

Bray-Curtis similarity

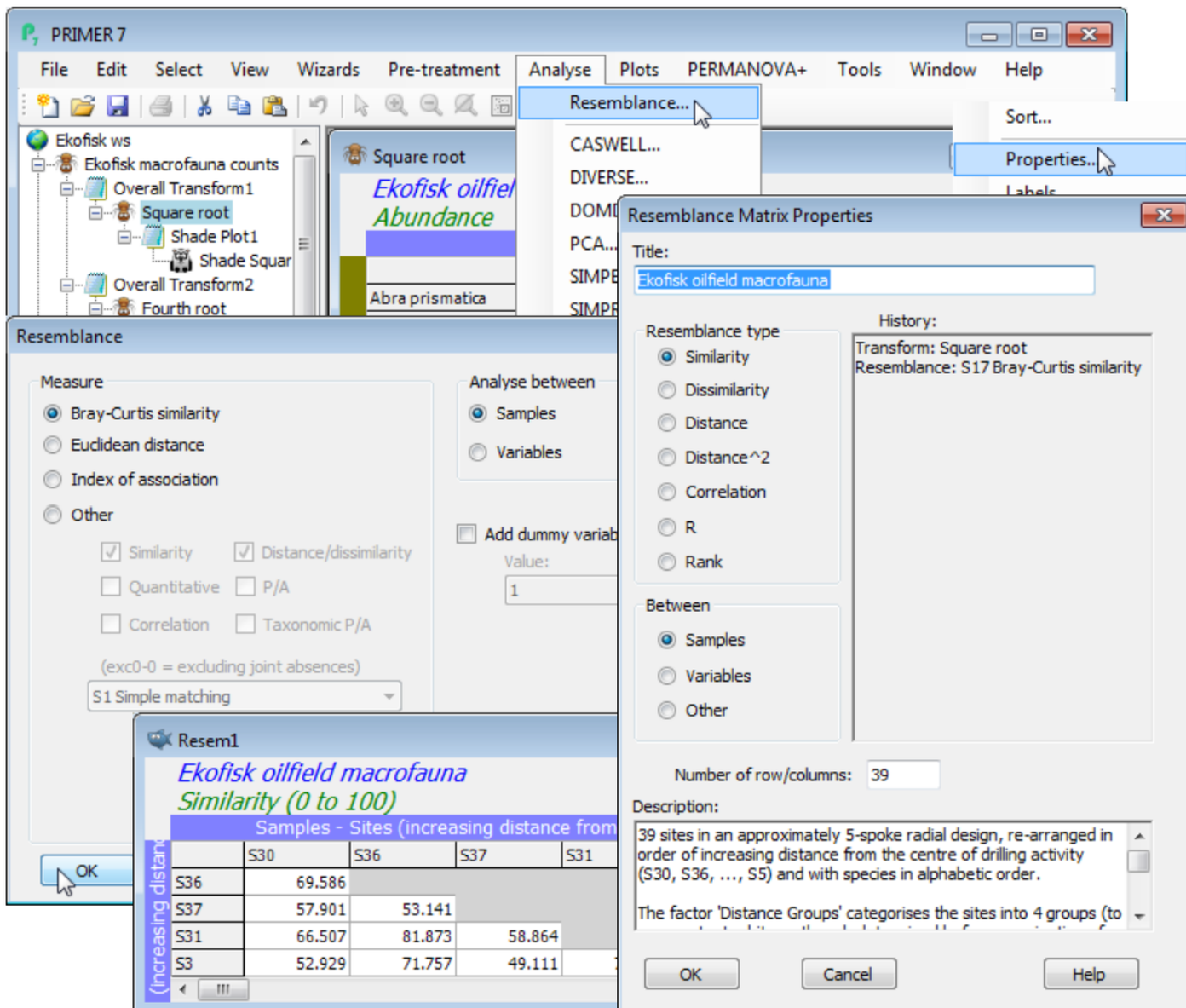
The most commonly-used similarity coefficient for biological community analysis, because it obeys many of the 'natural' biological guidelines in a way that most other coefficients do not (see CiMC), is the Bray-Curtis similarity, defined between samples 1 and 2 as:

$$S_{17} = 100 \left[1 - \frac{\sum_i |y_{i1} - y_{i2}|}{\sum_i y_{i1} + \sum_i y_{i2}} \right] \equiv 100 \frac{\sum_i \min(y_{i1}, y_{i2})}{\left(\sum_i y_{i1} + \sum_i y_{i2} \right) / 2} .$$

The two forms may not look identical but they are! Here y_{i1} is the count (or biomass, % cover, ...) for the i th (of p) species from sample 1, and $\sum_i (...)$ denotes summation over those species. Original references to coefficient definitions are not given here (nomenclature is always a source of debate!) - see [L&L](#), whose numbering scheme is followed where possible, hence S_{17} for Bray-Curtis.

Open the workspace C:\Examples v7\Ekofisk macrofauna\Ekofisk ws from earlier, and click on the Square root counts sheet (obtained earlier with **Pre-treatment>Transform(overall)>Square root**). Take **Analyse>Resemblance>(Measure•Bray-Curtis similarity) & (Analyse between•Samples)**, which are the defaults for this data type. A lower triangular matrix is produced, Resem1, which you should rename **B-C on sq rt**. **Edit>Properties** (or right-clicking over the matrix to get **Properties**) shows it is of Resemblance type•Similarity from 39 samples. The History box carries through the knowledge of how it was created to a subsequent Cluster or MDS ordination plot. This box is not user-editable, though the Title and Description boxes can be altered; changes to the Title are carried forward to a subsequent plot but not backward to the data sheet **Square root**.

Now repeat **Resemblance** directly on the original **Ekofisk macrofauna counts**, without the **Pre-Treatment** transformation. PRIMER tries to help - a warning message appears that no transform has been applied; community matrices usually require some transformation before calculating Bray-Curtis (though you can happily ignore this warning if you are interested in the pattern of the few most dominant species only). **Cancel** the calculation and resave the **Ekofisk ws** workspace.



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