

COMPASS Synoptic CB Porewater: DOC

November 2024 Samples

2025-10-31

```
##Setup - Change things here & write any notes
```

```
#identify section  
cat("Setup Information")
```

```
## Setup Information
```

```
##### Run information - PLEASE CHANGE  
Date_Run = "11/27/24" #Date that instrument was run  
Run_by = "Stephanie J. Wilson" #Instrument user  
Script_run_by = "Stephanie J. Wilson" #Code user  
run_notes = "TN Standard Curve stopped then restarted, but looks fine." #any notes from the run  
samples <- c("GCW", "GWI", "MSM", "SWH") #whatever identifies your samples within the same names  
samples_pattern <- paste(samples, collapse = "|")  
#samples_pattern <- "GCW" #use this instead of the line above if you have only one site code  
chks_name = "Chk_Std_50C_2N" #what did you name your check standards?
```

```
##### File Names - PLEASE CHANGE
```

```
#file path and name for raw summary data file  
#raw_file_name = "Raw Data/COMPASS_SynopticCB_PW_DOC_202505.txt" #example  
raw_file_name = "Raw Data/COMPASS_SynopticCB_PW_DOC_202411.txt"
```

```
#file path and name for raw all peaks file
```

```
#raw_allpeaks_name = "Raw Data/COMPASS_SynopticCB_PW_DOC_202505_allpeaks.txt" #example  
raw_allpeaks_name = "Raw Data/COMPASS_SynopticCB_PW_DOC_202411_allpeaks.txt"
```

```
#file path and name of processed data file
```

```
#processed_file_name = "Processed Data/COMPASS_SynopticCB_PW_Processed_DOC_202505.csv" #example  
processed_file_name = "Processed Data/COMPASS_SynopticCB_PW_Processed_DOC_202411.csv" #example
```

```
##### Log Files - PLEASE CHECK
```

```
#downloaded metadata csv - downloaded from Google drive as csv for this year  
Raw_Metadata = "Raw Data/COMPASS_SynopticCB_PW_SampleLog_2024.csv"
```

```
#qaqc log file path for this year
```

```
Log_path = "Raw Data/COMPASS_Synoptic_TOCTN_QAQClog_2024.csv"
```

```
##Set Up Code
```

```
##Read in metadata and create similar sample IDs for matching to samples
```

Import Data Functions

Import Sample Data

```
## Import Sample Data

## New names:
## * `` -> '...14'

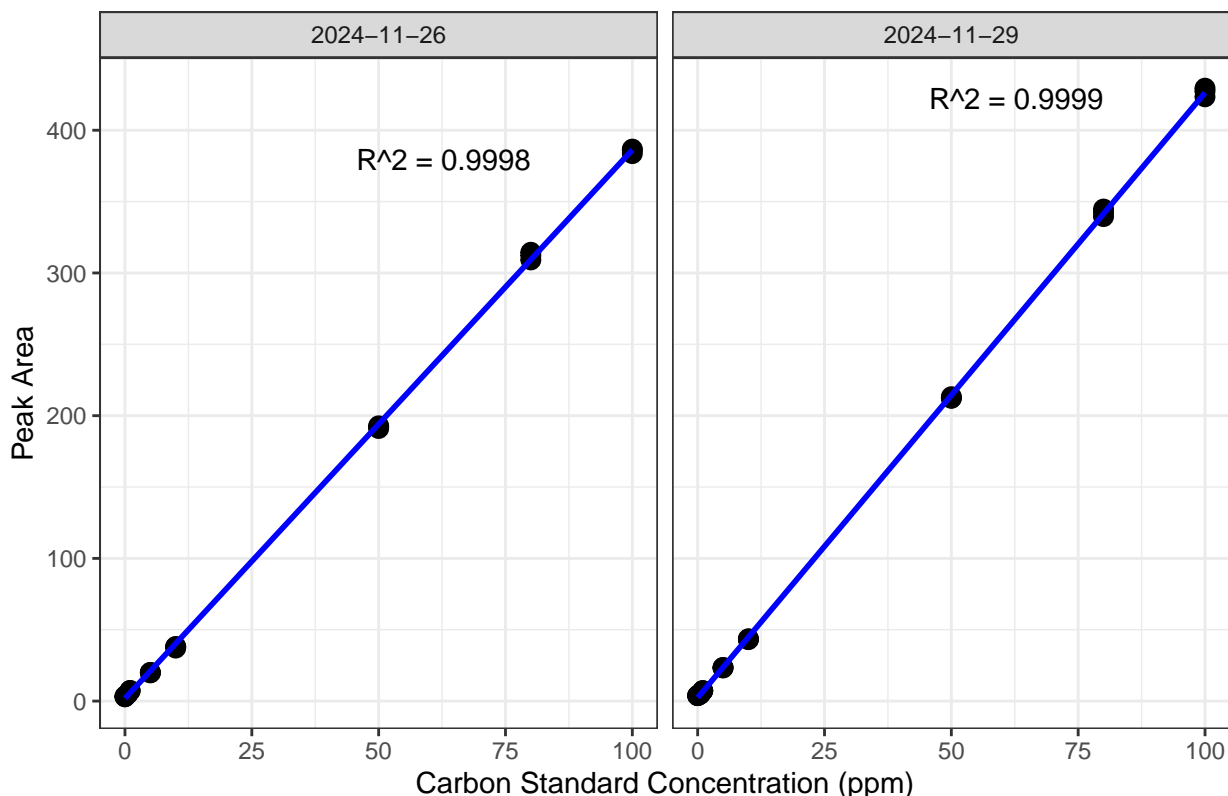
## # A tibble: 6 x 4
##   sample_name      npoc_raw    tdn_raw run_datetime
##   <chr>          <dbl>     <dbl>   <chr>
## 1 202411_SWH_UP_LysA_10cm 10.5     0.406  11/27/2024 3:11:21 AM
## 2 202411_SWH_UP_LysA_20cm  8.73    0.328  11/27/2024 3:38:16 AM
## 3 202411_SWH_UP_LysA_45cm 13.5     0.466  11/27/2024 4:05:40 AM
## 4 202411_SWH_UP_LysB_10cm 15.0     0.521  11/27/2024 4:22:06 AM
## 5 202411_SWH_UP_LysB_20cm  9.97    0.397  11/27/2024 4:49:42 AM
## 6 202411_SWH_UP_LysB_45cm 33.3     0.928  11/27/2024 5:17:47 AM
```

