lamp quick start

October 14, 2024

1 Quick Start

In this vignette we will demonstrate how to use lamp python package. The input data and reference files are located in https://github.com/wanchanglin/lamp/tree/master/examples/data.

1.1 Setup

To use lamp, the first step is to import some python libraries including lamp.

```
[1]: import sqlite3
import pandas as pd
from lamp import anno, stats, utils
```

1.2 Data Loading

lamp supports text files separated by comma (,) or tab (\t). The Microsoft's XLSX is also supported, provided that data set is in the first sheet.

Here we use a small example data set with tsv format. Load it into python and check its format:

```
[2]: d_data = "./data/df_pos_2.tsv"
data = pd.read_table(d_data, header=0, sep="\t")
data
```

```
[2]:
              name
                        namecustom
                                                       mzmin
                                                                                    rt
                                             mz
                                                                    mzmax
     0
           M151T34
                      M150.8867T34
                                     150.886715
                                                  150.886592
                                                              150.886863
                                                                             34.152700
     1
           M151T40
                      M151.0402T40
                                                  151.040092
                                                               151.040350
                                                                             39.838172
                                     151.040235
     2
           M152T40
                      M152.0436T40
                                     152.043607
                                                  152.043451
                                                               152.043737
                                                                             40.303700
     3
                                                               152.883959
           M153T34
                      M152.8838T34
                                     152.883824
                                                  152.883678
                                                                             34.174647
     4
           M153T36
                      M153.0195T36
                                     153.019474
                                                  153.019331
                                                               153.019633
                                                                             35.785847
     395
          M283T339
                     M283.2646T339
                                     283.264583
                                                  283.264341
                                                               283.264809
                                                                           338.763489
                                                  284.194939
     396
                      M284.1953T60
                                     284.195294
                                                              284.195536
                                                                            59.593561
           M284T60
     397
          M284T108
                     M284.2235T108
                                     284.223499
                                                  284.223156
                                                              284.223692
                                                                           108.406389
     398
          M284T339
                      M284.268T339
                                     284.267962
                                                  284.267634
                                                               284.268204
                                                                           338.725056
     399
           M285T34
                       M284.775T34
                                     284.775031
                                                  284.774635
                                                               284.775287
                                                                             34.079641
               rtmin
                            rtmax
                                    npeaks
                                                            X210
                                                                           X209
     0
           33.637595
                        35.465548
                                        97
                                            97
                                                    4.224942e+06
                                                                   3.946599e+06
```

```
1
      37.556072
                   40.532315
                                   95
                                       95
                                               1.419062e+06
                                                              1.251606e+06
2
      38.092678
                   40.909428
                                       81
                                               1.203919e+05
                                                              9.970442e+04
                                   81
3
      33.637595
                   35.465548
                                   98
                                       98
                                               5.592065e+06
                                                              5.761380e+06
4
      34.130244
                   36.287354
                                   98
                                       98
                                               7.284938e+06
                                                              1.083289e+07
                  339.165948
     338.398380
                                               3.509767e+05
                                                              4.117633e+05
395
                                   94
                                       94
396
      58.844217
                   60.107058
                                   59
                                       59
                                                        NaN
                                                                       NaN
397
     107.880510
                  108.971046
                                   72
                                       72
                                               7.477652e+04
                                                              7.482219e+04
398
     338.268300
                  339.370098
                                       84
                                               3.697604e+04
                                                              5.398264e+04
                                   84
399
      33.667172
                   35.198181
                                       97
                                               3.439330e+06
                                                              3.359842e+06
             X208
                            X207
                                           X206
                                                          X205
                                                                         X204
     3.668948e+06
0
                    3.754321e+06
                                   3.853724e+06
                                                  3.787350e+06
                                                                 3.584464e+06
1
     1.214826e+06
                    8.143028e+05
                                   5.331963e+05
                                                  1.930928e+06
                                                                 1.479001e+06
2
     9.384000e+04
                    4.186335e+04
                                                  2.115447e+05
                                                                 1.285713e+05
                                            NaN
3
     5.845419e+06
                    5.576013e+06
                                   5.552878e+06
                                                  6.132789e+06
                                                                 5.891378e+06
4
     1.140072e+07
                    8.220552e+06
                                   9.255154e+06
                                                  7.648211e+06
                                                                 7.723814e+06
. .
395
     3.948000e+05
                    4.338804e+05
                                   5.335221e+05
                                                  6.224684e+05
                                                                 7.009340e+05
396
                                                  2.558004e+04
              NaN
                             NaN
                                            NaN
                                                                 4.020517e+04
397
     3.399667e+04
                    7.233564e+04
                                   1.043879e+05
                                                  2.506785e+04
                                                                 2.753769e+04
     5.340109e+04
398
                    6.557698e+04
                                   7.656575e+04
                                                  1.040606e+05
                                                                 1.063727e+05
399
                    3.789056e+06
                                   3.478506e+06
     3.375577e+06
                                                  3.391588e+06
                                                                 5.067802e+06
                                           X201
             X203
                            X202
0
     3.499711e+06
                    3.623205e+06
                                   4.145770e+06
1
     1.076354e+06
                    9.293218e+05
                                   5.298062e+05
2
     9.389346e+04
                    7.163655e+04
                                   4.916483e+04
3
     5.418082e+06
                    5.036840e+06
                                   5.733794e+06
4
     5.571163e+06
                    5.362560e+06
                                   9.259675e+06
395
     3.005173e+05
                    3.133173e+05
                                   8.204783e+05
396
                                   5.446684e+04
               NaN
                    3.162670e+04
397
               NaN
                             NaN
                                            NaN
398
                    3.059370e+04
              NaN
                                   1.358056e+05
399
     3.497546e+06
                    3.316025e+06
                                   3.906000e+06
```

[400 rows x 110 columns]

This data set includes peak list and intensity data matrix. lamp requires peak list's name, m/z value and retention time. User needs to indicate the locations of peak name, m/z value, retention time and starting points of data matrix from data. Here they are 1, 3, 6 and 11, respectively.

