# DTarray\_pro-1.8.0 User Guide

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## 1 Introduction

DTarray\_pro extracts Uniprot ID numbers, molecular weights, and spectral counts from DTASelect-filter files. Protein data is combined into one dataset and written to the working directory as a tab delimitated text file (.tsv). This document will describe how to install and use the latest version of DTarray\_pro step by step. Some experience using a unix shell is assumed.

## 2 Installation

### 2.1 Downloading DTarray\_pro source code.

• DTarray\_pro expects to be installed in \$HOME/local. The program needs files stored in \$HOME/local/DTarray\_pro/db for some features to work. First, make \$HOME/local if it doesn't exist.

```
$ mkdir -p ~/local
```

- If you do not wish to build the program in \$HOME/local, you can run the configure script from the desired location with the --program-directory <path> option.
- DTarray\_pro is hosted at GitHub, which is a free hosting service for distributed version control in software development. To clone the latest version of the stable branch from the GitHub repo into your current working directory, run:

```
$ git clone --recurse-submodules --single-branch --
branch stable https://github.com/ajmaurais/
DTarray_pro
```

# 2.2 Compiling DTarray\_pro

- DTarray\_pro is written in vanilla c++ 11 and can be compiled on linux or macOS systems with gcc. Other compilers may work but have not been tested.
- To configure and build DTarray\_pro, run:

```
$ cd ~local/DTarray_pro
$ ./configure
$ make
```

• After you have run make, there should be several new files in the DTarray\_pro directory. If everything worked, two executable files should be located at DTarray\_pro/bin/

```
$ ls bin
DTarray DTsetup
```

- DTarray is the executable for DTarray\_pro
- DTsetup is the executable for a script which will automatically setup DTASelect-filter to be read into DTarray\_pro.

# 3 Usage

## 3.1 Setting up filter files with DTsetup

DTsetup searches all directories one level below where it is run for a file named DTASelect-filter.txt. If a matching file is found, it is copied into a folder named <date>\_<time>\_dtarraySetup with the name <sample\_name>.dtafilter. Once DTsetup is run, DTarray\_pro can be run from the newly created directory to compare spectral counts for proteins found in multiple runs.

To run DTsetup, first navigate to a directory containing subdirectories with DTASelect-filter.txt files and run the command:

### 3.2 DTarray\_pro Input format options

Users have two options for how the input DTASelect-filter files read in by DTarray\_pro are structured.

#### 3.2.1 std mode

In std mode, DTASelect-filter are stored in a common directory and the files are named with the format <sample\_name>.dtafilter (This is the default input mode for DTarray\_pro.)

```
$ 1s
sample_1.dtafilter sample_2.dtafilter
sample_3.dtafilter sample_4.dtafilter
```

The user can either manually setup the folder, or use DTsetup to automatically generate the folder. To run DTarray\_pro in std mode, run the commands:

```
$ cd <path_to_directory_with_filterfiles>
$ DTarray

DTarray_pro v1.7

Adding sample_1...
Adding sample_2...
Adding sample_3...
Adding sample_4...

4 files combined.

Writing protein data... done!
Protein data written in wide format to: DTarray_pro.tsv
```

If DTarray\_pro was successful, two files should have been generated. DTarray\_pro.tsv is a tab delimitated text file (.tsv) which contains a row for each protein and a column for the spectral counts of that protein in each sample. dtarray\_pro\_flist.txt is a list of valid .dtafilter files found in the directory. If a file list already exists in the folder, the existing file list is used. The user can edit the file list to change the order of the columns in the output file and add additional files to the list.

#### 3.2.2 subidr mode

In subdir mode, DTASelect-filter files are stored in a separate directory, the directory name is the sample name, and the filter files are named DTASelect-filter.txt (This is the default input mode for dtarray.pl.)

```
$ find */*.txt
sample_1/DTASelect-filter.txt
sample_2/DTASelect-filter.txt
sample_3/DTASelect-filter.txt
sample_4/DTASelect-filter.txt
```

To run DTarray\_pro in subdir mode, the user has to include the -i subdir option, because subdir is not the default behavior.

```
$ cd <path_to_parent_directory>
$ DTarray -i subdir
```

Upon completion, a files named DTarray\_pro.tsv and dtarray\_pro\_flist.txt are generated formatted the same as in std mode.

## 3.3 DTarray\_pro Output file options

This document will not provide an exhaustive list of options for DTarray\_pro. Instead, this section will provide examples of options will likely find most useful. For the full list of optional arguments see DTarray\_pro/helpFile.pdf or use DTarray -h to see the help file from the terminal.