Assessing Standard Curves

```
## Assess the Standard Curves

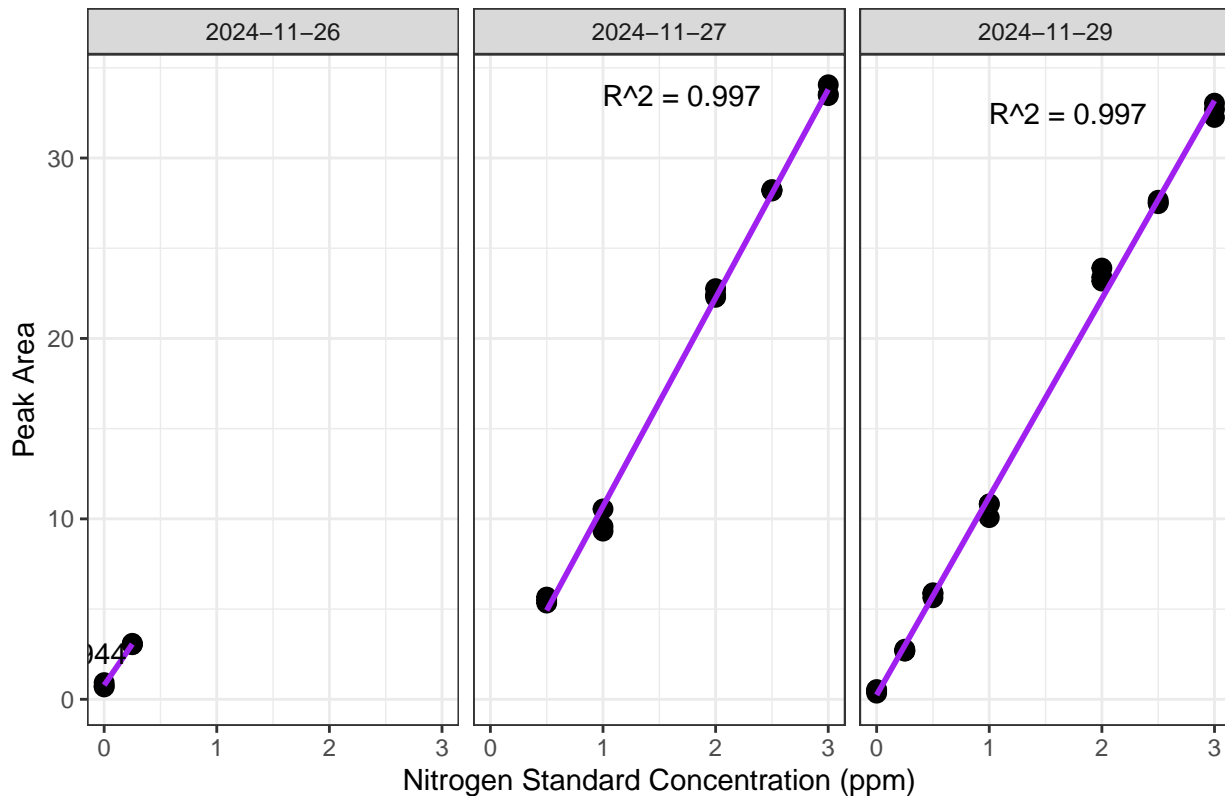
## New names:
## `geom_smooth()` using formula = 'y ~ x'
## * `` -> '...18'
```

