



# AlphaMap Tutorial

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This step-by-step guide helps you to get started with our software AlphaMap.

## Table of Contents

.....	- 1 -
<b><i>Program Description</i></b> .....	- 1 -
<b><i>Installation</i></b> .....	- 2 -
<b>Windows</b> .....	- 2 -
<b>MacOS</b> .....	- 4 -
<b><i>How to use the AlphaMap GUI</i></b> .....	- 7 -

## Program Description

AlphaMap enables the exploration of proteomic datasets on the peptide level. It is possible to evaluate the sequence coverage of any identified protein and its post-translational modifications (PTMs). AlphaMap further integrates all available UniProt sequence annotations as well as information about proteolytic cleavage sites.

# Installation

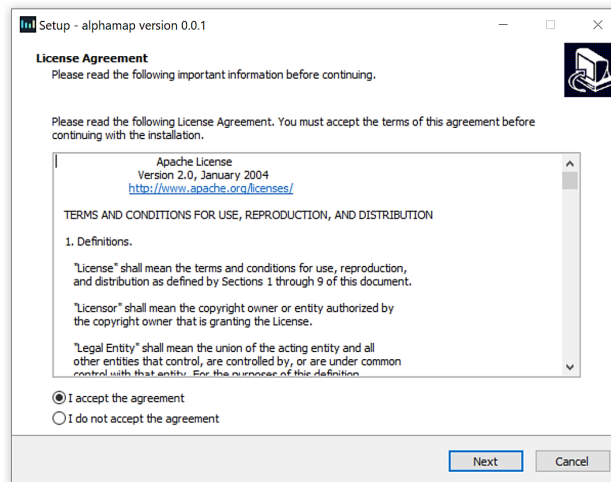
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## Windows

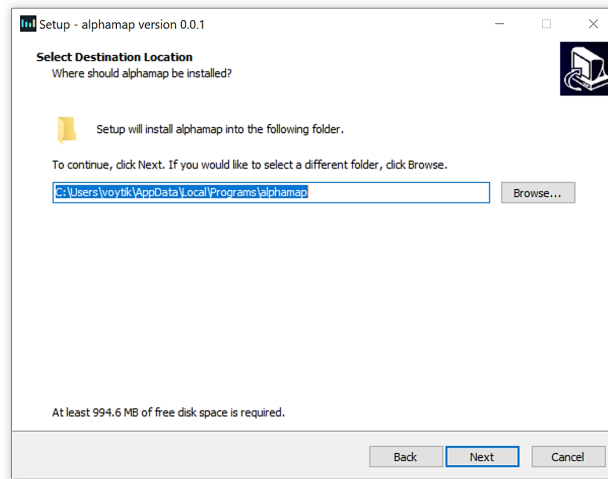
**Prerequisites:** **Windows 10** (a system update might be necessary in case older versions do not work)

**Important:** To prevent installation errors on Windows, we recommend uninstalling any previous AlphaMap version before installing a new one.

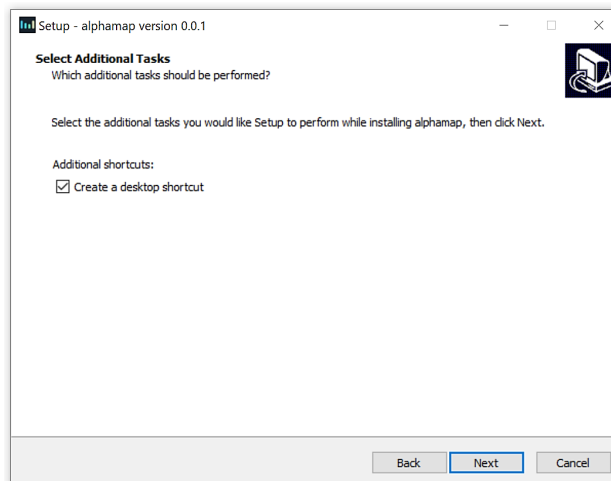
1. Download [the latest release](#) for Windows (alphamap\_installer\_windows.exe) from the GitHub repository and open the .exe file.
2. In the “User Account Control” dialog asking about permission for the app to make changes to your device press the “Yes” button.
3. In the appearing “Setup – alphamap version X.X.X” dialog window accept the License Agreement and press the “Next” button.