Load input data with xlsx format for 'lamp':

```
[3]: cols = [1, 3, 6, 11]

# d_data = "./data/df_pos_2.tsv"

# df = anno.read_peak(d_data, cols, sep='\t')
```

```
df = anno.read_peak(d_data, cols)
[3]:
              name
                                         rt
                                                      QC9
                                                                     QC5 \
                            mz
     0
           M151T34
                    150.886715
                                  34.152700
                                             3.664879e+06
                                                           3.735147e+06
     1
           M151T40
                    151.040235
                                  39.838172
                                            7.406381e+05
                                                           7.524075e+05
     2
           M152T40
                    152.043607
                                  40.303700
                                             6.105241e+04
                                                           5.335546e+04
     3
           M153T34
                    152.883824
                                  34.174647
                                             5.141479e+06
                                                           5.496344e+06
     4
           M153T36
                    153.019474
                                  35.785847
                                             5.336758e+06
                                                           5.558265e+06
     . .
               •••
                         •••
     395
         M283T339
                    283.264583
                                 338.763489
                                             7.330602e+05
                                                           8.243956e+05
          M284T60
                    284.195294
                                             2.310932e+04
     396
                                  59.593561
                                                                     NaN
     397
          M284T108
                    284.223499
                                 108.406389
                                             3.748444e+04
                                                           2.993283e+04
     398
          M284T339
                    284.267962
                                 338.725056
                                             1.161886e+05
                                                           1.476514e+05
     399
          M285T34
                    284.775031
                                  34.079641
                                             4.063268e+06
                                                           3.807148e+06
                   QC4
                                  QC3
                                               QC26
                                                             QC25
                                                                            QC24
     0
          5.190263e+06
                       2.742966e+06
                                      3.824723e+06
                                                    3.722932e+06 3.804188e+06
     1
                       6.429245e+05
                                      1.167016e+06
                                                     1.175981e+06
                                                                   1.122533e+06
                   NaN
     2
                                  NaN
                                       6.875157e+04
                                                     7.807399e+04
                                                                   8.943068e+04
     3
          8.335846e+06
                        3.860588e+06
                                       5.316874e+06
                                                     5.988232e+06
                                                                   5.844917e+06
     4
          1.118557e+07
                        6.876715e+06
                                       9.967314e+06
                                                     9.073822e+06
                                                                   9.328573e+06
                                       4.294760e+05
                                                                   4.570657e+05
     395
                   NaN
                        1.159506e+06
                                                     4.641813e+05
                                 NaN
                                       1.759336e+04
     396
                   NaN
                                                     2.645392e+04
                                                                    2.727266e+04
     397
                                       3.175596e+04
                                                     3.879604e+04
                                                                   4.299529e+04
                   NaN
                                  NaN
     398
                   NaN
                                  NaN
                                                NaN
                                                     6.753490e+04
                                                                    5.436219e+04
     399
          4.645099e+06 2.232221e+06
                                      4.576754e+06
                                                    4.533339e+06 4.559356e+06
                     X210
                                   X209
                                                  X208
                                                                X207
     0
             4.224942e+06
                          3.946599e+06
                                         3.668948e+06
                                                        3.754321e+06
     1
             1.419062e+06
                           1.251606e+06
                                          1.214826e+06
                                                        8.143028e+05
     2
                           9.970442e+04 9.384000e+04
             1.203919e+05
                                                        4.186335e+04
     3
             5.592065e+06
                           5.761380e+06
                                          5.845419e+06
                                                        5.576013e+06
             7.284938e+06
     4
                           1.083289e+07
                                          1.140072e+07
                                                        8.220552e+06
     . .
     395
             3.509767e+05
                           4.117633e+05
                                          3.948000e+05
                                                        4.338804e+05
     396
                      NaN
                                     NaN
                                                   NaN
                                                                  NaN
                                                        7.233564e+04
     397
             7.477652e+04
                           7.482219e+04
                                          3.399667e+04
     398
             3.697604e+04
                          5.398264e+04
                                          5.340109e+04
                                                        6.557698e+04
     399
             3.439330e+06 3.359842e+06
                                         3.375577e+06
                                                        3.789056e+06
                  X206
                                 X205
                                               X204
                                                             X203
                                                                            X202
     0
          3.853724e+06 3.787350e+06
                                       3.584464e+06
                                                    3.499711e+06 3.623205e+06
     1
          5.331963e+05 1.930928e+06
                                      1.479001e+06
                                                     1.076354e+06 9.293218e+05
     2
                   NaN 2.115447e+05
                                     1.285713e+05 9.389346e+04 7.163655e+04
```

use xlsx file

d_data = "./data/df_pos_2.xlsx"

```
3
     5.552878e+06 6.132789e+06
                                   5.891378e+06
                                                  5.418082e+06
                                                                 5.036840e+06
4
     9.255154e+06
                   7.648211e+06
                                   7.723814e+06
                                                  5.571163e+06
                                                                 5.362560e+06
. .
395
     5.335221e+05
                    6.224684e+05
                                   7.009340e+05
                                                  3.005173e+05
                                                                 3.133173e+05
396
                    2.558004e+04
                                   4.020517e+04
                                                                 3.162670e+04
              NaN
                                                           {\tt NaN}
397
     1.043879e+05
                    2.506785e+04
                                   2.753769e+04
                                                           NaN
                                                                          NaN
398
     7.656575e+04
                    1.040606e+05
                                   1.063727e+05
                                                           {\tt NaN}
                                                                 3.059370e+04
399
     3.478506e+06
                    3.391588e+06
                                   5.067802e+06
                                                  3.497546e+06
                                                                 3.316025e+06
             X201
0
     4.145770e+06
1
     5.298062e+05
2
     4.916483e+04
3
     5.733794e+06
4
     9.259675e+06
395
    8.204783e+05
396
     5.446684e+04
397
              NaN
398
    1.358056e+05
     3.906000e+06
399
```

[400 rows x 103 columns]

Data frame df now includes only name, mz, rt and intensity data matrix.

1.3 Metabolite Annotation

To performance metabolite annotation, users should provide their own reference file. Otherwise, lamp will use its default reference file for annotation.

```
[4]: ref_path = "" # if empty, use default reference file for matching
# load reference library
cal_mass = False
ref = anno.read_ref(ref_path, calc=cal_mass)
ref
```

[4]:	compound_id	molecular_formula	compound_name	\
0	1638	C10C1100	Chlordecone	
1	38485	C10H10Br202	Dibromothymoquinone	
2	32427	C10H10BrNO2	Brofoxine (USAN/INN)	
3	39834	C10H10Cl2N2O	Fenmetozole (USAN)	
4	10156	C10H10Cl203	4-(2,4-Dichlorophenoxy)butyric acid	
•••	•••	•••		
31639	80256	H5010P3	PPPi	
31640	37374	H6N09P3	(Diphosphono)Aminophosphonic Acid	
31641	32626	H9N2O4P	Ammonium phosphate (NF)	
31642	735	HNO3	Nitrate	

31643	40762	HNO3	Peroxynitrite
	exact_mass		
0	485.683441		
1	319.904755		
2	254.989491		
3	244.017018		
4	248.000700		
	•••		
31639	257.909557		
31640	256.925542		
31641	132.029994		
31642	62.995643		
31643	62.995643		

[31644 rows x 4 columns]

The reference file must have two columns: molecular_formula and compound_name (or name). The exact_mass is optional. if absent, lamp will calculates it based on NIST database. If your reference file has exact_mass and want to calculate it using NIST database, set calc as True. The exact_mass is used to match against a range of mz, controlled by ppm, in data frame df.

Another reference file is HMDB database for urine:

```
[5]: ref_path = "./data/hmdb_urine_v4_0_20200910_v1.tsv"
cal_mass = True # there is no exact mass in reference file, so calculate them
ref = anno.read_ref(ref_path, calc=cal_mass)
ref
```

	id	molecular_formula	molecular_name	\
0	HMDB0000001	C7H11N3O2	1-Methylhistidine	
1	HMDB0000002	C3H10N2	1,3-Diaminopropane	
2	HMDB0000005	C4H6O3	2-Ketobutyric acid	
3	HMDB0000008	C4H8O3	2-Hydroxybutyric acid	
4	HMDB0000010	C19H24O3	2-Methoxyestrone	
	•••	•••		
1606	HMDB0012308	C8H8O3	Vanillin	
1607	HMDB0012322	C10H80	2-Naphthol	
1608	HMDB0012325	C5H10O5	Arabinofuranose	
1609	HMDB0012451	C20H28O3	all-trans-5,6-Epoxyretinoic acid	
1610	HMDB0012467	C15H1309S	(-)-Epicatechin sulfate	
			inchi \	
0	InChI=1S/C7H	H11N3O2/c1-10-3-5(9	-4-10)2-6(8)7(11	
1		InChI=1S/C3H10N	2/c4-2-1-3-5/h1-5H2	
2	InChI=1S/C	C4H6O3/c1-2-3(5)4(6)7/h2H2,1H3,(H,6,7)	
3	InChI=1S/C4H	I803/c1-2-3(5)4(6)7	/h3,5H,2H2,1H3,(
4	InChI=1S/C19	0H24O3/c1-19-8-7-12	-13(15(19)5-6-18	
	1 2 3 4 1606 1607 1608 1609 1610	0 HMDB0000001 1 HMDB0000002 2 HMDB0000005 3 HMDB0000008 4 HMDB0000010 1606 HMDB0012308 1607 HMDB0012322 1608 HMDB0012325 1609 HMDB0012451 1610 HMDB0012467 0 InChI=1S/C7H 1 2 InChI=1S/C4H 3 InChI=1S/C4H	1 HMDB0000002 C3H10N2 2 HMDB0000005 C4H603 3 HMDB0000008 C4H803 4 HMDB0000010 C19H2403 1606 HMDB0012308 C8H803 1607 HMDB0012322 C10H80 1608 HMDB0012325 C5H1005 1609 HMDB0012451 C20H2803 1610 HMDB0012467 C15H1309S 0 InChI=1S/C7H11N302/c1-10-3-5(9) 1 InChI=1S/C3H10N 2 InChI=1S/C4H603/c1-2-3(5)4(6) 3 InChI=1S/C4H803/c1-2-3(5)4(6)7	O HMDB0000001 C7H11N302 1-Methylhistidine 1 HMDB0000002 C3H10N2 1,3-Diaminopropane 2 HMDB0000005 C4H603 2-Ketobutyric acid 3 HMDB0000008 C4H803 2-Hydroxybutyric acid 4 HMDB0000010 C19H2403 2-Methoxyestrone 1606 HMDB0012308 C8H803 Vanillin 1607 HMDB0012322 C10H80 2-Naphthol 1608 HMDB0012325 C5H1005 Arabinofuranose 1609 HMDB0012451 C20H2803 all-trans-5,6-Epoxyretinoic acid 1610 HMDB0012467 C15H1309S (-)-Epicatechin sulfate

```
1606
      InChI=1S/C8H8O3/c1-11-8-4-6(5-9)2-3-7(8)10/h2-...
      InChI=1S/C10H80/c11-10-6-5-8-3-1-2-4-9(8)7-10/...
1607
1608
     InChI=1S/C5H1005/c6-1-2-3(7)4(8)5(9)10-2/h2-9H...
1609
     InChI=1S/C20H28O3/c1-15(8-6-9-16(2)14-17(21)22...
1610
     InChI=1S/C15H1409S/c16-9-3-8-5-13(24-25(20,21)...
                        inchi_key
                                    exact_mass
0
      BRMWTNUJHUMWMS-LURJTMIESA-N
                                    169.085127
1
                                     74.084398
      XFNJVJPLKCPIBV-UHFFFAOYSA-N
2
      TYEYBOSBBBHJIV-UHFFFAOYSA-N
                                    102.031694
3
      AFENDNXGAFYKQO-VKHMYHEASA-N
                                    104.047344
4
      WHEUWNKSCXYKBU-QPWUGHHJSA-N
                                    300.172545
1606 MWOOGOJBHIARFG-UHFFFAOYSA-N
                                    152.047344
1607
     JWAZRIHNYRIHIV-UHFFFAOYSA-N
                                    144.057515
1608 HMFHBZSHGGEWLO-HWQSCIPKSA-N
                                    150.052823
1609
     KEEHJLBAOLGBJZ-WEDZBJJJSA-N
                                    316.203845
1610 WTXWEAXATVSZQX-AFYYWNPRSA-M
                                    369.028028
```

