
pTop User Guide

Version 1.0



pFind.ict.ac.cn

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1 Installation

1.1 Installation requirements

Hardware requirements

2 GB or higher recommended memory

Software requirements

Windows 7 or above

microsoft .NET Framework 4.5 or above

Xcalibur (2.1 or above) or MSFileReader

1.2 Installation steps

The Windows setup package of pTop can be downloaded from the website <http://pfind.ict.ac.cn>. Before installation, please fill in a registered table and send it to rxsun@ict.ac.cn to get a registration key.

The pTop setup package includes not only pTop, but also pXtract, pParseTD, pConfig and pLabel. pXtract creates MS1 and MS2 input files directly from Thermo Scientific RAW LC-MS/MS data files. pParseTD converts the MS1 and MS2 files to MGF files, in which detecting the relative accurate mono mass of the precursors and deconvoluting and deisotoping the MS/MS. pConfig is a tool that can add or change the basic configurations, such as amino acids, modifications. pLabel is a spectra labeling tool that can visualize the global- and local-view proteoform-spectrum matches, given the results of pTop or any other search engines. pLabel can label both CID and ETD spectra, and implement the manual de novo sequencing.

To install pTop on windows, the following simple steps are needed.

Step 1: Select the installer language (**Figure 1**). Now it only supports English and Chinese (Simplified).

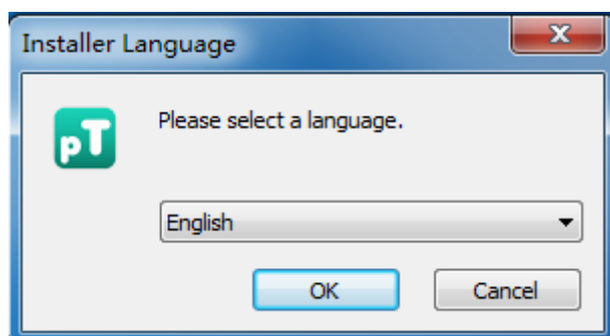


Figure 1. Installer language

Step2: Click Next to start the setup (**Figure 2**).



Figure 2. Welcome to the setup wizard

Step 3: Choose the install Location (Figure 3). And D drive disk is recommended.

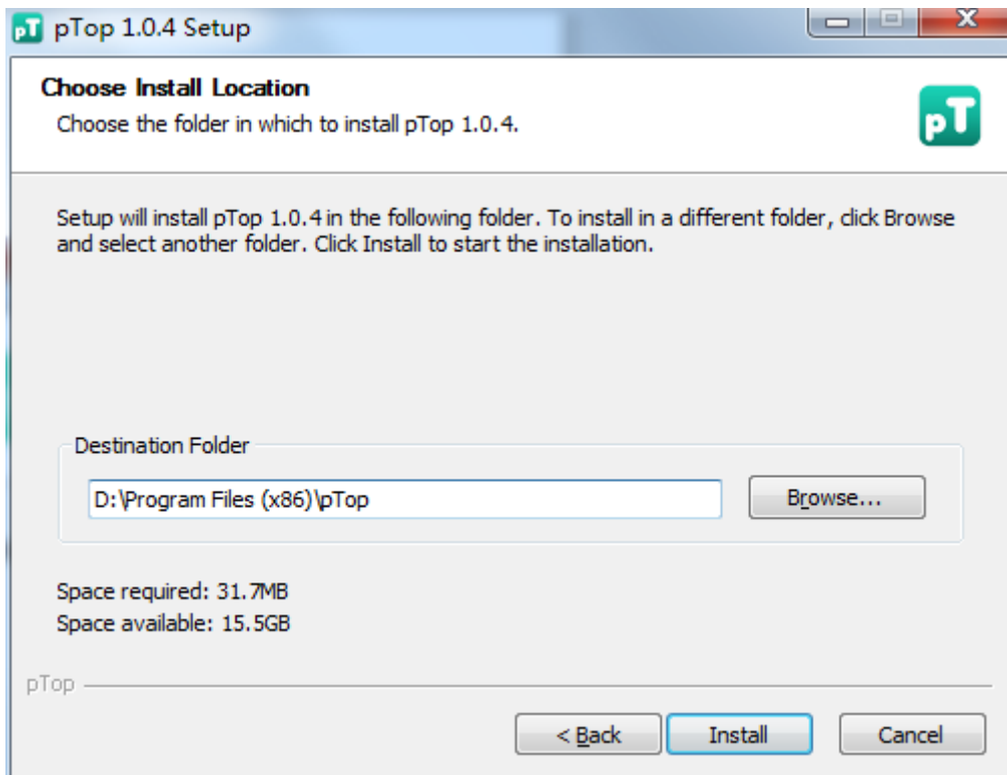


Figure 3. Choose install location

Step 4: Just wait a few seconds, the Installation will be finished (Figure 4).

