Preprocessing

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Import data

Import the bioactivity data obtained from the ChEMBL database as a dataframe. Select the IC50 data subset.

```
library(readxl)
## Warning: package 'readxl' was built under R version 3.2.4
df_raw <- suppressWarnings(read_excel("Human_AChE.xlsx"))</pre>
names <- c("CMPD_CHEMBLID", "CANONICAL_SMILES", "STANDARD_TYPE",
           "RELATION", "STANDARD_VALUE", "STANDARD_UNITS",
           "PROTEIN_ACCESSION", "PREF_NAME",
      "PUBMED_ID",
      "JOURNAL", "YEAR", "VOLUME",
      "ISSUE", "FIRST_PAGE", "MOLWEIGHT",
      "ALOGP",
           "PSA", "NUM_RO5_VIOLATIONS")
df <- df raw[, names]</pre>
df_1 <- subset(df, STANDARD_TYPE == "IC50")</pre>
df_2 <- subset(df_1, RELATION == "=")</pre>
df_4 <- subset(df_2, STANDARD_UNITS == "nM")</pre>
bioactivity <- paste0("Bioactivity data points: ", nrow(df))
IC50 <- paste0("IC50 Bioactivity data points: ", nrow(df_1))</pre>
IC50_omit <- paste0("Omit IC50 with lesser/greater than symbol: ", nrow(df_2))
final <- paste0("Final number of data points: ", nrow(df_4))</pre>
message <- c(bioactivity, IC50, IC50_omit, final)</pre>
print(message)
## [1] "Bioactivity data points: 9242"
## [2] "IC50 Bioactivity data points: 4910"
## [3] "Omit IC50 with lesser/greater than symbol: 3609"
## [4] "Final number of data points: 3596"
```

Remove redundant rows

```
ID <- df_4$CMPD_CHEMBLID
IC50_nm <- df_4$STANDARD_VALUE
smiles <- df_4$CANONICAL_SMILES
data_1 <- data.frame(ID, smiles, IC50_nm)

duplicate <- smiles[duplicated(smiles)]
unique <- unique(duplicate)
unique <- as.character(unique)
results <- data.frame()</pre>
```

```
for (i in unique) {
    pre_data <- subset(data_1, smiles == i)</pre>
    results <- rbind(results, pre_data)</pre>
}
results_2 <- data.frame()
for (i in unique) {
    class <- subset(results, smiles == i)</pre>
    IC50 <- class$IC50_nm</pre>
    mean_IC50_nm <- mean(IC50)</pre>
    sd \leftarrow sd(IC50)
    data_frame <- data.frame(i, mean_IC50_nm, sd)</pre>
    results_2 <- rbind(data_frame, results_2)</pre>
}
keep <- subset(results_2, sd < 2)</pre>
names(keep) <- c("CANONICAL_SMILES", "STANDARD_VALUE", "SD")</pre>
non_redundant <- df_4[!duplicated(df_4$CANONICAL_SMILES), ]</pre>
non_redundant <- non_redundant[!non_redundant$CANONICAL_SMILES %in% unique,
    ]
non_redundant <- non_redundant[, c("CANONICAL_SMILES", "STANDARD_VALUE")]</pre>
redundant <- df_4[df_4$CANONICAL_SMILES %in% unique, ]</pre>
redundant <- subset(redundant, !duplicated(redundant[, c("CANONICAL_SMILES")]))</pre>
redundant <- subset(redundant, select = -STANDARD_VALUE)</pre>
STANDARD_VALUE <- keep$STANDARD_VALUE
CANONICAL_SMILES <- keep$CANONICAL_SMILES
cleaned_redundant <- data.frame(CANONICAL_SMILES, STANDARD_VALUE)</pre>
curated_data <- rbind(non_redundant, cleaned_redundant)</pre>
curated_data_na_removed <- na.omit(curated_data)</pre>
unique <- paste0("number of unique redundant compounds: ", length(unique))
sd_2 <- paste0("Compounds left that are kept", " which has a sd of lower than two: ",</pre>
    nrow(keep))
combined <- paste0("Combined data points: ", nrow(curated_data))</pre>
non_redundant_compound <- paste0("number of non-redundant compounds: ", nrow(non_redundant))</pre>
curated <- paste0("number of curated data points where missing smiles are removed: ",</pre>
    nrow(curated_data_na_removed))
message <- c(non_redundant_compound, unique, sd_2, combined, curated)
print(message)
## [1] "number of non-redundant compounds: 2426"
## [2] "number of unique redundant compounds: 340"
## [3] "Compounds left that are kept which has a sd of lower than two: 145"
## [4] "Combined data points: 2571"
## [5] "number of curated data points where missing smiles are removed: 2570"
```