



BIONUMERICS Tutorial:

Creating summary spectra in a database with levels

1 Introduction

This tutorial describes how to combine several spectra to create summary spectra, according to the levels and dependencies present in the BIONUMERICS database.

2 Preparing the database

1. Create a new database and import the example raw spectra files as described in the tutorial: "Importing spectrum data in a database with levels". Make sure the *Database design* panel is docked above the *Database entries* panel.

Entries belonging to a selected level in the *Database design* panel are displayed in the *Database entries* panel.

2. Click on the "Isolate" level in the *Database design* panel (see Figure 1).

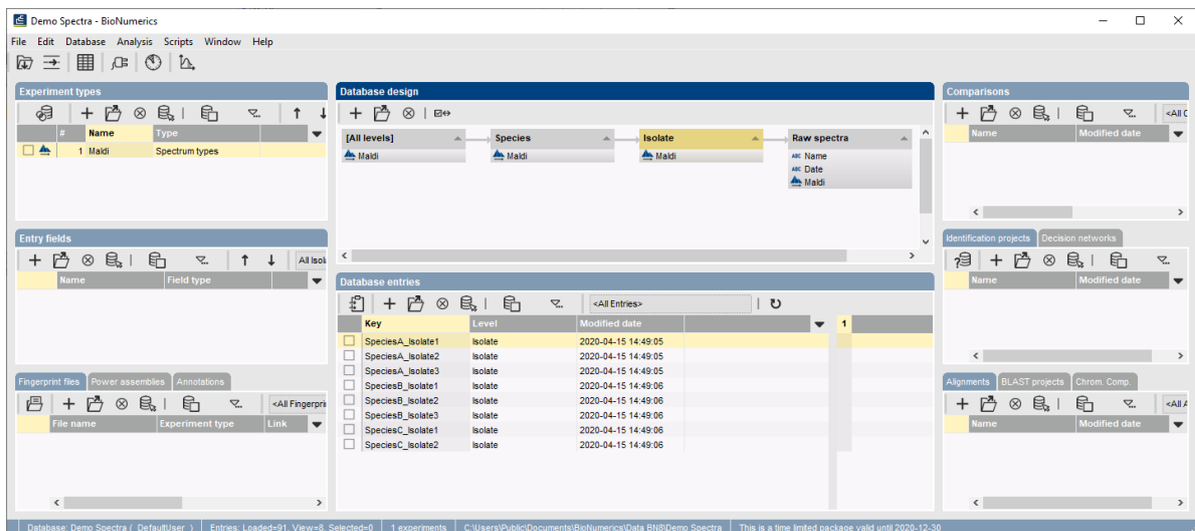


Figure 1: *Database design* panel shown above the *Database entries* panel: the entries belonging to the level "Isolate" are shown in the *Database entries* panel.

3. Verify the creation of the correct dependencies in each level, by double-clicking on an entry to open the *Entry* window. Click the tab of the *Dependencies* panel to display the dependencies (see Figure 2 for an example).

