

Install Quanformer

1. Create conda environment (Preinstall conda/miniconda)

```
conda create -n quanformer python=3.8
```

2. Activate environment

```
conda activate quanformer
```

3. Clone quanformer

Note: Make sure *checkpoint0029.pth* in */resources/* is normal (>300MB)。

```
git clone https://github.com/LinShuhaiLAB/QuanFormer.git
```

4. Install pytorch

Windows/Linux with NVIDIA GPU.

```
pip install torch==1.13.1+cu117 torchvision==0.14.1+cu117 torchaudio==0.13.1 --extra-index-url https://download.pytorch.org/whl/cu117
```

Linux without NVIDIA GPU.

```
pip install torch torchvision torchaudio --index-url https://download.pytorch.org/whl/cpu``
```

Windows/mac OS without NVIDIA GPU.

```
pip install torch torchvision torchaudio
```

5. Install requirements

```
cd QuanFormer
pip install -r requirements.txt
```

Use Quanformer

1. Parameter Descriptions

General parameters

- `--type`
 - **Default Value:** `'mzML'`
 - **Description:** Type of raw data files, currently only supports the mzML format.
- `--ppm`
 - **Default Value:** `10`
 - **Description:** PPM value for ROI extraction.
- `--source`
 - **Default Value:** `"resources/example"`
 - **Description:** Path to the raw data directory.
- `--feature`
 - **Default Value:** `"resources/test_feature.csv"`
 - **Description:** The path of the feature file. If it is not empty, the targeted mode will be used. If it is empty, it is the untargeted mode, and the parameters required for the untargeted mode need to be set.
- `--images_path`
 - **Default Value:** `"resources/example/output"`
 - **Description:** Path to the output ROI files.
- `--output`
 - **Default Value:** `"resources/example/output/area.csv"`
 - **Description:** Path to the output files.
- `--threshold`

- **Default Value:** 0.99
- **Description:** Keep only predictions with 0.99 confidence.
- `--model`
 - **Default Value:** "resources/checkpoint0029.pth"
 - **Description:** Path to the peak detection model.
- `--roi_plot`
 - **Default Value:** True
 - **Description:** Whether to plot ROIs. Must be set to True on the first use.
- `--plot`
 - **Default Value:** True
 - **Description:** Whether to plot predictions.
- `--num_classes`
 - **Default Value:** 1
 - **Description:** Number of classes.
- `--smooth_sigma`
 - **Default Value:** 0
 - **Description:** Sigma value for smoothing.
- `--processes_number`
 - **Default Value:** 1
 - **Description:** Number of processes.

Untargeted mode parameters for centWave algorithm.

- `--polarity`
 - **Default Value:** 'positive'
 - **Description:** Polarity.
- `--minWidth`
 - **Default Value:** 5

- **Description:** Min peak width
- `--maxWidth`
 - **Default Value:** 50
 - **Description:** Max peak width.
- `--s2n`
 - **Default Value:** 5
 - **Description:** Signal-to-noise ratio.
- `--noise`
 - **Default Value:** 100
 - **Description:** Noise level.
- `--mzDiff`
 - **Default Value:** 0.005
 - **Description:** m/z difference.
- `--prefilter`
 - **Default Value:** 3
 - **Description:** Pre-filtering parameter.

2. Run in command line mode.

2.1 Targeted Mode

Here is an example command showing how to use these parameters in **targeted mode**, quanformer can run in both centroided and profile data:

2.1.1 Centroided data

```
python main.py --ppm 10 --source resources/example/centroided --feature
resources/example/centroided_feature.csv --images_path
resources/example/centroided_output --output
resources/example/centroided_output/area.csv --model
resources/checkpoint0029.pth
```

```
(quanformer) zzy@zzy-AI:~/testQuanFormer/QuanFormer$ python main.py --ppm 10 --source resources/example/centroided --feature resources/example/centroided_feature.csv --images_path resources/example/centroided_output --output resources/example/centroided_output/area.csv --model resources/checkpoint0029.pth
build took 1.6138 seconds
build_predictor took 4.4070 seconds
quantify took 0.0024 seconds
Successfully exported results to resources/example/centroided_output/area.csv
```

