Aligning Sequences

The Alignment Editor integrates several popular multiple sequence alignment algorithms. Below is the list of available algorithms and links to the documentation:

- Port of the popular MUSCLE3 algorithm.
- KAlign plugin: effective work with huge alignments.
- ClustalW and MAFFT: these algorithms appeared in the version 1.7.2 of UGENE with the External Tools plugin.
- T-Coffee: this alignment algorithm is available since version 1.8.1 of UGENE with the External Tools plugin.

To align sequences choose a preferred alignment method in the Actions main menu, in the context menu or by Align main toolbar button.

Also you may find useful the following video tutorials devoted to the multiple sequence alignment:

- Making a multiple sequence alignment from FASTA file
- Working with large alignments in UGENE
- Performing profile-to-profile and profile-to-sequence MUSCLE alignments
- Running remote MUSCLE task