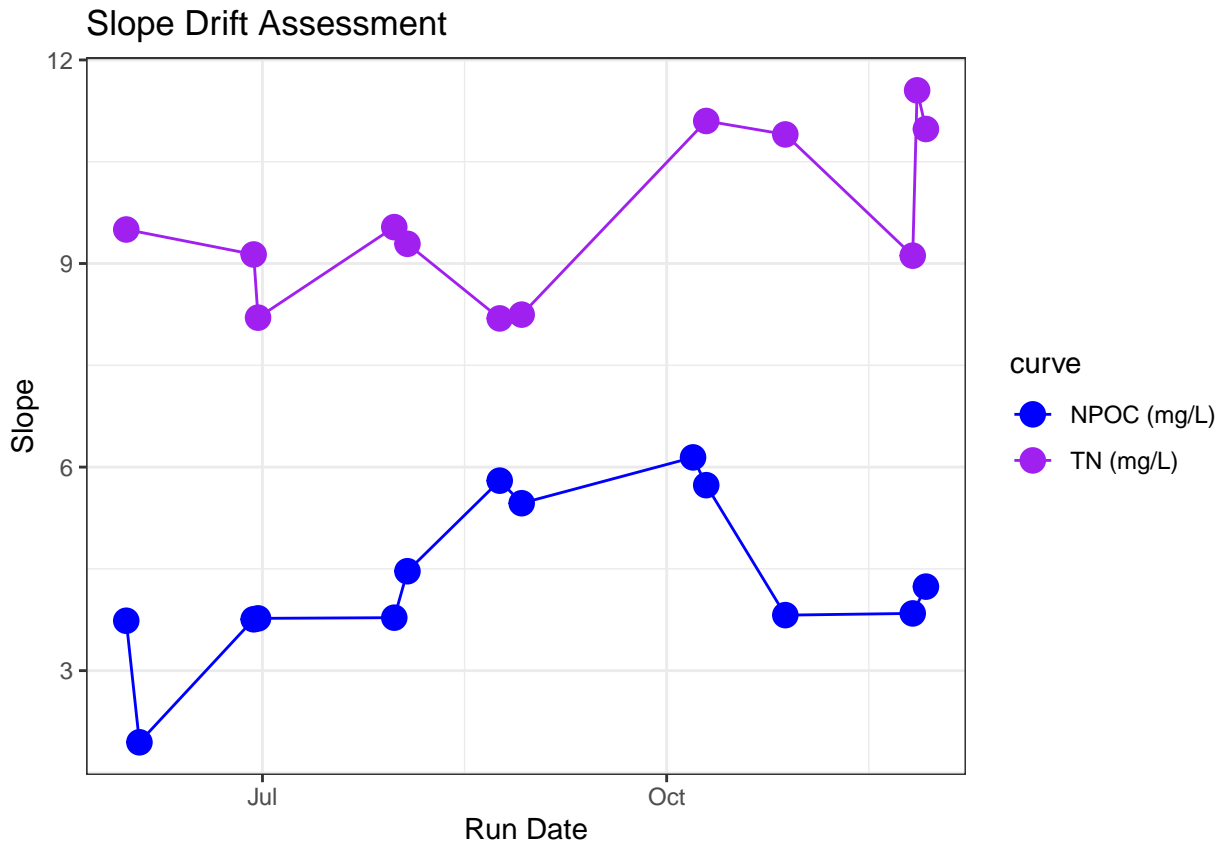
NPOC Std Curve by Date



```
## `geom_smooth()` using formula = 'y ~ x'
```

TN Std Curve by Date





```
## [1] "NPOC Curve r2 GOOD"
```

```
## [1] "TN Curve r2 GOOD"
```

