Guide to Turning an RDP File into a Data Set

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Below we will define the basic steps required to generate a data set from an RDP file

1. Take an RDP file such as this example:

; Root: 1.0; Bacteria: 1.0; Firmicutes: 1.0; Bacilli: 1.0; Bacillales: 1.0; Staphylococcaceae: 1.0; ; Root: 1.0; Bacteria: 1.0; Firmicutes: 1.0; Bacilli: 1.0; Lactobacillales: 1.0; Carnobacteriaceae: 0.8; ; Root: 1.0; Bacteria: 1.0; Bacteroidetes: 1.0; Bacteroidia: 1.0; Bacteroidales: 1.0; Prevotellaceae: 1.0; ; Root: 1.0; Bacteria: 1.0; Bacteroidetes: 1.0; Bacteroidia: 1.0; Bacteroidales: 1.0; Prevotellaceae: 1.0; Bacteroidetes: 1.0; Bacteroidia: 1.0; Bacteroidales: 1.0; Prevotellaceae: 1.0; Bacteroidia: 1.0

;Root:1.0;Bacteria:1.0;Bacteroidetes:1.0;Bacteroidia:1.0;Unclassified:0.5;Unclassified:0.5;

2. Take the first entry and seperate each level into its own row, while seper-

ating levels by a period:

Root, 1

Root.Bacteria, 1

Root.Bacteria.Firmicutes, 1

Root.Bacteria.Firmicutes.Bacilli, 1

Root.Bacteria.Firmicutes.Bacilli.Bacillales, 1

Root.Bacteria.Firmicutes.Bacilli.Bacillales.Staphylococcaceae, 1

3. Do the same with each following row, adding to the number at the end if it is the same:

Root, 5

Root.Bacteria, 5

Root.Bacteria.Firmicutes, 2

Root.Bacteria.Firmicutes.Bacilli, 2

Root.Bacteria.Firmicutes.Bacilli.Bacillales, 1

Root.Bacteria.Firmicutes.Bacilli.Bacillales.Staphylococcaceae, 1

Root.Bacteria.Firmicutes.Bacilli.Lactobacillales, 1

Root.Bacteria.Firmicutes.Bacilli.Lactobacillales.Carnobacteriaceae, .8

Root.Bacteria.Bacteroidetes, 3

Root.Bacteria.Bacteroidetes.Bacteroidia, 3

Root.Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales, 2

Root.Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Prevotellaceae, 2

Root.Bacteria.Bacteroidetes.Bacteroidia.Unclassified, .5

Root.Bacteria.Bacteroidetes.Bacteroidia.Unclassified.Unclassified, .5

4. Change any unclassifieds or anything else that could appear under multi-

ple different parents (we prefer adding a 'U' to the parents name):

Root.Bacteria.Bacteroidetes.Bacteroidia.BacteroidiaU, .5

Root.Bacteria.Bacteroidetes.Bacteroidia.BacteroidiaU.BacteroidiaUU, .5

5. Remove the 'Root' level since we are only looking at Bacteria in this example:

Bacteria, 5

Bacteria. Firmicutes, 2

Bacteria. Firmicutes. Bacilli, 2

Bacteria. Firmicutes. Bacilli. Bacillales, 1

Bacteria. Firmicutes. Bacilli. Bacillales. Staphylococcaceae, 1

Bacteria. Firmicutes. Bacilli. Lactobacillales, 1

Bacteria. Firmicutes. Bacilli. Lactobacillales. Carnobacteriaceae, 1

Bacteria.Bacteroidetes, 3

Bacteria.Bacteroidetes.Bacteroidia, 3

Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales, 2

 ${\bf Bacteroide tes. Bacteroidia. Bacteroidales. Prevotellaceae,\ 2}$

Bacteria.Bacteroidetes.Bacteroidia.BacteroidiaU, .5

Bacteria.Bacteroidetes.Bacteroidia.BacteroidiaU.BacteroidiaUU, .5

6. Repeat for any additional RDP files and add them as new columns (and rows as needed) in our data set:

Taxa, Sample 1, Sample 2

Bacteria, 5, 5

Bacteria. Firmicutes, 2, 5

Bacteria. Firmicutes. Bacilli, 2, 5

Bacteria. Firmicutes. Bacilli. Bacillales, 1, 5

Bacteria. Firmicutes. Bacilli. Bacillales. Staphylococcaceae, 1, 5

Bacteria. Firmicutes. Bacilli. Lactobacillales, 1, 0

Bacteria. Firmicutes. Bacilli. Lactobacillales. Carnobacteriaceae, 1, 0

Bacteria.Bacteroidetes, 3, 0

Bacteria.Bacteroidetes.Bacteroidia, 3, 0

Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales, 2, 0

Bacteria.Bacteroidetes.Bacteroidia.Bacteroidales.Prevotellaceae, 2, 0

Bacteria.Bacteroidetes.Bacteroidia.BacteroidiaU, .5, 0

Bacteria.Bacteroidetes.Bacteroidia.BacteroidiaU.BacteroidiaUU, .5, 0

Bacteria. Firmicutes. Bacilli. Lactobacillales. Streptococcaceae, 0, 5

7. Save the file as a .csv (or anything other file readable by R) and load it in R:

Notes:

- Any symbol can be used as a taxa level seperator except colons.
- There can only be one top level node, if we were looking at Archaea too we would need to keep the Root level in.