

Comparison of chromatographic stationary phases using Bayesian-based multilevel modeling

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1 Introduction

In this work we applied the previously developed Bayesian multilevel framework Kubik, Kaliszan, and Wiczling (2018) to characterize chromatographic gradient retention time datasets collected using a multicomponent mixtures of analytes, five stationary phases, and a wide range of chromatographic conditions (pH, organic modifier, temperature, gradient program). Such datasets carry much information about chromatographic retention that, if extracted, can provide useful predictive information, i.e. a detailed multidimensional characterization of chromatographic stationary phases and ability to predict retention (along with uncertainty) based on various number of preliminary experiments (e.g. to predict retention time for a set of analytes given no, or several measurements collected using a different stationary phase).

In this case study, we compared five RP-HPLC stationary phases (XBridge Shield RP18, XTerra MS C18, XBridge Phenyl, XBridge C8, Xterra MS C8) based on LC-MS/TOF data.

2 Experimental design

The data was collected using a mixture of 300 analytes and 84 gradient liquid chromatography experiments for each column. The experiments differed in gradient duration (30, 90, and 270 min) and pH of the mobile phase (from 2.5 to 10.5). Experiments were conducted in MeOH or ACN as organic modifiers and at two temperatures (25^0C and 35^0C).

The molecular structure of the analytes was converted from SMILE format to MDL mol format using OpenBabel. The input molecules were then analyzed for the presence of approximately 204 functional groups and structural elements using Checkmol (version 0.5b N. Haider, University of Vienna, 2003-2018). Functional groups that were not present on any analyte and functional groups merging other simpler functional groups were excluded from the analysis. The lipophilicity ($\log P$), dissociation constant ($pK_{a,1}$) were added to the dataset. They were calculated using ACD/Labs program based on the structures of analytes generated from smiles strings.

3 Setup

The packages we will use are listed below.

```
library(pracma)
library(dplyr)
library(ggplot2)
require(gridExtra)
```

```

library(cmdstanr)
library(knitr)
library(reshape2)
library(bayesplot)
library(posterior)
library(GGally)
library(kableExtra)

set.seed(10271998)

```

4 Data

Data can be accessed via github (data folder) or osf.io repositories Kubik et al. (2022b).

4.1 Prepare the data

We will begin with loading and merging all the required datasets:

```

data1 = read.csv('data/1-X_Bridge_Shield_C18_5cm.csv')
data2 = read.csv('data/1-XTerra_MS_C18.csv')
data3 = read.csv('data/1-X_Bridge_Phenyl.csv')
data4 = read.csv('data/1-XBridge_C8.csv')
data5 = read.csv('data/1-Xterra_C8.csv')

dataNames = read.csv('data/4-compounds-names.csv')
dataACD = read.csv('data/2-ACD-pKas-logP.csv')
dataACD$R = rowSums(dataACD[,3:7]<14) # No of dissociation steps

functional_groups = read.csv('data/6-checkmol-functional-groups.csv')
functional_groups_names = read.csv('data/Legend-checkmol-functional-group-names.csv')

# merge the data:

data1['Column'] <- 1
data2['Column'] <- 2
data3['Column'] <- 3
data4['Column'] <- 4
data5['Column'] <- 5

```

```

data1['ColumnName'] <- 'XBridge Shield RP18'
data2['ColumnName'] <- 'XTerra MS C18'
data3['ColumnName'] <- 'XBridge Phenyl'
data4['ColumnName'] <- 'XBridge C8'
data5['ColumnName'] <- 'Xterra MS C8'

data = rbind(data1, data2, data3, data4, data5)

data$Mod  = as.character(data$Mod)
data$Mod2 = ifelse(data$Mod=="MeOH",1,2)

data<-data %>%
  left_join(select(dataACD,METID,R))

rm(data1, data2, data3, data4, data5)

```

The data requires some cleaning:

1. we remove measurements with low score, analytes with less than 42 measurements per column, analytes with less than 210 measurements out of total 420, and use only measurements with the highest score (if several are present)

```

data <- data %>%
  subset(Score>95) %>%
  group_by(METID,EXPID,Column) %>%
  slice(which.max(Score)) %>%
  group_by(METID,Column) %>%
  add_count() %>%
  subset(n>42) %>%
  group_by(METID) %>%
  add_count() %>%
  subset(nn>210) %>%
  select(-n,-nn) %>%
  ungroup()

```

2. we remove some outlying measurements (mismatch between literature pKa and observed data)

```

data <- data %>%
  subset(METID!=72) %>% # dilevalol is repeated in the dataset
  subset(!(tg==270 & Temp==25 & Mod2==1 & pH==5)) %>%

```

```

subset(!(tg==270 & Temp==25 & Mod2==1 & pH==9)) %>%
subset(R<=2) %>%
subset(!(METID %in% c(11,91,154,204,227,274,102,235,131,158,257,138,281,248,110,135,78,
select(-R)

```

3. we also prepare available predictors: pKa, logP and functional_groups

```

pKaslit = dataACD[,3:7]           # pKa values as predicted by ACD
pKasliterror = dataACD[,25:29]    # pKa error as predicted by ACD
chargesA = abs(dataACD[,13:18])   # number of ionized groups (anions)
chargesB = abs(dataACD[,19:24])   # number of ionized groups (cations)
charges = chargesA+chargesB      # absolute charge
groupsA = (chargesA[,2:5] - chargesA[,1:4])      # acidic group
groupsB = -(chargesB[,2:5] - chargesB[,1:4])      # basic group
R = dataACD$R                   # number of dissociation steps
groups = groupsB-groupsA
logPobs = dataACD$logP

functional_groups=functional_groups[,2:ncol(functional_groups)] 

# combine nr of carboxylic acid and carboxylic acid salt functional groups
# heterocyclic compounds with more than 6 heterocycles are treated as if they have six
functional_groups[,76]=functional_groups[,76]+functional_groups[,77]
functional_groups[which(functional_groups[,202]>5.5),202] = 6;

# exclude some functional groups
idx_included <- c(4,5,9,11,14,18,19,20,21,24,26,29,30,31,32,33,34,38,39,40,41,45,49,50,52,
functional_groups_names <- functional_groups_names[idx_included,]
functional_groups <- functional_groups[,idx_included]

rm(idx_included)

```

4. finally, we select analytes with <=2 dissociation steps for the analysis

```

maxR <- 2
idx <- which(dataACD$METID %in% data$METID)
# max two dissociation steps
pKaslit <- pKaslit[idx,1:maxR]           # pKa values as predicted by ACD
pKasliterror <- pKasliterror[idx,1:maxR]  # pKa error as predicted by ACD
chargesA <- chargesA[idx,1:(maxR+1)]     # number of ionized groups (anions)
chargesB <- chargesB[idx,1:(maxR+1)]     # number of ionized groups (cations)

```

```

charges <- charges[idx,1:(maxR+1)]           # absolute charge
groupsA <- groupsA[idx,1:maxR]                # acidic group
groupsB <- groupsB[idx,1:maxR]                # basic group
R <- R[idx]                                    # number of dissociation steps
logPobs <- logPobs[idx]                      # logP
nrfungroups=functional_groups[idx,]
totalnrgroups <- summarise_each(nrfungroups, funs(sum))

# remove functional groups not present in the dataset:
nrfungroups <- nrfungroups[,which(totalnrgroups!=0)]
functionalgroupsnames<- functional_groups_names[which(totalnrgroups!=0),]
totalnrgroups <- summarise_each(nrfungroups, funs(sum))
K <- ncol(nrfungroups)

# identify acidic groups
idxGroupsA <-which(groupsA!=0,arr.ind = T)
nGroupsA <- nrow(idxGroupsA)
pKaslitA<-pKaslit[idxGroupsA]

# identify basic groups
idxGroupsB <-which(groupsB!=0,arr.ind = T)
nGroupsB <- nrow(idxGroupsB)
pKaslitB<-pKaslit[idxGroupsB]

# groups dissociated in the whole pH range

idxB = which(chargesB[,3]==1)
idxA = which(chargesA[,1]==1)

nObs <- length(data$METID)
nAnalytes <- length(unique(data$METID))
nColumns<-length(unique(data$Column))

# variables used later to annotate graphs legend
temp.labs <- c("25\u00b0C","35\u00b0C")
names(temp.labs) <- c('25','35')

mod.labs <- c("ACN","MeOH")
names(mod.labs) <- c('2','1')

col.labs <- c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xter

```

```

names(col.labs) <- c('1','2','3','4','5')

diss.labs <- c("r=1","r=2","r=3")
names(diss.labs) <- c('1','2','3')

```

4.2 Exploratory data analysis

During the exploratory data analysis phase, we create a series of plots to better understand our data.

4.2.1 Plots

The following graphs present retention time profiles for 6 analytes:

1. acridine (monoprotic acid)
2. baclofen (zwitterion: acidic and basic group)
3. hydrocortison (neutral)
4. pioglitazone (zwitterion: basic and acidic group)
5. quinine (diprotic: 2 basic groups),
6. tolbutamide (monoprotic base).

The vertical lines show ACD-based pKa values (solid line: acidic group; dotted line: basic group)

```

analyte_ID_sample <- c(9,17,33,58,140,180)
#analyte_ID_sample <-unique(data$METID) # if all

idx <- which(unique(data$METID) %in% analyte_ID_sample)
groups <- groupsB-groupsA

for(i in 1:length(analyte_ID_sample)){
  p <- ggplot(data[which(data$METID %in% analyte_ID_sample[i]),])+ 
    geom_jitter(aes(x = pHs, y = RT, color = as.factor(tg), shape=as.factor(ColumnName)))+
    facet_grid(Temp~Mod2, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+ 
    geom_vline(xintercept = pKaslit[idx[i],1],linetype=groups[idx[i],1]+2)+ 
    geom_vline(xintercept = pKaslit[idx[i],2],linetype=groups[idx[i],2]+2)+ 
    labs(title=paste(dataNames>Name[analyte_ID_sample[i]]), 
         x ="pH", y = "Retention time, min",
         color = "tg, min",

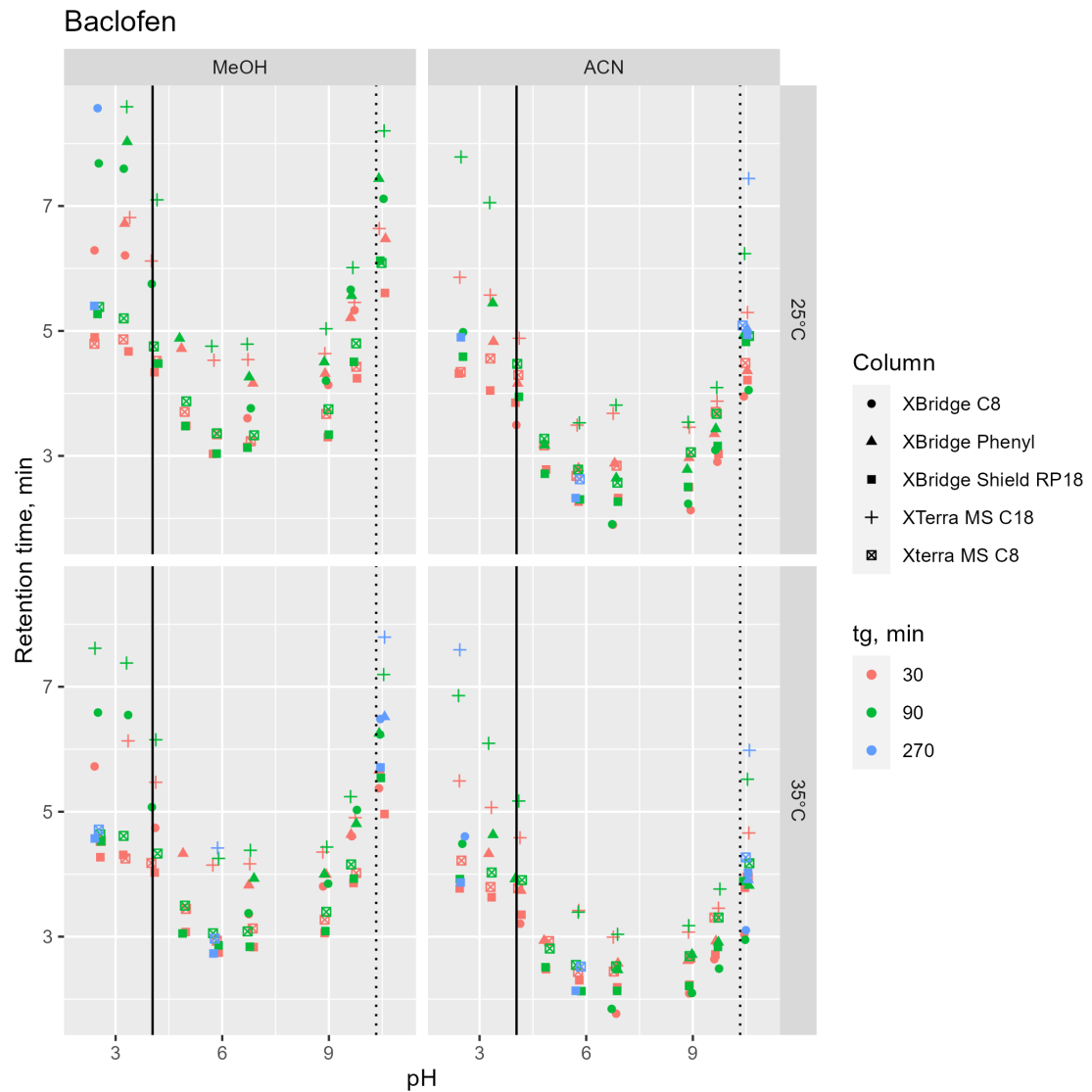
```

```

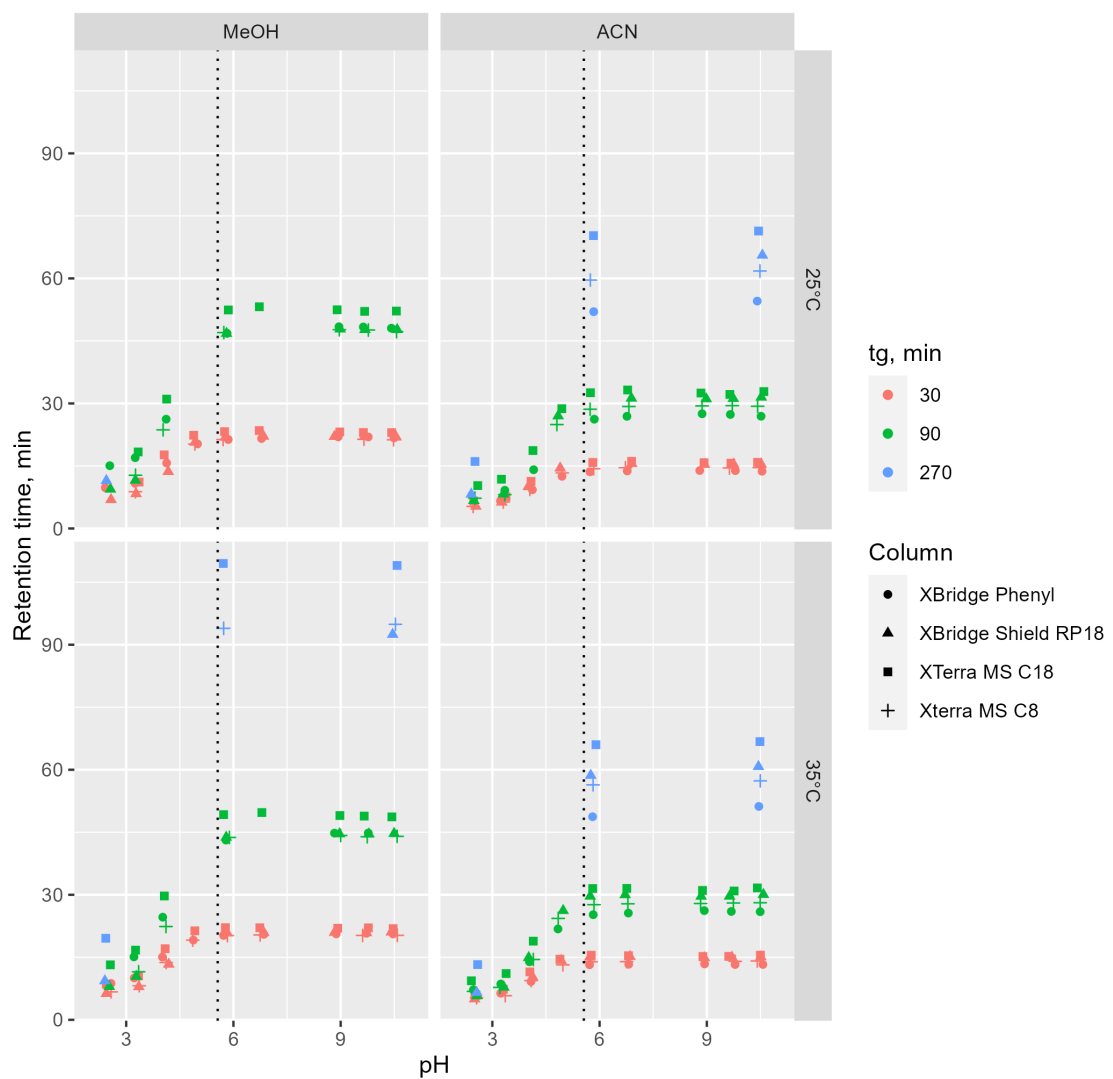
shape = 'Column')+
xlim(2,11)
print(p)

ggsave(paste0("figures\\rawdata\\", paste(dataNames>Name[analyte_ID_sample[i]]), ".png"),
}

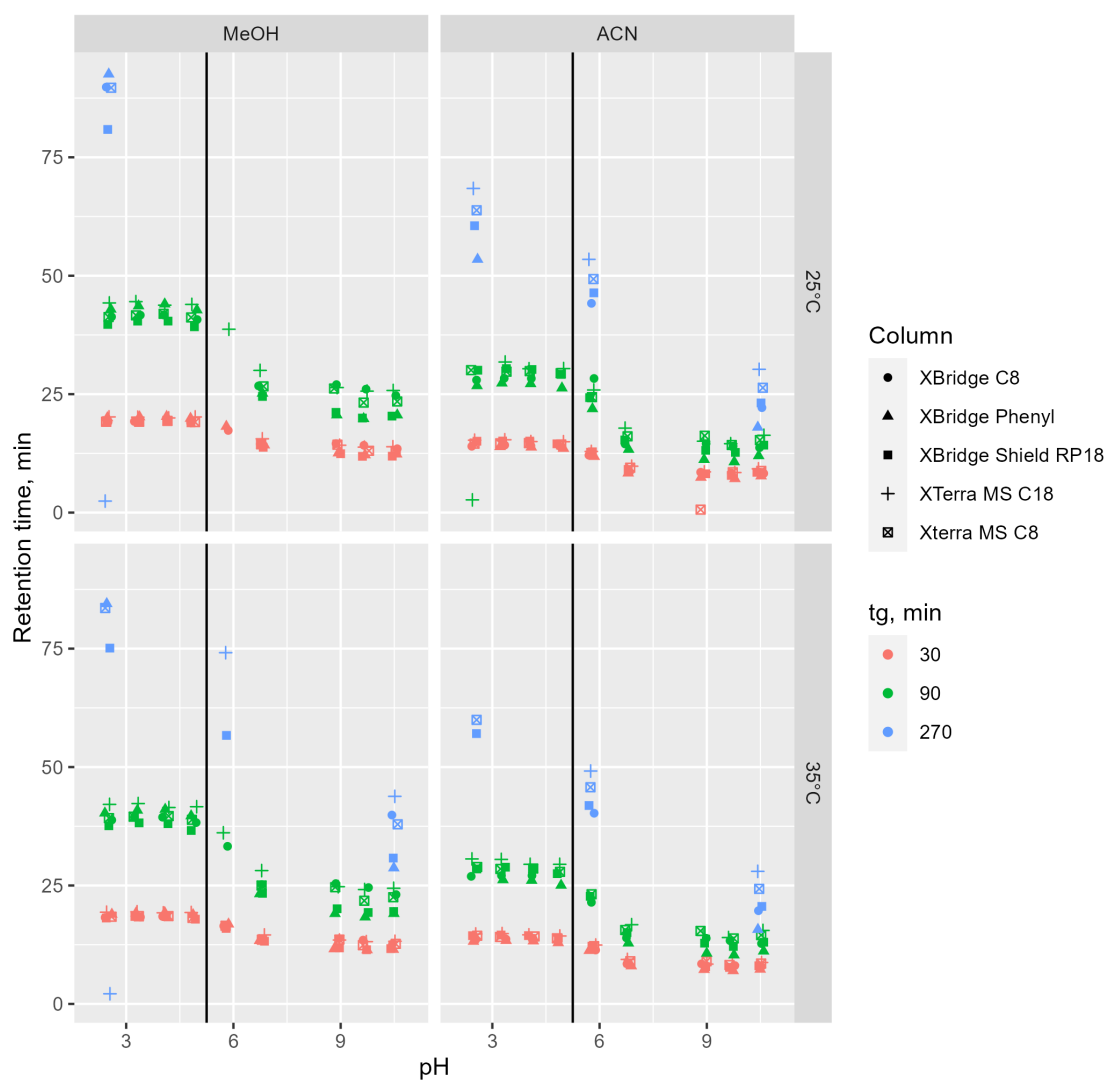
```



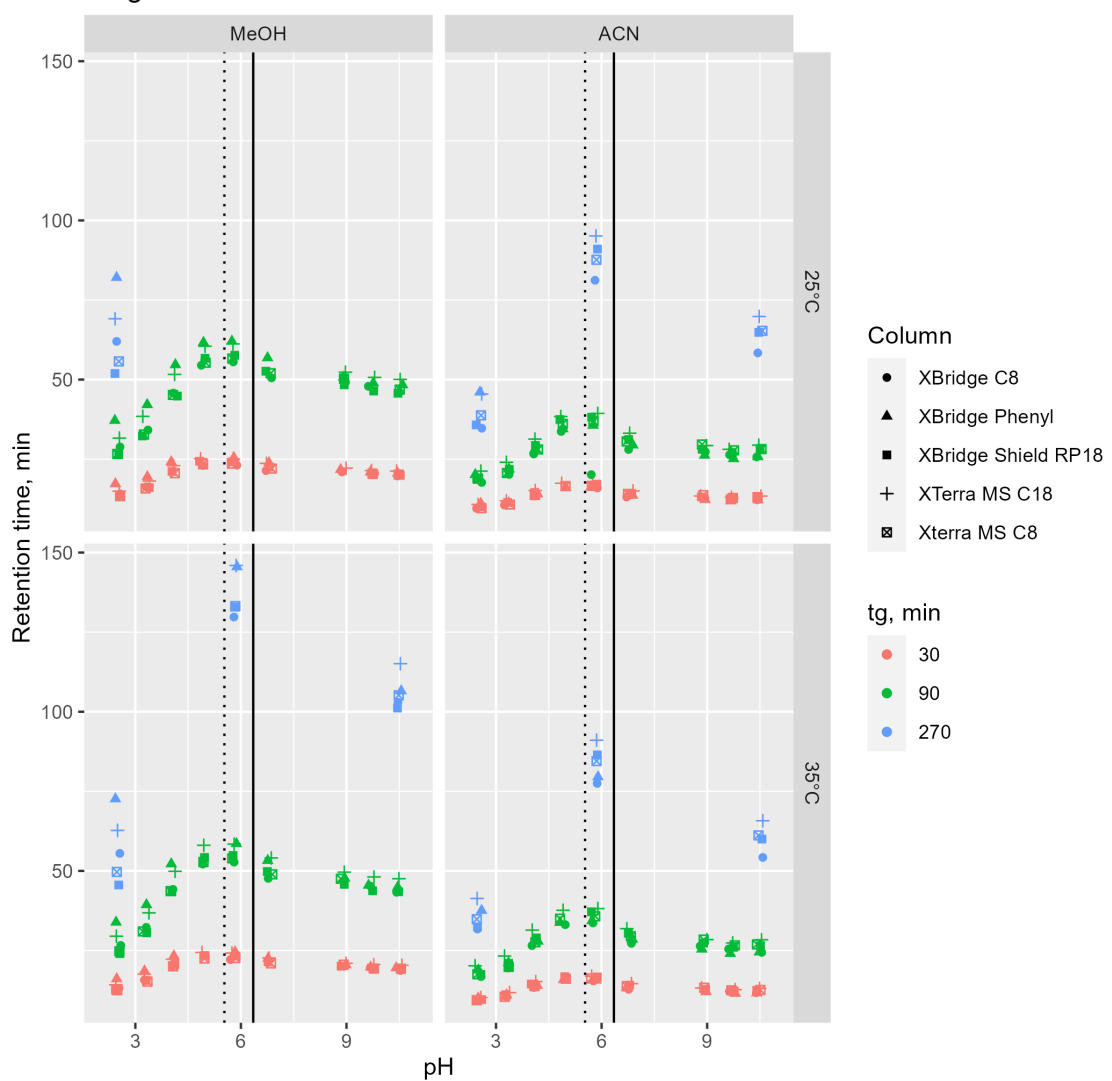
Acridine

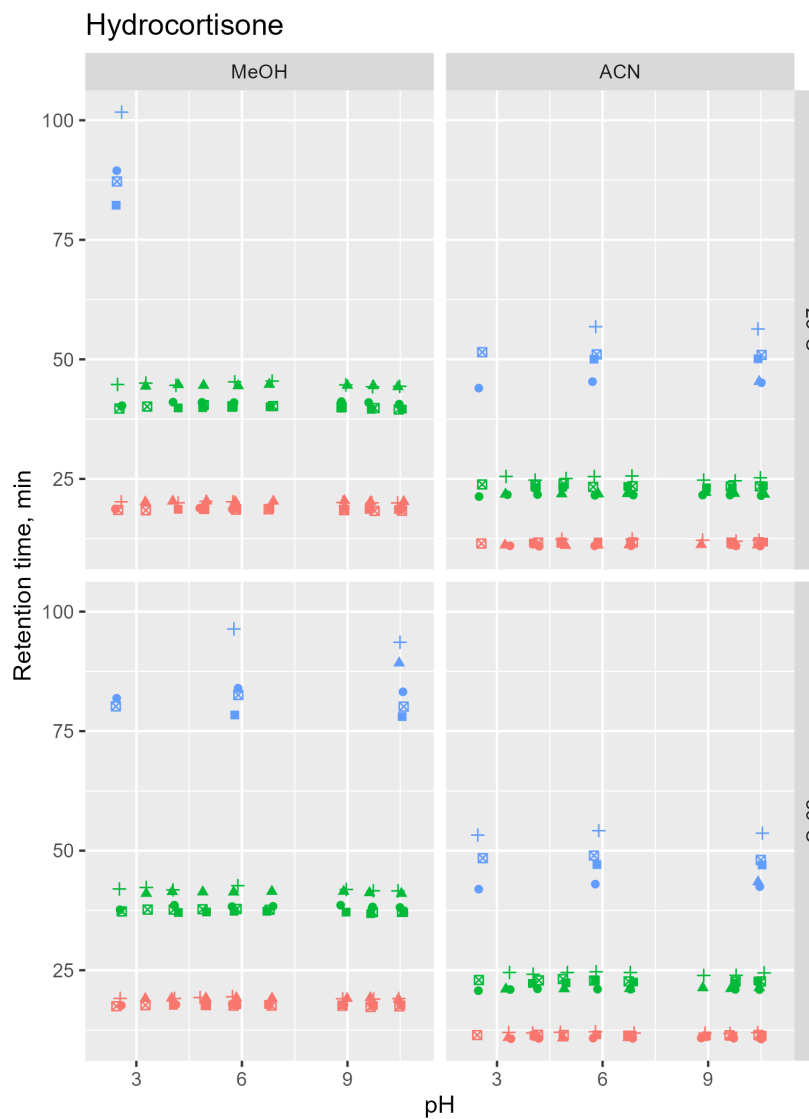


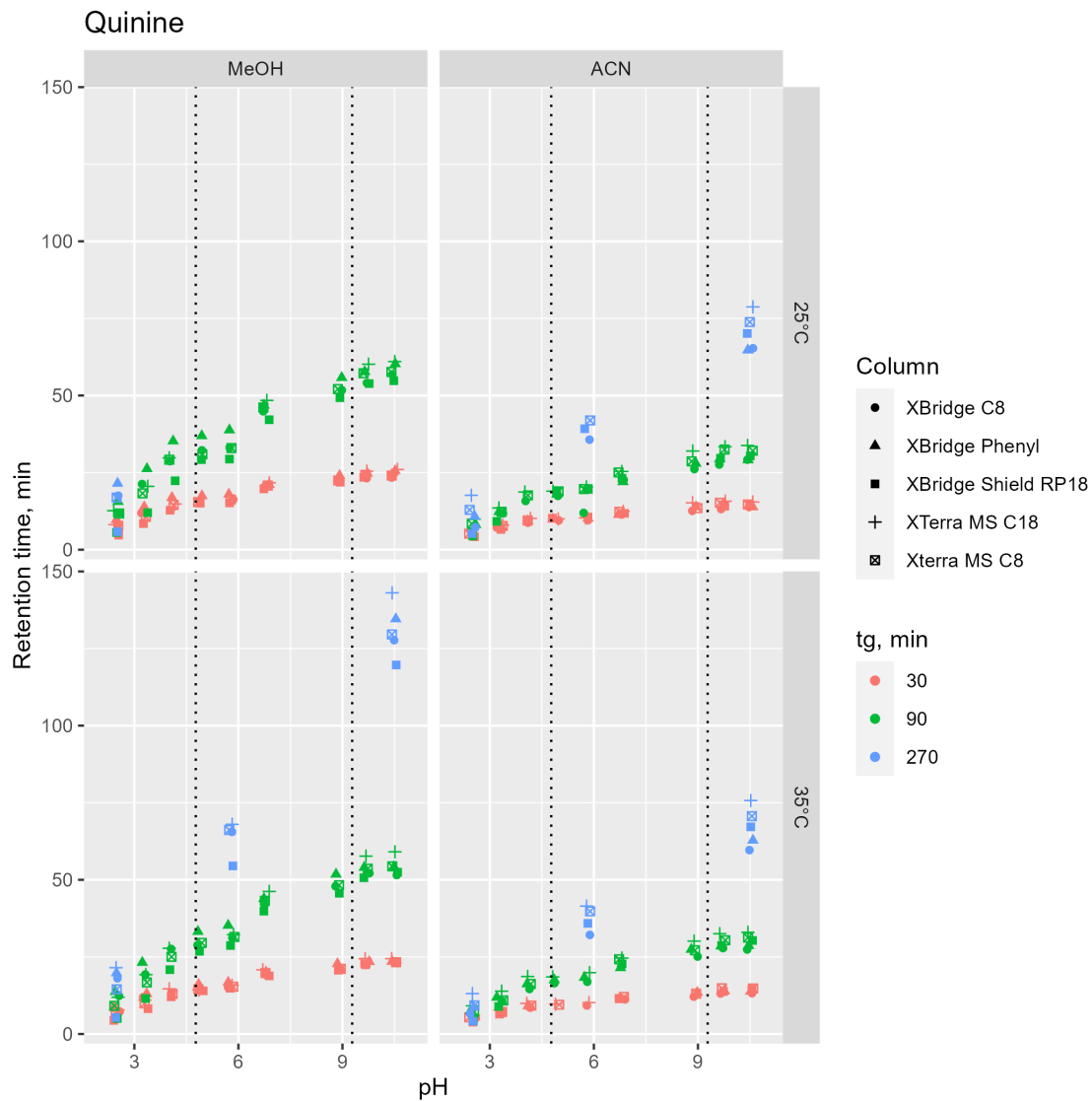
Tolbutamide



Pioglitazone







4.2.2 Summary of the data

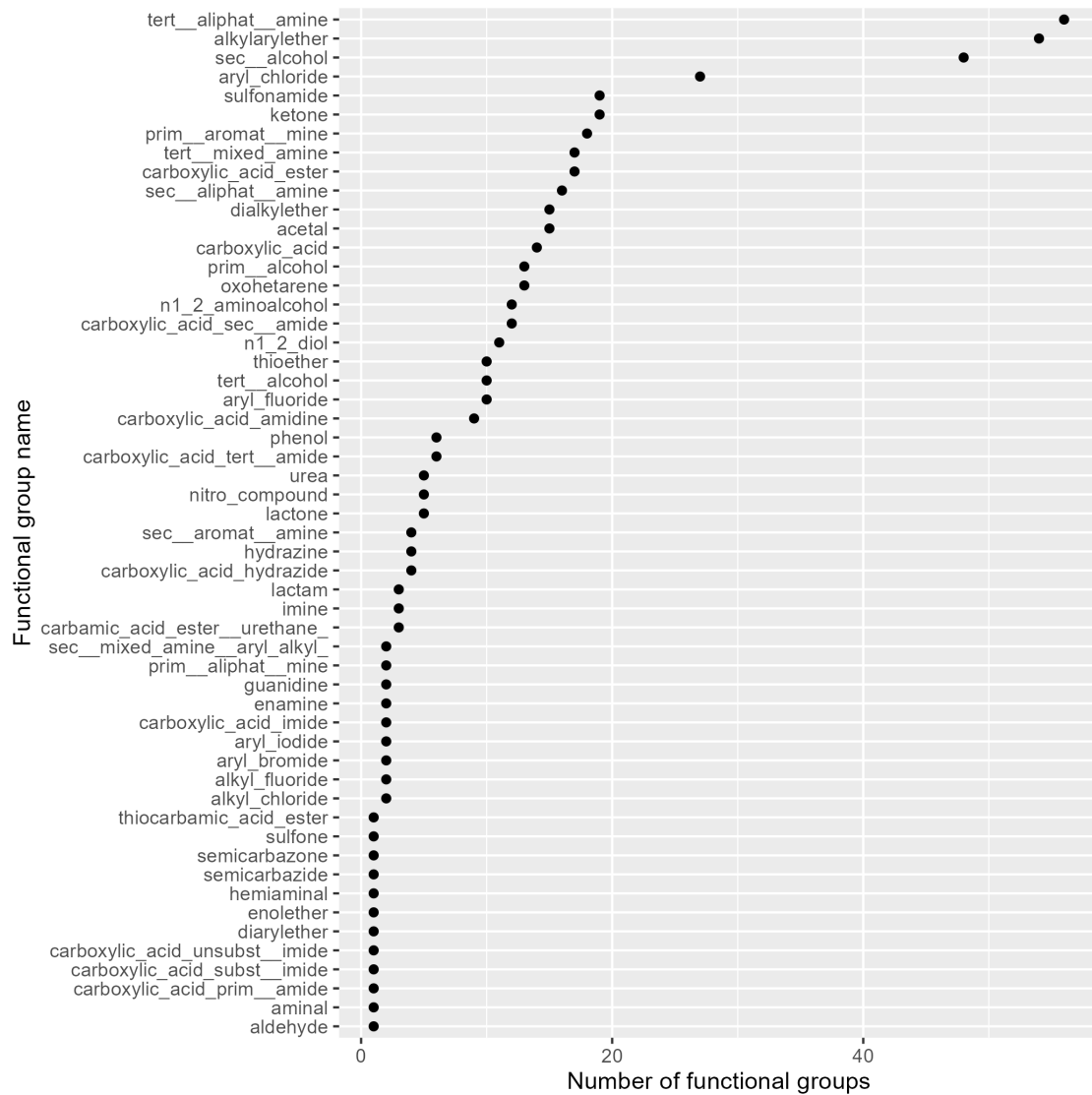
1. Number of identified analytes: 141
2. Number of observations: 51530
3. $\log P$: 2.5 ± 1.82
4. Number of analytes with 0, 1 and 2 dissociation steps: 14, 88, 39
5. Number of acidic and basic groups: 46 and 120
6. Number of functional groups across all analytes included in the analysis:

```

p<-totalnrgroups %>% tidyrr::gather("Name", "count", 1:54) %>%
  ggplot(., aes(x=count, y=reorder(Name, count)))+
  geom_point()+
  labs(x="Number of functional groups", y="Functional group name")

```

```
print(p)
```



```
ggsave(paste0("figures\\predictors\\", "functionalgroups", ".png"), plot=p, width = 20, height = 10)
```

5 Methods

5.1 Model

In this work $z = 1..51530$ denotes observation, $i=1..141$ denotes analyte, $col=1..5$ denotes column, $m=1..2$ denotes organic modifier and $r=1..R[i]$ denotes dissociation step for i -th analyte. The observed retention times ($t_{Robs,z}$) were described using the following model:

$$t_{Robs,z} \sim student_t(\nu, t_{R,z}, \sigma_{col[z],i[z]})$$

where z denotes z -th observation and $student_t$ denotes the Student's t-distribution with the mean given by the predicted retention time $t_{R,z}$, scale $\sigma_{i,col}$ (analyte and column-specific), and normality parameter ν (set to 3).

Gradient retention time $t_{R,z}$ was calculated utilizing the well-known integral equation:

$$\int_0^{t_{R,z}-t_{0,z}-t_e} \frac{dt}{t_{0,z} \cdot ki_z(t)} = 1,$$

where $ki_z(t)$ denotes instantaneous isocratic retention factor corresponding to the mobile phase composition at time t at column inlet for analyte and conditions corresponding to the z -th observation, $t_{0,z}$ denotes column hold-up (dead) time and t_e denotes extra column-time. The numerical solution of this integral equation was carried out using method of steps with 4 and 10 steps for methanol and acetonitrile gradients using method proposed by Nikitas et al. (Nikitas and Pappa-Louisi 2002) The following function described the relationship between the isocratic retention factor and pH for an i th-analyte with $R[i]$ dissociation steps and $R[i] + 1$ forms.

$$ki_z(t) = \frac{k_{z,i[z],1}(t) + \sum_{r=1}^{R[i]} k_{z,i[z],r+1}(t) \cdot 10^{r \cdot pH_z(t) - \sum_{r=1}^{R[i]} pKa_{z,i[z],r}(t)}}{1 + \sum_{r=1}^R 10^{r \cdot pH_z(t) - \sum_{r=1}^{R[i]} pKa_{z,i[z],r}(t)}}$$

$\log(k_{z,i,r})$ was assumed to depend on the organic modifier content based on the Neue equation, on temperature assuming linear equation, and on the pH of the mobiles phase (for ionized forms of analytes).

$$\begin{aligned} \log(k_{z,i,r}(t)) &= \log kw_{col[z],i,r} - \frac{S1_{m[z],col[z],i,r} \cdot (1 + S2_{m[z]}) \cdot \varphi_z(t)}{1 + S2_{m[z]} \cdot \varphi_z(t)} + \dots \\ &apH_{col[z],m[z],i,r} \cdot (pH_z(t) - 7) + dlogkT_{col[z],i} \cdot (T_z - 25)/10 \end{aligned}$$

where $\log k_{w, col, i, r}$ represents logarithm of retention factors extrapolated to 0% of organic modifier content for column col , i -th analyte and r -th analyte form; $S1_{i, m, col, r}$ and $S2_m$ are the slopes in the Neue equation for column c , modifier m , i -th analyte and r -th analyte form. In this parametrization of the Neue equation, $S1$ reflects the difference between logarithm of retention factors corresponding to water (0% of organic modifier content) and MeOH or ACN (100% of organic modifier content) as eluents. $d\log k_{T, col, i}$ denotes the change in $\log k_w$ due to the increase in temperature by $10^\circ C$. $apH_{col, m, i, r}$ denotes the pH effects on $\log k_w$;

Further a linear relationship between pKa values and the organic modifier content was assumed:

$$pKa_{z, i, r}(t) = pKa_{w, i, r} + \alpha_{m[z], i, r} \cdot \varphi_z(t)$$

where $pKa_{z, i, r}(t)$ denotes dissociation constant of an i -th analyte and r -th dissociation step form and chromatographic conditions corresponding the z -th observation, $pKa_{w, i, r}$ denotes aqueous pKa , and $\alpha_{m, i, r}$ denotes the slope for m -th modifier, i -th analyte and r -th form. The linear relationships is generally valid for $\varphi < 0.8$.

The relationship between pH and the organic modifier content for various combinations of organic modifier and buffer was experimentally determined prior to the chromatographic analysis. The obtained relationships was then described using quadratic equations for each nominal pH, temperature and organic modifier:

$$pH_z(t) = pH_{o, z} + \alpha_{1, z} \cdot \varphi_z(t) + \alpha_{2, z} \cdot \varphi_z(t)^2$$

First, individual values of $\log k_w$, $S1$ are defined for the neutral form of an analyte in MeOH for the Xbridge Shield RP18 column (denoted as $\log k_w N_i$ and $S1mN_i$). The effect of ACN was described as $(dS1N_i)$, the effect of column ($c = 1..4$) was described by $c\log k_w N_{c, i}$, $cS1mN_{c, i}$, and $cdS1N_{c, i}$). The individual parameters for the neutral forms were described using the following equations:

$$\begin{aligned}
\begin{bmatrix} \log kw N_i \\ S1m N_i \end{bmatrix} &\sim MVN\left(\begin{bmatrix} \hat{\log kw} + \beta_1 \cdot (\log P_i - 2.2) \\ \hat{S1m} + \beta_2 \cdot (\log P_i - 2.2) \end{bmatrix}, \Omega\right) \\
dS1N_i &\sim N(d\hat{S1}, \omega_3) \\
\Omega &= diag(\omega_{1:2}) \cdot \begin{bmatrix} 1 & \rho \\ \rho & 1 \end{bmatrix} \cdot diag(\omega_{1:2}) \\
\begin{bmatrix} c\log kw N_{1,i} \\ c\log kw N_{2,i} \\ c\log kw N_{3,i} \\ c\log kw N_{4,i} \end{bmatrix} &\sim MVN\left(\begin{bmatrix} \hat{c\log kw}_1 + c\beta_{1,1} \cdot (\log P_i - 2.2) \\ \hat{c\log kw}_2 + c\beta_{2,1} \cdot (\log P_i - 2.2) \\ \hat{c\log kw}_3 + c\beta_{3,1} \cdot (\log P_i - 2.2) \\ \hat{c\log kw}_4 + c\beta_{4,1} \cdot (\log P_i - 2.2) \end{bmatrix}, c\Omega\right) \\
c\Omega &= diag(c\omega) \cdot \begin{bmatrix} 1 & c\rho_{12} & c\rho_{13} & c\rho_{14} \\ c\rho_{21} & 1 & c\rho_{23} & c\rho_{24} \\ c\rho_{31} & c\rho_{32} & 1 & c\rho_{34} \\ c\rho_{41} & c\rho_{42} & c\rho_{43} & 1 \end{bmatrix} \cdot diag(c\omega) \\
cS1m N_{c,i} &\sim N(c\hat{S1m} + c\beta_{c,2} \cdot (\log P_i - 2.2), c\omega_{c,2}) \text{ for } c=1\dots4 \\
cdS1N_{c,i} &\sim N(cd\hat{S1}_c, c\omega_{c,3}) \text{ for } c=1\dots4 \\
d\log kT_{c,i} &\sim N(d\hat{\log kT}_c, \omega_{T,c}) \text{ for } c=1\dots4
\end{aligned}$$

were MVN denotes the multivariate normal distribution; $\log kw$, $S1m$, $d\hat{S1}$ are the mean values of individual chromatographic parameters that correspond to a typical neutral analyte with $\log P = 2.2$ at $25^\circ C$ on Xbridge Shield RP18 stationary phase. β s are regression coefficients between the individual chromatographic parameters and the $\log P_i$. ω denotes the standard deviation for between analyte variability (BAV). $d\log kT$ denotes the effect of temperature for a typical analyte and ω_T the standard deviation for between analyte variability for temperature effects. Similar set of equations was used for column effects. Here c denoted the column effect (4 differences with respect to the reference column).

The difference in retention between the ionized form of an analyte and the neutral form of an analyte was separately estimated for acids and bases for $\log kw$, $S1m$, $dS1$ parameters. Similar set of equations was used for column effects.

$$\begin{aligned}
d\log k w A_a &\sim N(d\hat{\log k w}_1, \kappa_1), \\
d\log k w B_b &\sim N(d\hat{\log k w}_2, \kappa_1), \\
dS1mA_a &\sim N(d\hat{S1m}_1, \kappa_2), \\
dS1mB_b &\sim N(d\hat{S1m}_2, \kappa_2), \\
ddS1A_a &\sim N(dd\hat{S1}_1, \kappa_3), \\
ddS1B_b &\sim N(dd\hat{S1}_2, \kappa_3), \\
cd\log k w A_{c,a} &\sim N(cd\log \hat{k w}_{c,1}, c\kappa_{c,1}) \text{ for } c=1\dots 4, \\
cd\log k w B_{c,b} &\sim N(cd\log \hat{k w}_{c,2}, c\kappa_{c,1}) \text{ for } c=1\dots 4, \\
cdS1mA_{c,a} &\sim N(cd\hat{S1m}_{c,1}, c\kappa_{c,2}) \text{ for } c=1\dots 4, \\
cdS1mB_{c,b} &\sim N(cd\hat{S1m}_{c,2}, c\kappa_{c,2}) \text{ for } c=1\dots 4, \\
cddS1A_{c,a} &\sim N(cdd\hat{S1}_{c,1}, c\kappa_{c,3}) \text{ for } c=1\dots 4, \\
cddS1B_{c,b} &\sim N(cdd\hat{S1}_{c,2}, c\kappa_{c,3}) \text{ for } c=1\dots 4.
\end{aligned}$$

where $a=1..46$ and $b=1..120$ denote the indexes of acidic and basic groups.

Similarly pK_a and α parameters were described separately for acids and bases:

$$\begin{aligned}
pKawA_a &\sim N(pKawA_{lit,a}, \tau_1), \\
pKawB_b &\sim N(pKawB_{lit,b}, \tau_1), \\
\alpha mA_a &\sim N(\alpha \hat{m}_1, \tau_2), \\
\alpha mB_b &\sim N(\alpha \hat{m}_2, \tau_2), \\
d\alpha A_a &\sim N(d\hat{\alpha}_1, \tau_3), \\
d\alpha B_b &\sim N(d\hat{\alpha}_2, \tau_3).
\end{aligned}$$

Further, we created the matrices containing the value of parameters for i -th analyte, col -th column, m -th modifier and r -th dissociation step based on the value of neutral form and effects of column, organic modifier, and dissociation. This transformation was denoted as $f(\cdot)$. The exact procedure can be found in the model code displayed later.

$$\begin{aligned}
logkw_{col,i,r} &= f(logkwN_i, clogkwN_{c,i}, dlogkwA_a, cdlogkwA_{c,a}, dlogkwB_b, cdlogkwB_{c,b}, \dots) \\
S1_{m,col,i,r} &= f(S1mN_i, cS1mN_{c,i}, dS1mA_a, cdS1mA_{c,a}, S1mB_b, cdS1mB_{c,b}, \dots) \\
dS1N_i, cdS1N_{c,i}, ddS1A_a, cddS1A_{c,a}, ddS1B_b, cddS1B_{c,b}, \dots) \\
apH_{m,col,i,r} &= f(ap\hat{H}_1, cap\hat{H}_{c,1}, ap\hat{H}_2, cap\hat{H}_{c,2}, \dots) \\
S2_m &= 10^{f(log\hat{S}2m, dlog\hat{S}2, \dots)} \\
pKaw_{i,r} &= f(pKawA_a, pKawB_b, \dots) \\
\alpha_{m,i,r} &= f(\alpha mA_a, d\alpha A_a, \alpha mB_b, d\alpha B_b, \dots)
\end{aligned}$$

Residual error model assumes different parameters for each column and analyte:

$$\begin{aligned} \log(\sigma_{col,i}) &= f(\log\sigma_i, c\log\sigma_{c,i}) \\ \log\sigma_i &\sim N(\log(m\sigma), s\sigma) \\ c\log\sigma_{c,i} &\sim N(c\log m\sigma_c, cs\sigma_c) \text{ for } c=1\dots 4, \end{aligned}$$

The detailed description of parameters and used priors is provided in the following table (BAV denotes between analyte variability):

```
# to properly render in quarto: https://github.com/quarto-dev/quarto-cli/issues/3340

Name = c('$\\hat{logkw}$',
          '$\\hat{S1m}$',
          '$\\hat{dS1}$',
          '$\\beta_1$',
          '$\\beta_2$',
          '$\\hat{dlogkw_1}$',
          '$\\hat{dlogkw_2}$',
          '$\\hat{dS1m_1}$',
          '$\\hat{dS1m_2}$',
          '$\\hat{ddS1_1}$',
          '$\\hat{ddS1_2}$',
          '$\\hat{apH_1}$',
          '$\\hat{apH_2}$',
          '$\\hat{dlogkT}$',
          '$\\omega_1$',
          '$\\omega_2$',
          '$\\omega_3$',
          '$\\rho$',
          '$\\omega_T$',
```

```

'$\\kappa_1$',
'$\\kappa_2$',
'$\\kappa_3$',
'$\\hat{clogkw_c}$',
'$\\hat{cS1m_c}$',
'$\\hat{cdS1_c}$',
'$c\\beta_{c,1}$',
'$c\\beta_{c,2}$',
'$\\hat{cdlogkw_{c,1}}$',
'$\\hat{cdlogkw_{c,2}}$',
'$\\hat{cdS1m_{c,1}}$',
'$\\hat{cdS1m_{c,2}}$',
'$\\hat{cdS1_{c,1}}$',
'$\\hat{cdS1_{c,2}}$',
'$\\hat{cddS1_{c,1}}$',
'$\\hat{cddS1_{c,2}}$',
'$\\hat{cdlogkT_c}$',
'$capH_{c,1}$',
'$capH_{c,2}$',
'$c\\omega_{c,1}$',
'$c\\omega_{c,2}$',
'$c\\omega_{c,3}$',
'$c\\kappa_{c,1}$',
'$c\\kappa_{c,2}$',
'$c\\kappa_{c,3}$',
'$c\\omega_{T,c}$',
'$\\hat{logS2m}$',
'$\\hat{dlogS2}$',
'$\\hat{\\alpha_m_1}$',
'$\\hat{\\alpha_m_2}$',
'$\\hat{d\\alpha_1}$',
'$\\hat{d\\alpha_2}$',
'$\\tau_1$',
'$\\tau_2$',
'$\\tau_3$',
'$m\\sigma$',
'$s\\sigma$',
'$clogmsigma_c$',
'$c\\sigma_c$')

Namecode = c('logkwHat',
           'S1mHat',
           'dS1Hat',

```

```
'beta[1]',
'beta[2]',
'dlogkwHat[1]',
'dlogkwHat[2]',
'dS1mHat[1]',
'dS1mHat[2]',
'ddS1Hat[1]',
'ddS1Hat[2]',
'apH[1]',
'apH[2]',
'dlogkTHat',
'omega[1]',
'omega[2]',
'omega[3]',
'rho[2,1]',
'omegaT',
'kappa[1]',
'kappa[2]',
'kappa[3]',
'clogkwHat[c]',
'cS1mHat[c]',
'cdS1Hat[c]',
'cbeta[c,1]',
'cbeta[c,2]',
'cdlogkwHat[c,1]',
'cdlogkwHat[c,2]',
'cdS1mHat[c,1]',
'cdS1mHat[c,2]',
'cddS1Hat[c,1]',
'cddS1Hat[c,2]',
'cdlogkTHat[c]',
'capH[c,1]',
'capH[c,2]',
'comega[c,1]',
'comega[c,2]',
'comega[c,3]',
'ckappa[c,1]',
'ckappa[c,2]',
'ckappa[c,3]',
'comegaT[c]',
'logS2mHat',
```

```

'dlogS2Hat',
'alphamHat[1]',
'alphamHat[2]',
'dalphaHat[1]',
'dalphaHat[2]',
'tau[1]',
'tau[2]',
'tau[3]',
'msigma',
:ssigma',
'clogmsigma[c]',
'cssigma[c]')

Description = c('typical logkw [Neutral]',
  'effect of MeOH on logkw [Neutral]',
  'effect of ACN on S1m [Neutral]',
  'effect of logP on logkw [Neutral]',
  'effect of logP on S1m [Neutral]',
  'effect of dissociation on logkw [Acids]',
  'effect of dissociation on logkw [Bases]',
  'effect of dissociation on S1m [Acids]',
  'effect of dissociation on S1m [Bases]',
  'effect of dissociation on dS1 [Acids]',
  'effect of dissociation on dS1 [Bases]',
  'effect of pH on logkw [Acids]',
  'effect of pH on logkw [Bases]',
  'effect of temperature on logkw',
  'sd of BAV for logkw [Neutral]',
  'sd of BAV for S1 [Neutral]',
  'sd of BAV for dS1 [Neutral]',
  'correlation logkw vs S1 [Neutral]',
  'sd of BAV for dlogkT [Neutral]',
  'sd of BAV for dlogkw [Acids and Bases]',
  'sd of BAV for dS1m [Acids and Bases]',
  'sd of BAV for ddS1 [Acids and Bases]',
  'effect of column c on logkw [Neutral]',
  'effect of column c on S1m [Neutral]',
  'effect of column c on dS1 [Neutral]',
  'effect of column c on beta[1] [Neutral]',
  'effect of column c on beta[2] [Neutral]',
  'effect of column c on dlogkw [Acids]',
```

```

'effect of column c on dlogkw [Bases]',
'effect of column c on dS1m [Acids]',
'effect of column c on dS1m [Bases]',
'effect of column c on ddS1 [Acids]',
'effect of column c on ddS1 [Bases]',
'effect of column c on dlogkwT',
'effect of column c on apH [Acids]',
'effect of column c on apH [Bases]',
'sd of BAV for clogkw [Neutral]',
'sd of BAV for cS1 [Neutral]',
'sd of BAV for cdS1 [Neutral]',
'sd of BAV for cdlogkw [Acids and Bases]',
'sd of BAV for cdS1m [Acids and Bases]',
'sd of BAV for cddS1 [Acids and Bases]',
'sd of BAV for dlogkT',
'typical value of S2m (log10 scale)',
'effect of ACN on logS2m',
'effect of MeOH on pKa [Acids]',
'effect of MeOH on pKa [Bases]',
'effect of ACN on alpham [Acids]',
'effect of ACN on alpham [Bases]',
'sd of BAV for pKalit',
'sd of BAV for alpham',
'sd of BAV for dalpha',
'typical sd of residuals for XBridge',
'sd of BAV of residuals for XBridge',
'effect of column c on msigma (log scale)',
'sd of BAV of residuals for column c'
)

```

```

Priors = c('N(2.2,2)',
          'N(4, 1)',
          'N(1, 1)',
          'N(1, 0.125)',
          'N(0.5, 0.5)',
          'N(-1, 0.125)',
          'N(-1, 0.125)',
          'N(0, 0.5)',
          'N(0, 0.5)',
          'N(0, 0.25)',
```

```
'N(0, 0.25)',  
'N(0, 0.1)',  
'N(0, 0.1)',  
'N(-0.087, 0.022)',  
'N+(0, 2)',  
'N+(0, 2)',  
'N+(0, 2)',  
'LKJCORRN(0.75, 0.125)',  
'N+(0, 0.022)',  
'N+(0, 0.25)',  
'N+(0, 0.25)',  
'N+(0, 0.25)',  
'N(0, 1)',  
'N(0, 0.5)',  
'N(0, 0.5)',  
'N(0, 0.25)',  
'N(0, 0.25)',  
'N(0, 0.0625)',  
'N(0, 0.0625)',  
'N(0, 0.25)',  
'N(0, 0.25)',  
'N(0, 0.125)',  
'N(0, 0.125)',  
'N(0, 0.011)',  
'N(0, 0.05)',  
'N(0, 0.05)',  
'N+(0, 1)',  
'N+(0, 1)',  
'N+(0, 1)',  
'N+(0, 0.125)',  
'N+(0, 0.125)',  
'N+(0, 0.125)',  
'N+(0, 0.011)',  
'N(-0.7, 0.125);',  
'N(1, 0.125);',  
'N(2, 0.25)',  
'N(-1, 0.25)',  
'N(0, 0.125)',  
'N(0, 0.125)',  
'N+(0, 0.25)',  
'N+(0, 0.125)',
```

```

'N+(0, 0.125)',  

'N+(0,1)',  

'N(0,1)',  

'N+(0,125)',  

'N+(0,125)')

```

```

table_of_parameters = data.frame(Name,Namecode,Description,Priors)

```

```

table_of_parameters %>%
kableExtra::kbl(caption = "Description of model paramters", escape = FALSE, longtable=TRUE)
kableExtra::kable_classic(full_width = F, html_font = "Cambria") %>%
kableExtra::kable_styling(latex_options = c("repeat_header")) %>%
kableExtra::pack_rows("XBridge Shield RP18 parameters", 1, 22) %>%
kableExtra::pack_rows("between column differences", 23, 43) %>%
kableExtra::pack_rows("S2", 44, 45) %>%
kableExtra::pack_rows("pKa", 46, 52) %>%
kableExtra::pack_rows("Residuals", 53, 56) %>%
unclass() %>%
cat()

```

Table 1: Description of model parameters

Name	Namecode	Description	Priors
XBridge Shield RP18 parameters			
$\hat{\logkw}$	logkwHat	typical logkw [Neutral]	N(2.2,2)
$\hat{S1m}$	S1mHat	effect of MeOH on logkw [Neutral]	N(4, 1)
$\hat{dS1}$	dS1Hat	effect of ACN on S1m [Neutral]	N(1, 1)
β_1	beta[1]	effect of logP on logkw [Neutral]	N(1, 0.125)
β_2	beta[2]	effect of logP on S1m [Neutral]	N(0.5, 0.5)
$\hat{dlogkw_1}$	dlogkwHat[1]	effect of dissociation on logkw [Acids]	N(-1, 0.125)
$\hat{dlogkw_2}$	dlogkwHat[2]	effect of dissociation on logkw [Bases]	N(-1, 0.125)
$\hat{dS1m_1}$	dS1mHat[1]	effect of dissociation on S1m [Acids]	N(0, 0.5)
$\hat{dS1m_2}$	dS1mHat[2]	effect of dissociation on S1m [Bases]	N(0, 0.5)
$\hat{ddS1_1}$	ddS1Hat[1]	effect of dissociation on dS1 [Acids]	N(0, 0.25)
$\hat{ddS1_2}$	ddS1Hat[2]	effect of dissociation on dS1 [Bases]	N(0, 0.25)
$\hat{apH_1}$	apH[1]	effect of pH on logkw [Acids]	N(0, 0.1)
$\hat{apH_2}$	apH[2]	effect of pH on logkw [Bases]	N(0, 0.1)
\hat{dlogkT}	dlogkTHat	effect of temperature on logkw	N(-0.087, 0.022)
ω_1	omega[1]	sd of BAV for logkw [Neutral]	N+(0, 2)

Table 1: Description of model parameters (*continued*)

Name	Namecode	Description	Priors
ω_2	omega[2]	sd of BAV for S1 [Neutral]	N+(0, 2)
ω_3	omega[3]	sd of BAV for dS1 [Neutral]	N+(0, 2)
ρ	rho[2,1]	correlation logkw vs S1 [Neutral]	LKJCORRN(0.75, 0.125)
ω_T	omegaT	sd of BAV for dlogkT [Neutral]	N+(0, 0.022)
κ_1	kappa[1]	sd of BAV for dlogkw [Acids and Bases]	N+(0, 0.25)
κ_2	kappa[2]	sd of BAV for dS1m [Acids and Bases]	N+(0, 0.25)
κ_3	kappa[3]	sd of BAV for ddS1 [Acids and Bases]	N+(0, 0.25)
between column differences			
$c\hat{\log}kw_c$	clogkwHat[c]	effect of column c on logkw [Neutral]	N(0, 1)
$c\hat{S1m}_c$	cS1mHat[c]	effect of column c on S1m [Neutral]	N(0, 0.5)
$c\hat{dS1}_c$	cdS1Hat[c]	effect of column c on dS1 [Neutral]	N(0, 0.5)
$c\beta_{c,1}$	cbeta[c,1]	effect of column c on beta[1] [Neutral]	N(0, 0.25)
$c\beta_{c,2}$	cbeta[c,2]	effect of column c on beta[2] [Neutral]	N(0, 0.25)
$c\hat{dlogkw}_{c,1}$	cdlogkwHat[c,1]	effect of column c on dlogkw [Acids]	N(0, 0.0625)
$c\hat{dlogkw}_{c,2}$	cdlogkwHat[c,2]	effect of column c on dlogkw [Bases]	N(0, 0.0625)
$c\hat{dS1m}_{c,1}$	cdS1mHat[c,1]	effect of column c on dS1m [Acids]	N(0, 0.25)
$c\hat{dS1m}_{c,2}$	cdS1mHat[c,2]	effect of column c on dS1m [Bases]	N(0, 0.25)
$c\hat{dS1}_{c,1}$	cddS1Hat[c,1]	effect of column c on ddS1 [Acids]	N(0, 0.125)
$c\hat{dS1}_{c,2}$	cddS1Hat[c,2]	effect of column c on ddS1 [Bases]	N(0, 0.125)
$c\hat{dlogkT}_c$	cdlogkTHat[c]	effect of column c on dlogkwT	N(0, 0.011)
$c\hat{apH}_{c,1}$	capH[c,1]	effect of column c on pH [Acids]	N(0, 0.05)
$c\hat{apH}_{c,2}$	capH[c,2]	effect of column c on pH [Bases]	N(0, 0.05)
$c\hat{\omega}_{c,1}$	comega[c,1]	sd of BAV for clogkw [Neutral]	N+(0, 1)
$c\hat{\omega}_{c,2}$	comega[c,2]	sd of BAV for cS1 [Neutral]	N+(0, 1)
$c\hat{\omega}_{c,3}$	comega[c,3]	sd of BAV for cdS1 [Neutral]	N+(0, 1)
$c\hat{\kappa}_{c,1}$	ckappa[c,1]	sd of BAV for cdlogkw [Acids and Bases]	N+(0, 0.125)
$c\hat{\kappa}_{c,2}$	ckappa[c,2]	sd of BAV for cdS1m [Acids and Bases]	N+(0, 0.125)
$c\hat{\kappa}_{c,3}$	ckappa[c,3]	sd of BAV for cddS1 [Acids and Bases]	N+(0, 0.125)
$c\hat{\omega}_{T,c}$	comegaT[c]	sd of BAV for dlogkT	N+(0, 0.011)
S2			
$\log\hat{S2m}$	logS2mHat	typical value of S2m (log10 scale)	N(-0.7, 0.125);
$d\hat{\log}S2$	dlogS2Hat	effect of ACN on logS2m	N(1, 0.125);
pKa			
$\hat{\alpha m}_1$	alphamHat[1]	effect of MeOH on pKa [Acids]	N(2, 0.25)
$\hat{\alpha m}_2$	alphamHat[2]	effect of MeOH on pKa [Bases]	N(-1, 0.25)
$\hat{d\alpha}_1$	dalphaHat[1]	effect of ACN on alpham [Acids]	N(0, 0.125)
$\hat{d\alpha}_2$	dalphaHat[2]	effect of ACN on alpham [Bases]	N(0, 0.125)

Table 1: Description of model parameters (*continued*)

Name	Namecode	Description	Priors
τ_1	tau[1]	sd of BAV for pKalit	$N+(0, 0.25)$
τ_2	tau[2]	sd of BAV for alpham	$N+(0, 0.125)$
τ_3	tau[3]	sd of BAV for dalpha	$N+(0, 0.125)$
Residuals			
$m\sigma$	msigma	typical sd of residuals for XBridge	$N+(0,1)$
$s\sigma$	ssigma	sd of BAV of residuals for XBridge	$N(0,1)$
$c\log m\sigma_c$	clogmsigma[c]	effect of column c on msigma (log scale)	$N+(0,125)$
$c\sigma_c$	cssigma[c]	sd of BAV of residuals for column c	$N+(0,125)$

5.2 Stan

Multilevel modeling was performed in [Stan software](#) linked with R/ [cmdstanr](#). For the inference we used the following settings: number of iterations = 500, warmup = 1000, and number of Markov chains = 8. The `reduce_sum` function, which was selected to accelerate the calculations, works by parallelizing the execution of a single Stan chain across multiple cores. Convergence diagnostics were checked using Gelman–Rubin statistics and trace plots.