[1611 rows x 6 columns]

Next we use HMDB reference file for compounds match. Here function argument ppm is used to control the m/z matching tolerance(range).

```
[6]:
                             mz molecular_formula
                                                                molecular name
                id
                     154.062402
     0
           M154T37
                                           C8H10O3
                                                                Hydroxytyrosol
          M164T119
                     164.046774
                                                            Phenylpyruvic acid
     1
                                            C9H8O3
     2
          M164T119
                     164.046774
                                            C9H8O3
                                                               m-Coumaric acid
     3
          M164T119
                     164.046774
                                                        4-Hydroxycinnamic acid
                                            C9H8O3
     4
          M164T119
                     164.046774
                                            C9H8O3
                                                        2-Hydroxycinnamic acid
     5
                                                            Phenylpyruvic acid
          M164T233
                     164.046832
                                            C9H8O3
     6
          M164T233
                     164.046832
                                            C9H8O3
                                                               m-Coumaric acid
     7
          M164T233
                    164.046832
                                            C9H8O3
                                                        4-Hydroxycinnamic acid
     8
                                                        2-Hydroxycinnamic acid
          M164T233
                     164.046832
                                            C9H8O3
     9
           M164T53
                    164.046825
                                            C9H8O3
                                                            Phenylpyruvic acid
                                                               m-Coumaric acid
     10
           M164T53
                     164.046825
                                            C9H8O3
                                                        4-Hydroxycinnamic acid
     11
           M164T53
                    164.046825
                                            C9H8O3
     12
                                                        2-Hydroxycinnamic acid
           M164T53
                     164.046825
                                            C9H8O3
     13
           M167T35
                     167.021095
                                           C7H5N04
                                                               Quinolinic acid
     14
         M173T36_3
                     173.104423
                                          C8H15N03
                                                               Hexanovlglycine
     15
           M174T35
                                                                  Suberic acid
                     174.088395
                                           C8H14O4
                                                    8-Hydroxy-7-methylguanine
     16
           M181T36
                    181.060407
                                          C6H7N502
     17
           M212T39
                    212.067866
                                          C10H12O5
                                                              Vanillactic acid
```

```
inchi
0
    InChI=1S/C8H1003/c9-4-3-6-1-2-7(10)8(11)5-6/h1...
    InChI=1S/C9H8O3/c10-8(9(11)12)6-7-4-2-1-3-5-7/...
1
2
    InChI=1S/C9H8O3/c10-8-3-1-2-7(6-8)4-5-9(11)12/...
    InChI=1S/C9H8O3/c10-8-4-1-7(2-5-8)3-6-9(11)12/...
3
4
    InChI=1S/C9H8O3/c10-8-4-2-1-3-7(8)5-6-9(11)12/...
5
    InChI=1S/C9H8O3/c10-8(9(11)12)6-7-4-2-1-3-5-7/...
    InChI=1S/C9H8O3/c10-8-3-1-2-7(6-8)4-5-9(11)12/...
6
7
    InChI=1S/C9H8O3/c10-8-4-1-7(2-5-8)3-6-9(11)12/...
8
    InChI=1S/C9H8O3/c10-8-4-2-1-3-7(8)5-6-9(11)12/...
9
    InChI=1S/C9H8O3/c10-8(9(11)12)6-7-4-2-1-3-5-7/...
10
    InChI=1S/C9H8O3/c10-8-3-1-2-7(6-8)4-5-9(11)12/...
    InChI=1S/C9H8O3/c10-8-4-1-7(2-5-8)3-6-9(11)12/...
11
12
    InChI=1S/C9H8O3/c10-8-4-2-1-3-7(8)5-6-9(11)12/...
13
    InChI=1S/C7H5N04/c9-6(10)4-2-1-3-8-5(4)7(11)12...
    InChI=1S/C8H15N03/c1-2-3-4-5-7(10)9-6-8(11)12/...
14
15
    InChI=1S/C8H14O4/c9-7(10)5-3-1-2-4-6-8(11)12/h...
    InChI=1S/C6H7N5O2/c1-11-2-3(9-6(11)13)8-5(7)10...
16
17
    InChI=1S/C10H12O5/c1-15-9-5-6(2-3-7(9)11)4-8(1...
    InChI=1S/C10H16N2O5S/c13-8(14)4-2-1-3-7-9-6(5-...
18
                       inchi key
                                  exact_mass
                                               ppm_error
0
    JUUBCHWRXWPFFH-UHFFFAOYSA-N
                                      154.06
                                                   -3.84
1
    BTNMPGBKDVTSJY-UHFFFAOYSA-N
                                      164.05
                                                   -3.47
    KKSDGJDHHZEWEP-SNAWJCMRSA-N
2
                                      164.05
                                                   -3.47
3
    NGSWKAQJJWESNS-ZZXKWVIFSA-N
                                      164.05
                                                   -3.47
4
    PMOWTIHVNWZYFI-AATRIKPKSA-N
                                      164.05
                                                   -3.47
5
    BTNMPGBKDVTSJY-UHFFFAOYSA-N
                                      164.05
                                                   -3.12
    KKSDGJDHHZEWEP-SNAWJCMRSA-N
6
                                      164.05
                                                   -3.12
7
    NGSWKAQJJWESNS-ZZXKWVIFSA-N
                                                   -3.12
                                      164.05
8
    PMOWTIHVNWZYFI-AATRIKPKSA-N
                                      164.05
                                                   -3.12
    BTNMPGBKDVTSJY-UHFFFAOYSA-N
                                      164.05
                                                   -3.16
   KKSDGJDHHZEWEP-SNAWJCMRSA-N
10
                                                   -3.16
                                      164.05
11
    NGSWKAQJJWESNS-ZZXKWVIFSA-N
                                      164.05
                                                   -3.16
   PMOWTIHVNWZYFI-AATRIKPKSA-N
                                                   -3.16
12
                                      164.05
13
   GJAWHXHKYYXBSV-UHFFFAOYSA-N
                                      167.02
                                                   -4.57
14
   UPCKIPHSXMXJOX-UHFFFAOYSA-N
                                      173.11
                                                   -4.45
15
   TYFQFVWCELRYAO-UHFFFAOYSA-N
                                      174.09
                                                   -4.67
16
   VHPXSVXJBWZORQ-UHFFFAOYSA-N
                                      181.06
                                                    2.39
17
    SVYIZYRTOYHQRE-UHFFFAOYSA-N
                                      212.07
                                                   -2.86
    QPFQYMONYBAUCY-ZKWXMUAHSA-N
                                                   -2.16
                                      276.08
```

match gives the compound matching results. lamp also provides a mass adjust option by adduct library. You can provide your own adducts library otherwise lamp uses its default adducts library.