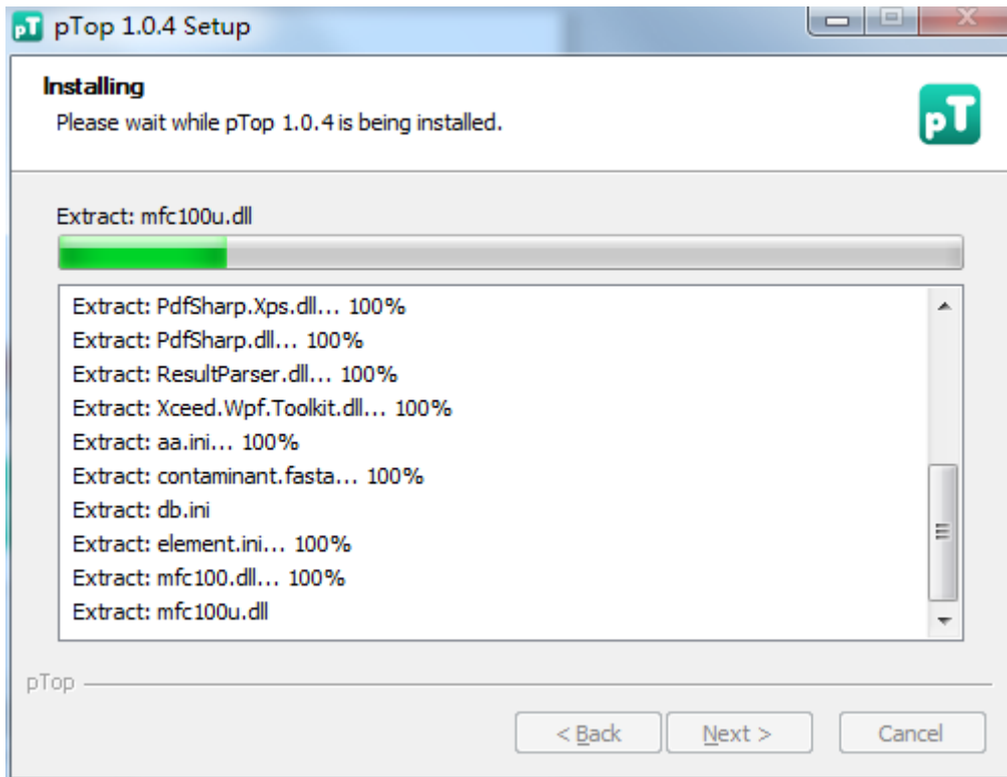


Figure 4. Installing

Finally, you can check the box of run pTop and then click Finish to start pTop (**Figure 5**).

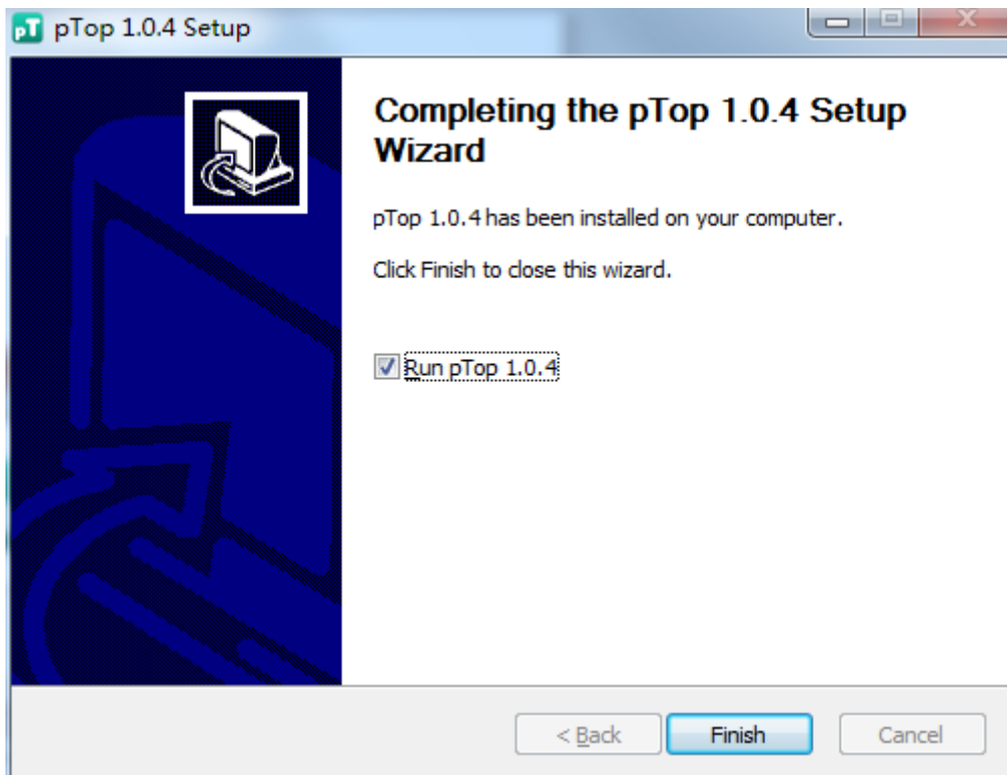
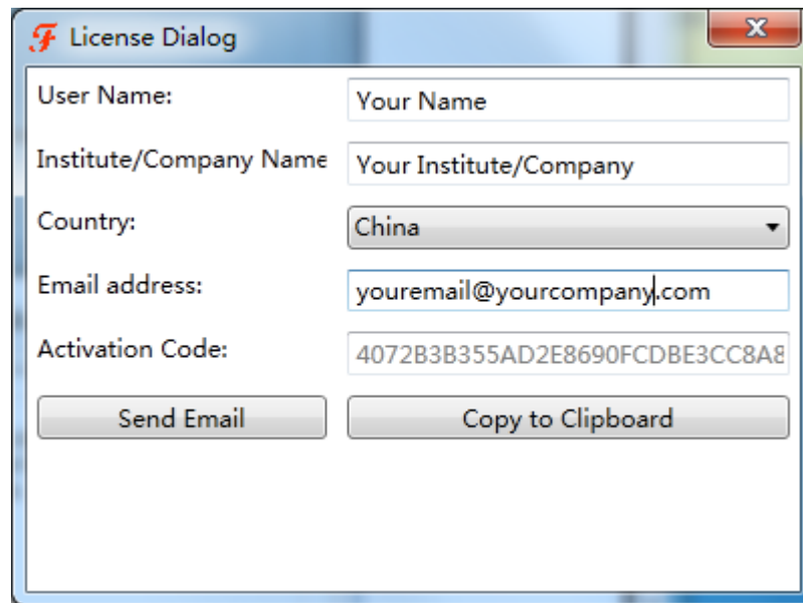


Figure 5. Installation finished

2 Activation

All users are required to go through a software activation process in order to use pTop1.0. A license wizard will appear to guide users through the activation process the first time pTop1.0 is launched.