5.2.1 Initialize variables and parameters

```
# create Stan data set:

datastruct <- with(data,
  list(nAnalytes=length(unique(data$METID)),
       nModifiers=length(unique(data$Mod2)),
       nColumns=length(unique(data$Column)),
       nObs=length(data$METID),
       analyte=match(data$METID, unique(data$METID)),
       modifier=match(data$Mod2, sort(unique(data$Mod2))),
       column=match(data$Column, unique(data$Column)),
       steps=4*(2-data$Mod2) + 10*(data$Mod2-1),
       hplcparam=cbind(data$tg,data$td,data$to,data$te,data$fio,data$fik,
                        data$alpha1,data$alpha2,(data$Temp-25)/10,data$Colu
       logPobs=logPobs,
       maxR=maxR,
       R=R,
       nGroupsA=nGroupsA,
       nGroupsB=nGroupsB,
```

```

pKaslitA=pKaslitA,
pKaslitB=pKaslitB,
idxGroupsA=idxGroupsA,
idxGroupsB=idxGroupsB,
trobs=data$RT,
run_estimation=1))

# initialize the values for each variable in each chain:

init <- function(){
  list(  logkwHat  = rnorm(1,2.2,2),
         S1mHat    = rnorm(1,4,1),
         dS1Hat    = rnorm(1,1,0.5),
         dlogkwHat = rnorm(2,-1,0.125),
         dS1mHat   = rnorm(2,0,0.5),
         ddS1Hat   = rnorm(2,0,0.25),
         logS2mHat = rnorm(1,-0.7,0.125),
         dlogS2Hat = rnorm(1,1,0.125),
         beta     = rnorm(2,c(1,0.5),c(0.125,0.5)),
         dlogkTHat = rnorm(1,-0.087,0.0022),
         apH      = rnorm(2,0,0.1),
         alphamHat = rnorm(2,c(2,-1),0.25),
         dalphaHat = rnorm(2,0,0.125),
         tau      = c(0.25,0.125,0.125)*exp(rnorm(3, 0, 0.2)),
         omega    = c(1,1,1)*exp(rnorm(3, 0, 0.5)),
         rho      = matrix(c(1, 0.75, 0.75, 1), nrow=2),
         omegaT   = rlnorm(1,log(0.022),0.2),
         kappa    = 0.25* exp(rnorm(3, 0, 0.2)),

         clogkwHat = rnorm(nColumns-1,0,1),
         cS1mHat   = rnorm(nColumns-1,0,0.5),
         cdS1Hat   = rnorm(nColumns-1,0,0.25),
         cdlogkwHat = matrix(rnorm(2*(nColumns-1),0,0.0625),nrow=(nColumns-1)),
         cdS1mHat = matrix(rnorm(2*(nColumns-1),0,0.25),nrow=(nColumns-1)),
         cddS1Hat = matrix(rnorm(2*(nColumns-1),0,0.125),nrow=(nColumns-1)),
         cbeta   = matrix(rnorm(2*(nColumns-1),0,0.25),nrow=(nColumns-1)),
         cdlogkTHat = rnorm(nColumns-1,0,0.011),

         capH    = matrix(rnorm(2*(nColumns-1),0,0.05),nrow=(nColumns-1)),
         comega  = 0.1 * exp(matrix(rnorm(3*(nColumns-1),0,1),nrow=(nColumns-1))),
         ckappa  = 0.1 * exp(matrix(rnorm(3*(nColumns-1),0,0.125),nrow=(nColumns-1))),
```

```

comegaT = 0.1 * exp(matrix(rnorm(1*(nColumns-1),0,0.011),nrow=(nColumns-1))),
corr_L = diag(nColumns-1),
paramN = cbind(2+0.75*(logPobs-2.2), 4*matrix(1,nAnalytes,1)+0.5*(logPobs-2.2))
dS1N = matrix(0,nAnalytes,1),
dlogkT = rnorm(nAnalytes,-0.0868,0.0217),
dlogkwA = matrix(-1,nGroupsA,1),
dlogkwB = matrix(-1,nGroupsB,1),
dS1mA = matrix(0,nGroupsA,1),
dS1mB = matrix(0,nGroupsB,1),
dS1A = matrix(0,nGroupsA,1),
dS1B = matrix(0,nGroupsB,1),

etaclogkwNStd = matrix(0,nColumns-1,nAnalytes),
etacS1mN = matrix(0,nColumns-1,nAnalytes),
etacdS1N = matrix(0,nColumns-1,nAnalytes),
etacdlogkT = matrix(0,nColumns-1,nAnalytes),
etacdlogkwA = matrix(0,nColumns-1,nGroupsA),
etacdlogkwB = matrix(0,nColumns-1,nGroupsB),
etacdS1mA = matrix(0,nColumns-1,nGroupsA),
etacdS1mB = matrix(0,nColumns-1,nGroupsB),
etacdS1A = matrix(0,nColumns-1,nGroupsA),
etacdS1B = matrix(0,nColumns-1, nGroupsB),
pKawA = pKaslitA,
pKawB = pKaslitB,
etaalphamA = matrix(0,nGroupsA,1),
etaalphamB = matrix(0,nGroupsB,1),
etadalphaA = matrix(0,nGroupsA,1),
etadalphaB = matrix(0,nGroupsB,1),

msigma = rlnorm(1,log(1),0.2),
ssigma = rlnorm(1,log(0.2),0.2),
logsigma = rnorm(nAnalytes,0,0.2),

clogmsigma = rnorm(nColumns-1,0,0.125),
cssigma = rlnorm(nColumns-1,log(0.125),0.125),
etaclogsigma = matrix(rnorm(nAnalytes*(nColumns-1),0,0.125),nrow=(nColumns-1))

)
}

```

5.2.2 The Stan model:

```
writeLines(readLines("stan/hplc-gra-fivecolumns.stan"))

functions {
  // credit http://srmart.in/informative-priors-for-correlation-matrices-an-easy-approach/
  real lkj_corr_point_lower_tri_lpdf(matrix rho, real point_mu_lower,
                                     real point_scale_lower) {
    // works only for [2x2 matrix]
    real lpdf = lkj_corr_lpdf(rho | 1)
        + normal_lpdf(rho[2, 1] | point_mu_lower, point_scale_lower);
    return lpdf;
  }

  // pH and fi at a given time at column inlet
  vector gra_state(real t, vector hplcparam) {
    vector[2] sol;
    real tg = hplcparam[1];
    real td = hplcparam[2];
    real fio = hplcparam[5];
    real fik = hplcparam[6];
    real pHo = hplcparam[8];
    real alpha1 = hplcparam[9];
    real alpha2 = hplcparam[10];
    real fi;

    fi = fio + (fik - fio) / tg * (t - td);

    if (t < td) {
      fi = fio;
    } else if (t > tg + td) {
      fi = fik;
    }

    sol[1] = fi;
    sol[2] = pHo + alpha1 * fi + alpha2 * fi ^ 2;

    return sol;
  }

  real funlnki(vector logkw, vector apH, vector S1, real S2, vector pKaw, vector alpha,
```

```

        int nDiss, real fi, real pH) {
real lnki;
vector[3] logkix;
vector[2] pHmpKa;

logkix = log(10) *(logkw - S1*(1+S2) * fi / (1 + S2 * fi) + apH * (pH - 7));
pHmpKa = log(10) *(pH - (pKaw + alpha * fi));

if (nDiss == 0) {
    lnki = logkix[1];
} else if (nDiss == 1) {
    lnki = logkix[1] +
        log1p_exp(pHmpKa[1] + logkix[2] - logkix[1]) -
        log1p_exp(pHmpKa[1]);
} else if (nDiss == 2) {
    lnki = logkix[1] +
        log1p_exp(pHmpKa[1]+logkix[2]-logkix[1] + log1p_exp(pHmpKa[2]+logkix[3]-logkix[2]) -
        log1p_exp(pHmpKa[1] + log1p_exp(pHmpKa[2]));
}

return lnki;
}

vector areaandslope(real dt, real lnki1, real lnki2, real invki1, real invki2) {
    vector[2] cki_b;
    real bo;
    real cki;

    if (invki2 > 1.001 * invki1) {
        bo = (lnki1 - lnki2) / dt;
        cki = (invki2 - invki1) / bo;
    }

    else if (invki1 > 1.001 * invki2) {
        bo = (lnki2 - lnki1) / dt;
        cki = (invki1 - invki2) / bo;
    }
    else {
        bo = 0.001 / dt;
        cki = dt * (invki2 + invki1) / 2;
    }

    cki_b[1] = cki;
}

```

```

    cki_b[2] = bo;

    return cki_b;
}

real chromgratrapz(int steps, vector logkw, vector apH, vector S1,  real S2, vector pKaw,
                    vector alpha, int nDiss, vector hplcparam) {

    real tg = hplcparam[1];
    real td = hplcparam[2];
    real to = hplcparam[3];
    real te = hplcparam[4];

    vector[1] sol;
    real time1;
    real time2;
    vector[2] fipH1;
    vector[2] fipH2;
    real lnki1;
    real lnki2;
    real invki1;
    real invki2;
    vector[2] cki_b;
    real cumki1;
    real cumki2;
    real bo;
    real tr;
    real dt;

    dt = tg / steps;

    time1 = 0;
    time2 = td;

    fipH1 = gra_state(time1, hplcparam);
    lnki1 = funlnki(logkw, apH, S1, S2, pKaw, alpha, nDiss, fipH1[1], fipH1[2]);
    lnki2=lnki1;

    invki1 = exp(-lnki1)/to;
    invki2 = invki1;

    cumki1 = 0;
    cumki2 = td*invki1;
}

```

```

bo = 0.001 / td;

for (x in 1 : steps) {
  if (cumki2 >= 1) {
    continue;
  }
  time1 = time2;
  time2 += dt;
  fipH2 = gra_state(time2, hplcparam);
  lnki1 = lnki2;
  lnki2 = funlnki(logkw, apH, S1, S2, pKaw, alpha, nDiss, fipH2[1], fipH2[2]);
  invki1 = invki2;
  invki2 = exp(-lnki2)/to;
  cki_b = areaandslope(dt, lnki1, lnki2, invki1, invki2);
  cumki1 = cumki2;
  cumki2 += cki_b[1];
  bo = cki_b[2];
}

if (cumki2 >= 1 && cumki1==0) {
  tr = te+to+1/invki2;
} else if (cumki2 >= 1) {
  tr = te+to+time1+log1p_exp(log((1-cumki1)*bo*to) + lnki1)/bo;
} else if (cumki2 < 1) {
  tr = te+to+time2+(1-cumki2)/invki2;
}

return tr;
}

real partial_sum(array[] int ind, int start, int end,
                vector trobs,
                array[] int steps,
                array[] vector hplcparam,
                array[] int analyte,
                array[] int column,
                array[] int modifier,
                array[] int R,
                array[,] vector logkw,
                array[,] vector apH,
                array[, ,] vector S1,

```

```

        array[,] real S2,
        array[] vector pKaw,
        array[,] vector alpha,
        array[,] real dlogkT,
        array[] vector sigma) {

real lp = 0;

for (z in start : end) {

real y_hat = chromgratrapz(steps[z],
                           logkw[analyte[z], column[z], : ] + dlogkT[analyte[z], column
                           apH[analyte[z], column[z], : ],
                           S1[analyte[z], modifier[z], column[z], : ],
                           S2[modifier[z], column[z]],
                           pKaw[analyte[z], : ],
                           alpha[analyte[z], modifier[z], : ],
                           R[analyte[z]],
                           hplcparam[z]);

lp = lp + student_t_lpdf(trobs[z] | 3, y_hat, sigma[analyte[z], column[z]]);

}
return lp;
}
}

data {
int nAnalytes;           // number of analytes
int nColumns;            // number of columns
int nModifiers;          // number of org. modifiers
int nObs;                // number of observations
array[nObs] int analyte; // analyte indexes
array[nObs] int column;  // column indexes
array[nObs] int modifier; // modifier indexes
array[nObs] int<lower=1> steps; // steps for gradient retention time approximation
array[nObs] vector[12] hplcparam; // [tg, td, to, te, fio, fik, org modifier, pHo, alpha1,
vector[nAnalytes] logPobs;
int<lower=0, upper=2> maxR; //
array[nAnalytes] int<lower=0, upper=2> R;
int<lower=1> nGroupsA;
int<lower=1> nGroupsB;
vector[nGroupsA] pKaslitA;
}

```

```

vector[nGroupsB] pKaslitB;
array[nGroupsA,2] int idxGroupsA;
array[nGroupsB,2] int idxGroupsB;

vector[nObs] trobs; // observed retention factors
int<lower=0, upper=1> run_estimation; // 0 for prior predictive, 1 for estimation
}

transformed data {
  int grainsize = 1;
  array[nObs] int ind = rep_array(1, nObs);
}

parameters {

  real logkwHat; // typical logkw [Neutral]
  real S1mHat; // effect of MeOH on logkw [Neutral]
  real dS1Hat; // effect of ACN on S1m [Neutral]
  array[2] real dlogkwHat; // effect of dissociation on logkw [Acids, Bases]
  array[2] real dS1mHat; // effect of dissociation on S1m [Acids, Bases]
  array[2] real ddS1Hat; // effect of dissociation on dS1 [Acids, Bases]
  real logS2mHat; // typical value of S2m (log10 scale)
  real dlogS2mHat; // effect of ACN on logS2m
  vector[2] beta; // effect of logP on logkw and S1m
  real dlogkTHat; // effect of temperature on logkw
  vector[2] apH; // effect of pH on logkw [Acids, Bases]
  array[2] real alphamHat; // effect of MeOH on pKa [Acids, Bases]
  array[2] real dalphaHat; // effect of ACN on alpham [Acids, Bases]
  vector<lower=0>[3] tau; // sd for between analyte variability of pKa's
  vector<lower=0>[3] omega; // sd of BAV [logkw, S1m, dS1]
  corr_matrix[2] rho; // correlation matrix [logkw vs. S1m]
  real<lower=0> omegaT; // sd of BAV [dlogkT]
  vector<lower=0>[3] kappa; // sd of BAV [dlogkw, dS1m, ddS1]

  // 2nd column
  vector[nColumns-1] clogkwHat; // effect of column on logkw [Neutral]
  vector[nColumns-1] cS1mHat; // effect of column on S1m [Neutral]
  vector[nColumns-1] cdS1Hat; // effect of column on dS1 [Neutral]
  matrix[nColumns-1,2] cdlogkwHat; // effect of column on logkw [Acids, Bases]
  matrix[nColumns-1,2] cdS1mHat; // effect of column on dS1m [Acids, Bases]
  matrix[nColumns-1,2] cddS1Hat; // effect of column on ddS1 [Acids, Bases]
  matrix[nColumns-1,2] cbeta;
  vector[nColumns-1] cdlogkTHat; // effect of column on dlogkTHat
  matrix[nColumns-1,2] capH; // effect of column on apH [Acids, Bases]
}

```

```

matrix<lower=0>[nColumns-1,3] comega; // sd of BAV [clogkw,cS1m,cdS1]
vector<lower=0>[nColumns-1] comegaT; // sd of BAV [cdlogkT]
matrix<lower=0>[nColumns-1,3] ckappa; // sd of BAV [cdlogkw,cdS1m,cddS1]
cholesky_factor_corr[nColumns-1] corr_L; // cholesky factor correlation matrix

// 1st column
array[nAnalytes] vector[2] paramN;
vector[nAnalytes] dS1N;
vector[nAnalytes] dlogkT;
vector[nGroupsA] dlogkwA;
vector[nGroupsB] dlogkwB;
vector[nGroupsA] dS1mA;
vector[nGroupsB] dS1mB;
vector[nGroupsA] dS1A;
vector[nGroupsB] dS1B;

// 2nd column
matrix[nColumns-1, nAnalytes] etaclogkwNStd;
array[nColumns-1] vector[nAnalytes] etacS1mN;
array[nColumns-1] vector[nAnalytes] etacdS1N;
array[nColumns-1] vector[nAnalytes] etacdlogkT;
array[nColumns-1] vector[nGroupsA] etacdlogkwA;
array[nColumns-1] vector[nGroupsB] etacdlogkwB;
array[nColumns-1] vector[nGroupsA] etacdS1mA;
array[nColumns-1] vector[nGroupsB] etacdS1mB;
array[nColumns-1] vector[nGroupsA] etacdS1A;
array[nColumns-1] vector[nGroupsB] etacdS1B;

// Dissociation
vector[nGroupsA] pKawA;
vector[nGroupsB] pKawB;
vector[nGroupsA] etaalphanA;
vector[nGroupsB] etaalphanB;
vector[nGroupsA] etadalphaA;
vector[nGroupsB] etadalphaB;

// residual variability for the 1st and 2nd column
real<lower=0> msigma; // typical sigma for the 1st column]
real<lower=0> ssigma;
vector[nAnalytes] logsigma;

vector[nColumns-1] clogmsigma; // effect of column on log(msigma)]
vector<lower=0>[nColumns-1] cssigma; ; //sd of residual [1st,2nd column]

```

```

array[nColumns-1] vector[nAnalytes] etaclogsigma;
}
transformed parameters {
cov_matrix[2] Omega;

array[nAnalytes] vector[3] miu;
array[nAnalytes,3] vector[nColumns-1] cmiu;
array[nAnalytes,nColumns] vector[maxR + 1] logkwx;
array[nAnalytes,nModifiers, nColumns] vector[maxR + 1] S1x;
array[nModifiers,nColumns] real S2x;
array[nAnalytes,nColumns] vector[maxR + 1] apHx;
array[nAnalytes,nModifiers] vector[maxR] alphax;
array[nAnalytes, nColumns] real dlogkTx;
array[nAnalytes] vector[maxR] pKawx;
array[nAnalytes] vector[nColumns] sigmax;

array[nColumns-1] vector[nAnalytes] clogkwN;
matrix[nColumns-1, nAnalytes] etaclogkwN;
array[nColumns-1] vector[nAnalytes] cS1mN;
array[nColumns-1] vector[nAnalytes] cdS1N;
array[nColumns-1] vector[nGroupsA] cdlogkwA;
array[nColumns-1] vector[nGroupsB] cdlogkwB;
array[nColumns-1] vector[nGroupsA] cdS1mA;
array[nColumns-1] vector[nGroupsB] cdS1mB;
array[nColumns-1] vector[nGroupsA] cdS1A;
array[nColumns-1] vector[nGroupsB] cdS1B;
array[nColumns-1] vector[nAnalytes] cdlogkT;
array[nColumns-1] vector[nAnalytes] clogsigma;
vector[nGroupsA] alphamA;
vector[nGroupsB] alphamB;
vector[nGroupsA] dalphaA;
vector[nGroupsB] dalphaB;

Omega = quad_form_diag(rho, omega[1 : 2]); // diag_matrix(omega) * rho * diag_matrix(omega)

for (i in 1 : nAnalytes) {
  miu[i, 1] = logkwHat + beta[1] * (logPobs[i] - 2.2);
  miu[i, 2] = S1mHat + beta[2] * (logPobs[i] - 2.2);
  miu[i, 3] = dS1Hat;
}

for (i in 1 : nAnalytes) {

```

```

for (c in 1 : (nColumns-1)) {
  cmiu[i, 1, c] = clogkwHat[c] + cbeta[c,1] * (logPobs[i] - 2.2);
  cmiu[i, 2, c] = cS1mHat[c] + cbeta[c,2] * (logPobs[i] - 2.2);
  cmiu[i, 3, c] = cdS1Hat[c];
}

// Matt's trick to use unit scale
etaclogkwN = diag_pre_multiply(comega[,1], corr_L * etaclogkwNStd);

for (i in 1 : nAnalytes) {
  logkwx[i, 1, :] = paramN[i,1]*[1,1,1]';
  for (c in 1 : (nColumns-1)) {
    clogkwN[c,i] = cmiu[i, 1, c] + etaclogkwN[c,i];
    logkwx[i, c+1, :] = (paramN[i,1]+clogkwN[c,i])*[1,1,1]';
  }
}

for (d in 1 : nGroupsA) {
  logkwx[idxGroupsA[d,1], 1, 1+idxGroupsA[d,2]] += dlogkwA[d];
  if (idxGroupsA[d,2]==1) {
    logkwx[idxGroupsA[d,1], 1, 3] += dlogkwA[d];
  }
  for (c in 1 : (nColumns-1)) {
    cdlogkwA[c,d] = cdlogkwHat[c,1] + ckappa[c,1]*etacdlogkwA[c,d];
    logkwx[idxGroupsA[d,1], c+1, 1+idxGroupsA[d,2]] += dlogkwA[d]+cdlogkwA[c,d];
    if (idxGroupsA[d,2]==1) {
      logkwx[idxGroupsA[d,1], c+1, 3] += dlogkwA[d]+cdlogkwA[c,d];
    }
  }
}

for (d in 1 : nGroupsB) {
  logkwx[idxGroupsB[d,1], 1, idxGroupsB[d,2]] += dlogkwB[d];
  if (idxGroupsB[d,2]==2) {
    logkwx[idxGroupsB[d,1], 1, 1] += dlogkwB[d];
  }
  for (c in 1 : (nColumns-1)) {
    cdlogkwB[c,d] = cdlogkwHat[c,2]+ ckappa[c,1]*etacdlogkwB[c,d];
    logkwx[idxGroupsB[d,1], c+1, idxGroupsB[d,2]] += dlogkwB[d]+cdlogkwB[c,d];
    if (idxGroupsB[d,2]==2) {
      logkwx[idxGroupsB[d,1], c+1, 1] += dlogkwB[d]+cdlogkwB[c,d];
    }
  }
}

for (i in 1 : nAnalytes) {
  apHx[i, 1, :] = [0,0,0]';
  for (c in 1 : (nColumns-1)) {

```

```

    apHx[i, c+1, : ] = [0,0,0] ';
  } }

  for (d in 1 : nGroupsA) {
    apHx[idxGroupsA[d,1], 1, 1+idxGroupsA[d,2]] += apH[1];
  if (idxGroupsA[d,2]==1) {
    apHx[idxGroupsA[d,1], 1, 3] += apH[1];
  }
  for (c in 1 : (nColumns-1)) {
    apHx[idxGroupsA[d,1], c+1, 1+idxGroupsA[d,2]] += apH[1]+capH[c,1];
  if (idxGroupsA[d,2]==1) {
    apHx[idxGroupsA[d,1], c+1, 3] += apH[1]+capH[c,1];
  }}}

  for (d in 1 : nGroupsB) {
    apHx[idxGroupsB[d,1], 1, idxGroupsB[d,2]] += apH[2];
    if (idxGroupsB[d,2]==2) {
      apHx[idxGroupsB[d,1], 1, 1] += apH[2];
    }
    for (c in 1 : (nColumns-1)) {
      apHx[idxGroupsB[d,1], c+1, idxGroupsB[d,2]] += apH[2]+capH[c,2];
      if (idxGroupsB[d,2]==2) {
        apHx[idxGroupsB[d,1], c+1, 1] += apH[2]+capH[c,2];
      }}}

  for (i in 1 : nAnalytes) {
    S1x[i, 1, 1, : ] = paramN[i,2]*[1,1,1]';
    S1x[i, 2, 1, : ] = paramN[i,2]*[1,1,1]' + dS1N[i];
  for (c in 1 : (nColumns-1)) {
    cS1mN[c,i] = cmiu[i,2,c] + comega[c,2]*etacS1mN[c,i];
    cdS1N[c,i] = cmiu[i,3,c] + comega[c,3]*etacdS1N[c,i];
    S1x[i, 1, c+1, : ] = (paramN[i,2]+cS1mN[c,i])*[1,1,1]';
    S1x[i, 2, c+1, : ] = (paramN[i,2]+cS1mN[c,i])*[1,1,1]' + (dS1N[i]+cdS1N[c,i]);
  }}}

  for (d in 1 : nGroupsA) {
    S1x[idxGroupsA[d,1], 1, 1, 1+idxGroupsA[d,2]] += dS1mA[d];
    S1x[idxGroupsA[d,1], 2, 1, 1+idxGroupsA[d,2]] += dS1mA[d]+dS1A[d];

  if (idxGroupsA[d,2]==1) {
    S1x[idxGroupsA[d,1], 1, 1, 3] += dS1mA[d];
    S1x[idxGroupsA[d,1], 2, 1, 3] += dS1mA[d]+dS1A[d];
  }}}

```

```

}

for (c in 1 : (nColumns-1)) {
  cdS1mA[c,d] = cdS1mHat[c,1] + ckappa[c,2]*etacdS1mA[c,d];
  cdS1A[c,d] = cddS1Hat[c,1] + ckappa[c,3]*etacdS1A[c,d];
  S1x[idxGroupsA[d,1], 1, c+1, 1+idxGroupsA[d,2]] += dS1mA[d] +cdS1mA[c,d];
  S1x[idxGroupsA[d,1], 2, c+1, 1+idxGroupsA[d,2]] += dS1mA[d]+dS1A[d]+cdS1mA[c,d]+cdS1A[c,d];
  if (idxGroupsA[d,2]==1) {
    S1x[idxGroupsA[d,1], 1, c+1, 3] += dS1mA[d] +cdS1mA[c,d];
    S1x[idxGroupsA[d,1], 2, c+1, 3] += dS1mA[d]+dS1A[d]+cdS1mA[c,d]+cdS1A[c,d];
  }}}

for (d in 1 : nGroupsB) {
  S1x[idxGroupsB[d,1], 1, 1, idxGroupsB[d,2]] += dS1mB[d];
  S1x[idxGroupsB[d,1], 2, 1, idxGroupsB[d,2]] += dS1mB[d]+dS1B[d];
  if (idxGroupsB[d,2]==2) {
    S1x[idxGroupsB[d,1], 1, 1, 1] += dS1mB[d];
    S1x[idxGroupsB[d,1], 2, 1, 1] += dS1mB[d]+dS1B[d];
  }
  for (c in 1 : (nColumns-1)) {
    cdS1mB[c,d] = cdS1mHat[c,2] + ckappa[c,2]*etacdS1mB[c,d];
    cdS1B[c,d] = cddS1Hat[c,2] + ckappa[c,3]*etacdS1B[c,d];

    S1x[idxGroupsB[d,1], 1, c+1, idxGroupsB[d,2]] += dS1mB[d] +cdS1mB[c,d];
    S1x[idxGroupsB[d,1], 2, c+1, idxGroupsB[d,2]] += dS1mB[d]+dS1B[d]+cdS1mB[c,d]+cdS1B[c,d];
    if (idxGroupsB[d,2]==2) {
      S1x[idxGroupsB[d,1], 1, c+1, 1] += dS1mB[d] +cdS1mB[c,d];
      S1x[idxGroupsB[d,1], 2, c+1, 1] += dS1mB[d]+dS1B[d]+cdS1mB[c,d]+cdS1B[c,d];
    }}}

S2x[1,1] = 10^(logS2mHat);
S2x[2,1] = 10^(logS2mHat + dlogS2Hat);

for (c in 1 : (nColumns-1)) {
  S2x[1,c+1] = S2x[1,1];
  S2x[2,c+1] = S2x[2,1];
}

for (i in 1 : nAnalytes) {
  dlogkTx[i, 1] = dlogkT[i];
  for (c in 1 : (nColumns-1)) {
    cdlogkT[c,i] = cdlogkTHat[c]+comegaT[c]*etacdlogkT[c,i];
    dlogkTx[i, c+1] = dlogkT[i] + cdlogkT[c,i];
  }
}

```

```

  } }

  for (i in 1 : nAnalytes) {
    pKawx[i, : ] = [0,0]';
    alphax[i, 1, : ] = [0,0]';
    alphax[i, 2, : ] = [0,0]';
  }

  for (d in 1 : nGroupsA) {

    alphamA[d] = alphamHat[1] + tau[2]*etaalphamA[d];
    dalphaA[d] = dalphaHat[1] + tau[3]*etadalphaA[d];

    pKawx[idxGroupsA[d,1], idxGroupsA[d,2]] = pKawA[d];
    alphax[idxGroupsA[d,1], 1, idxGroupsA[d,2]]= alphamA[d];
    alphax[idxGroupsA[d,1], 2, idxGroupsA[d,2]]= alphamA[d]+dalphaA[d];
  }

  for (d in 1 : nGroupsB) {
    alphamB[d] = alphamHat[2] + tau[2]*etaalphamB[d];
    dalphaB[d] = dalphaHat[2] + tau[3]*etadalphaB[d];
    pKawx[idxGroupsB[d,1], idxGroupsB[d,2]] = pKawB[d];
    alphax[idxGroupsB[d,1], 1, idxGroupsB[d,2]]= alphamB[d];
    alphax[idxGroupsB[d,1], 2, idxGroupsB[d,2]]= alphamB[d]+dalphaB[d];
  }

  for (i in 1 : nAnalytes) {
    sigmax[i, 1] = exp(logsigma[i]);
    for (c in 1 : (nColumns-1)) {
      clogmsigma[c,i] = clogmsigma[c] + cssigma[c]*etaclogmsigma[c,i];
      sigmax[i, c+1] = exp(logsigma[i]+clogmsigma[c,i]);
    }
  }

  model {
    logkwHat ~ normal(2.2, 2);
    S1mHat ~ normal(4, 1);
    dS1Hat ~ normal(1, 0.5);
    dlogkwHat ~ normal(-1, 0.125);
    dS1mHat ~ normal(0, 0.5);
    ddS1Hat ~ normal(0, 0.25);
    logS2mHat ~ normal(-0.7, 0.125);
    dlogS2Hat ~ normal(1, 0.125);
  }
}

```

```

beta[{1}] ~ normal(1, 0.125);
beta[{2}] ~ normal(0.5, 0.5);
dlogkTHat ~ normal(-0.087, 0.022);
apH ~ normal(0, 0.1);

alphamHat[{1}] ~ normal(2, 0.25);
alphamHat[{2}] ~ normal(-1, 0.25);
dalphaHat ~ normal(0, 0.125);
tau[{1}] ~ normal(0, 0.25);
tau[{2,3}] ~ normal(0, 0.125);
omega ~ normal(0, 2);
rho ~ lkj_corr_point_lower_tri(0.75, 0.125);
omegaT ~ normal(0, 0.022);
kappa ~ normal(0, 0.25);

clogkwHat ~ normal(0, 1);
cS1mHat ~ normal(0, 0.5);
cdS1Hat ~ normal(0, 0.25);
to_vector(cdlogkwHat) ~ normal(0, 0.0625);
to_vector(cdS1mHat) ~ normal(0, 0.25);
to_vector(cddS1Hat) ~ normal(0, 0.125);
to_vector(cbeta) ~ normal(0, 0.25);
cdlogkTHat ~ normal(0, 0.011);
to_vector(capH) ~ normal(0, 0.05);
to_vector(comega) ~ normal(0, 0.5);
comegaT ~ normal(0, 0.011);
to_vector(ckappa) ~ normal(0, 0.125);

corr_L~lkj_corr_cholesky(2.0);

for (i in 1 : nAnalytes) {
  paramN[i] ~ multi_normal(miu[i,1:2], Omega);
}

dS1N ~ normal(miu[,3], omega[3]);
dlogkT ~ normal(dlogkTHat, omegaT);
dlogkwA ~ normal(dlogkwHat[1], kappa[1]);
dlogkwB ~ normal(dlogkwHat[2], kappa[1]);
dS1mA ~ normal(dS1mHat[1], kappa[2]);
dS1mB ~ normal(dS1mHat[2], kappa[2]);
dS1A ~ normal(ddS1Hat[1], kappa[3]);
dS1B ~ normal(ddS1Hat[2], kappa[3]);

```

```

to_vector(etaclogkwNStd) ~ normal(0, 1);

for (c in 1 : (nColumns-1)) {
  etacS1mN[c] ~ normal(0,1);
  etacdS1N[c] ~ normal(0,1);
  etacdlogkT[c] ~ normal(0,1);
  etacdlogkwA[c] ~ normal(0,1);
  etacdlogkwB[c] ~ normal(0,1);
  etacdS1mA[c] ~ normal(0,1);
  etacdS1mB[c] ~ normal(0,1);
  etacdS1A[c] ~ normal(0,1);
  etacdS1B[c] ~ normal(0,1);
}

pKawA ~ normal(pKaslitA, tau[1]);
pKawB ~ normal(pKaslitB, tau[1]);
etaalphamA ~ normal(0,1);
etaalphamB ~ normal(0,1);
etadalphaA ~ normal(0,1);
etadalphaB ~ normal(0,1);

msigma ~ normal(0,1);
ssigma ~ normal(0,1);
logsigma ~ normal(log(msigma),ssigma);

clogmsigma ~ normal(0,0.125);
cssigma ~ normal(0,0.125);

for (c in 1 : (nColumns-1)) {
  etaclogsigma[c] ~ normal(0,1);
}

if (run_estimation == 1) {

  target += reduce_sum(partial_sum, ind, grainsize, trobs, steps,
                        hplcparam, analyte, column, modifier, R, logkwx, apHx,
                        S1x, S2x, pKawx, alphax, dlogkTx, sigmax);
}

generated quantities {
  corr_matrix[nColumns-1] crho;
  crho = corr_L * corr_L';
}

```

```
}
```

5.2.3 Fitting the model

Compile the model:

```
mod1 <- cmdstan_model("stan/hplc-gra-fivecolumns.stan",
                      stanc_options = list("01"),
                      cpp_options = list(stan_threads = TRUE))

# for optimization
modode <- cmdstan_model("stan/hplc-gra-fivecolumns.stan",
                        stanc_options = list("01"))
```

The optimization was used for initial testing:

```
fit_ode <- modode$optimize(
  data = datastruct,
  output_dir = "stanfiles",
  init = init
)

fit_ode$print(max_rows=100)
```

For local computations one can use cmdstanr:

```
# fit <- mod1$sample(
#   data = datastruct,
#   output_dir = "stanfiles",
#   init = init,
#   iter_warmup = 1000,
#   iter_sampling = 1000,
#   chains = 4,
#   parallel_chains = 4,
#   threads_per_chain = 6,
#   refresh = 100,
#   adapt_delta=0.9
# )
```

We performed computations at the Academic Computer Center in Gdańsk, [Tryton Cluster](#).
In this case:

1. we dumped the necessary data to .json format

```

write_stan_json(datastruct, "stan/standata.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-1.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-2.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-3.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-4.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-5.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-6.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-7.json", always_decimal = FALSE)
write_stan_json(init(), "stan/init-8.json", always_decimal = FALSE)

```

2. we run the model using the batch file:
3. After calculations, we loaded the output files using cmdstanr. The files are accessible through [stan output files](#).

```

fit <- cmdstanr::as_cmdstan_fit(c(
  'stanfiles/output_1.csv',
  'stanfiles/output_2.csv',
  'stanfiles/output_3.csv',
  'stanfiles/output_4.csv',
  'stanfiles/output_5.csv',
  'stanfiles/output_6.csv',
  'stanfiles/output_7.csv',
  'stanfiles/output_8.csv'
))

```

4. finally we checked the diagnostics of Monte Carlo inferences based on the Stan documentation described here .

```

#fit$cmdstan_diagnose()
setwd("stanfiles")
str = paste0(cmdstan_path(), "/bin/diagnose  output_*.csv")
system(str,intern=TRUE)

```

The diagnostics are reasonable given model complexity. Output copied here to save time:

6 Results

6.1 Summary of model parameters (table):

```
# fit$print(max_rows=200) # for first 200 parameters

fit$print(c("logkwHat", "S1mHat", "dS1Hat",
           "dlogkwHat", "dS1mHat", "ddS1Hat",
           "logS2mHat", "dlogS2Hat",
           "beta",
           "dlogkTHat", "apH",
           "omega", "omegaT", "kappa",
           "rho[2,1]",
           "msigma", "ssigma"), max_rows = 26)
```

variable	mean	median	sd	mad	q5	q95	rhat	ess_bulk	ess_tail
logkwHat	3.60	3.60	0.08	0.08	3.47	3.72	1.00	5947	3030
S1mHat	4.92	4.92	0.08	0.08	4.79	5.05	1.00	5251	3063
dS1Hat	0.61	0.61	0.05	0.05	0.53	0.69	1.00	3259	2365
dlogkwHat[1]	-0.79	-0.79	0.07	0.07	-0.91	-0.67	1.00	7672	3143
dlogkwHat[2]	-0.97	-0.97	0.05	0.05	-1.05	-0.88	1.00	5300	2998
dS1mHat[1]	0.17	0.17	0.12	0.12	-0.03	0.36	1.00	4231	3375
dS1mHat[2]	0.12	0.11	0.07	0.07	0.00	0.24	1.00	2482	2783
ddS1Hat[1]	0.28	0.28	0.08	0.08	0.15	0.41	1.00	6200	3217
ddS1Hat[2]	-0.67	-0.67	0.05	0.06	-0.76	-0.58	1.00	6993	3444
logS2mHat	-0.31	-0.31	0.01	0.01	-0.33	-0.28	1.02	432	986
dlogS2Hat	0.42	0.42	0.01	0.01	0.41	0.43	1.01	552	1093
beta[1]	0.84	0.84	0.04	0.04	0.77	0.91	1.00	7278	3170
beta[2]	0.51	0.51	0.05	0.05	0.43	0.58	1.00	3976	2814
dlogkTHat	-0.09	-0.09	0.00	0.00	-0.09	-0.08	1.00	4047	2966
apH[1]	-0.03	-0.03	0.00	0.00	-0.03	-0.03	1.00	4583	3724
apH[2]	0.08	0.08	0.00	0.00	0.08	0.08	1.00	3114	3288
omega[1]	0.92	0.92	0.06	0.06	0.83	1.02	1.00	5132	2669
omega[2]	0.93	0.93	0.06	0.06	0.84	1.03	1.00	4993	3096
omega[3]	0.55	0.55	0.03	0.03	0.50	0.61	1.00	8536	3214
omegaT	0.03	0.03	0.00	0.00	0.03	0.04	1.00	7750	3273
kappa[1]	0.59	0.58	0.03	0.03	0.53	0.65	1.00	4404	3301
kappa[2]	0.69	0.69	0.05	0.05	0.62	0.77	1.00	3416	3502
kappa[3]	0.55	0.55	0.04	0.04	0.49	0.61	1.00	3604	3242
rho[2,1]	0.87	0.87	0.02	0.02	0.83	0.91	1.00	3500	3223
msigma	0.39	0.39	0.03	0.03	0.35	0.44	1.00	9144	3009

```
ssigma      0.81  0.81 0.05 0.05  0.74  0.90 1.00      12125      2859
```

```
fit$print(c("alphamHat", "dalphaHat",
           "tau"), max_rows = 7)
```

variable	mean	median	sd	mad	q5	q95	rhat	ess_bulk	ess_tail
alphamHat[1]	2.22	2.22	0.15	0.15	1.96	2.47	1.00	1721	2559
alphamHat[2]	-1.35	-1.35	0.10	0.10	-1.51	-1.18	1.01	1282	2346
dalphaHat[1]	0.22	0.22	0.10	0.10	0.06	0.38	1.00	2171	2544
dalphaHat[2]	-0.20	-0.20	0.07	0.07	-0.32	-0.08	1.01	1514	2412
tau[1]	0.88	0.88	0.05	0.05	0.81	0.97	1.00	6372	3337
tau[2]	0.96	0.96	0.06	0.06	0.87	1.06	1.00	2193	2683
tau[3]	0.79	0.79	0.05	0.05	0.71	0.88	1.00	1967	2870

```
fit$print(c("clogkwHat", "cS1mHat", "cdS1Hat",
           "cdlogkwHat", "cdS1mHat", "cddS1Hat",
           "cbeta",
           "cdlogkTHat",
           "capH",
           "comega", "ckappa", "comegaT",
           "crho",
           "clogmsigma", "cssigma"), max_rows = 108)
```

variable	mean	median	sd	mad	q5	q95	rhat	ess_bulk	ess_tail
clogkwHat[1]	0.43	0.43	0.01	0.01	0.41	0.45	1.01	554	1048
clogkwHat[2]	0.18	0.18	0.01	0.01	0.16	0.21	1.01	734	1437
clogkwHat[3]	0.11	0.11	0.01	0.01	0.09	0.13	1.01	612	1192
clogkwHat[4]	0.18	0.18	0.01	0.01	0.16	0.19	1.01	726	1350
cS1mHat[1]	0.59	0.59	0.02	0.02	0.56	0.62	1.00	2127	2781
cS1mHat[2]	-0.10	-0.10	0.02	0.02	-0.14	-0.06	1.00	2010	2704
cS1mHat[3]	0.37	0.37	0.02	0.02	0.34	0.40	1.00	1308	3004
cS1mHat[4]	0.50	0.50	0.02	0.02	0.47	0.52	1.00	1837	3007
cdS1Hat[1]	0.15	0.15	0.01	0.01	0.13	0.17	1.01	928	1643
cdS1Hat[2]	0.81	0.81	0.03	0.02	0.77	0.85	1.01	379	872
cdS1Hat[3]	0.51	0.51	0.04	0.04	0.44	0.58	1.05	153	462
cdS1Hat[4]	0.04	0.04	0.02	0.02	0.01	0.06	1.00	607	1094
cdlogkwHat[1,1]	0.04	0.04	0.02	0.02	0.02	0.07	1.00	2224	2870
cdlogkwHat[2,1]	-0.05	-0.05	0.02	0.02	-0.08	-0.02	1.00	1549	2160
cdlogkwHat[3,1]	-0.03	-0.03	0.02	0.02	-0.06	0.00	1.00	1668	2740
cdlogkwHat[4,1]	0.04	0.04	0.02	0.02	0.02	0.07	1.00	1319	2124

cdlogkwHat[1,2]	0.00	0.00	0.01	0.01	-0.02	0.02	1.00	1667	2364
cdlogkwHat[2,2]	-0.04	-0.04	0.02	0.02	-0.07	-0.01	1.01	1538	2256
cdlogkwHat[3,2]	-0.06	-0.06	0.01	0.01	-0.08	-0.04	1.01	1175	1869
cdlogkwHat[4,2]	-0.03	-0.03	0.01	0.01	-0.05	-0.01	1.00	1137	2304
cdS1mHat[1,1]	-0.01	-0.01	0.04	0.04	-0.09	0.06	1.00	3059	3209
cdS1mHat[2,1]	-0.19	-0.19	0.06	0.06	-0.29	-0.10	1.00	2963	2774
cdS1mHat[3,1]	-0.49	-0.49	0.05	0.05	-0.56	-0.41	1.00	2231	2921
cdS1mHat[4,1]	-0.23	-0.23	0.04	0.04	-0.30	-0.16	1.00	2450	3229
cdS1mHat[1,2]	0.36	0.36	0.03	0.03	0.31	0.41	1.00	2707	3167
cdS1mHat[2,2]	-0.41	-0.41	0.04	0.04	-0.47	-0.35	1.00	2232	3005
cdS1mHat[3,2]	-0.09	-0.09	0.03	0.03	-0.14	-0.04	1.00	1905	2758
cdS1mHat[4,2]	0.29	0.29	0.03	0.03	0.24	0.34	1.00	2106	2691
cddS1Hat[1,1]	0.05	0.05	0.03	0.03	0.01	0.09	1.00	4704	3379
cddS1Hat[2,1]	0.13	0.13	0.05	0.05	0.06	0.21	1.00	2515	2663
cddS1Hat[3,1]	0.46	0.46	0.09	0.09	0.32	0.61	1.00	847	1433
cddS1Hat[4,1]	0.00	0.00	0.02	0.02	-0.04	0.04	1.00	4056	3302
cddS1Hat[1,2]	0.17	0.17	0.02	0.02	0.14	0.19	1.00	2144	3206
cddS1Hat[2,2]	0.57	0.57	0.03	0.03	0.52	0.62	1.00	1653	2285
cddS1Hat[3,2]	0.50	0.51	0.06	0.06	0.41	0.60	1.03	359	938
cddS1Hat[4,2]	-0.04	-0.04	0.02	0.01	-0.06	-0.01	1.00	3877	3310
cbeta[1,1]	0.00	0.00	0.01	0.01	-0.01	0.01	1.01	886	1478
cbeta[2,1]	-0.02	-0.02	0.01	0.01	-0.03	0.00	1.01	1160	1953
cbeta[3,1]	-0.03	-0.03	0.01	0.01	-0.04	-0.02	1.01	945	1688
cbeta[4,1]	-0.02	-0.02	0.01	0.01	-0.03	-0.01	1.01	1001	1807
cbeta[1,2]	-0.07	-0.07	0.01	0.01	-0.09	-0.05	1.00	2265	2809
cbeta[2,2]	0.10	0.10	0.01	0.01	0.08	0.12	1.00	2647	2838
cbeta[3,2]	-0.03	-0.03	0.01	0.01	-0.04	-0.01	1.00	2162	2837
cbeta[4,2]	-0.02	-0.02	0.01	0.01	-0.03	0.00	1.00	2452	2849
cdlogkTHat[1]	-0.01	-0.01	0.00	0.00	-0.01	0.00	1.00	3880	3363
cdlogkTHat[2]	-0.02	-0.02	0.00	0.00	-0.02	-0.02	1.00	4513	3484
cdlogkTHat[3]	-0.01	-0.01	0.00	0.00	-0.01	0.00	1.00	3368	3541
cdlogkTHat[4]	0.00	0.00	0.00	0.00	-0.01	0.00	1.00	4096	3513
capH[1,1]	-0.01	-0.01	0.00	0.00	-0.02	-0.01	1.00	4697	3790
capH[2,1]	-0.02	-0.02	0.00	0.00	-0.02	-0.02	1.00	5050	3925
capH[3,1]	0.00	0.00	0.00	0.00	0.00	0.01	1.00	4877	3408
capH[4,1]	-0.02	-0.02	0.00	0.00	-0.02	-0.02	1.00	4981	3932
capH[1,2]	-0.03	-0.03	0.00	0.00	-0.04	-0.03	1.00	3850	3894
capH[2,2]	-0.04	-0.04	0.00	0.00	-0.05	-0.04	1.00	3629	3707
capH[3,2]	-0.05	-0.05	0.00	0.00	-0.05	-0.05	1.00	3402	3568
capH[4,2]	-0.01	-0.01	0.00	0.00	-0.02	-0.01	1.00	3709	3544
comega[1,1]	0.12	0.12	0.01	0.01	0.10	0.13	1.01	862	1591
comega[2,1]	0.13	0.13	0.01	0.01	0.12	0.15	1.00	1315	2135
comega[3,1]	0.12	0.12	0.01	0.01	0.11	0.13	1.00	1157	2056

comega[4,1]	0.10	0.10	0.01	0.01	0.09	0.11	1.00	1212	1998
comega[1,2]	0.06	0.06	0.02	0.02	0.02	0.09	1.03	315	289
comega[2,2]	0.15	0.15	0.02	0.02	0.12	0.18	1.00	2256	2851
comega[3,2]	0.05	0.05	0.01	0.01	0.03	0.07	1.01	426	343
comega[4,2]	0.02	0.02	0.01	0.01	0.00	0.04	1.00	760	1516
comega[1,3]	0.14	0.14	0.01	0.01	0.13	0.16	1.01	1387	2315
comega[2,3]	0.29	0.29	0.02	0.02	0.26	0.32	1.00	864	1617
comega[3,3]	0.47	0.47	0.03	0.03	0.42	0.52	1.02	701	1289
comega[4,3]	0.16	0.16	0.01	0.01	0.15	0.19	1.00	1393	1907
ckappa[1,1]	0.07	0.07	0.01	0.01	0.06	0.08	1.00	1256	2247
ckappa[2,1]	0.11	0.11	0.01	0.01	0.10	0.13	1.00	1718	2961
ckappa[3,1]	0.08	0.08	0.01	0.01	0.07	0.09	1.01	1764	2733
ckappa[4,1]	0.08	0.08	0.01	0.01	0.07	0.09	1.00	1491	2505
ckappa[1,2]	0.07	0.07	0.03	0.04	0.02	0.13	1.01	417	852
ckappa[2,2]	0.20	0.20	0.03	0.03	0.15	0.25	1.00	1664	2343
ckappa[3,2]	0.11	0.11	0.03	0.03	0.06	0.15	1.01	862	830
ckappa[4,2]	0.04	0.03	0.02	0.03	0.00	0.08	1.03	479	1426
ckappa[1,3]	0.07	0.07	0.02	0.02	0.04	0.11	1.00	865	1522
ckappa[2,3]	0.27	0.27	0.02	0.02	0.24	0.31	1.00	1741	2782
ckappa[3,3]	0.72	0.72	0.04	0.04	0.65	0.78	1.01	1281	2450
ckappa[4,3]	0.09	0.09	0.02	0.02	0.06	0.11	1.01	745	676
comegaT[1]	0.00	0.00	0.00	0.00	0.00	0.00	1.00	1656	2300
comegaT[2]	0.01	0.01	0.00	0.00	0.01	0.01	1.00	2194	2711
comegaT[3]	0.00	0.00	0.00	0.00	0.00	0.00	1.01	1408	2289
comegaT[4]	0.00	0.00	0.00	0.00	0.00	0.00	1.00	2595	2298
crho[1,1]	1.00	1.00	0.00	0.00	1.00	1.00	NA	NA	NA
crho[2,1]	0.57	0.57	0.07	0.07	0.44	0.68	1.00	1083	1700
crho[3,1]	0.78	0.78	0.04	0.04	0.71	0.84	1.01	1015	1731
crho[4,1]	0.72	0.72	0.05	0.05	0.64	0.79	1.00	1300	2030
crho[1,2]	0.57	0.57	0.07	0.07	0.44	0.68	1.00	1083	1700
crho[2,2]	1.00	1.00	0.00	0.00	1.00	1.00	NA	NA	NA
crho[3,2]	0.55	0.55	0.07	0.07	0.42	0.67	1.00	1242	1933
crho[4,2]	0.55	0.56	0.07	0.07	0.43	0.66	1.00	1383	2292
crho[1,3]	0.78	0.78	0.04	0.04	0.71	0.84	1.01	1015	1731
crho[2,3]	0.55	0.55	0.07	0.07	0.42	0.67	1.00	1242	1933
crho[3,3]	1.00	1.00	0.00	0.00	1.00	1.00	NA	NA	NA
crho[4,3]	0.92	0.92	0.02	0.02	0.89	0.94	1.00	1324	2429
crho[1,4]	0.72	0.72	0.05	0.05	0.64	0.79	1.00	1300	2030
crho[2,4]	0.55	0.56	0.07	0.07	0.43	0.66	1.00	1383	2292
crho[3,4]	0.92	0.92	0.02	0.02	0.89	0.94	1.00	1324	2429
crho[4,4]	1.00	1.00	0.00	0.00	1.00	1.00	NA	NA	NA
clogmsigma[1]	0.20	0.20	0.02	0.02	0.17	0.22	1.00	2901	3087
clogmsigma[2]	-0.01	-0.01	0.02	0.02	-0.05	0.02	1.00	3536	3376

clogmsigma[3]	-0.26	-0.26	0.02	0.02	-0.29	-0.22	1.00	3010	3356
clogmsigma[4]	-0.10	-0.10	0.02	0.02	-0.13	-0.07	1.00	3325	3444
cssigma[1]	0.07	0.07	0.03	0.03	0.02	0.11	1.01	574	1063
cssigma[2]	0.17	0.17	0.02	0.02	0.14	0.20	1.00	1761	2633
cssigma[3]	0.16	0.16	0.02	0.02	0.13	0.19	1.00	1919	2850
cssigma[4]	0.08	0.08	0.03	0.02	0.03	0.12	1.01	560	769

The summary can also be extracted for individual parameters. Here presented for analyte 9 (Baclofen)

```
# which(unique(data$METID)==9) - 3rd compound in stan
fit$print(c("logkwx[3,1,1]", "logkwx[3,1,2]", "logkwx[3,1,3]",
           "logkwx[3,2,1]", "logkwx[3,2,2]", "logkwx[3,2,3]",
           "S1x[3,1,1,1]", "S1x[3,1,1,2]", "S1x[3,1,1,3]",
           "S1x[3,2,1,1]", "S1x[3,2,1,2]", "S1x[3,2,1,3]",
           "S1x[3,1,2,1]", "S1x[3,1,2,2]", "S1x[3,1,2,3]",
           "S1x[3,2,2,1]", "S1x[3,2,2,2]", "S1x[3,2,2,3]",
           "apHx[3,1,1]", "apHx[3,1,2]", "apHx[3,1,3]",
           "pKawx[3,1]", "pKawx[3,2]",
           "alphax[3,1,1]", "alphax[3,1,2]",
           "alphax[3,2,1]", "alphax[3,2,2]",
           "S2x[1,1]", "S2x[2,1]",
           "sigmax[3,1]", "sigmax[3,2]"), max_rows = 31)
```

