01_TNBC_Diagnostics

2022-11-18

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Overview

The document will perform the following diagnostics:

- 1. Check the unique number and values of the metadata, and constuct a tibble of metadata for each patient
- 2. Checking the basic properties such as dimensions and column names of the dataset
- 3. Check if sample_id and patient_id have the same length, and check if sample_id and ImageNb have the same values
- 4. Check if one value in one column is unique to one value in the other two columns
- 5. Check if cell_type and mm have the same length, and if true then check if one cell_type is unique to one mm value
- 6. Check the names of the mm cell type and construct a tibble for the count of different cell types present in each sample
- 7. Find number of unique cluster number and investigate number of different clusters in each sample with graphs
- 8. Investigate the range, median and standard deviation of the all spatial coordinates, for each cell type and for each cell type in each sample

Libraries

```
library(readr)
library(dplyr)
library(ggplot2)
library(formattable)
```

Read CSV

Read in the MATLAB revised TNBC output CSV files.

```
## Rows: 179194 Columns: 64
## -- Column specification ------
## Delimiter: ","
## chr (3): SITE_02, RECURRENCE_LABEL, mm
## dbl (61): sample_id, patient_id, AGE_AT_DX, STAGE, LATERAL, GRADE, Survival_...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
```

Dataset Diagnostics

The aim of this rmarkdown file is to perform disgnostics on the TNBC dataset.

Basic Properties

There are 179194 rows and 64 columns in the TNBC dataset, 44 of the columns are proteins markers and 8 coulmns are the patients metadata.

```
# dimensions
dim(TNBC)

## [1] 179194 64
```

```
# names of the columns
names(TNBC)
```

```
"patient_id"
                                                         "AGE_AT_DX"
##
   [1] "sample_id"
   [4] "STAGE"
                                 "SITE_02"
##
                                                         "LATERAL"
   [7] "GRADE"
                                 "RECURRENCE LABEL"
##
                                                         "Survival_days_capped"
## [10] "cluster_id"
                                                         "cell_type"
## [13] "ImageNb"
                                 "cellLabelInImage"
                                                         "cellSize"
## [16] "cellRadius"
                                 "centroidX"
                                                         "centroidY"
## [19] "majoraxis"
                                                         "Au"
                                 "eccentricity"
## [22] "Background"
                                 "betaCatenin"
                                                         "Ca"
## [25] "CD11b"
                                 "CD11c"
                                                         "CD138"
## [28] "CD16"
                                 "CD20"
                                                         "CD209"
## [31] "CD3"
                                 "CD31"
                                                         "CD4"
## [34] "CD45"
                                 "CD45R0"
                                                         "CD56"
## [37] "CD63"
                                 "CD68"
                                                         "CD8"
## [40] "dsDNA"
                                                         "Fe"
                                 "EGFR"
## [43] "FoxP3"
                                 "H3K27me3"
                                                         "H3K9ac"
## [46] "HLA_Class_1"
                                 "HLADR"
                                                         "IDO"
                                                         "Ki67"
## [49] "Keratin17"
                                 "Keratin6"
## [52] "Lag3"
                                 "MPO"
                                                         "Na"
## [55] "P"
                                 "p53"
                                                         "panKeratin"
                                                         "pS6"
## [58] "PD1"
                                 "PDL1"
                                                         "Ta"
## [61] "Si"
                                 "SMA"
## [64] "Vimentin"
```