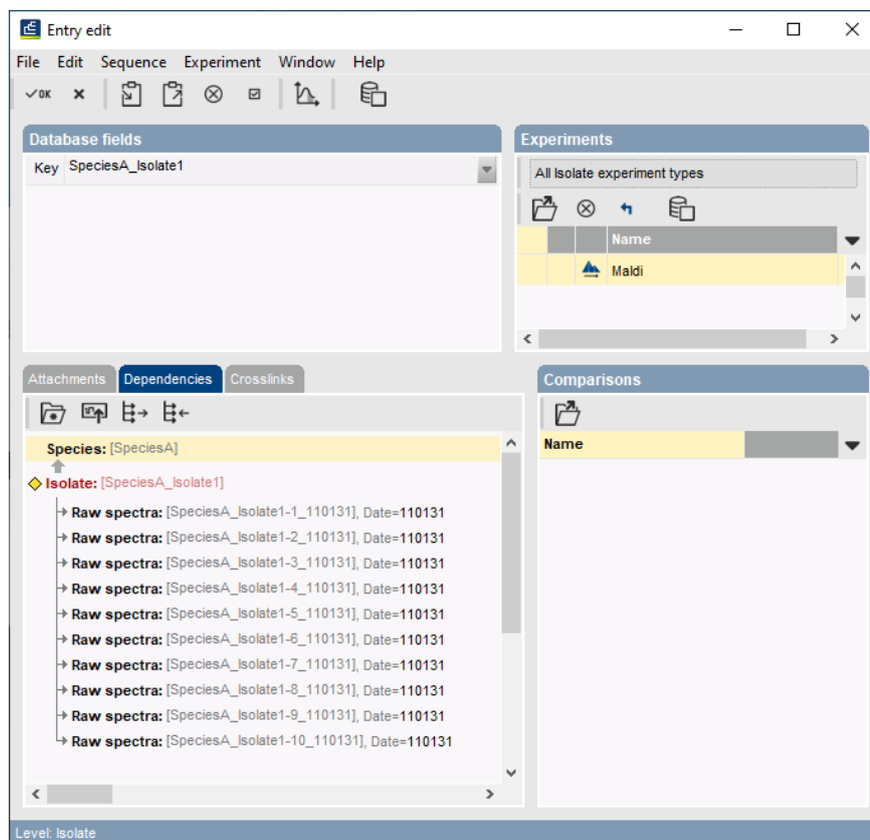


Figure 2: Example of dependencies at a database level.

3 Creating summary spectra from raw spectra

1. At the lowest level, "Raw spectra", select all entries (**Edit** > **Select all (Ctrl+A)**) and select **Analysis** > **Spectrum types** > **Summarize...** to open the *Create summary spectra* wizard.
2. In the first page of the *Create summary spectra* wizard, select the option *Higher level* and press <Next>.
3. In the second page, select the level "Isolate" as target level and press <Next> twice.

In the fourth step of the wizard (see Figure 3), the user can select an appropriate workflow for the summary of the spectra.

Three predefined summary templates are present: one for technical replicates (**Replicates (technical)**), one for biological replicates (**Replicates (biological)**) and one general template for processing for instance several strains of the same subgroup or the same species (**Related spectra**). The settings of each template can be changed in the *Summary Spectrum* window and saved to the database (see further).

4. Select the predefined summary workflow **Replicates (technical)** and press <Next> and <Finish>.

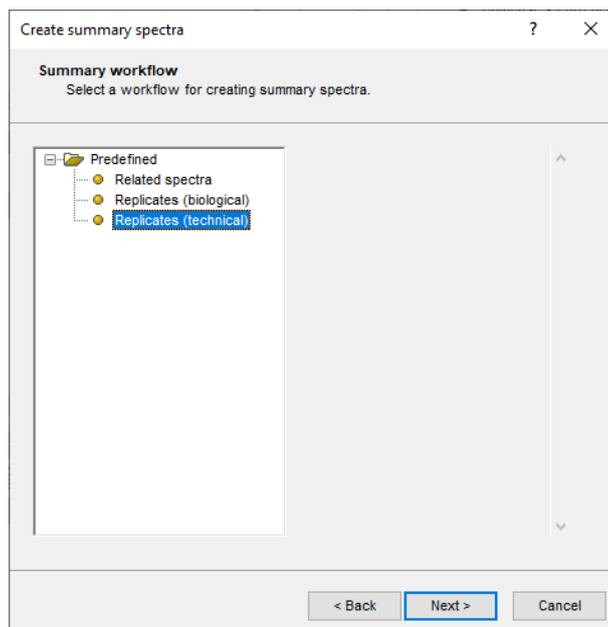


Figure 3: Summary templates.

The *Summary Spectrum* window will now open with the raw spectra organized into 8 summary spectra (see Figure 5).

5. Select a summary spectrum from the list in the *Summary Spectra* panel.

The member spectra are updated in the *Members* panel and in the *Peak List* panel. The average similarity of all member spectra is calculated and the similarity of each individual spectrum with this average similarity is indicated in the **Simil** column. Member spectra that are grayed out have been inactivated because their similarity (**Simil**) is below the similarity threshold that is specified. The similarity threshold depends on the selected summary template:

- **Replicates (technical)**: similarity threshold placed at 95%
- **Replicates (biological)**: similarity threshold placed at 80%
- **Related spectra**: placed at 40%

6. To check (and optionally update) the similarity threshold, double-click on the **Import & Similarity filter** operator in the *Workflow* panel to call the *Workflow* window. Select the **Similarity filter** operator from the flow chart, and choose **File > Operator settings...** (⚙️). The similarity threshold can be updated by the user and the updated template can be saved to the database with **File > Save workflow as template**. Close the *Workflow* window.