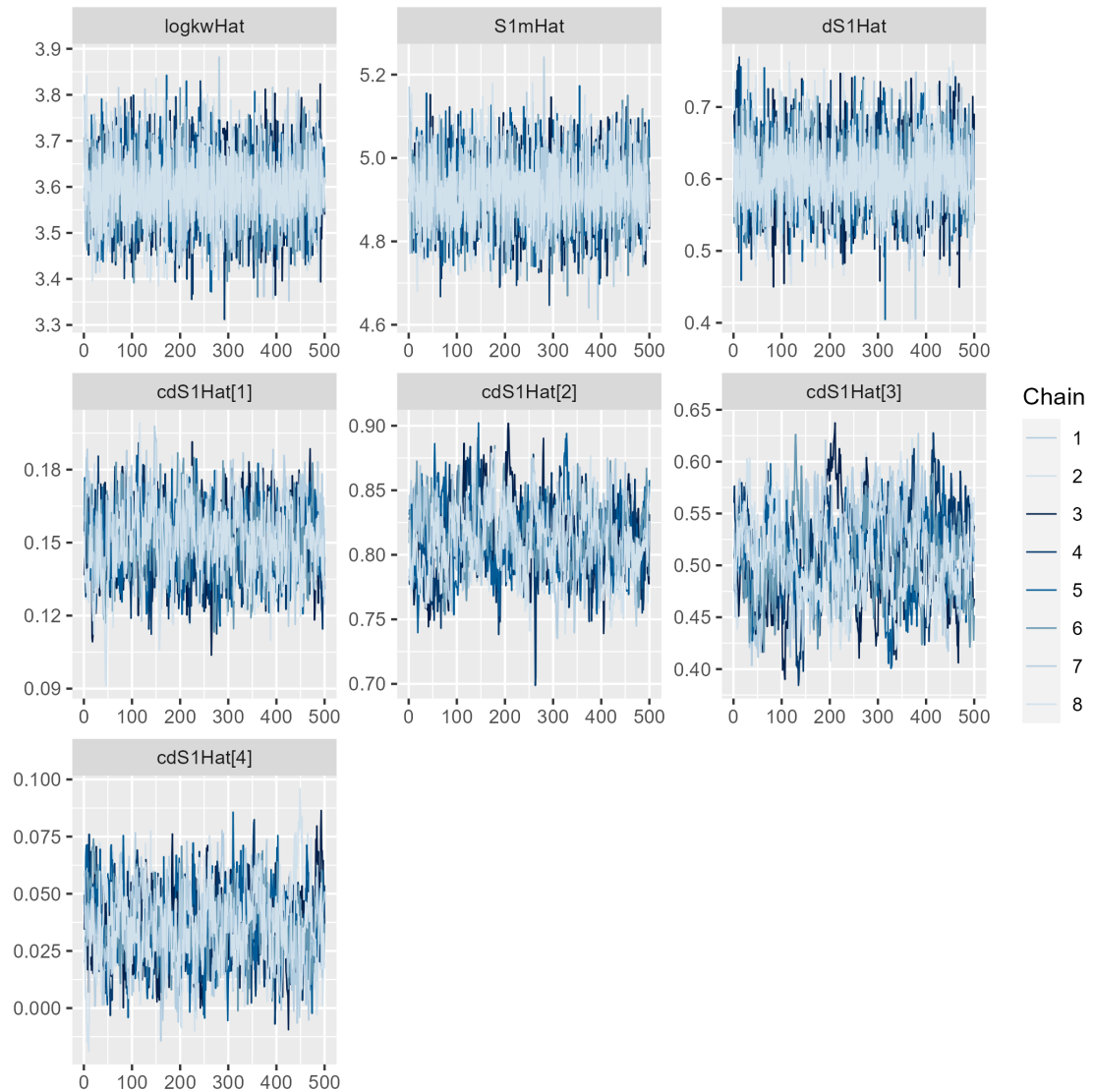
variable	mean	median	sd	mad	q5	q95	rhat	ess_bulk	ess_tail
logkwx[3,1,1]	1.33	1.33	0.03	0.03	1.28	1.38	1.00	6062	3190
logkwx[3,1,2]	0.68	0.68	0.07	0.07	0.58	0.79	1.00	5550	3342
logkwx[3,1,3]	0.68	0.68	0.07	0.07	0.58	0.79	1.00	5550	3342
logkwx[3,2,1]	1.64	1.64	0.04	0.04	1.58	1.70	1.00	6117	3234
logkwx[3,2,2]	1.04	1.04	0.06	0.06	0.94	1.14	1.00	6694	3794
logkwx[3,2,3]	1.04	1.04	0.06	0.06	0.94	1.14	1.00	6694	3794
S1x[3,1,1,1]	3.78	3.79	0.31	0.31	3.28	4.28	1.00	5248	2966
S1x[3,1,1,2]	4.74	4.72	0.71	0.71	3.59	5.95	1.00	6928	3011
S1x[3,1,1,3]	4.74	4.72	0.71	0.71	3.59	5.95	1.00	6928	3011
S1x[3,2,1,1]	3.35	3.35	0.26	0.27	2.92	3.78	1.00	5914	3292
S1x[3,2,1,2]	5.74	5.74	0.63	0.63	4.73	6.78	1.00	6900	3283
S1x[3,2,1,3]	5.74	5.74	0.63	0.63	4.73	6.78	1.00	6900	3283
S1x[3,1,2,1]	4.53	4.53	0.31	0.32	4.02	5.03	1.00	5363	3001
S1x[3,1,2,2]	5.48	5.46	0.72	0.71	4.30	6.69	1.00	7097	3287
S1x[3,1,2,3]	5.48	5.46	0.72	0.71	4.30	6.69	1.00	7097	3287
S1x[3,2,2,1]	4.20	4.20	0.27	0.28	3.76	4.62	1.00	5462	3331
S1x[3,2,2,2]	6.62	6.62	0.64	0.64	5.58	7.68	1.00	6955	3383

S1x[3,2,2,3]	6.62	6.62	0.64	0.64	5.58	7.68	1.00	6955	3383
apHx[3,1,1]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
apHx[3,1,2]	-0.03	-0.03	0.00	0.00	-0.03	-0.03	1.00	4583	3724
apHx[3,1,3]	-0.03	-0.03	0.00	0.00	-0.03	-0.03	1.00	4583	3724
pKawx[3,1]	6.74	6.74	0.08	0.08	6.61	6.86	1.00	8439	3301
pKawx[3,2]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
alphax[3,1,1]	2.14	2.14	0.90	0.90	0.67	3.64	1.00	8395	3357
alphax[3,1,2]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
alphax[3,2,1]	2.49	2.47	1.14	1.11	0.59	4.39	1.00	7846	3292
alphax[3,2,2]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
S2x[1,1]	0.49	0.49	0.02	0.02	0.47	0.52	1.02	432	986
S2x[2,1]	1.30	1.30	0.03	0.03	1.26	1.34	1.02	355	754
sigmax[3,1]	0.39	0.39	0.03	0.03	0.34	0.45	1.00	4626	3538
sigmax[3,2]	0.45	0.45	0.04	0.04	0.38	0.52	1.00	2663	3556

6.2 Trace plots

Several trace plots are shown.

```
bayesplot::mcmc_trace(fit$draws(c("logkwHat","S1mHat","dS1Hat","cdS1Hat")))
```



6.3 Summary of model parameters (figures)

First plot presents parameter specific for XBridge Shield RP18 column and parameters common for all the columns:

```
p1<-bayesplot::mcmc_intervals(fit$draws(c("logkwHat", "S1mHat", "dS1Hat", "dlogkwHat",
                                             "dlogkTHat", "dS1mHat", "ddS1Hat", "logS2mHat", "dlogS2Hat",
                                             "apH", "alphamHat", "dalphahat")), point_size = 2)
```

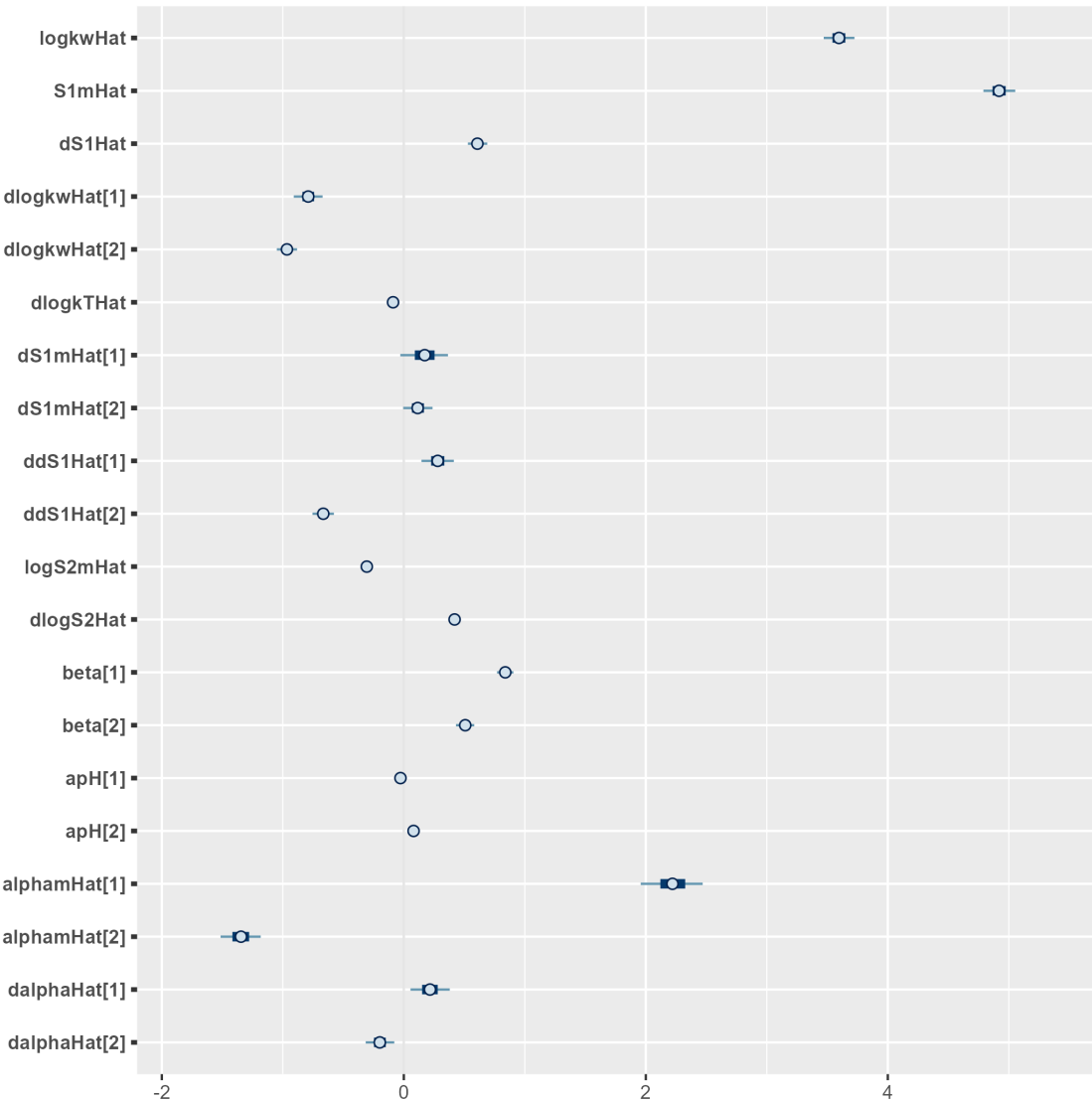
```

p2<-bayesplot::mcmc_intervals(fit$draws(c("omega","omegaT","kappa","msigma","ssigma", "tau

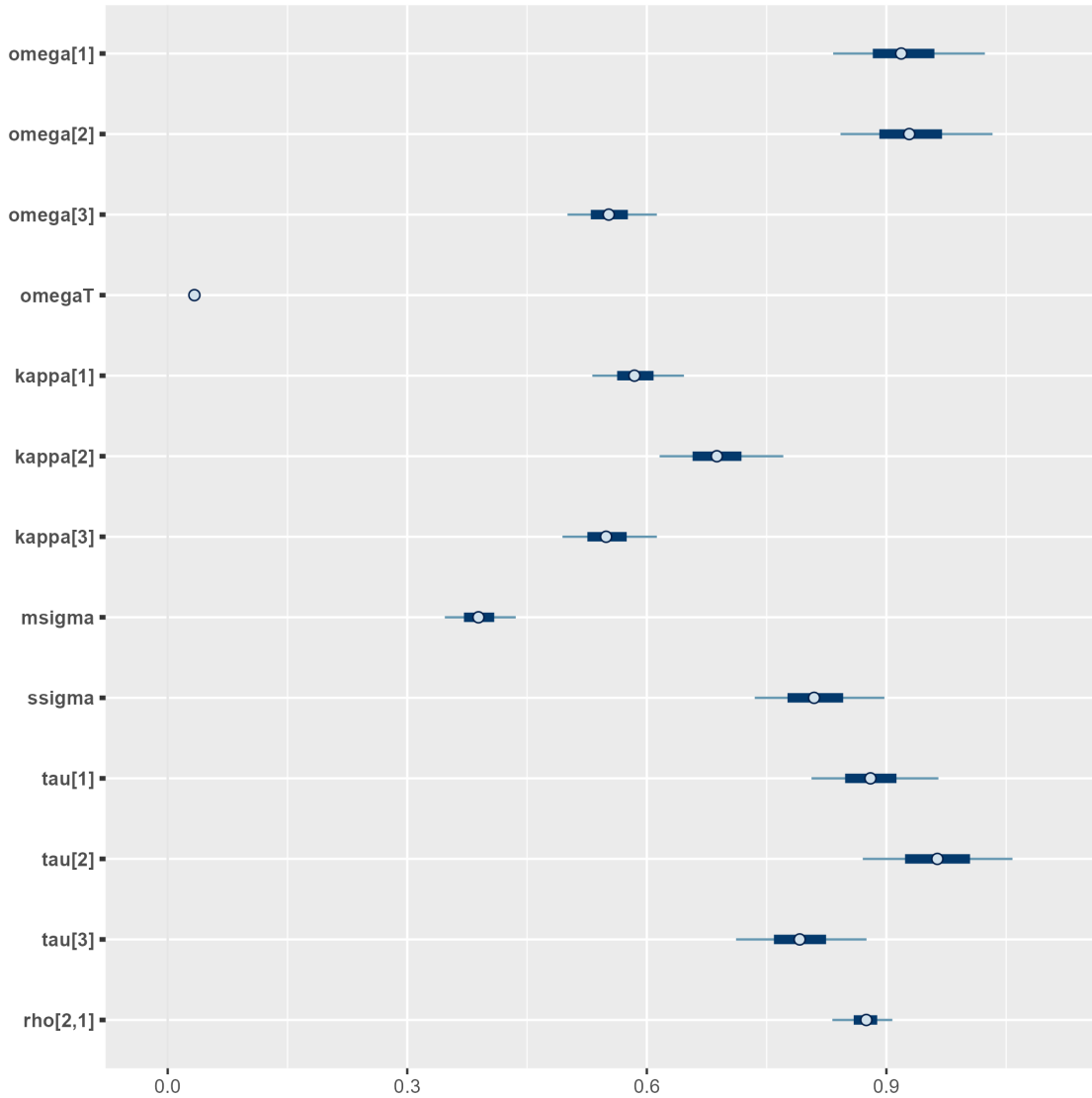
ggsave(paste0("figures\\param\\", "Paramteres.XBridgeShieldRP18_1", ".png"), plot=p1, width=10, height=10)
ggsave(paste0("figures\\param\\", "Paramteres.XBridgeShieldRP18_2", ".png"), plot=p2, width=10, height=10)

print(p1)

```



```
print(p2)
```



Column effects relative to XBridge Shield RP18 column:

```
p1<-bayesplot::mcmc_intervals(fit$draws(c("clogkwHat[1]", "cS1mHat[1]", "cdS1Hat[1]", "cdlogkwHat[1,1]", "cdlogkwHat[1,2]", "cdS1mHat[1,1]", "cdS1mHat[1,2]", "cddS1Hat[1,1]", "cddS1Hat[1,2]", "cbeta[1,1]", "cbeta[1,2]", "cdlogkTHat[1]", "capH[1,1]", "capH[1,2]")), point_size = 2)+
```

```

  theme(plot.title = element_text(size = 8))+  

  scale_y_discrete(labels = c("clogkwHat[1]"=="clogkwHat[c]",  

    "cS1mHat[1]"=="cS1mHat[c]",  

    "cdS1Hat[1]"=="cdS1Hat[c]",  

    "cdlogkwHat[1,1]"=="cdlogkwHat[c,1]",  

    "cdlogkwHat[1,2]"=="cdlogkwHat[c,2]",  

    "cdS1mHat[1,1]"= "cdS1mHat[c,1]",  

    "cdS1mHat[1,2]"=="cdS1mHat[c,2]",  

    "cddS1Hat[1,1]"=="cddS1Hat[c,1]",  

    "cddS1Hat[1,2]"=="cddS1Hat[c,2]",  

    "cbeta[1,1]"=="cbeta[c,1]",  

    "cbeta[1,2]"=="cbeta[c,2]",  

    "cdlogkTHat[1]"=="cdlogkTHat[c]",  

    "capH[1,1]"=="capH[c,1]",  

    "capH[1,2]"=="capH[c,2]"),  

  limits = c("capH[1,2]",  

    "capH[1,1]",  

    "cdlogkTHat[1]",  

    "cbeta[1,2]",  

    "cbeta[1,1]",  

    "cddS1Hat[1,2]",  

    "cddS1Hat[1,1]",  

    "cdS1mHat[1,2]",  

    "cdS1mHat[1,1]",  

    "cdlogkwHat[1,2]",  

    "cdlogkwHat[1,1]",  

    "cdS1Hat[1]",  

    "cS1mHat[1]",  

    "clogkwHat[1]"))+  

  xlim(-0.5,1)  

p2<-bayesplot::mcmc_intervals(fit$draws(c("clogkwHat[2]", "cS1mHat[2]", "cdS1Hat[2]",  

  "cdlogkwHat[2,1]", "cdlogkwHat[2,2]",  

  "cdS1mHat[2,1]", "cdS1mHat[2,2]",  

  "cddS1Hat[2,1]", "cddS1Hat[2,2]",  

  "cbeta[2,1]", "cbeta[2,2]",  

  "cdlogkTHat[2]",  

  "capH[2,1]", "capH[2,2]")), point_size = 2)+  

  theme(axis.text.y=element_blank(), plot.title = element_text(size = 8))+  

  xlim(-0.5,1)

```

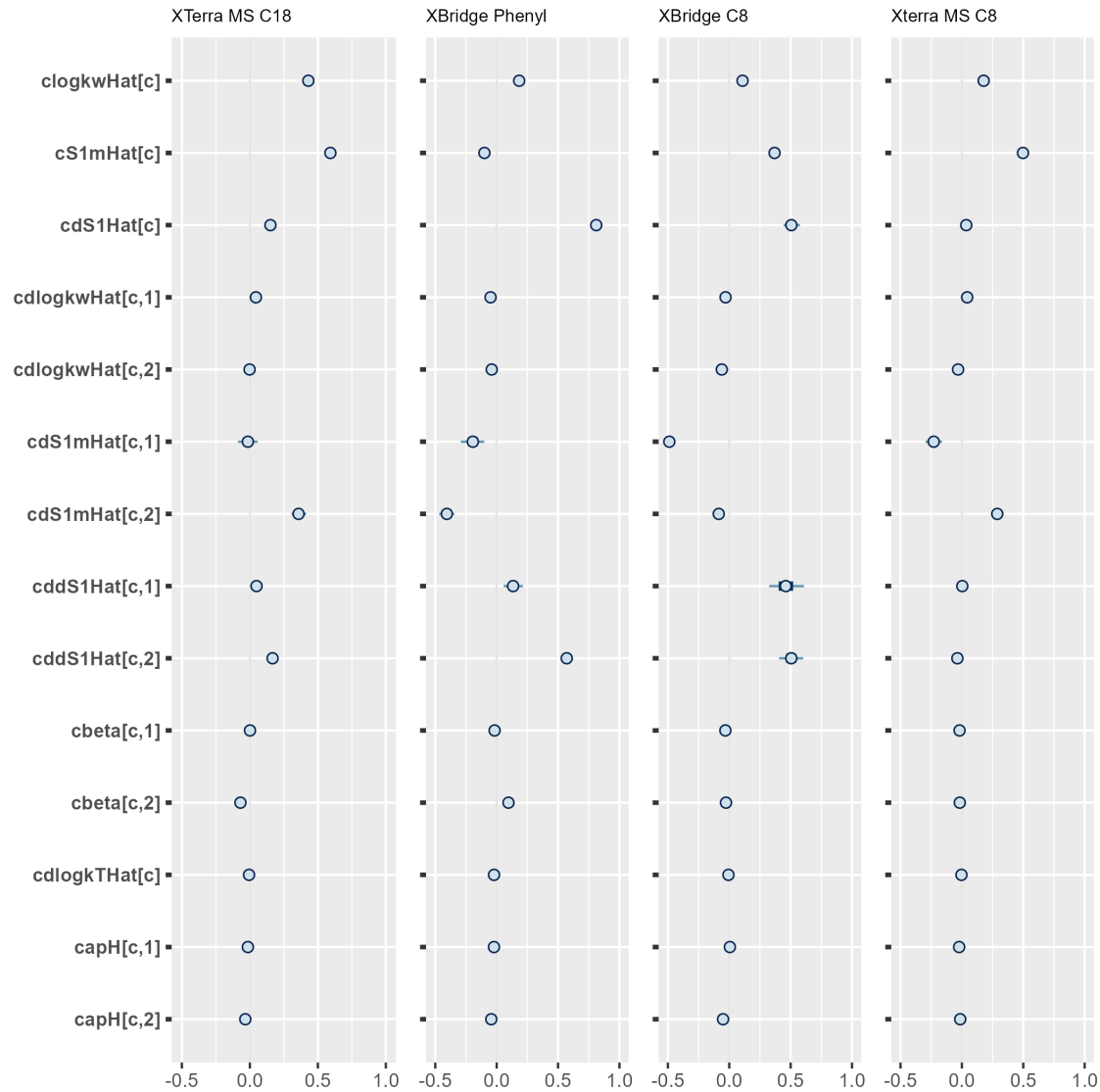
```

p3<-bayesplot::mcmc_intervals(fit$draws(c("clogkwHat[3]", "cS1mHat[3]", "cdS1Hat[3]",
  "cdlogkwHat[3,1]", "cdlogkwHat[3,2]",
  "cdS1mHat[3,1]", "cdS1mHat[3,2]",
  "cddS1Hat[3,1]", "cddS1Hat[3,2]",
  "cbeta[3,1]", "cbeta[3,2]",
  "cdlogkTHat[3]",
  "capH[3,1]", "capH[3,2]")), point_size = 2) +
  theme(axis.text.y=element_blank(), plot.title = element_text(size = 8)) +
  xlim(-0.5,1)

p4<-bayesplot::mcmc_intervals(fit$draws(c("clogkwHat[4]", "cS1mHat[4]", "cdS1Hat[4]",
  "cdlogkwHat[4,1]", "cdlogkwHat[4,2]",
  "cdS1mHat[4,1]", "cdS1mHat[4,2]",
  "cddS1Hat[4,1]", "cddS1Hat[4,2]",
  "cbeta[4,1]", "cbeta[4,2]",
  "cdlogkTHat[4]",
  "capH[4,1]", "capH[4,2]")), point_size = 2) +
  theme(axis.text.y=element_blank(), plot.title = element_text(size = 8)) +
  xlim(-0.5,1)

p=grid.arrange(p1+ggtitle("XTerra MS C18"),
  p2+ggtitle("XBridge Phenyl"),
  p3+ggtitle("XBridge C8"),
  p4+ggtitle("Xterra MS C8"), ncol=4, widths=c(1.75,1,1,1))

```



```
ggsave(paste0("figures\\param\\", "Difference_1", ".png"), plot=p, width = 20, height = 20)
```

```
p1<-bayesplot::mcmc_intervals(fit$draws(c("comega[1,1]",
"comega[1,2]",
"comega[1,3]",
"comegaT[1]",
"ckappa[1,1]",
"ckappa[1,2]",
```

```

"ckappa[1,3]")), point_size = 2) +
theme(plot.title = element_text(size = 8))+
scale_y_discrete(labels = c("comega[1,1]"=="comega[c,1]",
                            "comega[1,2]"=="comega[c,2]",
                            "comega[1,3]"=="comega[c,3]",
                            "comegaT[1]" = "comegaT[c]",
                            "ckappa[1,1]"=="ckappa[c,1]",
                            "ckappa[1,2]"=="ckappa[c,2]",
                            "ckappa[1,3]"=="ckappa[c,3]"),
limits = c("ckappa[1,3]",
           "ckappa[1,2]",
           "ckappa[1,1]",
           "comegaT[1]",
           "comega[1,3]",
           "comega[1,2]",
           "comega[1,1]"))+
xlim(0,0.8)

p2<-bayesplot::mcmc_intervals(fit$draws(c("comega[2,1]",
                                             "comega[2,2]",
                                             "comega[2,3]",
                                             "comegaT[2]",
                                             "ckappa[2,1]",
                                             "ckappa[2,2]",
                                             "ckappa[2,3]")), point_size = 2)+
theme(axis.text.y=element_blank(),plot.title = element_text(size = 8))+
xlim(0,0.8)

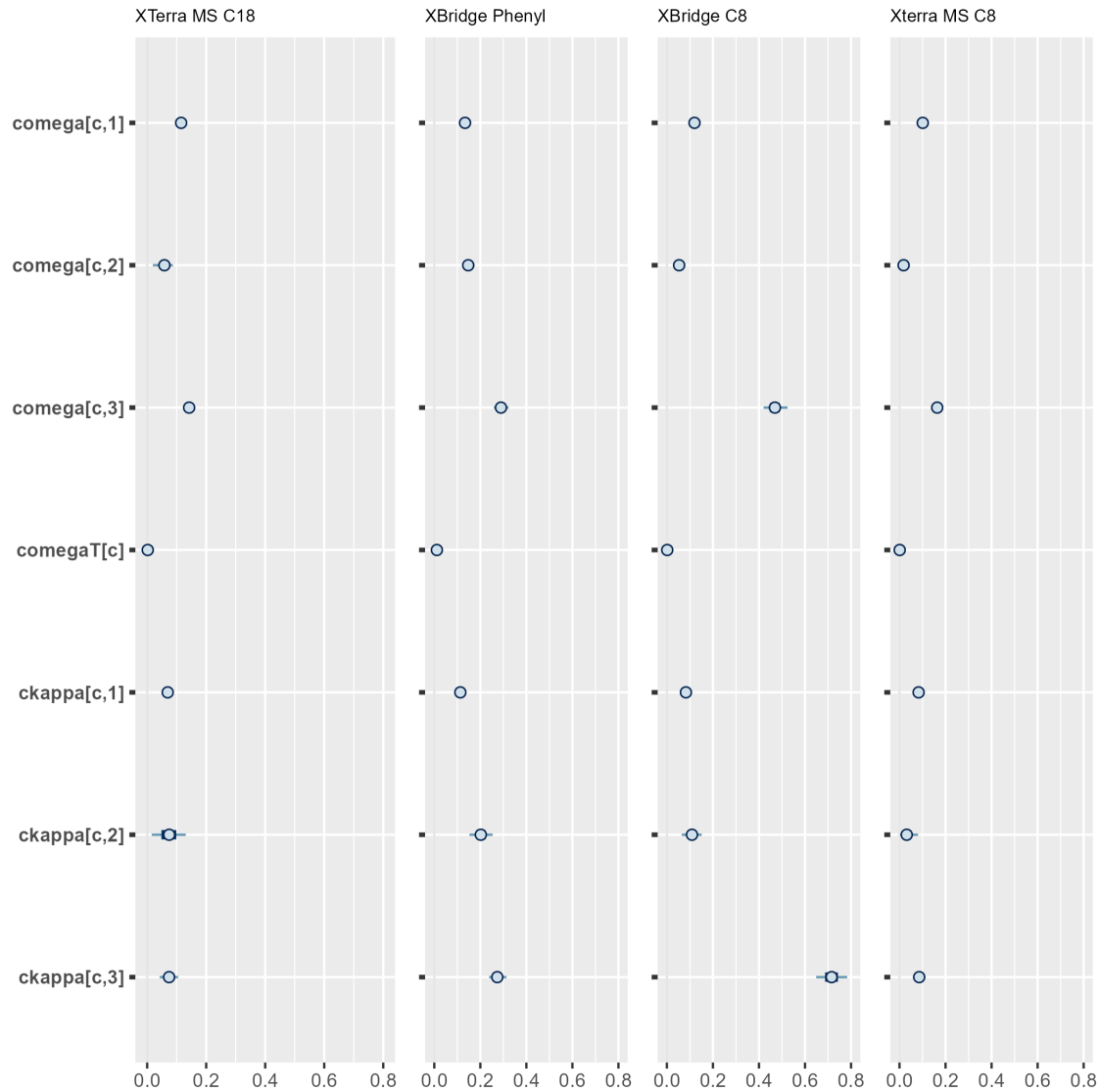
p3<-bayesplot::mcmc_intervals(fit$draws(c("comega[3,1]",
                                             "comega[3,2]",
                                             "comega[3,3]",
                                             "comegaT[3]",
                                             "ckappa[3,1]",
                                             "ckappa[3,2]",
                                             "ckappa[3,3]")), point_size = 2)+
theme(axis.text.y=element_blank(),plot.title = element_text(size = 8))+
xlim(0,0.8)

p4<-bayesplot::mcmc_intervals(fit$draws(c("comega[4,1]",
                                             "comega[4,2]",
                                             "comega[4,3]",
                                             "comega[4,4]")), point_size = 2)+
theme(axis.text.y=element_blank(),plot.title = element_text(size = 8))+
xlim(0,0.8)

```

```
"comegaT[4]",
"ckappa[4,1]",
"ckappa[4,2]",
"ckappa[4,3]"), point_size = 2) +
theme(axis.text.y=element_blank(), plot.title = element_text(size = 8)) +
xlim(0,0.8)

p=grid.arrange(p1+ggtitle("XTerra MS C18"),
                p2+ggtitle("XBridge Phenyl"),
                p3+ggtitle("XBridge C8"),
                p4+ggtitle("Xterra MS C8"), ncol=4, widths=c(1.75,1,1,1))
```



```
ggsave(paste0("figures\\param\\", "Difference_2", ".png"), plot=p, width = 20, height = 20)
```

Parameters characterizing columns:

```
draws_df_subset <- fit$draws(format = "df", variable = c("clogkwHat", "cdlogkwHat"))

Neutral_results <- draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(clogkwHat[c]) %>%
```

```

  mutate(ColumnName=recode(c,
                           '1' = 'XTerra MS C18',
                           '2' = 'XBridge Phenyl',
                           '3' = 'XBridge C8',
                           '4' = 'Xterra MS C8'))%>%
  group_by(ColumnName)%>%
  tidybayes::median_qi(logkwHat = clogkwHat) %>%
  mutate(Type = "N")

Acids_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(clogkwHat[c],cdlogkwHat[c,r]) %>%
  filter(r==1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(xlogkwHat=clogkwHat+cdlogkwHat)%>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnName=recode(c,
                           '1' = 'XTerra MS C18',
                           '2' = 'XBridge Phenyl',
                           '3' = 'XBridge C8',
                           '4' = 'Xterra MS C8'))%>%
  group_by(ColumnName)%>%
  tidybayes::median_qi(logkwHat = xlogkwHat) %>%
  mutate(Type = "A")

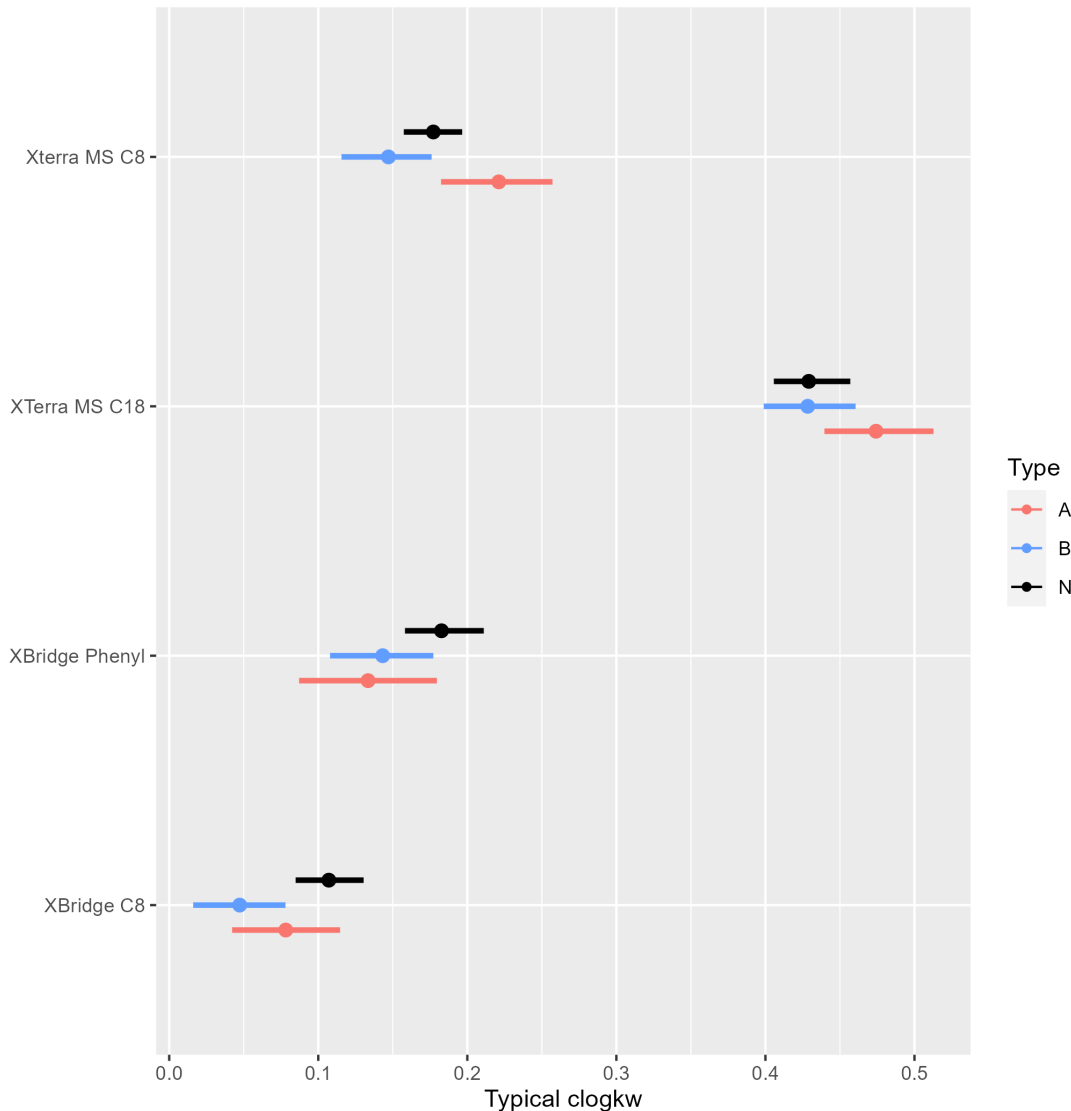
Basic_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(clogkwHat[c],cdlogkwHat[c,r]) %>%
  filter(r==2)%>%
  ungroup() %>%
  select(-r) %>%
  mutate(xlogkwHat=clogkwHat+cdlogkwHat)%>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnName=recode(c,
                           '1' = 'XTerra MS C18',
                           '2' = 'XBridge Phenyl',
                           '3' = 'XBridge C8',
                           '4' = 'Xterra MS C8'))%>%
  group_by(ColumnName)%>%
  tidybayes::median_qi(logkwHat = xlogkwHat) %>%

```

```
mutate(Type = "B")

p1<-bind_rows(Neutral_results, Acids_results, Basic_results) %>%
  ggplot(aes(y = ColumnNames, x = logkwHat, xmin = .lower, xmax = .upper,color = Type)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("A", "B", "N"), values = c("#F8766D", "#619CFF", "black")) +
  ylab(' ') +
  xlab("Typical clogkw")

print(p1)
```



```

'1' = 'XTerra MS C18',
'2' = 'XBridge Phenyl',
'3' = 'XBridge C8',
'4' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%
tidybayes::median_qi(S1Hat = xS1mHat) %>%
mutate(Type = "N")%>%
mutate(Modifier="MeOH")

MeOH_Acids_results <-draws_df_subset %>%
slice_sample(n=1000) %>%
tidybayes::spread_draws(cS1mHat[c],cdS1mHat[c,r]) %>%
filter(r==1) %>%
ungroup() %>%
select(-r) %>%
mutate(xS1mHat=cS1mHat+cdS1mHat)%>%
mutate_if(is.character,as.factor) %>%
mutate(ColumnName=recode(c,
'1' = 'XTerra MS C18',
'2' = 'XBridge Phenyl',
'3' = 'XBridge C8',
'4' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%
tidybayes::median_qi(S1Hat = xS1mHat) %>%
mutate(Type = "A")%>%
mutate(Modifier="MeOH")

MeOH_Basic_results <-draws_df_subset %>%
slice_sample(n=1000) %>%
tidybayes::spread_draws(cS1mHat[c],cdS1mHat[c,r]) %>%
filter(r==2) %>%
ungroup() %>%
select(-r) %>%
mutate(xS1mHat=cS1mHat+cdS1mHat)%>%
mutate_if(is.character,as.factor) %>%
mutate(ColumnName=recode(c,
'1' = 'XTerra MS C18',
'2' = 'XBridge Phenyl',
'3' = 'XBridge C8',
'4' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%

```

```

tidybayes::median_qi(S1Hat = xS1mHat) %>%
  mutate(Type = "B") %>%
  mutate(Modifier="MeOH")

ACN_Neutral_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(cS1mHat[c],cdS1Hat[c]) %>%
  mutate(xS1aHat=cS1mHat+cdS1Hat)%>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(c,
                            '1' = 'XTerra MS C18',
                            '2' = 'XBridge Phenyl',
                            '3' = 'XBridge C8',
                            '4' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(S1Hat = xS1aHat) %>%
  mutate(Type = "N") %>%
  mutate(Modifier="ACN")

ACN_Acids_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(cS1mHat[c],cdS1Hat[c],cdS1mHat[c,r],cddS1Hat[c,r]) %>%
  filter(r==1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(xS1aHat= cS1mHat +cdS1mHat +cdS1Hat + cddS1Hat)%>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(c,
                            '1' = 'XTerra MS C18',
                            '2' = 'XBridge Phenyl',
                            '3' = 'XBridge C8',
                            '4' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(S1Hat = xS1aHat) %>%
  mutate(Type = "A") %>%
  mutate(Modifier="ACN")

ACN_Basic_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(cS1mHat[c],cdS1Hat[c],cdS1mHat[c,r],cddS1Hat[c,r]) %>%

```

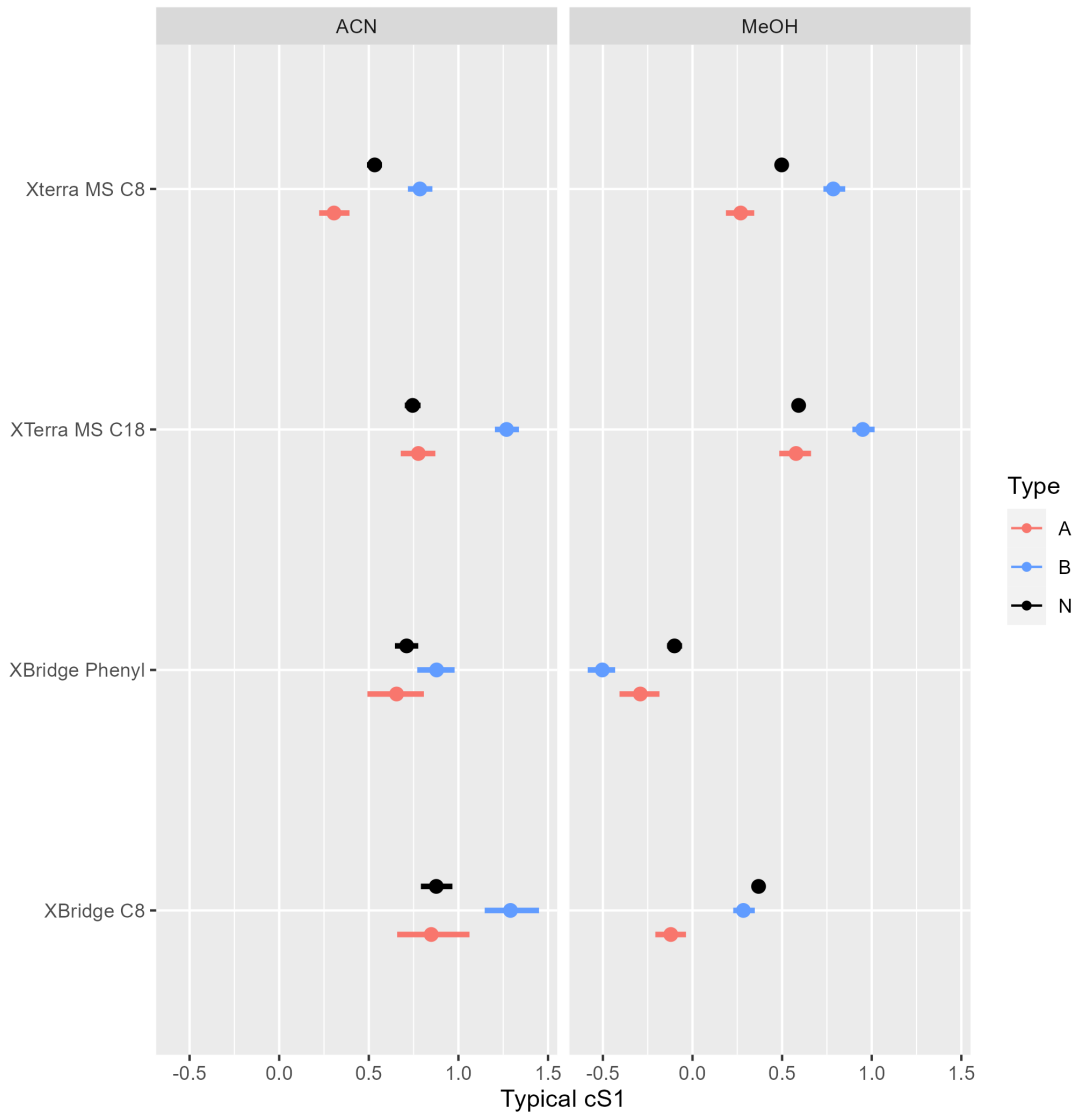
```

filter(r==2) %>%
ungroup() %>%
select(-r) %>%
mutate(xS1aHat= cS1mHat+cdS1mHat+cdS1Hat + cddS1Hat)%>%
mutate_if(is.character,as.factor) %>%
mutate(ColumnNames=recode(c,
  '1' = 'XTerra MS C18',
  '2' = 'XBridge Phenyl',
  '3' = 'XBridge C8',
  '4' = 'Xterra MS C8'))%>%
group_by(ColumnNames)%>%
tidybayes::median_qi(S1Hat = xS1aHat) %>%
mutate(Type = "B")%>%
mutate(Modifier="ACN")

p2<-bind_rows(MeOH_Neutral_results, MeOH_Acids_results, MeOH_Basic_results,
  ACN_Neutral_results,ACN_Acids_results, ACN_Basic_results) %>%
ggplot(aes(y = ColumnNames, x = S1Hat, xmin = .lower, xmax = .upper,color = Type)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("A", "B", "N"), values = c("#F8766D", "#619CFF", "black")) +
  ylab(' ')+
  xlab("Typical cS1")+
  facet_wrap(.~Modifier)

print(p2)

```



```

draws_df_subset <- fit$draws(format = "df", variable = c("beta", "cbeta"))

logkw_results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(cbeta[c, r]) %>%
  filter(r == 1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(xbeta = cbeta) %>%

```

```

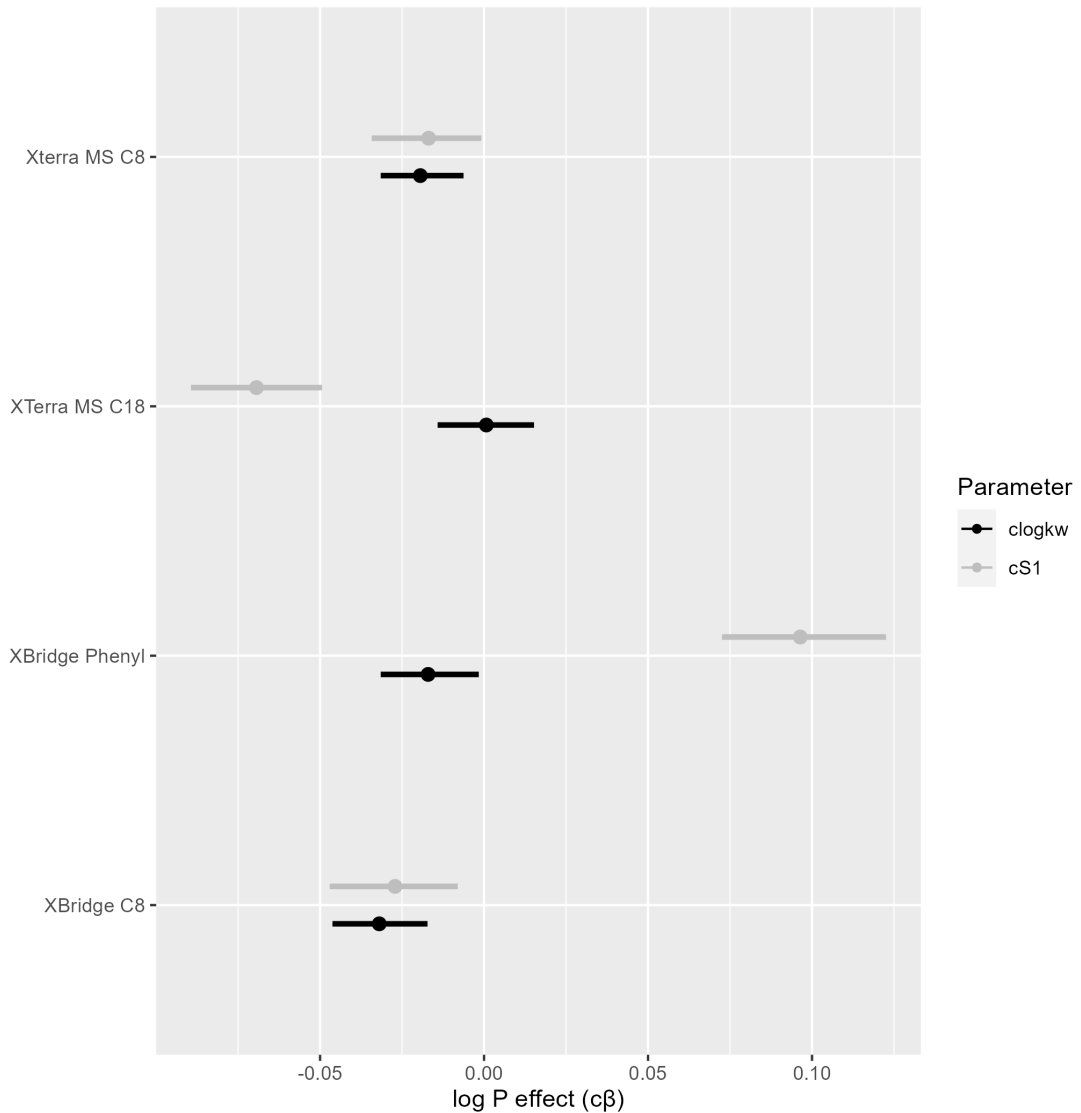
  mutate_if(is.character, as.factor) %>%
  mutate(ColumnName=recode(c,
    '1' = 'XTerra MS C18',
    '2' = 'XBridge Phenyl',
    '3' = 'XBridge C8',
    '4' = 'Xterra MS C8')) %>%
  group_by(ColumnName) %>%
  tidybayes::median_qi(beta = xbeta) %>%
  mutate(Parameter="logkw")

S1_results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(cbeta[c,r]) %>%
  filter(r==2) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(xbeta=cbeta) %>%
  mutate_if(is.character, as.factor) %>%
  mutate(ColumnName=recode(c,
    '1' = 'XTerra MS C18',
    '2' = 'XBridge Phenyl',
    '3' = 'XBridge C8',
    '4' = 'Xterra MS C8')) %>%
  group_by(ColumnName) %>%
  tidybayes::median_qi(beta = xbeta) %>%
  mutate(Parameter="S1")

p3<-bind_rows(logkw_results, S1_results) %>%
  ggplot(aes(y = ColumnName, x = beta, xmin = .lower, xmax = .upper, color = Parameter)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("clogkw", "cS1"), values = c("black", "gray")) +
  ylab(' ') +
  xlab("log P effect (c\u03b22)")

print(p3)

```



```

draws_df_subset <- fit$draws(format = "df", variable = c("apH", "capH"))

Acids_results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(capH[c, r]) %>%
  filter(r==1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(xapH=capH)%>%

```

```

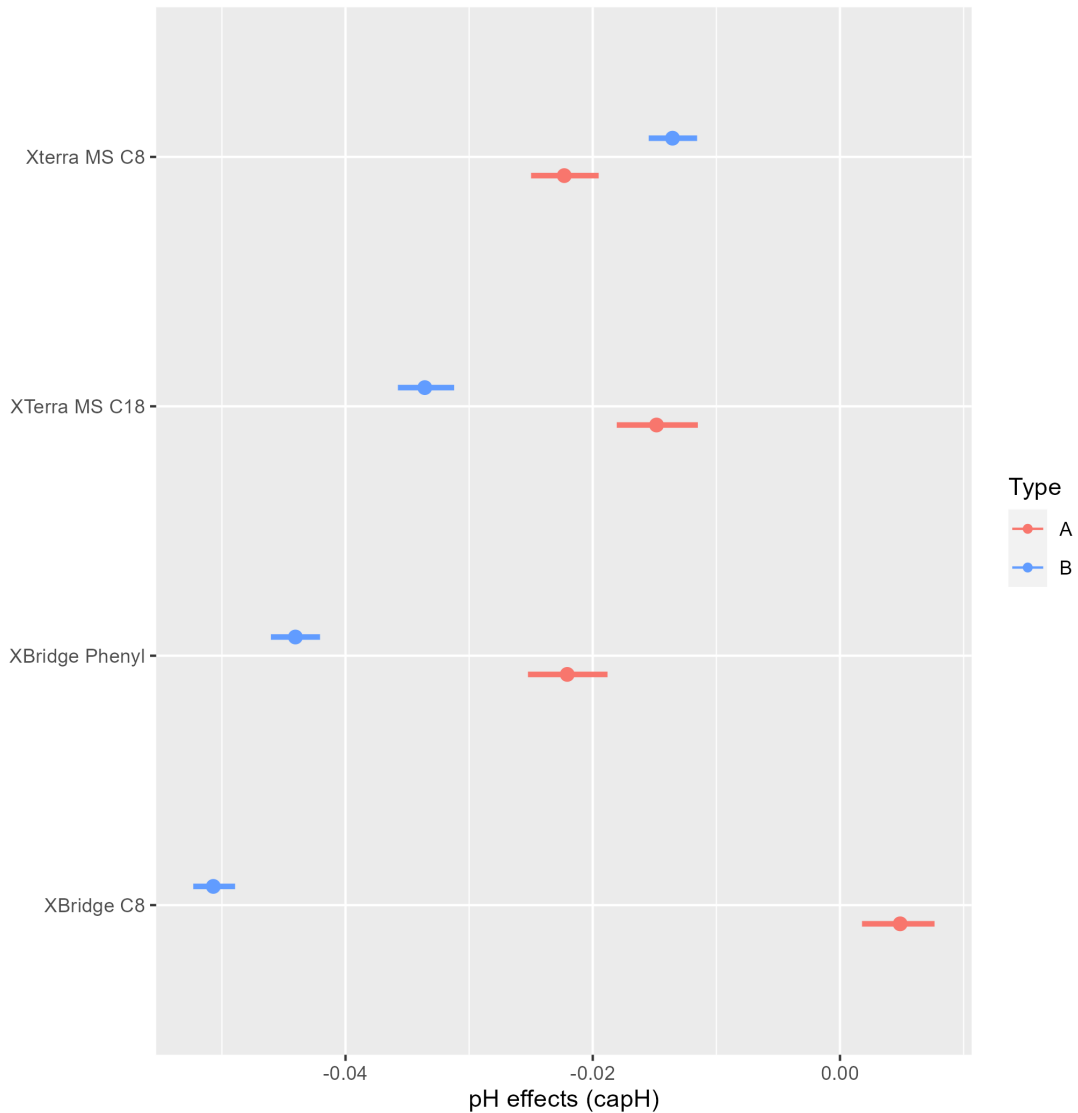
  mutate_if(is.character, as.factor) %>%
  mutate(ColumnNames=recode(c,
    '1' = 'XTerra MS C18',
    '2' = 'XBridge Phenyl',
    '3' = 'XBridge C8',
    '4' = 'Xterra MS C8')) %>%
  group_by(ColumnNames) %>%
  tidybayes::median_qi(apH = xapH) %>%
  mutate(Type="A")

Bases_results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(capH[c, r]) %>%
  filter(r==2) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(xapH=capH) %>%
  mutate_if(is.character, as.factor) %>%
  mutate(ColumnNames=recode(c,
    '1' = 'XTerra MS C18',
    '2' = 'XBridge Phenyl',
    '3' = 'XBridge C8',
    '4' = 'Xterra MS C8')) %>%
  group_by(ColumnNames) %>%
  tidybayes::median_qi(apH = xapH) %>%
  mutate(Type="B")

p4<-bind_rows(Acids_results, Bases_results) %>%
  ggplot(aes(y = ColumnNames, x = apH, xmin = .lower, xmax = .upper, color = Type)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("A", "B"), values = c("#F8766D", "#619cff")) +
  ylab(' ') +
  xlab("pH effects (capH)")

print(p4)

```



```

draws_df_subset <- fit$draws(format = "df", variable = c("dlogkTHat", "cdlogkTHat"))

Results <-draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(cdlogkTHat[c]) %>%
  ungroup() %>%
  mutate(xdlogkTHat=cdlogkTHat)%>%
  mutate_if(is.character, as.factor) %>%
  mutate(ColumnName=recode(c,

```

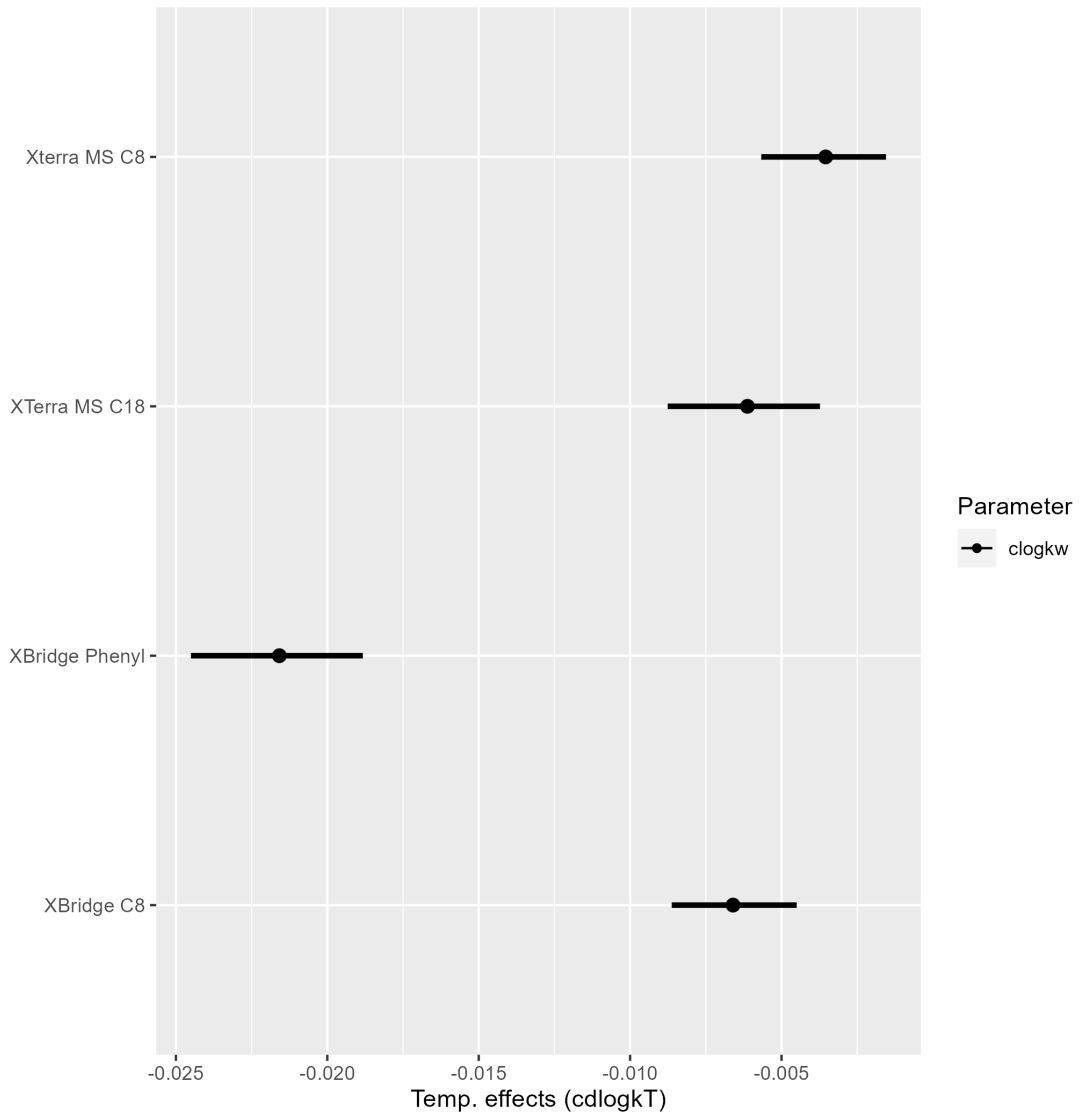
```

'1' = 'XTerra MS C18',
'2' = 'XBridge Phenyl',
'3' = 'XBridge C8',
'4' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%
tidybayes::median_qi(dlogkTHat = xdlogkTHat)%>%
mutate(Parameter="clogkw")

p5<-Results %>%
  ggplot(aes(y = ColumnName, x = dlogkTHat, xmin = .lower, xmax = .upper,color = Parameter,
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("clogkw"), values = c("black"))+
  ylab(' ')+
  xlab("Temp. effects (cdlogkT)"))

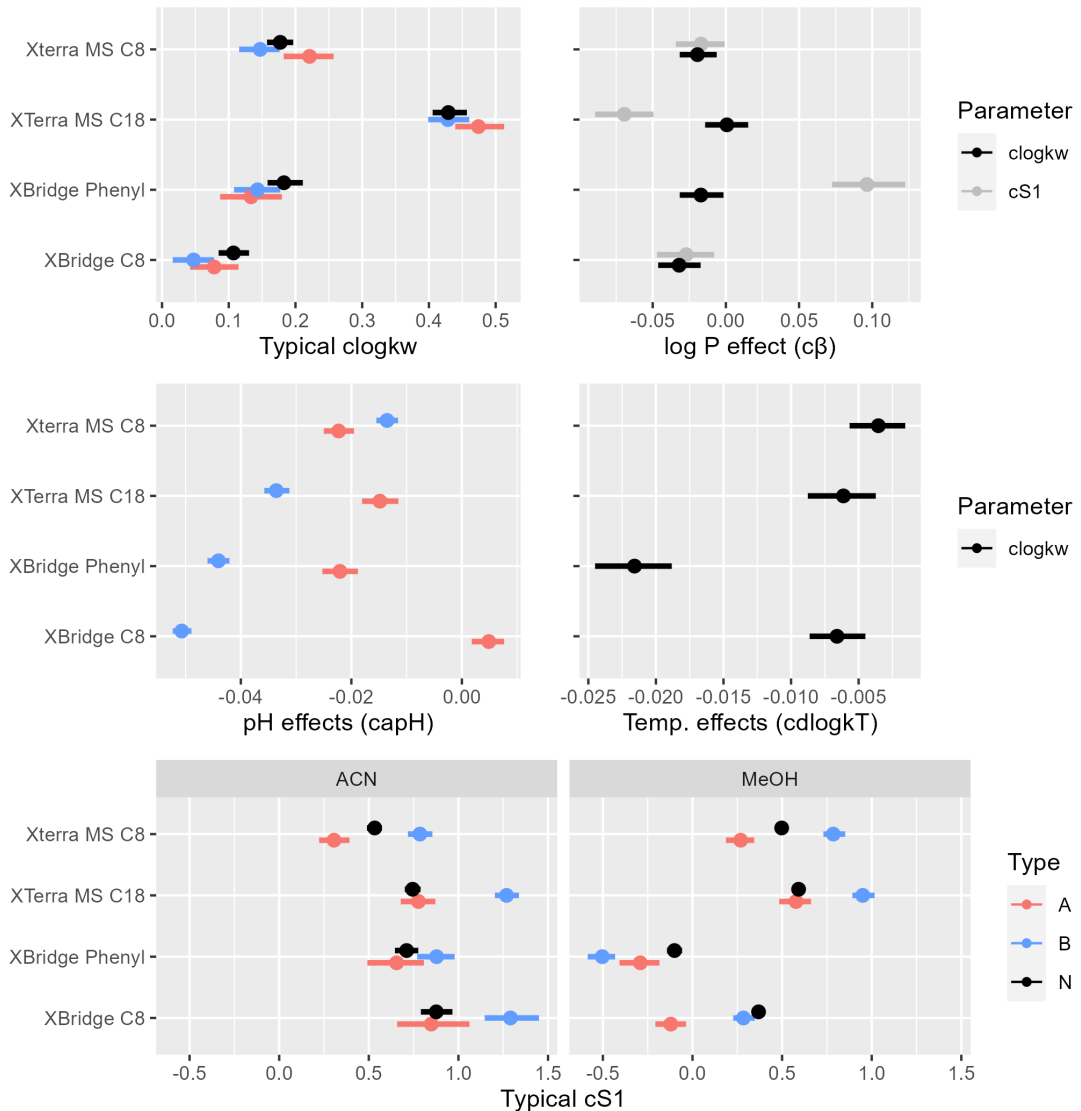
print(p5)