The adducts library's format looks like:

```
[7]: add_path = './data/adducts_short.tsv'
lib_df = pd.read_csv(add_path, sep="\t")
lib_df
```

```
[7]:
                         exact_mass
                                       charge ion_mode
                 label
                 [M+H]+
                            1.007276
     0
                                             1
                                                     pos
     1
              [M+NH4]+
                           18.033826
                                             1
                                                     pos
     2
                [M+Na]+
                           22.989221
                                             1
                                                     pos
     3
                [M+Mg]+
                           23.984493
                                             1
                                                     pos
     4
                 [M+K]+
                           38.963158
                                             1
                                                     pos
     5
                [M+Fe]+
                           55.934388
                                             1
                                                     pos
     6
                [M+Cu]+
                           62.929049
                                             1
                                                     pos
     7
                [M+2H] +
                            2.015101
                                             1
                                                     pos
     8
                [M+3H] +
                            3.022926
                                             1
                                                     pos
     9
                 [M-H]-
                           -1.007276
                                             1
                                                     neg
                                             1
     10
             [M+35C1]-
                           34.969401
                                                     neg
     11
          [M+Formate] -
                           44.998203
                                             1
                                                     neg
     12
          [M+Acetate]-
                           59.013853
                                             1
                                                     neg
```

The adducts library must have columns of label, exact_mass, charge and ion_mode.

We use this adducts file to adjust mass:

```
[8]: ion_mode = "pos"
# if empty, use default adducts library
add_path = "./data/adducts_short.tsv"
lib_add = anno.read_lib(add_path, ion_mode)
lib_add
```

```
[8]:
            label
                                  charge
                    exact_mass
           [M+H]+
     0
                      1.007276
                                       1
     1
         [M+NH4]+
                     18.033826
                                       1
     2
          [M+Na]+
                     22.989221
                                       1
     3
          [M+Mg]+
                     23.984493
                                       1
     4
           [M+K]+
                     38.963158
                                       1
          [M+Fe]+
     5
                     55.934388
                                       1
     6
          [M+Cu]+
                     62.929049
                                       1
     7
          [M+2H] +
                      2.015101
                                       1
     8
          [M+3H] +
                      3.022926
                                       1
```

Now use function comp_match_mass_add to match compounds:

```
[9]: match_1 = anno.comp_match_mass_add(df, ppm, ref, lib_add)
match_1
```

```
[9]:
                id
                             mz molecular_formula
                                                                     molecular_name
                                                                     Dihydrothymine
     0
           M152T40
                    152.043607
                                          C5H8N2O2
     1
           M154T37
                    154.062402
                                            C8H8O3
                                                        p-Hydroxyphenylacetic acid
     2
                                                        3-Hydroxyphenylacetic acid
           M154T37
                    154.062402
                                            C8H8O3
```

3	M154T37	154.062402	C8H8O3	ortho-Hydroxyphenylacetic acid
4	M154T37	154.062402	C8H8O3	Mandelic acid
5	M154T37	154.062402	C8H8O3	3-Cresotinic acid
6	M154T37	154.062402	С8Н8ОЗ	4-Hydroxy-3-methylbenzoic acid
7	M154T37	154.062402	C8H8O3	Vanillin
8	M157T35	157.036819	C4H10N2O2	2,4-Diaminobutyric acid
9	M157T35	157.036819	C4H10N2O2	L-2,4-diaminobutyric acid
10	M167T35	167.021095	C5H8N2O2	-
				Dihydrothymine
11	M174T35	174.088395	C9H13NO	Phenylpropanolamine
12	M174T35	174.088395	C10H14O	Thymol
13	M174T35	174.088395	C10H140	(S)-Carvone
14	M174T35	174.088395	C8H12O4	2-Octenedioic acid
15	M174T35	174.088395	C8H12O4	cis-4-Octenedioic acid
16	M181T36	181.060407	C8H8N2O3	Nicotinuric acid
17	M184T38	184.097942	C10H13N2	Nicotine imine
18	M185T39_2	185.082034	C5H15NO4P	Phosphorylcholine
19	M186T36	186.045606	C6H14N2O	N-Acetylputrescine
				· · · · · · · · · · · · · · · · · · ·
20	M187T38	187.097642	C5H15NO4P	Phosphorylcholine
21	M193T40	193.050761	C5H14N4	Agmatine
22	M200T36	200.061328	C7H16N2O	N-Acetylcadaverine
23	M201T39_1	201.051849	C10H10O3	4-Methoxycinnamic acid
24	M203T36_1	203.002108	C9H9NO	Indole-3-carbinol
25	M212T39	212.067866	C8H15NO3	
				Hexanoylglycine
26	M212T39	212.067866	C10H10O5	Vanilpyruvic acid
27	M217T37_1	217.018279	C10H11NO	Tryptophol
28	M221T37	221.012328	C9H11NO2	L-Phenylalanine
29	M223T38	223.008162	C4H10NO6P	O-Phosphothreonine
30	M223T40	223.096863	C12H14O4	Monoisobutyl phthalic acid
				· -
31	M226T44	226.128007	C8H18N4O2	Asymmetric dimethylarginine
32	M226T44	226.128007	C8H18N4O2	Symmetric dimethylarginine
33	M227T36	227.066175	C9H10N2O5	3-Nitrotyrosine
34	M229T38	229.069418	C4H10N3O5P	Phosphocreatine
35	M233T38	233.043479	C8H10N4O2	Caffeine
36	M245T44	245.045772	C7H15N3O3	Homocitrulline
37	M245T37_2	245.093315	C13H18O2	Ibuprofen
38	M249T38	249.038309	C8H10N4O3	1,3,7-Trimethyluric acid
39	M261T43	260.972975	C10H7NO4	Xanthurenic acid
40	M269T37_2	269.088048	C10H12N4O5	Inosine
41	M275T168	275.201932	C18H24O2	Estradiol
				17a-Estradiol
42	M275T168	275.201932	C18H24O2	
43	M277T181	277.217564	C18H28O2	19-Norandrosterone
44	M277T181	277.217564	C18H28O2	19-Noretiocholanolone
45	M278T71	278.148195	C11H20N2O6	Saccharopine
46	M279T233	279.233232	C18H30O2	alpha-Linolenic acid
47	M279T233	279.233232	C18H28O2	19-Norandrosterone
48	M279T233	279.233232	C18H28O2	19-Noretiocholanolone
49	M281T287	281.248903	C18H32O2	Linoleic acid

```
50
     M281T287
               281.248903
                                    C18H3002
51
      M282T61
               282.070271
                                  C10H14N2O6
52
      M282T61
               282.070271
                                  C10H14N2O6
53
      M283T37
               283.103695
                                  C11H14N4O5
                                                  inchi \
0
    InChI=1S/C5H8N2O2/c1-3-2-6-5(9)7-4(3)8/h3H,2H2...
1
    InChI=1S/C8H8O3/c9-7-3-1-6(2-4-7)5-8(10)11/h1-...
2
    InChI=1S/C8H803/c9-7-3-1-2-6(4-7)5-8(10)11/h1-...
3
    InChI=1S/C8H8O3/c9-7-4-2-1-3-6(7)5-8(10)11/h1-...
4
    InChI=1S/C8H8O3/c9-7(8(10)11)6-4-2-1-3-5-6/h1-...
5
    InChI=1S/C8H8O3/c1-5-3-2-4-6(7(5)9)8(10)11/h2-...
6
    InChI=1S/C8H8O3/c1-5-4-6(8(10)11)2-3-7(5)9/h2-...
7
    InChI=1S/C8H8O3/c1-11-8-4-6(5-9)2-3-7(8)10/h2-...
    InChI=1S/C4H10N2O2/c5-2-1-3(6)4(7)8/h3H,1-2,5-...
8
9
    InChI=1S/C4H10N2O2/c5-2-1-3(6)4(7)8/h3H,1-2,5-...
10
    InChI=1S/C5H8N2O2/c1-3-2-6-5(9)7-4(3)8/h3H,2H2...
    InChI=1S/C9H13NO/c1-7(10)9(11)8-5-3-2-4-6-8/h2...
11
12
    InChI=1S/C10H14O/c1-7(2)9-5-4-8(3)6-10(9)11/h4...
13
    InChI=1S/C10H140/c1-7(2)9-5-4-8(3)10(11)6-9/h4...
14
    InChI=1S/C8H12O4/c9-7(10)5-3-1-2-4-6-8(11)12/h...
    InChI=1S/C8H12O4/c9-7(10)5-3-1-2-4-6-8(11)12/h...
15
16
   InChI=1S/C8H8N2O3/c11-7(12)5-10-8(13)6-2-1-3-9...
17
    InChI=1S/C10H13N2/c1-12-7-3-5-10(12)9-4-2-6-11...
18
    InChI=1S/C5H14N04P/c1-6(2,3)4-5-10-11(7,8)9/h4...
19
    InChI=1S/C6H14N2O/c1-6(9)8-5-3-2-4-7/h2-5,7H2,...
20
    InChI=1S/C5H14N04P/c1-6(2,3)4-5-10-11(7,8)9/h4...
21
    InChI=1S/C5H14N4/c6-3-1-2-4-9-5(7)8/h1-4,6H2,(...
22
    InChI=1S/C7H16N2O/c1-7(10)9-6-4-2-3-5-8/h2-6,8...
    InChI=1S/C10H1003/c1-13-9-5-2-8(3-6-9)4-7-10(1...
23
24
    InChI=1S/C9H9NO/c11-6-7-5-10-9-4-2-1-3-8(7)9/h...
25
    InChI=1S/C8H15N03/c1-2-3-4-5-7(10)9-6-8(11)12/...
    InChI=1S/C10H1005/c1-15-9-5-6(2-3-7(9)11)4-8(1...
26
27
    InChI=1S/C10H11NO/c12-6-5-8-7-11-10-4-2-1-3-9(...
28
    InChI=1S/C9H11N02/c10-8(9(11)12)6-7-4-2-1-3-5-...
29
    InChI=1S/C4H10N06P/c1-2(3(5)4(6)7)11-12(8,9)10...
    InChI=1S/C12H14O4/c1-8(2)7-16-12(15)10-6-4-3-5...
30
31
    InChI=1S/C8H18N4O2/c1-12(2)8(10)11-5-3-4-6(9)7...
32
    InChI=1S/C8H18N4O2/c1-10-8(11-2)12-5-3-4-6(9)7...
    InChI=1S/C9H10N2O5/c10-6(9(13)14)3-5-1-2-8(12)...
33
    InChI=1S/C4H10N3O5P/c1-7(2-3(8)9)4(5)6-13(10,1...
34
35
    InChI=1S/C8H10N4O2/c1-10-4-9-6-5(10)7(13)12(3)...
    InChI=1S/C7H15N3O3/c8-5(6(11)12)3-1-2-4-10-7(9...
36
37
    InChI=1S/C13H1802/c1-9(2)8-11-4-6-12(7-5-11)10...
    InChI=1S/C8H10N4O3/c1-10-4-5(9-7(10)14)11(2)8(...
38
    InChI=1S/C10H7N04/c12-7-3-1-2-5-8(13)4-6(10(14...
39
    InChI=1S/C10H12N4O5/c15-1-4-6(16)7(17)10(19-4)...
40
```