Metadata

There are 8 patient metadata in the dataset, they are patient_id, AGE_AT_DX, STAGE, SITE_02, LATERAL, GRADE, RECURRENCE_LABEL, Survival_days_capped. We want to investigate on their unique number and metadata information for each patient.

```
# unique number for each metadata
data.frame(patient_id = length(unique(TNBC$patient_id)),
           AGE AT DX = length(unique(TNBC$AGE AT DX)),
           STAGE = length(unique(TNBC$STAGE)),
           SITE 02 = length(unique(TNBC$SITE 02)),
           LATERAL = length(unique(TNBC$LATERAL)),
           GRADE = length(unique(TNBC$GRADE)),
           RECURRENCE_LABEL = length(unique(TNBC$RECURRENCE_LABEL)),
           Survival_days_capped = length(unique(TNBC$Survival_days_capped)))
     patient_id AGE_AT_DX STAGE SITE_02 LATERAL GRADE RECURRENCE_LABEL
##
## 1
             39
                       28
                              8
##
     Survival_days_capped
## 1
# unique values
# patient_id
print("patien_id")
## [1] "patien_id"
data.frame(patient_id = unique(TNBC$patient_id)) %>% t()
               [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11]
##
## patient_id 30824 30805 30812 30838 30865 30847 30846 30783 30781 30782 30753
              [,12] [,13] [,14] [,15] [,16] [,17] [,18] [,19] [,20] [,21] [,22]
## patient_id 30770 30766 30744 30742 30734 30786 30739 30762 30785 30789 30843
              [,23] [,24] [,25] [,26] [,27] [,28] [,29] [,30] [,31] [,32] [,33]
## patient_id 30868 30866 30821 30827 30818 30823 30851 30853 30799 30854 30732
              [,34] [,35] [,36] [,37] [,38] [,39]
##
## patient_id 30771 30738 30765 30860 30740 30754
# AGE AT DX
print("AGE_AT_DX")
## [1] "AGE_AT_DX"
data.frame(AGE_AT_DX = unique(TNBC$AGE_AT_DX)) %>% t()
             [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11] [,12] [,13]
##
## AGE_AT_DX
               77
                    67
                         42
                              41
                                   64
                                        53
                                             26
                                                  79
                                                       60
                                                              38
                                                                    31
                                                                          37
                                                                                39
             [,14] [,15] [,16] [,17] [,18] [,19] [,20] [,21] [,22] [,23] [,24]
                            35
                                        68
                                                                 75
## AGE_AT_DX
                      50
                                  52
                                              36
                                                    49
                                                           43
                                                                       63
                45
             [,25] [,26] [,27] [,28]
## AGE_AT_DX
                59
                      62
                            91
```

```
# STAGE
print("STAGE")
## [1] "STAGE"
data.frame(STAGE = unique(TNBC$STAGE)) %>% t()
        [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8]
## STAGE 33 32 21
                         22 11
                                 10
# SITE_02
print("SITE_02")
## [1] "SITE_02"
data.frame(SITE_02 = unique(TNBC$SITE_02)) %>% t()
##
          [,1]
                 [,2]
                        [,3]
                             [,4]
                                    [,5]
                                            [,6]
                                                   [,7]
## SITE_02 "C504" "C509" "C505" "C508" "C503" "C502" "C501"
# LATERAL
print("LATERAL")
## [1] "LATERAL"
data.frame(LATERAL = unique(TNBC$LATERAL)) %>% t()
          [,1] [,2]
## LATERAL 2 1
# GRADE
print("GRADE")
## [1] "GRADE"
data.frame(GRADE = unique(TNBC$GRADE)) %>% t()
        [,1] [,2] [,3] [,4] [,5]
## GRADE 1 3
# RECURRENCE_LABEL
print("RECURRENCE_LABEL")
## [1] "RECURRENCE_LABEL"
```

```
data.frame(RECURRENCE_LABEL = unique(TNBC$RECURRENCE_LABEL)) %>% t()
##
                    [,1]
                                [,2]
## RECURRENCE_LABEL "POSITIVE" "NEGATIVE"
# Survival_days_capped
print("Survival days capped")
## [1] "Survival_days_capped"
data.frame(Survival days capped = unique(TNBC$Survival days capped)) %>% t()
                        [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11]
## Survival_days_capped 2612 745 3130 2523 1683 2275 946 3767 3822 3774 4353
                        [,12] [,13] [,14] [,15] [,16] [,17] [,18] [,19] [,20]
## Survival_days_capped 1072 4145
                                       530 2842 5063 3725 4761
##
                        [,21] [,22] [,23] [,24] [,25] [,26] [,27] [,28] [,29]
## Survival_days_capped 2438 1568
                                    1738
                                           2832 2759
                                                       3063
                                                             2853
                                                                    2096 3573
##
                        [,30] [,31] [,32] [,33] [,34] [,35] [,36] [,37] [,38]
## Survival days capped 3355
                                584
                                       635
                                             194 4785
                                                       3658 1009
                                                                    1754 4430
# metadata information for each sample_id
# n_{1}(1,2,3,4,5,6,7) are the unique values of the metadata in one sample
TNBC %>%
  group by (sample id) %>%
  summarise(n 1 = length(unique(AGE AT DX)),
            AGE_AT_DX = unique(AGE_AT_DX),
            n_2 = length(unique(STAGE)),
            STAGE = unique(STAGE),