The image shows a Windows-style dialog box titled "License Dialog". It contains several input fields and two buttons. The fields are: "User Name:" with the text "Your Name"; "Institute/Company Name" with the text "Your Institute/Company"; "Country:" with a dropdown menu showing "China"; "Email address:" with the text "youremail@yourcompany.com"; and "Activation Code:" with the text "4072B3B355AD2E8690FCDBE3CC8A8". Below the fields are two buttons: "Send Email" and "Copy to Clipboard".

Figure 6 license wizard

Please carefully fill in the user information required to get the license file on the computer that will be running pTop1.0. Your information will be useful for developers and will be strictly confidential. Thank you.

If you've already installed Microsoft Outlook, and the email address you just filled in is registered in your Outlook, then just click "Send Email"; Otherwise, click "Copy to Clipboard", then your information is copied to the clipboard, what you need to do is paste the registration information into the body of your email and send it to ptop@ict.ac.cn.

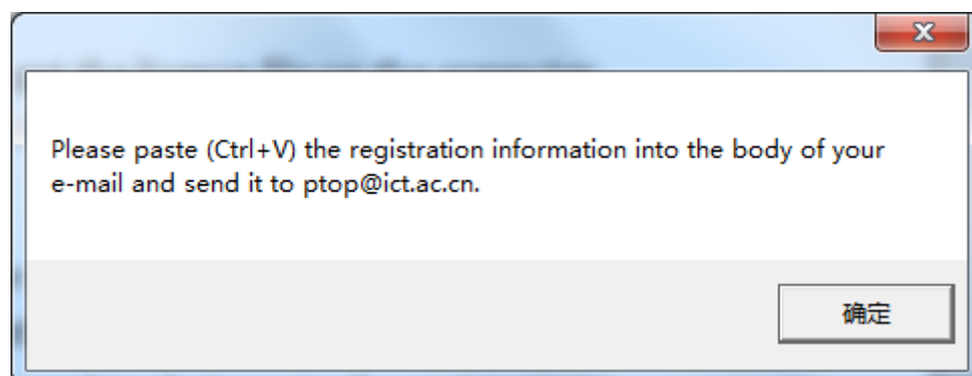


Figure 7 operation tips

Then you will get the license file. Put the license file *pTop.license* in the bin directory of your

installation directory, and you will be able to launch pTop1.0 successfully.

Important

Once the computer hardware upgraded, the license file also need to be updated.

3 Usage

3.1 Startup GUI

Double click the icon , then pTop will start up. You will see the main dialog window of pTop (**Figure 8**).

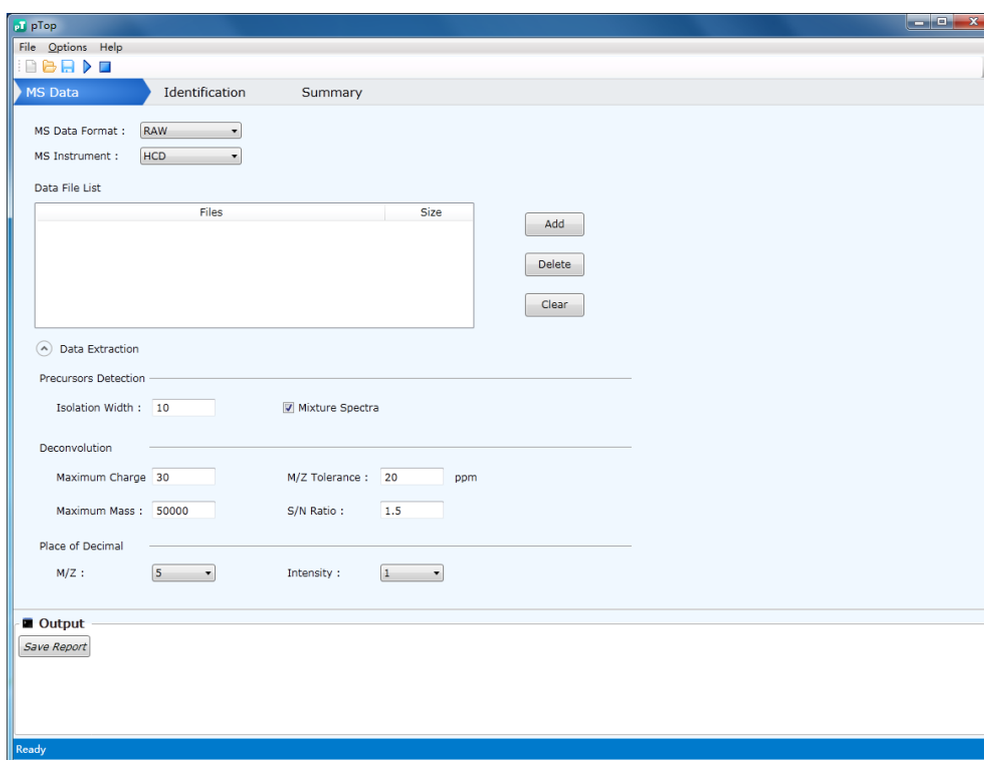


Figure 8. Main dialog window of pTop

3.2 Setting common parameters

The common parameters are listed in the 'MS Data' panel and the 'Identification' panel. How to set the common parameters will be detailed introduced as follows.

3.2.1 Spectra

The important parameters of the input spectra data are 'MS Data Format', 'MS Instrument' and 'Data File List'. (**Figure 9**)

MS Data Format

Following formats are supported by pTop: RAW, MGF and PF.

MS Instrument

Instrument determines which fragment ion series will be used for scoring. Now HCD, CID, ETD and UVPD are supported.

Data File List

Click Add to put the paths of input files in the list, the path or folder containing the tandem mass spectra.