```



```

p=grid.arrange(p1+ theme(legend.position='none'),
               p3+theme(axis.text.y=element_blank()),
               p4+ theme(legend.position='none'),
               p5+theme(axis.text.y=element_blank()),
               p2, ncol=2, widths = c(1,1), layout_matrix = rbind(c(1, 2),c(3, 4), c(5, 5))
  )
  
```



```
ggsave(paste0("figures\\param\\", "Joined_Effects", ".png"), plot=p, width = 20, height =
```

Parameters characterizing columns:

```
draws_df_subset <- fit$draws(format = "df", variable = c("logkwHat", "clogkwHat", "dlogkwHat"))

Neutral_results <- draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwHat, clogkwHat[c]) %>%
```

```

  mutate(c=c+1)%>%
  mutate(clogkwHat=logkwHat+clogkwHat)%>%
  select(.draw,logkwHat,clogkwHat,c) %>%
  tidyrr::pivot_wider(names_from = c, values_from = clogkwHat) %>%
  rename(`1`=logkwHat) %>%
  select(-.draw)%>%
  tidyrr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xlogkwHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(logkwHat = xlogkwHat) %>%
  mutate(Type = "N")

Acids_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwHat,clogkwHat[c],dlogkwHat[r],cdlogkwHat[c,r]) %>%
  filter(r==1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(clogkwHat=logkwHat+clogkwHat+dlogkwHat+cdlogkwHat)%>%
  mutate(logkwHat=logkwHat+dlogkwHat)%>%
  select(.draw,logkwHat,clogkwHat,c) %>%
  tidyrr::pivot_wider(names_from = c, values_from = clogkwHat) %>%
  rename(`1`=logkwHat) %>%
  select(-.draw)%>%
  tidyrr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xlogkwHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(logkwHat = xlogkwHat) %>%

```

```

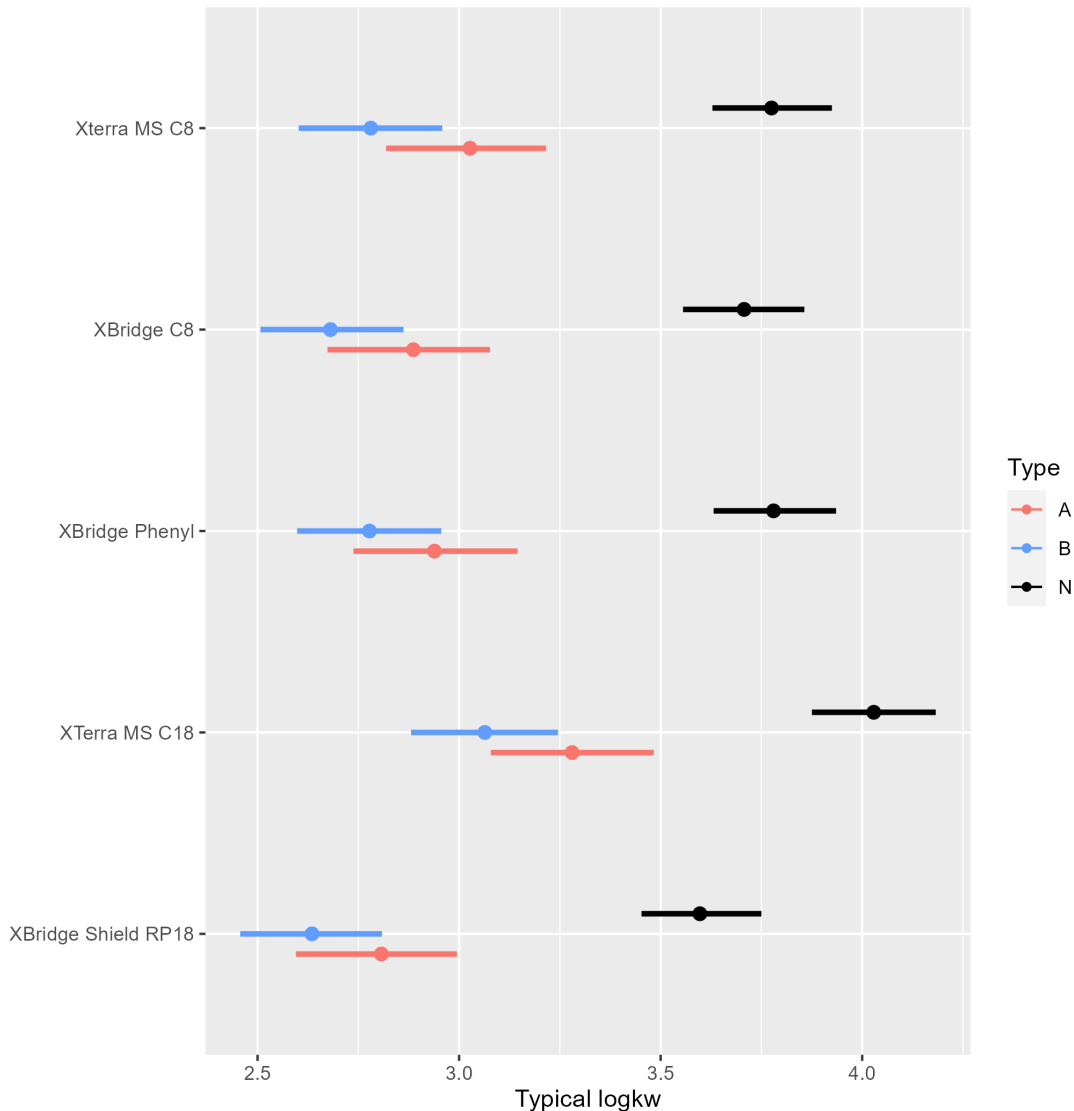
  mutate(Type = "A")

Basic_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwHat,clogkwHat[c],dlogkwHat[r],cdlogkwHat[c,r]) %>%
  filter(r==2)%>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(clogkwHat=logkwHat+clogkwHat+dlogkwHat+cdlogkwHat)%>%
  mutate(logkwHat=logkwHat+dlogkwHat)%>%
  select(.draw,logkwHat,clogkwHat,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = clogkwHat) %>%
  rename(`1`=logkwHat) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xlogkwHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(logkwHat = xlogkwHat) %>%
  mutate(Type = "B")

p1<-bind_rows(Neutral_results, Acids_results, Basic_results) %>%
  ggplot(aes(y = ColumnNames, x = logkwHat, xmin = .lower, xmax = .upper,color = Type)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("A", "B", "N"), values = c("#F8766D", "#619CFF", "black")) +
  ylab(' ') +
  xlab("Typical logkw")

print(p1)

```



```

tidyr::pivot_wider(names_from = c, values_from = cS1mHat) %>%
  rename(`1`=S1mHat) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xS1mHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(S1Hat = xS1mHat) %>%
  mutate(Type = "N")%>%
  mutate(Modifier="MeOH")

MeOH_Acids_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(S1mHat,cS1mHat[c],dS1mHat[r],cdS1mHat[c,r]) %>%
  filter(r==1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(cS1mHat=S1mHat+cS1mHat+dS1mHat+cdS1mHat)%>%
  mutate(S1mHat=S1mHat+dS1mHat)%>%
  select(.draw,S1mHat,cS1mHat,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = cS1mHat) %>%
  rename(`1`=S1mHat) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xS1mHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(S1Hat = xS1mHat) %>%
  mutate(Type = "A")%>%
  mutate(Modifier="MeOH")

```

```

MeOH_Basic_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(S1mHat,cS1mHat[c],dS1mHat[r],cdS1mHat[c,r]) %>%
  filter(r==2) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(cS1mHat=S1mHat+cS1mHat+dS1mHat+cdS1mHat)%>%
  mutate(S1mHat=S1mHat+dS1mHat)%>%
  select(.draw,S1mHat,cS1mHat,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = cS1mHat) %>%
  rename(`1`=S1mHat) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xS1mHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(S1Hat = xS1mHat) %>%
  mutate(Type = "B")%>%
  mutate(Modifier="MeOH")

ACN_Neutral_results <-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(S1mHat,cS1mHat[c],dS1Hat,cdS1Hat[c]) %>%
  mutate(c=c+1)%>%
  mutate(cS1aHat=S1mHat+cS1mHat+dS1Hat+cdS1Hat)%>%
  mutate(S1aHat =S1mHat+dS1Hat)%>%
  select(.draw,S1aHat,cS1aHat,c)%>%
  tidyr::pivot_wider(names_from = c, values_from = cS1aHat) %>%
  rename(`1`=S1aHat) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xS1aHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',

```

```

'2' = 'XTerra MS C18',
'3' = 'XBridge Phenyl',
'4' = 'XBridge C8',
'5' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%
tidybayes::median_qi(S1Hat = xS1aHat) %>%
mutate(Type = "N")%>%
mutate(Modifier="ACN")

ACN_Acids_results <-draws_df_subset %>%
slice_sample(n=1000) %>%
tidybayes::spread_draws(S1mHat,cS1mHat[c],dS1Hat,cdS1Hat[c],dS1mHat[r],cdS1mHat[c,r],ddS
filter(r==1) %>%
ungroup() %>%
select(-r) %>%
mutate(c=c+1)%>%
mutate(cS1aHat= S1mHat+cS1mHat + dS1mHat+cdS1mHat + dS1Hat+cdS1Hat + ddS1Hat+ cddS1Hat)%>%
mutate(S1aHat = S1mHat+dS1mHat+dS1Hat+ddS1Hat)%>%
select(.draw,S1aHat,cS1aHat,c) %>%
tidyr::pivot_wider(names_from = c, values_from = cS1aHat) %>%
rename(`1`=S1aHat) %>%
select(-.draw)%>%
tidyr::pivot_longer(c(1:5),names_to = "ColumnName", values_to = "xS1aHat") %>%
mutate_if(is.character,as.factor) %>%
mutate(ColumnName=recode(ColumnName,
'1' = 'XBridge Shield RP18',
'2' = 'XTerra MS C18',
'3' = 'XBridge Phenyl',
'4' = 'XBridge C8',
'5' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%
tidybayes::median_qi(S1Hat = xS1aHat) %>%
mutate(Type = "A")%>%
mutate(Modifier="ACN")

ACN_Basic_results <-draws_df_subset %>%
slice_sample(n=1000) %>%
tidybayes::spread_draws(S1mHat,cS1mHat[c],dS1Hat,cdS1Hat[c],dS1mHat[r],cdS1mHat[c,r],ddS
filter(r==2) %>%
ungroup() %>%
select(-r) %>%

```

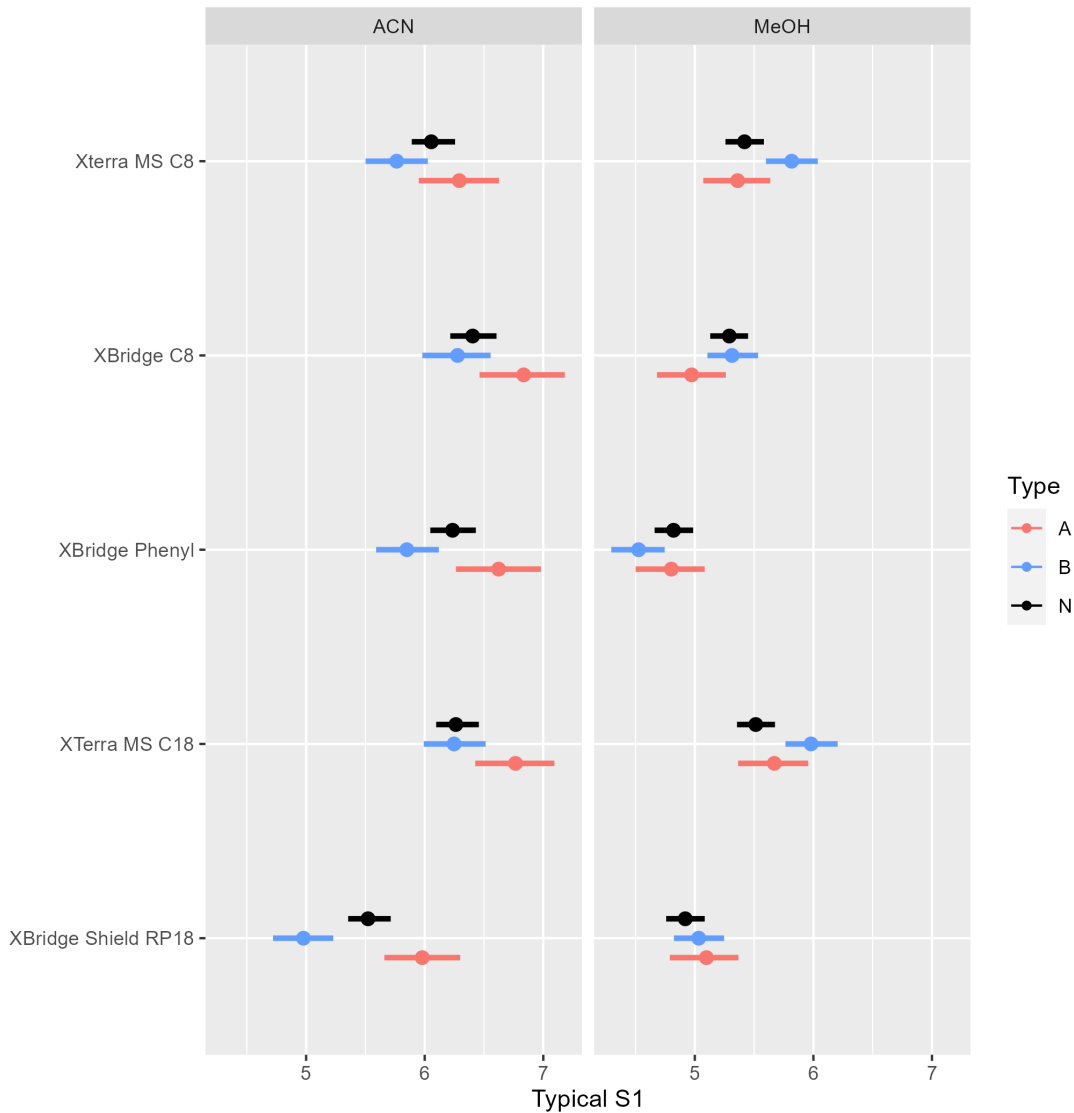
```

  mutate(c=c+1)%>%
  mutate(cS1aHat= S1mHat+cS1mHat + dS1mHat+cdS1mHat + dS1Hat+cdS1Hat + ddS1Hat+cddS1Hat)%>%
  mutate(S1aHat = S1mHat+dS1mHat+dS1Hat+ddS1Hat)%>%
  select(.draw,S1aHat,c)%>%
  tidyr::pivot_wider(names_from = c, values_from = cS1aHat) %>%
  rename(`1`=S1aHat) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xS1aHat") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(S1Hat = xS1aHat) %>%
  mutate(Type = "B")%>%
  mutate(Modifier="ACN")

p2<-bind_rows(MeOH_Neutral_results, MeOH_Acids_results, MeOH_Basic_results,
               ACN_Neutral_results,ACN_Acids_results, ACN_Basic_results) %>%
  ggplot(aes(y = ColumnNames, x = S1Hat, xmin = .lower, xmax = .upper,color = Type)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("A", "B", "N"), values = c("#F8766D", "#619cff", "black")) +
  ylab(' ')+
  xlab("Typical S1")+
  facet_wrap(~Modifier)

print(p2)

```



```

draws_df_subset <- fit$draws(format = "df", variable = c("beta", "cbeta"))

logkw_results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(beta[r], cbeta[c, r]) %>%
  filter(r == 1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c = c + 1) %>%

```

```

  mutate(cbeta=beta+cbeta)%>%
  select(.draw,beta,cbeta,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = cbeta) %>%
  rename(`1`=beta) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xbeta") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(beta = xbeta)%>%
  mutate(Parameter="logkw")

S1_results <-draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(beta[r],cbeta[c,r]) %>%
  filter(r==2) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(cbeta=beta+cbeta)%>%
  select(.draw,beta,cbeta,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = cbeta) %>%
  rename(`1`=beta) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xbeta") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(beta = xbeta)%>%
  mutate(Parameter="S1")

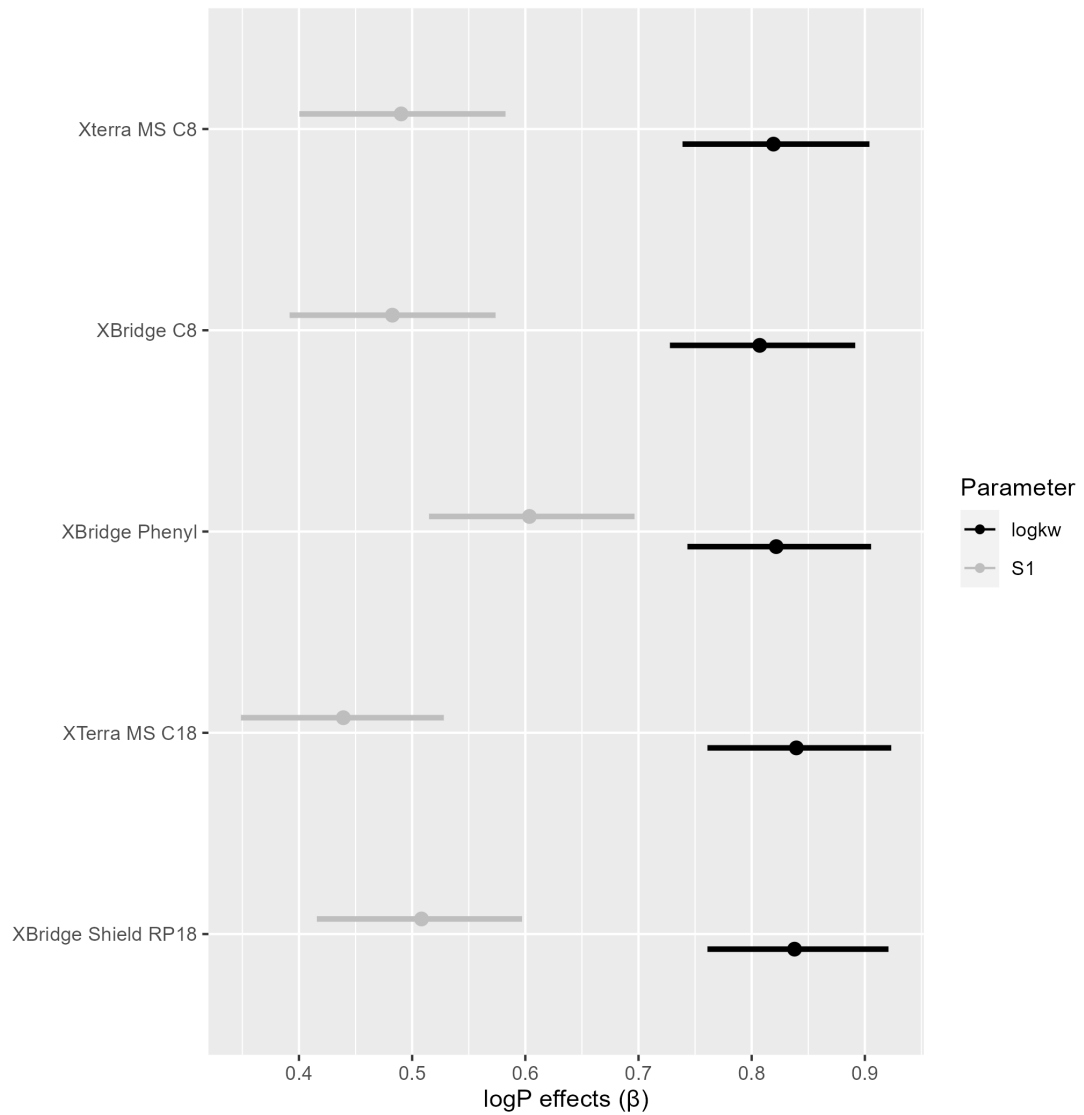
```

```

p3<-bind_rows(logkw_results, S1_results) %>%
  ggplot(aes(y = ColumnNames, x = beta, xmin = .lower, xmax = .upper, color = Parameter)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("logkw", "S1"), values = c("black", "gray")) +
  ylab(' ') +
  xlab("logP effects (\u03b2)")

print(p3)

```



```

draws_df_subset <- fit$draws(format = "df", variable = c("apH", "capH"))

Acids_results <-draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(apH[r], capH[c,r]) %>%
  filter(r==1) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(capH=apH+capH)%>%
  select(.draw,apH,capH,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = capH) %>%
  rename(`1` = apH) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xapH") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
                            '1' = 'XBridge Shield RP18',
                            '2' = 'XTerra MS C18',
                            '3' = 'XBridge Phenyl',
                            '4' = 'XBridge C8',
                            '5' = 'Xterra MS C8'))%>%
  group_by(ColumnNames)%>%
  tidybayes::median_qi(apH = xapH)%>%
  mutate(Type="A")

Bases_results <-draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(apH[r], capH[c,r]) %>%
  filter(r==2) %>%
  ungroup() %>%
  select(-r) %>%
  mutate(c=c+1)%>%
  mutate(capH=apH+capH)%>%
  select(.draw,apH,capH,c) %>%
  tidyr::pivot_wider(names_from = c, values_from = capH) %>%
  rename(`1` = apH) %>%
  select(-.draw)%>%
  tidyr::pivot_longer(c(1:5),names_to = "ColumnNames", values_to = "xapH") %>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,

```

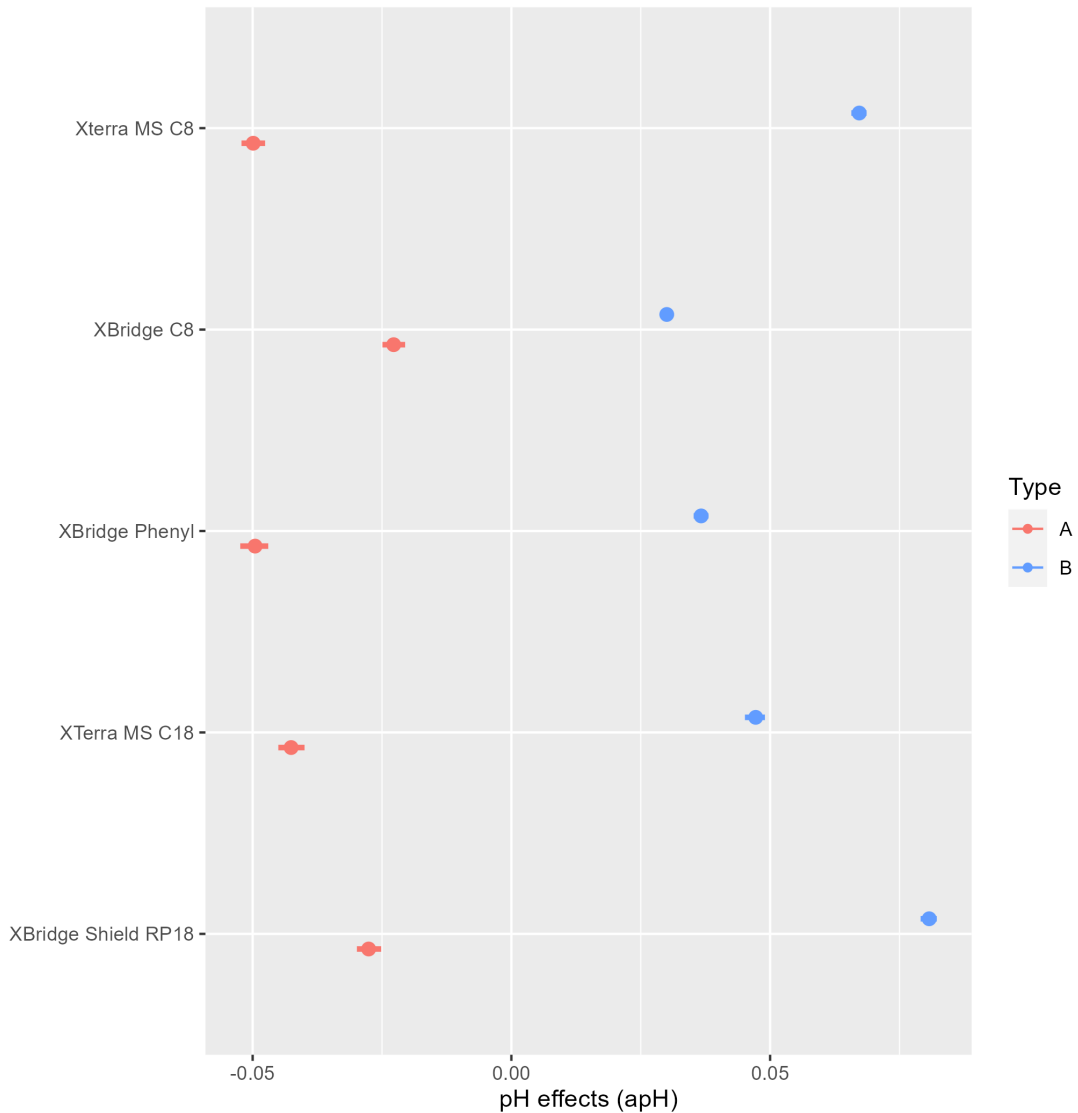
```

'1' = 'XBridge Shield RP18',
'2' = 'XTerra MS C18',
'3' = 'XBridge Phenyl',
'4' = 'XBridge C8',
'5' = 'Xterra MS C8'))%>%
group_by(ColumnName)%>%
tidybayes::median_qi(apH = xapH)%>%
mutate(Type="B")

p4<-bind_rows(Acids_results, Bases_results) %>%
  ggplot(aes(y = ColumnName, x = apH, xmin = .lower, xmax = .upper,color = Type)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("A", "B"), values = c("#F8766D", "#619cff"))+
  ylab(' ')+
  xlab("pH effects (apH)")

print(p4)

```



```

draws_df_subset <- fit$draws(format = "df", variable = c("dlogkTHat", "cdlogkTHat"))

Results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(dlogkTHat, cdlogkTHat[c]) %>%
  ungroup() %>%
  mutate(c=c+1)%>%
  mutate(cdlogkTHat=dlogkTHat+cdlogkTHat)%>%
  select(.draw,dlogkTHat,cdlogkTHat,c) %>%

```

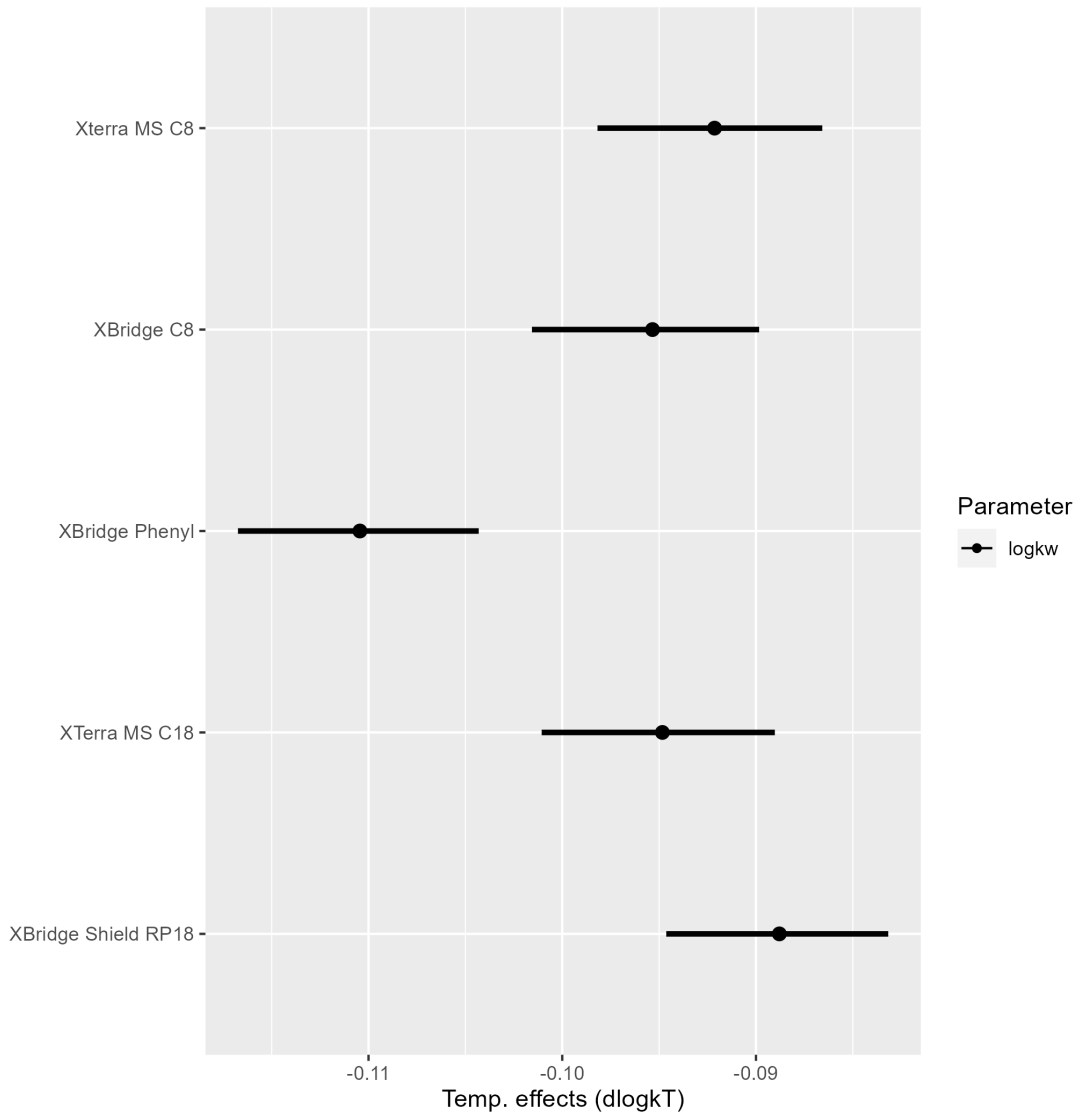
```

tidyr::pivot_wider(names_from = c, values_from = cdlogkTHat) %>%
  rename(`1` = dlogkTHat) %>%
  select(-.draw) %>%
  tidyr::pivot_longer(c(1:5), names_to = "ColumnNames", values_to = "xdlogkTHat") %>%
  mutate_if(is.character, as.factor) %>%
  mutate(ColumnNames=recode(ColumnNames,
    '1' = 'XBridge Shield RP18',
    '2' = 'XTerra MS C18',
    '3' = 'XBridge Phenyl',
    '4' = 'XBridge C8',
    '5' = 'Xterra MS C8')) %>%
  group_by(ColumnNames) %>%
  tidybayes::median_qi(dlogkTHat = xdlogkTHat) %>%
  mutate(Parameter="logkw")

p5<-Results %>%
  ggplot(aes(y = ColumnNames, x = dlogkTHat, xmin = .lower, xmax = .upper, color = Parameter))
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +
  scale_color_manual(labels = c("logkw"), values = c("black"))+
  ylab(' ')+
  xlab("Temp. effects (dlogkT)")

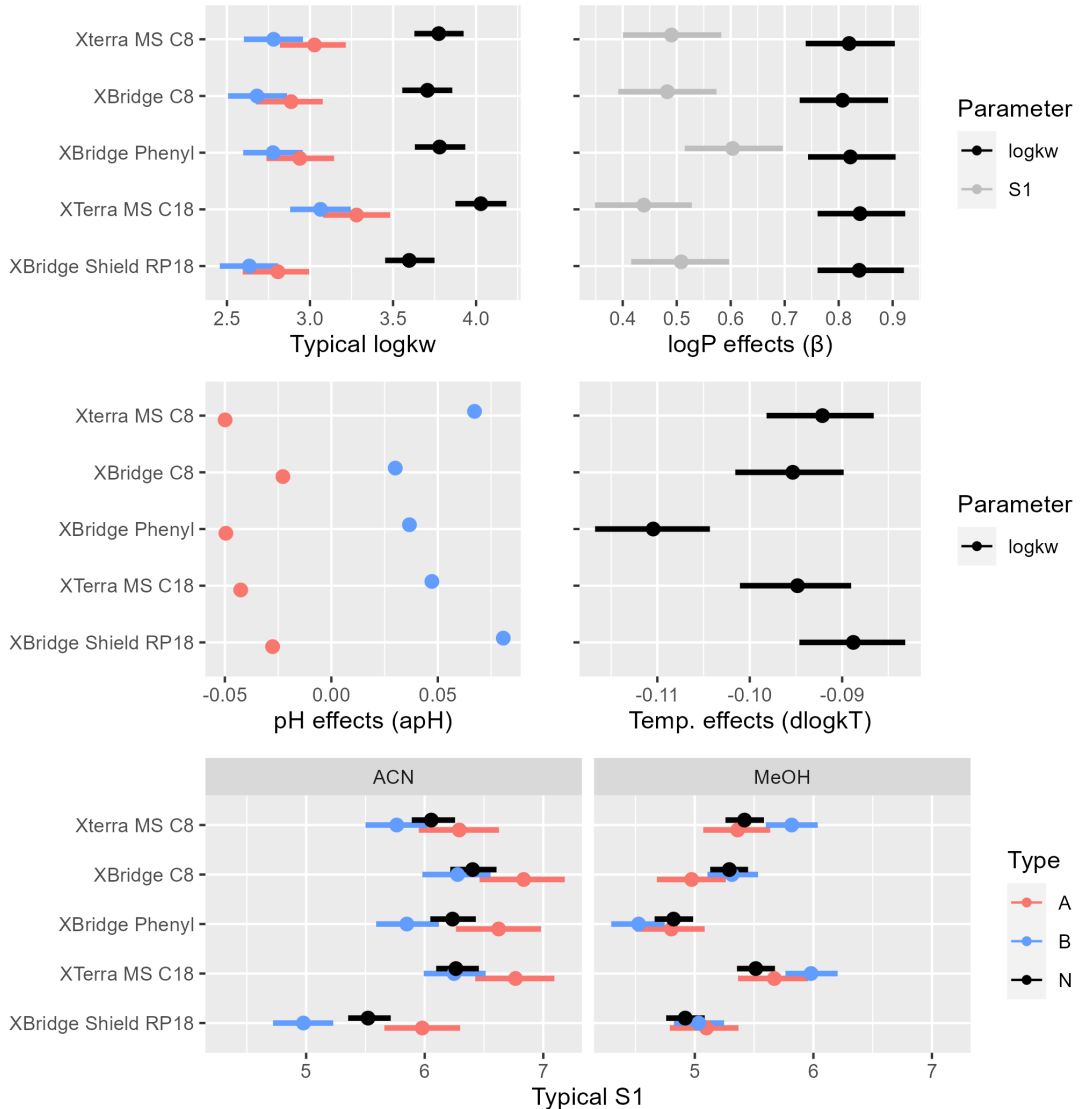
print(p5)

```



```

p=grid.arrange(p1+ theme(legend.position='none'),
  p3+theme(axis.text.y=element_blank()),
  p4+ theme(legend.position='none'),
  p5+theme(axis.text.y=element_blank()),
  p2, ncol=2, widths = c(1,1), layout_matrix = rbind(c(1, 2),c(3, 4), c(5, 5))
)
  
```



```
ggsave(paste0("figures\\param\\", "Joined", ".png"), plot=p, width = 20, height = 20, unit
```

Standard deviations:

```
draws_df_subset <- fit$draws(format = "df", variable = c("comega", "ckappa"))

comega_results <- draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(comega[c,r]) %>%
```

```

ungroup() %>%
  mutate(xparam=comega)%>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(c,
    '1' = 'XTerra MS C18',
    '2' = 'XBridge Phenyl',
    '3' = 'XBridge C8',
    '4' = 'Xterra MS C8'))%>%
  mutate(ParamNames=recode(r,
    '1' = 'clogkw',
    '2' = 'cS1m',
    '3' = 'cdS1'))%>%
  group_by(ColumnNames,ParamNames)%>%
  tidybayes::median_qi(param = xparam)%>%
  mutate(Parameter="comega")

ckappa_results <-draws_df_subset %>%
  slice_sample(n = 1000) %>%
  tidybayes::spread_draws(ckappa[c,r]) %>%
  ungroup() %>%
  mutate(xparam=ckappa)%>%
  mutate_if(is.character,as.factor) %>%
  mutate(ColumnNames=recode(c,
    '1' = 'XTerra MS C18',
    '2' = 'XBridge Phenyl',
    '3' = 'XBridge C8',
    '4' = 'Xterra MS C8'))%>%
  mutate(ParamNames=recode(r,
    '1' = 'clogkw',
    '2' = 'cS1m',
    '3' = 'cdS1'))%>%
  group_by(ColumnNames,ParamNames)%>%
  tidybayes::median_qi(param = xparam)%>%
  mutate(Parameter="ckappa")

p1<-bind_rows(comega_results, ckappa_results) %>%
  mutate(ParamNames=factor(ParamNames,levels=c("clogkw","cS1m","cdS1")))%>%
  ggplot(aes(y = ColumnNames, x = param, xmin = .lower, xmax = .upper, color=ParamNames)) +
  tidybayes::geom_pointinterval(position = position_dodge(width = .3)) +

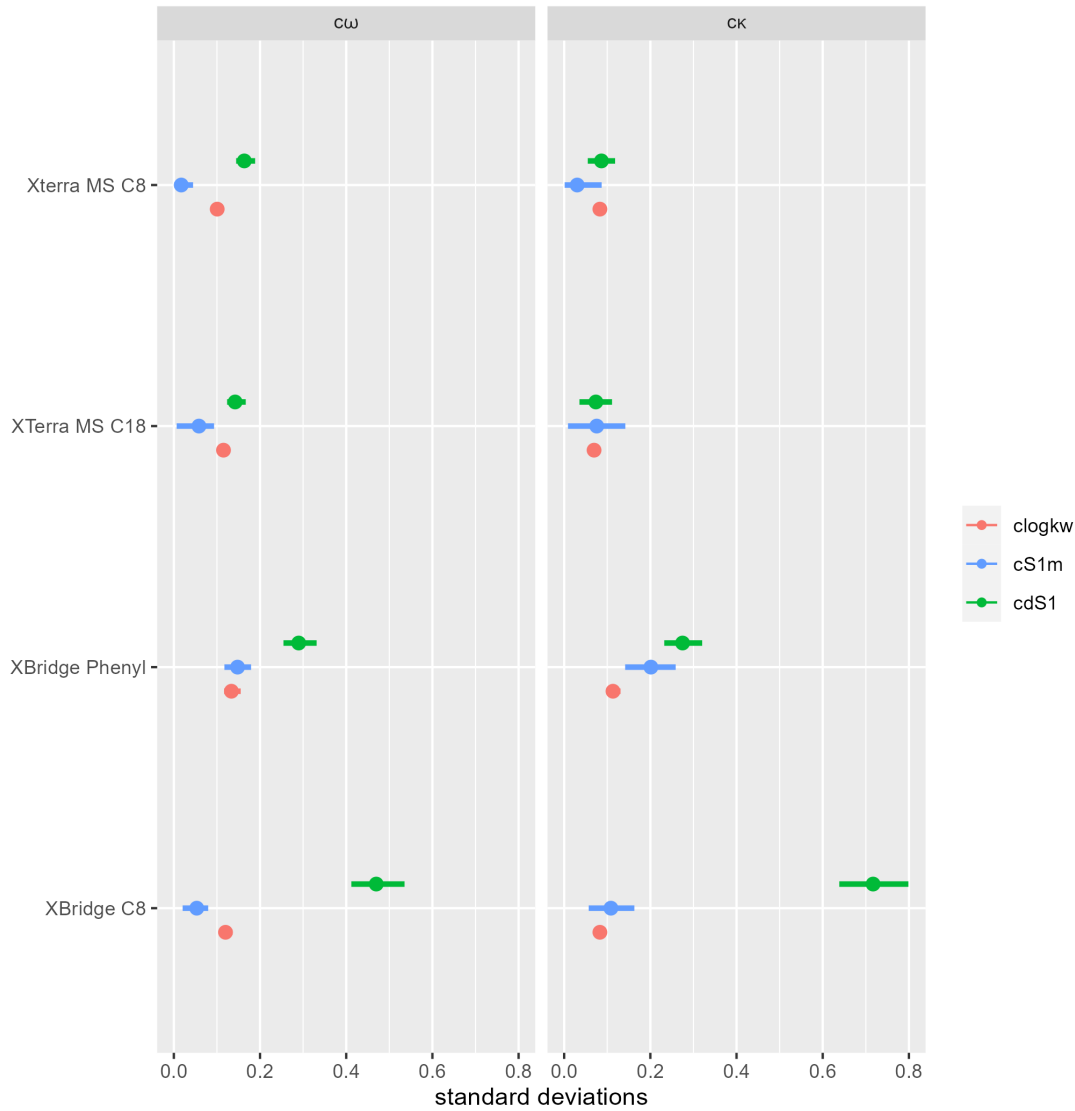
```

```

scale_colour_manual(name = " ", values=c("clogkw" = "#F8766D", "cS1m" = "#619CFF", "cdS1" = "#008000"))
facet_wrap(~factor(Parameter, levels=c("comega", "ckappa"), labels=c(expression(paste("c", "omega")), expression(paste("c", "kappa")))))
ylab(' ')
xlab("standard deviations")

print(p1)

```



```

ggsave(paste0("figures\\param\\", "Joined_SD", ".png"), plot=p1, width = 20, height = 10,

```

6.4 Isocratic predictions

To better asses the impact of parameters on retention we created graphs presenting the isocratic logarithm of retention factor vs. φ for selected analytes. Separate graphs are shown for each dissociation form (r=1,r=2,r=3). Here the individual predictions (given all the data) are shown:

```
analyte_ID_sample <-c(9,17,33,58,140,180)

for(i in 1:length(analyte_ID_sample)){
  idx_analyte = which(unique(data$METID)==analyte_ID_sample[i])
  draws_df_subset <- fit$draws(format = "df", variable = c(
    sprintf("logkwx[%s,1,1]",idx_analyte),
    sprintf("logkwx[%s,1,2]",idx_analyte),
    sprintf("logkwx[%s,1,3]",idx_analyte),
    sprintf("logkwx[%s,2,1]",idx_analyte),
    sprintf("logkwx[%s,2,2]",idx_analyte),
    sprintf("logkwx[%s,2,3]",idx_analyte),
    sprintf("logkwx[%s,3,1]",idx_analyte),
    sprintf("logkwx[%s,3,2]",idx_analyte),
    sprintf("logkwx[%s,3,3]",idx_analyte),
    sprintf("logkwx[%s,4,1]",idx_analyte),
    sprintf("logkwx[%s,4,2]",idx_analyte),
    sprintf("logkwx[%s,4,3]",idx_analyte),
    sprintf("logkwx[%s,5,1]",idx_analyte),
    sprintf("logkwx[%s,5,2]",idx_analyte),
    sprintf("logkwx[%s,5,3]",idx_analyte),
    sprintf("S1x[%s,1,1,1]",idx_analyte),
    sprintf("S1x[%s,1,1,2]",idx_analyte),
    sprintf("S1x[%s,1,1,3]",idx_analyte),
    sprintf("S1x[%s,2,1,1]",idx_analyte),
    sprintf("S1x[%s,2,1,2]",idx_analyte),
    sprintf("S1x[%s,2,1,3]",idx_analyte),
    sprintf("S1x[%s,1,2,1]",idx_analyte),
    sprintf("S1x[%s,1,2,2]",idx_analyte),
    sprintf("S1x[%s,1,2,3]",idx_analyte),
    sprintf("S1x[%s,2,2,1]",idx_analyte),
    sprintf("S1x[%s,2,2,2]",idx_analyte),
    sprintf("S1x[%s,2,2,3]",idx_analyte),
    sprintf("S1x[%s,1,3,1]",idx_analyte),
    sprintf("S1x[%s,1,3,2]",idx_analyte),
```

```

sprintf("S1x[%s,1,3,3]",idx_analyte),
sprintf("S1x[%s,2,3,1]",idx_analyte),
sprintf("S1x[%s,2,3,2]",idx_analyte),
sprintf("S1x[%s,2,3,3]",idx_analyte),
sprintf("S1x[%s,1,4,1]",idx_analyte),
sprintf("S1x[%s,1,4,2]",idx_analyte),
sprintf("S1x[%s,1,4,3]",idx_analyte),
sprintf("S1x[%s,2,4,1]",idx_analyte),
sprintf("S1x[%s,2,4,2]",idx_analyte),
sprintf("S1x[%s,2,4,3]",idx_analyte),
sprintf("S1x[%s,1,5,1]",idx_analyte),
sprintf("S1x[%s,1,5,2]",idx_analyte),
sprintf("S1x[%s,1,5,3]",idx_analyte),
sprintf("S1x[%s,2,5,1]",idx_analyte),
sprintf("S1x[%s,2,5,2]",idx_analyte),
sprintf("S1x[%s,2,5,3]",idx_analyte),
"S2x[1,1]",
"S2x[2,1])
```

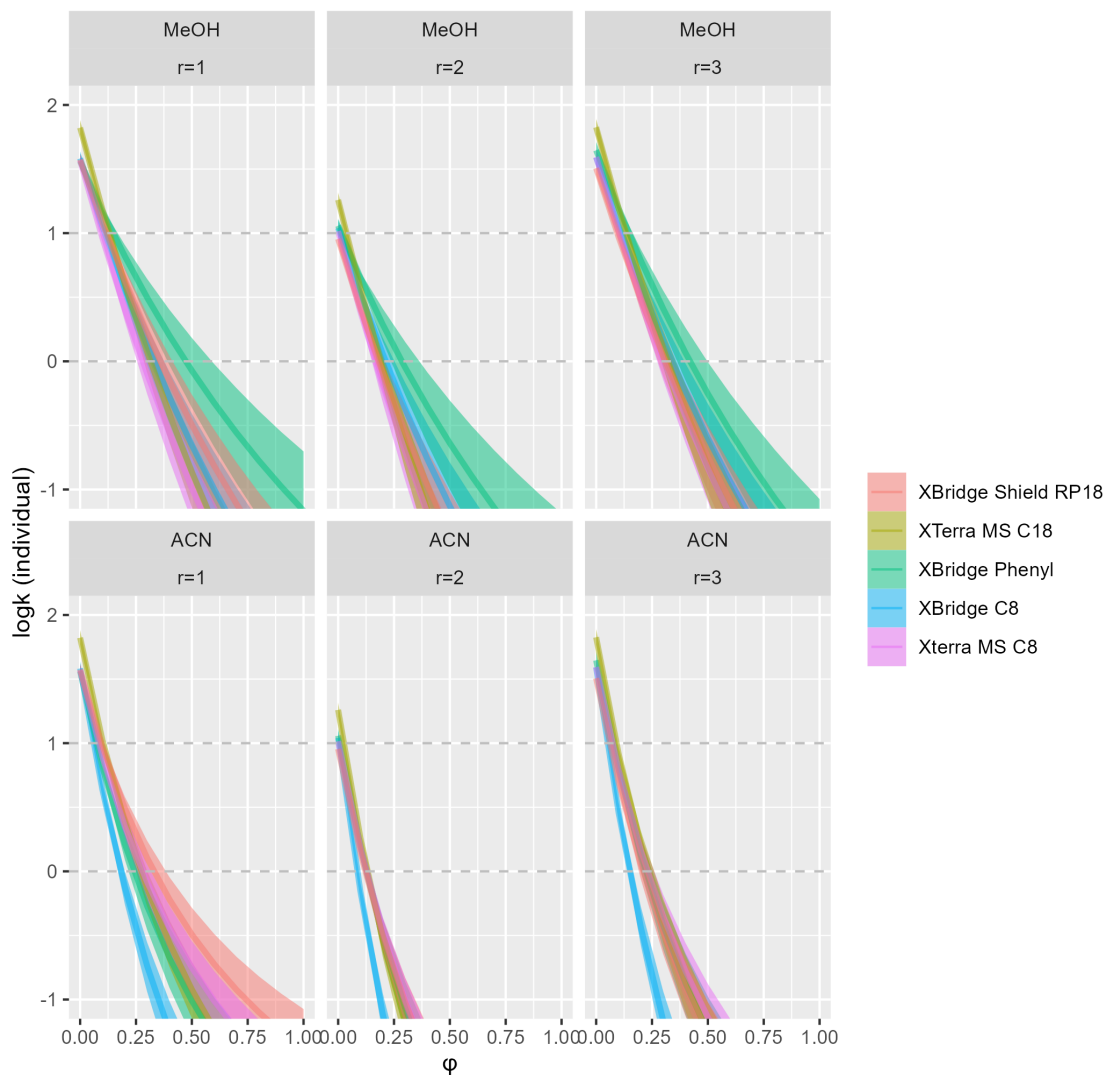
p <-draws_df_subset %>%
 slice_sample(n=1000) %>%
 tidybayes::spread_draws(logkwx[, c, r], S1x[, m, c, r], S2x[m,]) %>%
 filter(r<=R[idx_analyte]+1) %>%
 tidyrr::expand_grid(fi = seq(0,1,0.1)) %>%
 mutate(logk = logkwx-S1x*(1+S2x)*fi/(1+S2x*fi)) %>%
 ggplot(aes(x = fi, y = logk, color = as.factor(c), fill = as.factor(c))) +
 ggdist::stat_lineribbon(.width = c(.90), alpha = 1/2) +
 facet_wrap(m~r, nrow = 2, labeller = labeller(m=mod.labs,r=diss.labs)) +
 labs(title=paste(dataNames>Name[analyte_ID_sample[i]]),color= " ",fill= " ", x="\u03C6",
 scale_fill_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl",
 scale_color_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl",
 geom_hline(yintercept= c(0,1), linetype="dashed",color="gray") +
 coord_cartesian(xlim=c(0,1),ylim=c(-1,2))

print(p)

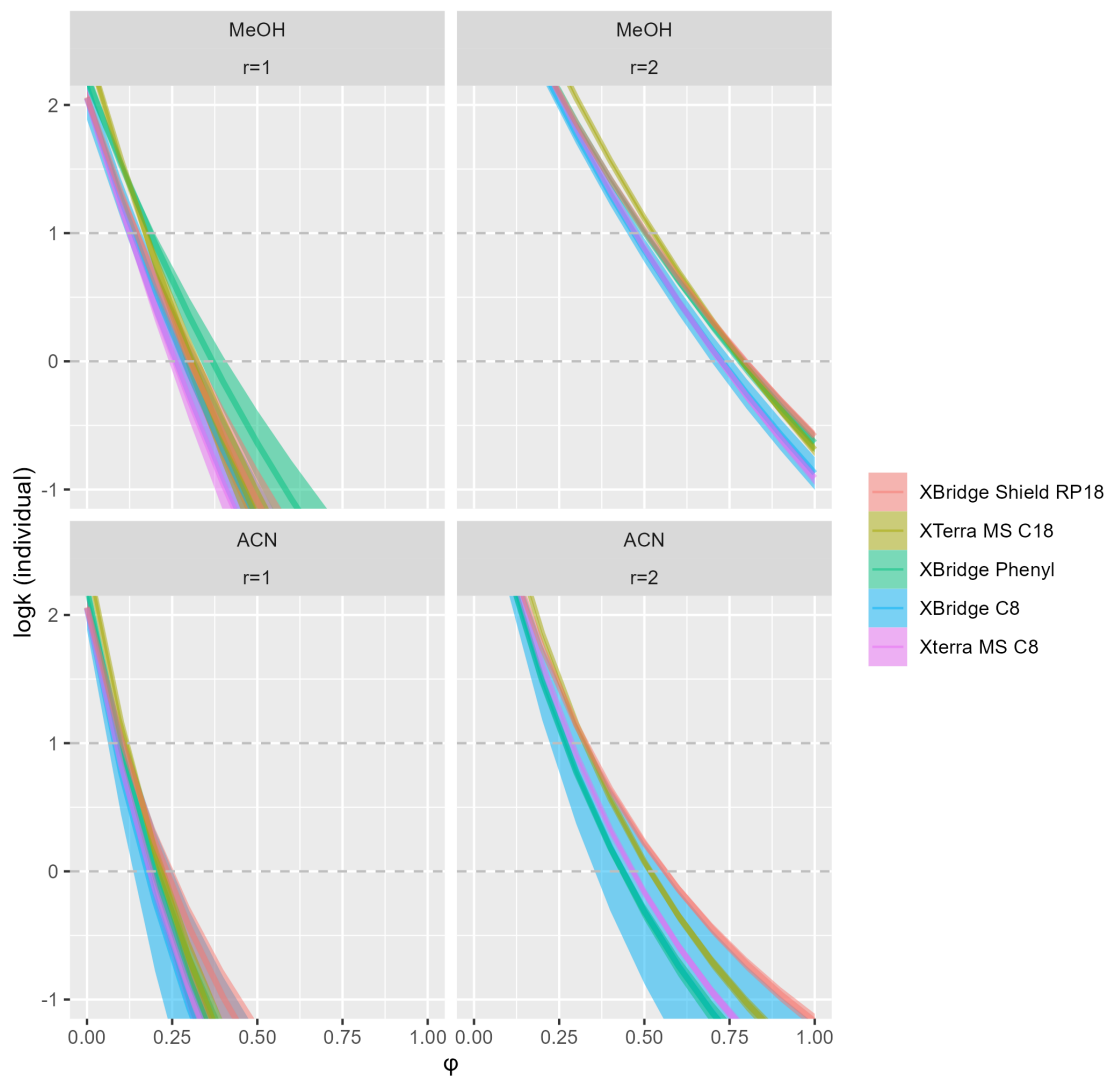
ggsave(paste0("figures\\izoparam\\", paste(dataNames>Name[analyte_ID_sample[i]]), ".Isocra

