

INSTRUCTION SHEET OF THE SOFTWARE :

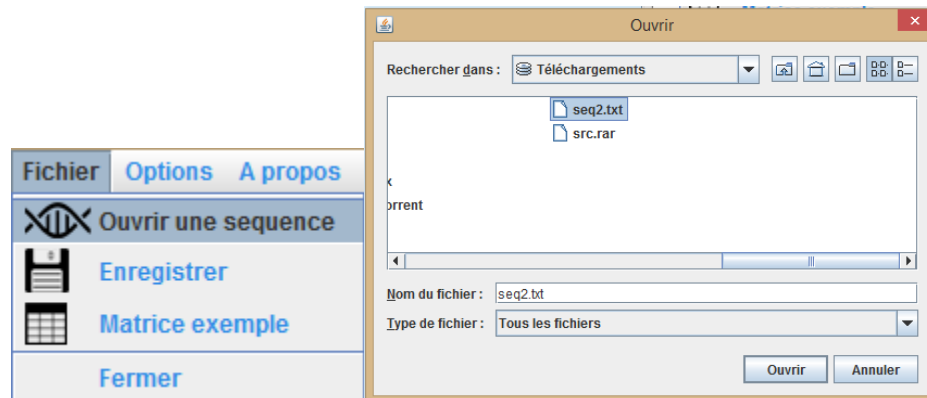
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Start :

- 1) Open a sequence from a text file or a FASTA file.

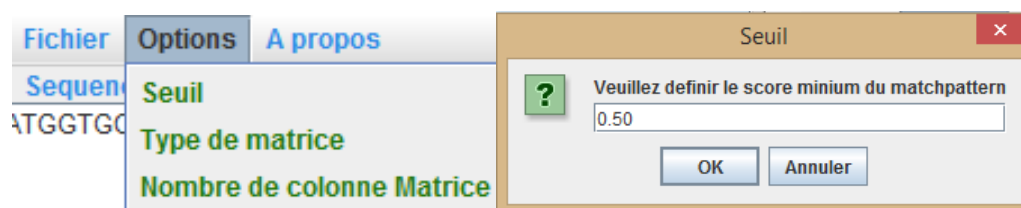


- 2) An example matrix of 5 columns can be used.
You can also generate the matrix you want.



Parameters :

The options menu allow you to chose the treshold, the type of the matrix (Entropy ou LogOdd) and the number of columns of your matrix.



The parameters chosen are visible on the launch button.



Clic on **Lancer** to launch the analysis of the sequence.
The analysis can be turned off with the **Stop** button.

The **Clear** button allow you to clear the results generated and delete the sequence analysed.

The results can be saved in a file : Clic on **Fichier** then in the menu select **Enregistrer** to save the results.