#### 3.3.1 Get spectral counts for unique peptides by protein

the -u option is used to include a column for the total counts for unique peptides in the DTarray\_pro.tsv output file.

```
$ DTarray -u
```

By default, the columns for spectral counts and unique peptide spectral counts are grouped by sample. See section 3.3.2 to change this behavior.

# 3.3.2 Specify how to group supplementary information columns in output file

The examples in this section all use the -u option (section 3.3.1), but the -s option is also compatible with other DTarray\_pro options including the -lr

option (see section 3.3.5)

By default, the columns for sample specific supplementary information are grouped by sample as shown in table 1.

	S	ample_1	$Sample\_2$		
Protein	SC	$Unique\_SC$	SC	$Unique\_SC$	
ALBU TRFE IGHG1	2149 661 573	2149 661 152	3092 698 382	3092 698 52	

Table 1: Default column arrangement

The -s option can be used to control this behavior. To group the columns by sample, then observation (table 2) add the -s 1 option.

	S	С	${\bf Unique\_SC}$		
Protein	Sample_1	Sample_2	Sample_1	Sample_2	
ALBU TRFE IGHG1	2149 661 573	3092 698 382	2149 $661$ $152$	3092 698 52	

Table 2: Grouping columns by sample

#### 3.3.3 Get spectral counts for peptides

By default no peptide file is generated. To also generate a file containing spectral counts for peptides in each sample, use the -p 1 option.

```
$ DTarray -p 1
```

An additional file will be generated named peptideList.tsv containing peptide data.

#### 3.3.4 Modify how peptides are grouped in output file

By default, peptides are grouped by sequence and parent protein. A separate entry for each charge state of a given peptide will be included in peptide output files. If the -g 2 option is set, peptides will also be grouped by charge; i.e., the spectral counts for each peptide will be the sum of all charge states identified for that peptide.

```
$ DTarray -p 1 -g 2
```

The <code>-modG <group\_method></code> specifies how to group modified peptides in <code>peptideList.tsv</code>. By default peptides with the same sequence, but different modification status will not be grouped. A separate entry will be included for each modification status found for a peptide. To ignore modification status when grouping peptides, use the <code>-modG 1</code> option.

```
$ DTarray -p 1 -modG 1
```

If the -p 1 option is not set, the -g and -modG will be ignored. The -g and -modG options can also be combined as desired.

#### 3.3.5 Get subcelluar location data for proteins

DTarray\_pro can use DTarray\_pro/db/humanLoc.tsv to lookup subcelluar localization information for proteins. humanLoc.tsv contains Uniprot annotations for subcelluar localization by Uniprot ID, updated as of Apr 19, 2019. Currently, sub cell location information is available for human proteins only.

There are two ways in which subcelluar location information can be compiled.

1. The -loc option will add a column for the location of each protein in DTarray\_pro.tsv. To include the subcelluar location column in DTarray\_pro.tsv run the command:

```
$ DTarray -loc
```

2. The -lr 1 option will create a file named loc\_summary.tsv with the sum of spectral counts, sequences, and proteins identified for each subcelluar location. To generate loc\_summary.tsv run the command:

```
$ DTarray -lr 1
```

By default, columns are arranged by sample. The -s 1 option can be used to arrange the columns by observation. See section 3.3.2 for an explanation of the -s option.

#### 3.3.6 Calculate molecular weights for peptides and proteins

DTarray\_pro will calculate protein/peptide molecular weights and molecular formulas when the -mw option is provided. Columns will be included in output files for average mass, monoisotopic mass and molecular formula.

Three files are required for the calculation:

- 1. An atom count table, named atomCountTable.txt which contains the number and types of atoms found in each amino acid (similar to Cimage table).
- 2. An atom mass table, located at DTarray\_pro/db/atomMasses.txt, containing the masses of each atom.
- 3. A .fasta file to lookup protein sequences located at DTarray\_pro/db/humanProteome.fasta (required for proteins only) Currently, the -mw option is supported for human proteins only.

By default the atom count table located at DTarray\_pro/db/atomCountTable.txt is used. The default atom count table includes a static modification on cysteine for iodoacetamide alkylation. To calculate peptide and protein masses with default residue masses, run:

```
$ DTarray -p 1 -mw
```

The user can also supply a custom atomCountTable.txt file with the -act <file\_name> option. A copy of the default atom count table can be generated in the working directory with the -mact option.

```
# make default atomCountTable.txt in working directory
$ DTarray -mact
# run DTarray with custom atomCountTable.txt
$ DTarray -act ./atomCountTable.txt -p 1 -mw
```

The user can also edit the default atom count table as at DTarray\_pro/db/atomCountTable.txt as desired, but editing DTarray\_pro/db/atomMasses.txt is not recommended.