2.1.2 Profile data

example download link(https://drive.google.com/drive/folders/1JopRY0mgMxRGg45iXiBgbTi7uG3M3tS?usp=drive_link)

```
python main.py --ppm 10 --source resources/example/profile --feature resources/example/profile_feature.csv --images_path resources/example/profile_output --output resources/example/profile_output/area.csv --model resources/checkpoint0029.pth
```

```
(quanformer) zzy@zzy-AI:~/testQuanFormer/QuanFormer$ python main.py --ppm 10 --source resources/example/profile --feature resources/example/profile_feature.csv --images_path resources/example/profile_output --output resources/example/profile_output/area.csv --model resources/checkpoint0029.pth
build took 40.7478 seconds
build_predictor took 24.4392 seconds
quantify took 0.1401 seconds
Successfully exported results to resources/example/profile_output/area.csv
```

2.2 Install R before running in untargeted mode

R version 4.4.2, xcms version 4.4.0 In view of the possible problems in downloading R packages, I have packaged my R dependency packages and put them in the following link: (https://drive.google.com/file/d/1oEIANtyXztyRkKUcWpUh3jznCG2trHwv/view?usp=drive_link)

Before using untargeted mode, you should check whether R is installed, open the terminal and input:

```
R --version
```

If R and Rscript are not installed, they can be installed through the following commands. (<https://cran.r-project.org/bin/linux/ubuntu/>)

```
sudo apt-get update
# update indices
sudo apt update -qq
# install two helper packages we need
sudo apt install --no-install-recommends software-properties-common dirmngr
# add the signing key (by Michael Rutter) for these repos
# To verify key, run gpg --show-keys /etc/apt/trusted.gpg.d/cran_ubuntu_key.asc
# Fingerprint: E298A3A825C0D65DFD57CBB651716619E084DAB9
wget -q0- https://cloud.r-project.org/bin/linux/ubuntu/marutter_pubkey.asc |
sudo tee -a /etc/apt/trusted.gpg.d/cran_ubuntu_key.asc
# add the R 4.0 repo from CRAN -- adjust 'focal' to 'groovy' or 'bionic' as
needed
sudo add-apt-repository "deb https://cloud.r-project.org/bin/linux/ubuntu
$(lsb_release -cs)-cran40/"

sudo apt install --no-install-recommends r-base
sudo apt-get install libxml2-dev
```

Then, run the following commands to install packages.

(<https://www.bioconductor.org/packages/release/bioc/html/xcms.html>)

```
sudo R

if (!require("BiocManager", quietly = TRUE))
  install.packages("BiocManager")

BiocManager::install("xcms")
BiocManager::install("MSnbase")
install.packages("dplyr")
```

2.3 Untargeted Mode

Finally, to run the untargeted mode, the feature parameter needs to be set to empty or not set the feature parameter (default is empty), and at the same time, additional parameters required by the centWave algorithm such as polarity, peakWidth, s2n, noise, mzDiff, and prefilter need to be set. A complete command for running the untargeted mode is as follows:

```
python main.py --ppm 10 --source resources/example/centroided --polarity  
positive --minWidth 5 --maxWidth 50 --s2n 5 --noise 100 --mzDiff 0.005 --  
prefilter 3 --images_path resources/example/untargeted_centroided_output --  
output resources/example/untargeted_centroided_output/area.csv --model  
resources/checkpoint0029.pth --processes_number 2
```

```
(quanformer) zzy@zzy-AI:~/testQuanFormer/QuanFormer$ python main.py --ppm 10 --source resources/example/centroided --polarity positive --minWidth 5 --maxWidth 50 --s2n 5 --noise 100 --mzDiff 0.005 --prefilter 3 --images_path resources/example/untargeted_centroided_output --output resources/example/untargeted_centroided_output/area.csv --model resources/checkpoint0029.pth --processes_number 2
```

[1] "已设置北大阿里云镜像"

载入需要的程序包: BiocGenerics

载入程序包: 'BiocGenerics'

The following objects are masked from 'package:stats':

IQR, mad, sd, var, xtabs

The following objects are masked from 'package:base':

anyDuplicated, aperm, append, as.data.frame, basename, cbind, colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find, get, grep, grepl, intersect, is.unsorted, lapply, Map, mapply, match, mget, order, paste, pmax, pmax.int, pmin, pmin.int, Position, rank, rbind, Reduce, rownames, sapply, saveRDS, setdiff, table, tapply, union, unique, unsplit, which.max, which.min

载入需要的程序包: Biobase

Welcome to Bioconductor

Vignettes contain introductory material; view with 'browseVignettes()'. To cite Bioconductor, see 'citation("Biobase")', and for packages 'citation("pkgname")'.