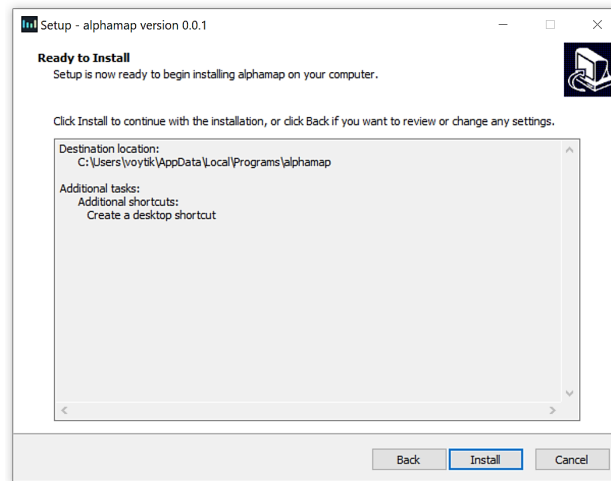
4. Select the destination location for the installation of AlphaMap software (the size of the whole package is 994.6 MB) and press the “Next” button.



5. In the next dialog window mark the "Create a desktop shortcut" check box and press the "Next" button.



6. Check the setting and if everything is correct, press "Install" button. You may go back to change some settings using the "Back" button or "Cancel" the installation.



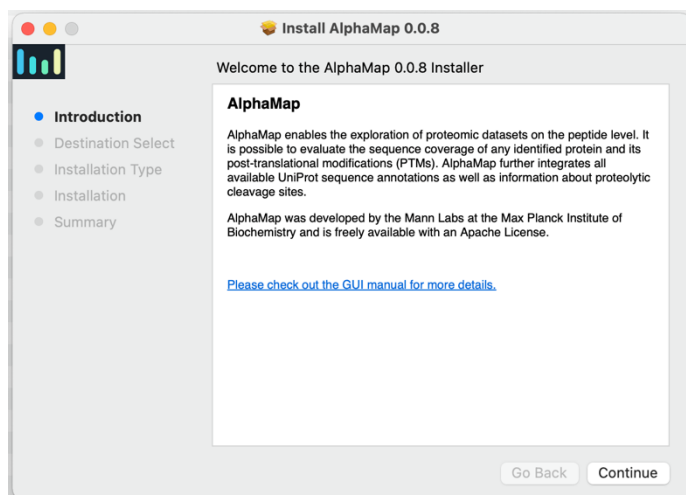
7. Wait till the installation process is finished and with the marked “Launch alphamap” check box press the “Finish” button.
8. In the appearing “Windows Security Alert” dialog window press the “Allow access” button that will prevent the Windows Defender Firewall from blocking the AlphaMap tool on your PC.
9. Check your default browser (Google Chrome or Mozilla Firefox are suggested for the fast running of the AlphaMap) and start working with the tool.

\* If you install AlphaMap for all users, you might need admin privileges to run it (right-click on the AlphaMap logo on your desktop and select "Run as administrator").

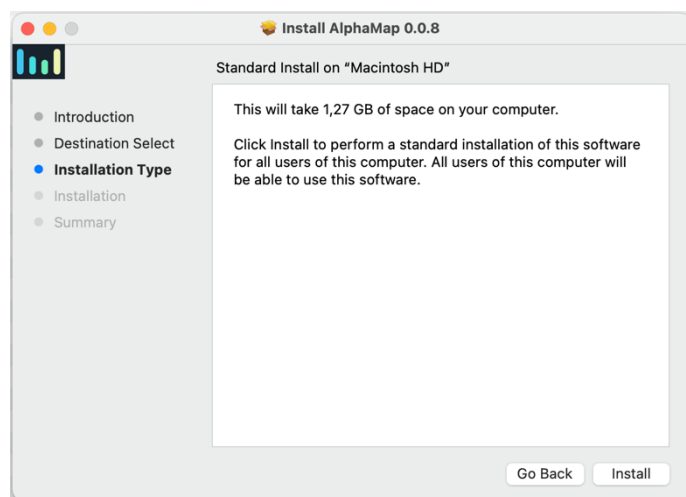
## MacOS

**Prerequisites:** at least **macOS Catalina (10.15) or higher** (a system update might be necessary in case older versions do not work)