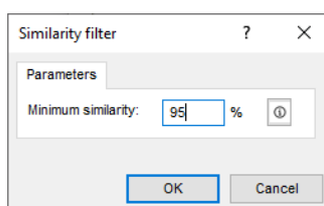


Figure 4: The similarity threshold for the technical replicates template.

7. Press **<Peak Matching>** to calculate the summary spectra and to identify summary peaks.

After execution of the workflow, the *Summary Spectrum* window will look like Figure 6. The summary spectra will appear as a thick red line. Each peak from the summary spectrum appears as thin vertical line, adjacent peaks are marked with a different color.

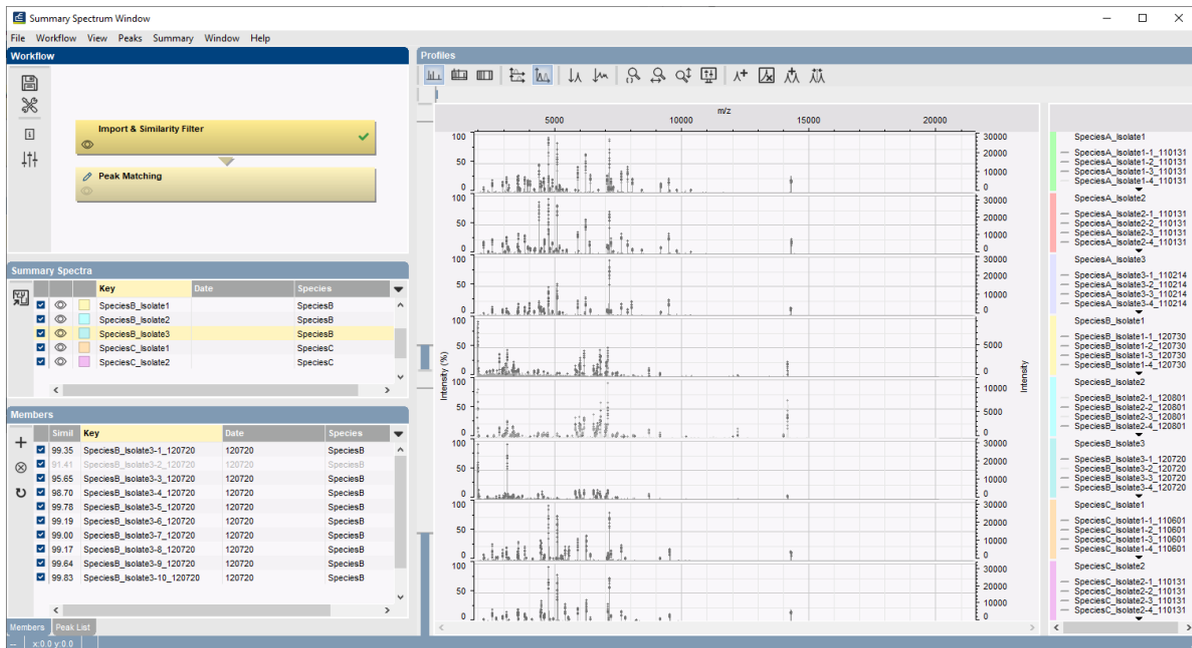


Figure 5: *Summary Spectrum* window before running the workflow.