}

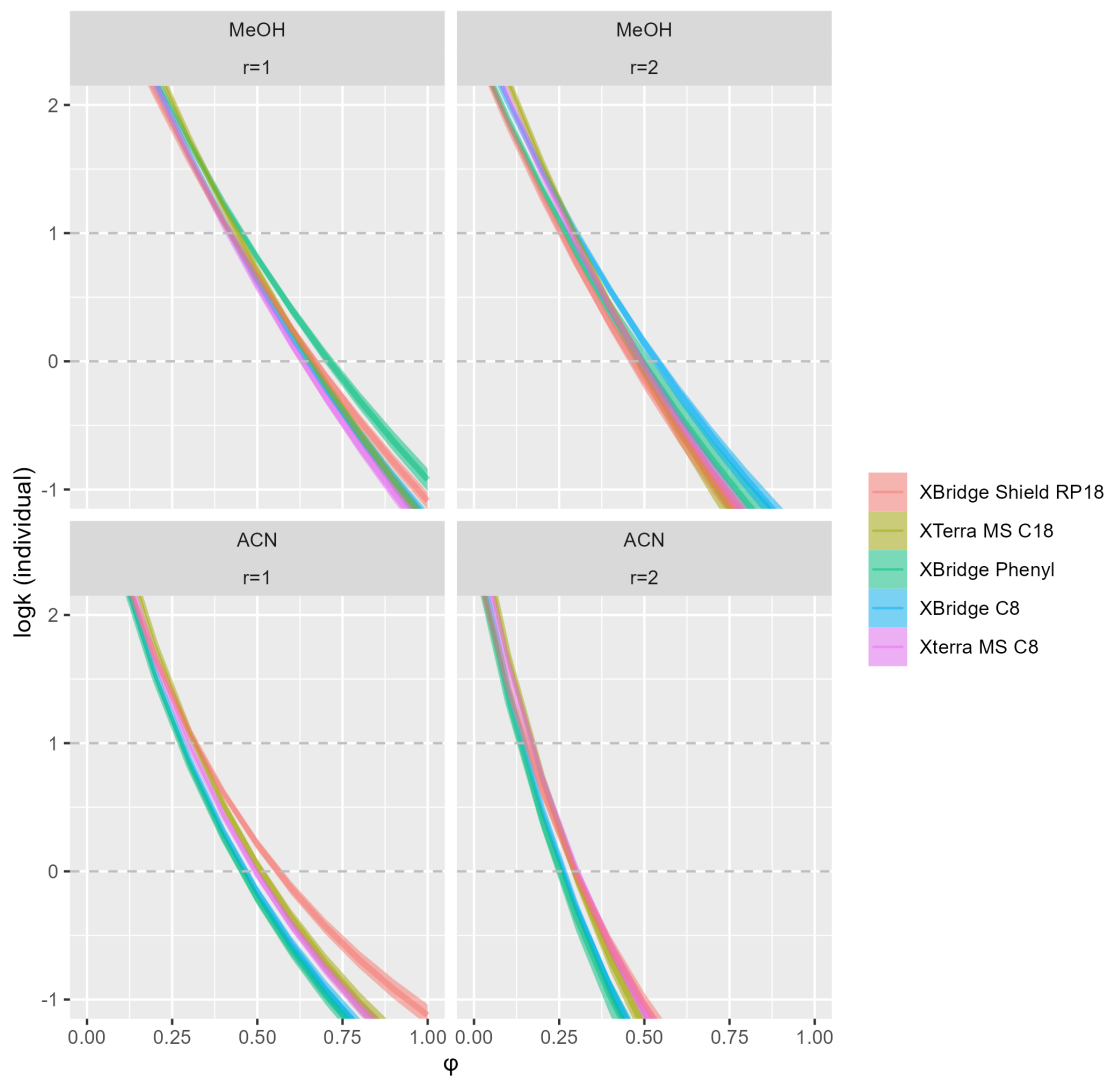
Baclofen



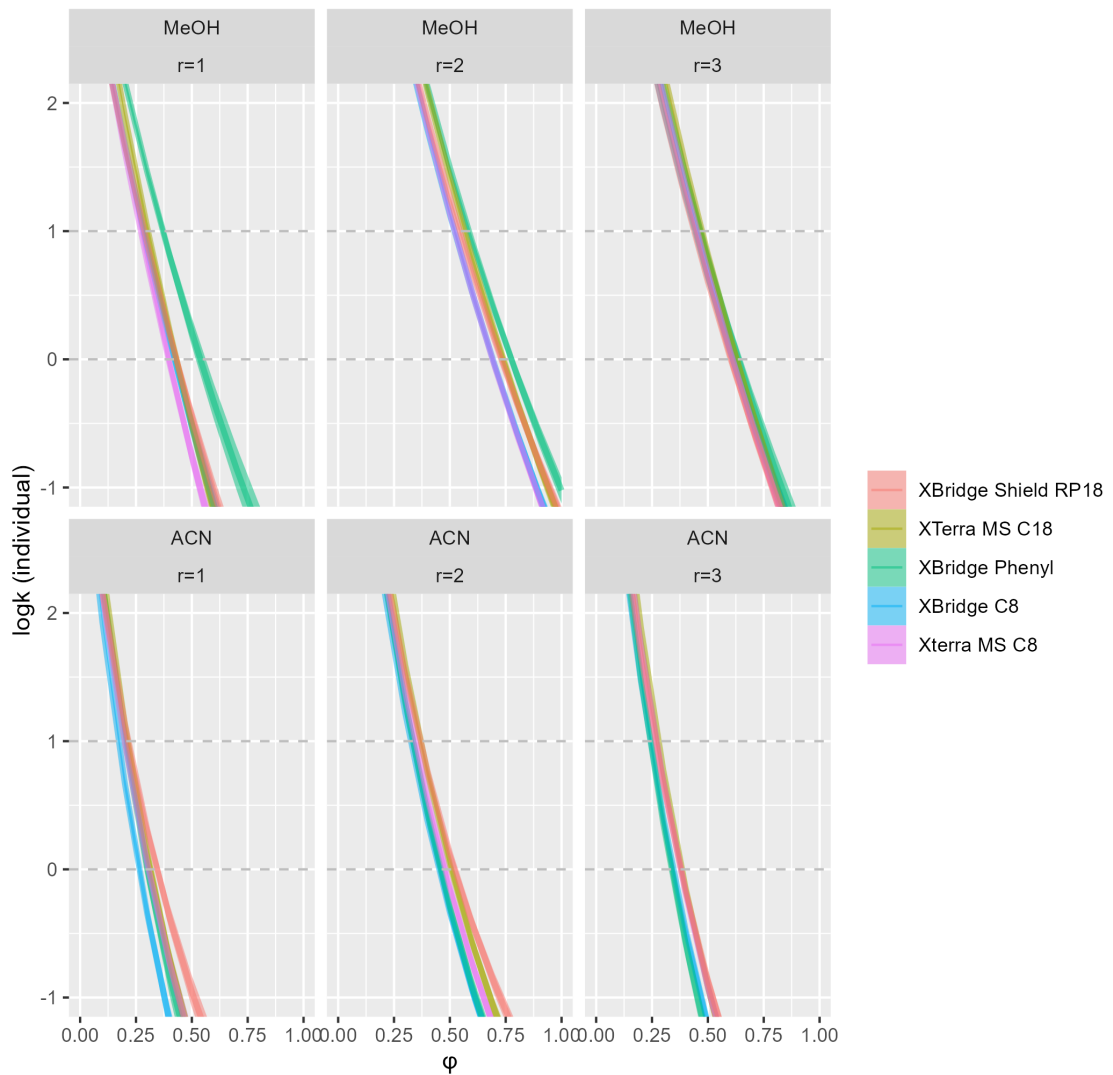
Acridine



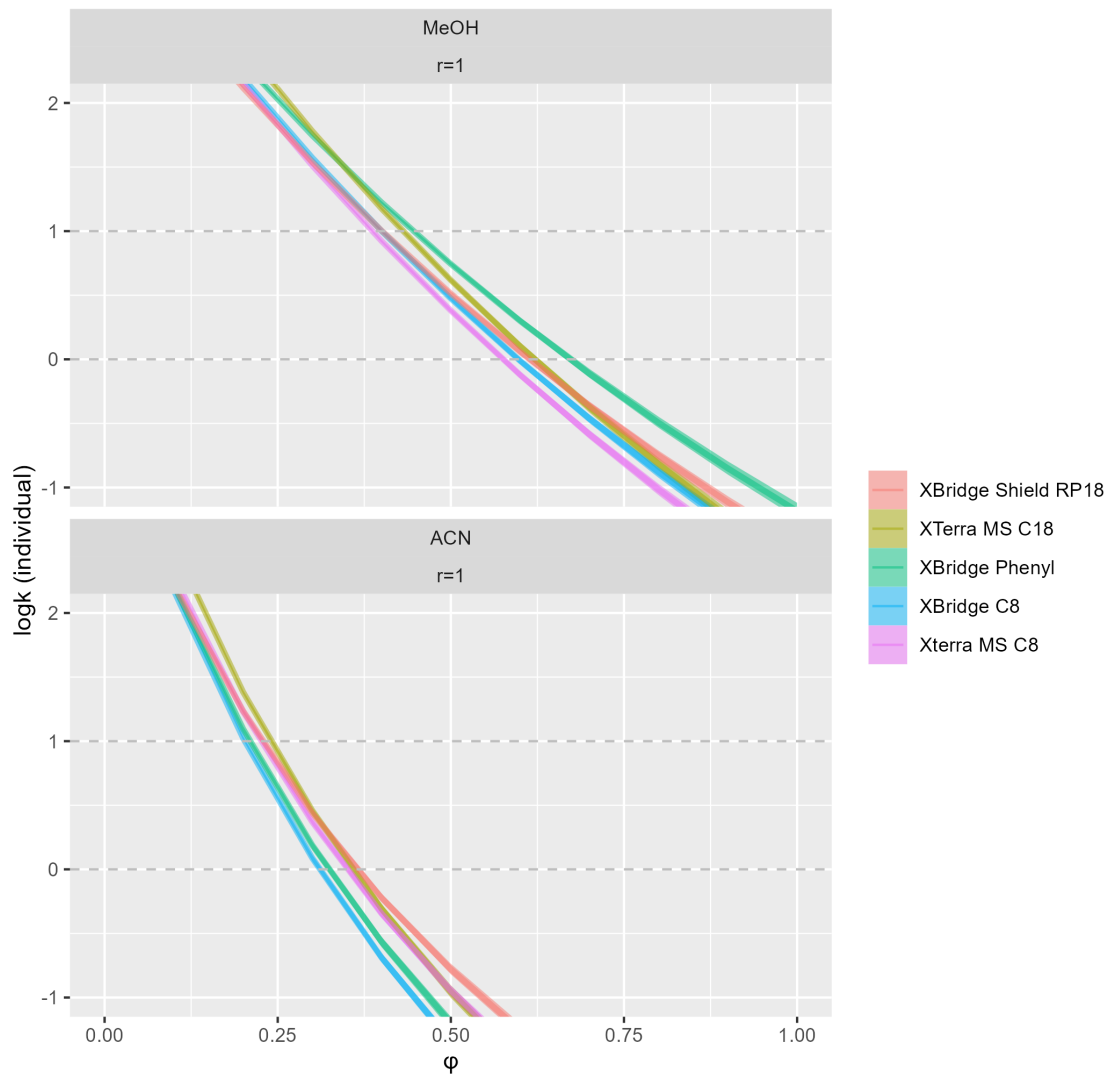
Tolbutamide



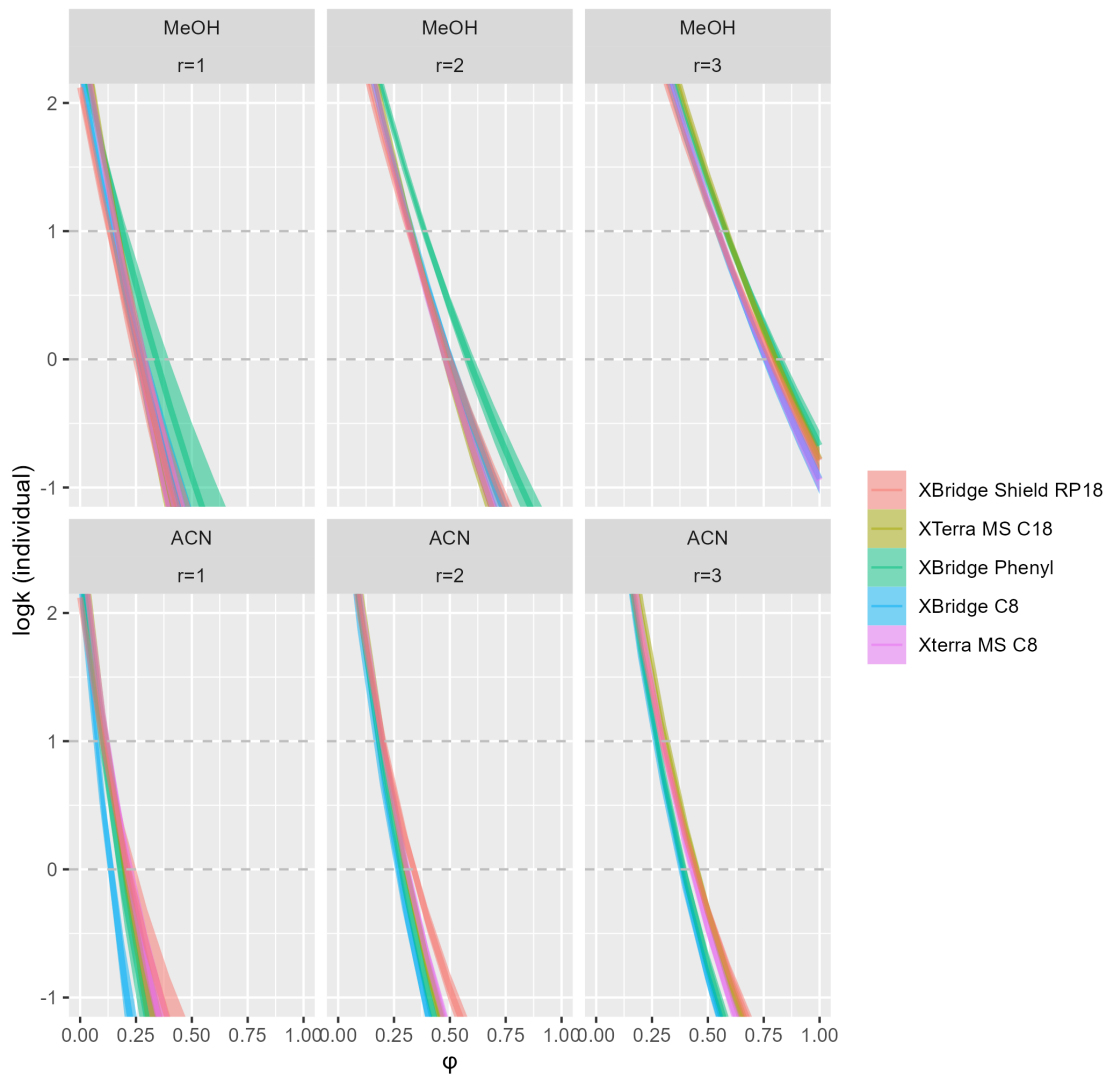
Pioglitazone



Hydrocortisone



Quinine



```
# AGG as the graphic backend of RStudio. (Tools -> Global Options -> General -> Graphics)
# https://github.com/tidyverse/ggplot2/issues/4661
# otherwise coord_cartesian does not work
```

Similarly we can quantify the column effects (between column differences in $\log K$ using XBridge Shield RP18 as a reference column):

```
analyte_ID_sample <- c(9, 17, 33, 58, 140, 180)
```

```

for(i in 1:length(analyte_ID_sample)){
  idx_analyte = which(unique(data$METID)==analyte_ID_sample[i])
  draws_df_subset <- fit$draws(format = "df", variable = c(
    sprintf("logkwx[%s,1,1]",idx_analyte),
    sprintf("logkwx[%s,1,2]",idx_analyte),
    sprintf("logkwx[%s,1,3]",idx_analyte),
    sprintf("logkwx[%s,2,1]",idx_analyte),
    sprintf("logkwx[%s,2,2]",idx_analyte),
    sprintf("logkwx[%s,2,3]",idx_analyte),
    sprintf("logkwx[%s,3,1]",idx_analyte),
    sprintf("logkwx[%s,3,2]",idx_analyte),
    sprintf("logkwx[%s,3,3]",idx_analyte),
    sprintf("logkwx[%s,4,1]",idx_analyte),
    sprintf("logkwx[%s,4,2]",idx_analyte),
    sprintf("logkwx[%s,4,3]",idx_analyte),
    sprintf("logkwx[%s,5,1]",idx_analyte),
    sprintf("logkwx[%s,5,2]",idx_analyte),
    sprintf("logkwx[%s,5,3]",idx_analyte),
    sprintf("S1x[%s,1,1,1]",idx_analyte),
    sprintf("S1x[%s,1,1,2]",idx_analyte),
    sprintf("S1x[%s,1,1,3]",idx_analyte),
    sprintf("S1x[%s,2,1,1]",idx_analyte),
    sprintf("S1x[%s,2,1,2]",idx_analyte),
    sprintf("S1x[%s,2,1,3]",idx_analyte),
    sprintf("S1x[%s,1,2,1]",idx_analyte),
    sprintf("S1x[%s,1,2,2]",idx_analyte),
    sprintf("S1x[%s,1,2,3]",idx_analyte),
    sprintf("S1x[%s,2,2,1]",idx_analyte),
    sprintf("S1x[%s,2,2,2]",idx_analyte),
    sprintf("S1x[%s,2,2,3]",idx_analyte),
    sprintf("S1x[%s,1,3,1]",idx_analyte),
    sprintf("S1x[%s,1,3,2]",idx_analyte),
    sprintf("S1x[%s,1,3,3]",idx_analyte),
    sprintf("S1x[%s,2,3,1]",idx_analyte),
    sprintf("S1x[%s,2,3,2]",idx_analyte),
    sprintf("S1x[%s,2,3,3]",idx_analyte),
    sprintf("S1x[%s,1,4,1]",idx_analyte),
    sprintf("S1x[%s,1,4,2]",idx_analyte),
    sprintf("S1x[%s,1,4,3]",idx_analyte),
    sprintf("S1x[%s,2,4,1]",idx_analyte),
  )
)
}

```

```

sprintf("S1x[%s,2,4,2]",idx_analyte),
sprintf("S1x[%s,2,4,3]",idx_analyte),
sprintf("S1x[%s,1,5,1]",idx_analyte),
sprintf("S1x[%s,1,5,2]",idx_analyte),
sprintf("S1x[%s,1,5,3]",idx_analyte),
sprintf("S1x[%s,2,5,1]",idx_analyte),
sprintf("S1x[%s,2,5,2]",idx_analyte),
sprintf("S1x[%s,2,5,3]",idx_analyte),
"S2x[1,1]",
"S2x[2,1]"))

p<-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwx[, c, r], S1x[, m, c, r], S2x[m, ]) %>%
  filter(r<=R[idx_analyte]+1) %>%
  tidyr::expand_grid(fi = seq(0,1,0.05)) %>%
  mutate(k = (logkwx-S1x*(1+S2x)*fi/(1+S2x*fi))) %>%
  select(.draw,c,r,m,k,fi) %>%
  tidyr::pivot_wider(names_from = c, values_from = k) %>%
  mutate(cdk2 = `2`-`1` ) %>%
  mutate(cdk3 = `3`-`1` ) %>%
  mutate(cdk4 = `4`-`1` ) %>%
  mutate(cdk5 = `5`-`1` ) %>%
  tidyr::pivot_longer(cdk2:cdk5,names_to = "names_cdk", values_to = "cdk")%>%
  ggplot(aes(x = fi, y = cdk, color = as.factor(names_cdk), fill = as.factor(names_cdk))) +
  ggdist::stat_lineribbon(.width = c(.90), alpha = 1/2) +
  facet_wrap(m~r, nrow = 2,labeller = labeller(m=mod.labs,r=diss.labs)) +
  labs(y = "Between column difference in logk (individual)", title=paste(dataNames>Name[analyte_ID_sample[i]])) +
  scale_fill_discrete(labels= c("XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xterra M
  scale_color_discrete(labels= c("XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xterra M
  geom_hline(yintercept= c(0), linetype="dashed",color="gray") +
  coord_cartesian(xlim=c(0,1),ylim=c(-1,1))

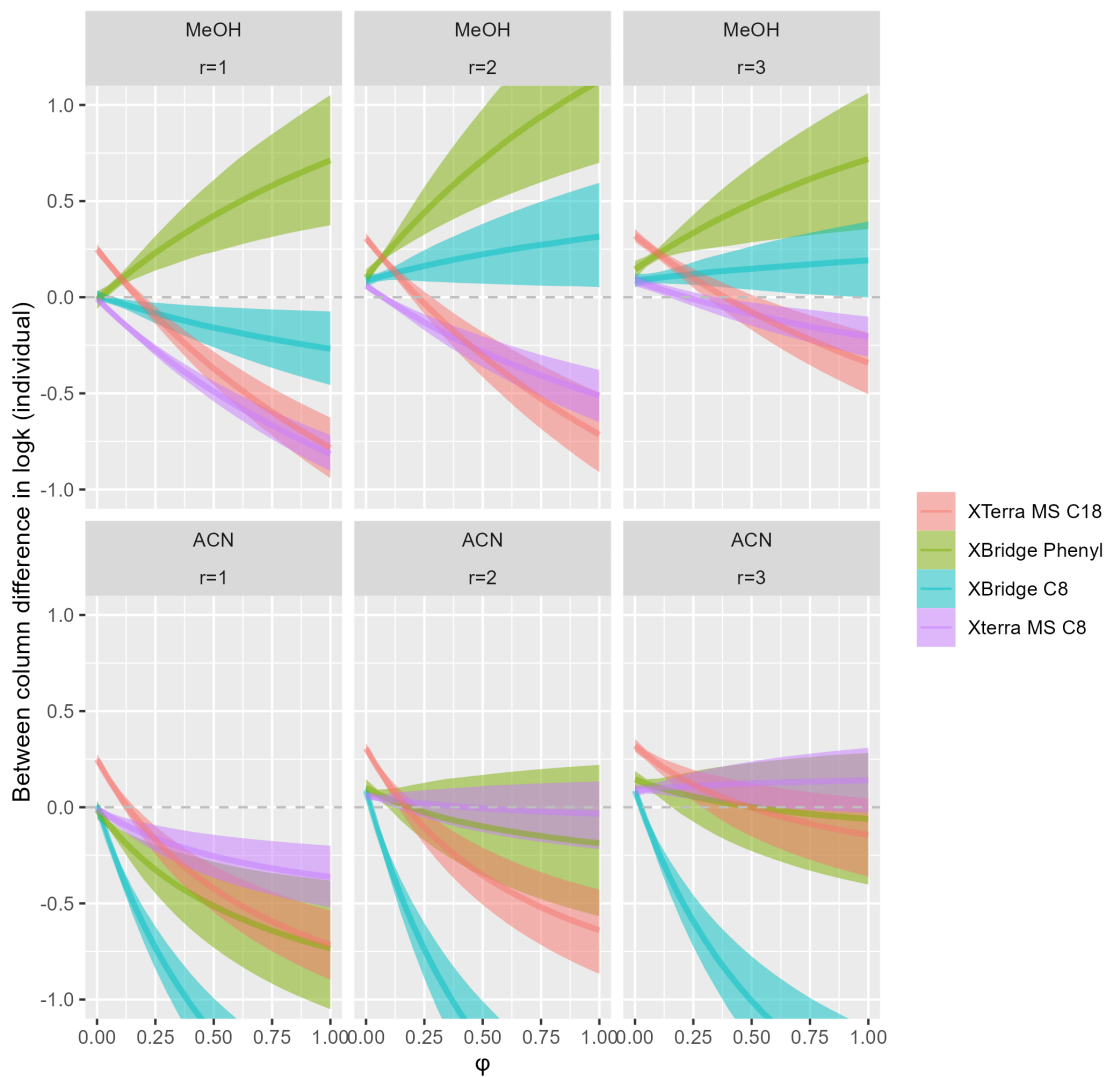
print(p)

ggsave(paste0("figures\\izoparam\\", paste(dataNames>Name[analyte_ID_sample[i]]), ".Isocr

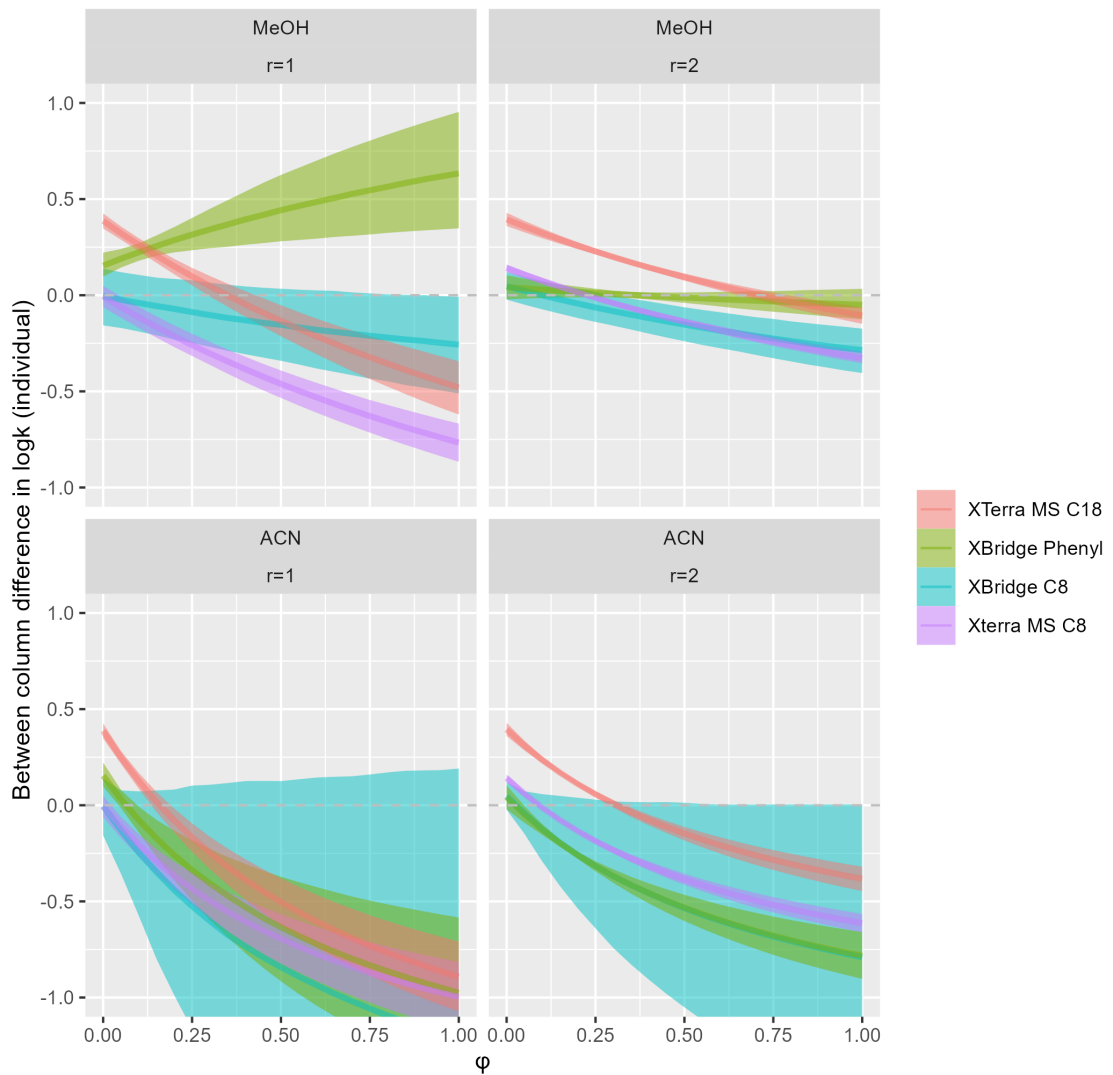
}

```

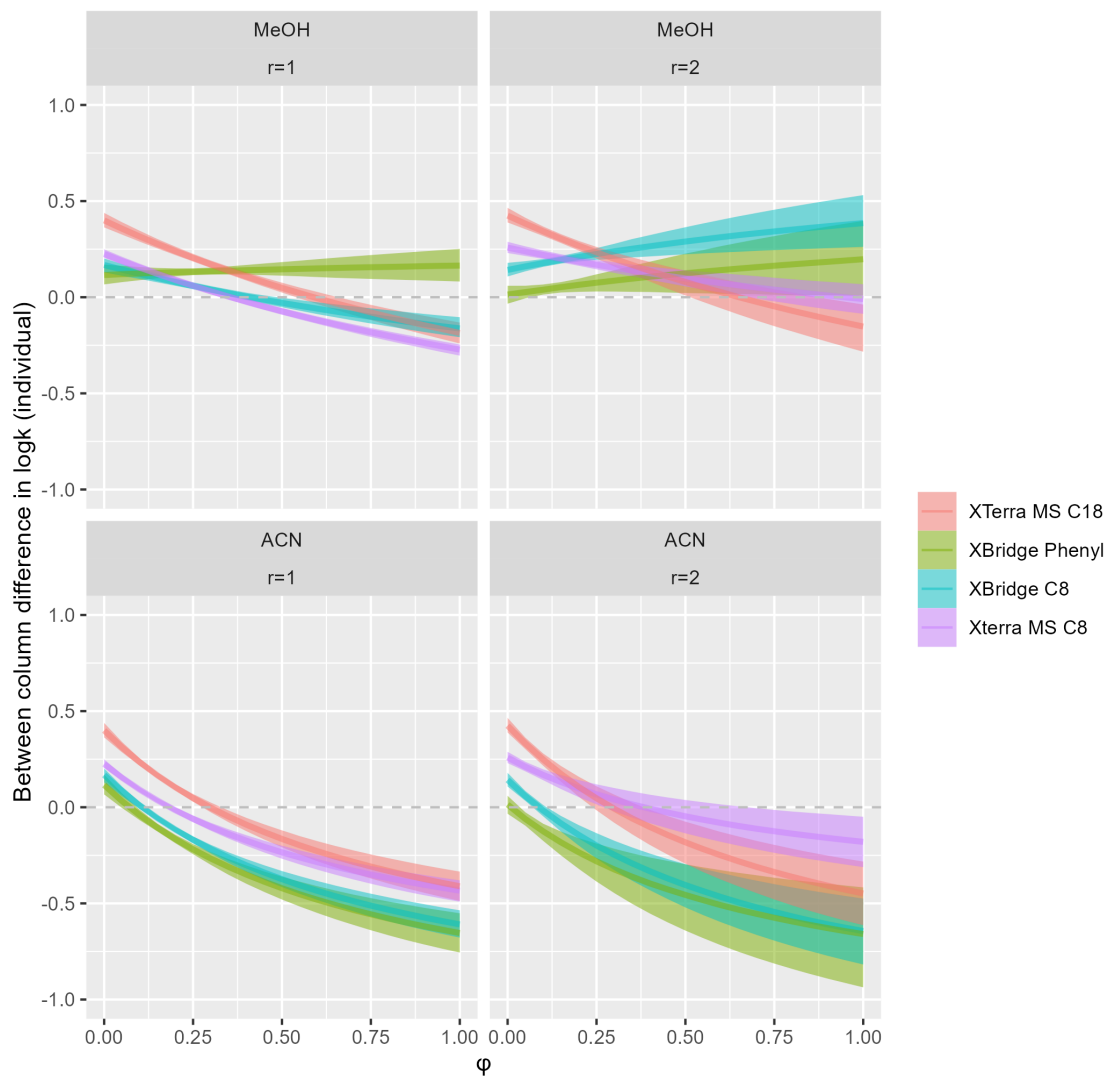
Baclofen



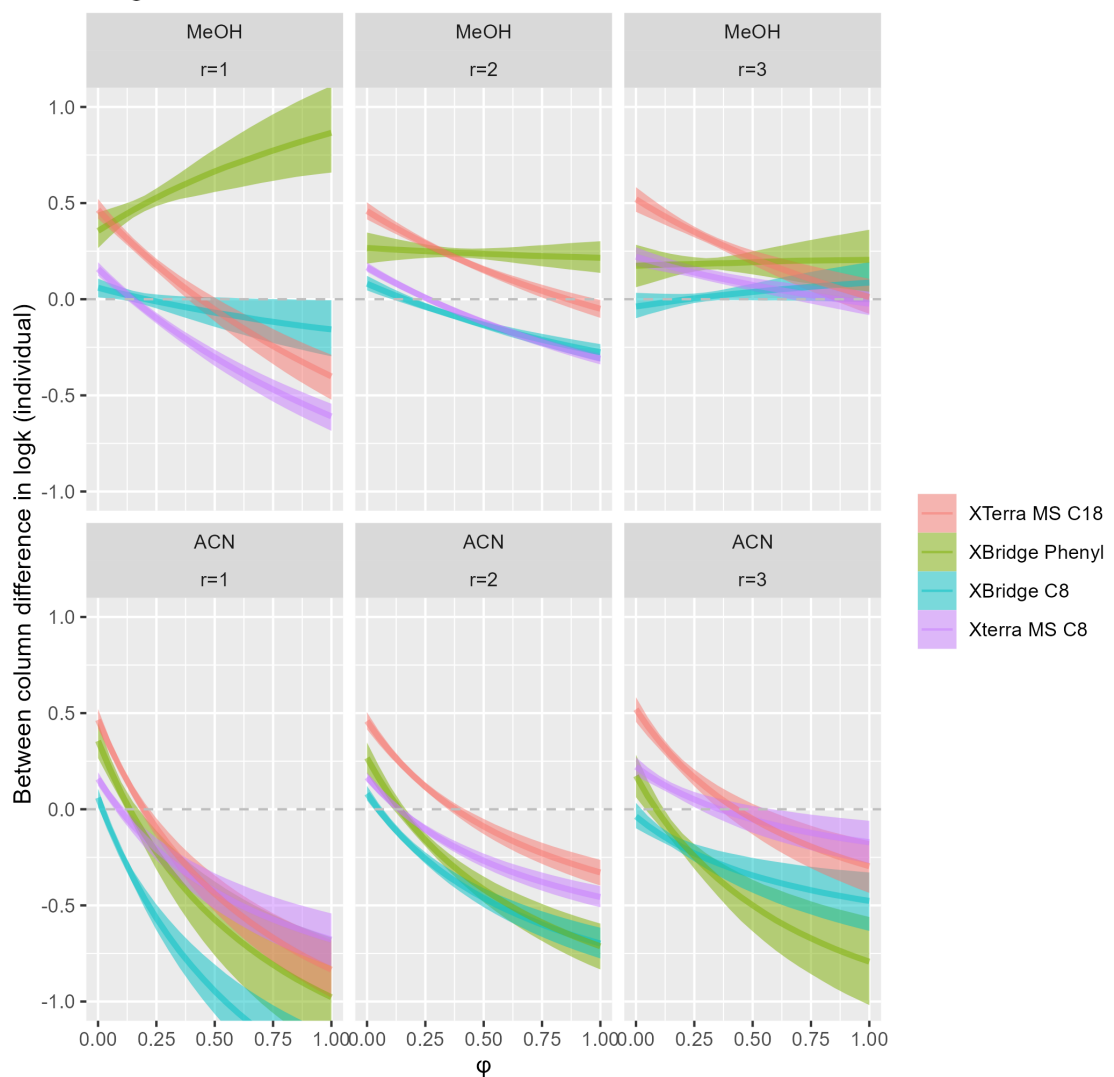
Acridine



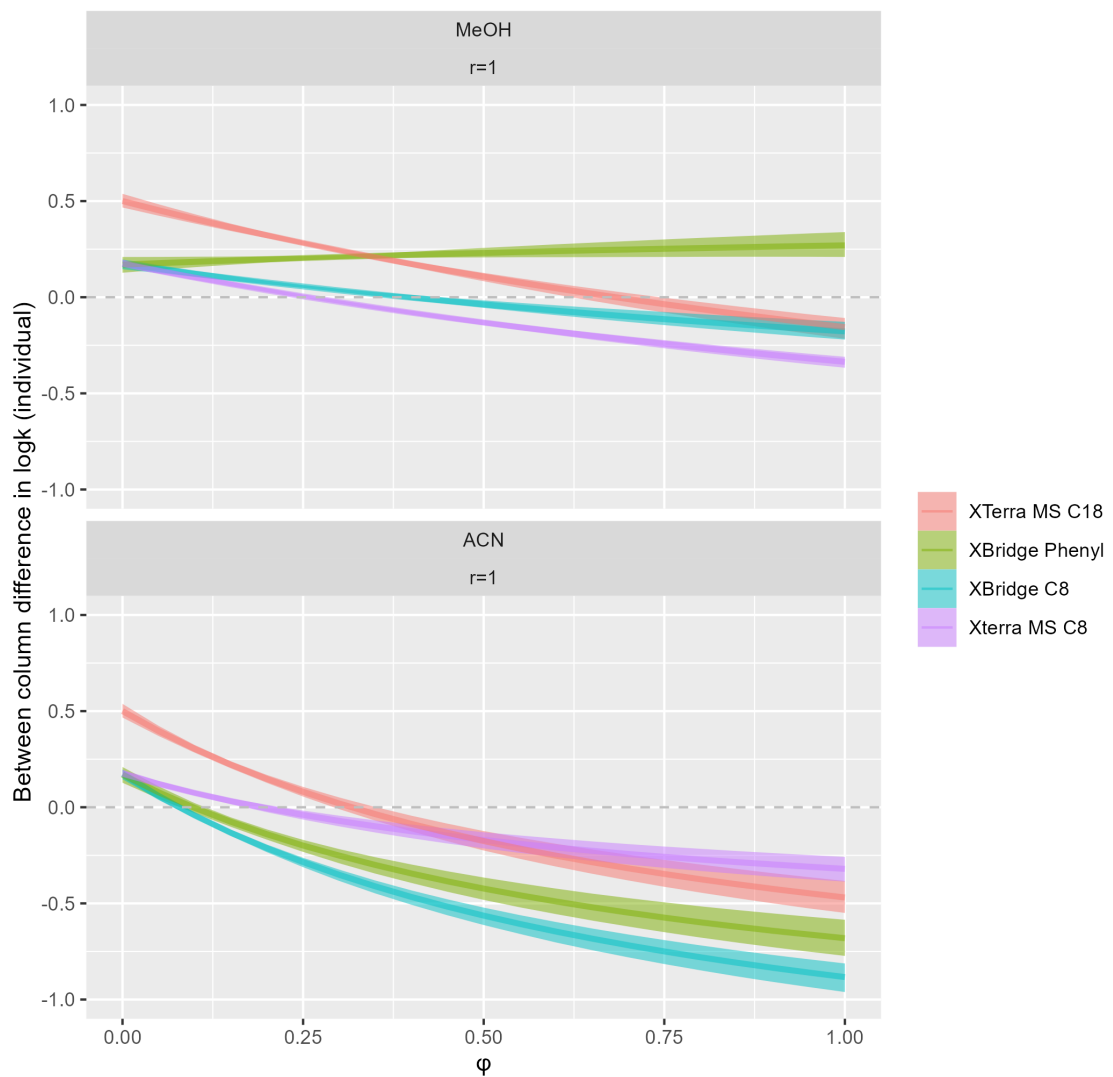
Tolbutamide

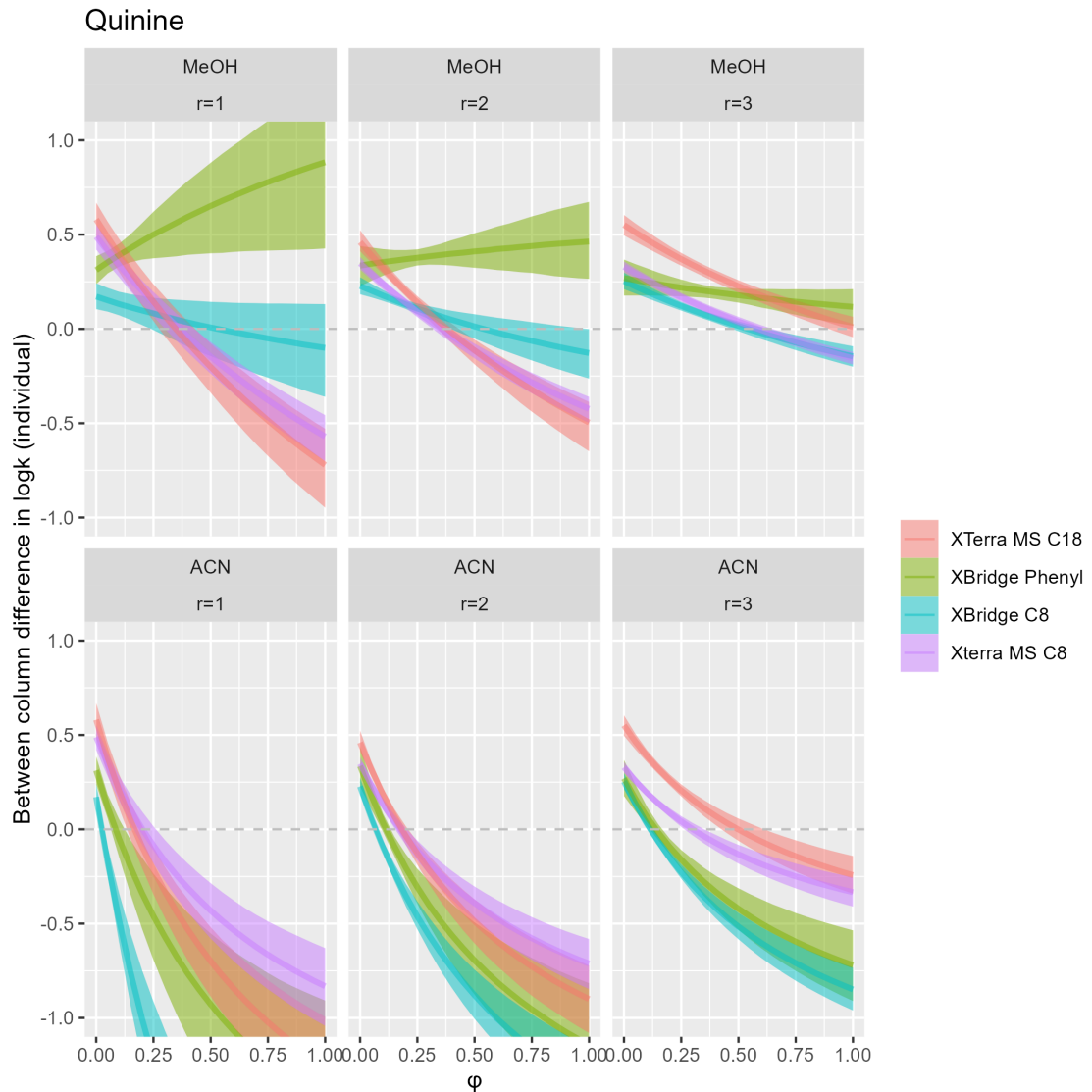


Pioglitazone



Hydrocortisone





or predict the organic modifier content leading to $\log k$ of 1:

```
analyte_ID_sample <-c(9,17,33,58,140,180)

for(i in 1:length(analyte_ID_sample)){

  idx_analyte = which(unique(data$METID)==analyte_ID_sample[i])
  draws_df_subset <- fit$draws(format = "df", variable = c(
```

```
    sprintf("logkwx[%s,1,1]",idx_analyte),
    sprintf("logkwx[%s,1,2]",idx_analyte),
    sprintf("logkwx[%s,1,3]",idx_analyte),
    sprintf("logkwx[%s,2,1]",idx_analyte),
    sprintf("logkwx[%s,2,2]",idx_analyte),
    sprintf("logkwx[%s,2,3]",idx_analyte),
    sprintf("logkwx[%s,3,1]",idx_analyte),
    sprintf("logkwx[%s,3,2]",idx_analyte),
    sprintf("logkwx[%s,3,3]",idx_analyte),
    sprintf("logkwx[%s,4,1]",idx_analyte),
    sprintf("logkwx[%s,4,2]",idx_analyte),
    sprintf("logkwx[%s,4,3]",idx_analyte),
    sprintf("logkwx[%s,5,1]",idx_analyte),
    sprintf("logkwx[%s,5,2]",idx_analyte),
    sprintf("logkwx[%s,5,3]",idx_analyte),
    sprintf("S1x[%s,1,1,1]",idx_analyte),
    sprintf("S1x[%s,1,1,2]",idx_analyte),
    sprintf("S1x[%s,1,1,3]",idx_analyte),
    sprintf("S1x[%s,2,1,1]",idx_analyte),
    sprintf("S1x[%s,2,1,2]",idx_analyte),
    sprintf("S1x[%s,2,1,3]",idx_analyte),
    sprintf("S1x[%s,1,2,1]",idx_analyte),
    sprintf("S1x[%s,1,2,2]",idx_analyte),
    sprintf("S1x[%s,1,2,3]",idx_analyte),
    sprintf("S1x[%s,2,2,1]",idx_analyte),
    sprintf("S1x[%s,2,2,2]",idx_analyte),
    sprintf("S1x[%s,2,2,3]",idx_analyte),
    sprintf("S1x[%s,1,3,1]",idx_analyte),
    sprintf("S1x[%s,1,3,2]",idx_analyte),
    sprintf("S1x[%s,1,3,3]",idx_analyte),
    sprintf("S1x[%s,2,3,1]",idx_analyte),
    sprintf("S1x[%s,2,3,2]",idx_analyte),
    sprintf("S1x[%s,2,3,3]",idx_analyte),
    sprintf("S1x[%s,1,4,1]",idx_analyte),
    sprintf("S1x[%s,1,4,2]",idx_analyte),
    sprintf("S1x[%s,1,4,3]",idx_analyte),
    sprintf("S1x[%s,2,4,1]",idx_analyte),
    sprintf("S1x[%s,2,4,2]",idx_analyte),
    sprintf("S1x[%s,2,4,3]",idx_analyte),
    sprintf("S1x[%s,1,5,1]",idx_analyte),
    sprintf("S1x[%s,1,5,2]",idx_analyte),
```

```

sprintf("S1x[%s,1,5,3]",idx_analyte),
sprintf("S1x[%s,2,5,1]",idx_analyte),
sprintf("S1x[%s,2,5,2]",idx_analyte),
sprintf("S1x[%s,2,5,3]",idx_analyte),
"S2x[1,1]",
"S2x[2,1])"

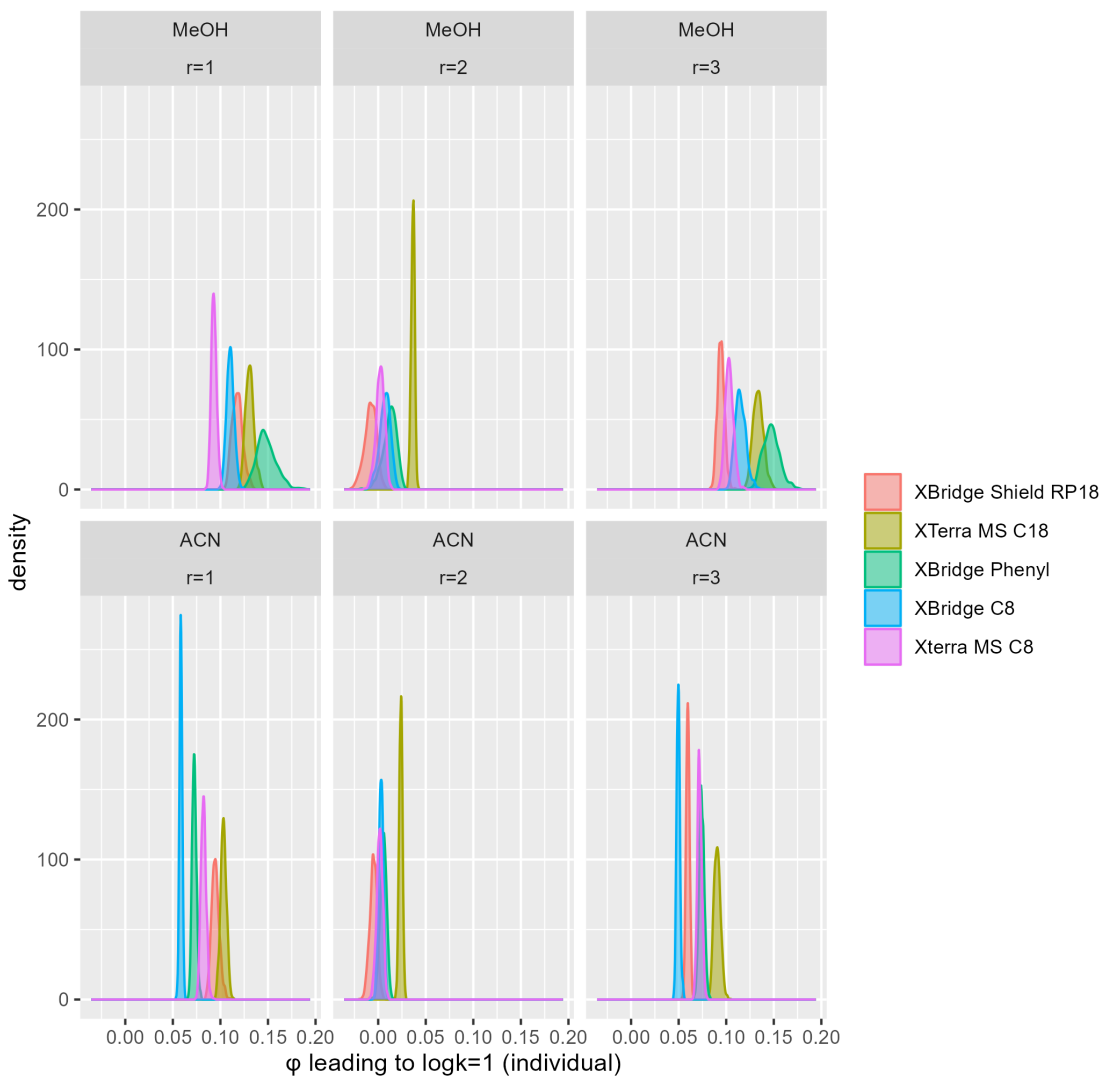
p<-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwx[, c, r], S1x[, m, c, r], S2x[m, ]) %>%
  filter(r<=R[idx_analyte]+1) %>%
  mutate(foo = (logkwx-1)/S1x/(1+S2x)) %>%
  mutate(fix = foo/(1-S2x*foo)) %>%
  ggplot(aes(x = fix, color = as.factor(c), fill = as.factor(c))) +
  geom_density(alpha = 1/2) +
  #coord_cartesian(xlim=c(0,1))+
  facet_wrap(m~r, nrow = 2,labeller = labeller(m=mod.labs,r=diss.labs))+
  labs(title=paste(dataNames>Name[analyte_ID_sample[i]]),
       color= " ",
       fill= " ",
       x = "\u03c6 leading to log=1 (individual)")+
  scale_fill_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl",
  scale_color_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl",
  facet_wrap(m~r, nrow = 2,labeller = labeller(m=mod.labs,r=diss.labs))

print(p)

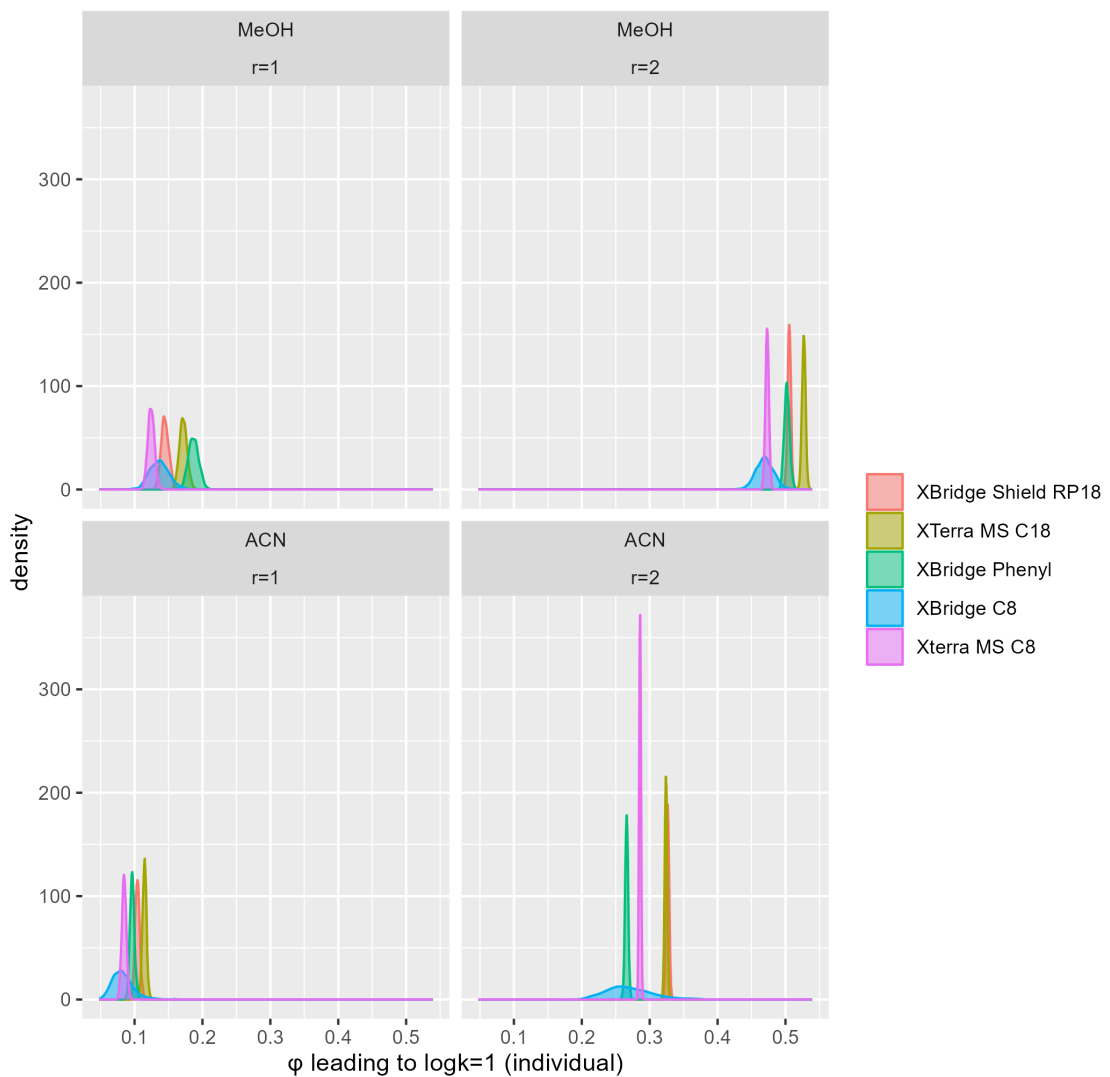
ggsave(paste0("figures\\izoparam\\", paste(dataNames>Name[analyte_ID_sample[i]]), ".filog"
}

```

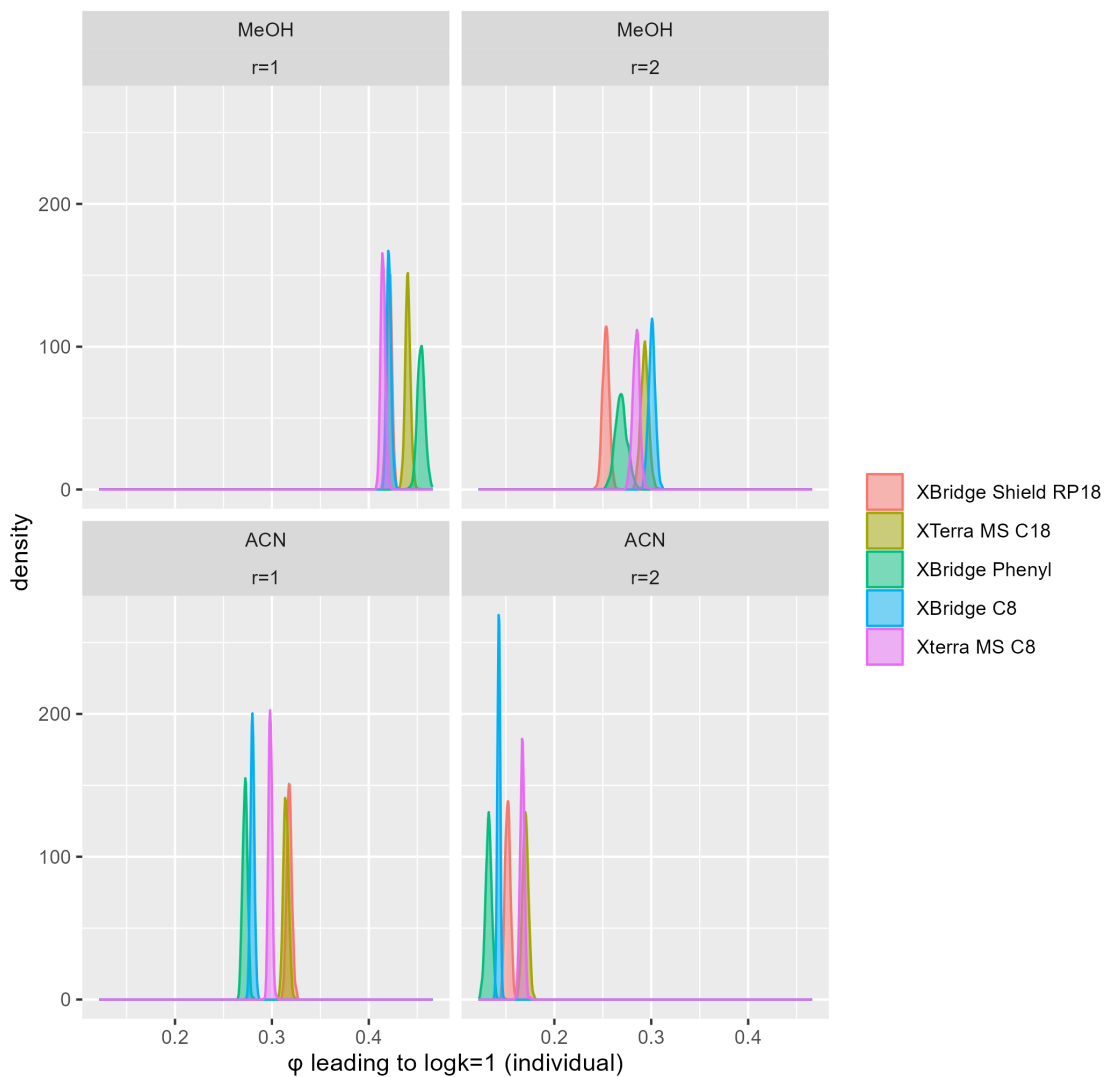
Baclofen



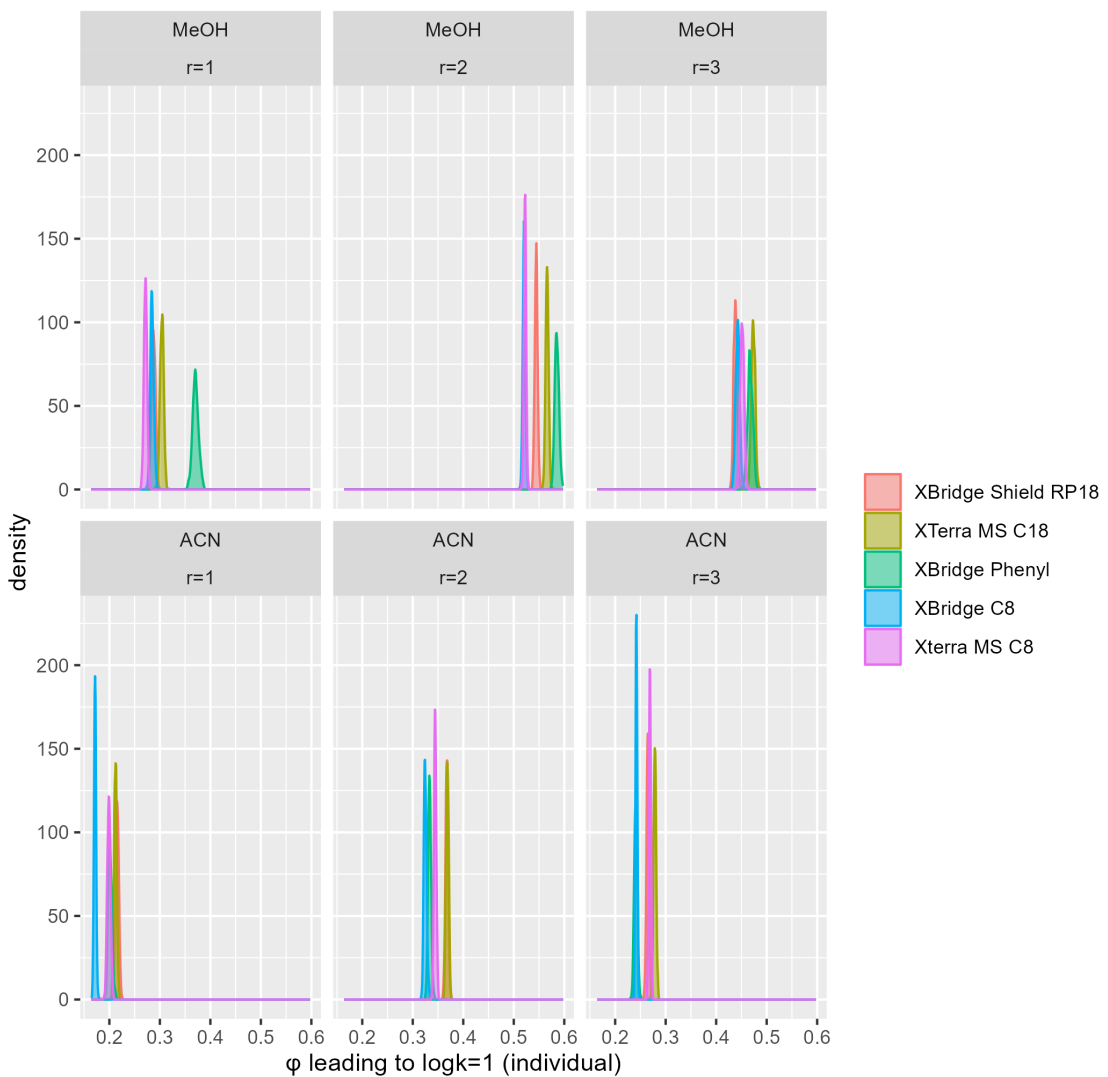
Acridine



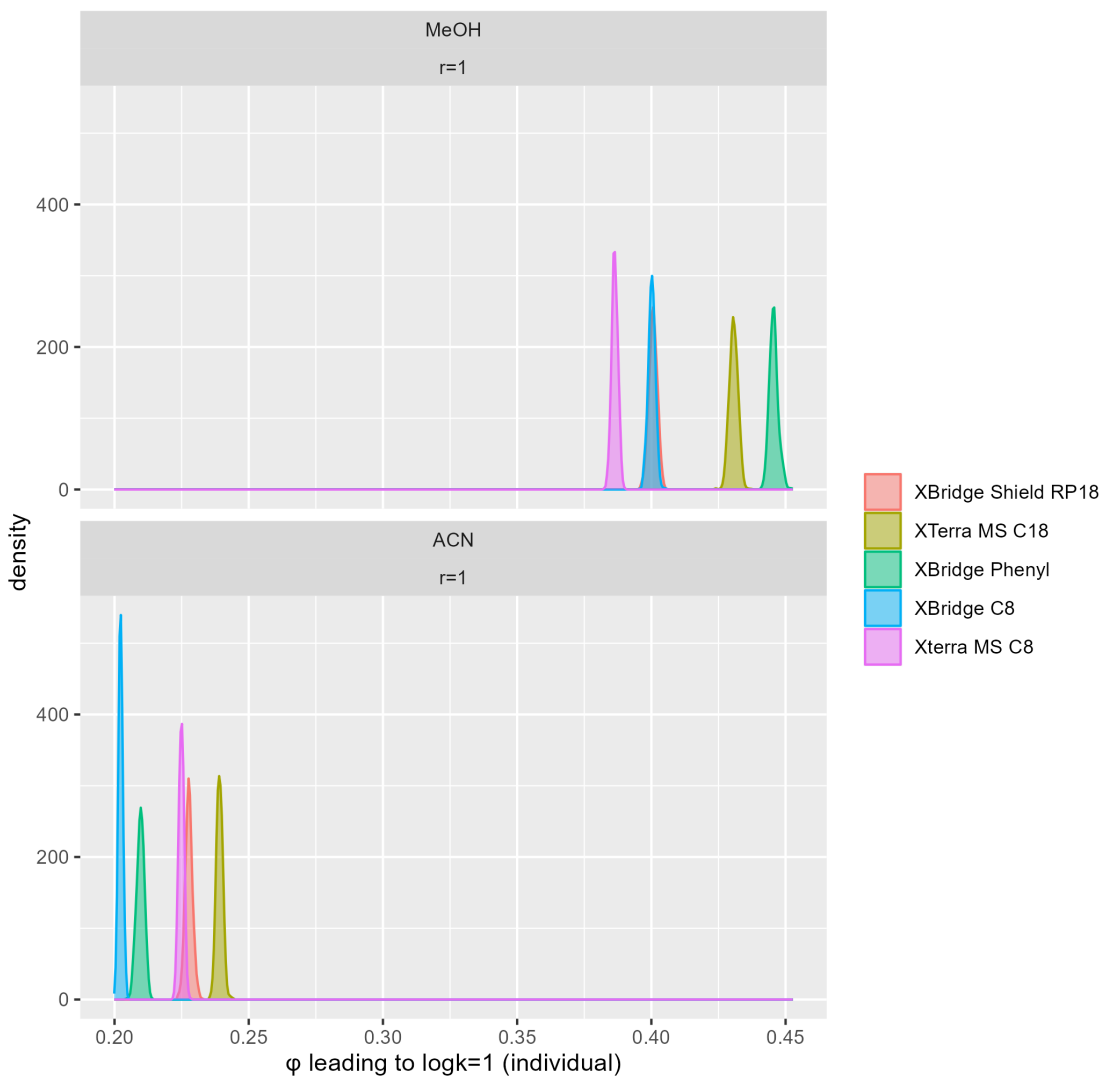
Tolbutamide

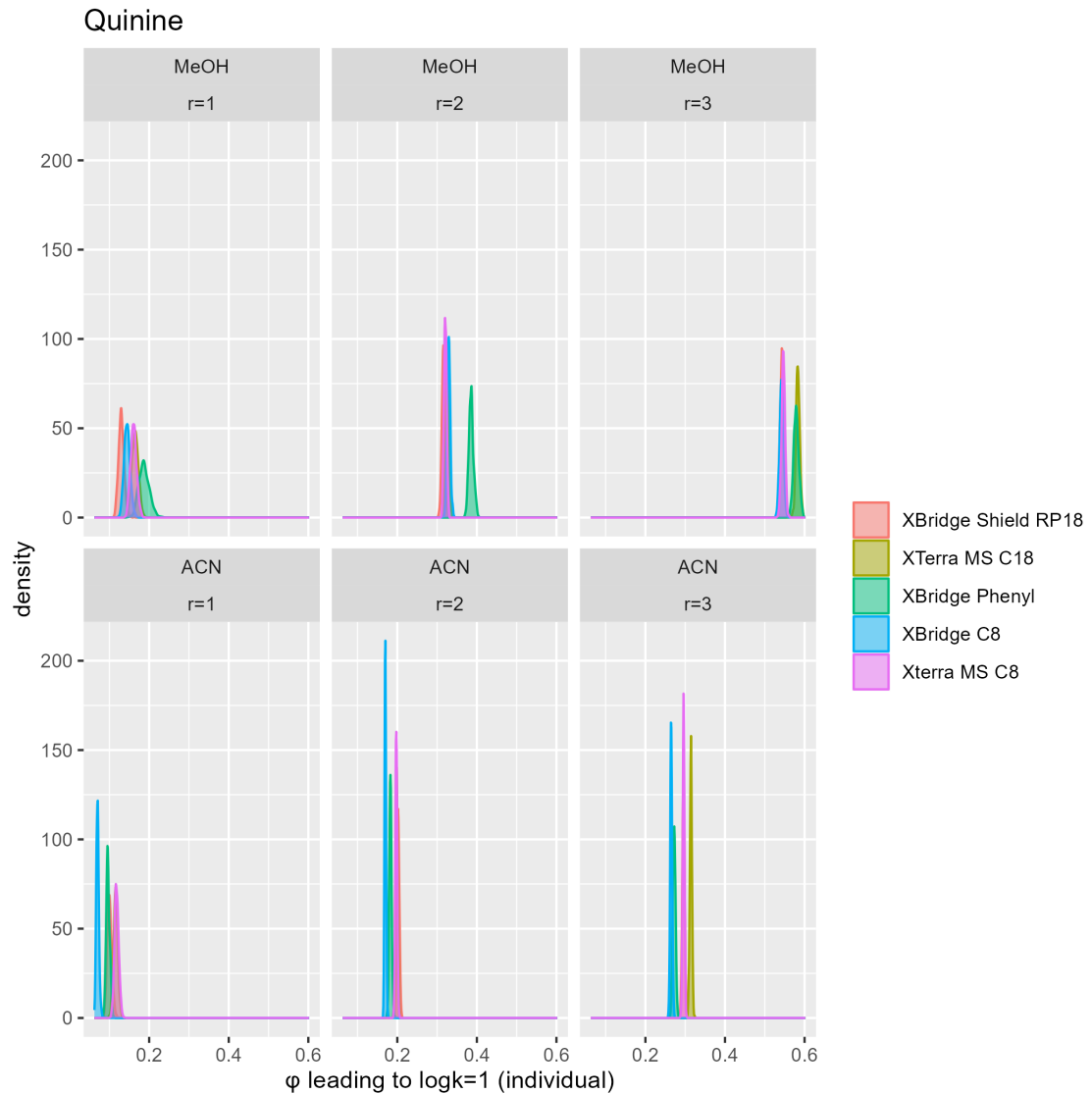


Pioglitazone



Hydrocortisone





6.5 Individual Parameters

Individual parameter are the analyte-specific parameters estimated by the model. The following plots allow to assess the correlations between these parameters.