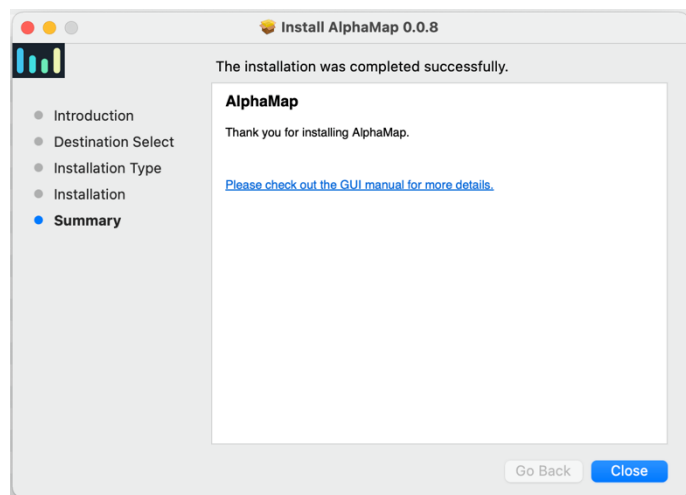
1. Download [the latest release](#) for macOS (alphamap\_gui\_installer\_macos.pkg) from the GitHub repository and open the .pkg file.
2. Click “Continue” on the appearing “Install AlphaMap X.X.X” dialog window.



3. Click “Install” to start the installation. This might take a few minutes.

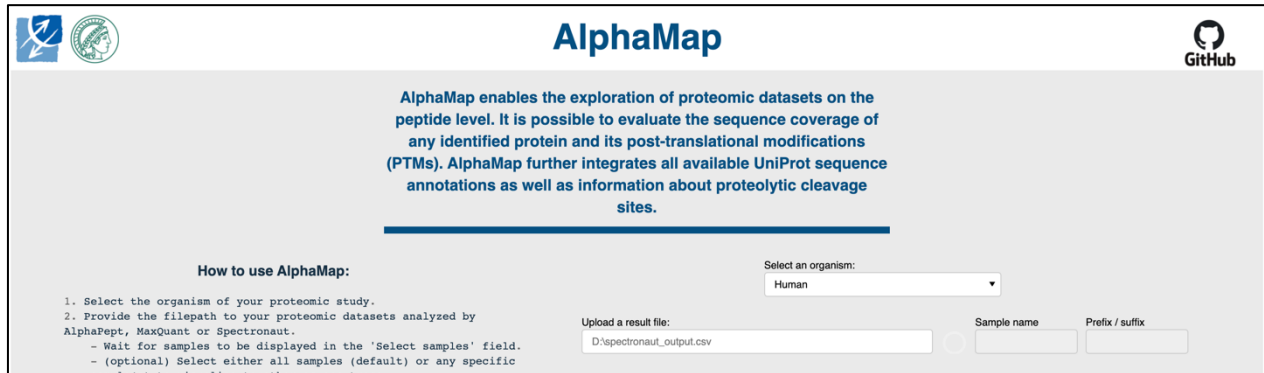


4. Click “Close” to quit the installation menu. AlphaMap is now available in your applications folder on your MacOS.



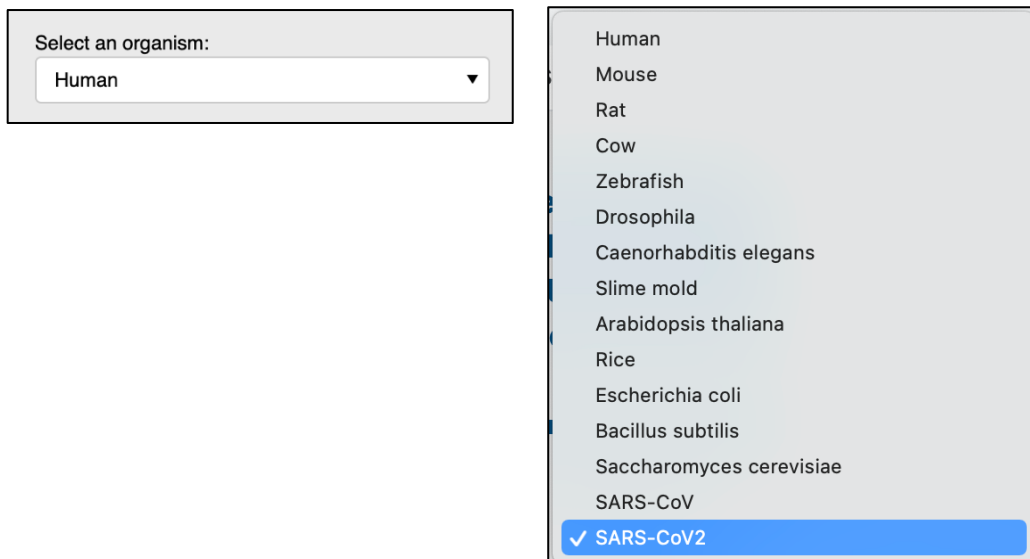
\* In rare cases, you might get an error message that the “alphamap\_gui\_install\_macos.pkg” cannot be opened because it is coming from an unidentified developer. This error can be avoided by pressing the OPTION key or by going to ‘Security and Privacy’ under ‘System Preferences’ and enable AlphaMap installation.

