## **Align with ClustalW Element**

Aligns multiple sequence alignments (MSAs) supplied with ClustalW.

ClustalW is a general purpose multiple sequence alignment program for DNA or proteins. Visit http://www.clustal.org/ to learn more about it.



Clustal is used as an external tool from UGENE and it must be installed on your system. To learn more about the external tools, please, read main UGENE User Manual.

## Parameters in GUI

Parameter	Description	Default value
Weight matrix	For proteins it is a scoring table which describes the similarity of each amino acid to each other. For DNA it is the scores assigned to matches and mismatches.	default
End gaps	The penalty for closing a gap.	False
Gap distance	The gap separation penalty. Tries to decrease the chances of gaps being too close to each other.	4.42
Gap extension penalty	The penalty for extending a gap.	8.52
Gap open penalty	The penalty for opening a gap.	53.90
Hydrophilic gaps off	Hydrophilic gap penalties are used to increase the chances of a gap within a run (5 or more residues) of hydrophilic amino acids.	False
Residue-specific gaps off	Residue-specific penalties are amino specific gap penalties that reduce or increase the gap opening penalties at each position in the alignment.	False
Iteration type	Alignment improvement iteration type.	None
Number of iterations	The maximum number of iterations to perform.	3
Tool path (require d)	Path to the ClustalW tool. The default path can be set in the UGENE Application Settings.	default
Temporary directory	Directory to store temporary files.	default

## Parameters in Workflow File

Type: clustalw

Parameter	Parameter in the GUI	Туре
matrix	Weight matrix	numeric
		Available values are:  • 0 - for IUB  • 1 - for ClustalW  • 2 - for BLOSUM  • 3 - for PAM  • 4 - for GONNET  • 5 - for ID  • -1 - for default matrix
close-gap-penalty	End gaps	boolean
gap-distance	Gap distance	numeric
gap-ext-penalty	Gap extension penalty	numeric
gap-open-penalty	Gap open penalty	numeric

no-hydrophilic-gaps	Hydrophilic gaps off	boolean
no-residue-specific-gaps	Residue-specific gaps off	boolean
iteration-type	Iteration type	numeric
		Available values are:
		<ul><li>0 - for None</li><li>1 - for Tree</li><li>2 - for Alignment</li></ul>
iterations-max-num Number of iterations		numeric
path	Tool path	string
temp-dir	Temporary directory	string

## Input/Output Ports The element has 1 input port.

Name in GUI: Input MSA

Name in Workflow File: in-msa

Slots:

Slot In GUI	Slot in Workflow File	Type
MSA	msa	msa

And 1 output port.

Name in GUI: ClustalW result MSA Name in Workflow File: out-msa

Slots:

Slot In GUI	Slot in Workflow File	Туре
MSA	msa	msa