

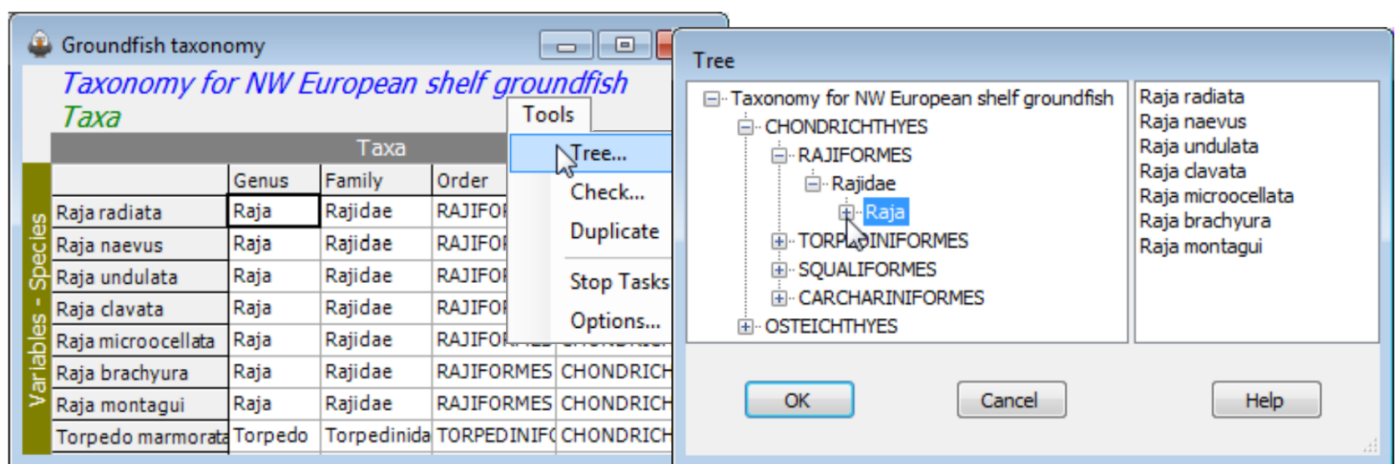
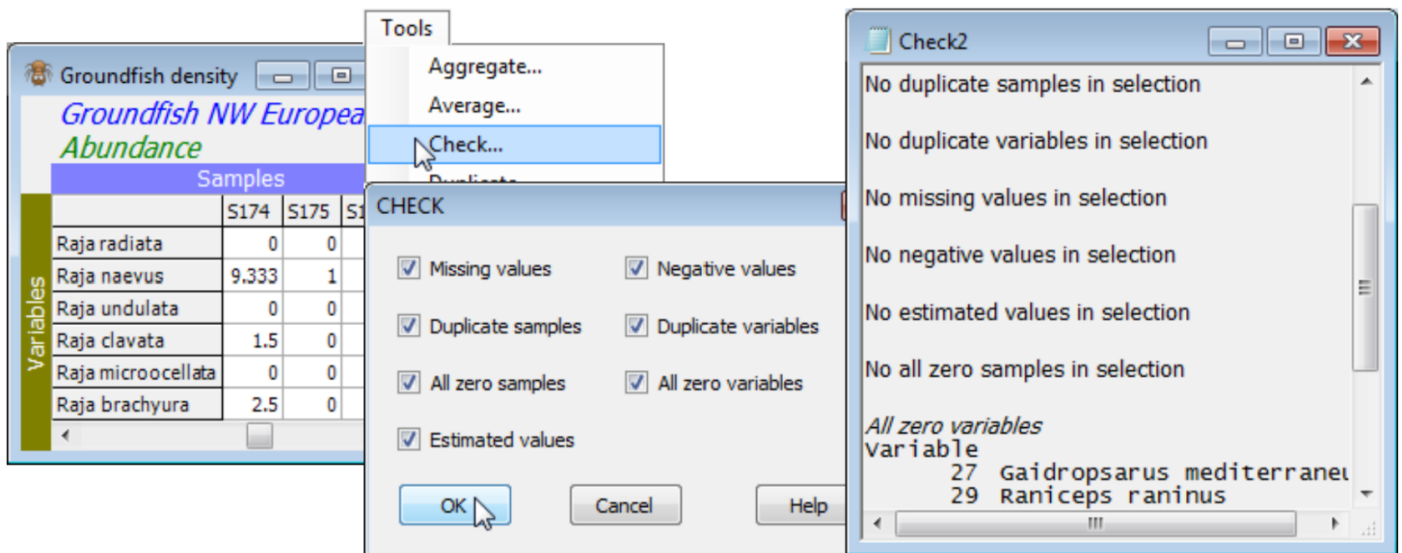


Tree menu; Check on datasheets & resemblances; Undefined resemblances

The other **Tools** menu item for aggregation sheets is distinctive to this case, namely **Tools>Tree**; it simply displays the hierarchical structure of an aggregation file in the same way as the Explorer tree, in a left-hand panel. Successive clicking on the  icons unroll the taxonomic structure, and it can be rolled back with . No operations can be performed on the display in this state.



When the active window is a datasheet, **Tools>Check** can check for the following: a) ✓ Missing values, identified in the sheet by 'Missing!', and which might have been read in as blank cells in an Excel worksheet for example; b) ✓ Negative values, which are not appropriate for abundance-type data analysed by Bray-Curtis, though common for environmental variables (especially normalised) input to Euclidean distance; c) ✓ Duplicate sample (and/or) variable labels, which are tolerated for some analyses (warnings are usually given) but are best avoided wherever possible; d) ✓ All zero samples (and/or) variables; and e) ✓ Estimated values, displayed in red type in the matrix. The latter come from applying **Tools>Missing** (seen shortly) to environmental variables – or to other normally distributed data – containing **Missing!** cells, which otherwise might not be tolerated by some analysis routines requiring complete data. All or any of the 7 boxes can be ticked. Whether it is important to check for a particular attribute depends on the analysis. For example, species which are zero over all samples will be ignored when Bray-Curtis similarity is computed among samples, and can safely be left in the matrix, but all-zero samples are potentially more of a problem since Bray-Curtis similarity between two blank samples is set to 'Undefined!'. Dependent on the context, these samples might best be omitted, or a different similarity used (e.g. zero-adjusted Bray-Curtis, Section 5), or the entry left as 'Undefined!', i.e. treated as unknown.



When the active window is a resemblance sheet, **Tools>Check** looks for only three data attributes: a) ✓ Undefined values, arising as suggested above; b) ✓ Out of bounds values, for distance coefficients (or transformations) that return very large or small values (NaN); and c) ✓ Duplicate labels, as above. Blanking a cell in a resemblance matrix sets it to **Undefined!** status, and several of the core routines using resemblances (e.g. MDS, Cluster, ANOSIM) are carefully written in PRIMER to tolerate a few **Undefined!** entries, treating them as unknown. (You can appreciate that knowing the similarities SS_{12} , SS_{13} , SS_{14} , SS_{23} , SS_{24} might enable you to place four samples in relation to each other without knowing similarity SS_{34}). Blanking out NaN (Not a Number) entries, to **Undefined!**, is one possibility therefore, but others may be equally good or better (replacing by a large, but finite value, modifying the coefficient or transformation which generated them etc.).

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