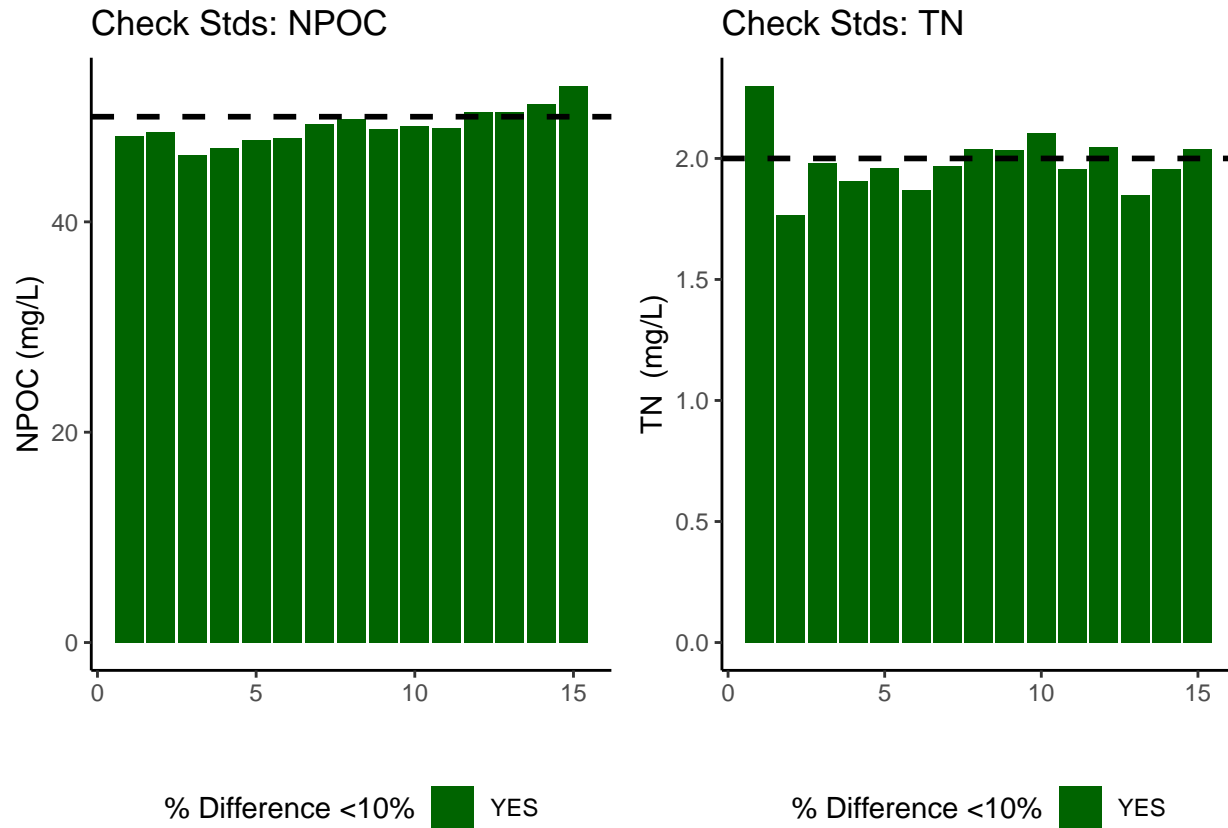
Assess Check Standards

```
## Assess the Check Standards
```

```
## New names:  
## * ` ` -> '...14'
```

```
## [1] "Carbon Check Standard RSD within Range"
```

```
## [1] "Nitrogen Check Standard RSD within Range"
```