            n_3 = length(unique(SITE_02)),
            SITE 02 = unique(SITE 02),
            n_4 = length(unique(LATERAL)),
            LATERAL = unique(LATERAL),
            n_5 = length(unique(GRADE)),
            GRADE = unique(GRADE),
            n_6 = length(unique(RECURRENCE_LABEL)),
            RECURRENCE LABEL = unique(RECURRENCE LABEL),
            n_7 = length(unique(Survival_days_capped)),
            Survival_days_capped = unique(Survival_days_capped))
## # A tibble: 39 x 15
##
      sample_id
                  n_1 AGE_AT_DX
                                  n_2 STAGE
                                               n_3 SITE_02
                                                             n_4 LATERAL
                                                                            n_5 GRADE
                          <dbl> <int> <dbl> <int> <chr>
                                                                    <dbl> <int> <dbl>
##
          <dbl> <int>
                                                           <int>
##
   1
              1
                    1
                             77
                                     1
                                          33
                                                 1 C504
                                                                1
                                                                        2
                                                                              1
                                                                                    1
              2
                             67
                                          32
                                                 1 C509
                                                                        2
                                                                              1
                                                                                    3
##
   2
                    1
                                     1
                                                               1
##
   3
              3
                    1
                             42
                                          21
                                                 1 C509
                                                               1
                                                                        2
                                                                              1
                                                                                    3
                                     1
                                                 1 C505
   4
              4
                             41
                                          22
                                                                        2
                                                                              1
                                                                                    3
##
                    1
                                     1
                                                                1
##
   5
              5
                    1
                             64
                                    1
                                          11
                                                 1 C508
                                                                1
                                                                        1
                                                                              1
                                                                                    3
   6
              6
                             53
                                                 1 C508
                                                                                    3
##
                    1
                                    1
                                          10
                                                               1
                                                                        1
                                                                              1
   7
              7
                    1
                             62
                                    1
                                          32
                                                 1 C509
                                                                        2
                                                                                    2
              8
                             26
                                          21
                                                 1 C503
                                                                                    3
##
   8
                    1
                                    1
                                                               1
                                                                        1
                                                                              1
```

```
##
                              79
                                      1
                                           21
                                                   1 C504
## 10
              10
                     1
                              60
                                      1
                                           22
                                                   1 C502
                                                                 1
                                                                          2
                                                                                1
                                                                                       3
## # ... with 29 more rows, and 4 more variables: n 6 <int>,
       RECURRENCE_LABEL <chr>, n_7 <int>, Survival_days_capped <dbl>
```