# How to use the AlphaMap GUI



The screenshot shows the AlphaMap web interface. At the top, there are logos for the project and GitHub. The main heading is 'AlphaMap'. Below it, a paragraph describes the tool's capabilities: 'AlphaMap enables the exploration of proteomic datasets on the peptide level. It is possible to evaluate the sequence coverage of any identified protein and its post-translational modifications (PTMs). AlphaMap further integrates all available UniProt sequence annotations as well as information about proteolytic cleavage sites.' Below this, a section titled 'How to use AlphaMap:' provides instructions: 1. Select the organism of your proteomic study. 2. Provide the filepath to your proteomic datasets analyzed by AlphaPept, MaxQuant or Spectronaut. It also includes sub-instructions: '- Wait for samples to be displayed in the 'Select samples' field.' and '- (optional) Select either all samples (default) or any specific samples to display in the 'Select samples' field.' To the right of the instructions, there is a 'Select an organism:' dropdown menu with 'Human' selected. Below this, there is an 'Upload a result file:' section with a text input field containing 'D:\spectronaut\_output.csv'. To the right of this field are two more input fields: 'Sample name' and 'Prefix / suffix'.

1. First select the organism of your proteomic study. Currently, the [13 most popular organisms based on UniProt](#) are available for selection, including: Human [Taxon identifier=9606], Mouse [10090], Rat [10116], Cow [9913], Zebrafish [7955], Drosophila [7227], Caenorhabditis elegans [6239], Slime mold [44689], Arabidopsis thaliana [3702], Rice [39947], Escherichia coli (strain K12) [83333], Bacillus subtilis (strain 168) [224308], Saccharomyces cerevisiae (strain ATCC 204508 / S288c) [559292]. Additionally, the reviewed proteomes of SARS-CoV [694009] and SARS-CoV2 [2697049] are available.



This image shows a close-up of the 'Select an organism:' dropdown menu. The menu is open, displaying a list of organisms. The organisms listed are: Human, Mouse, Rat, Cow, Zebrafish, Drosophila, Caenorhabditis elegans, Slime mold, Arabidopsis thaliana, Rice, Escherichia coli, Bacillus subtilis, Saccharomyces cerevisiae, SARS-CoV, and SARS-CoV2. The SARS-CoV2 option is highlighted with a blue background and a checkmark icon, indicating it is the selected organism.

2. Upload your proteomic datasets analyzed by AlphaPept, MaxQuant, Spectronaut, DIA-NN or FragPipe:
  - a) Provide the filepath to the result file in the “Upload a result file:” field, e.g. “D:\alphapept\_output.csv” (Windows) or “/Users/test/alphapept\_output.csv” (MacOS).
  - b) Wait for samples to be displayed in the “Select samples” field. The loading process is indicated by a spinner symbol.
  - c) (optional) Select either all samples (default) or any specific sample(s) to visualize together as one trace.
  - d) (optional) Choose a name by which the selected sample(s) will be displayed in the figure. If no name is provided, the original names of all selected samples will be concatenated by semicolon. If ‘all samples’ were selected, the filename will be the default name.
  - e) (optional) Provide a prefix or suffix to be removed from the original names of the selected samples. This option only applies if no user defined name is provided (see d).
  - f) Up to three datasets or sets of selected samples can be visualized together in the GUI. For this, use the “Upload additional result files” option.