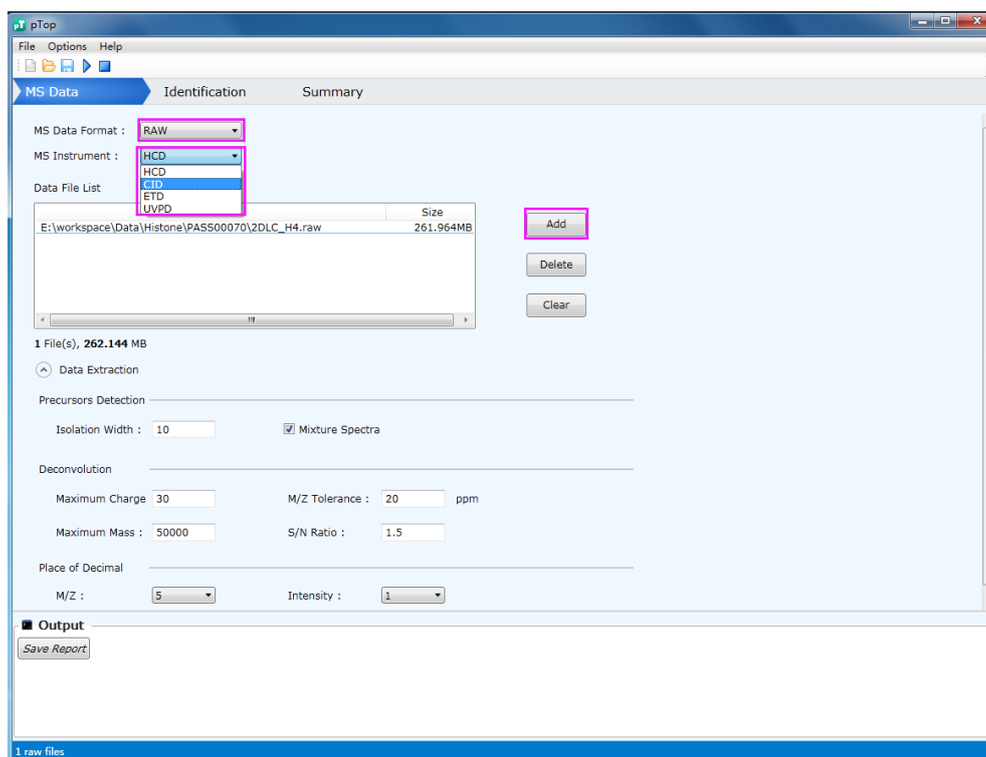


Figure 9. MS Data panel

3.2.2 Database and Mass Tolerance

For the first time you use a database, you should click 'Customize Database...' (Figure 10) to add and open the FASTA file (Figure 11). Then the database you choose will appear in the select box of database, and it will be directed chose in your subsequent search.

The precursor and fragment mass tolerance could be modified to fit the input data sets.