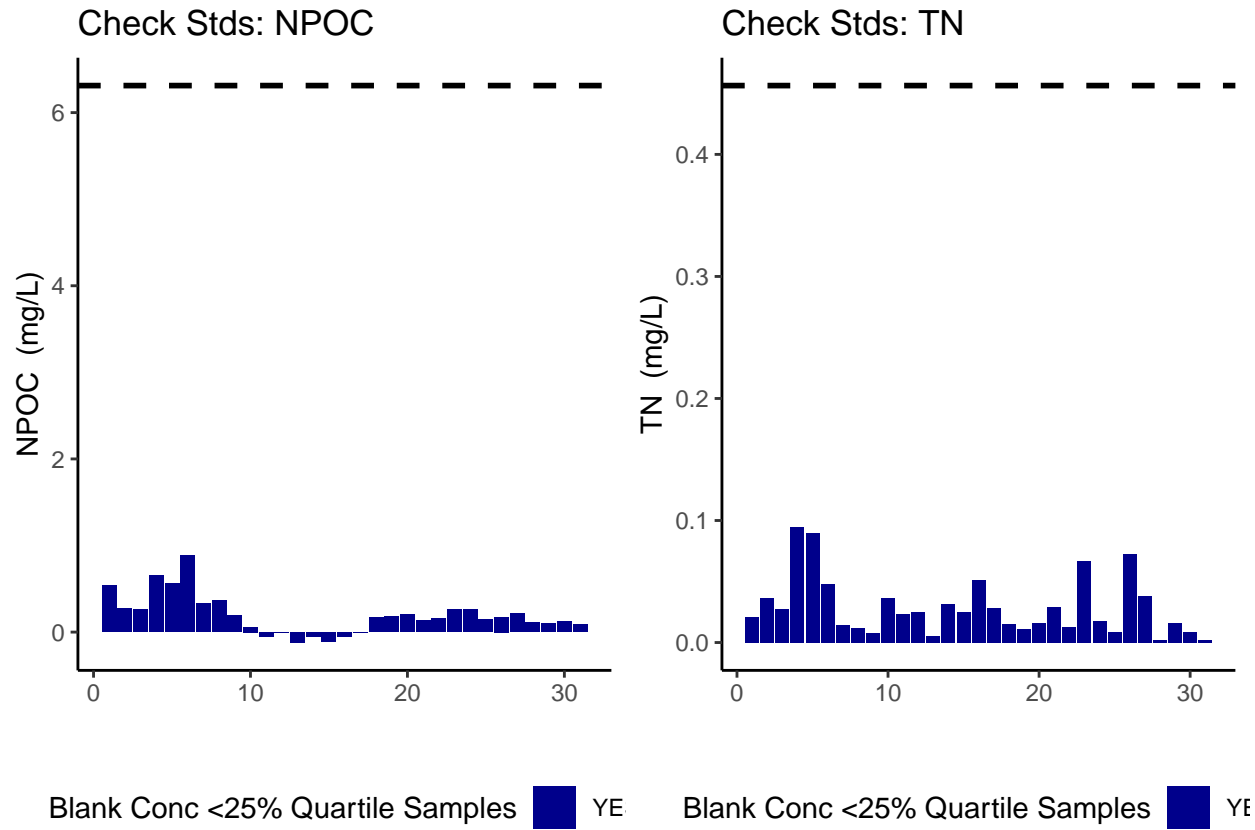
```
## [1] ">60% of Carbon Check Standards are within range of expected concentration"
## [1] ">60% of Nitrogen Check Standards are within range of expected concentration"
```

Assess Blanks

```
## Assess Blanks

## New names:
## * `` -> '...14'

## [1] ">60% of Carbon Blank concentrations are lower 25% quartile of samples"
## [1] ">60% of Nitrogen Blank concentrations are lower 25% quartile of samples"
```