The screenshot shows a web-based interface for uploading proteomic data. At the top, there is a dropdown menu labeled "Select an organism:" with "SARS-CoV2" selected. Below this, on the left, is a text input field labeled "Upload a result file:" (annotated with 'a') containing the path "/Users/isabell/Desktop/data/teome\_data.csv". To the right of this field is a spinner icon (annotated with 'b'). Further right is a text input field labeled "Sample name" (annotated with 'd') containing the word "teome". To the right of that is another text input field labeled "Prefix / suffix" (annotated with 'e'). Below the "Upload a result file:" field is a section labeled "Select samples:" (annotated with 'c'). This section contains a list box with the following items: "All samples", "20200814\_EXPL2\_OzKa\_SA\_COVID19\_DIA\_teome\_1", "20200814\_EXPL2\_OzKa\_SA\_COVID19\_DIA\_teome\_8", "20200814\_EXPL2\_OzKa\_SA\_COVID19\_DIA\_teome\_9", and "20200814\_EXPL2\_OzKa\_SA\_COVID19\_DIA\_teome\_10". At the bottom of the interface is a button labeled "Upload additional result files" (annotated with 'f') with a plus sign icon to its right.



f
Upload additional result files
—

Upload a result file:

☐

Sample name

Prefix / suffix

Select samples:

All samples

20200928\_EXPL0\_OzKa\_SA\_Covid\_Phospho\_DIA\_1

20200928\_EXPL0\_OzKa\_SA\_Covid\_Phospho\_DIA\_2

20200928\_EXPL0\_OzKa\_SA\_Covid\_Phospho\_DIA\_3

20200928\_EXPL0\_OzKa\_SA\_Covid\_Phospho\_DIA\_4

Upload a result file:

☐

Sample name

Prefix / suffix

Select samples:

All samples

20200830\_EXPL2\_FyHa\_SA\_COVID19\_DIA\_diGly\_1

20200830\_EXPL2\_FyHa\_SA\_COVID19\_DIA\_diGly\_8

20200830\_EXPL2\_FyHa\_SA\_COVID19\_DIA\_diGly\_9

20200830\_EXPL2\_FyHa\_SA\_COVID19\_DIA\_diGly\_10

- \* If you would like to choose different samples from the same result file, you need to provide the same filepath and select the different samples.
- \* To remove the result file that was previously uploaded, just remove its path from the "Upload a result file:" field (2a step).
- \* If you cannot upload the selected file, please take a look at the detailed instructions for AlphaPept, MaxQuant, Spectronaut, DIA-NN and FragPipe input formats.

Spectronaut instructions
—

The data needs to be exported in the **normal long** format as .tsv or .csv file.

It needs to include the following columns:

- PEP.AllOccurringProteinAccessions
- EG.ModifiedSequence
- R.FileName

To ensure the correct export format from Spectronaut, you can download and apply the provided export scheme "spectronaut\_export\_scheme.rs".

MaxQuant instructions
—

To visualize the proteins which were analyzed by the MaxQuant software please use the **evidence.txt** file.

The following columns from the file are used for visualization:

- Proteins
- Modified sequence
- Raw file

- 9 -

AlphaPept instructions

To visualize the proteins which were analyzed by the AlphaPept software please use the **results.csv** file.

The following columns from the file are used for visualization:

- protein\_group
- sequence
- shortname

DIA-NN instructions

To visualize the proteins which were analyzed by the DIA-NN software please use the **{experiment\_name}.tsv** file.

The following columns from the file are used for visualization:

- Protein.Ids
- Modified.Sequence
- Run

FragPipe instructions

There are two options to visualize data analyzed by FragPipe:

1) Upload individual **"peptide.tsv"** files for single MS runs. In this case, the following columns from the original file are used for visualization:

- Protein ID
- Peptide
- Assigned Modifications

2) Upload the **"combined\_peptide.tsv"** file with the joint information about peptides identified in all runs (there is an option to select the experiment(s)). Be aware that the combined\_peptide.tsv does not provide information about PTM localization. PTMs are therefore not shown for this option. Following columns are used for visualization:

- Protein ID
- Sequence
- All 'Spectral Count' columns containing information about individual experiments

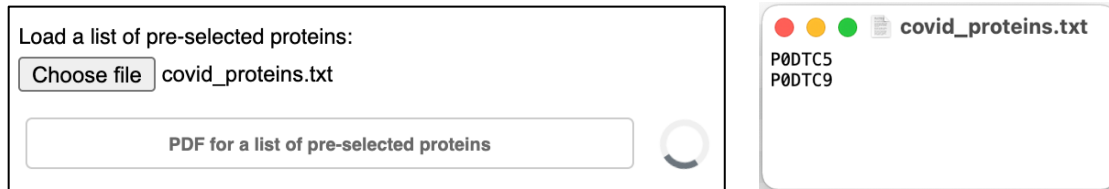
- Press the "Upload Data" button. The loading process is indicated by a spinner symbol.
- Select a protein of interest. Per default, you can choose from all UniProt accessions. Click the "Search by a gene name" option to select proteins by their gene name.