Uniqueess of patient information

There are total 39 patients in the dataset with their corresponding 39 images. The original study from Keren et al. in 2018 have 41 patients however the images for patient 22 and 38 failed the quality check and were left out in the analysis.

The following code chunk will check the uniquness of sample_id, patient_id and ImageNb.

```
# unique values
unique_sample <- unique(TNBC$sample_id)
unique_patient <- unique(TNBC$patient_id)
unique_image <- unique(TNBC$ImageNb)

# if sample_id and patient_id have same length
dplyr::setequal(unique_sample, unique_patient)</pre>
## [1] FALSE
```

```
# if sample_id and ImageNb have same values
all.equal(unique_sample, unique_image)
```

```
## [1] TRUE
```

```
## # A tibble: 39 x 6
##
      sample_id cell_count num_unique_image ImageNb num_unique_patient patient_id
           <dbl>
##
                                                    <dbl>
                                                                                     <dbl>
                       <int>
                                           <int>
                                                                         <int>
##
    1
               1
                        5199
                                               1
                                                        1
                                                                              1
                                                                                     30824
##
    2
               2
                        3033
                                               1
                                                        2
                                                                              1
                                                                                     30805
               3
                                                        3
##
    3
                        5671
                                               1
                                                                              1
                                                                                     30812
               4
                                               1
                                                        4
                                                                                     30838
##
    4
                        5381
                                                                              1
               5
                                                        5
##
    5
                        4252
                                               1
                                                                              1
                                                                                     30865
##
    6
               6
                        4894
                                               1
                                                        6
                                                                              1
                                                                                     30847
##
    7
               7
                        3308
                                               1
                                                        7
                                                                              1
                                                                                     30854
               8
                                               1
                                                        8
##
    8
                        3786
                                                                              1
                                                                                     30846
                                                        9
##
    9
               9
                        5105
                                               1
                                                                              1
                                                                                     30783
                                               1
                                                       10
## 10
              10
                        4066
                                                                              1
                                                                                     30781
## # ... with 29 more rows
```

Uniqueness of cell type

The columns cell_type and mm are the cell types for each cells, where cell_type is the factor level number and mm is the names of the cell_type. There are 16 different cell types in the dataset.

The following code chunk will perform the diagnostics of these two columns.

```
# unique values
unique_celltype <- unique(TNBC$cell_type)</pre>
unique_mm <- unique(TNBC$mm)</pre>
# if cell_type and mm have the same number
dplyr::setequal(length(unique_celltype), length(unique_mm))
## [1] TRUE
# since true, if cell_type and mm are unique to each other
TNBC %>%
  group_by(mm) %>%
  summarise(cell_count = n(),
            num_unique_cellType = length(unique(cell_type)), cell_type = unique(cell_type))
## # A tibble: 16 x 4
##
                   cell_count num_unique_cellType cell_type
      mm
                                                        <dbl>
##
      <chr>>
                         <int>
                                              <int>
                         17084
                                                            5
##
   1 B
                                                  1
   2 CD3 T
##
                          1135
                                                  1
                                                            4
                                                            2
##
   3 CD4 T
                          9918
                                                  1
##
  4 CD8 T
                         13376
                                                  1
                                                            3
## 5 DC
                          2381
                                                  1
                                                           10
## 6 DC/Mono
                          1280
                                                  1
                                                           11
## 7 Endothelial
                           279
                                                  1
                                                           13
## 8 Epithelial
                         31871
                                                  1
                                                           14
## 9 Mac
                          5552
                                                  1
                                                            7
## 10 Mesenchymal
                         27698
                                                  1
                                                           15
## 11 Mono/Neu
                           835
                                                  1
                                                            9
## 12 Neu
                                                            8
                          1365
                                                  1
## 13 NK
                           285
                                                  1
                                                            6
## 14 Other
                         56603
                                                  1
                                                           16
## 15 Other immune
                          8669
                                                  1
                                                           12
                           863
## 16 T reg
                                                  1
                                                            1
## Conclusion: cell_type and mm are unique to each other(they are redunant)
# names of the cell types
unique_mm
                                        "CD4 T"
                                                                       "DC"
##
    [1] "B"
                        "CD3 T"
                                                       "CD8 T"
   [6] "DC/Mono"
                                                       "Mac"
                        "Endothelial" "Epithelial"
                                                                       "Mesenchymal"
## [11] "Mono/Neu"
                        "Neu"
                                        "NK"
                                                       "Other"
                                                                       "Other immune"
## [16] "T reg"
```

```
# tibble of number of samples each cell type present
TNBC %>%
  group_by(mm) %>%
  summarise(cell_count = n(), NumOfSamples = n_distinct(sample_id))
```

```
## # A tibble: 16 x 3
##
      mm
                    cell_count NumOfSamples
##
      <chr>
                          <int>
                                       <int>
##
    1 B
                         17084
                                           31
    2 CD3 T
                                           22
##
                          1135
##
   3 CD4 T
                          9918
                                           36
##
   4 CD8 T
                          13376
                                           36
##
  5 DC
                                           34
                          2381
##
   6 DC/Mono
                          1280
                                           28
  7 Endothelial
                                           25
##
                            279
##
    8 Epithelial
                         31871
                                           39
## 9 Mac
                                           38
                          5552
## 10 Mesenchymal
                         27698
                                           39
                                           36
## 11 Mono/Neu
                           835
## 12 Neu
                                           36
                          1365
## 13 NK
                            285
                                           26
## 14 Other
                         56603
                                           39
## 15 Other immune
                                           36
                          8669
## 16 T reg
                           863
                                           22
```