alpha-Linolenic acid

Ribothymidine

3-Methyluridine

1-Methylinosine

```
41 InChI=1S/C18H24O2/c1-18-9-8-14-13-5-3-12(19)10...
```

- 42 InChI=1S/C18H24O2/c1-18-9-8-14-13-5-3-12(19)10...
- 43 InChI=1S/C18H28O2/c1-18-9-8-14-13-5-3-12(19)10...
- 14 InChI=1S/C18H28O2/c1-18-9-8-14-13-5-3-12(19)10...
- 45 InChI=1S/C11H20N2O6/c12-7(10(16)17)3-1-2-6-13-...
- 46 InChI=1S/C18H3002/c1-2-3-4-5-6-7-8-9-10-11-12-...
- 47 InChI=1S/C18H28O2/c1-18-9-8-14-13-5-3-12(19)10...
- 48 InChI=1S/C18H28O2/c1-18-9-8-14-13-5-3-12(19)10...
- 49 InChI=1S/C18H32O2/c1-2-3-4-5-6-7-8-9-10-11-12-... 50 InChI=1S/C18H30O2/c1-2-3-4-5-6-7-8-9-10-11-12-...
- 51 InChI=1S/C10H14N2O6/c1-4-2-12(10(17)11-8(4)16)...
- 52 InChI=1S/C10H14N2O6/c1-11-6(14)2-3-12(10(11)17...
- 53 InChI=1S/C11H14N4O5/c1-14-3-13-9-6(10(14)19)12...