Select a protein of interest:

P0DTC5

☒ Search by UniProt accession
☐ Search by a gene name

5. (optional) Load a list of pre-selected proteins. This can be a .txt file containing either UniProt accessions or gene names containing one identifier per line. This will reduce the options available in the selection of proteins of interest (step 4). When the list of proteins is loaded, there is an option to generate a pdf report with plots for all pre-defined proteins. The report is intended for fast screening of the data. Note that we generally recommend the interactive visualization and the download of single .svg files as described in step 7a. The report generation process is indicated by a spinner symbol.



**IMPORTANT!** Once you uploaded the list of pre-defined proteins, the original list of proteins in the uploaded result file(s) will be filtered. Proteins that are not contained in the list of pre-defined proteins will no longer be available for visualization. If you would like to use the whole list again, repeat from step 2.

6. Select annotation options for the sequence visualization.
- All sequence annotations from UniProt are available and displayed per default. You can choose a customized set of displayed annotations in the “UniProt annotation” selection.

UniProt annotations							—
Molecule processing							—
Chain	Initiator methionine	Peptide	Propeptide	Signal peptide	Transit peptide		
Post-translational modification							—
Cross-link	Disulfide bond	Glycosylation	Lipidation	Modified residue			
Family & Domain							—
Coiled coil	Compositional bias	Domain	Motif	Region	Repeat	Zinc finger	
Subcellular location							—
Intramembrane		Topological domain		Transmembrane			
Function							—
Active site	Binding site	Calcium binding	DNA binding	Metal binding	Nucleotide binding	Site	
Sequence							—
Alternative sequence	Natural variant	Non-adjacent residues	Non-standard residue	Non-terminal residue	Sequence conflict	Sequence uncertainty	
Other options							—
Secondary structure			Mutagenesis				
<input type="checkbox"/> Select all <input type="checkbox"/> Clear all							

- All theoretical cleavage sites for the most common proteases can be shown. Trypsin is selected by default. Alternative or additional proteases can be selected in the “Protease cleavage sites” selection.

Protease cleavage sites —

☐ arg-c  
☐ asp-n  
☐ bnps-skatole  
☐ caspase 1  
☐ caspase 2  
☐ caspase 3  
☐ caspase 4  
☐ caspase 5  
☐ caspase 6  
☐ caspase 7  
☐ caspase 8  
☐ caspase 9  
☐ caspase 10  
☐ chymotrypsin high specificity  
☐ chymotrypsin low specificity  
☐ clostripain  
☐ cnbr  
☐ enterokinase  
☐ factor xa  
☐ formic acid  
☐ glutamyl endopeptidase  
☐ granzyme b  
☐ hydroxylamine  
☐ iodosobenzoic acid  
☐ lysc  
☐ ntcb  
☐ pepsin ph1.3  
☐ pepsin ph2.0  
☐ proline endopeptidase  
☐ proteinase k  
☐ staphylococcal peptidase i  
☐ thermolysin  
☐ thrombin  
☐ trypsin\_full  
☐ trypsin\_exception  
☐ non-specific  
☒ trypsin

☐ Select all
☐ Clear all

\* You can use “Select all” or “Clear all” checkboxes to speed up the selection process.