6.5.1 Neutral Form

```
#Extract sample for plots
draws_df <- fit$draws(format = "df")
```

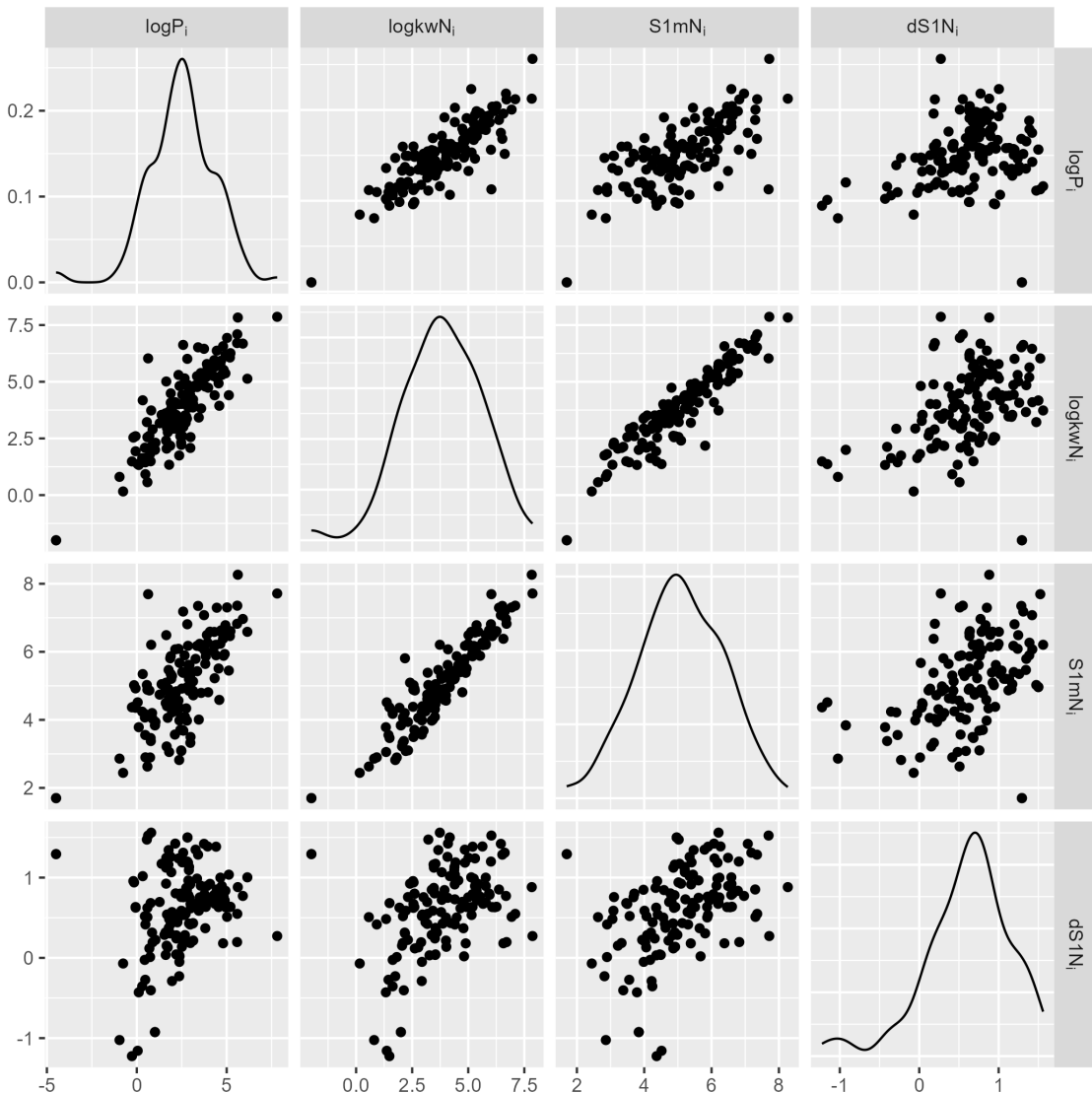
Individual parameters for the reference column (XBridge Shield RP18):

```
param <- apply(draws_df[,which(colnames(draws_df) %in% grep("^param", names(draws_df), val
param <- melt(param)
param1 <- param[1:nAnalytes,]
param2 <- param[(nAnalytes+1):(2*nAnalytes),]
param3 <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dS1N", names(draws_df), val

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)],param1,para
colnames(data_to_plot_param) <- c(expression('logP'[i]),expression('logkWN'[i]),expression

p<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param)
            labeller = "label_parsed",upper = list(continuous = "points"))

print(p)
```



```
ggsave(paste0("figures\\iparam\\", "XBridgeShieldRP18.NeutralForm", ".png"), plot=p, width=10, height=10)
```

Individual parameters for column effects

```
param <- apply(draws_df[,which(colnames(draws_df) %in% grep("clogkwN", names(draws_df), value=TRUE))], 2, mean)
param <- melt(param)
param <- matrix(param$value, nrow = nAnalytes, byrow = TRUE)
param1 <- param[,1]
param2 <- param[,2]
```

```

param3 <- param[,3]
param4 <- param[,4]

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)],param1,para
colnames(data_to_plot_param) <- c(expression('logP'[i]),expression('clogkwN1'[i]),expression

p1<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param)
             labeller = "label_parsed",upper = list(continuous = "points"))

param <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cS1mN", names(draws_df), val
param <- melt(param)
param <- matrix(param$value,nrow = nAnalytes, byrow = TRUE)
param1 <- param[,1]
param2 <- param[,2]
param3 <- param[,3]
param4 <- param[,4]

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)],param1,para
colnames(data_to_plot_param) <- c(expression('logP'[i]),expression('cS1mN1'[i]),expression

p2<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param)
             labeller = "label_parsed",upper = list(continuous = "points"))

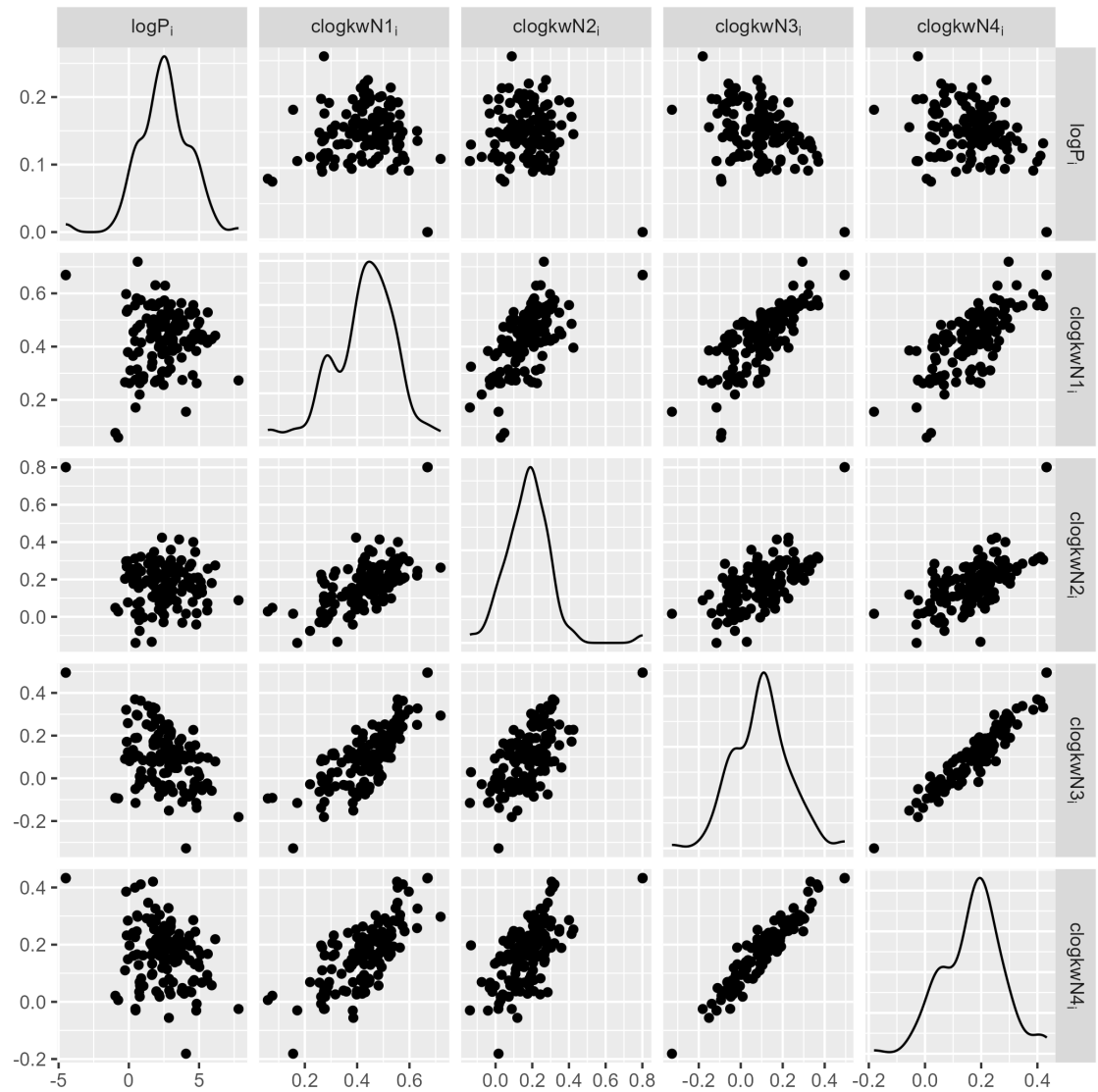
param <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cdS1N", names(draws_df), val
param <- melt(param)
param <- matrix(param$value,nrow = nAnalytes, byrow = TRUE)
param1 <- param[,1]
param2 <- param[,2]
param3 <- param[,3]
param4 <- param[,4]

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)],param1,para
colnames(data_to_plot_param) <- c(expression('logP'[i]),expression('cdS1N1'[i]),expression

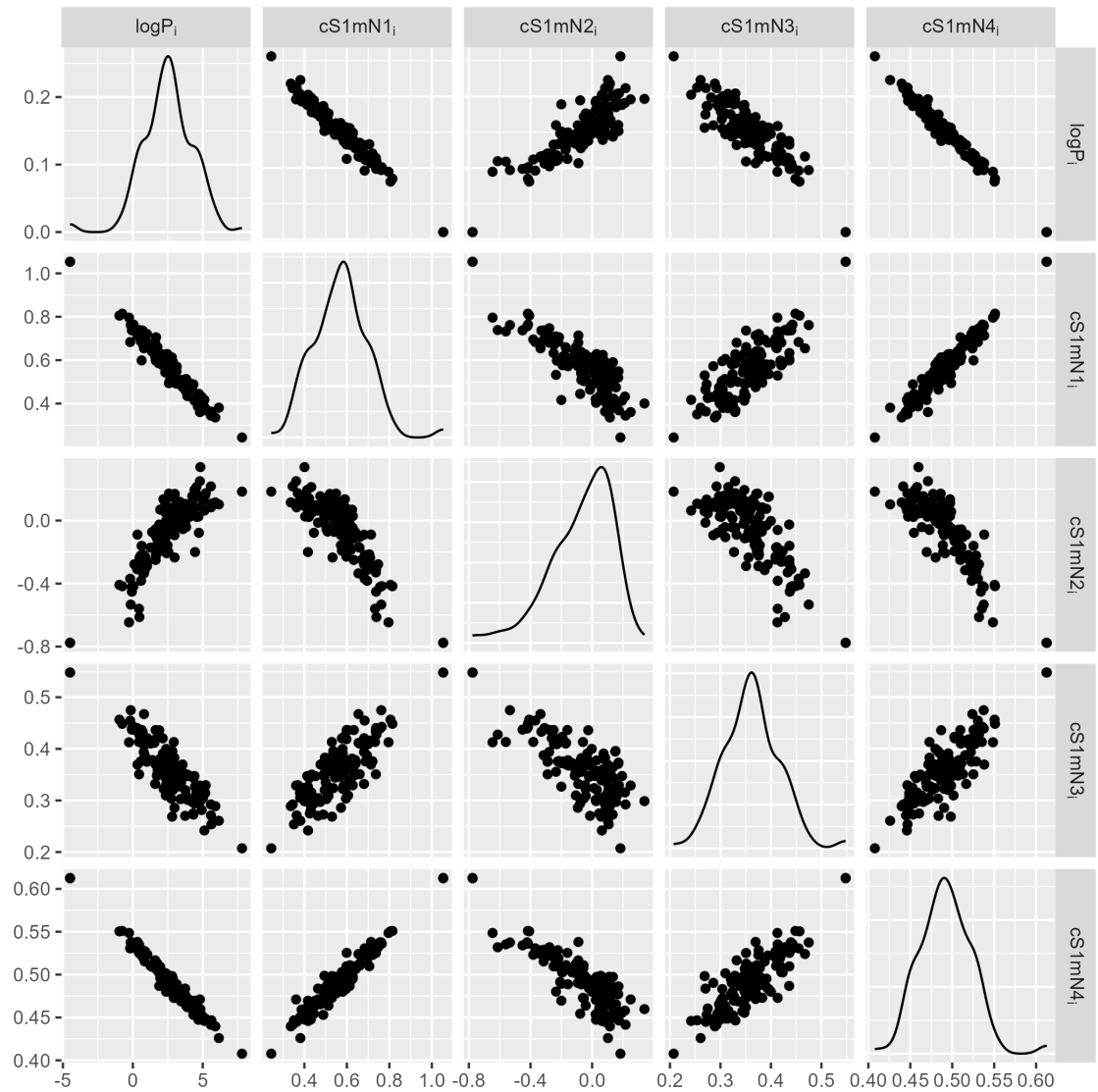
p3<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param)
             labeller = "label_parsed",upper = list(continuous = "points"))

print(p1)

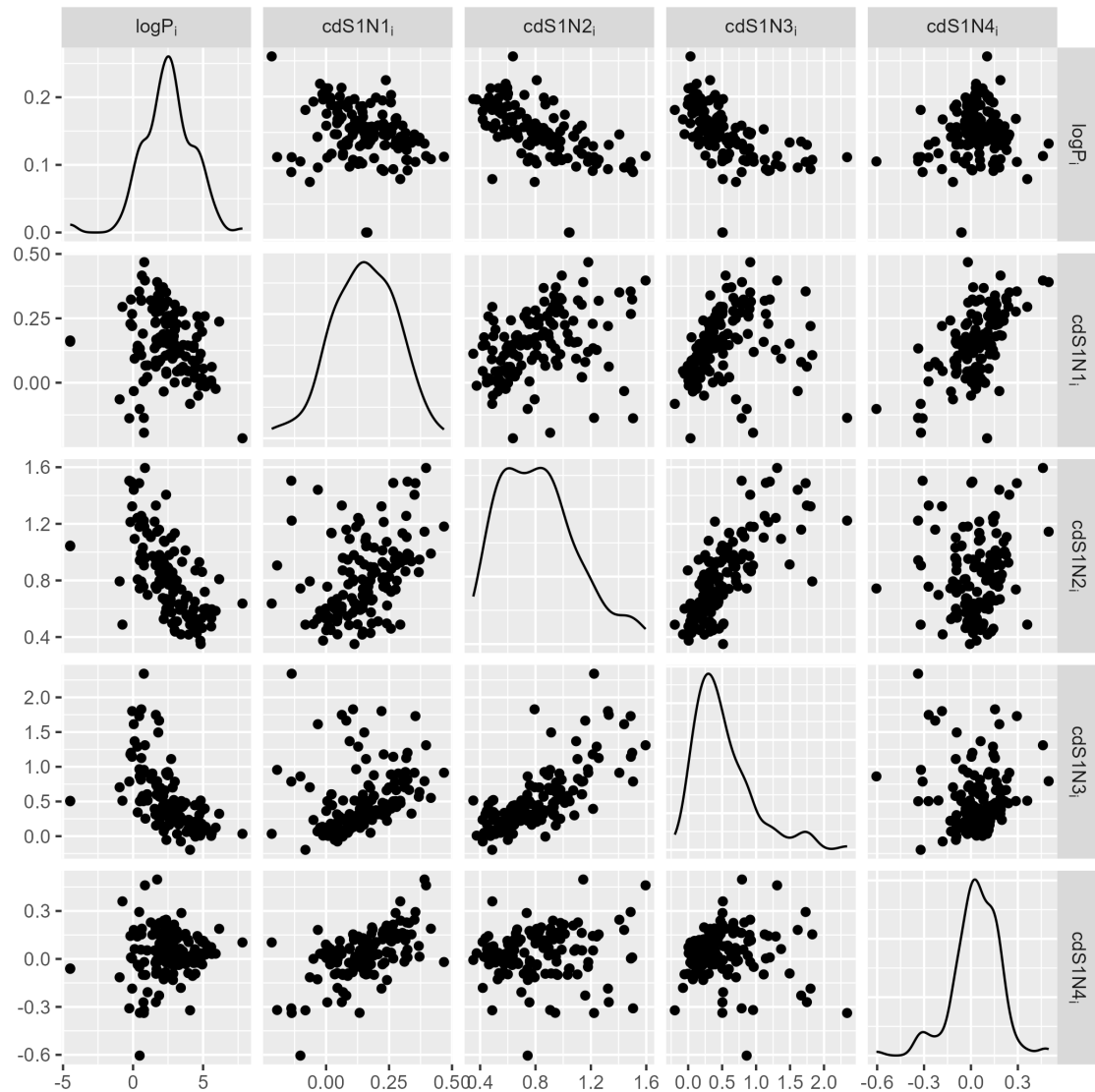
```



```
print(p2)
```



```
print(p3)
```



```

ggsave(paste0("figures\\iparam\\", "coleffects.NeutralForm_1", ".png"), plot=p1, width =
ggsave(paste0("figures\\iparam\\", "coleffects.NeutralForm_2", ".png"), plot=p2, width =
ggsave(paste0("figures\\iparam\\", "coleffects.NeutralForm_3", ".png"), plot=p3, width =

```

6.5.2 Effect of dissociation

Individual parameters for the reference column (XBridge Shield RP18)

```

dlogkwA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dlogkwA", names(draws_df),
dlogkwA <- melt(dlogkwA)[,1]

dlogkwB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dlogkwB", names(draws_df),
dlogkwB <- melt(dlogkwB)[,1]

dlogkw <- c(dlogkwA,dlogkwB)

dS1mA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dS1mA", names(draws_df), value
dS1mA <- melt(dS1mA)[,1]
dS1mB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dS1mB", names(draws_df), value
dS1mB <- melt(dS1mB)[,1]

dS1m <- c(dS1mA,dS1mB)

dS1A <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dS1A", names(draws_df), value
dS1A <- melt(dS1A)[,1]
dS1B <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dS1B", names(draws_df), value
dS1B <- melt(dS1B)[,1]

dS1 <- c(dS1A,dS1B)

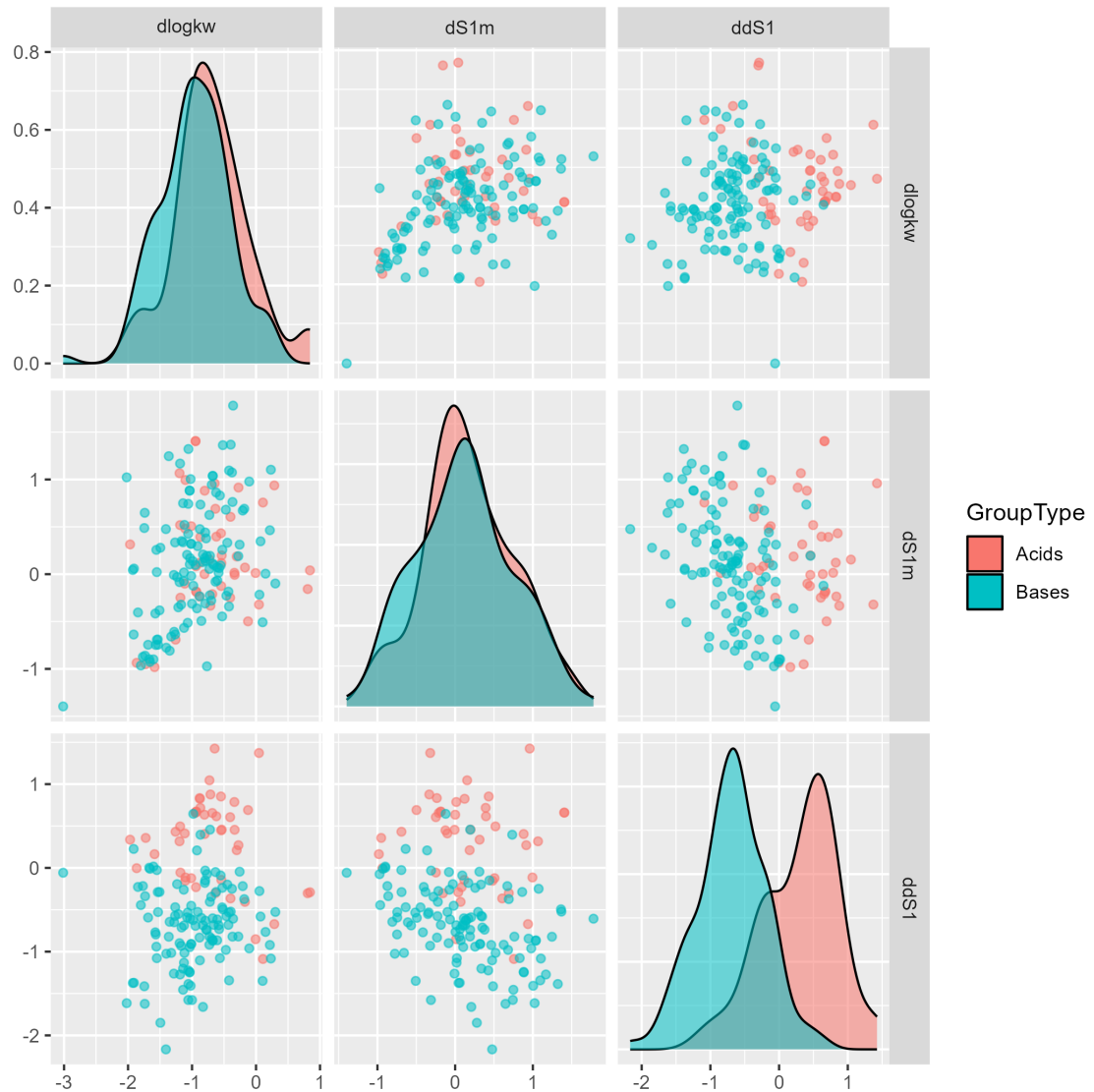
GroupType = c(rep("Acids", length(dlogkwA)),rep("Bases", length(dlogkwB)))

data_to_plot_diss <- data.frame(dlogkw, dS1m,dS1,GroupType)

p<-ggpairs(data_to_plot_diss,columns = 1:3, columnLabels = c("dlogkw","dS1m","ddS1"),
            labeller = "label_parsed",
            legend =1,
            aes(color = GroupType, alpha = 0.5),
            upper = list(continuous = "points"))+
            scale_alpha(guide = "none")

print(p)

```



```
ggsave(paste0("figures\\iparam\\", "XBridgeShieldRP18.DissForm", ".png"), plot=p, width =
```

Individual parameters for the particular column (1-4) relative to XBridge Shield RP18

```
paramA <- apply(draws_df[,which(colnames(draws_df) %in% grep("cdlogkwA", names(draws_df),
paramA <- melt(paramA)
paramA <- matrix(paramA$value,nrow = datastruct$nGroupsA, byrow = TRUE)
paramA1 <- paramA[,1]
paramA2 <- paramA[,2]
```

```

paramA3 <- paramA[,3]
paramA4 <- paramA[,4]

paramB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cdlogkwB", names(draws_df),
paramB <- melt(paramB)
paramB <- matrix(paramB$value,nrow = datastruct$nGroupsB, byrow = TRUE)
paramB1 <- paramB[,1]
paramB2 <- paramB[,2]
paramB3 <- paramB[,3]
paramB4 <- paramB[,4]

param1 <- c(paramA1,paramB1)
param2 <- c(paramA2,paramB2)
param3 <- c(paramA3,paramB3)
param4 <- c(paramA4,paramB4)

GroupType = c(rep("Acids", length(paramA1)),rep("Bases", length(paramB1)))

data_to_plot_diss <- data.frame(param1,param2,param3,param4,GroupType)

p1<-ggpairs(data_to_plot_diss,columns = 1:4, columnLabels = c("cdlogkw1","cdlogkw2","cdlogkw3"),
             labeller = "label_parsed",
             legend =1,
             aes(color = GroupType, alpha = 0.5),
             upper = list(continuous = "points"))+
             scale_alpha(guide = "none")

paramA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cdS1mA", names(draws_df),
paramA <- melt(paramA)
paramA <- matrix(paramA$value,nrow = datastruct$nGroupsA, byrow = TRUE)
paramA1 <- paramA[,1]
paramA2 <- paramA[,2]
paramA3 <- paramA[,3]
paramA4 <- paramA[,4]

paramB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cdS1mB", names(draws_df),
paramB <- melt(paramB)
paramB <- matrix(paramB$value,nrow = datastruct$nGroupsB, byrow = TRUE)
paramB1 <- paramB[,1]
paramB2 <- paramB[,2]
paramB3 <- paramB[,3]

```

```

paramB4 <- paramB[,4]

param1 <- c(paramA1,paramB1)
param2 <- c(paramA2,paramB2)
param3 <- c(paramA3,paramB3)
param4 <- c(paramA4,paramB4)

GroupType = c(rep("Acids", length(paramA1)),rep("Bases", length(paramB1)))

data_to_plot_diss <- data.frame(param1,param2,param3,param4,GroupType)

p2<-ggpairs(data_to_plot_diss,columns = 1:4, columnLabels = c("cdS1m1","cdS1m2","cdS1m3","
  labeller = "label_parsed",
  legend =1,
  aes(color = GroupType, alpha = 0.5),
  upper = list(continuous = "points"))+
  scale_alpha(guide = "none")

paramA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cdS1A", names(draws_df), value=TRUE))], 1, c)
paramA <- melt(paramA)
paramA <- matrix(paramA$value,nrow = datastruct$nGroupsA, byrow = TRUE)
paramA1 <- paramA[,1]
paramA2 <- paramA[,2]
paramA3 <- paramA[,3]
paramA4 <- paramA[,4]

paramB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^cdS1B", names(draws_df), value=TRUE))], 1, c)
paramB <- melt(paramB)
paramB <- matrix(paramB$value,nrow = datastruct$nGroupsB, byrow = TRUE)
paramB1 <- paramB[,1]
paramB2 <- paramB[,2]
paramB3 <- paramB[,3]
paramB4 <- paramB[,4]

param1 <- c(paramA1,paramB1)
param2 <- c(paramA2,paramB2)
param3 <- c(paramA3,paramB3)
param4 <- c(paramA4,paramB4)

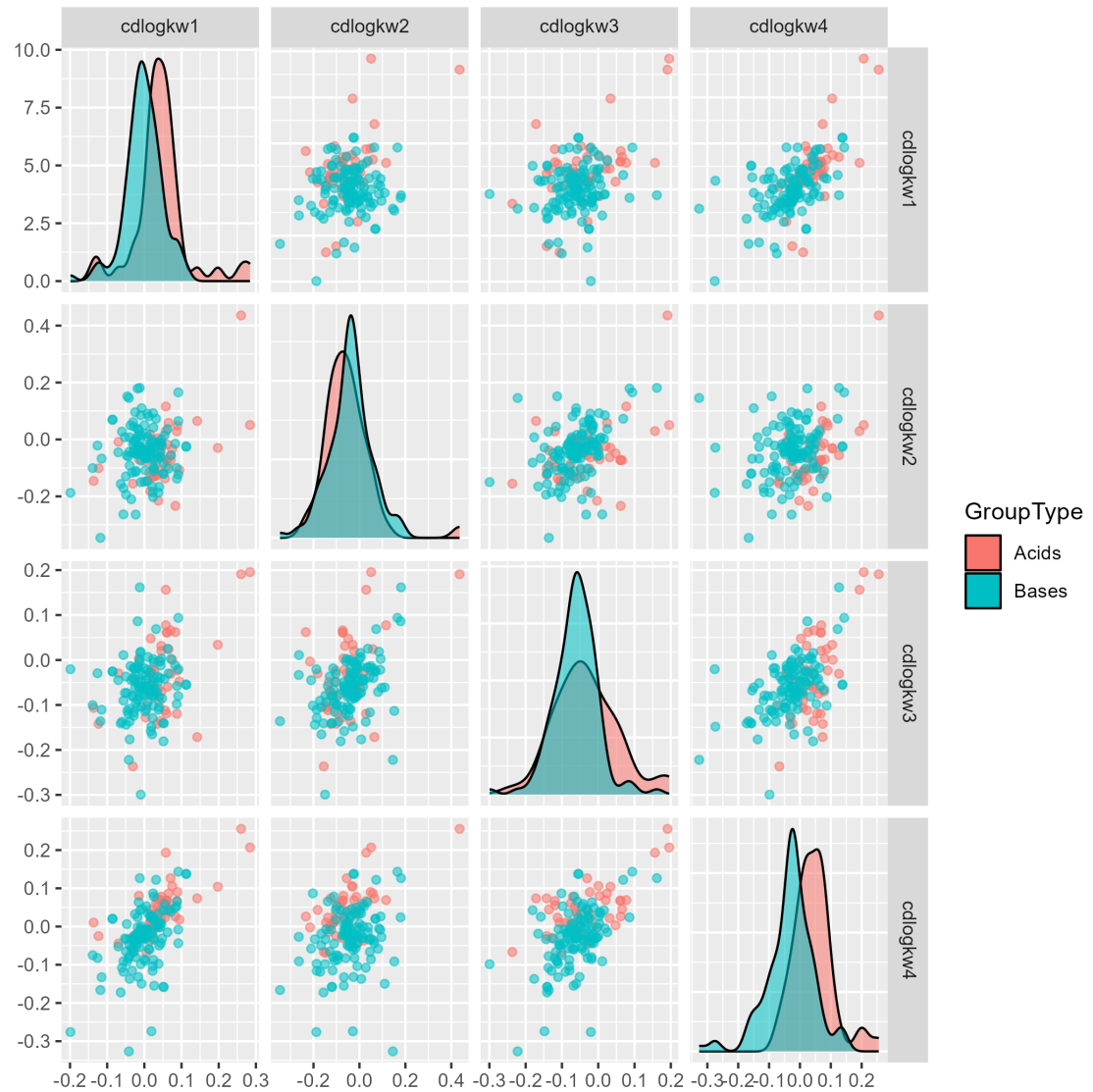
GroupType = c(rep("Acids", length(paramA1)),rep("Bases", length(paramB1)))

```

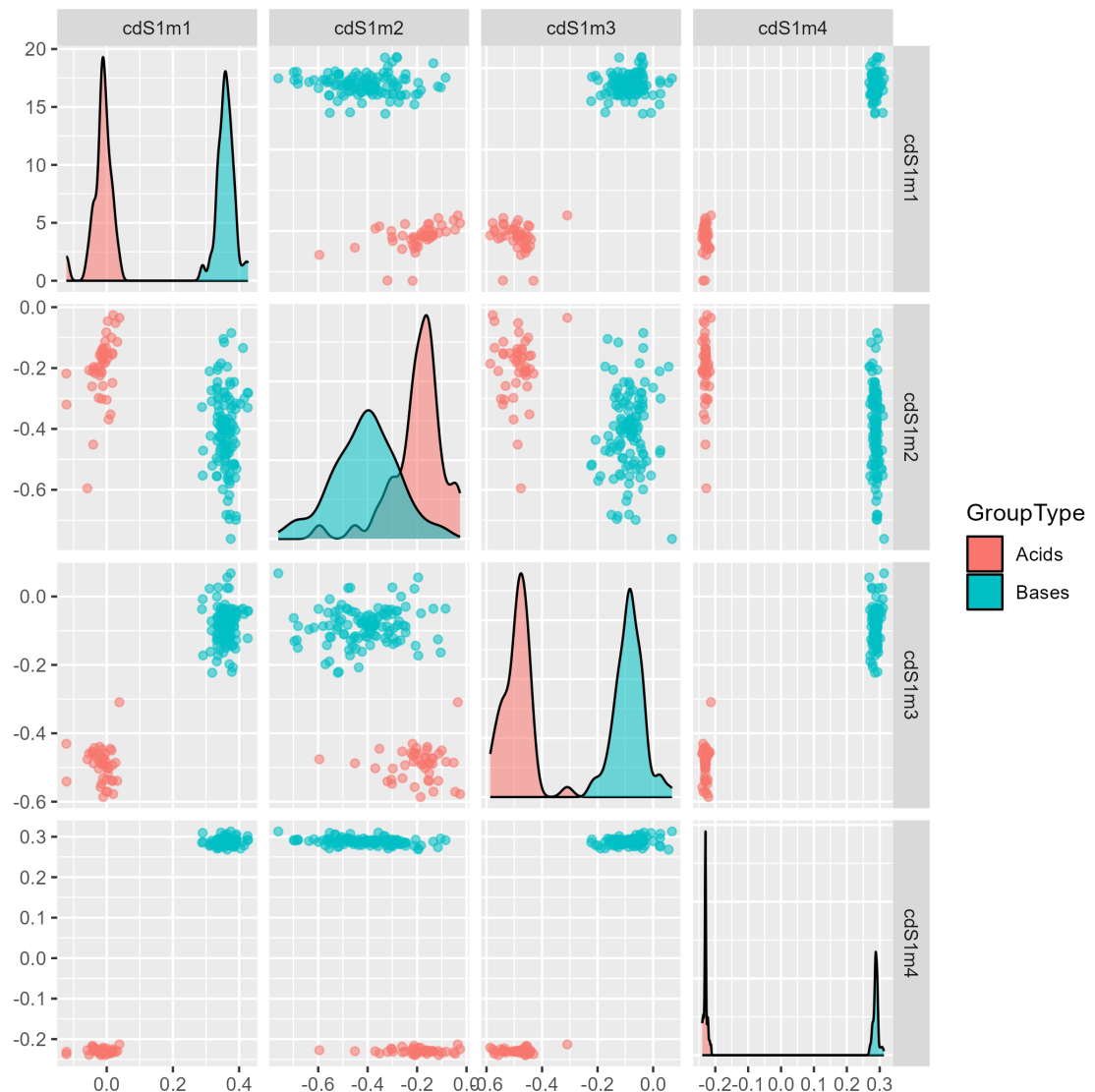
```
data_to_plot_diss <- data.frame(param1,param2,param3,param4,GroupType)

p3<-ggpairs(data_to_plot_diss,columns = 1:4, columnLabels = c("cdS11","cdS12","cdS13","cdS14"),
             labeller = "label_parsed",
             legend =1,
             aes(color = GroupType, alpha = 0.5),
             upper = list(continuous = "points"))+
             scale_alpha(guide = "none")

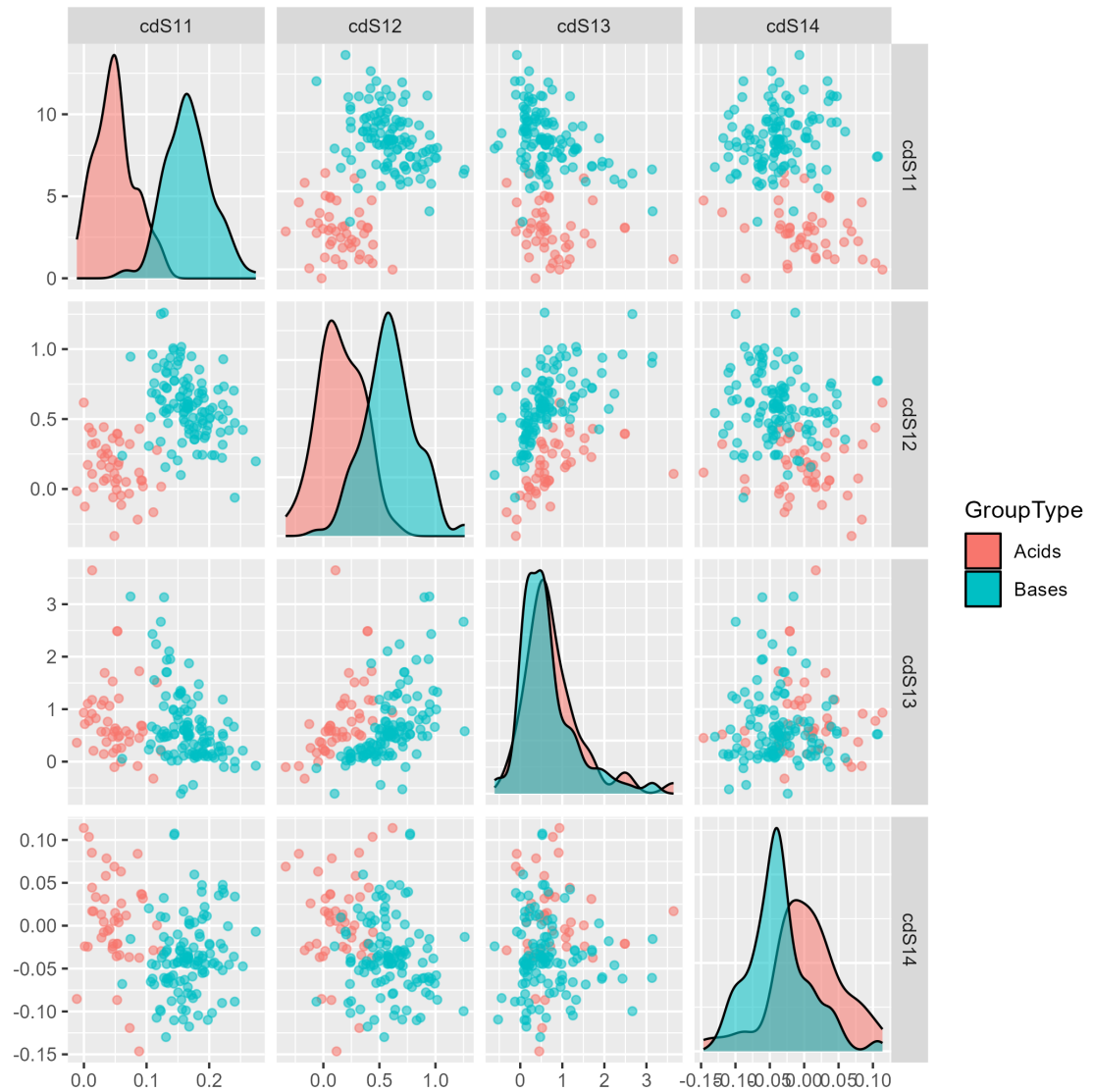
print(p1)
```



```
print(p2)
```



```
print(p3)
```



```
ggsave(paste0("figures\\iparam\\", "Differences.DissForm_1", ".png"), plot=p1, width = 20,
ggsave(paste0("figures\\iparam\\", "Differences.DissForm_2", ".png"), plot=p2, width = 20,
ggsave(paste0("figures\\iparam\\", "Differences.DissForm_3", ".png"), plot=p3, width = 20,
```

6.5.3 pKa-related paraemters

```
pKawA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^pKawA", names(draws_df), val
pKawA <- melt(pKawA)[,1]
pKawB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^pKawB", names(draws_df), val
pKawB <- melt(pKawB)[,1]

alphamA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^alphamA", names(draws_df),
alphamA <- melt(alphamA)[,1]
alphamB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^alphamB", names(draws_df),
alphamB <- melt(alphamB)[,1]

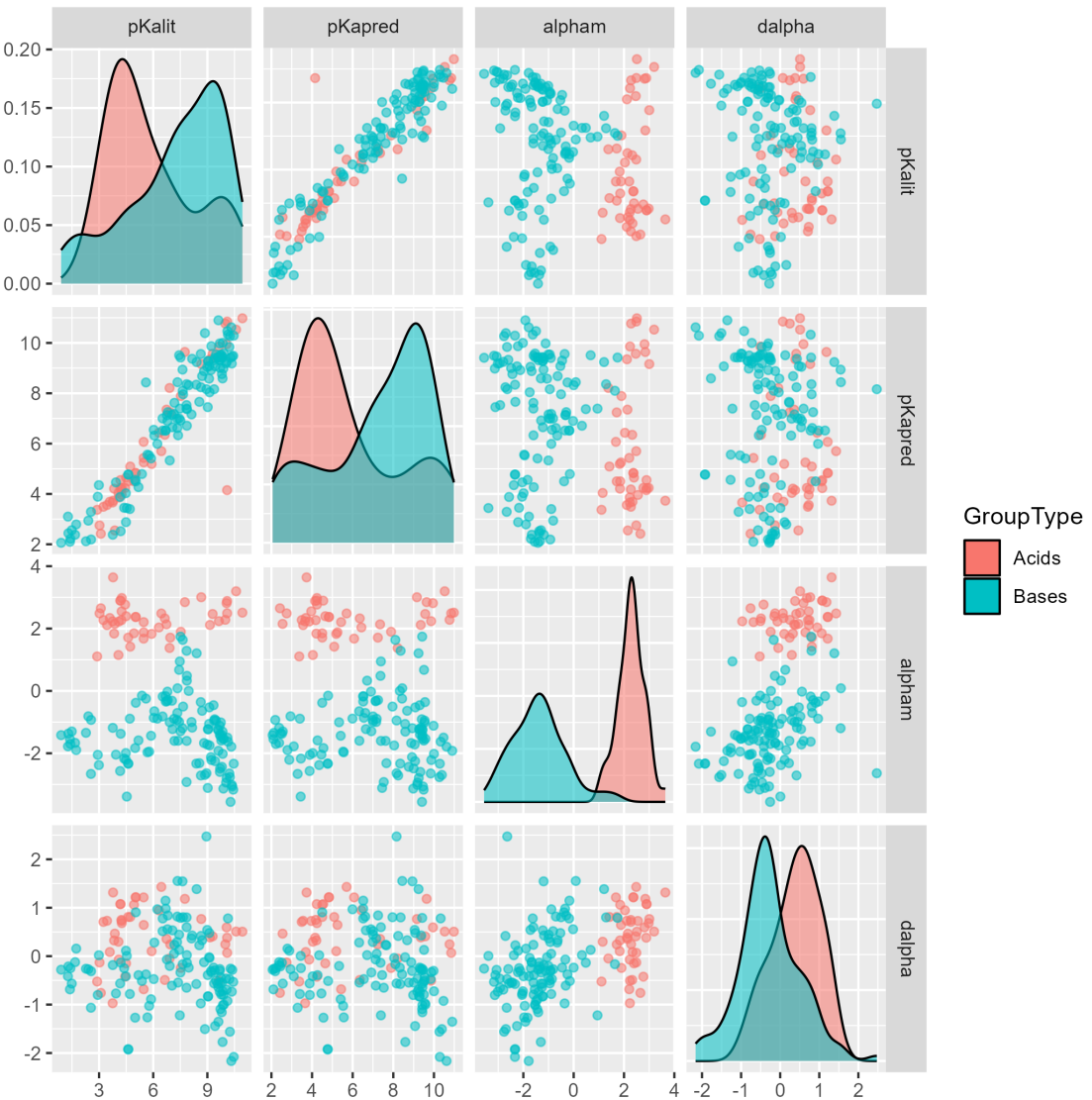
dalpA <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dalpA", names(draws_df),
dalpA <- melt(dalpA)[,1]
dalpB <- apply(draws_df[,which(colnames(draws_df) %in% grep("^dalpB", names(draws_df),
dalpB <- melt(dalpB)[,1]

pKapred <- c(pKawA, pKawB)
pKaslit <- c(pKaslitA, pKaslitB)
alpham <- c(alphamA, alphamB)
dalpA <- c(dalpA, dalpB)

GroupType = c(rep("Acids", length(alphamA)),rep("Bases", length(alphamB)))

data_to_plot_param <- data.frame(pKapred, pKaslit, alpham, dalpA, GroupType)

p<-ggpairs(data_to_plot_param,columns = 1:4, columnLabels = c("pKalit", "pKapred", "alpham",
    labeller = "label_parsed",
    legend =1,
    aes(color = GroupType, alpha = 0.5),
    upper = list(continuous = "points"))+
    scale_alpha(guide = "none")
print(p)
```



```
ggsave(paste0("figures\\iparam\\", "pKas", ".png"), plot=p, width = 20, height = 20, units
```

6.6 Eta plots

Eta plots shows the centered and standardized individual parameters (e.g. $\eta_{logkwN,i} = (logkwN_i - (\hat{logkwN} + \beta_1 \cdot \log P_i)) / \omega_1$). They allow to visualize the unexplained between-analyte variability of chromatographic parameters.

```

model_etas <- cmdstan_model("stan/hplc-gra-fivecolumns-etas.stan")
fit_etas  <- model_etas$generate_quantities(fit,
                                              data = datastruct,
                                              seed = 123,
                                              parallel_chains = 4,
                                              output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-etas-202308081442-1-6560',
  'stanfiles/hplc-gra-fivecolumns-etas-202308081442-2-6560',
  'stanfiles/hplc-gra-fivecolumns-etas-202308081442-3-6560',
  'stanfiles/hplc-gra-fivecolumns-etas-202308081442-4-6560
))

draws_etas_df <- as_draws_df(x$generated_quantities)

# draws_etas_df <- fit_etas$draws(format = "df")

```

6.6.1 Neutral Forms

etalogkwN:

```

param1 <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etalogkwN", names
  draws_etas_df))], 1)

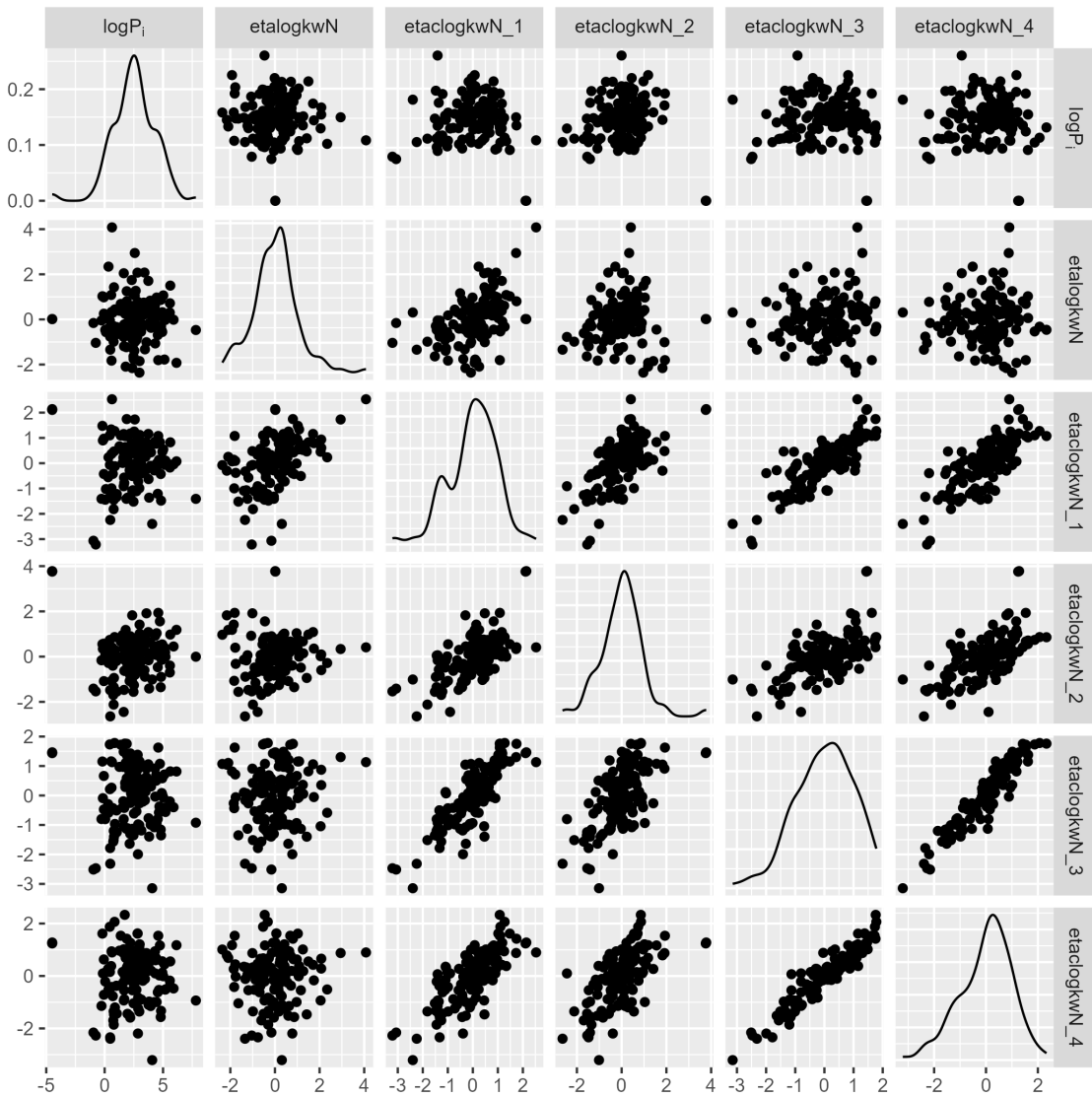
cparam <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etaclogkwNc", names
  draws_etas_df))], 1)
cparam <- melt(cparam)
cparam <- matrix(cparam$value, nrow = nAnalytes, byrow = TRUE)
cparam1 <- cparam[,1]
cparam2 <- cparam[,2]
cparam3 <- cparam[,3]
cparam4 <- cparam[,4]

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)], param1, cparam)
colnames(data_to_plot_param) <- c(expression('logP'[i]), expression('etalogkwN'), expression('etaclogkwNc'))

p<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param),
            labeller = "label_parsed", upper = list(continuous = "points"))

print(p)