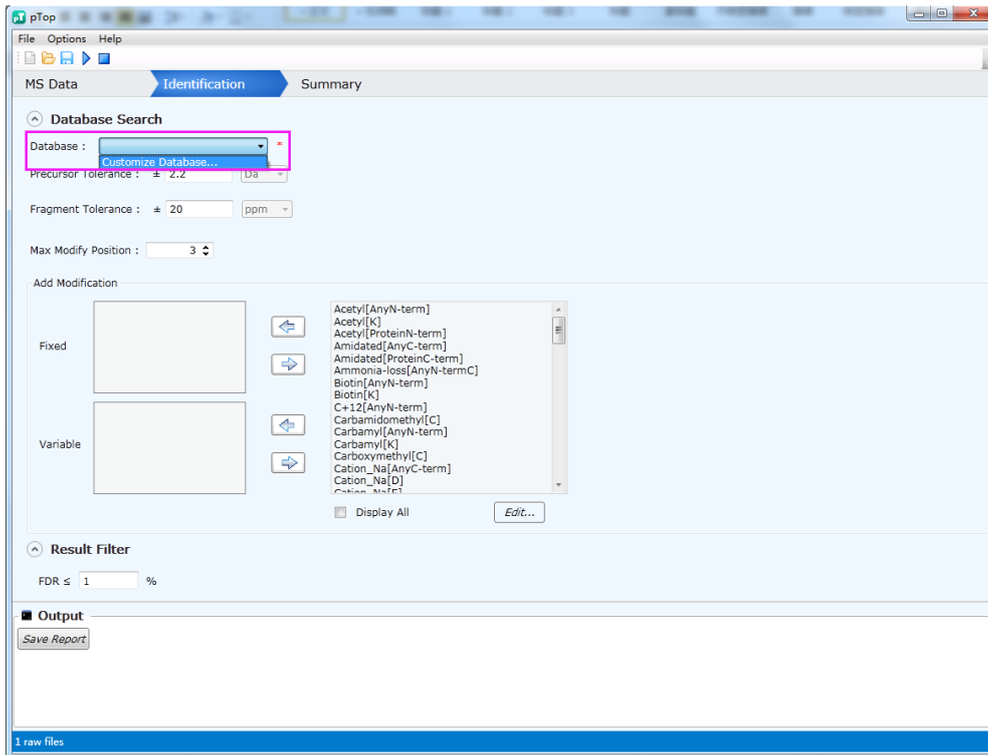


Figure 10. Choose the database

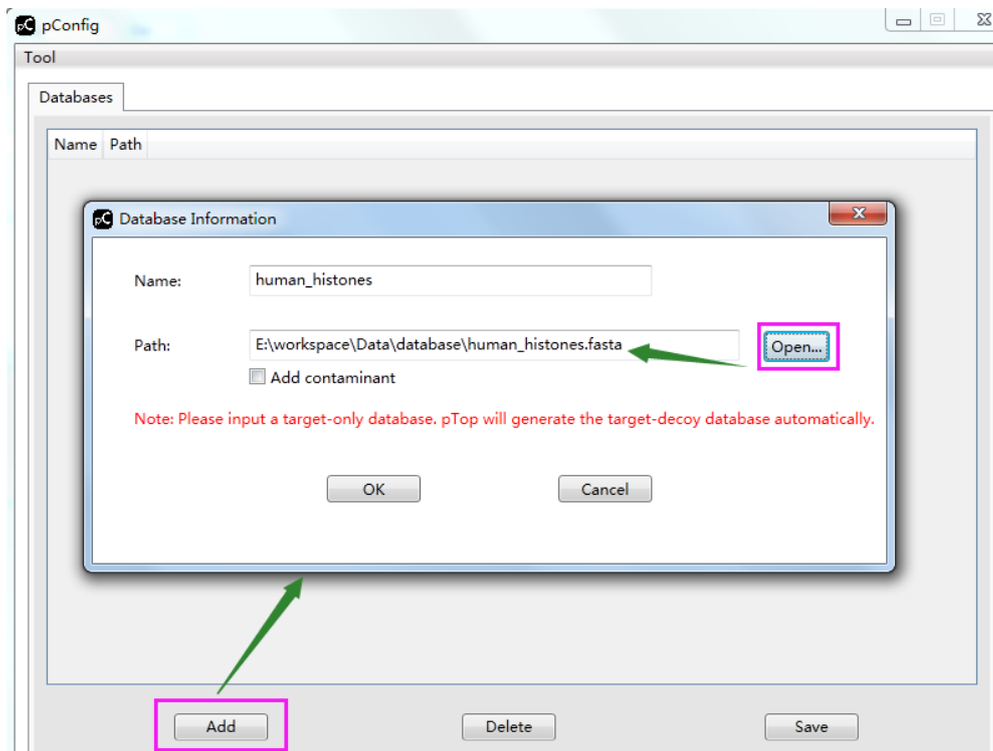


Figure 11. Add a new database

3.2.3 Modifications

pTop supports two types of modifications. Fixed modifications are applied universally, to every instance of the specified residues or terminus. Variable modifications are those which may or may not be present. A search with many variable modifications can take a much longer time than the same search with fixed modifications. The left or right arrows mean add or delete the fixed or variable modifications to the fixed and variable boxes. And you can choose the 'Max Modify Position' to set the maximum variable modifications allowed on each protein in the search. (Figure 12)

The modifications on the right side are those common ones. You can check the box of 'display all' to show all the modifications in the modification.ini file. If you still cannot find the modifications you have to add, please click 'Edit...' to add your modifications.

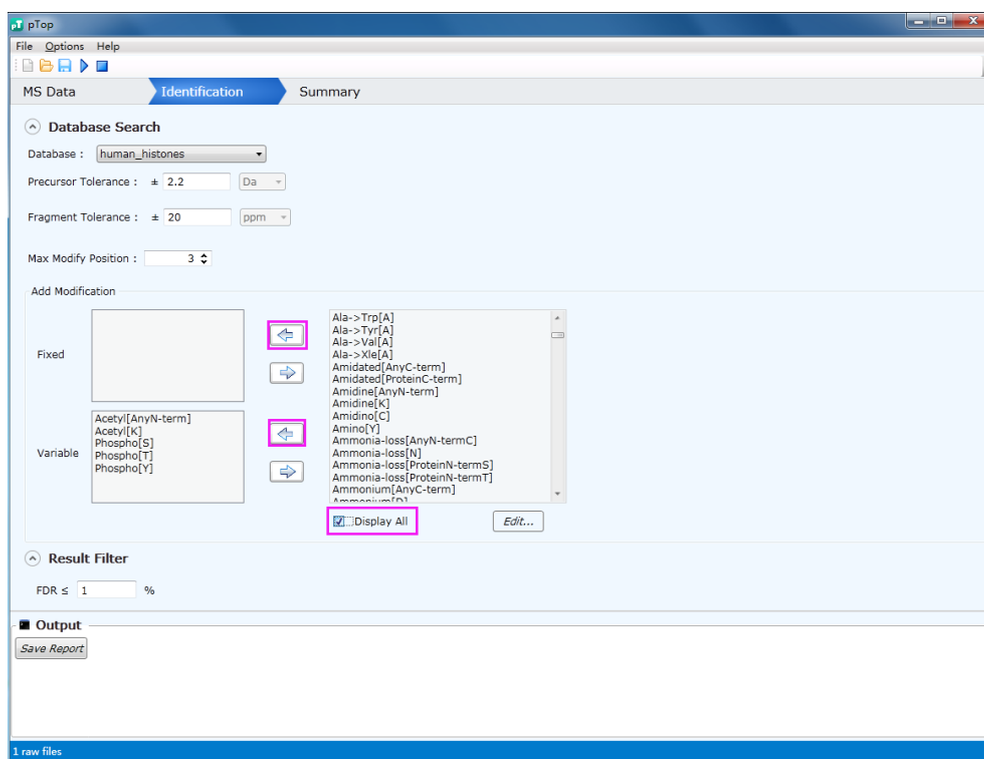


Figure 12. Select modifications

To add a modification, you have to type in the name, choose its composition and then the mono mass will be calculated automated. You also have to choose the positions that it might occur. And then type in the neutral loss of the modification if it have, and do nothing if not. (Figure 13)

If you choose the 'Common' box, the modification you add will appear in the modification list even if the 'Display All' box in not checked.