Inchi_key					
1 XQXPVVBIMDBYFF-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 2 FVMDYYGIDFPZAX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 3 CCVYRRGZDBSHFU-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 4 IWYDHOAUDWTVEP-ZETCQYMHSA-N 154.06 [M+2H]+ -0.28 5 WHSXTWFYRGDBGO-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 6 LTFHNKUKQYVHDX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 7 MWOOGOJBHIARFG-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 9 OGNSCSPNOLGXSM-UKFMYHEASA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 157.09 [M+K]+ -3.23 <t< td=""><td></td><td>inchi_key</td><td>-</td><td></td><td></td></t<>		inchi_key	-		
2 FVMDYYGIDFPZAX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 3 CCVYRRGZDBSHFU-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 4 IWYDHOAUDWTVEP-ZETCQYMHSA-N 154.06 [M+2H]+ -0.28 5 WHSXTWFYRGOBGO-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 6 LTFHNKUKQYVHDX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 7 MWOOGJBHIARFG-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 9 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 174.09 [M+K]+ -3.83 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+M]+ -3.23 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+M]+ -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+B]+ -1.52 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+B]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 181.06 [M+H]+ -1.52 <tr< td=""><td></td><td></td><td></td><td>•</td><td></td></tr<>				•	
3 CCVYRRGZDBSHFU-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 4 IWYDHOAUDWTVEP-ZETCQYMHSA-N 154.06 [M+2H]+ -0.28 5 WHSXTWFYRGOBGO-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 6 LTFHNKUKQYVHDX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 7 MWOOGOJBHIARFG-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 9 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 174.09 [M+Ma]+ -3.10 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg]+ -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg]+ -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 <td>_</td> <td>XQXPVVBIMDBYFF-UHFFFAOYSA-N</td> <td>154.06</td> <td></td> <td>-0.28</td>	_	XQXPVVBIMDBYFF-UHFFFAOYSA-N	154.06		-0.28
4 IWYDHOAUDWTVEP-ZETCQYMHSA-N 154.06 [M+2H]+ -0.28 5 WHSXTWFYRGOBGO-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 6 LTFHNKUKQYVHDX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 7 MWOOGOJBHIARFG-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 8 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 9 OGNSCSPNOLGXSM-VKHMYHEASA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 174.09 [M+Mg]+ -3.23 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Mg]+ -3.23 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg]+ -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Hg]+ -1.52 15 LQYYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 <td></td> <td>FVMDYYGIDFPZAX-UHFFFAOYSA-N</td> <td>154.06</td> <td></td> <td>-0.28</td>		FVMDYYGIDFPZAX-UHFFFAOYSA-N	154.06		-0.28
5 WHSXTWFYRGOBGO-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 6 LTFHNKUKQYVHDX-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 7 MWOOGOJBHIARFG-UHFFFAOYSA-N 154.06 [M+2H]+ -0.28 8 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 9 OGNSCSPNOLGXSM-VKHMYHEASA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K]+ -3.83 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Mg]+ -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg]+ -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 186.05 [M+H]+ 4.61 18 YHBONZFOIEMCP-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20		CCVYRRGZDBSHFU-UHFFFAOYSA-N			
6 LTFHNKUKQYVHDX-UHFFFAOYSA-N 154.06 [M+2H] + -0.28 7 MWOOGOJBHIARFG-UHFFFAOYSA-N 154.06 [M+2H] + -0.28 8 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K] + -3.61 9 OGNSCSPNOLGXSM-VKHMYHEASA-N 157.04 [M+K] + -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K] + -3.83 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Mg] + -3.23 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg] + -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Hg] + -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H] + -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H] + -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na] + 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-N 186.05 [M+Fe] + 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 193.05 [M+Cu] + -0.69		IWYDHOAUDWTVEP-ZETCQYMHSA-N			
7 MWOOGOJBHIARFG-UHFFFAOYSA-N 154.06 [M+2H] + -0.28 8 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K] + -3.61 9 OGNSCSPNOLGXSM-VKHMYHEASA-N 157.04 [M+K] + -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K] + -3.83 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Ma] + -3.10 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg] + -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg] + -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H] + -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H] + -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H] + -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na] + 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-N 185.08 [M+H] + 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe] + 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 193.05 [M+Cu] + -0.69 <td>5</td> <td>WHSXTWFYRGOBGO-UHFFFAOYSA-N</td> <td>154.06</td> <td>[M+2H]+</td> <td>-0.28</td>	5	WHSXTWFYRGOBGO-UHFFFAOYSA-N	154.06	[M+2H]+	-0.28
8 OGNSCSPNOLGXSM-UHFFFAOYSA-N 157.04 [M+K]+ -3.61 9 OGNSCSPNOLGXSM-VKHMYHEASA-N 157.04 [M+K]+ -3.61 10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K]+ -3.83 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Ma]+ -3.10 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg]+ -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg]+ -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na]+ 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-N 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 201.05 [M+Fe]+ -3.42	6	LTFHNKUKQYVHDX-UHFFFAOYSA-N	154.06	[M+2H]+	-0.28
9	7	MWOOGOJBHIARFG-UHFFFAOYSA-N	154.06	[M+2H]+	-0.28
10 NBAKTGXDIBVZOO-VKHMYHEASA-N 167.02 [M+K]+ -3.83 11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Na]+ -3.10 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg]+ -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg]+ -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na]+ 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-N 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 212.07 [M+K]+ -0.28 <td>8</td> <td>OGNSCSPNOLGXSM-UHFFFAOYSA-N</td> <td>157.04</td> <td>[M+K] +</td> <td>-3.61</td>	8	OGNSCSPNOLGXSM-UHFFFAOYSA-N	157.04	[M+K] +	-3.61
11 DLNKOYKMWOXYQA-VXNVDRBHSA-N 174.09 [M+Na] + -3.10 12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg] + -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg] + -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H] + -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H] + -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H] + -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na] + 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-N 185.08 [M+H] + 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe] + 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 187.10 [M+3H] + 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu] + -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe] + 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na] + -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe] + -3.4	9	OGNSCSPNOLGXSM-VKHMYHEASA-N	157.04	[M+K] +	-3.61
12 MGSRCZKZVOBKFT-UHFFFAOYSA-N 174.09 [M+Mg] + -3.23 13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg] + -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H] + -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H] + -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H] + -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na] + 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-N 185.08 [M+H] + 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe] + 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 187.10 [M+3H] + 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu] + -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe] + 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na] + -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe] + -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K] + -0.28	10	NBAKTGXDIBVZOO-VKHMYHEASA-N	167.02	[M+K] +	-3.83
13 ULDHMXUKGWMISQ-VIFPVBQESA-N 174.09 [M+Mg]+ -3.23 14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na]+ 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-O 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 <td>11</td> <td>DLNKOYKMWOXYQA-VXNVDRBHSA-N</td> <td>174.09</td> <td>[M+Na]+</td> <td>-3.10</td>	11	DLNKOYKMWOXYQA-VXNVDRBHSA-N	174.09	[M+Na]+	-3.10
14 BNTPVRGYUHJFHN-HWKANZROSA-N 174.09 [M+2H]+ -1.52 15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na]+ 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-O 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-N 187.10 [M+3H]+ 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -0.28 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+H]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.06	12	MGSRCZKZVOBKFT-UHFFFAOYSA-N	174.09	[M+Mg]+	-3.23
15 LQVYKEXVMZXOAH-UPHRSURJSA-N 174.09 [M+2H]+ -1.52 16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na]+ 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-O 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-O 187.10 [M+3H]+ 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -0.28 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+E]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	13	ULDHMXUKGWMISQ-VIFPVBQESA-N	174.09	[M+Mg]+	-3.23
16 ZBSGKPYXQINNGF-UHFFFAOYSA-N 181.06 [M+H]+ -1.99 17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na]+ 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-O 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-O 187.10 [M+3H]+ 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+E]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 217.02 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	14	BNTPVRGYUHJFHN-HWKANZROSA-N	174.09	[M+2H] +	-1.52
17 GTQXYYYOJZZJHL-UHFFFAOYSA-N 184.10 [M+Na] + 4.61 18 YHHSONZFOIEMCP-UHFFFAOYSA-O 185.08 [M+H] + 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe] + 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-O 187.10 [M+3H] + 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu] + -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe] + 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na] + -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe] + -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K] + -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+E] + -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe] + -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe] + -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.01 [M+H] + 1.69	15	LQVYKEXVMZXOAH-UPHRSURJSA-N	174.09	[M+2H] +	-1.52
18 YHHSONZFOIEMCP-UHFFFAOYSA-O 185.08 [M+H]+ 4.80 19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-O 187.10 [M+3H]+ 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+ZH]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	16	ZBSGKPYXQINNGF-UHFFFAOYSA-N	181.06	[M+H] +	-1.99
19 KLZGKIDSEJWEDW-UHFFFAOYSA-N 186.05 [M+Fe]+ 3.25 20 YHHSONZFOIEMCP-UHFFFAOYSA-O 187.10 [M+3H]+ 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+ZH]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	17	GTQXYYYOJZZJHL-UHFFFAOYSA-N	184.10	[M+Na]+	4.61
20 YHHSONZFOIEMCP-UHFFFAOYSA-O 187.10 [M+3H]+ 4.52 21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	18	YHHSONZFOIEMCP-UHFFFAOYSA-O	185.08	[M+H] +	4.80
21 QYPPJABKJHAVHS-UHFFFAOYSA-N 193.05 [M+Cu]+ -0.69 22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	19	KLZGKIDSEJWEDW-UHFFFAOYSA-N	186.05	[M+Fe]+	3.25
22 RMOIHHAKNOFHOE-UHFFFAOYSA-N 200.06 [M+Fe]+ 3.39 23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na]+ -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	20	YHHSONZFOIEMCP-UHFFFAOYSA-O	187.10	[M+3H]+	4.52
23 AFDXODALSZRGIH-QPJJXVBHSA-N 201.05 [M+Na] + -1.82 24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe] + -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K] + -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H] + -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe] + -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe] + -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg] + -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H] + 1.69	21	QYPPJABKJHAVHS-UHFFFAOYSA-N	193.05	[M+Cu]+	-0.69
24 IVYPNXXAYMYVSP-UHFFFAOYSA-N 203.00 [M+Fe]+ -3.42 25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	22	RMOIHHAKNOFHOE-UHFFFAOYSA-N	200.06	[M+Fe]+	3.39
25 UPCKIPHSXMXJOX-UHFFFAOYSA-N 212.07 [M+K]+ -2.29 26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	23	AFDXODALSZRGIH-QPJJXVBHSA-N	201.05	[M+Na]+	-1.82
26 YGQHQTMRZPHIBB-UHFFFAOYSA-N 212.07 [M+2H]+ -0.28 27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	24	IVYPNXXAYMYVSP-UHFFFAOYSA-N	203.00	[M+Fe]+	-3.42
27 MBBOMCVGYCRMEA-UHFFFAOYSA-N 217.02 [M+Fe]+ -0.79 28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	25	UPCKIPHSXMXJOX-UHFFFAOYSA-N	212.07	[M+K]+	-2.29
28 COLNVLDHVKWLRT-QMMMGPOBSA-N 221.01 [M+Fe]+ -4.70 29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	26	YGQHQTMRZPHIBB-UHFFFAOYSA-N	212.07	[M+2H]+	-0.28
29 USRGIUJOYOXOQJ-GBXIJSLDSA-N 223.01 [M+Mg]+ -4.06 30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	27	MBBOMCVGYCRMEA-UHFFFAOYSA-N	217.02	[M+Fe]+	-0.79
30 RZJSUWQGFCHNFS-UHFFFAOYSA-N 223.10 [M+H]+ 1.69	28	COLNVLDHVKWLRT-QMMMGPOBSA-N	221.01	[M+Fe]+	-4.70
,	29	USRGIUJOYOXOQJ-GBXIJSLDSA-N	223.01	[M+Mg]+	-4.06
31 YDGMGEXADBMOMJ-LURJTMIESA-N 226.13 [M+Mg]+ 2.38	30	RZJSUWQGFCHNFS-UHFFFAOYSA-N	223.10	[M+H]+	1.69
	31	YDGMGEXADBMOMJ-LURJTMIESA-N	226.13	[M+Mg]+	2.38

```
32
   HVPFXCBJHIIJGS-LURJTMIESA-N
                                       226.13
                                                [M+Mg]+
                                                               2.38
                                                 [M+H]+
                                                              -0.32
33
   FBTSQILOGYXGMD-LURJTMIESA-N
                                       227.07
34
    DRBBFCLWYRJSJZ-UHFFFAOYSA-N
                                       229.07
                                               [M+NH4]+
                                                              -0.94
35
    RYYVLZVUVIJVGH-UHFFFAOYSA-N
                                       233.04
                                                 [M+K]+
                                                              -0.23
    XIGSAGMEBXLVJJ-YFKPBYRVSA-N
                                                               0.17
36
                                       245.05
                                                [M+Fe]+
37
    HEFNNWSXXWATRW-UHFFFAOYSA-N
                                       245.09
                                                 [M+K]+
                                                              -2.13
   BYXCFUMGEBZDDI-UHFFFAOYSA-N
                                                              -0.56
38
                                       249.04
                                                 [M+K]+
39
    FBZONXHGGPHHIY-UHFFFAOYSA-N
                                       260.97
                                                [M+Fe]+
                                                               4.13
40
   UGQMRVRMYYASKQ-KQYNXXCUSA-N
                                       269.09
                                                 [M+H]+
                                                               0.01
41
    VOXZDWNPVJITMN-ZBRFXRBCSA-N
                                                               5.00
                                       275.20
                                                [M+3H] +
                                                               5.00
42
    VOXZDWNPVJITMN-SFFUCWETSA-N
                                       275.20
                                                [M+3H] +
   UOUIARGWRPHDBX-CQZDKXCPSA-N
                                       277.22
                                                 [M+H]+
                                                               4.90
   UOUIARGWRPHDBX-DHMVHTBWSA-N
                                       277.22
                                                 [M+H]+
                                                               4.90
45
    ZDGJAHTZVHVLOT-YUMQZZPRSA-N
                                       278.15
                                                [M+2H] +
                                                               3.44
    DTOSIQBPPRVQHS-PDBXOOCHSA-N
                                                               4.93
46
                                       279.23
                                                 [M+H]+
47
    UOUIARGWRPHDBX-CQZDKXCPSA-N
                                       279.23
                                                [M+3H]+
                                                               4.93
48
   UOUIARGWRPHDBX-DHMVHTBWSA-N
                                       279.23
                                                               4.93
                                                [M+3H] +
49
    OYHQOLUKZRVURQ-HZJYTTRNSA-N
                                       281.25
                                                               4.97
                                                 [M+H]+
50
    DTOSIQBPPRVQHS-PDBXOOCHSA-N
                                       281.25
                                                [M+3H] +
                                                               4.97
    DWRXFEITVBNRMK-JXOAFFINSA-N
                                       282.07
                                                [M+Mg]+
                                                               2.10
51
                                                [M+Mg]+
52
   UTQUILVPBZEHTK-UHFFFAOYSA-N
                                       282.07
                                                               2.10
   WJNGQIYEQLPJMN-IOSLPCCCSA-N
                                                 [M+H]+
                                                              -0.01
53
                                       283.10
```