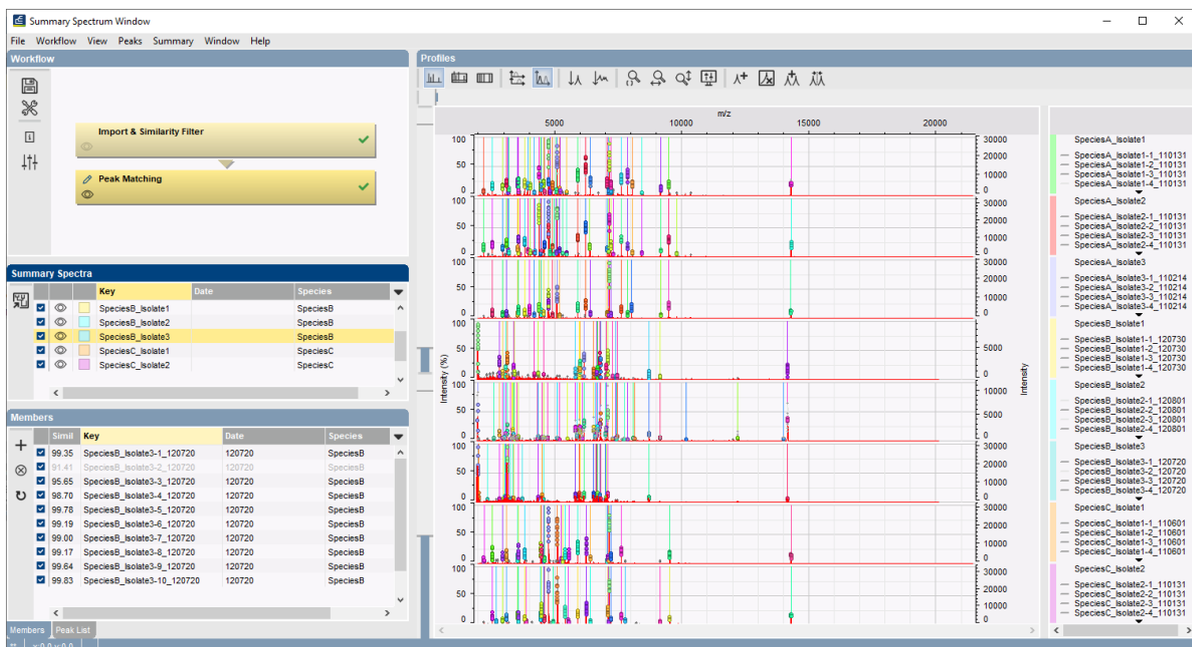


Figure 6: *Summary Spectrum* window after running the workflow.

8. Save the summary spectra to the database with **File > Save spectrum data** ().

Each entry at the level **Isolate** will now have an experiment present for the spectrum type **Maldi**.

4 Comparison of member spectra

To review the quality of the raw spectra and the subsequent summary, it is possible to create a comparison from the *Summary Spectrum* window with all the member spectra:

1. In the *Summary Spectrum* window, select **Summary** > **Create comparisons** (🔍) and select the option to create a comparison from the member spectra and press <OK>.

This action will open a comparison window including all member spectra grouped according to their summary spectra.

2. Select **Clustering** > **Calculate** > **Cluster analysis (similarity matrix)...**, choose the curve-based **Pearson correlation** coefficient, click <Next>, make sure **UPGMA** is selected and press <Finish>.

In the resulting dendrogram the member spectra are clustered together with the other members. If the member spectra of several summary spectra are clustered together, either the preprocessing or summary workflow need to be reviewed. In case of closely related isolates, it might also be that they cannot be distinguished from one another based on their spectra. In this case, the two isolates from Species C are very similar and it is likely that the distinction between them will be less reliable than the distinction between the isolates from Species A (see Figure 7).

