

Software for isometric gene tree reconciliation

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Progress

First chapter

- Introduction to the topic
- Explanation of basic terms, essential terminology from bioinformatics; gene tree, species tree as trees from graph theory
- Definition of gene tree reconciliation – mapping gene tree to species tree
- Introduction of existing software with different gene tree reconciliation algorithms; three types:
 - Scoring gene tree reconciliation
 - Probabilistic gene tree reconciliation
 - Isometric gene tree reconciliation
- GitHub of project: https://github.com/mihalova/diploma_thesis

Software

- Goal: software for isometric reconciliation of unrooted gene tree and rooted species tree
- Source code based on master's thesis of Radoslav Chládek (Algorithms for isometric gene tree reconciliation. Univerzita Komenského v Bratislave, 2019)
- Programming language: Java (SDK 11)
- Libraries: Ipsolve (linear programming solver – used in reconciliation algorithm), junit-jupiter-api (for running unit tests)

My work

- Changing parser to parse unrooted gene tree with intervals or add intervals to gene tree
- Parse mapping of leaves from gene tree to species tree
- Separating one Reconciliator class to three (Main, Printer and Reconciliator)

```
if (tolerance != null && !time.contains("-")){
    //nastavenie tolerance pre hrany
    double t = Double.parseDouble(time);
    double[] interval = makeInterval(t, tolerance);
    min_time = interval[0];
    max_time = interval[1];
} else if (time.contains("-")) {
    //parovanie intervalov 2.02-3.01 = double "-" double
    String[] interval = time.split( regex: "-");
    min_time = Double.parseDouble(interval[0]);
    max_time = Double.parseDouble(interval[1]);
} else
    throw new IllegalArgumentException("Insert tree with intervals or set tolerance.");

str = str.substring(0, colonIndex);
if(!str.startsWith("(")){
    //tu bude parovanie leafmap G0_S1 = nodeG string "_" nodeS string
    String[] mappingG = str.split( regex: "_");
    G.addMapping(str, mappingG[1]);
}
```

Input and output

Input: (path to directory with trees and tolerance - optional)

1. Path to directory with rooted species tree and unrooted gene tree with intervals
2. Path to directory with rooted species tree, unrooted gene tree with intervals on some edges and tolerance
3. Path to directory with rooted species tree, unrooted gene tree without intervals and tolerance

Output: most parsimonious reconciliation of gene tree to species tree (Radoslav Chládek's algorithm)

Example of reconciliation

```
Total DL of solution: (3,1)
Number of the most parsimonious solutions: 1
Minimal DL: (3,1)

#[min depth] [max depth]
root 0.0162725 0.07937
G11_S20G12_S21 0.13371 0.17307
G11_S20 0.2301 0.2301
G12_S21 0.23484 0.23484
G0_S15G1_S16G3_S18G4_S18G5_S18G7_S18 0.110875 0.13432
G0_S15G1_S16 0.19174 0.19917
G0_S15 0.22723 0.22723
G1_S16 0.22835 0.22835
G3_S18G4_S18G5_S18G7_S18 0.1125 0.139195
G3_S18 0.23102 0.23102
G4_S18G5_S18G7_S18 0.13721 0.16969
G4_S18G5_S18 0.22064 0.224865
G4_S18 0.23102 0.23102
G5_S18 0.23102 0.23102
G7_S18 0.23102 0.23102
```

Next work

Software finds most parsimonious rooting of unrooted gene tree considering species tree from different isometric reconciliations (different algorithms)

Validation:

- Comparison of results from software with considered branch lengths/without considered branch lengths
- How much good/bad was the result?
- Dependency change of parameters (length of branch intervals, tolerance of edge) to the result

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