1.4 Correlation Analysis

Next step is correlation analysis, based on intensity data matrix along all peaks. All results are filtered by the correlation coefficient, p-values and retention time difference. That is: keep correlation results in an retention time differences/windows(such as 1 seconds) with correlation coefficient larger than a threshold(such as 0.5) and their correlation p-values less than a threshold (such as 0.05).

lamp supports two correlation methods, pearson and spearman. Also parameter positive allows user to select only positive correlation results.

Two functions, _tic and _toc, record the correlation computation time in seconds.

```
[10]: thres_rt = 1.0
    thres_corr = 0.5
    thres_pval = 0.05
    method = "spearman" # "pearson"
    positive = True
```

Elapsed time: 1.373020887374878 seconds.

```
[11]:
                                                 p_value rt_diff
                          name_b r_value
               name_a
                                           1.267076e-23
                                                             0.02
      0
              M151T34
                         M153T34
                                      0.80
                                           1.752854e-16
      1
                                      0.71
                                                             0.20
              M151T34
                         M155T34
      2
                                      0.78 1.869949e-21
                                                             0.14
              M151T34
                         M161T34
      3
                                      0.69 3.239594e-15
              M151T34
                         M163T34
                                                             0.20
      4
                                      0.51 5.776482e-08
                                                             0.73
              M151T34
                         M167T35
      1783
            M283T34_1
                       M283T34_2
                                      0.62
                                           4.214876e-12
                                                             0.29
      1784
            M283T34_1
                         M285T34
                                      0.82 5.937139e-26
                                                             0.08
      1785
            M283T34_2
                         M285T34
                                      0.66 7.898957e-14
                                                             0.37
      1786
                                                             0.15
              M283T60
                                      0.86 1.033010e-29
                         M284T60
      1787
             M283T339
                                      0.91 4.031333e-39
                                                             0.04
                        M284T339
```

[1788 rows x 5 columns]

corr gives results of correlation coefficient(r_value), correlation p-values(p_value) and retention time difference(rt_diff).

Based on the correlation analysis, we can extract the groups and their sizes by:

```
[12]: # get correlation group and size
    corr_df = stats.corr_grp_size(corr)
    corr_df
```

[12]:		name	cor_grp_size	cor_grp
	0	M215T35	52	M216T35::M217T35::M218T35::M219T34::M219T35::M
	1	M219T35	52	M221T34::M223T34::M225T35::M226T35::M229T34::M
	2	M216T35	52	M217T35::M218T35::M219T34::M219T35::M221T34::M
	3	M217T35	52	M218T35::M219T34::M219T35::M221T34::M223T34::M
	4	M218T35	51	M219T34::M219T35::M221T34::M223T34::M225T35::M
		•••	•••	
	335	M265T415	1	M281T415
	336	M266T66	1	M265T66
	337	M258T51	1	M257T51
	338	M271T197	1	M272T197
	339	M243T287	1	M163T287

[340 rows x 3 columns]

1.5 Summarize Results

The final step gets the summary table in different format and save for the further analysis.

```
[13]: # get summary of metabolite annotation
sr, mr = anno.comp_summ(df, match)
```

This function combines peak table with compound matching results and returns two results in different formats. sr is single row results for each peak id in peak table df:

[14]: sr [14]: name ${\tt exact_mass}$ ppm_error mzrt 0 M151T34 150.886715 34.152700 ${\tt NaN}$ ${\tt NaN}$ 1 M151T40 151.040235 39.838172 NaN NaN2 M152T40 152.043607 40.303700 ${\tt NaN}$ ${\tt NaN}$ 3 152.883824 34.174647 ${\tt NaN}$ M153T34 ${\tt NaN}$ 4 M153T36 153.019474 35.785847 NaN NaN395 M283T61 283.068474 60.739869 ${\tt NaN}$ ${\tt NaN}$ M284T108 284.223499 108.406389 396 ${\tt NaN}$ ${\tt NaN}$ 397 M284T339 284.267962 338.725056 NaN NaN 398 M284T60 284.195294 59.593561 ${\tt NaN}$ ${\tt NaN}$ 399 M285T34 284.775031 34.079641 NaN NaN molecular_formula molecular_name inchi inchi_key 0 NaN NaN NaN NaN 1 NaN NaN NaN NaN2 ${\tt NaN}$ NaNNaN ${\tt NaN}$ 3 NaN NaN NaN NaN 4 NaNNaN ${\tt NaN}$ $\tt NaN$. . ••• ••• 395 NaN NaN ${\tt NaN}$ $\tt NaN$ 396 NaN NaN NaN NaN 397 NaN NaN NaN NaN 398 ${\tt NaN}$ ${\tt NaN}$ ${\tt NaN}$ ${\tt NaN}$ 399 NaN NaN NaN NaN