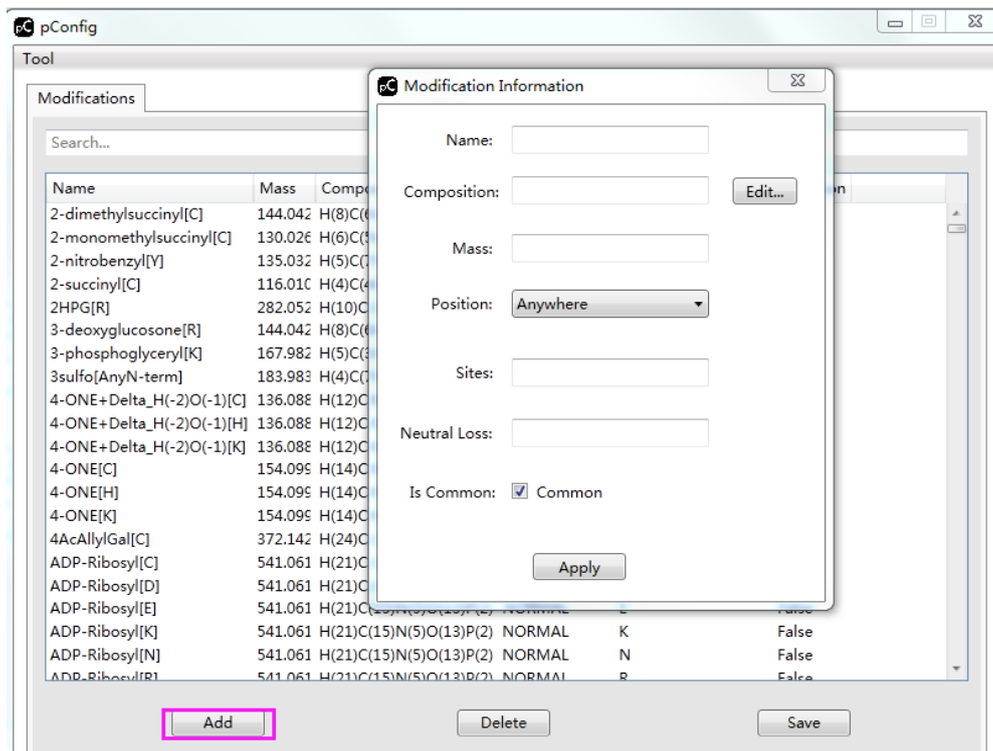


Figure 13. Add a custom modification

3.3 Run pTop

In the summary panel, you can see all the configuration information. And the red rows stand for those you must fill in but you haven't and the green rows mean you did not fill in while it does not matter. After check all the settings in the summary panel, you can click 'Start' to run pTop. (Figure 14)

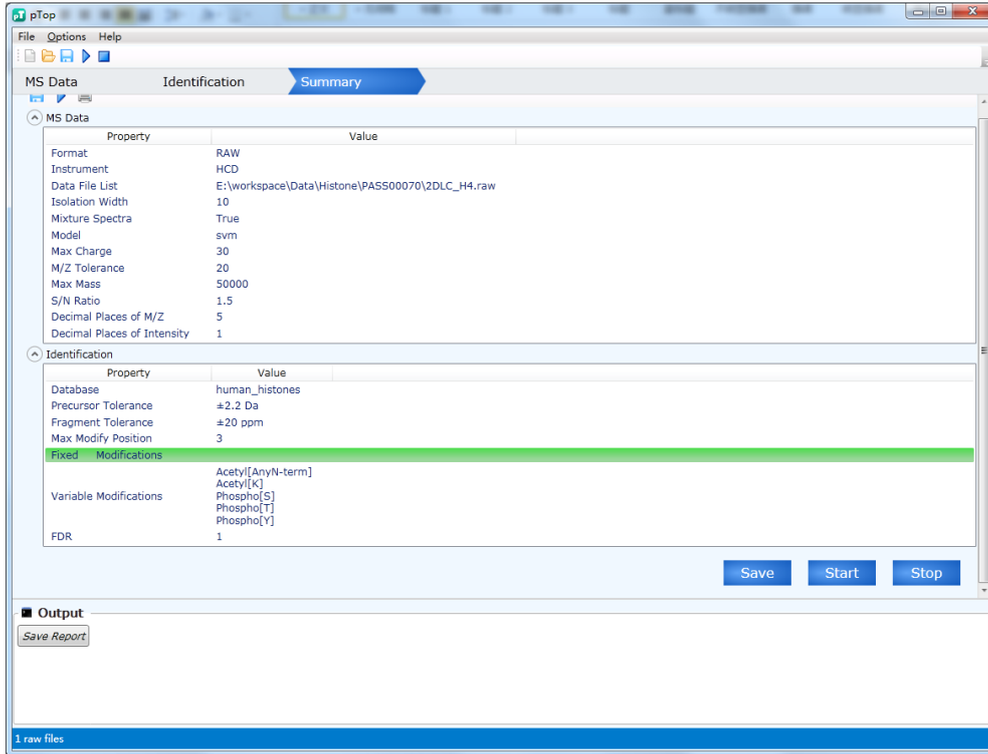


Figure 14. Summary panel

When pTop is running, you can see the progress information in the 'Output' box. (Figure 15)