载入需要的程序包: mzR

载入需要的程序包: Rcpp

载入需要的程序包: S4Vectors

载入需要的程序包: stats4

载入程序包: 'S4Vectors'

The following object is masked from 'package:utils':


```
findMatches
```

The following objects are masked from 'package:base':

```
expand.grid, I, unname
```

载入需要的程序包: ProtGenerics

载入程序包: 'ProtGenerics'

The following object is masked from 'package:stats':

```
smooth
```

This is MSnbase version 2.32.0

Visit <https://lgatto.github.io/MSnbase/> to get started.

Consider switching to the 'R for Mass Spectrometry'

packages - see <https://RforMassSpectrometry.org> for details.

载入程序包: 'MSnbase'

The following object is masked from 'package:base':

```
trimws
```

载入需要的程序包: BiocParallel

This is xcms version 4.4.0

载入程序包: 'xcms'

The following object is masked from 'package:stats':

```
sigma
```

```
Detecting mass traces at 10 ppm ... OK
Detecting chromatographic peaks in 13602 regions of interest ... OK: 3691 found.
Detecting mass traces at 10 ppm ... OK
Detecting chromatographic peaks in 11854 regions of interest ... OK: 3306 found.
Detecting mass traces at 10 ppm ... OK
Detecting chromatographic peaks in 11088 regions of interest ... OK: 3394 found.
[=====] 100/100 (100%) in 3s
Sample number 2 used as center sample.
Aligning B1.mzML against B2.mzML ... OK
Aligning B3.mzML against B2.mzML ... OK
Applying retention time adjustment to the identified chromatographic peaks ... OK
[=====] 100/100 (100%) in 3s
Defining peak areas for filling-in .... OK
Start integrating peak areas from original files
Requesting 244 peaks from B1.mzML ... got 191.
Requesting 247 peaks from B2.mzML ... got 217.
Requesting 272 peaks from B3.mzML ... got 238.
build took 228.6459 seconds
build_predictor took 345.1565 seconds
quantify took 0.6404 seconds
Successfully exported results to resources/example/untargeted_centroided_output/
area.csv
```