$Cluster_id$

Cells are clustered into different cell clusters by applying the technique FlowSOM, and it has 113 cluster for all cells in the TNBC dataset.

```
# unique cluster_id
unique_clusterId <- sort(unique(TNBC$cluster_id))
length(unique_clusterId)</pre>
```

[1] 113

unique_clusterId

```
6
                                      7
                                                                                     18
##
     [1]
            1
                2
                    3
                         4
                             5
                                          8
                                               9
                                                  10
                                                      11
                                                           12
                                                               13
                                                                   14
                                                                        15
                                                                            16
                                                                                 17
##
    [19]
          19
               20
                   21
                       22
                            23
                                24
                                     25
                                         26
                                              27
                                                  28
                                                      29
                                                           30
                                                               31
                                                                    32
                                                                        33
                                                                            34
                                                                                 35
                                                                                     36
##
    [37]
          37
               38
                   39
                       40
                            41
                                42
                                     43
                                         44
                                              45
                                                  46
                                                      47
                                                           48
                                                               49
                                                                    50
                                                                        51
                                                                            52
                                                                                 53
                                                                                     54
                                         62
##
    [55]
          55
               56
                   57
                       58
                            59
                                60
                                     61
                                              63
                                                  64
                                                      65
                                                           66
                                                               67
                                                                    68
                                                                        69
                                                                            70
                                                                                 71
                                                                                     72
##
   [73]
          73
               74
                   75
                       76
                            77
                                78
                                     79
                                         80
                                              81
                                                  82
                                                      83
                                                           84
                                                               85
                                                                    86
                                                                        87
                                                                            88
                                                                                     90
          91
                                             99 100 101 113 114 128 129 143 157 158
##
   [91]
              92
                   93
                            95
                                96 97
                                         98
                       94
## [109] 159 172 175 187 211
```

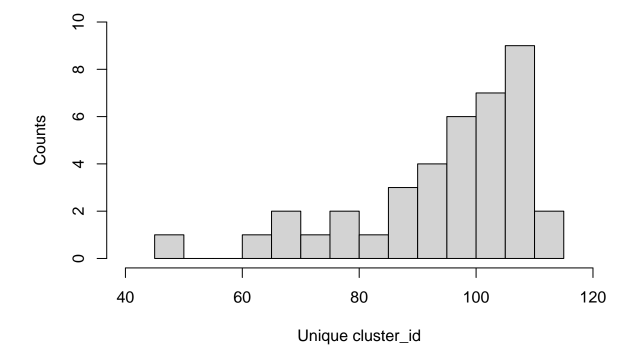
The following tibble and graphs show number of cluster_id presents in each sample.

```
# cluster_id for each sample_id
TNBC_clusterId <- TNBC %>%
  group_by(sample_id) %>%
```