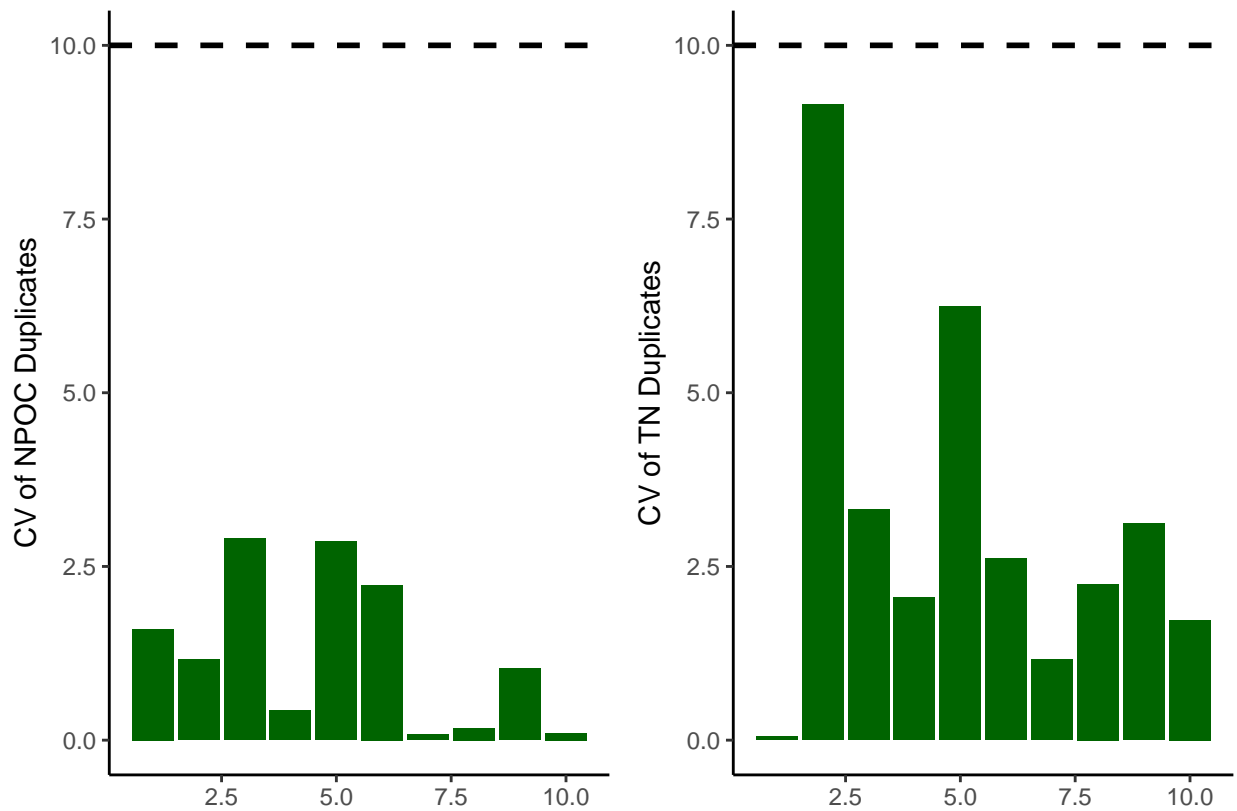
```
## carbon blanks:
## [1] 0.1949061

## nitrogen blanks:
## [1] 0.02876839

Assess Duplicates

## Assess Duplicates

## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use 'linewidth' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```



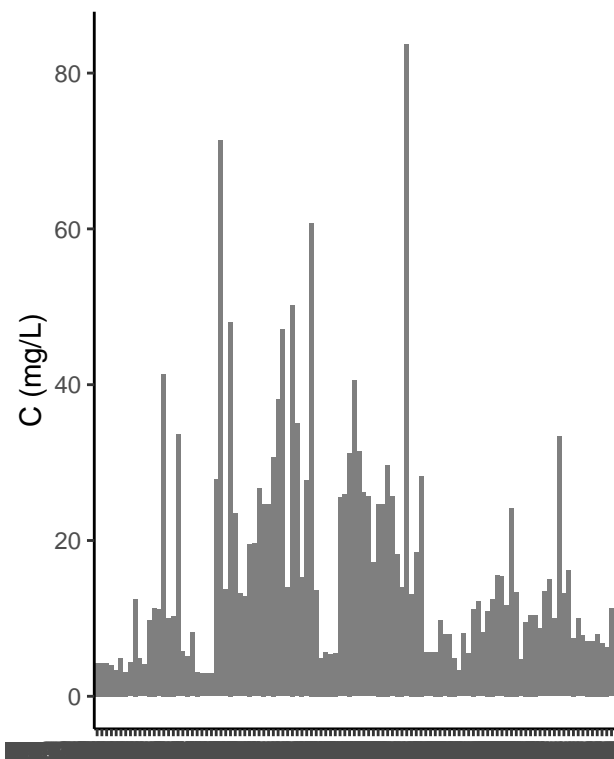
```
## [1] ">60% of Carbon Duplicates have a CV <10%"
```

```
## [1] ">60% of Nitrogen Duplicates have a CV <10%"
```

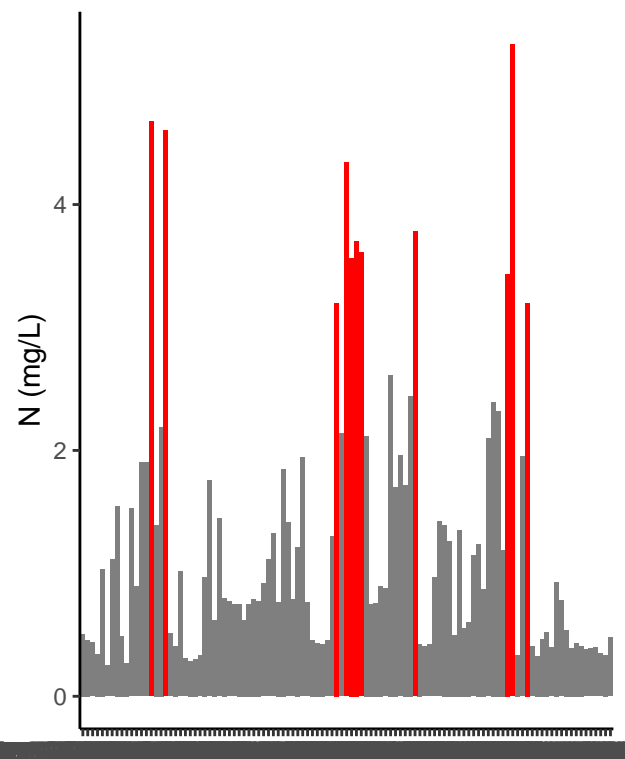
Sample Flagging - Are samples Within the range of the curve?