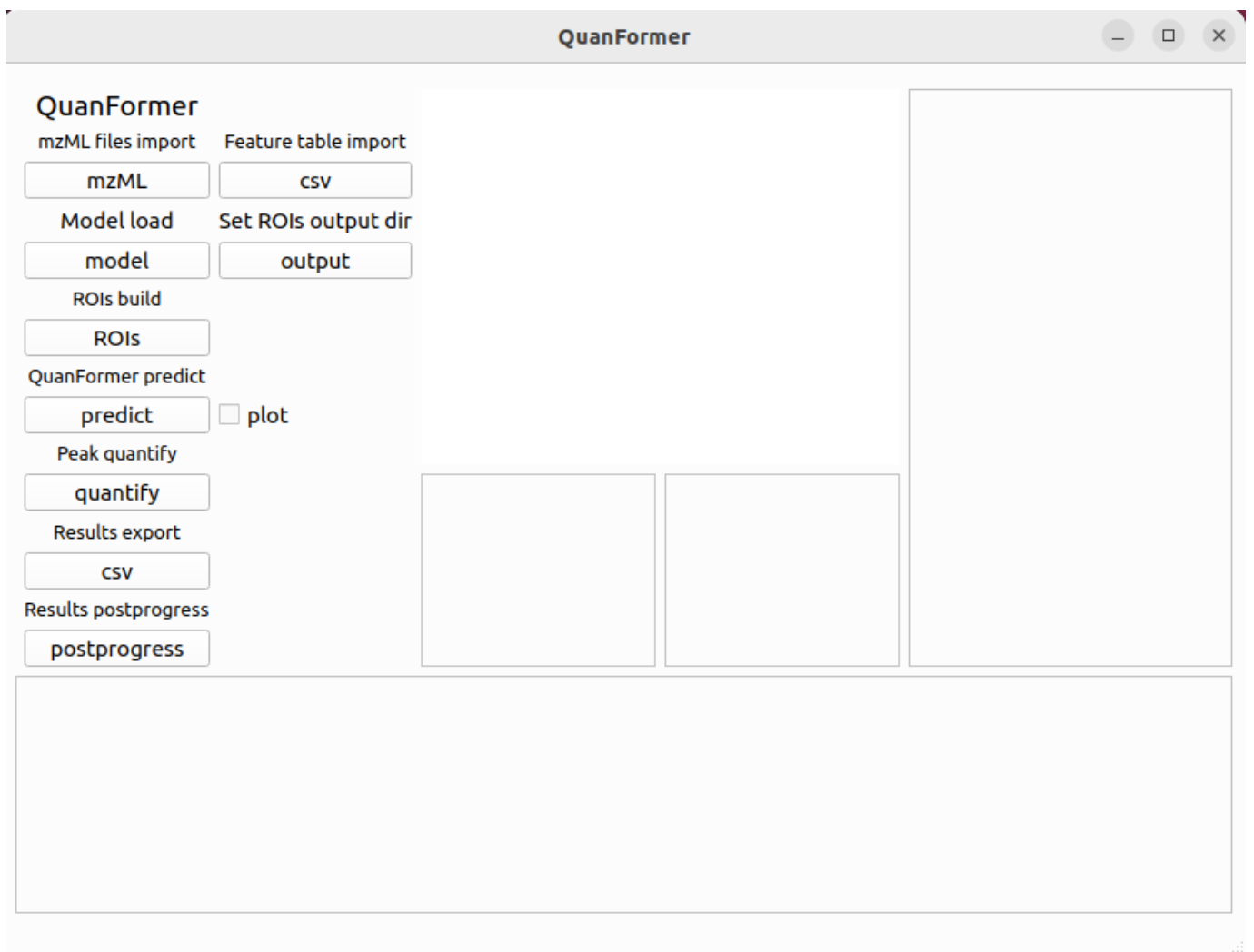
Note: if "FileNotFoundError: [Errno 2] No such file or directory:

'sources/example/xcms_peak_list.csv'" appears in the terminal window, it usually means that the R environment or the dependent packages are not installed correctly. **Please return Step 2.2.**

3. Run in GUI mode

3.1 Targeted Mode

```
python GUI/ms-main.py
```



1. Click mzML button to choose the directory of mzML data, like*
*/resources/example/centroided**.
2. Click csv button to choose the feature table, like
/resources/example/centroided_feature.csv.
3. Click model button to load the model weight, like */resources/checkpoint0029.pth*, make sure the weight is normal (>300MB), or download the weights separately from the Github repository can solve the above problem. Just go to <https://github.com/LinShuhaiLAB/QuanFormer/blob/main/resources/checkpoint0029.pth> and click Download raw file button.
4. Click output button and choose an empty output folder.
5. Click ROIs button and wait for a moment.
6. Check the plot option(or not, reduce time), after clicking the "predict" button, wait for the log column to output "Successfully predict ROIs with plot". Click a different choice in the list column to display the results.