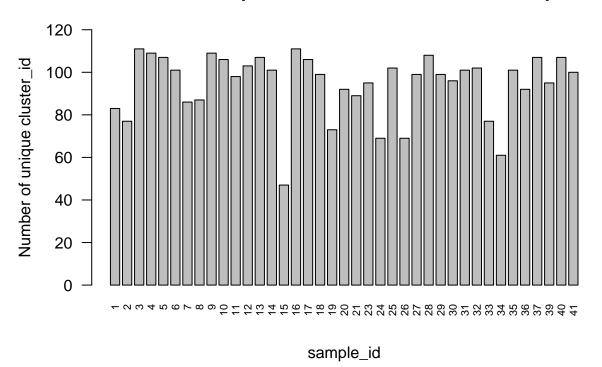
```
## # A tibble: 39 x 3
##
      sample_id cell_count num_unique_cluster
           <dbl>
##
                       <int>
                                           <int>
##
    1
               1
                       5199
                                              83
               2
                                              77
##
                       3033
##
               3
                       5671
                                              111
               4
##
                       5381
                                              109
    5
               5
                       4252
                                              107
##
    6
               6
                       4894
                                              101
    7
               7
##
                       3308
                                              86
##
                       3786
                                              87
    9
               9
                                              109
##
                       5105
## 10
              10
                        4066
                                              106
## # ... with 29 more rows
```

```
# Histogram
hist(TNBC_clusterId$num_unique_cluster,
    main = "Histogram of Unique Number Clusters for each sample_id",
    xlab = "Unique cluster_id", ylab = "Counts",
    breaks = 20, xlim = c(40, 120), ylim = c(0, 10))
```

Histogram of Unique Number Clusters for each sample_id



Count of the Unique Number Clusters for Each sample_id



Spatial Information

The two columns, centroidX and centroidY contain the spatial coordinates x and y for cells in each sample.

All spatial coordinates We first want to investigate the range of all the spatial coordinates by finding their range, median and standard deviation.

The range of all spatial coordinate x is (4.512, 2043.474), the median is 1054.774 and standard deviation is 577.792; the range of all spatial coordinates y is (3.514, 2044.483), the median is 1055.1165 and standard deviation is 580.025.

```
# subset spatial coordinates
centroidX <- TNBC$centroidX
centroidY <- TNBC$centroidY

# Range for the whole dataset
range_wholeX <- c(min(centroidX), max(centroidX), median(centroidX), sd(centroidX))
range_wholeX</pre>
```

[1] 4.511905 2043.474000 1054.774000 577.792180

```
range_wholeY <- c(min(centroidY), max(centroidY), median(centroidY), sd(centroidY))
range_wholeY</pre>
```

[1] 3.514286 2044.483000 1055.116500 580.024631

Each cell types We now want to investigate the spatial coordinates by cell types in TNBC dataset. The following tibbles include the counts for each cell type along with the range, median and standard deviation of their spatial coordinates in all samples.

mm	count	min	max	median	sd
В	17084	4.582417	2043.392	1068.6680	541.1996
CD3 T	1135	6.784946	2043.073	1424.7880	500.7704
CD4 T	9918	4.610000	2043.281	1323.7545	559.5474
CD8 T	13376	4.688073	2043.297	1110.1715	570.1150
DC	2381	4.566929	2043.363	948.9395	645.9324
DC/Mono	1280	8.995781	2043.230	1308.6955	499.2984
Endothelial	279	28.724440	2022.824	1309.8260	535.6648
Epithelial	31871	4.567416	2043.380	1118.1760	566.4550
Mac	5552	4.686047	2043.274	1194.2770	559.2432
Mesenchymal	27698	4.555556	2043.347	1028.6410	575.8002
Mono/Neu	835	10.019080	2042.503	1266.4960	546.5867
Neu	1365	8.695842	2042.541	1153.5900	569.1121
NK	285	15.819670	2029.213	1343.4280	523.0047
Other	56603	4.511905	2043.474	914.0930	587.5303
Other immune	8669	4.641791	2043.379	1056.2430	572.1199
T reg	863	9.922190	2039.853	1233.3820	511.7799

