# EEGpal: **Peark or Trace export module**

Version 2.0, 31.08.2025

The **Peak or Trace export** module has three possible functions. Firstly, the module enables the identification of the maximum (max) and minimum (min) positive and negative peaks within the specified time interval. The module generates an Excel or text file containing the amplitude and position.

The second functionality is to export the amplitude values for each time point, known as the timecourse. This enables statistics to be performed on the electrodes and times.

The third functionality of this module is to calculate the average of the signal in the specified time interval. This is useful for extracting values for plotting.

The module saves output files that can easily be loaded into your favourite statistical software, such as Excel, R or SPSS. By default, it saves .xlsx files, but you can also save .txt files with a tab as the delimiter.

This module is designed to work with epoched files (i.e. files processed by the**Epoching** module). Imported files must have an identical number of TFs, electrodes and sampling rate in the main EEGpal window.

### Find peaks

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1. Sampling rate of the data. This should normally be transferred automatically from the main window. You can adjust this if necessary.
2. Choose the action *‘Peaks detection’.*
3. Specify the time interval. Warning: the value 0 marks the start of the file. This module does not take into account a possible .mrk file that could define a different origin.  
   EoF=End of File.
4. Specify the electrode to be studied and the indices of the electrodes (1, 2, 64), rather than the names specified by the coordinate file (A1, A2, B32). A ‘\*’ means that all the electrodes will be study.
5. Indicate whether you want to detect positive peaks (i.e. looking for maxima) or negative peaks (i.e. looking for minima)
6. In addition, you can detect the peaks on the average of specified electrodes in **4**, on the global field power (GFP) or on the global map dissimilarity (GMD).   
   The result for GFP and GMD is independent of the parameter **5**. Indeed, these values are always positive so the module will always return the maxima.
7. Select if the output file format: .xlsx or .txt (with tab as delimiter).
8. Indicate whether the peak position should be specified in milliseconds (ms) or time frames (tf) in the output file.
9. Select the output folder where the output will be recorded.
10. You can specify manually a suffix in the output file name. For example: *PeakDetection\_Positive\_98-122ms\_P1.xlsx*
11. Record an additional check file. Read the section **How can I use the check file?** in the FAQ bellow**.**
12. There are three validation buttons:
    1. The **Run** button will carry out the processing parameterized in the module.
    2. The **Save in memory** button will store all the parameters in memory and close the module without performing the processing.
    3. The button **Cancel** closes the module without processing and without keeping the entered parameters in memory. The same effect will be achieved by closing the module window.

FAQ

**What the output looks like?**

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The first columns show the amplitude and the last columns show the position in either milliseconds or a time frame according to parameter **8**.

**How is the peak detection performed?**

The module uses the Matlab function *findpeaks*. It specifically looks for a high value surrounded by lower values. This detection is not affected by the boundaries of the interval (which are never identified as peaks). This function only detects local maxima. To detect local minima, the signal is inverted. This is why it is not possible to detect a positive peak (max) and a negative peak (min) at the same time. You must perform two separate runs, changing option **5**.

**What happens if there are several local peaks?**

The Find Peaks module will always return a single peak in the output file. For positive peaks, it will report the peak with the largest amplitude; for negative peaks, it will report the peak with the smallest amplitude. However, the check file informs you if more than one peak has been found within the specified time interval (point **3**).

**How can I use the check file?**

The optional check file allows you to know if only one peak is detected in the interval (**value=0**), no peak is detected (**value=1**) or several peaks are detected (**value=2**). You can use this information to look at the signal yourself in Cartool to possibly correct the position according to your own expertise.

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Description automatically generated

Timecourse export

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1. Sampling rate of the data. This should normally be transferred automatically from the main window. You can adjust this if necessary.
2. Choose the action *‘Timecourse extraction’.*
3. Specify the time interval. Warning: the value 0 marks the start of the file. This module does not take into account a possible .mrk file that could define a different origin.   
   EoF=End of File.
4. Specify the electrode to be studied and the indices of the electrodes (1, 2, 64), rather than the names specified by the coordinate file (A1, A2, B32). A ‘\*’ means that all the electrodes will be study.
5. In addition, you can compute the average of specified electrodes in **4**, on the global field power (GFP) or on the global map dissimilarity (GMD).
6. elect the output file format: .xlsx, .txt (with tabs as delimiters) or .ep (to open in Cartool without electrode names). This option is only available if you choose to save the 2D file in the parameters **7**.
7. You have two choices here:
   1. 2D files: Record the timecourse separately for each input file. This will produce a table with a header. Each line represents a temporal point (tf) and each column represents a channel (first the electrodes, then the average, the GFP and the GMP)
   2. 3D files. All timecourses will be recorded in a single file. This will create a .mat file without headers. The file is organized as follows:
      1. x=time (in tf)
      2. y=channels (first electrodes, then avg, GFP and GMP)
      3. z=input files
8. Select the output folder where the output will be recorded.
9. You can specify manually a suffix in the output file name. For example: *P032\_epoched\_Trigger\_1\_Timecourse\_100-1540tf\_POI2.xlsx for only one file* if you select ‘2D files’ in **7**, or *3D\_Timecourse\_100-1540tf\_POI2.mat* if you selected ‘3D file’ in **7**.
10. There are three validation buttons:
    1. The **Run** button will carry out the processing parameterized in the module.
    2. The **Save in memory** button will store all the parameters in memory and close the module without performing the processing.
    3. The button **Cancel** closes the module without processing and without keeping the entered parameters in memory. The same effect will be achieved by closing the module window.

FAQ

**What the output looks like?**

In the case of ‘2D files’:

A table of numbers and letters

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In case of a ‘3D file’:

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**Should I use a ‘2D files’ or ‘3D file’ ?**

depends on the programme you are going to use to open it. The 2D file is a standard matrix that is easier to work with using software such as Excel or SPSS. However, having multiple files could make performing statistics more difficult. This is why the option to have a 3D file containing all the data has been added to this module. This would be suitable for the R software, which can import .mat files using the R.matlab library:

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Average time interval

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1. Sampling rate of the data. This should normally be transferred automatically from the main window. You can adjust this if necessary.
2. Choose the action *‘Average time interval’.*
3. Specify the time interval. Warning: the value 0 marks the start of the file. This module does not take into account a possible .mrk file that could define a different origin.   
   EoF=End of File.
4. Specify the electrode to be studied and the indices of the electrodes (1, 2, 64), rather than the names specified by the coordinate file (A1, A2, B32). A ‘\*’ means that all the electrodes will be study.
5. In addition, you can compute the average of specified electrodes in **4**, on the global field power (GFP) or on the global map dissimilarity (GMD).
6. Select if the output file format: .xlsx or .txt (with tab as delimiter).
7. Select the output folder where the output will be recorded.
8. You can specify manually a suffix in the output file name. For example: *AverageTimeInterval\_200-250tf\_POI1.xlsx*
9. There are three validation buttons:
   1. The **Run** button will carry out the processing parameterized in the module.
   2. The **Save in memory** button will store all the parameters in memory and close the module without performing the processing.
   3. The button **Cancel** closes the module without processing and without keeping the entered parameters in memory. The same effect will be achieved by closing the module window.

FAQ

**What the output looks like?**

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