# pParse+ User Guide

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# pParse+ User Guide

### Introduction

pParse+ is a software tool for extracting MS\MS spectra from \*.RAW format. It inspects the isolation windows corresponding to each MS\MS spectra and exports the most possible precursors. Besides pParse+ recalibrates the mass of monoisotopic peak and could detect co-eluted precursors. Currently, users can set co-eluted as '0' to export only one recalibrated precursor monoisotopic peak for each MS\MS scan.

pParse+ do not need install. You can use it after decompressed it into a folder.

### **Recommended Minimum System Requirements**

CPU: Pentium III or higher

Main Memory: 1GB

Operating System: Windows XP/Vista/7/8

• Xcalibur 2.1 or higher

### Usage 1

a. Set the parameters. A template of parameter configure file named pParse.para can be found in the folder where pParse+ is decompressed.

b. Save the parameter configure file and open command line prompt in current folder, run pParse+ as follow.

pParse.exe pParse.para

### Usage 2

Now pParse has 21 options and each option has a default value. Users need configure a few of them to run pParse. For example, both

pParse.exe -D D:\mydata\First.raw

and

pParse.exe -D D:\mydata\

are legal.

The command line

#### pParse.exe

will print a detailed user guide about usage of more options.

### **How to Set the Parameters?**

To correctly run pParse+, users need to set a few parameters. Some of the parameters are very important. For example, if **co-eluted** is set as 1, pParse will export all co-eluted precursors.

Usually, users only need to set datapath to run pParse.exe.

```
[Basic Options]
```

```
datapath=E:\MannData\
# both folder or file is legal.
```

### [Advanced Option]

```
co-elute=1
# 0, do not export co-eluted precursors.
# 1, export co-eluted precursors.
input_format=raw
# raw/ms1
isolation_width=2
# default=2.
mars_threshold=-0.68
trainingset=.\TrainingSet.txt
ipv_file=.\IPV.txt.
```

## [0/1 Switches]

```
output_mgf=1
output_mars_y=0
delete_msn=0
rewrite_files=0
export_unchecked_mono=0
cut_similiar_mono=1
output_trainingdata=0
check_activationcenter=1
output_all_mars_y=0
debug_mode=0
```

# pParse+用户手册

### 软件简介

pParse+是一个支持从 RAW 格式导出 MGF 格式串联质谱图的软件。下载好压缩包后,解压即可使用。

pParse+目前具有导出共洗脱母离子和不导出共洗脱母离子两种模式,可以在参数文件中的 co-eluted 项目下修改。

### 最小系统配置要求

- 处理器: Pentium III 以上
- 内存: 1GB
- 操作系统: Windows XP/Vista/7/8
- Xcalibur 2.1 或者更高版本

### 使用方法1

首先配置好参数文件,解压后的文件夹中包含了一个参数文件模板:

pParse.para

配置好参数后,在当前目录下打开命令行,执行:

pParse.exe pParse.para

#### 使用方法2

pParse+现在有21个命令行开关可以控制,每一个开关都有默认参数。用户可以仅仅配置其中一小部分,就可以正常运行pParse+。比如说:

pParse.exe -D D:\mydata\First.raw

或者

pParse.exe -D D:\mydata\First.raw -O E:\mydata\ 分别处理单个 RAW 和批处理文件夹内的所有 RAW。

单独键入命令:

#### pParse.exe

会输出一个比较详细使用说明,此外,如果用户不小心删除了 pParse.para,这一命令还会生成一份参数文件模板。

### 参数文件配置说明

pParse+正常执行依赖于参数文件的正确配置,大部分参数不需要用户修改,常规使用,仅需配置 datapath 参数即可。

# This is standard pParse configure file

# Time: 2014.1.6

#'#'后面可以添加注释,文件中可以任意空行。

### [基本参数]

datapath=E:\MannData\

# 文件夹和文件均可.

### [高级选项]

co-elute=1

# 0, 不导出共洗脱母离子.

#1,导出共洗脱母离子.

input\_format=raw

# raw/ms1

isolation\_width=2

# 默认值 2.

mars\_threshold=-0.68

trainingset=.\TrainingSet.txt

ipv\_file=.\IPV.txt.

# [其他开关]

output\_mgf=1

output mars y=0

delete\_msn=0

rewrite\_files=0

export\_unchecked\_mono=0

cut\_similiar\_mono=1

output\_trainingdata=0

check\_activationcenter=1

output\_all\_mars\_y=0

debug\_mode=0