```
## Sample Flagging
```

C: Grey = Within Range of Curve



N: Grey = Within Range of Curve

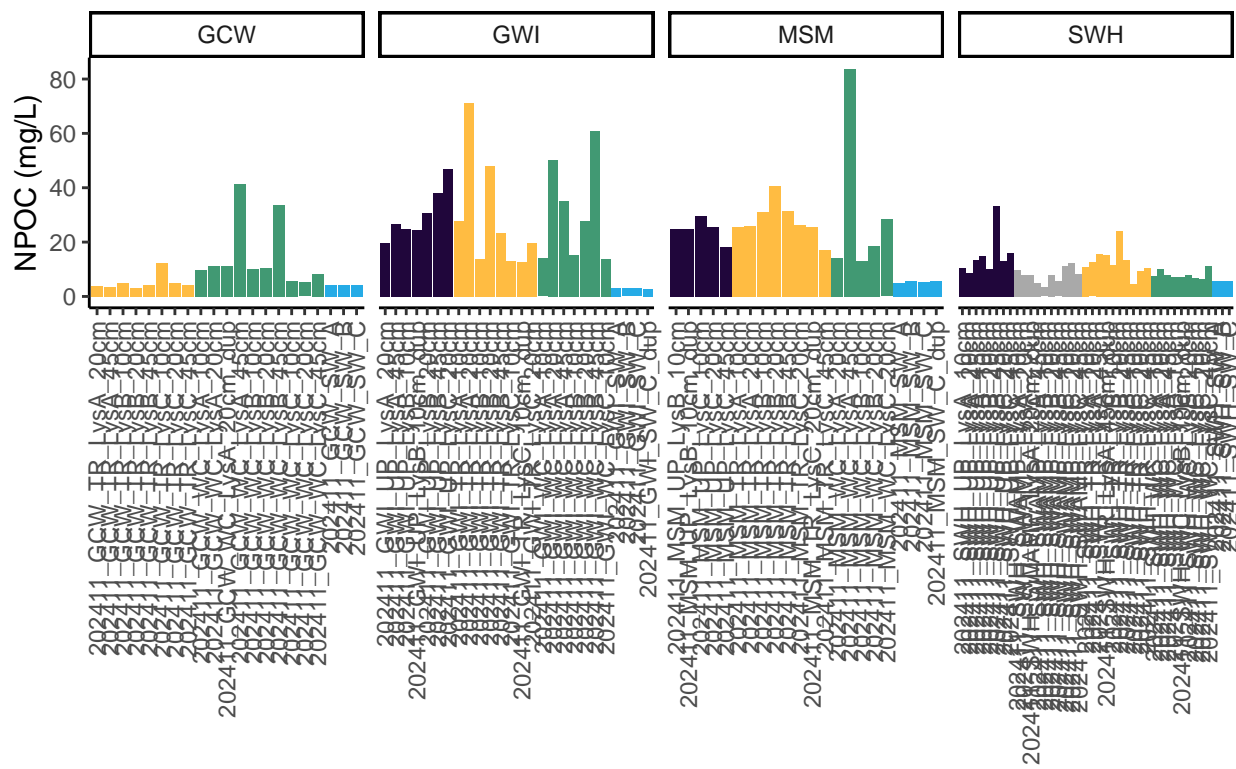


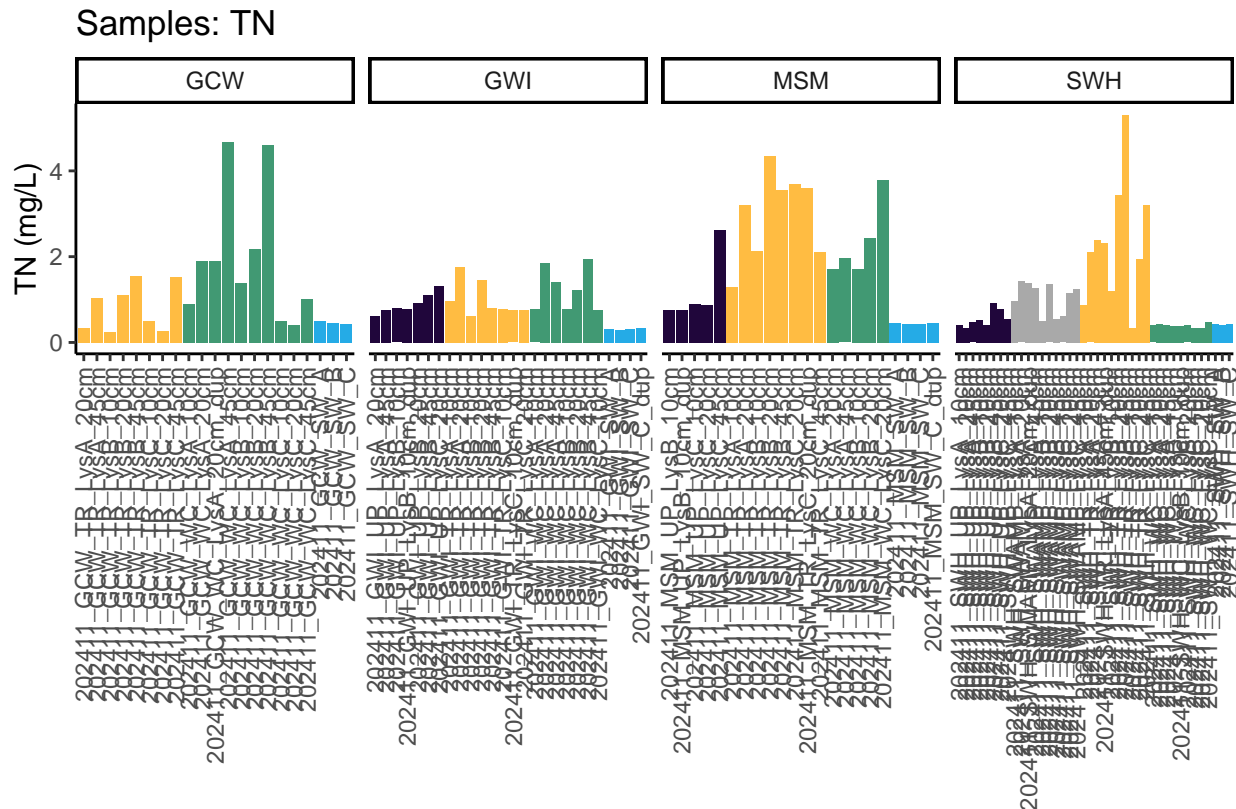
Visualize Data by Plot