[400 rows x 9 columns]

mr is multiple rows format if the match more than once from the reference file:

5]: mr							
5]:	name	mz	rt	molecular_formula	molecular_name	inchi	١
0	M151T34	150.886715	34.152700	NaN	NaN	NaN	
1	M151T40	151.040235	39.838172	NaN	NaN	NaN	
2	M152T40	152.043607	40.303700	NaN	NaN	NaN	
3	M153T34	152.883824	34.174647	NaN	NaN	NaN	
4	M153T36	153.019474	35.785847	NaN	NaN	NaN	
	•••	•••	•••	•••	•••		
404	M283T61	283.068474	60.739869	NaN	NaN	NaN	
405	M284T108	284.223499	108.406389	NaN	NaN	NaN	
406	M284T339	284.267962	338.725056	NaN	NaN	NaN	
407	M284T60	284.195294	59.593561	NaN	NaN	NaN	
408	M285T34	284.775031	34.079641	NaN	NaN	NaN	
	inchi_key	exact_mass	ppm_error				

0	NaN	NaN	NaN
1	NaN	NaN	NaN
2	NaN	NaN	NaN
3	NaN	NaN	NaN
4	NaN	NaN	${\tt NaN}$
• •		•••	
404	 NaN	NaN	NaN
			NaN NaN
404	NaN	NaN	
404 405	NaN NaN	NaN NaN	NaN

[409 rows x 9 columns]

Now we merges single format results with correlation results:

```
[16]: # merge summery table with correlation analysis
res = anno.comp_summ_corr(sr, corr_df)
res
```

	res						
[16]:		name	mz	rt	exact_mass	ppm_error	\
[20]	0	M167T35	167.021095		167.02	-4.57	`
	1		276.077397	36.385373	276.08	-2.16	
	2		154.062402	37.183625	154.06	-3.84	
	3	M181T36	181.060407	35.734801	181.06	2.39	
	4	M174T35	174.088395	35.001130	174.09	-4.67	
		•••	•••	•••			
	395	M279T50	279.159930	50.055451	NaN	NaN	
	396	M279T79	279.163910	78.758079	NaN	NaN	
	397	M282T85	282.207859	84.719202	NaN	NaN	
	398	M283T47	283.110871	46.822069	NaN	NaN	
	399	M284T108	284.223499	108.406389	NaN	NaN	
		molecular_	formula		<u>-</u>	\	
	0		C7H5NO4		linic acid		
	1	C10H	16N2O5S	Biot	in sulfone		
	2		C8H10O3	· · · · · · · · · · · · · · · · · · ·	roxytyrosol		
	3			Hydroxy-7-met			
	4		C8H14O4	Su	beric acid		
	• •		•••		•••		
	395		NaN		NaN		
	396		NaN		NaN		
	397		NaN		NaN		
	398		NaN		NaN		
	399		NaN		NaN		

inchi \

0 InChI=1S/C7H5NO4/c9-6(10)4-2-1-3-8-5(4)7(11)12...

```
1
     InChI=1S/C10H16N2O5S/c13-8(14)4-2-1-3-7-9-6(5-...
2
     InChI=1S/C8H10O3/c9-4-3-6-1-2-7(10)8(11)5-6/h1...
     InChI=1S/C6H7N5O2/c1-11-2-3(9-6(11)13)8-5(7)10...
3
4
     InChI=1S/C8H14O4/c9-7(10)5-3-1-2-4-6-8(11)12/h...
395
                                                       NaN
396
                                                       NaN
397
                                                       NaN
398
                                                       NaN
399
                                                       NaN
                         inchi_key
                                    cor_grp_size
0
     GJAWHXHKYYXBSV-UHFFFAOYSA-N
                                             25.0
1
     QPFQYMONYBAUCY-ZKWXMUAHSA-N
                                             13.0
2
     JUUBCHWRXWPFFH-UHFFFAOYSA-N
                                             12.0
3
     VHPXSVXJBWZORQ-UHFFFAOYSA-N
                                              9.0
4
     TYFQFVWCELRYAO-UHFFFAOYSA-N
                                              9.0
. .
395
                                              NaN
                               NaN
396
                               NaN
                                              NaN
397
                               NaN
                                              NaN
398
                                              NaN
                               NaN
399
                               NaN
                                              NaN
                                                   cor_grp
0
     M171T34::M197T36::M209T34::M211T34::M213T34::M...
1
     M277T36_2::M278T36::M173T36_2::M186T36::M187T3...
2
     M155T38::M158T37_2::M164T36::M171T37_2::M173T3...
3
     M224T36::M225T35::M226T35::M227T36::M269T37_2:...
     M211T34::M213T34::M219T34::M221T34::M229T35::M...
4
. .
395
                                                       NaN
396
                                                       NaN
397
                                                       NaN
398
                                                       NaN
399
                                                       NaN
```

[400 rows x 11 columns]

The result data frame res is re-arranged as four parts from top to bottom:

- 1st part: identified metabolites, satisfied with correlation analysis
- 2nd part: identified metabolites, not satisfied with correlation
- 3rd part: no identified metabolites, satisfied with correlation
- 4th part: no identified metabolites, not satisfied with correlation

The users should focus on the first part and perform their further analysis.

You can save all results in different forms, such as text format TSV or CSV. You can also save all

results into a sqlite3 database and use DB Browser for SQLite to view:

```
[17]: f_save = False  # here we do NOT save results
db_out = "test.db"
sr_out = "test_s.tsv"
```

```
[18]: if f save:
          # save all results into a sqlite3 database
          conn = sqlite3.connect(db out)
          df[["name", "mz", "rt"]].to_sql("peaklist",
                                          conn,
                                          if_exists="replace",
                                          index=False)
          corr_df.to_sql("corr_grp", conn, if_exists="replace", index=False)
          corr.to_sql("corr_pval_rt", conn, if_exists="replace", index=False)
          match.to_sql("match", conn, if_exists="replace", index=False)
          mr.to_sql("anno_mr", conn, if_exists="replace", index=False)
          res.to_sql("anno_sr", conn, if_exists="replace", index=False)
          conn.commit()
          conn.close()
          # save final results
          res.to_csv(sr_out, sep="\t", index=False)
```

1.6 End User Usages

For end users, lamp provides two computation options: command line interface(CLI) and graphical user interface (GUI).

To use GUI, you need to open a terminal and type in:

\$ lamp gui

To use CLI, open a terminal and type in command with required arguments, something like:

```
$ lamp cli \
    --sep "tab" \
    --input-data "./data/df_pos_3.tsv" \
    --col-idx "1, 2, 3, 4" \
    --add-path "" \
    --ref-path "" \
    --ion-mode "pos" \
    --cal-mass \
    --thres-rt "1.0" \
    --thres-corr "0.5" \
    --thres-pval "0.05" \
    --method "pearson" \
    --positive \
    --ppm "5.0" \
```

```
--save-db \
--save-mr \
--db-out "./res/test.db" \
--sr-out "./res/test_s.tsv" \
--mr-out "./res/test m.tsv"
```

For the best practice, you can create a bash script <code>.sh</code> (Linux and MacOS) or Windows script <code>.bat</code> to contain these CLI arguments. Change parameters in these files each time when processing new data set.

For example, there are lamp_cli.sh and lamp_cli.bat in https://github.com/wanchanglin/lamp/tree/master/examples. You can run them and check the results in directory examples/res:

• For Linux and MacOS terminal:

```
$ chmod +x lamp_cli.sh
$ ./lamp_cli.sh
```

• For Windows terminal:

```
$ lamp_cli.bat
```