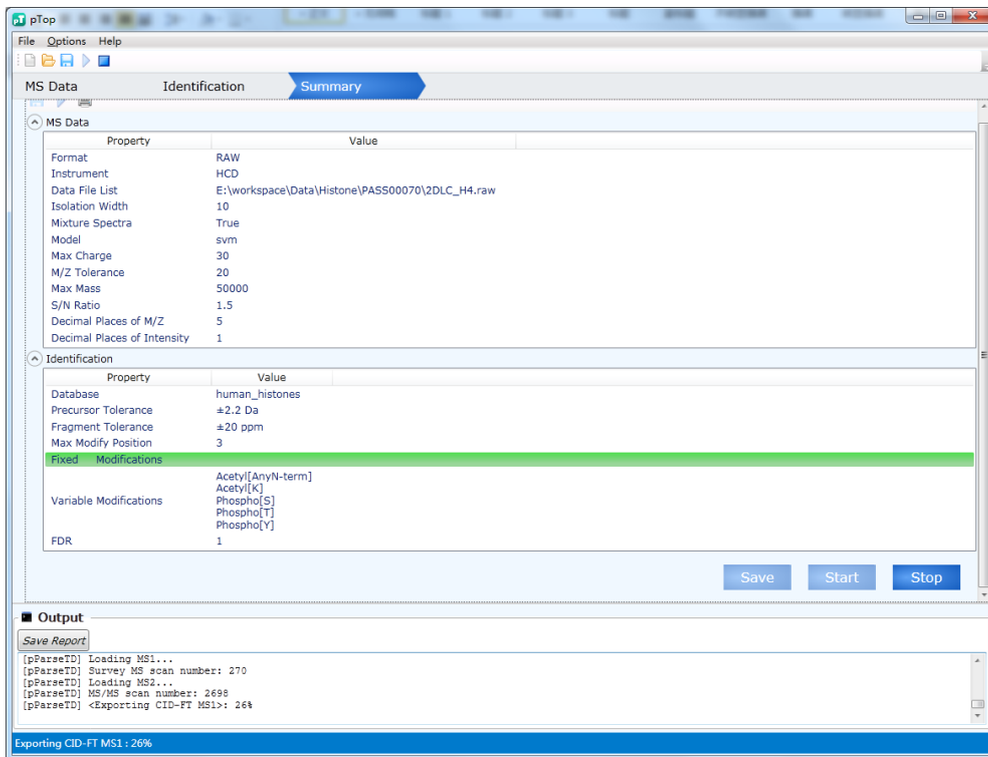


Figure 15. Run pTop

3.4 Results

In the same path of the input data, you can see a folder with the same name of MGF file. (Figure 16)

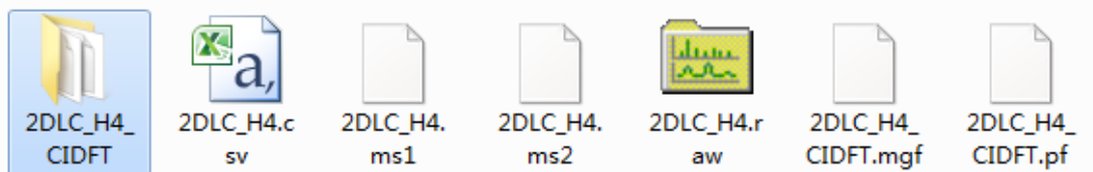


Figure 16. Output files

In the folder, there are 4 files for each search. They are .label, .cfg, filter.csv, query.txt and summary.txt (Figure 17). And the finally identification reports are list in the filter.csv file (Figure 18), in the summary.txt file, you can find the overall results about the total MS/MS, the identification rate for each input file. And pLabel could open the .label file to check the identified proteoform-spectrum-matching (PSM) (Figure 19).