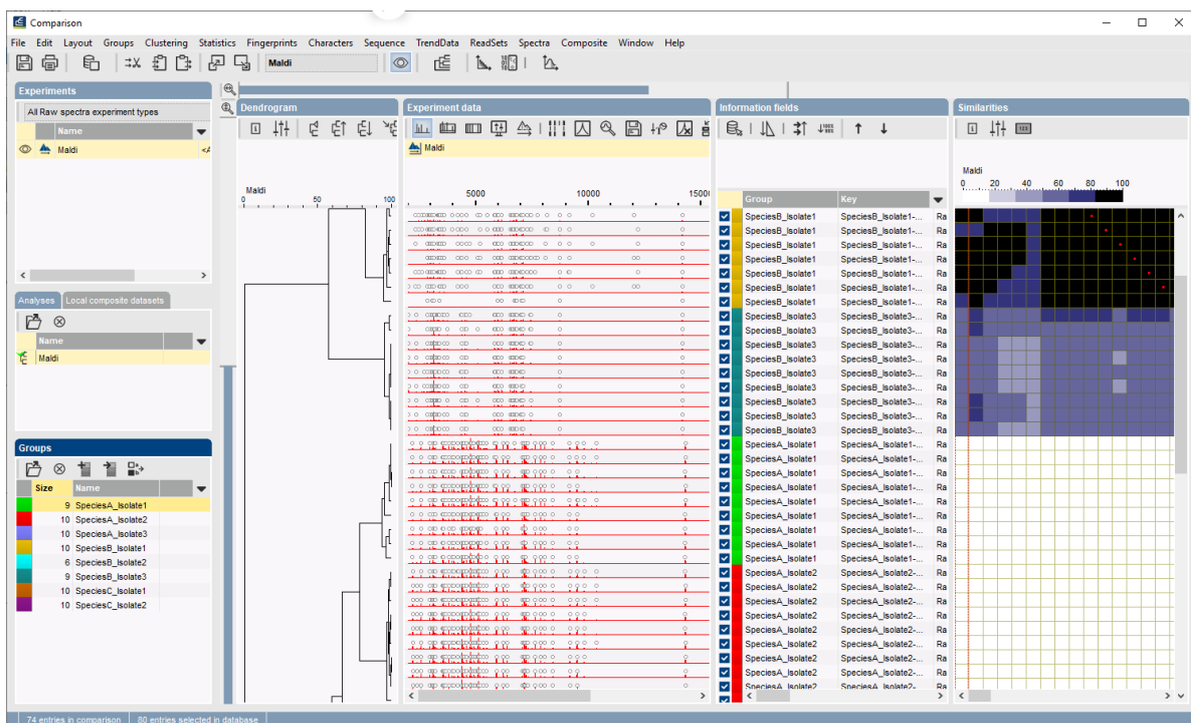


Figure 7: Comparison window of member spectra generated from the *Summary Spectrum* window.

5 Creating summary spectra from summary spectra

1. Close the *Comparison* window (save if desired) and the *Summary Spectrum* window.
2. Unselect all entries in the *Main* window by pressing the **F4** key.

The summary spectra of the isolates can themselves be summarized per species:

3. Select the level "Isolate" in the *Database design* panel to make it the active level.
4. Click somewhere in the *Database entries* panel to make it the active panel and select **Edit** > **Select all (Ctrl+A)** to select all 8 isolate entries in the "Isolate" level.
5. Select **Analysis** > **Spectrum types** > **Summarize...**, select the option *Higher level* and press <Next>. Choose "Species" as target level, press <Next> twice and choose the workflow "Related spectra". Press <Next> and <Finish>.
6. Press <**Peak Matching**> in the *Workflow* panel to execute the summary workflow.

For these spectra, the tolerance settings are a bit too strict, this is seen best in the summary spectrum of Species A (see Figure 8).

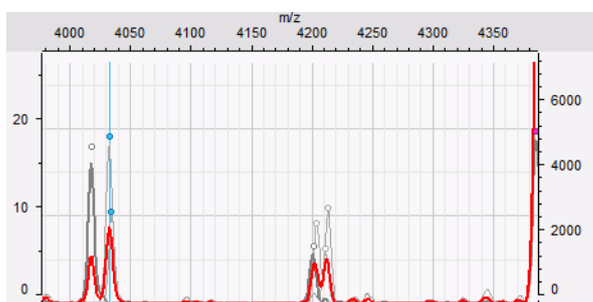


Figure 8: Summary spectra of Species A at default tolerance settings.

7. To change the tolerance settings, double-click on <**Peak Matching**> to open the *Workflow* window. Highlight peak matching and select **File** > **Operator settings...** (⚙️). This will open the dialog box with the parameters of the Peak matching operator. Increase the constant tolerance from 1 to 2 and increase the linear tolerance from 300 ppm to 400. Press <OK>.
8. Close the *Workflow* window, rerun the workflow by pressing <**Peak Matching**> (see Figure 9 for the updated peak matching for Species A).

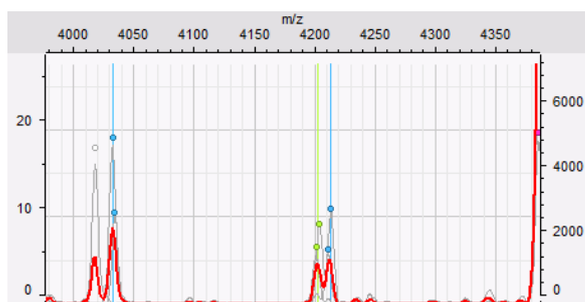


Figure 9: Summary spectra of Species A with less strict tolerance settings.

9. Save the results with **File** > **Save spectrum data** (💾).

The entries at the level **Species** will now also have an experiment present for **Maldi**. All summary spectra can be used for follow-up analysis such as peak matching, determining common or discriminative peaks, identifying unknown spectra against a database, etc..