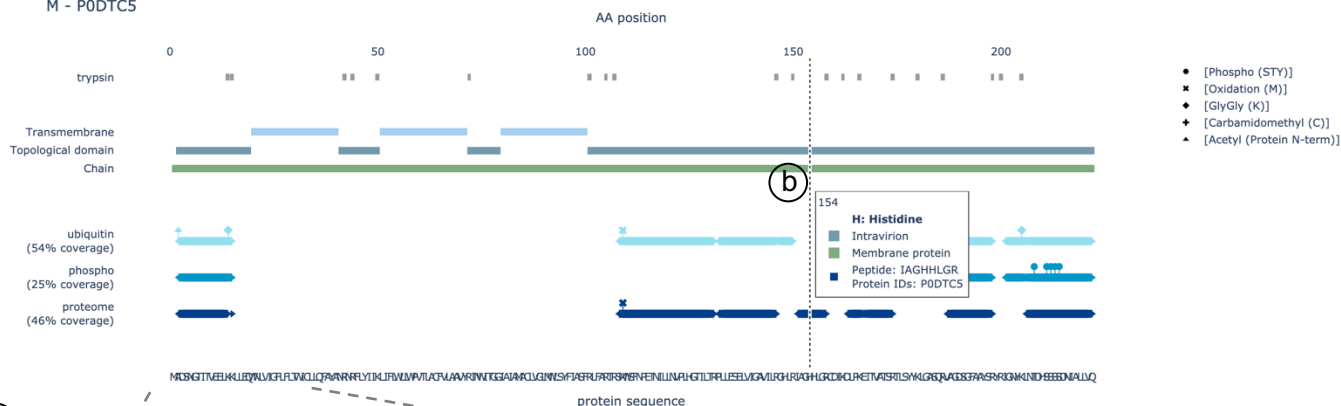
7. Press the "Visualize Protein" button. The loading process is indicated by a spinner symbol.

- a) You can use the interactive toolbar to for example zoom in and out or to highlight specific sequence regions. Press the little camera icon to download a high-resolution .svg image of the currently displayed protein and sequence region.
- b) If you hover over the sequence, all annotation information for the current sequence position of the cursor will be displayed.
- c) You can directly visit other websites for further exploration of details on the selected protein of interest. UniProt, PhosphoSitePlus, Protter, PDB an Peptide Atlas are available for direct access.

a

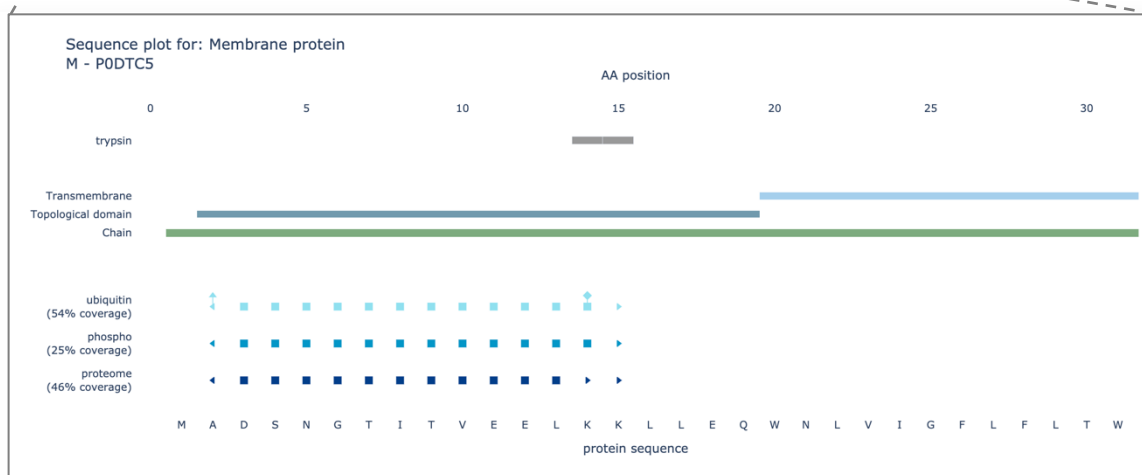


Sequence plot for: Membrane protein  
M - PODTC5



c

Inspect target protein on other platforms:



Here we show an example sequence representation of the 'Membrane protein' (PODTC5) of SARS-CoV-2. The full sequence is shown in the upper part of the figure and a zoom-in on is provided at in the bottom panel. Amino acids covered by an experimentally observed peptide are indicated by squares. N- and C-terminal peptide ends are illustrated by left- and right-pointed triangles instead. PTM modifications are indicated by stick representations on top of the of the amino acid (square or triangle) they are localized on. In the zoom-in you can see an N-terminal acetylation (hat) coinciding with a peptide start (left-pointed triangle) and a ubiquitination (diamond) on lysine 14 of the ubiquitin track. You can also see that two alternatively cleaved peptides were detected in the proteome track, one ending on lysine 14 and one ending with lysine 15.

In the example figure above, intraviron regions are well covered by the experimental data, whereas none of the observed peptides map to the helical trans-membrane domains. Both

phosphorylations ('Phospho (STY)', indicated by circles) and ubiquitinations ('GlyGly (K)', indicated by diamonds) could be identified on the viral protein. This is just one example of how to inspect proteomics data with AlphaMap.

**Now enjoy exploring your own data!**