QuanFormer

QuanFormer

mzML files import

mzML

Model load

model

ROIs build

ROIs

QuanFormer predict

predict

Peak quantify

quantify

Results export

csv

Results postprogress

postprogress

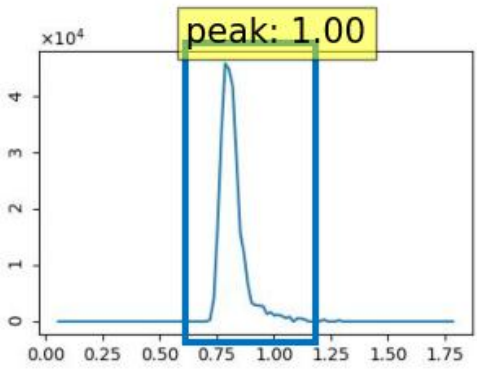
Feature table import

csv

Set ROIs output dir

output

☒ plot



ROI	File
B1	1.0.jpeg
B3	2.0.jpeg
B2	3.0.jpeg
	4.0.jpeg
	5.0.jpeg
	6.0.jpeg

Image

1	/home/zzy/testQuanFormer/QuanFo
2	/home/zzy/testQuanFormer/QuanFo
3	/home/zzy/testQuanFormer/QuanFo
4	/home/zzy/testQuanFormer/QuanFo
5	/home/zzy/testQuanFormer/QuanFo
6	/home/zzy/testQuanFormer/QuanFo
7	/home/zzy/testQuanFormer/QuanFo
8	/home/zzy/testQuanFormer/QuanFo
9	/home/zzy/testQuanFormer/QuanFo
10	/home/zzy/testQuanFormer/QuanFo
11	/home/zzy/testQuanFormer/QuanFo
12	/home/zzy/testQuanFormer/QuanFo

```

21:27:36 Start to build ROIs, wait for a moment please
21:27:36 Successfully built EIC
21:29:18 Start to predict ROIs, wait for a moment please
21:29:18 Successfully predict ROIs with plot
21:29:26 Successfully quantify ROIs
21:29:30 Successfully exported results to: /home/zzy/testQuanFormer/QuanFormer/resources/example/centroided_output/area.csv
21:29:57 Start to post process ROIs
21:29:57 Successfully completed post-processing

```

- Click quantify button.
- Click the csv button to create a new empty csv format file.
- Click the postprogress button to progress the csv.

3.2 Untargeted Mode

In view of the fact that the centWave module requires additional environmental configuration and is relatively time-consuming during operation, and after searching for ROIs, what is finally obtained is a feature table that can be read by the feature button in the GUI mode. Therefore, we suggest to first run the ROIs search module based on the centWave algorithm in the command-line mode:

```
python getFeature.py --source resources/example/centroided --polarity positive
--ppm 10 --minWidth 5 --maxWidth 50 --s2n 5 --noise 100 --mzDiff 0.015 --
prefilter 3
```

```
(quanformer) zzy@zzy-AI:~/testQuanFormer/QuanFormer$ python getFeature.py --source resources/example/centroided --polarity positive --ppm 10 --minWidth 5 --maxWidth 50 --s2n 5 --noise 100 --mzDiff 0.015 --prefilter 3
```

```
[1] "已设置北大阿里云镜像"
```

```
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colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find,  
get, grep, grepl, intersect, is.unsorted, lapply, Map, mapply,  
match, mget, order, paste, pmax, pmax.int, pmin, pmin.int,  
Position, rank, rbind, Reduce, rownames, sapply, saveRDS, setdiff,  
table, tapply, union, unique, unsplit, which.max, which.min
```

```
载入需要的程序包：Biobase
```

```
Welcome to Bioconductor
```

```
Vignettes contain introductory material; view with  
'browseVignettes()'. To cite Bioconductor, see  
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载入程序包: 'MSnbase'

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```

载入需要的程序包: BiocParallel

This is xcms version 4.4.0

载入程序包: 'xcms'

The following object is masked from 'package:stats':

```
The following object is masked from 'package:base':
```

```
trimws
```

```
载入需要的程序包：BiocParallel
```

```
This is xcms version 4.4.0
```

```
载入程序包：'xcms'
```

```
The following object is masked from 'package:stats':
```

```
sigma
```

```
Detecting mass traces at 10 ppm ... OK
```

```
Detecting chromatographic peaks in 13602 regions of interest ... OK: 3680 found.
```

```
Detecting mass traces at 10 ppm ... OK
```

```
Detecting chromatographic peaks in 11854 regions of interest ... OK: 3295 found.
```

```
Detecting mass traces at 10 ppm ... OK
```

```
Detecting chromatographic peaks in 11088 regions of interest ... OK: 3386 found.
```

```
[=====] 100/100 (100%) in 3s
```

```
Sample number 2 used as center sample.
```

```
Aligning B1.mzML against B2.mzML ... OK
```

```
Aligning B3.mzML against B2.mzML ... OK
```

```
Applying retention time adjustment to the identified chromatographic peaks ... OK
```

```
[=====] 100/100 (100%) in 3s
```

```
Defining peak areas for filling-in .... OK
```

```
Start integrating peak areas from original files
```

```
Requesting 246 peaks from B1.mzML ... got 193.
```

```
Requesting 247 peaks from B2.mzML ... got 216.
```

```
Requesting 273 peaks from B3.mzML ... got 238.
```

After the operation is completed, a feature table in.csv format is generated

(./resources/peak_list.csv). Then *python GUI/ms-main.py* and import it. Other operations are the same as targeted analysis (Step 3.1).