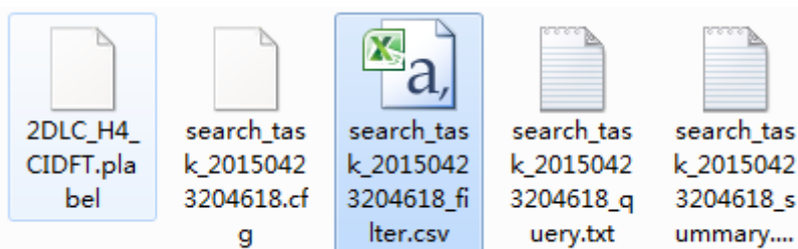


Figure 17. Output reports

A	B	C	D	E	F	G	H	I	J	K	L	M	
ID	Title	Charge State	Precursor Mass	Theory Mass	Mass Diff	Da	Mass Diff ppm	Number of Matched Peaks	Score	Protein AC	Sequence	PTMs	Evalue
1	2DLC_H4_1770.1770.12.0.dta	12	11342.42	11342.4004	0.017	1.5	101	149.78	101	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (5)Acetyl [K]		4.94E-63
2	2DLC_H4_1709.1709.12.0.dta	12	11341.42	11342.4004	-0.981	-86.5	98	147.07	98	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		7.14E-62
3	2DLC_H4_2092.2092.12.0.dta	12	11300.4	11300.3898	0.01	0.9	93	137.06	93	IPF100453-SRR022GELG1	(20)Diethyl [K], (5)Acetyl [K]		5.81E-58
4	2DLC_H4_2105.2105.12.0.dta	12	11300.4	11300.3898	0.01	0.8	91	136.04	91	IPF100453-SRR022GELG1	(23)Methyl [K], (20)Methyl [K], (5)Acetyl [K]		2.59E-57
5	2DLC_H4_1806.1806.12.0.dta	12	11342.41	11342.4004	0.011	1	91	136.01	91	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (5)Acetyl [K]		3.08E-57
6	2DLC_H4_1687.1687.12.0.dta	12	11342.41	11342.4004	0.009	0.8	89	132.09	89	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		6.52E-56
7	2DLC_H4_2100.2100.12.0.dta	12	11300.4	11300.3898	0.009	0.8	88	131.53	88	IPF100453-SRR022GELG1	(23)Methyl [K], (20)Methyl [K], (0)Acetyl [ProteinN-term]		1.57E-55
8	2DLC_H4_1962.1962.12.0.dta	12	11300.4	11300.3898	0.013	1.2	88	130	88	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		9.48E-55
9	2DLC_H4_2010.2010.12.0.dta	12	11300.4	11300.3898	0.014	1.3	85	126.95	85	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		1.46E-53
10	2DLC_H4_2063.2063.12.0.dta	12	11300.4	11300.3898	0.009	0.8	85	125.04	85	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		4.66E-52
11	2DLC_H4_1744.1744.11.0.dta	11	11342.41	11342.4004	0.012	1	84	124.68	84	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		3.39E-52
12	2DLC_H4_2169.2169.12.0.dta	12	11300.4	11300.3898	0.01	0.9	83	123.81	83	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		4.49E-52
13	2DLC_H4_1742.1742.12.0.dta	12	11341.43	11342.4004	-0.973	-86.8	83	123.61	83	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		1.39E-51
14	2DLC_H4_2144.2144.12.0.dta	12	11299.41	11300.3898	-0.981	-86.8	81	123.04	81	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		1.91E-51
15	2DLC_H4_1953.1953.11.0.dta	11	11300.38	11300.3898	-0.007	-0.6	80	120.98	80	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		8.19E-51
16	2DLC_H4_1854.1854.12.1.dta	12	11313.39	11314.3691	-0.986	-87.1	79	115.99	79	IPF100453-SRR022GELG1	(16)Acetyl [K], (0)Acetyl [ProteinN-term]		1.50E-48
17	2DLC_H4_2037.2037.11.0.dta	11	11300.39	11300.3898	0.004	0.4	77	115.44	77	IPF100453-SRR022GELG1	(23)Methyl [K], (19)Methyl [K], (0)Acetyl [ProteinN-term]		3.40E-48
18	2DLC_H4_1841.1841.12.0.dta	12	11341.42	11342.4004	-0.984	-86.8	77	114.87	77	IPF100453-SRR022GELG1	(20)Diethyl [K], (12)Acetyl [K], (0)Acetyl [ProteinN-term]		5.89E-48
19	2DLC_H4_1854.1854.12.0.dta	12	11342.41	11342.4004	0.005	0.5	76	113.39	76	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [K], (0)Acetyl [ProteinN-term]		1.92E-47
20	2DLC_H4_1693.1693.11.0.dta	11	11341.42	11342.4004	-0.983	-86.7	74	113.27	74	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		8.73E-47
21	2DLC_H4_1682.1693.11.1.dta	11	11344.41	11342.4004	2.007	177	74	113.27	74	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		8.73E-47
22	2DLC_H4_2184.2184.12.0.dta	12	11300.4	11300.3898	0.009	0.8	75	113.04	75	IPF100453-SRR022GELG1	(20)Diethyl [K], (5)Acetyl [K]		5.57E-47
23	2DLC_H4_2074.2074.11.0.dta	11	11300.39	11300.3898	0.005	0.5	75	111.24	75	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		2.18E-46
24	2DLC_H4_2221.2221.12.0.dta	12	11300.4	11300.3898	0.006	0.5	74	110.94	74	IPF100453-SRR022GELG1	(20)Diethyl [K], (5)Acetyl [K]		2.69E-46
25	2DLC_H4_2083.2083.13.0.dta	13	11300.39	11300.3898	0.002	0.2	74	110.18	74	IPF100453-SRR022GELG1	(20)Diethyl [K], (5)Acetyl [K]		1.63E-46
26	2DLC_H4_1973.1973.13.0.dta	13	11300.4	11300.3898	0.008	0.8	73	109.54	73	IPF100453-SRR022GELG1	(20)Diethyl [K], (0)Acetyl [ProteinN-term]		4.80E-46
27	2DLC_H4_1764.1764.11.0.dta	11	11341.42	11342.4004	-0.984	-86.7	72	108.33	72	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		3.92E-45
28	2DLC_H4_1764.1764.11.1.dta	11	11344.4	11342.4004	1.997	176.1	72	108.33	72	IPF100453-SRR022GELG1	(20)Diethyl [K], (16)Acetyl [K], (0)Acetyl [ProteinN-term]		3.92E-45

Figure 18. Identification list

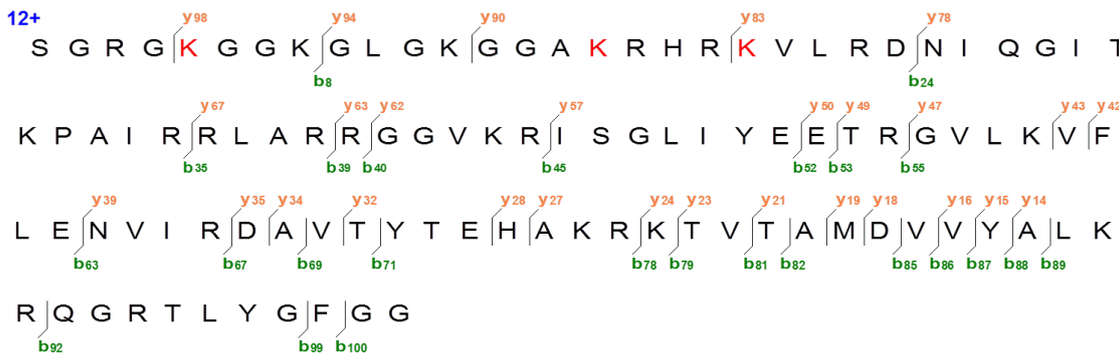
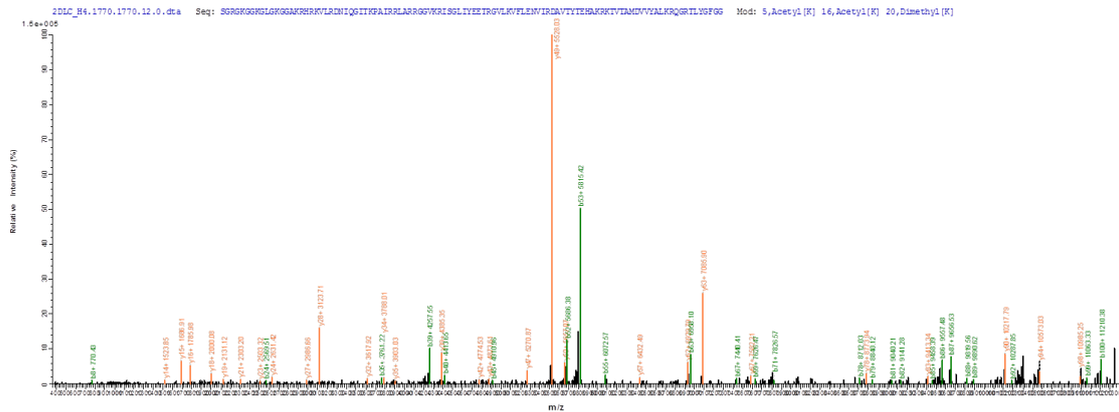


Figure 19. Matched MS/MS in pLabel

4 Contact information

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