-					
mm	count	min	max	median	sd
В	17084	4.600000	2044.441	1143.6310	577.0482
CD3 T	1135	14.928100	2043.847	1070.0920	479.8067
CD4 T	9918	3.855556	2044.359	1106.5920	553.1205

mm	count	min	max	median	sd
CD8 T	13376	4.039474	2044.379	1095.8965	574.7819
DC	2381	5.198718	2043.463	1122.3520	618.9403
DC/Mono	1280	3.650000	2043.193	1003.7210	534.9473
Endothelial	279	20.383230	2010.723	901.5644	493.5723
Epithelial	31871	4.493421	2044.347	949.2707	558.0874
Mac	5552	5.909605	2043.214	977.1770	569.4235
Mesenchymal	27698	3.777778	2044.483	979.0533	580.2001
Mono/Neu	835	10.988020	2044.450	1257.1100	531.7334
Neu	1365	7.002710	2042.952	1213.8160	538.2672
NK	285	18.128130	2042.485	1040.5380	540.3429
Other	56603	3.514286	2044.471	1121.4110	594.5656
Other immune	8669	4.250000	2044.372	1041.3730	587.8098
T reg	863	10.049020	2038.496	973.4490	525.8779

Each cell type in each sample We next want to group the TNBC data by sample number and cell type, and investigate the spatial coordinates. The following tibbles include <code>sample_id</code> and cell type information along with the range, median and standard deviation of their spatial coordinates in all samples.

```
## centroidX
TNBC %>%
  group_by(sample_id, mm) %>%
  summarise(n = n(),
            min = min(centroidX), max = max(centroidX),
            median = median(centroidX), sd = sd(centroidX))
## # A tibble: 523 x 7
  # Groups:
               sample_id [39]
##
      sample_id mm
                                              max median
                                                             sd
                                 n
                                       min
##
          <dbl> <chr>
                                      <dbl> <dbl>
                                                   <dbl> <dbl>
                             <int>
##
                                                   1062.
   1
              1 B
                               734
                                      34.6
                                            2022.
                                                           321.
##
    2
              1 CD4 T
                                            2043.
                                                   1566.
                               152
                                    100.
              1 CD8 T
                                                   1147.
##
    3
                               147
                                       9.66 2041.
                                                           615.
##
    4
              1 DC
                                 1
                                    238.
                                             238.
                                                    238.
                                     44.7
                                            2042.
                                                   1742.
                                                           584.
##
    5
              1 Epithelial
                                20
                                    347.
                                            2003.
                                                   1520.
    6
              1 Mac
                                10
                                                           528.
    7
              1 Mesenchymal
                               289
                                     36.4
                                            2042.
                                                   1074.
                                                           518.
##
                                                   1739.
##
    8
              1 Mono/Neu
                                 2 1540.
                                            1938.
                                                           281.
##
   9
              1 Neu
                                 2 1159.
                                            1572.
                                                   1365.
                                                           292.
## 10
              1 NK
                                 4 610.
                                            1141.
                                                    644.
                                                           255.
## # ... with 513 more rows
## centroidY
TNBC %>%
  group_by(sample_id, mm) %>%
  summarise(n = n(),
            min = min(centroidY), max = max(centroidY),
            median = median(centroidY), sd = sd(centroidY))
```

A tibble: 523 x 7

sample_id [39]

Groups:

##	sa	ample_id	mm	n	min	max	${\tt median}$	sd
##		<dbl></dbl>	<chr></chr>	<int></int>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
##	1	1	В	734	6.73	2043.	1793.	525.
##	2	1	CD4 T	152	12.7	2000.	1005.	635.
##	3	1	CD8 T	147	12.1	1998.	1074.	624.
##	4	1	DC	1	412.	412.	412.	NA
##	5	1	Epithelial	20	6.44	818.	240.	284.
##	6	1	Mac	10	418.	1767.	1100.	461.
##	7	1	Mesenchymal	289	8.05	2038.	596.	538.
##	8	1	Mono/Neu	2	170.	1415.	793.	880.
##	9	1	Neu	2	988.	1686.	1337.	494.
##	10	1	NK	4	301.	1086.	872.	349.
##	#	. with 5	13 more rows					