicates a gene name pattern (in this case a prefix) and the second column specifies a species name. Note: this example only gives a partial list of the species in Figure 2.

For example in the SMAP file given in Figure 4, the gene ID "YALI123" should match the species "ylip", instead of "yeast", because the pattern "YALI*" occurs before "Y*".

1.4 Reconciliation file format (*.recon)

When SPIMAP's "-r" option is used, the reconciliation found for the gene tree and species is saved to a file "OUTPUT_PREFIX.recon". The reconciliation file format is tab-delimited, where each line has three fields:

- 1. gene node ID.
- 2. species node ID.
- 3. event (one of the following: "gene", "spec", "dup")

Each line specifies the mapping of one node in the gene tree (field 1) to one node or branch in the species tree (field 2). Branches are indicate using the node ID directly below it (i.e. the younger of the two incident nodes). The lines can be given in any order.

If the gene node is a leaf, it will map to a leaf in the species tree and the event field will contain the event "gene". All internal nodes of the gene tree are marked either as speciations (event "spec") or duplications (event "dup"). Specaition nodes map directly to the indicated species node, and duplication nodes map to the indication species branch. The time of the duplication along the species branch is not indicated in this file format nor is it inferred by SPIMAP.

If gene IDs are not given to the ancestral nodes of a gene tree or species tree, SPIMAP will by default name them with "nXXX" where XXX is the preorder traversal of the internal nodes.

KLLA0C08239g	klac	gene
ADL072C	agos	gene
kwal_5828	kwal	gene
CAGL0J02970g	cgla	gene
scas_g715.48	scas	gene
smik_6662	smik	gene
sbay_7039	sbay	gene
smik_6659	smik	gene
sbay_7037	sbay	gene
YER061C	scer	gene
spar_6281	spar	gene
n10	n5	spec
n9	n7	spec
n8	n6	spec
n7	n5	spec
n6	n5	dup
n5	n3	spec
n4	n3	spec
n3	n9	spec
n2	n8	spec
n1	n2	spec

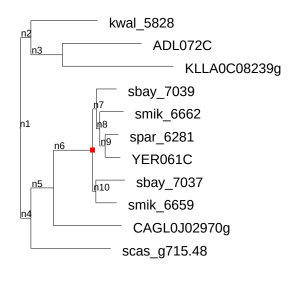


Figure 4: **Example *.recon file.** The reconciliation file format (left) specifies how all the nodes in a gene tree (right) map to the nodes and branches in the species tree (see Figure 2). Notice that gene node "n6" (red dot) represents a duplication event along species branch "n5" (shown Figure 2). The gene tree and species tree have their own name space ("n5" in the gene tree is not the same as "n5" in the species tree).

1.5 SPIMAP model parameters file format (*.params)

SPIMAP has several parameters for its substitution rates model. These parameters are learned by the spimap-train-rates program, which saves the parameters in a custom *.params file format (Figure 5). The spimap program reads these parameters using the "-p" option. Most uses of SPIMAP do not require understanding the contents of a *.params file.

The *.params file format is tab-delimited and each line is processed one at a time.

If the first field of a line is the word "baserate", then the remaining two fields are interpreted as floating point values α_G and β_G , which are the two parameters, shape and scale, of the inverse-gamma distributed gene-specific rate.

If the first field of the line does not match "baserate", then the first field indicates a species tree branch and the remaining two fields are interpreted as floating point values α_i and β_i , which are the two parameters, shape and scale, of the gamma distributed species-specific rate. Each branch is indicated by its more recent node. Ancestral nodes are indicated by an integer, where are assigned in pretraversal order.

<u>?</u> }	6.98457288742 3.28887700831 4.64684152603 1.13027572632	5.98457288742 394.209221588 551.109741211
1 2 3	4.64684152603	
3		551.109741211
	1 13027572632	
	1.10021312002	164.191940308
4	0.610769152641	75.0393371582
5	7.14405012131	927.631103516
6	2.96983885765	238.195861816
7	5.63683271408	632.264831543
8	0.974860072136	94.9837493896
9	0.856632292271	78.6899032593
10	4.64683914185	544.528686523
11	1.92581880093	271.891052246
12	3.84569692612	624.703308105
13	3.14617466927	335.446655273
14	0.699178874493	84.1814575195
15	0.746283352375	137.345901489
scer	8.42576217651	763.305847168
ctro	6.70220327377	999.845153809
scas	9.14448356628	1253.45031738
agos	8.84074497223	801.648925781
sbay	6.95680332184	1048.7590332
kwal	14.3321857452	1962.9083252
dhan	15.7483224869	2699.00878906
smik	10.2562847137	1143.78076172
cgla	9.81903266907	1015.43951416
spar	5.80616807938	799.18963623
calb	8.38038921356	1233.68322754
lelo	9.40990924835	973.772583008
cpar	9.43262672424	1184.28100586
klac	6.6709280014	767.418823242
clus	8.37989234924	881.762878418
cgui	11.9692239761	1187.47314453

Figure 5: **Example *.params file.** The *.params file contains the parameters for SPIMAP's substitution rate model.