```



```
ggsave(paste0("figures\\etaplots\\", "neutralforms_1", ".png"), plot=p, width = 20, height = 15)
```

etaS1mN and etadS1N:

```
param1 <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etaS1mN", names(draws_etas_df)))] , 2, mean)

cparam <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etacS1mNc", names(draws_etas_df)))] , 2, mean)
cparam <- melt(cparam)
cparam <- matrix(cparam$value, nrow = nAnalytes, byrow = TRUE)
```

```

cparam1 <- cparam[,1]
cparam2 <- cparam[,2]
cparam3 <- cparam[,3]
cparam4 <- cparam[,4]

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)],param1,cparam)
colnames(data_to_plot_param) <- c(expression('logP'[i]),expression('etaS1mN'),expression('

p2<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param)
               labeller = "label_parsed",upper = list(continuous = "points"))

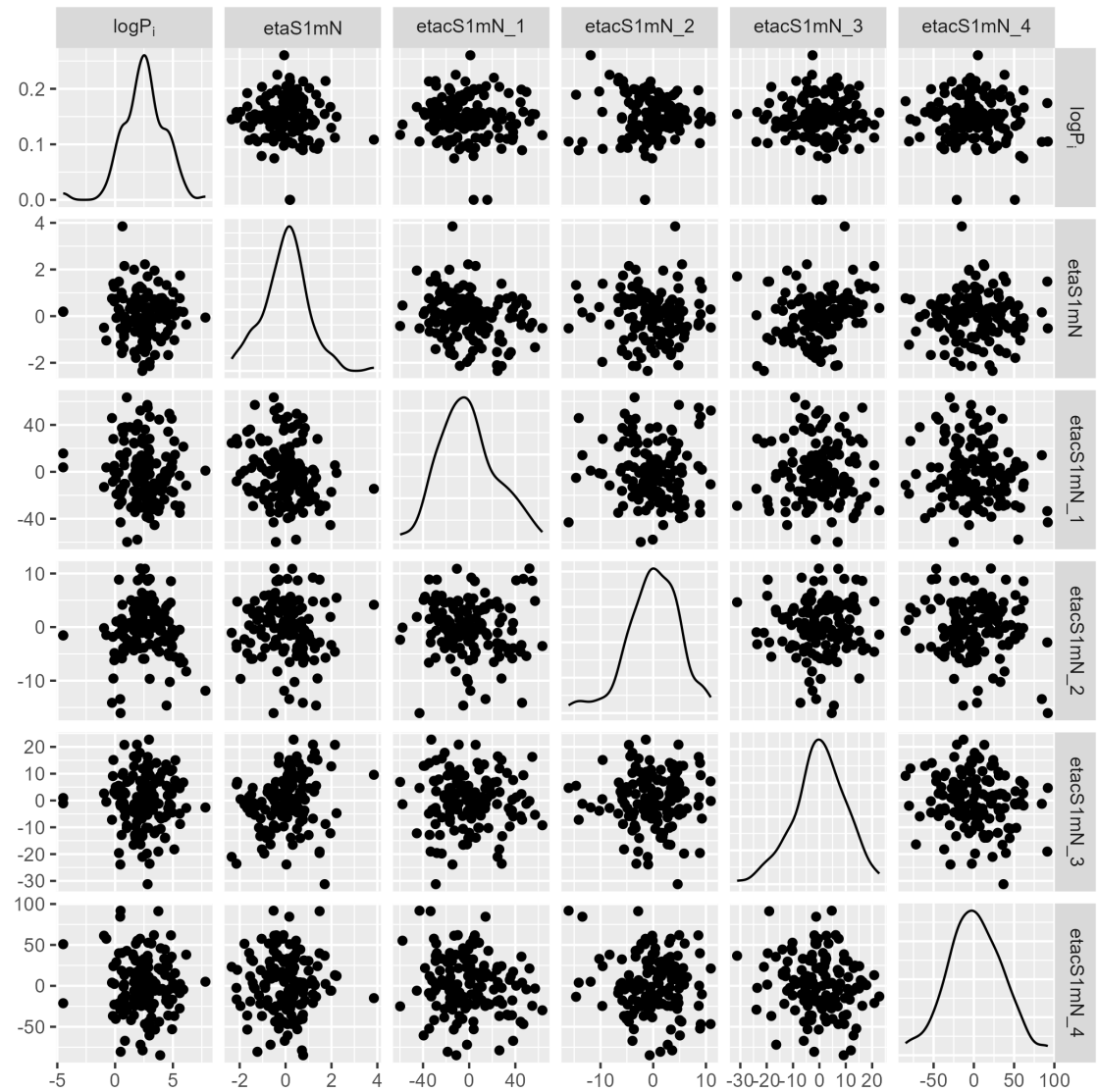
param1 <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etadS1N", names(draws_etas_df)))] ,1,mean)
cparam <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etacdS1Nc", names(draws_etas_df)))] ,1,mean)
cparam <- melt(cparam)
cparam <- matrix(cparam$value,nrow = nAnalytes, byrow = TRUE)
cparam1 <- cparam[,1]
cparam2 <- cparam[,2]
cparam3 <- cparam[,3]
cparam4 <- cparam[,4]

data_to_plot_param <- cbind(dataACD$logP[which(dataACD$METID %in% data$METID)],param1,cparam)
colnames(data_to_plot_param) <- c(expression('logP'[i]),expression('etadS1N'),expression('

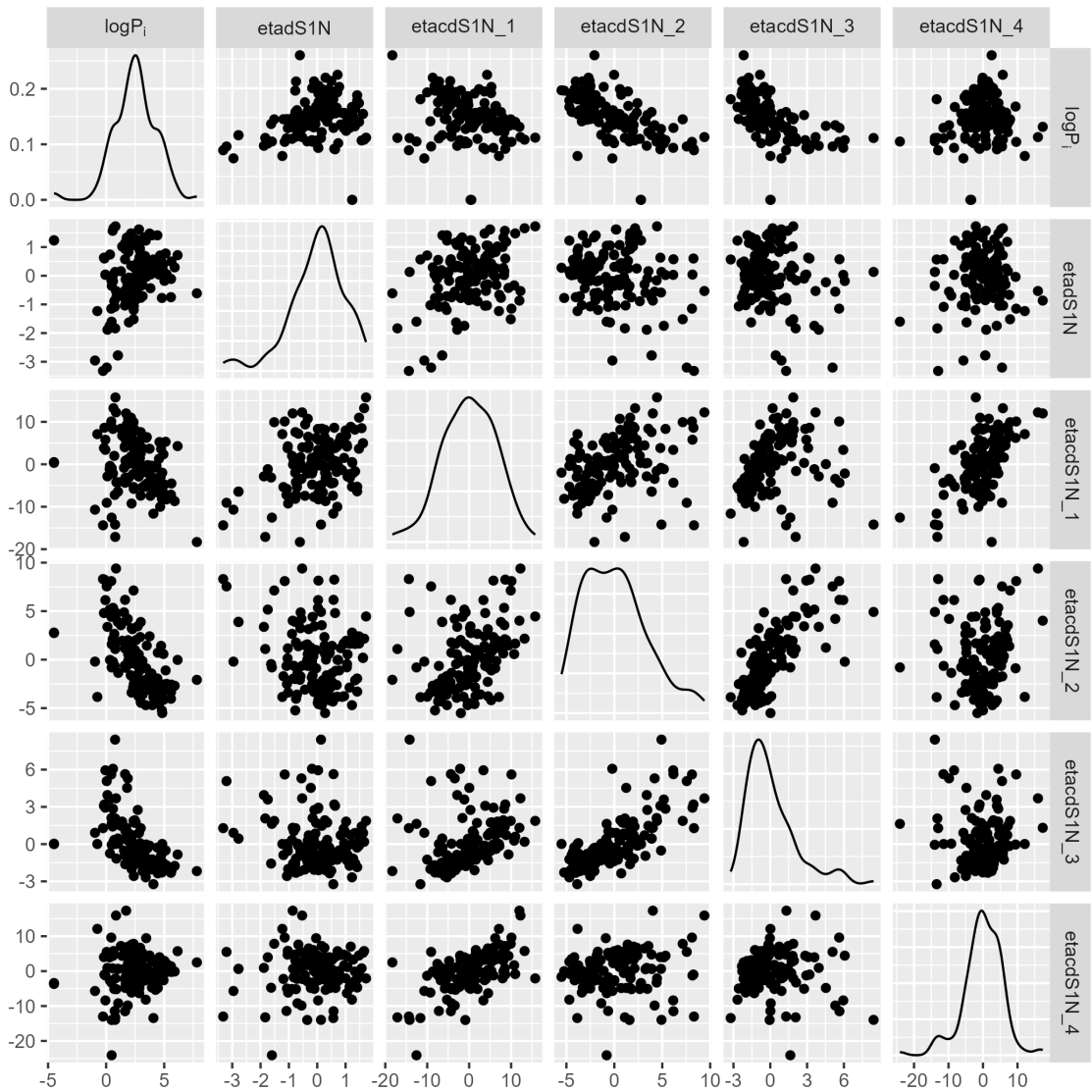
p3<-ggpairs(as.data.frame(data_to_plot_param), columnLabels = colnames(data_to_plot_param)
               labeller = "label_parsed",upper = list(continuous = "points"))

print(p2)

```



```
print(p3)
```



```
ggsave(paste0("figures\\etaplots\\", "neutralforms_2", ".png"), plot=p2, width = 20, height = 15)
ggsave(paste0("figures\\etaplots\\", "neutralforms_3", ".png"), plot=p3, width = 20, height = 15)
```

6.6.2 Effect of functional groups (exploratory)

The following ETA plots present the relationship between individual eta values for logkwN and number of functional groups. This part is exploratory. Graphs are shown if there are at least 10 functional groups present in the dataset.

```

param1 <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etalogkwN", names

cparam <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etaclogkwNc", nam
cparam <- melt(cparam)
cparam <- matrix(cparam$value,nrow = nAnalytes, byrow = TRUE)
cparam1 <- cparam[,1]
cparam2 <- cparam[,2]
cparam3 <- cparam[,3]
cparam4 <- cparam[,4]

for(i in which(totalnrgroups>10)){
nrgroups = as.factor(nrfungroups[,i])

data_to_plot_fungr <- data.frame(param1,cparam1,cparam2,cparam3,cparam4, nrgroups)

p1<-data_to_plot_fungr %>% tidyverse::gather("name", "count", 1:5) %>%
  mutate(etaname = case_when(name == "param1" ~ "etalogkwN",
                             name == "cparam1" ~ "etaclogkwN_1",
                             name == "cparam2" ~ "etaclogkwN_2",
                             name == "cparam3" ~ "etaclogkwN_3",
                             name == "cparam4" ~ "etaclogkwN_4"
                           )) %>%
  ggplot(., aes(y=count, x=nrgroups))+
  geom_boxplot()+
  facet_wrap(.~etaname)+
  labs(title=paste(functionalgroupsnames[i,2]), x="Nr of functional groups", y="eta")

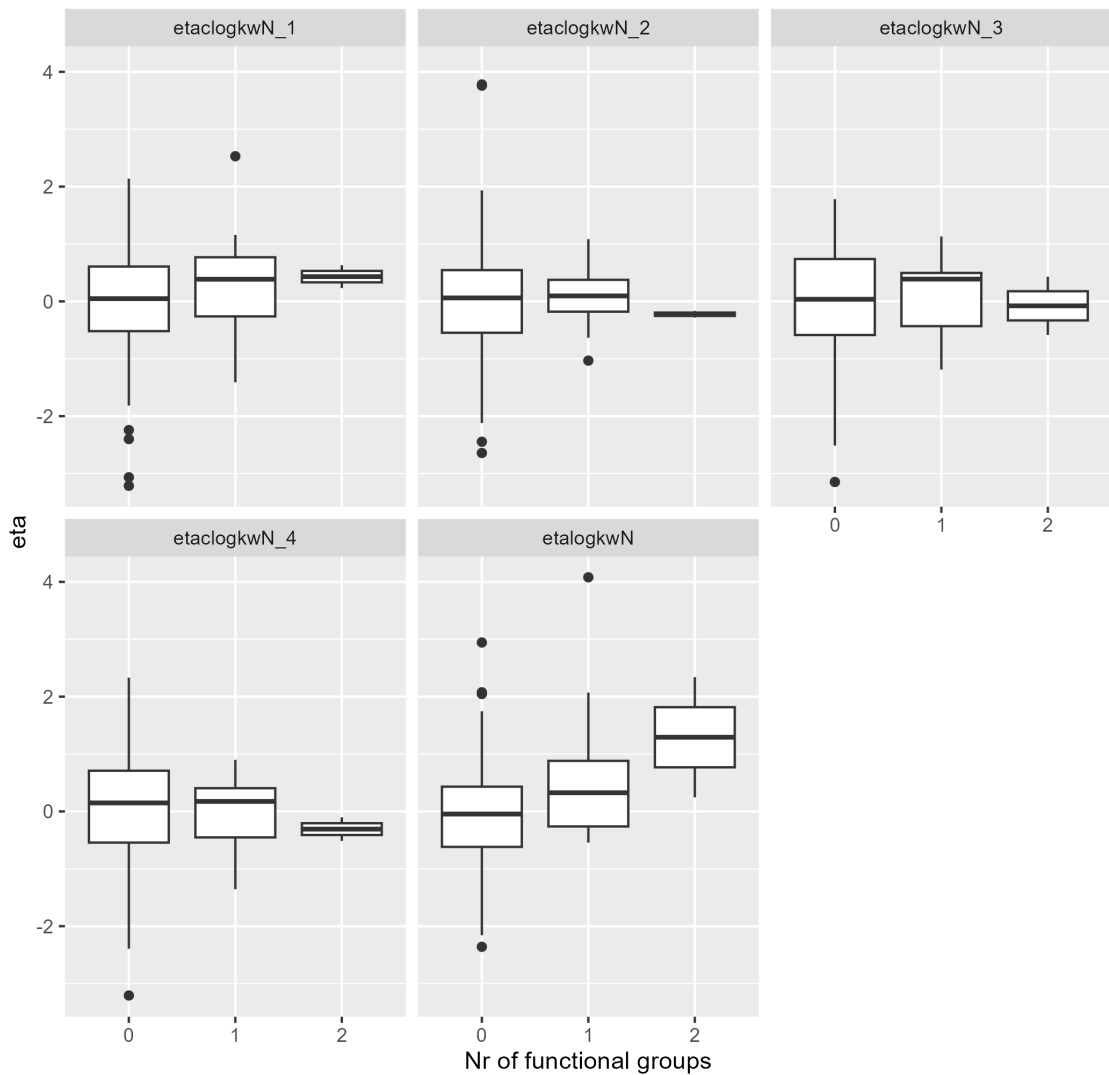
print(p1)

ggsave(paste0("figures\\etaplots\\", functionalgroupsnames[i,2], ".functionalgroupseffect

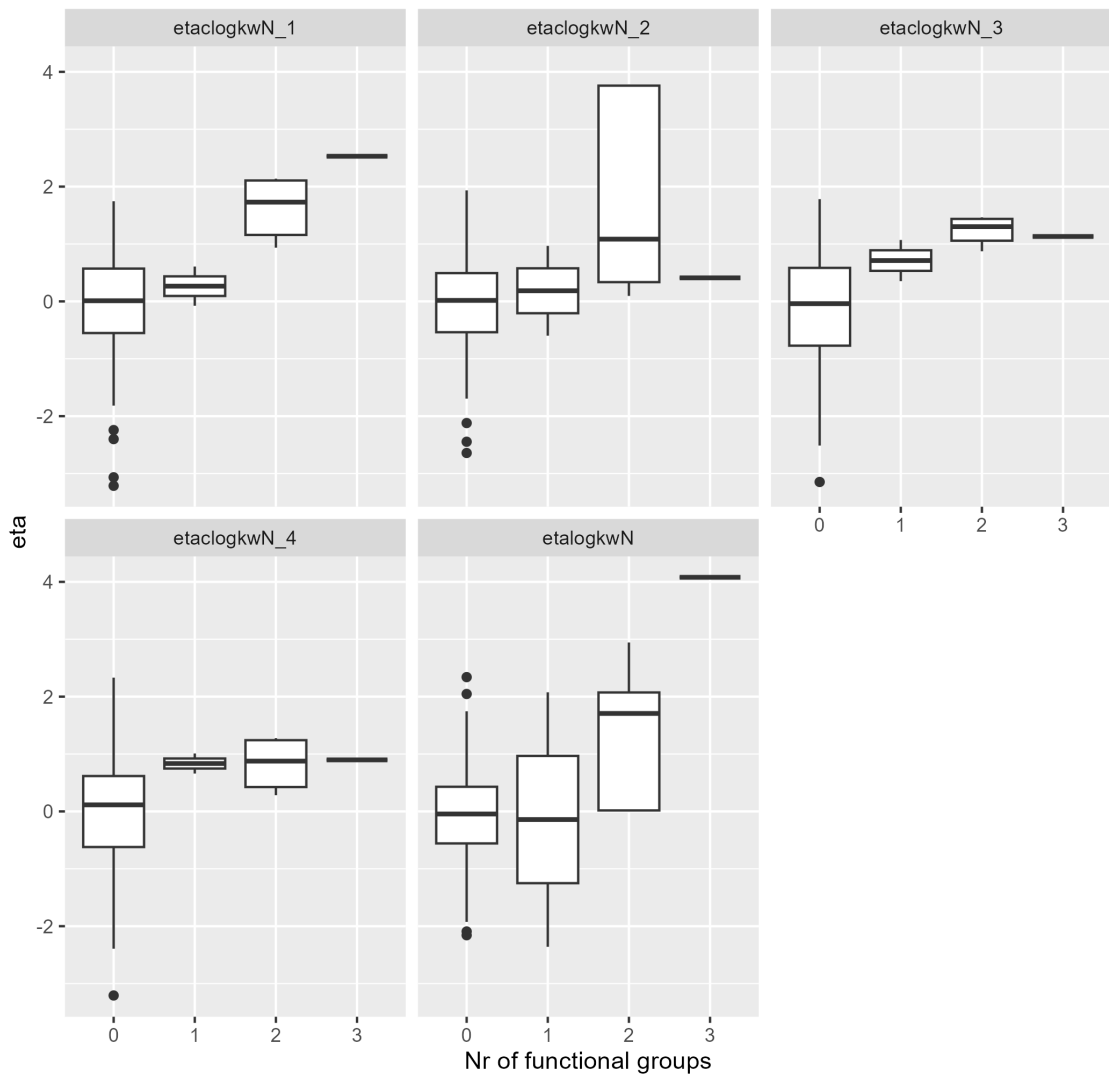
}