```
## Visualize Data

## Warning in rbind(c("202411", "SWH", "UP", "LysA", "10cm"), c("202411", "SWH", :
## number of columns of result is not a multiple of vector length (arg 1)
```

Samples: NPOC





Convert data from mg/L to uMoles/L

Check to see if samples run match metadata & merge info

```
## Check Sample IDs with Metadata  
## All sample IDs are present in metadata.
```

Export Processed Data

Export Processed Data

```

## # A tibble: 6 x 21
##   Project      Region Site  Zone  Replicate Depth_cm Sample_ID Year Month Day
##   <chr>        <chr>  <chr> <fct> <chr>       <int> <chr>     <int> <int> <int>
## 1 COMPASS: Sy~ CB     SWH    UP     A           10 202411_S~ 2024   11   12
## 2 COMPASS: Sy~ CB     SWH    UP     A           20 202411_S~ 2024   11   12
## 3 COMPASS: Sy~ CB     SWH    UP     A           45 202411_S~ 2024   11   12
## 4 COMPASS: Sy~ CB     SWH    UP     B           10 202411_S~ 2024   11   12
## 5 COMPASS: Sy~ CB     SWH    UP     B           20 202411_S~ 2024   11   12
## 6 COMPASS: Sy~ CB     SWH    UP     B           45 202411_S~ 2024   11   12
## # i 11 more variables: Time <chr>, Time_Zone <chr>, npoc_mgL <dbl>,
## # npoc_uM <dbl>, npoc_flag <chr>, tdn_mgL <dbl>, tdn_uM <dbl>,
## # tdn_flag <chr>, Analysis runtime <chr>, Run notes <chr>, Field notes <chr>

```

#end