```

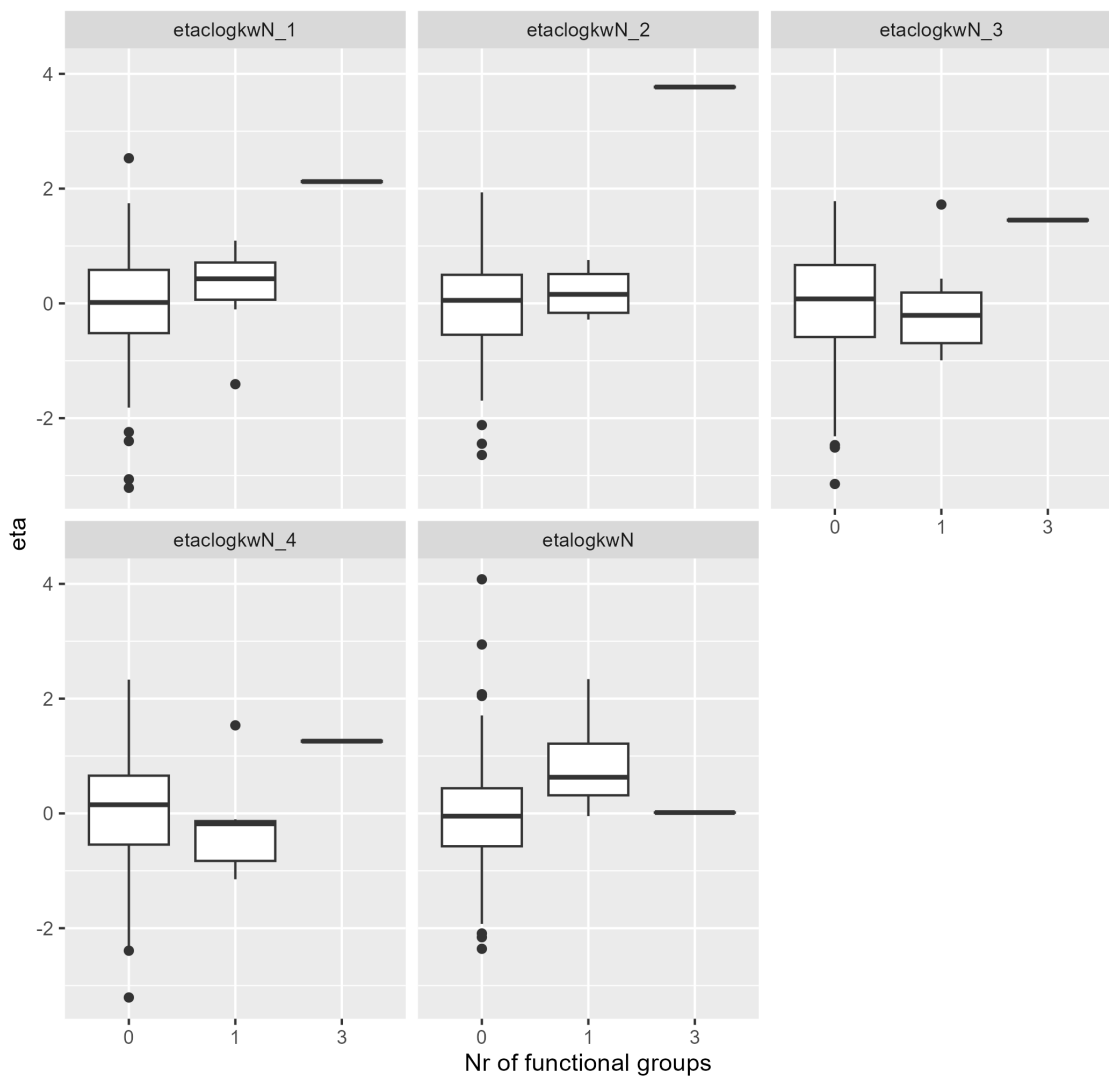
ketone



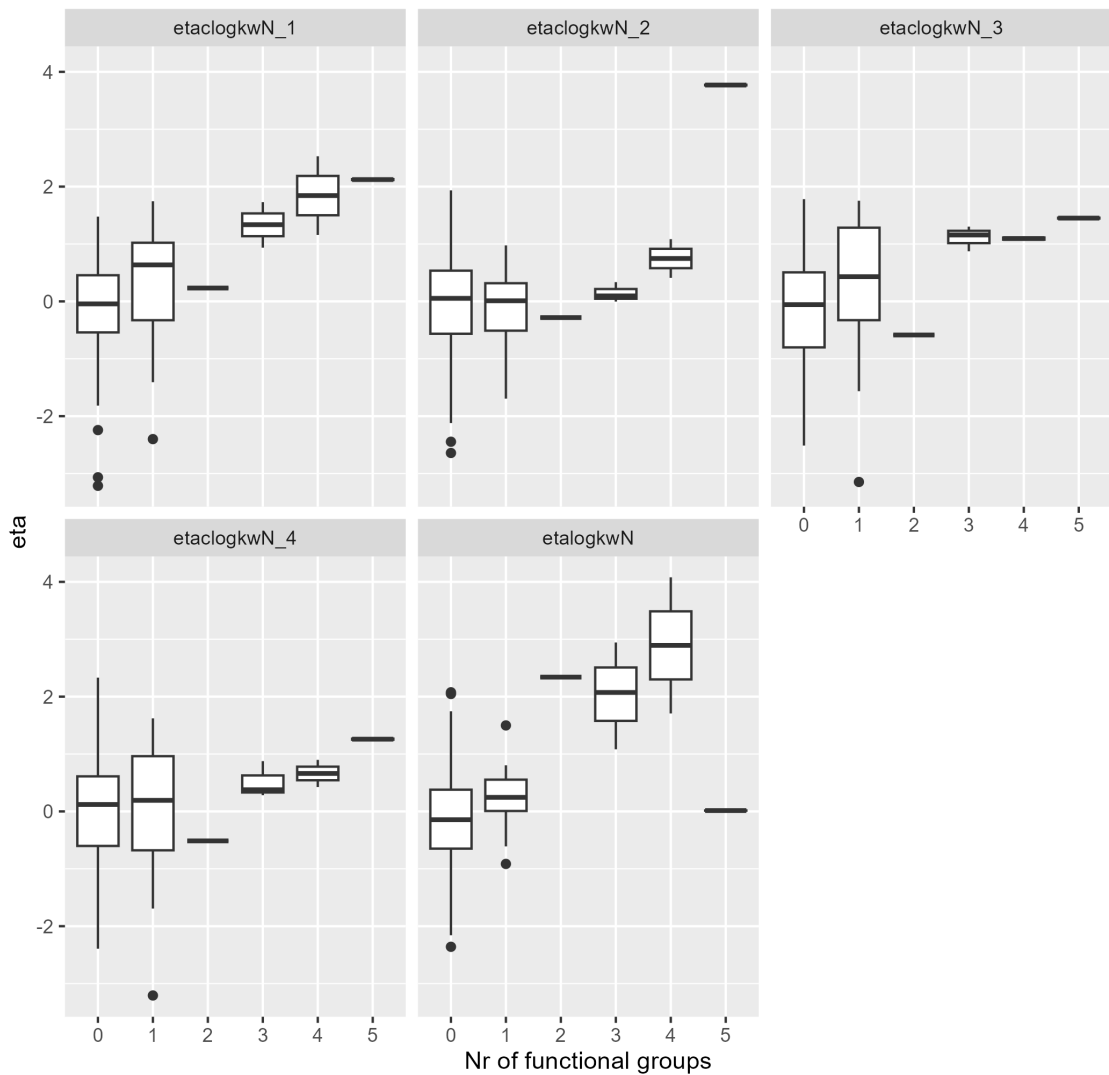
acetal



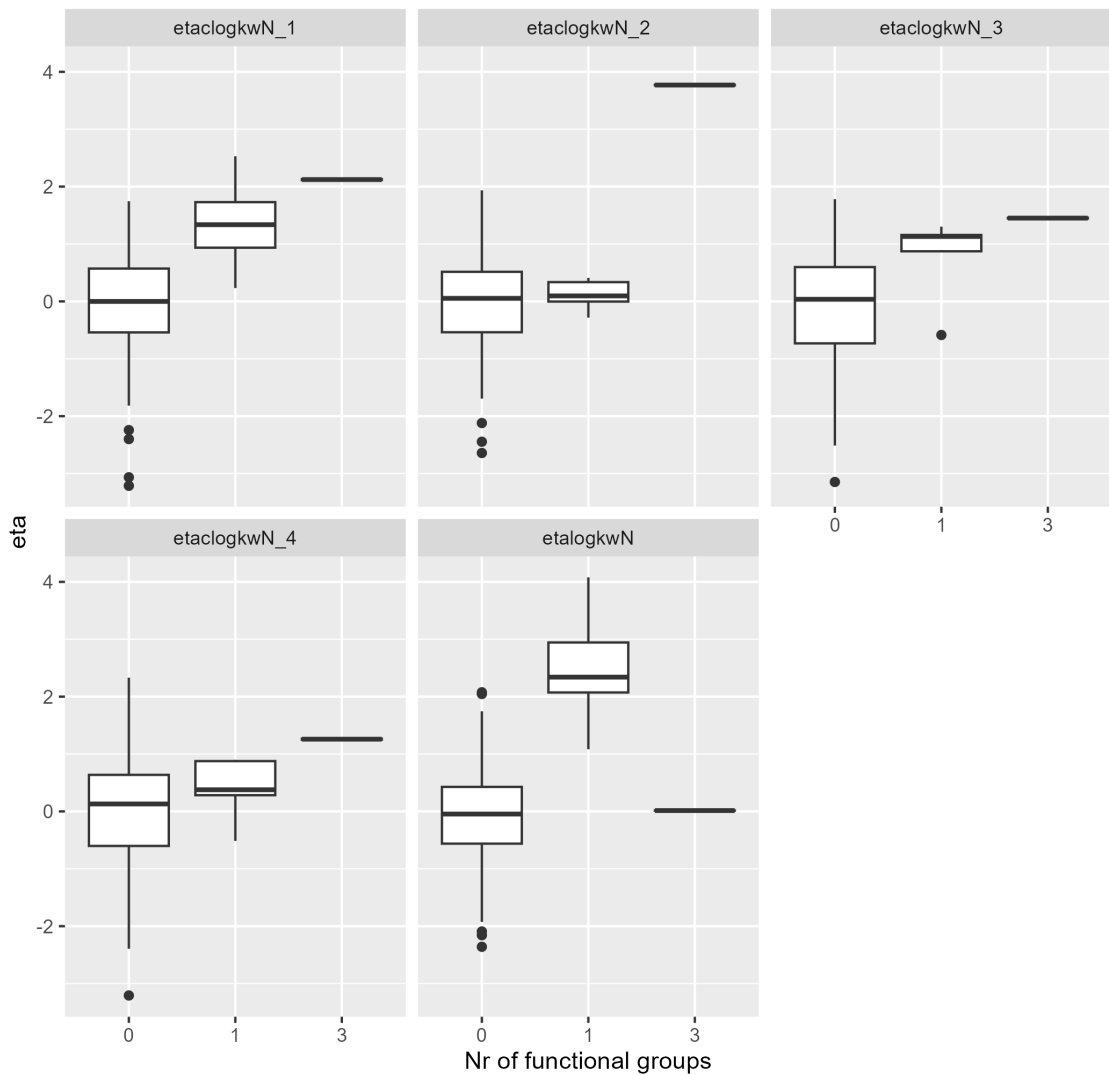
prim. alcohol



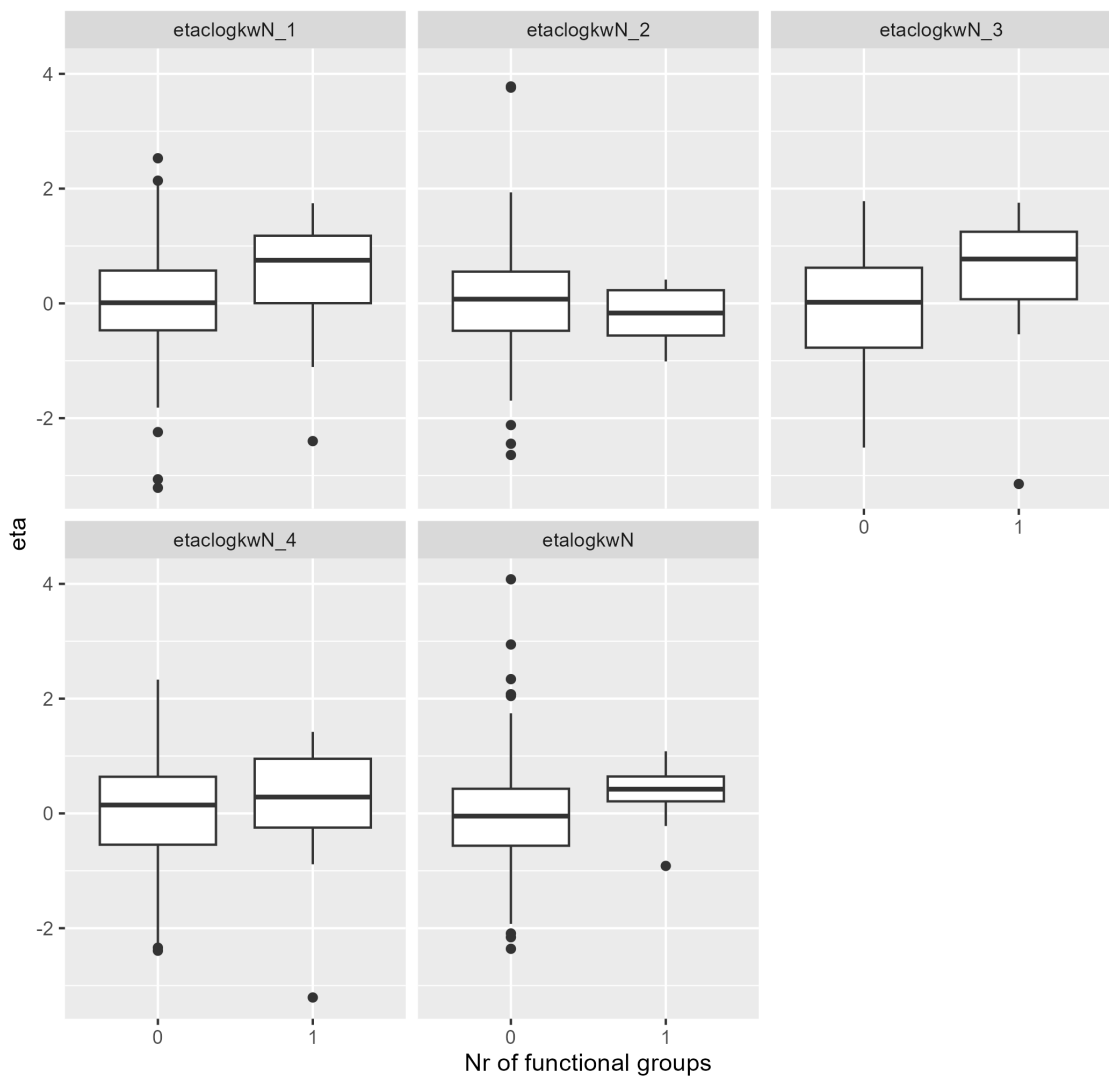
sec. alcohol



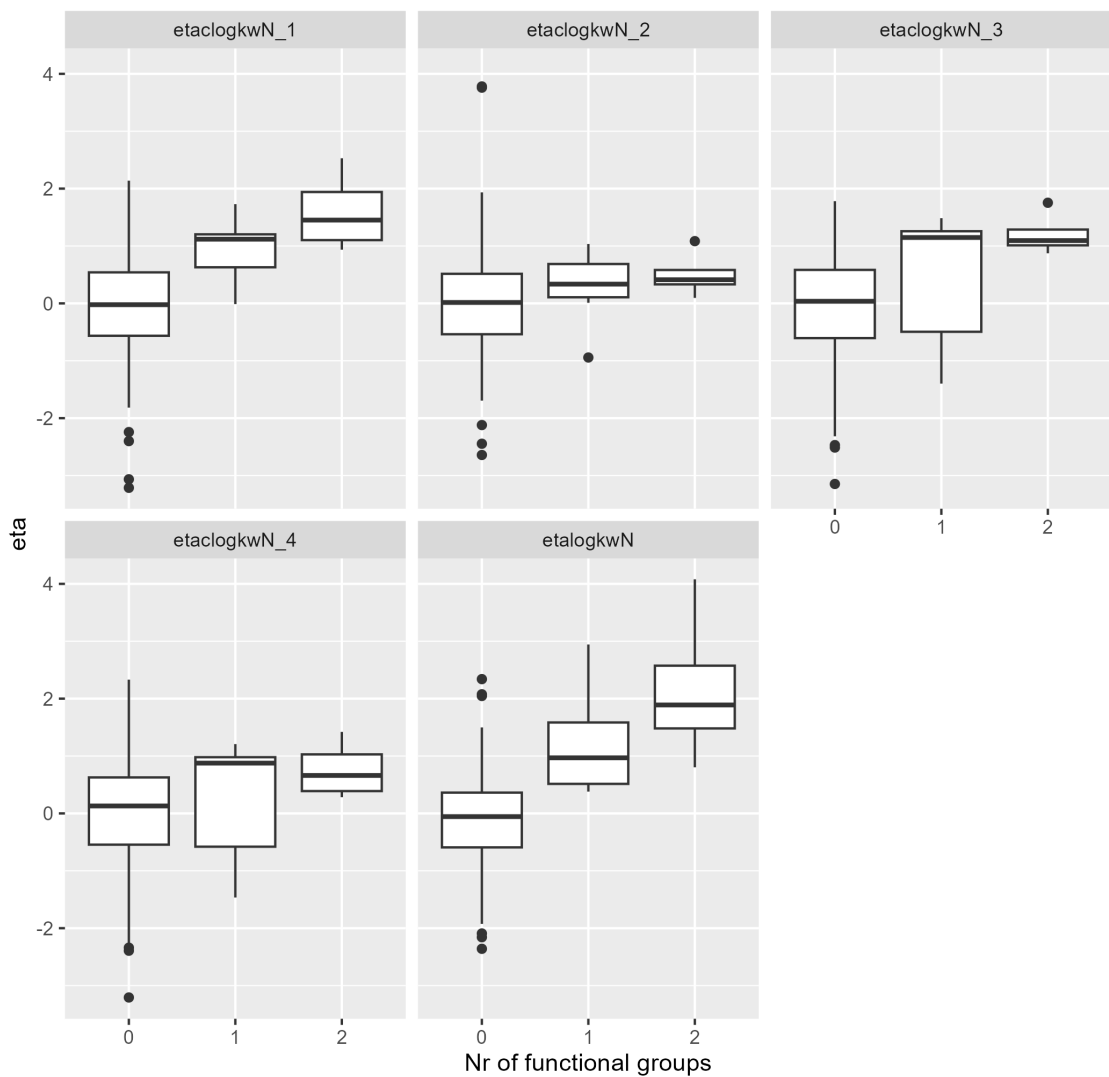
1,2-diol



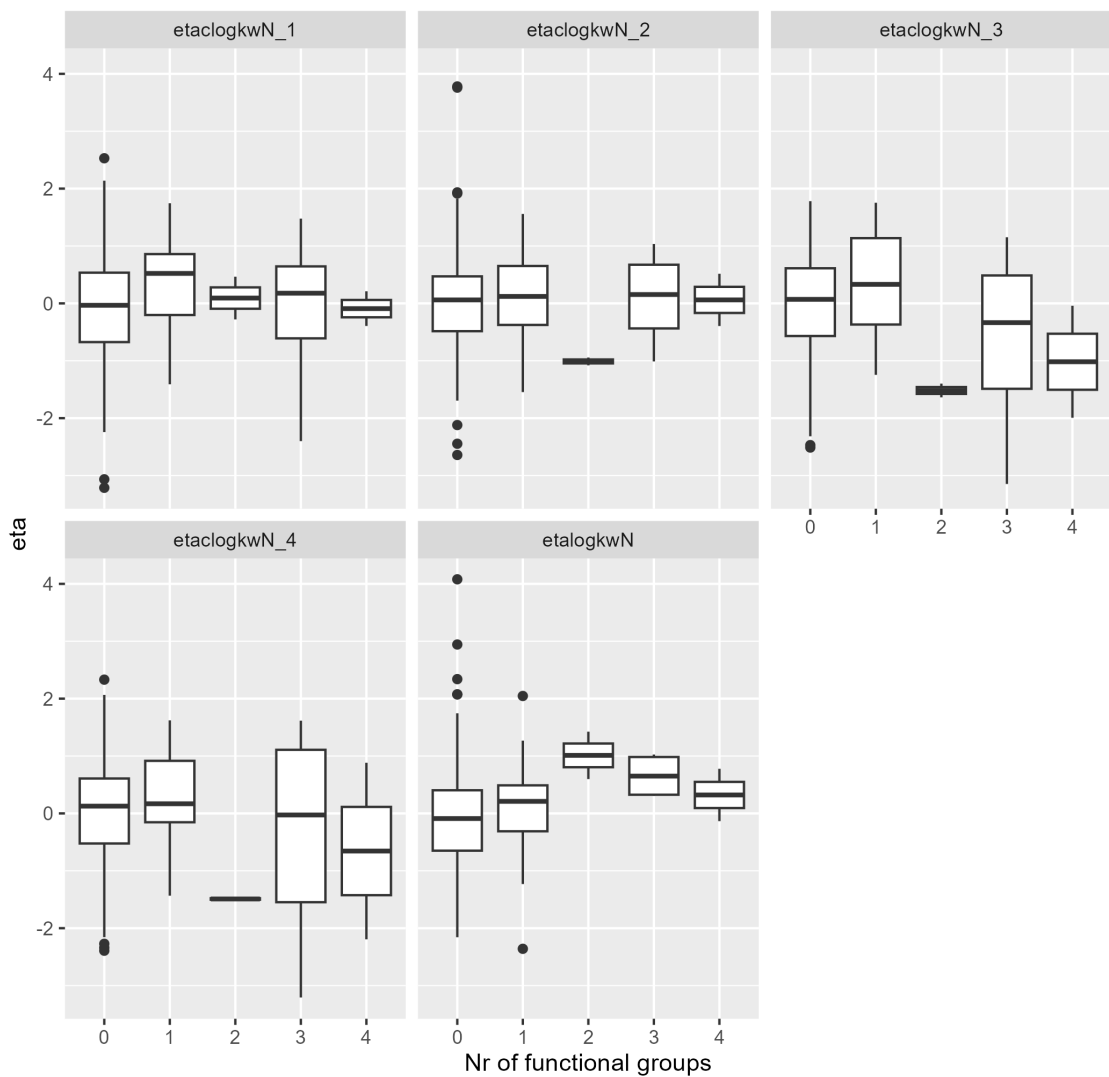
1,2-aminoalcohol



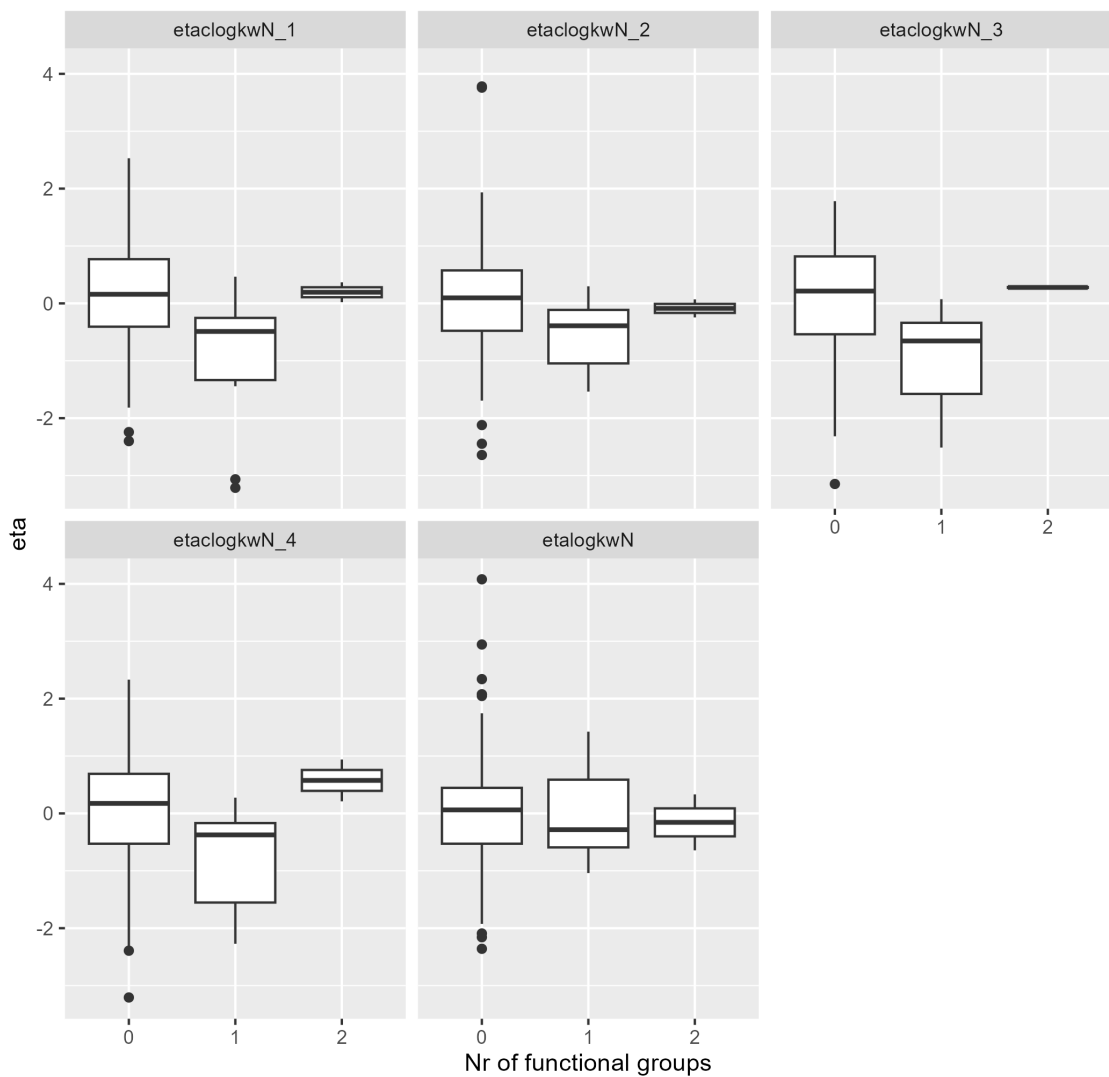
dialkylether



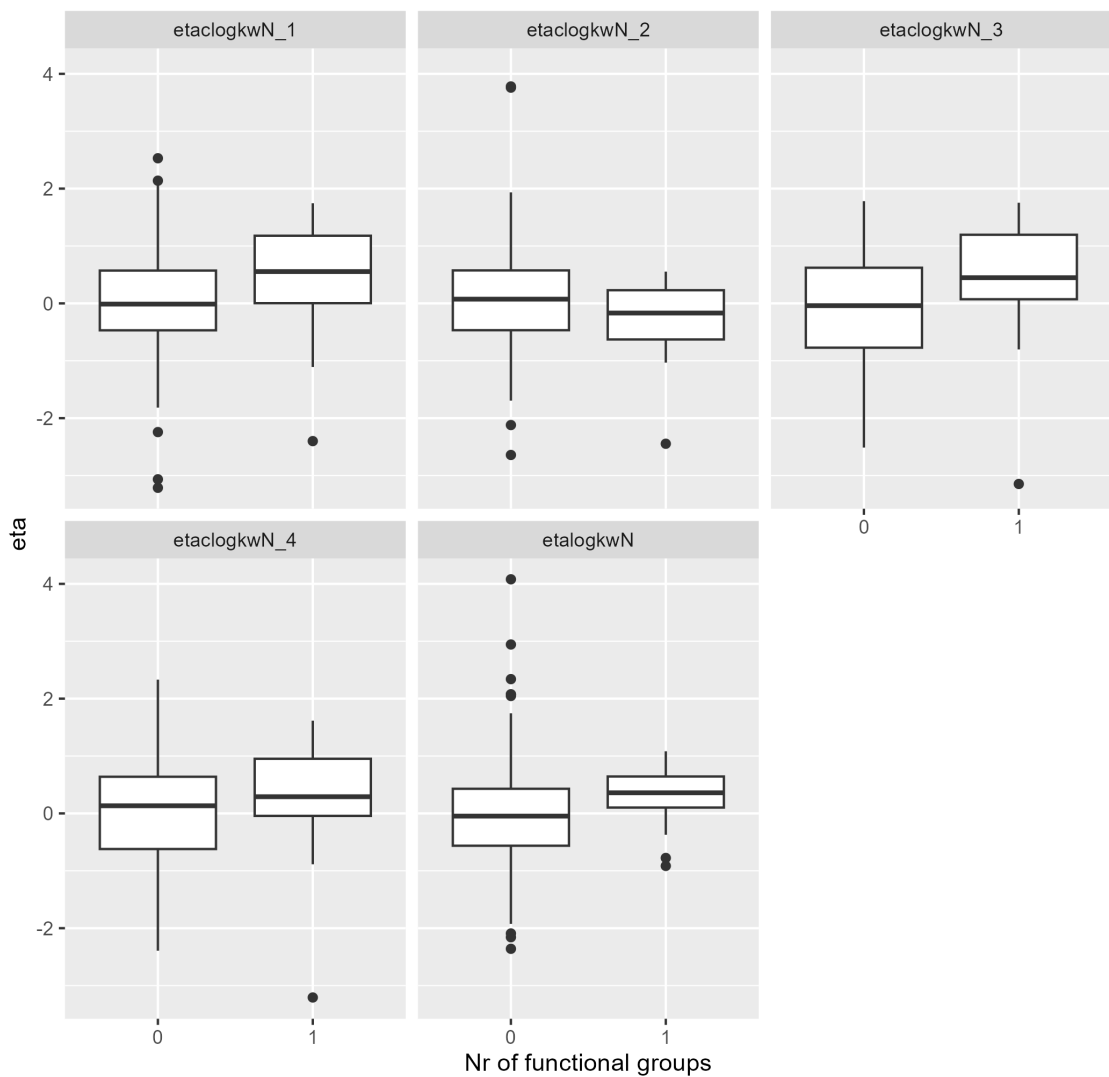
alkylarylether



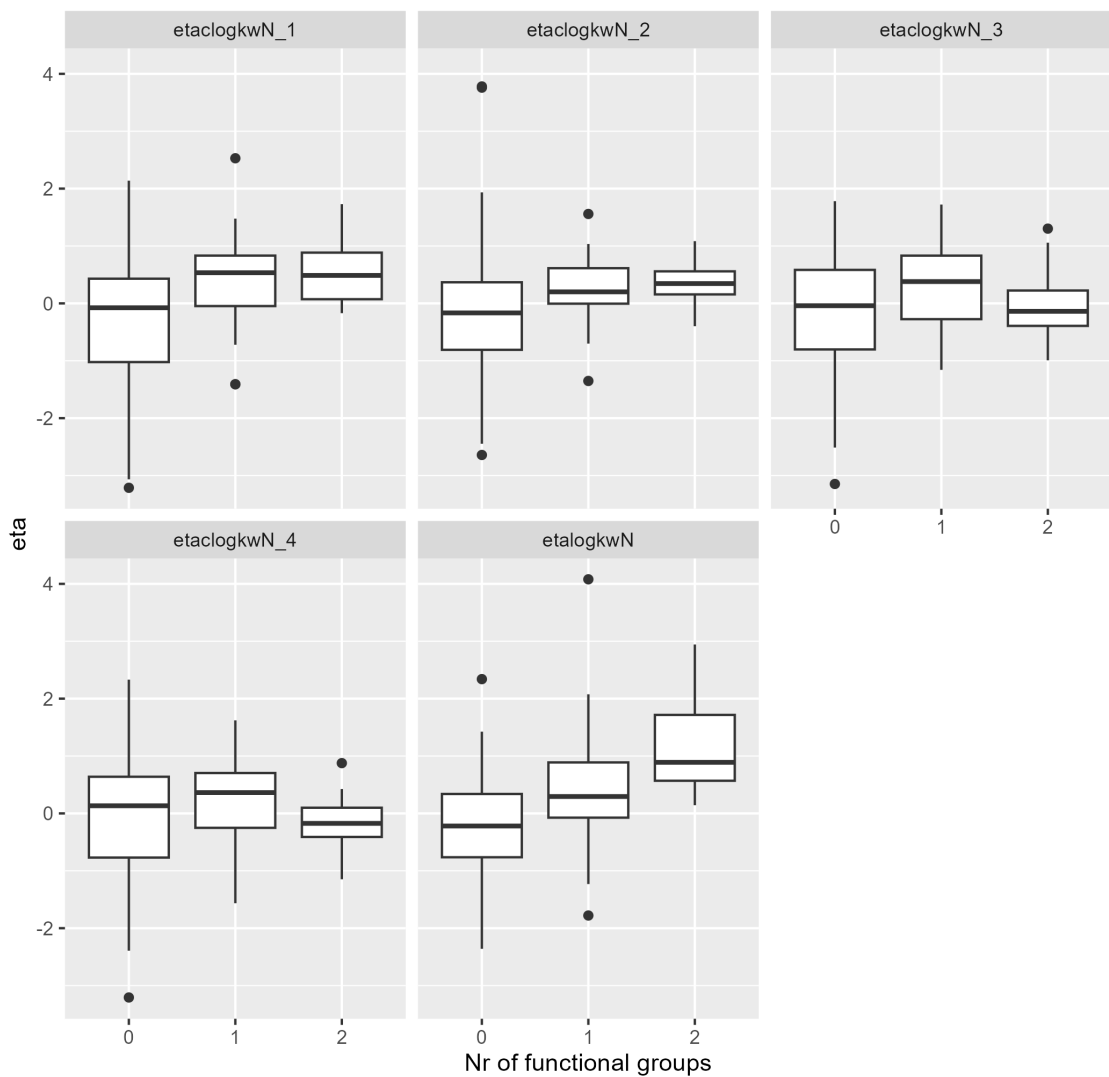
prim. aromat. mine



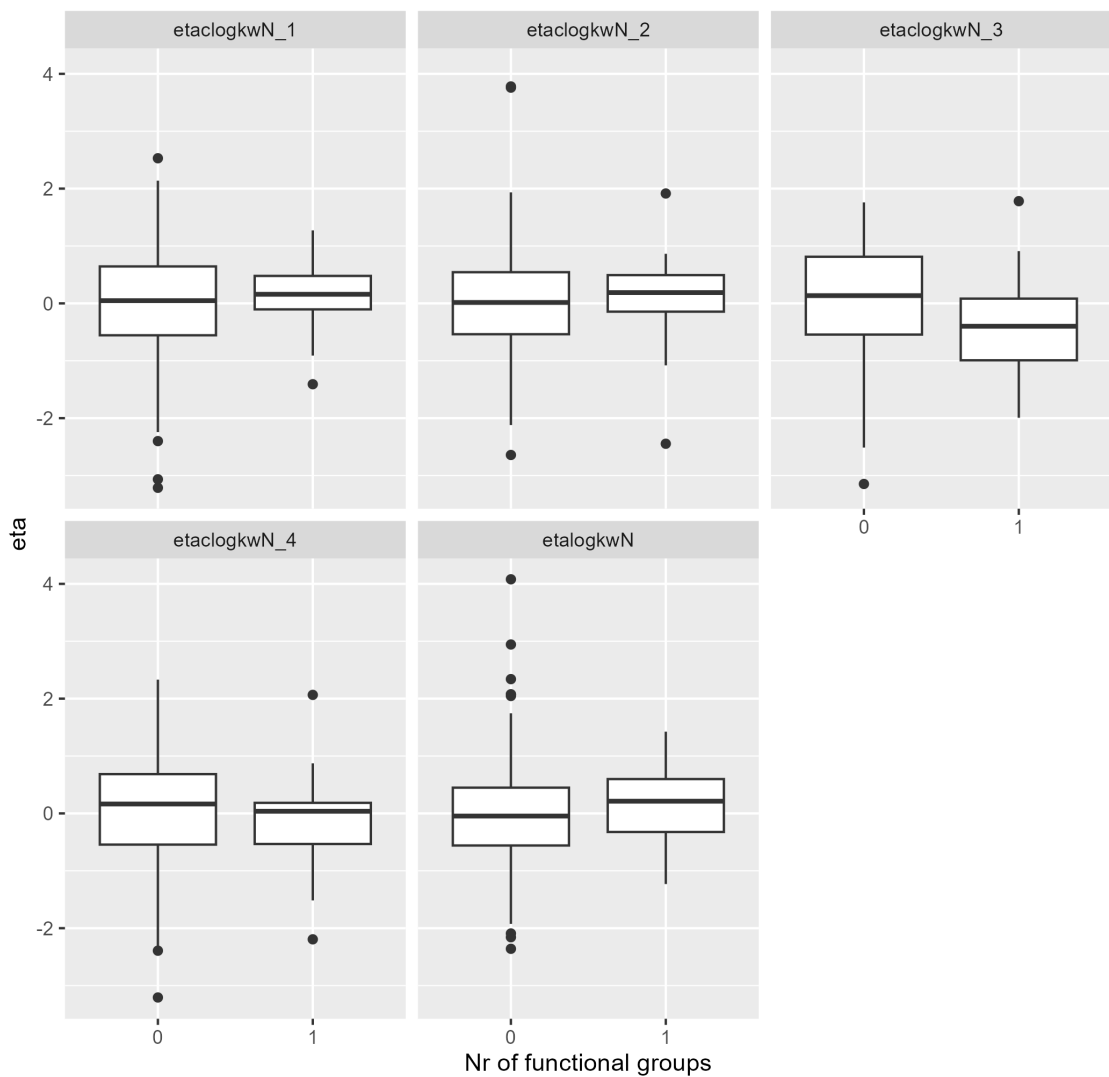
sec. aliphat. amine



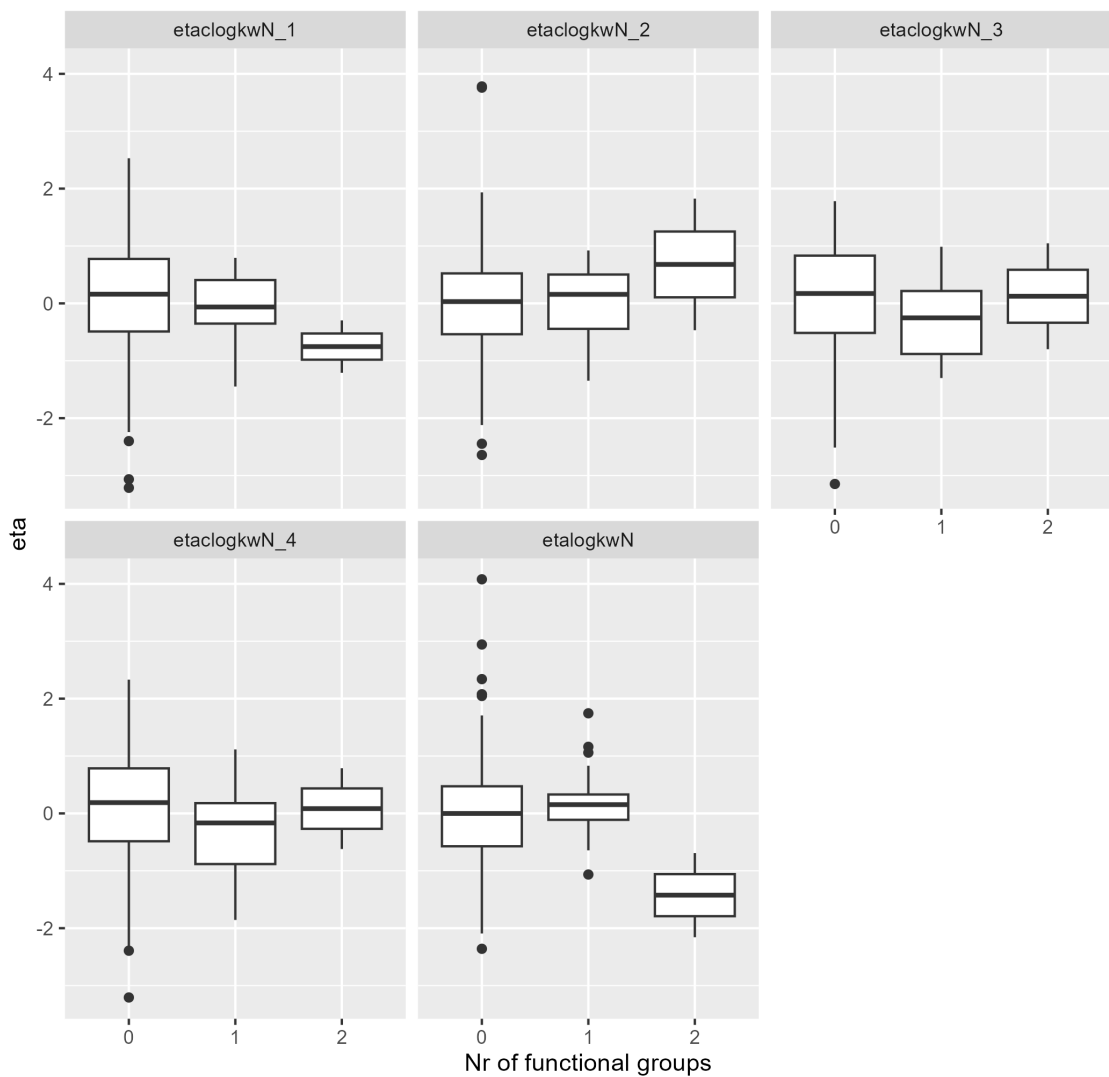
tert. aliphat. amine



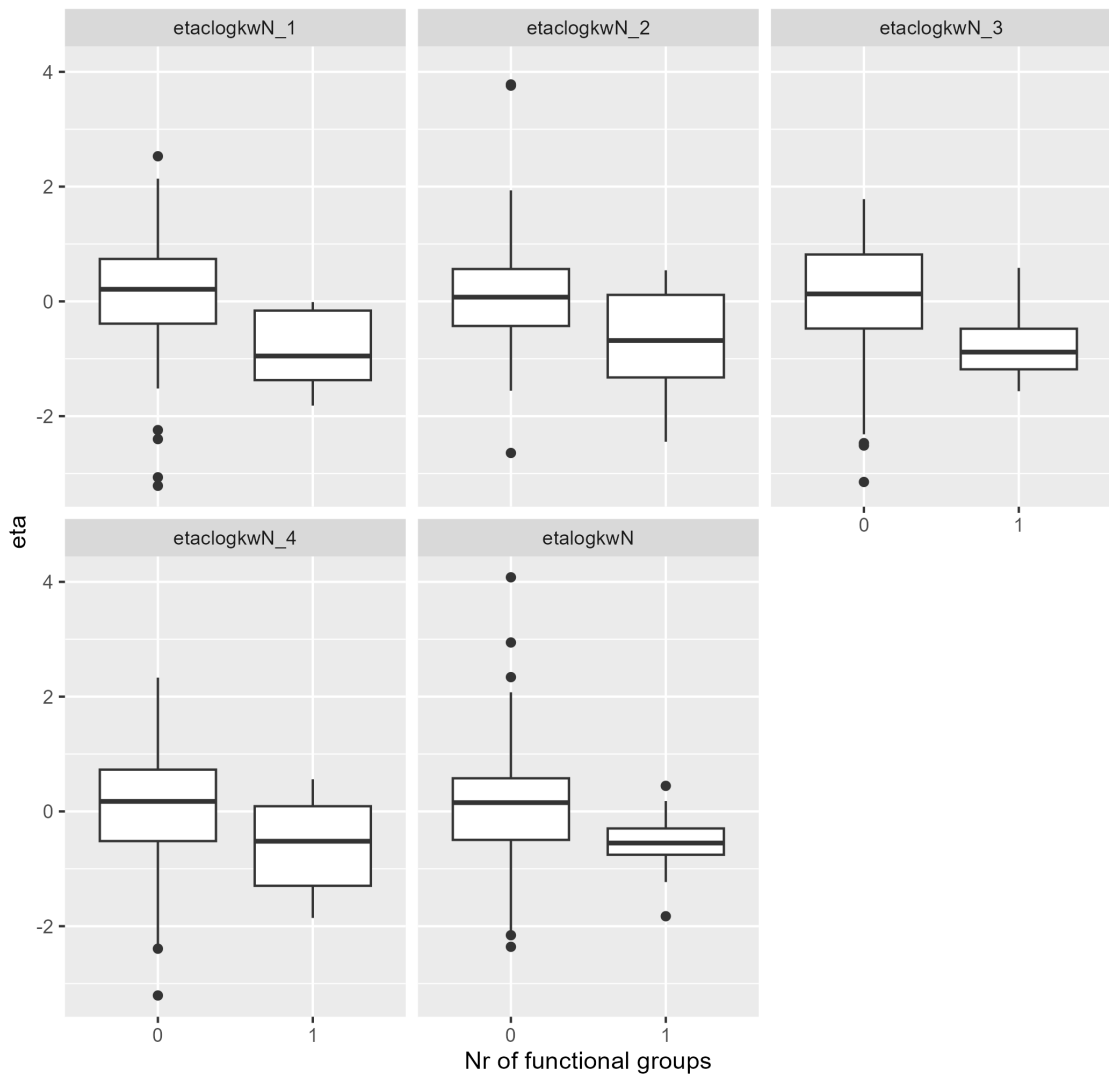
tert. mixed amine



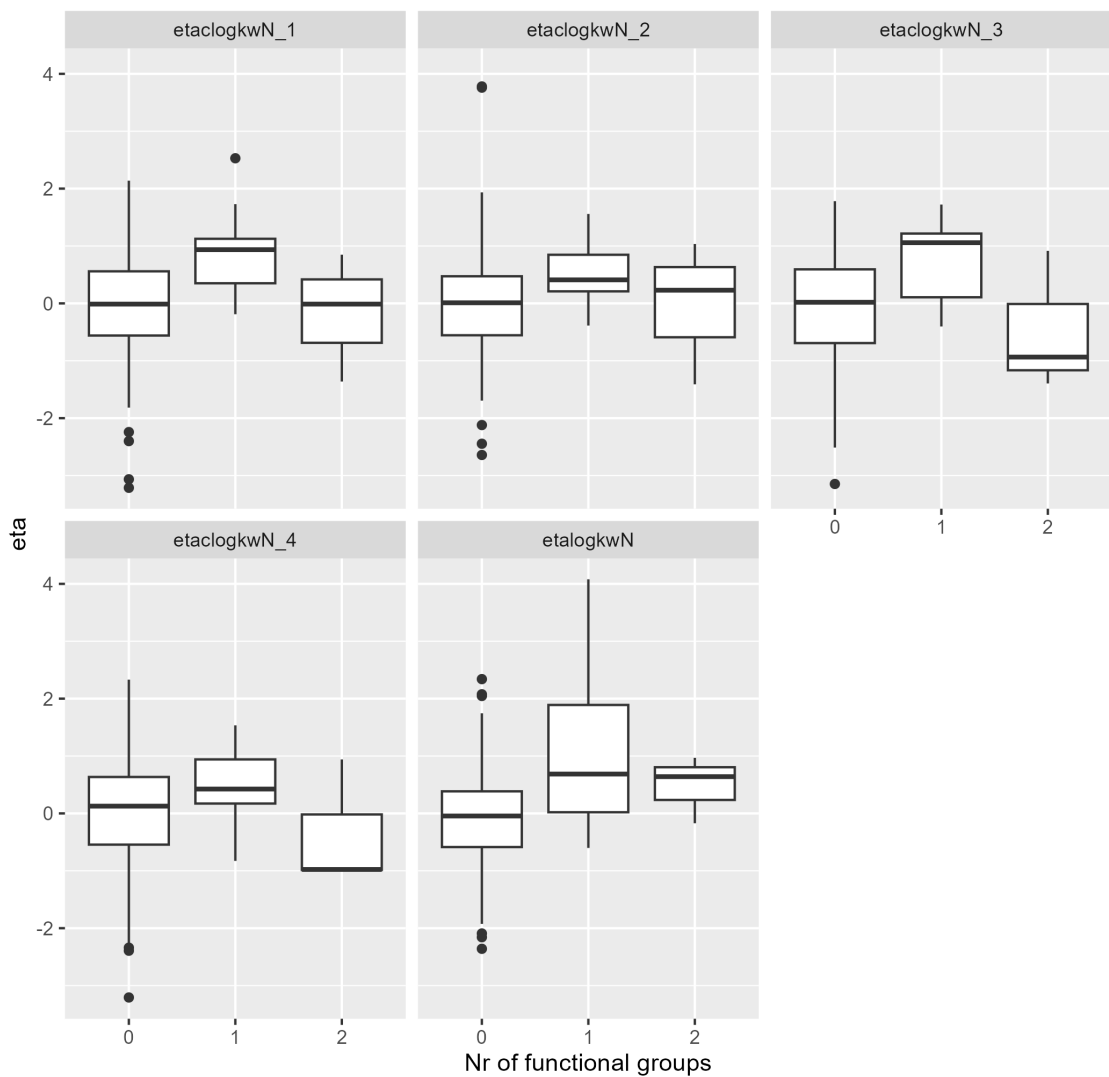
aryl chloride



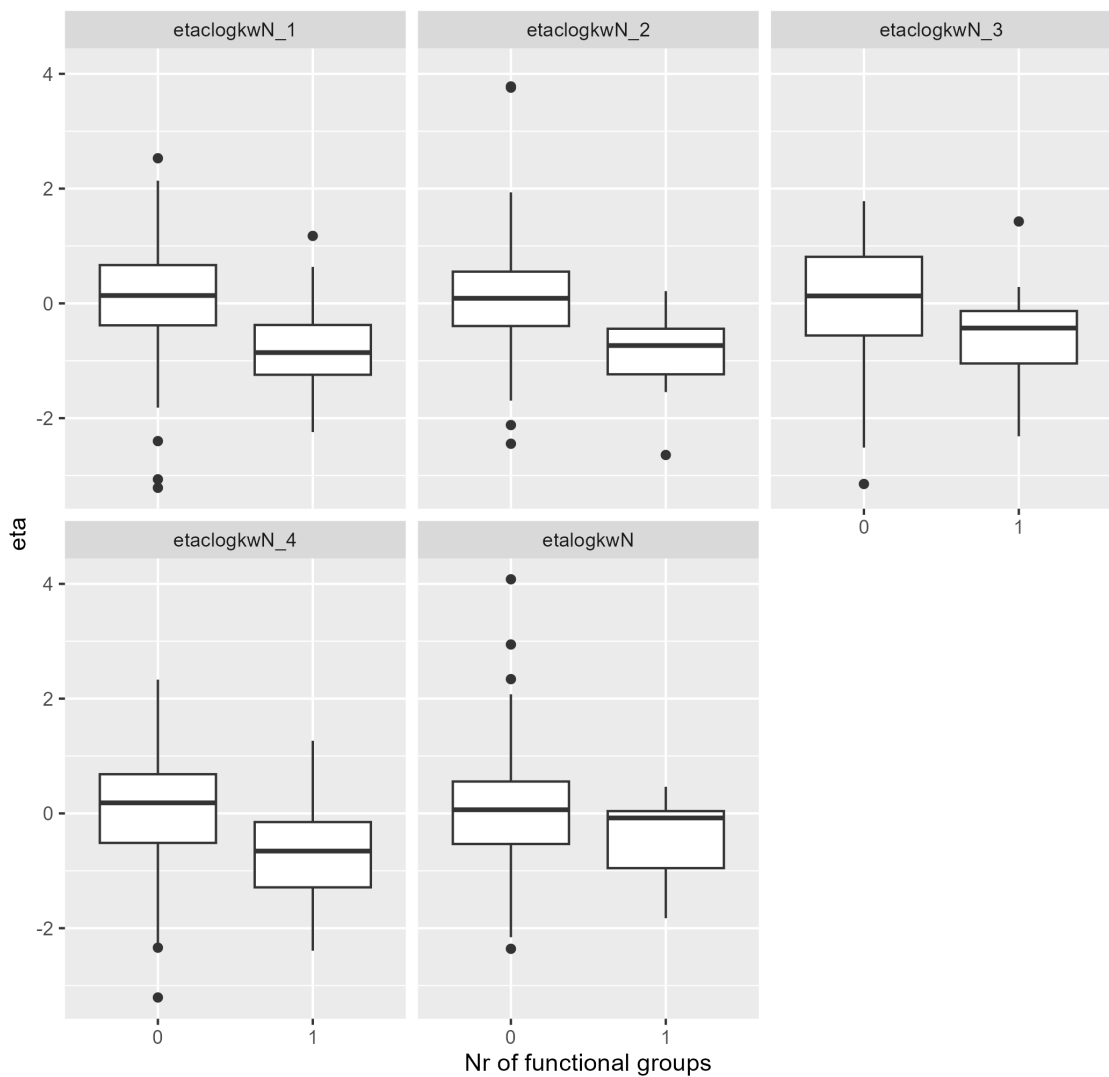
carboxylic acid



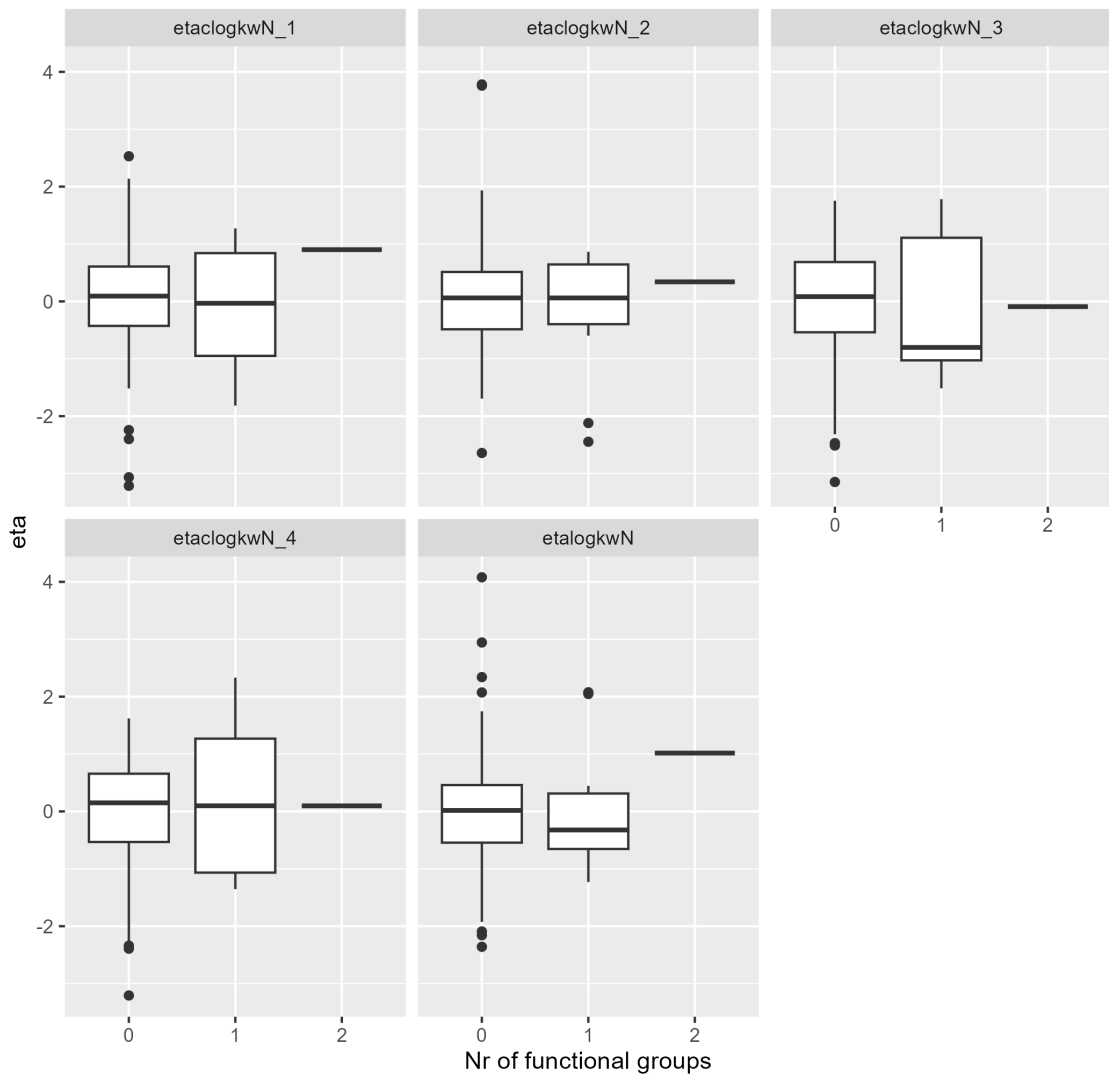
carboxylic acid ester



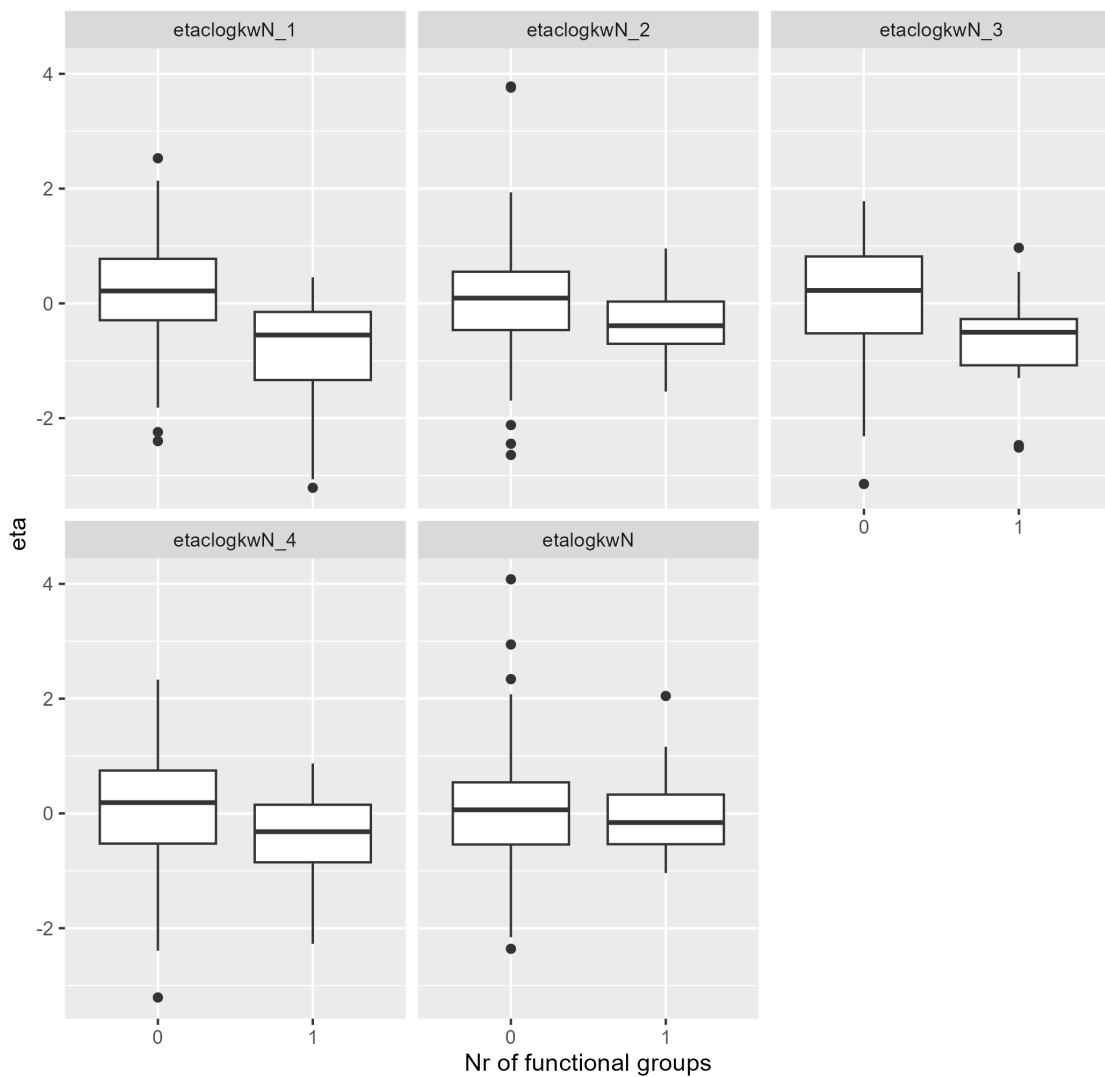
carboxylic acid sec. amide



oxohetarene



sulfonamide



6.6.3 Effect of dissociation

```
paramA <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("etacdlogkwAc", na.rm = TRUE))], 2, mean)
paramA <- melt(paramA)
paramA <- matrix(paramA$value, nrow = datastruct$nGroupsA, byrow = TRUE)
paramA1 <- paramA[, 1]
paramA2 <- paramA[, 2]
paramA3 <- paramA[, 3]
```

```

paramA4 <- paramA[,4]

paramB <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etacdlogkwBc", names(draws_etas_df)))], 1, mean)
paramB <- melt(paramB)
paramB <- matrix(paramB$value, nrow = datastruct$nGroupsB, byrow = TRUE)
paramB1 <- paramB[,1]
paramB2 <- paramB[,2]
paramB3 <- paramB[,3]
paramB4 <- paramB[,4]

param1 <- c(paramA1,paramB1)
param2 <- c(paramA2,paramB2)
param3 <- c(paramA3,paramB3)
param4 <- c(paramA4,paramB4)

GroupType = c(rep("Acids", length(paramA1)),rep("Bases", length(paramB1)))

data_to_plot_diss <- data.frame(param1,param2,param3,param4,GroupType)

p1<-ggpairs(data_to_plot_diss,columns = 1:4, columnLabels = c("etadlogkw1","etadlogkw2","etadlogkw3","etadlogkw4"),
             labeller = "label_parsed",
             legend =1,
             aes(color = GroupType, alpha = 0.5),
             upper = list(continuous = "points"))+
             scale_alpha(guide = "none")

paramA <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etacdS1mAc", names(draws_etas_df)))], 1, mean)
paramA <- melt(paramA)
paramA <- matrix(paramA$value, nrow = datastruct$nGroupsA, byrow = TRUE)
paramA1 <- paramA[,1]
paramA2 <- paramA[,2]
paramA3 <- paramA[,3]
paramA4 <- paramA[,4]

paramB <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etacdS1mBc", names(draws_etas_df)))], 1, mean)
paramB <- melt(paramB)
paramB <- matrix(paramB$value, nrow = datastruct$nGroupsB, byrow = TRUE)
paramB1 <- paramB[,1]
paramB2 <- paramB[,2]

```

```

paramB3 <- paramB[,3]
paramB4 <- paramB[,4]

param1 <- c(paramA1,paramB1)
param2 <- c(paramA2,paramB2)
param3 <- c(paramA3,paramB3)
param4 <- c(paramA4,paramB4)

GroupType = c(rep("Acids", length(paramA1)),rep("Bases", length(paramB1)))

data_to_plot_diss <- data.frame(param1,param2,param3,param4,GroupType)

p2<-ggpairs(data_to_plot_diss,columns = 1:4, columnLabels = c("etacdS1m1","etacdS1m2","eta
    labeller = "label_parsed",
    legend =1,
    aes(color = GroupType, alpha = 0.5),
    upper = list(continuous = "points"))+
    scale_alpha(guide = "none")

paramA <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etacdS1Ac", names
paramA <- melt(paramA)
paramA <- matrix(paramA$value,nrow = datastruct$nGroupsA, byrow = TRUE)
paramA1 <- paramA[,1]
paramA2 <- paramA[,2]
paramA3 <- paramA[,3]
paramA4 <- paramA[,4]

paramB <- apply(draws_etas_df[,which(colnames(draws_etas_df) %in% grep("^etacdS1Bc", names
paramB <- melt(paramB)
paramB <- matrix(paramB$value,nrow = datastruct$nGroupsB, byrow = TRUE)
paramB1 <- paramB[,1]
paramB2 <- paramB[,2]
paramB3 <- paramB[,3]
paramB4 <- paramB[,4]

param1 <- c(paramA1,paramB1)
param2 <- c(paramA2,paramB2)
param3 <- c(paramA3,paramB3)
param4 <- c(paramA4,paramB4)

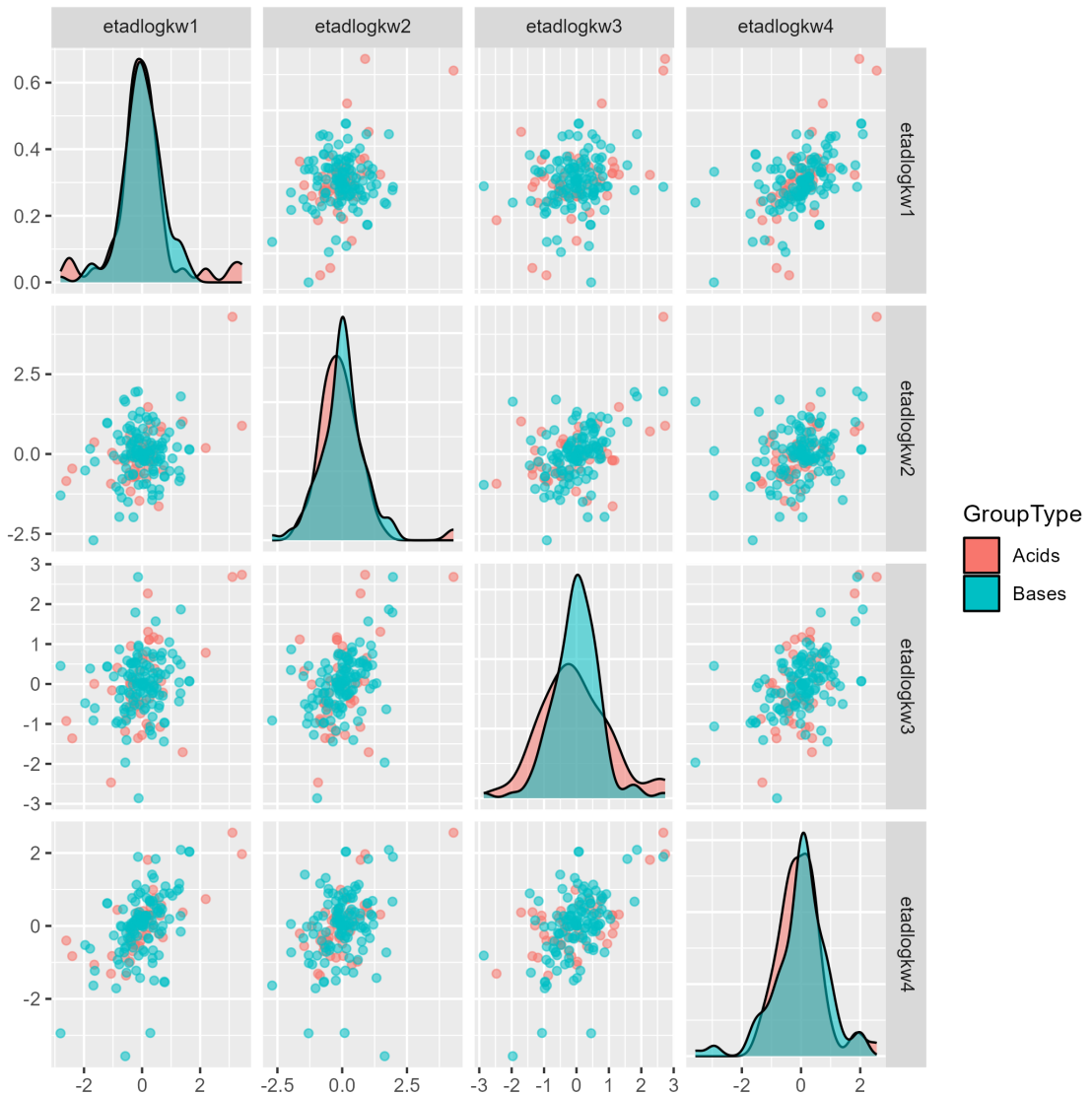
```

```
GroupType = c(rep("Acids", length(paramA1)),rep("Bases", length(paramB1)))

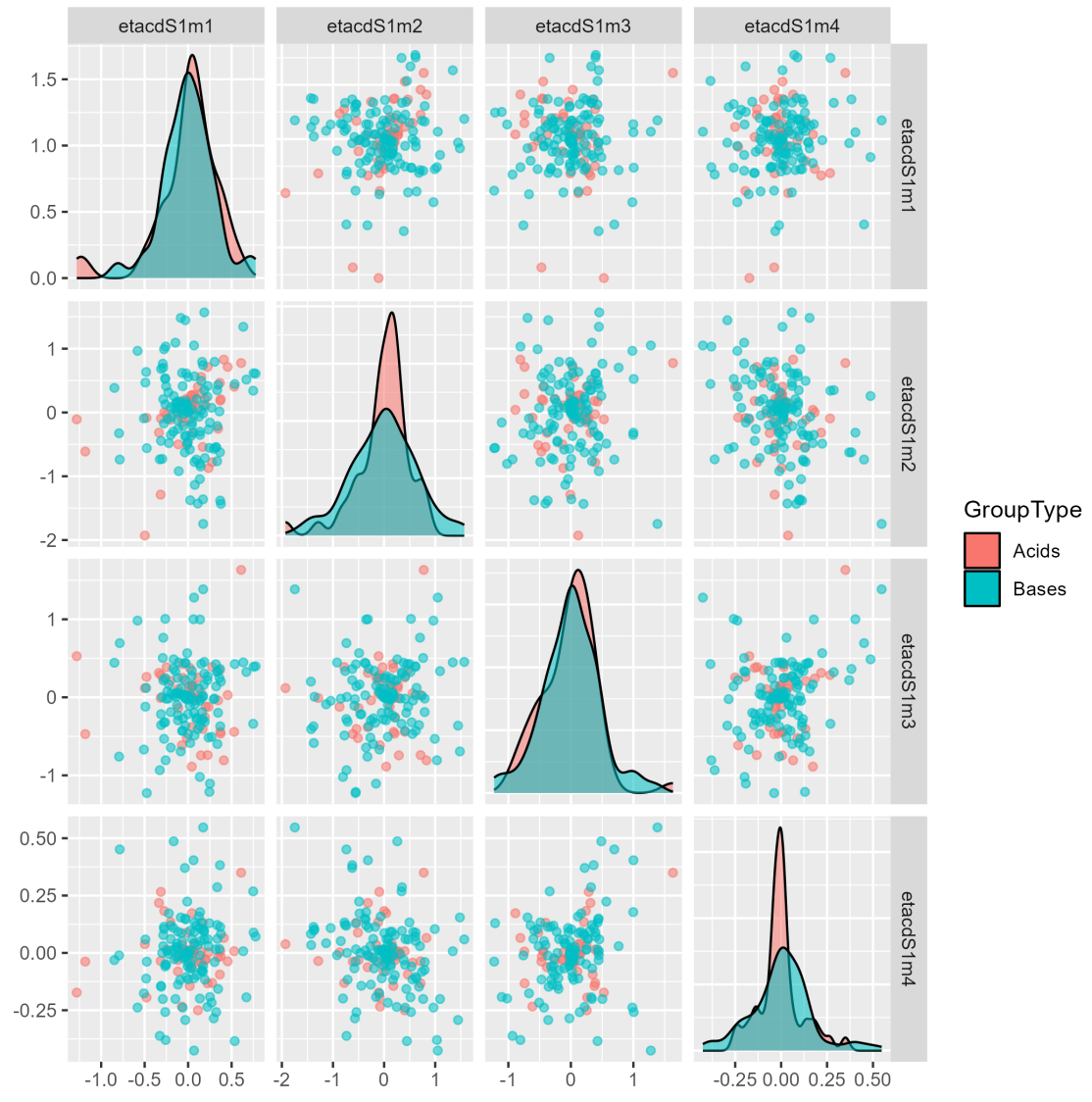
data_to_plot_diss <- data.frame(param1,param2,param3,param4,GroupType)

p3<-ggpairs(data_to_plot_diss,columns = 1:4, columnLabels = c("etacdS11","etacdS12","etacd
    labeller = "label_parsed",
    legend =1,
    aes(color = GroupType, alpha = 0.5),
    upper = list(continuous = "points"))+
    scale_alpha(guide = "none")

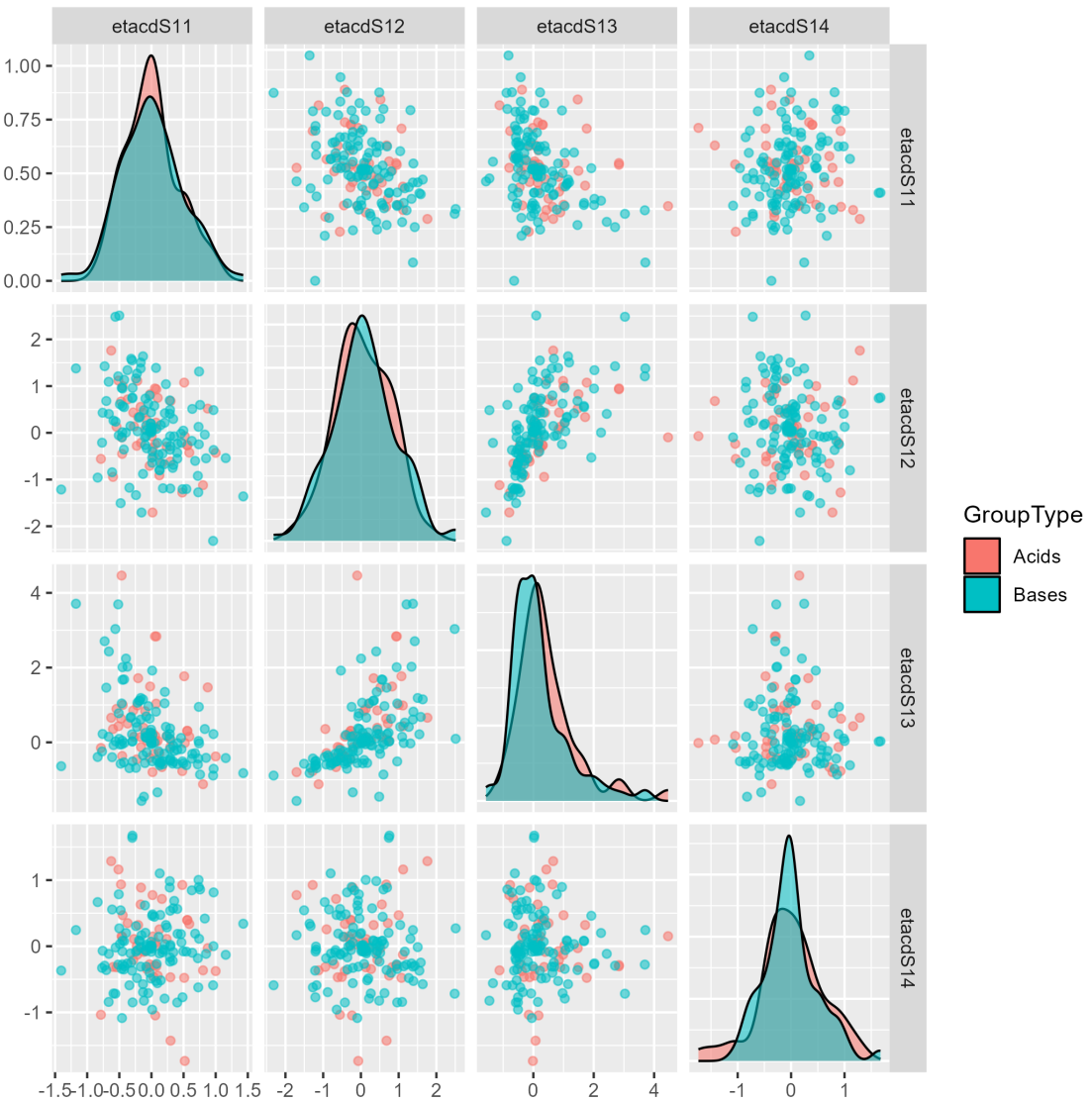
print(p1)
```



```
print(p2)
```



```
print(p3)
```



```

ggsave(paste0("figures\\etaplots\\", "dissociatedforms_1", ".png"), plot=p1, width = 20, height = 10)
ggsave(paste0("figures\\etaplots\\", "dissociatedforms_2", ".png"), plot=p2, width = 20, height = 10)
ggsave(paste0("figures\\etaplots\\", "dissociatedforms_3", ".png"), plot=p3, width = 20, height = 10)
  
```

7 Comparison of observed and model predicted retention times

The individual and population model predictions were simulated using a more dense design. The individual predictions are based on population-level parameters, predictors, and all of the

observed retention time measurements. The population predictions are based on population-level parameters and predictors.

```

fitsim <- cmdstanr::as_cmdstan_fit(c(
  #  'stanfiles/output_1.csv',
  #  'stanfiles/output_2.csv',
  #  'stanfiles/output_3.csv',
  #  'stanfiles/output_4.csv',
  #  'stanfiles/output_5.csv',
  #  'stanfiles/output_6.csv',
  #  'stanfiles/output_7.csv',
  'stanfiles/output_8.csv'
))

design <- read.csv('data/hplcparam_design.csv')
design$Mod  = as.character(design$Mod)
design$Mod2 = ifelse(design$Mod=="MeOH",1,2) # MeOH = 1, ACN = 2
design$expid = match(design$expid, unique(design$expid))
nAnalytes <- datastruct$nAnalytes
nColumns <- datastruct$nColumns
nExp <- nrow(design);
datasim <- design %>%
  slice(rep(row_number(), nAnalytes*nColumns)) %>%
  mutate(Column = rep(c(1,2,3,4,5), each = nExp*nAnalytes)) %>%
  mutate(METID = rep(rep(unique(data$METID), each=nExp), nColumns))
datasim$expid = datasim$expid*(2-datasim$Column) + (length(unique(datasim$expid))+datasim$expid)
datasim$ColumnName = ifelse(datasim$Column==1,"XBridge Shield RP18",
                            ifelse(datasim$Column==2,"XTerra MS C18",
                            ifelse(datasim$Column==3,"XBridge Phenyl",
                            ifelse(datasim$Column==4,"XBridge C8", "Xterra M

datasim$to[datasim$Column==1] = 0.532
datasim$to[datasim$Column==2] = 0.542
datasim$to[datasim$Column==3] = 0.542
datasim$to[datasim$Column==4] = 0.552
datasim$to[datasim$Column==5] = 0.566

datastructsim <-datastruct
datastructsim$nObs=length(datasim$METID)
datastructsim$analyte=match(datasim$METID, unique(datasim$METID))
datastructsim$modifier=match(datasim$Mod2, sort(unique(datasim$Mod2)))

```

```

datastructsim$steps=4*(2-datasim$Mod2) + 10*(datasim$Mod2-1)
datastructsim$column=match(datasim$Column, unique(datasim$Column))
datastructsim$hplcparam=cbind(datasim$tg,datasim$td,datasim$to,datasim$te,
                               datasim$fio,datasim$fik,datasim$Mod2-1,datasim$pHo,
                               datasim$alpha1,datasim$alpha2,(datasim$Temp-25)/10,
                               datasim$Column-1)
datastructsim$trobs = rep(0,datastructsim$nObs)

model_sim <- cmdstan_model("stan/hplc-gra-fivecolumns-sim.stan")

fit_sim <- model_sim$generate_quantities(fitsim,
                                           data = datastructsim,
                                           seed = 123,
                                           parallel_chains = 4,
                                           output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-sim-202308082137-1-2fe36
  '))

draws_sim_df <- as_draws_df(x$generated_quantities)

# draws_sim_df <- fit_sim$draws(format = "df")

tr_sim_Cond <- apply(draws_sim_df[,which(colnames(draws_sim_df) %in% grep("trCond", names
  draws_sim_df))], 1, mean)

tr_sim_Pred <- apply(draws_sim_df[,which(colnames(draws_sim_df) %in% grep("trPred", names
  draws_sim_df))], 1, mean)

tr_sim_Cond<-as.data.frame(t(tr_sim_Cond))

datasim$trCond_l=tr_sim_Cond`5%` 

datasim$trCond_m=tr_sim_Cond`50%` 

datasim$trCond_h=tr_sim_Cond`95%` 

tr_sim_Pred<-as.data.frame(t(tr_sim_Pred))

datasim$trPred_l=tr_sim_Pred`5%` 

```

```

datasim$trPred_m=tr_sim_Pred`50%
datasim$trPred_h=tr_sim_Pred`95%

```

7.1 Individual predictions:

```

analyte_ID_sample <- c(9,17,33,58,140,180)

#analyte_ID_sample <-unique(data$METID)

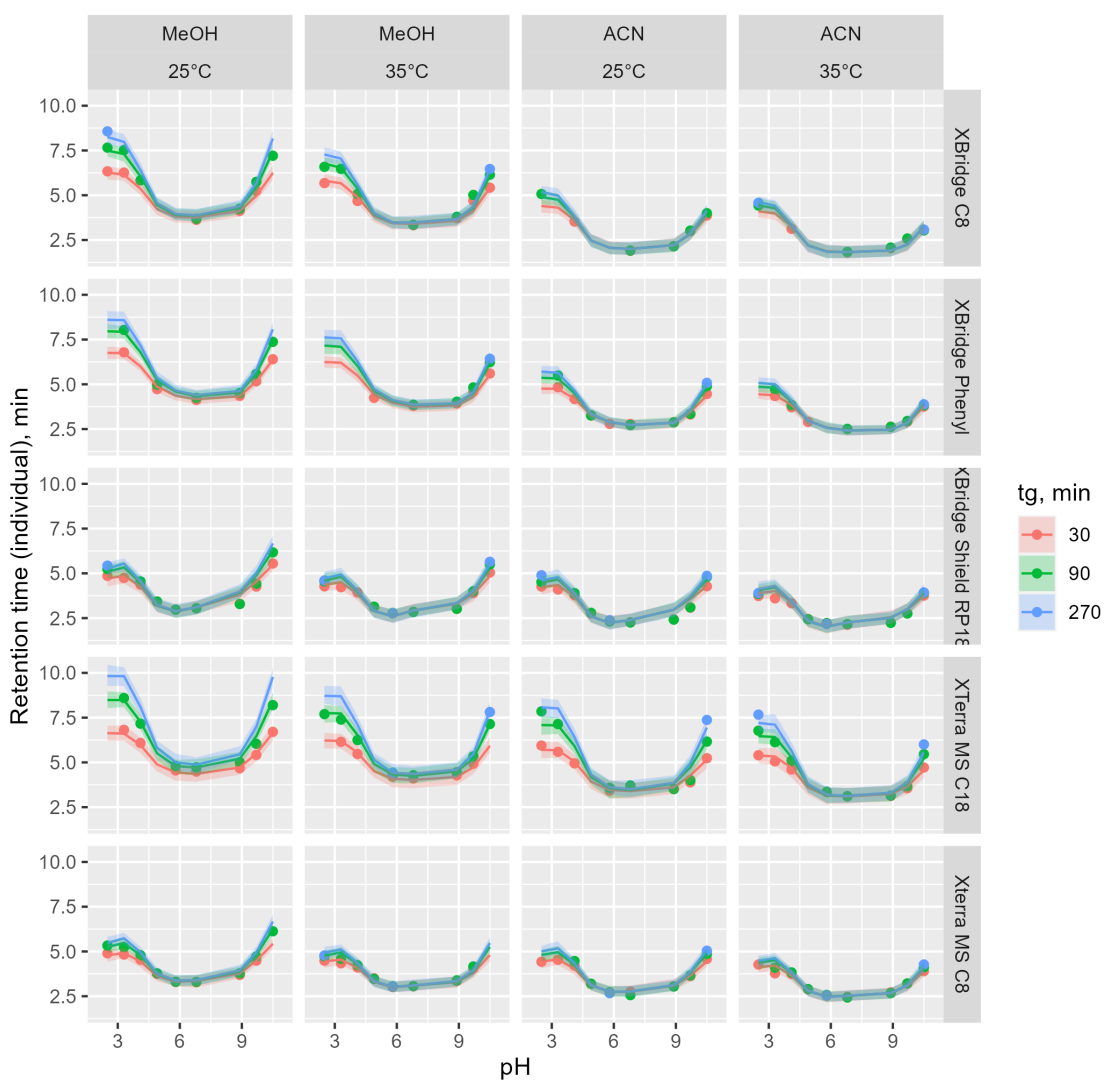
for(i in 1:length(analyte_ID_sample)){

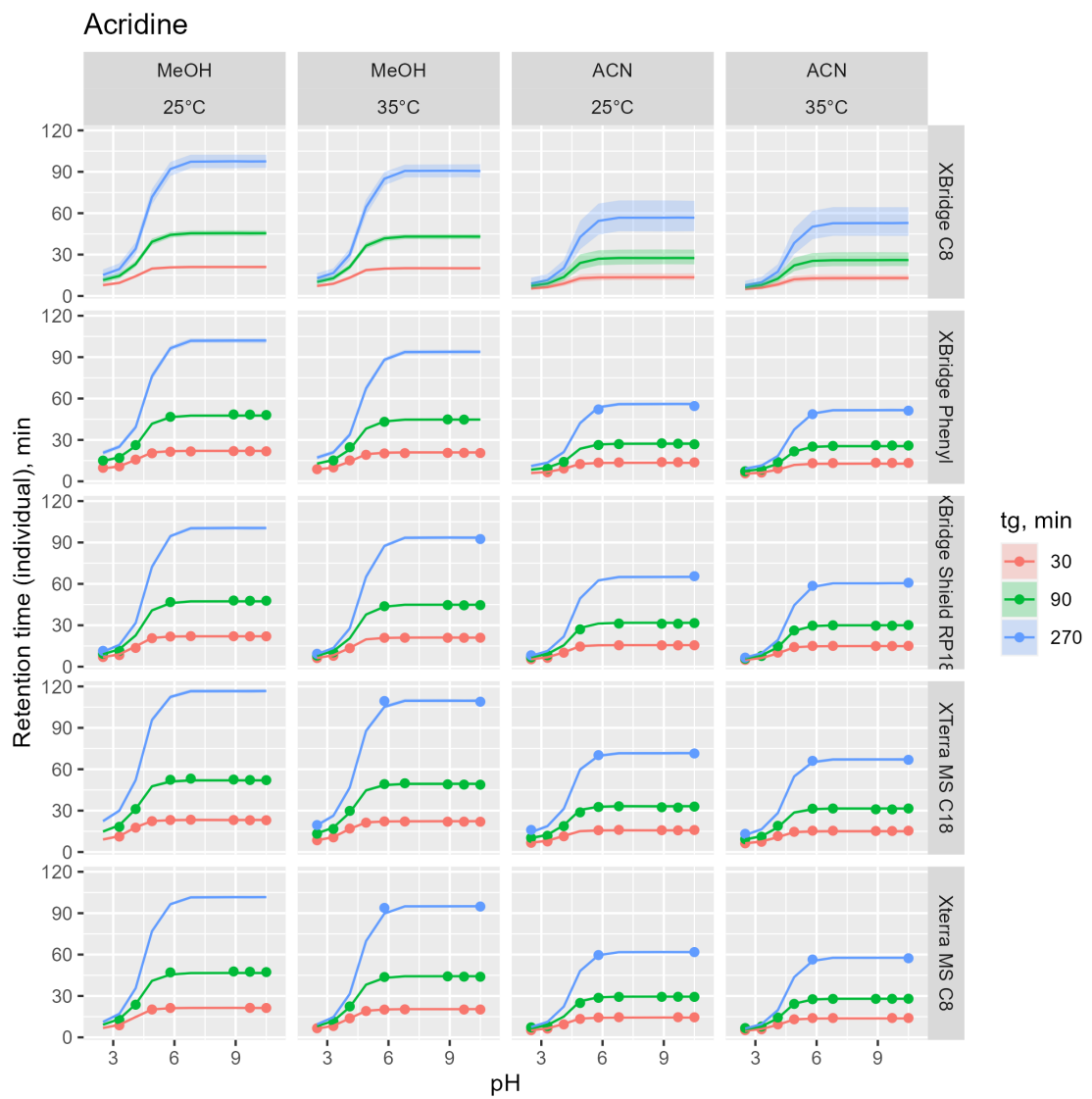
  p <- ggplot()+
    geom_point(data=data[which(data$METID %in% analyte_ID_sample[i]),],
               aes(x = pHs, y = RT, color = as.factor(tg)))+
    geom_line(data=datasim[which(datasim$METID %in% analyte_ID_sample[i]),],
              aes(x = pHs, y = trCond_m, color = as.factor(tg)))+
    geom_ribbon(data=datasim[which(datasim$METID %in% analyte_ID_sample[i]),],
                aes(x = pHs, ymin = trCond_l, ymax = trCond_h, fill = as.factor(tg)), alpha=0.5)+ 
    xlim(2,11)+ 
    facet_grid(ColumnNames~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+ 
    labs(title=paste(dataNames>Name[analyte_ID_sample[i]]),
         x ="pH",
         y = "Retention time (individual), min",
         color = "tg, min",
         fill = "tg, min")

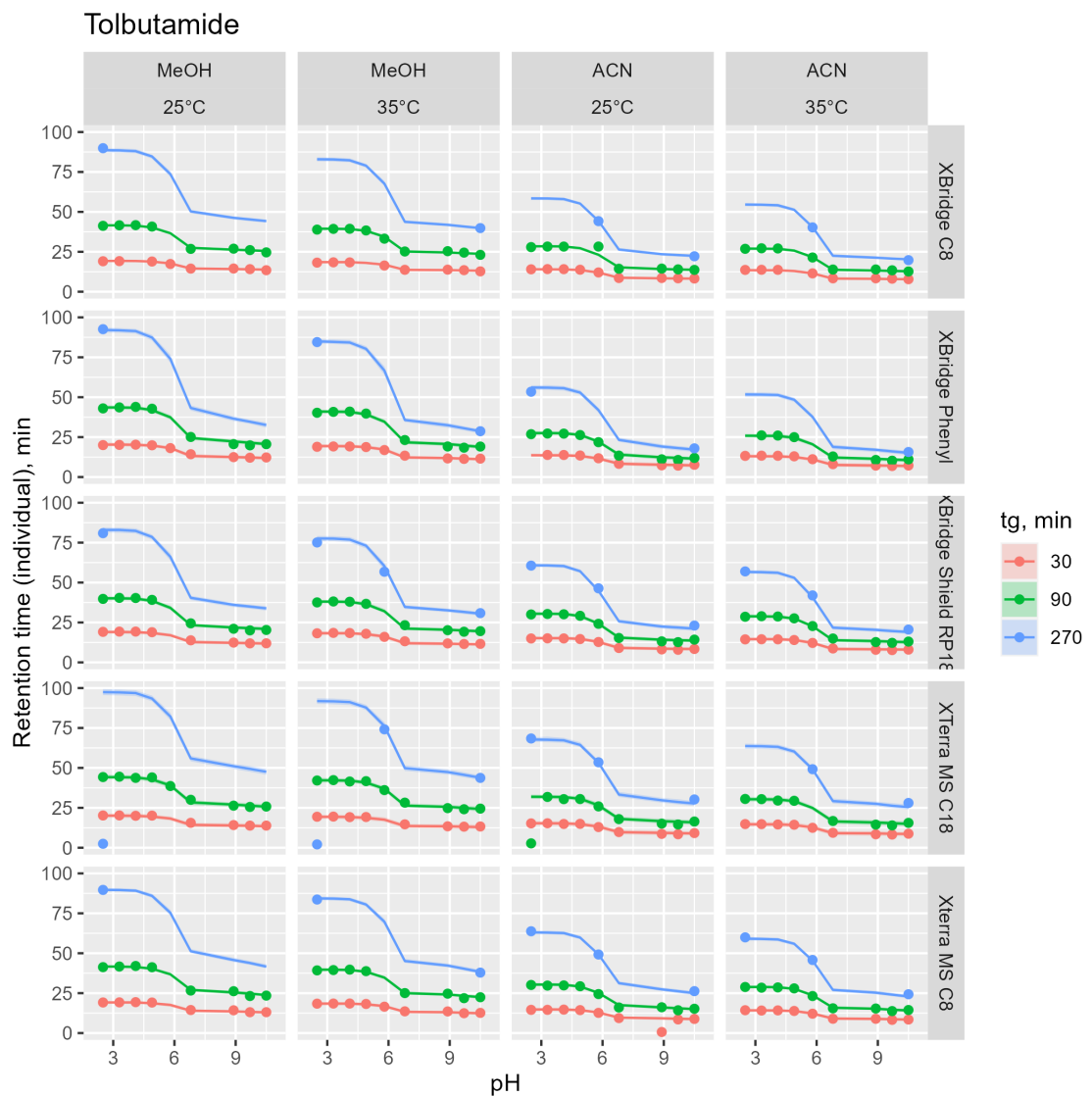
  print(p)

  ggsave(paste0("figures\\concordanceplots\\", paste(dataNames>Name[analyte_ID_sample[i]]),
}
```

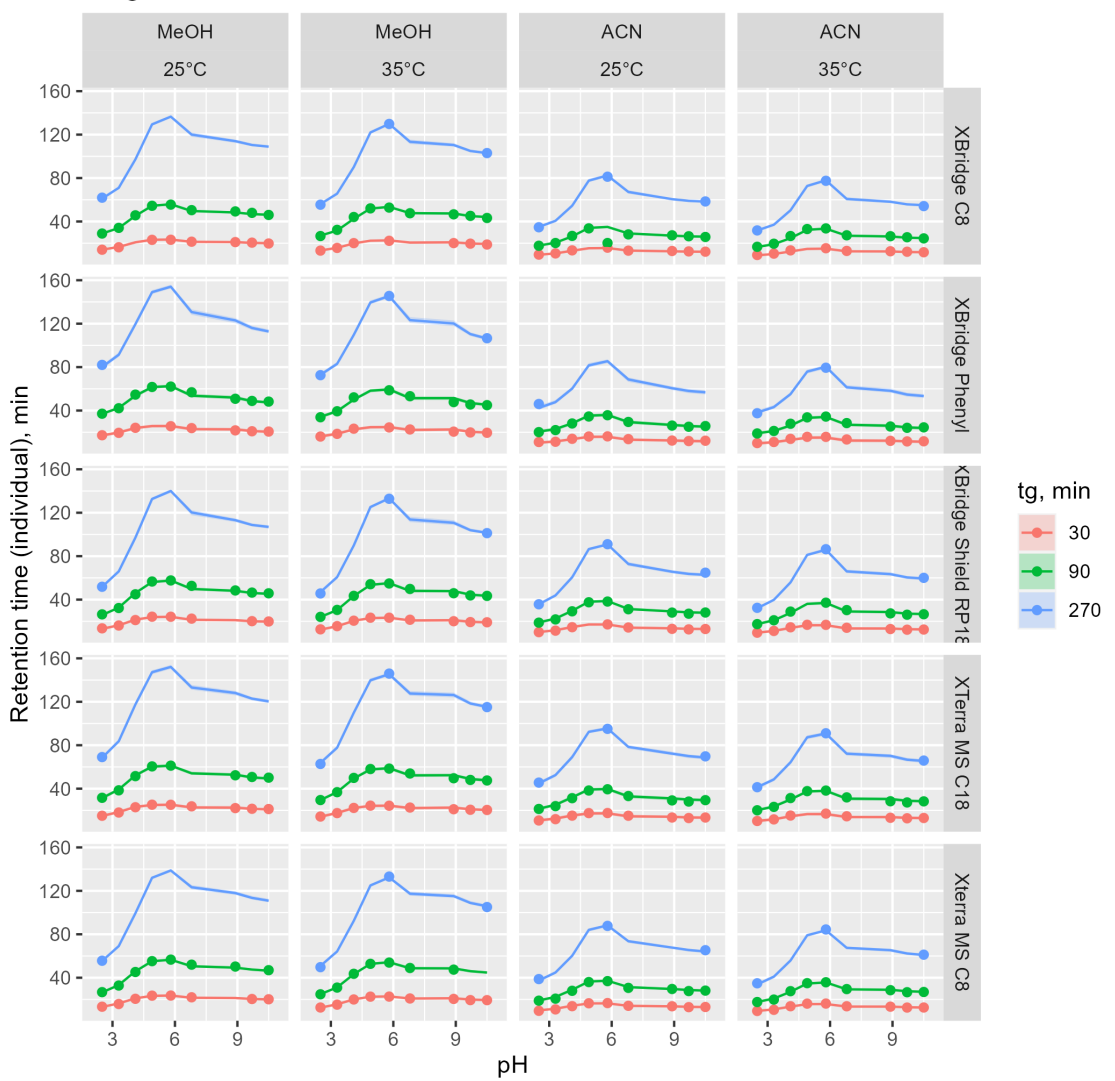
Baclofen



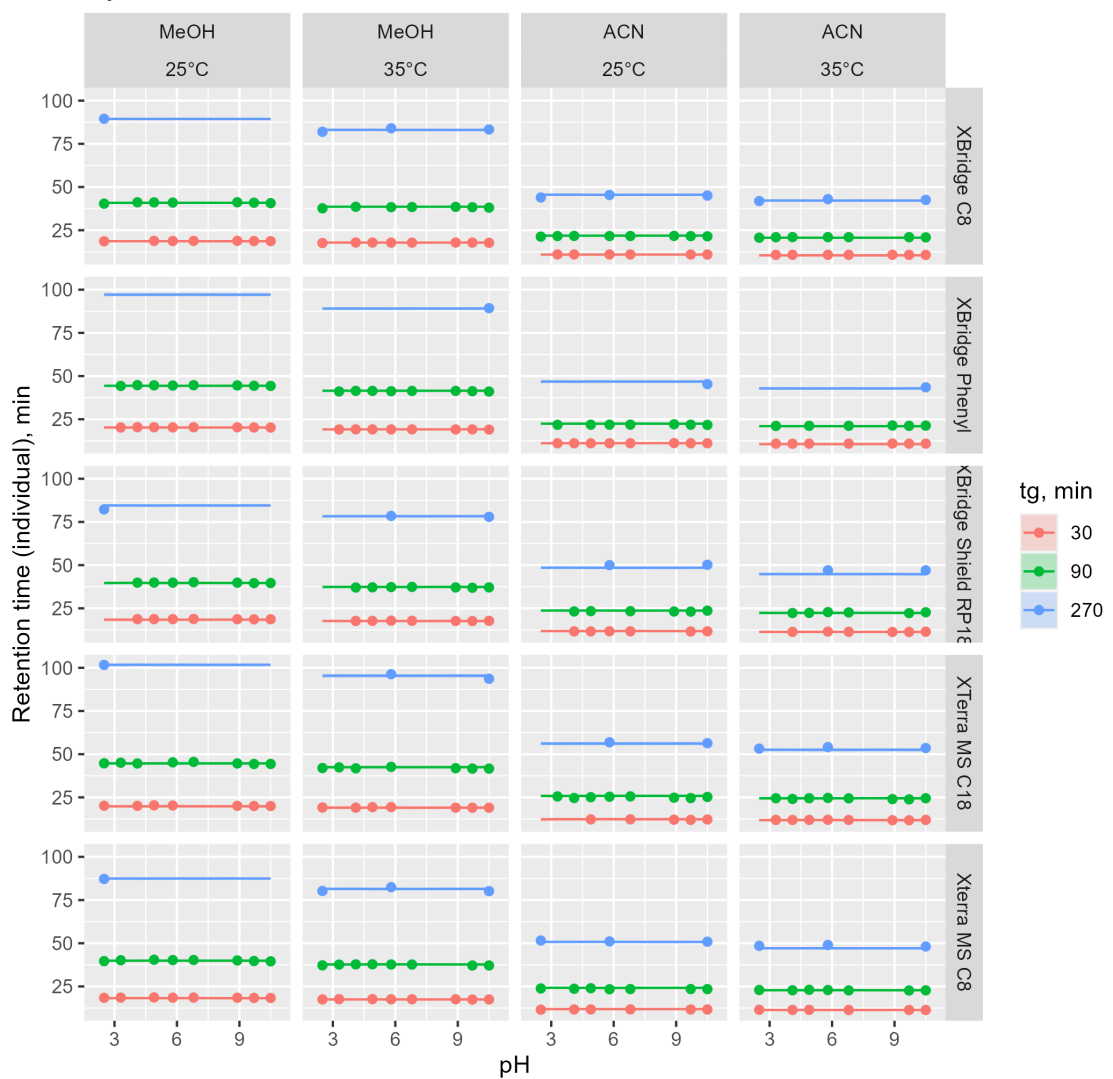


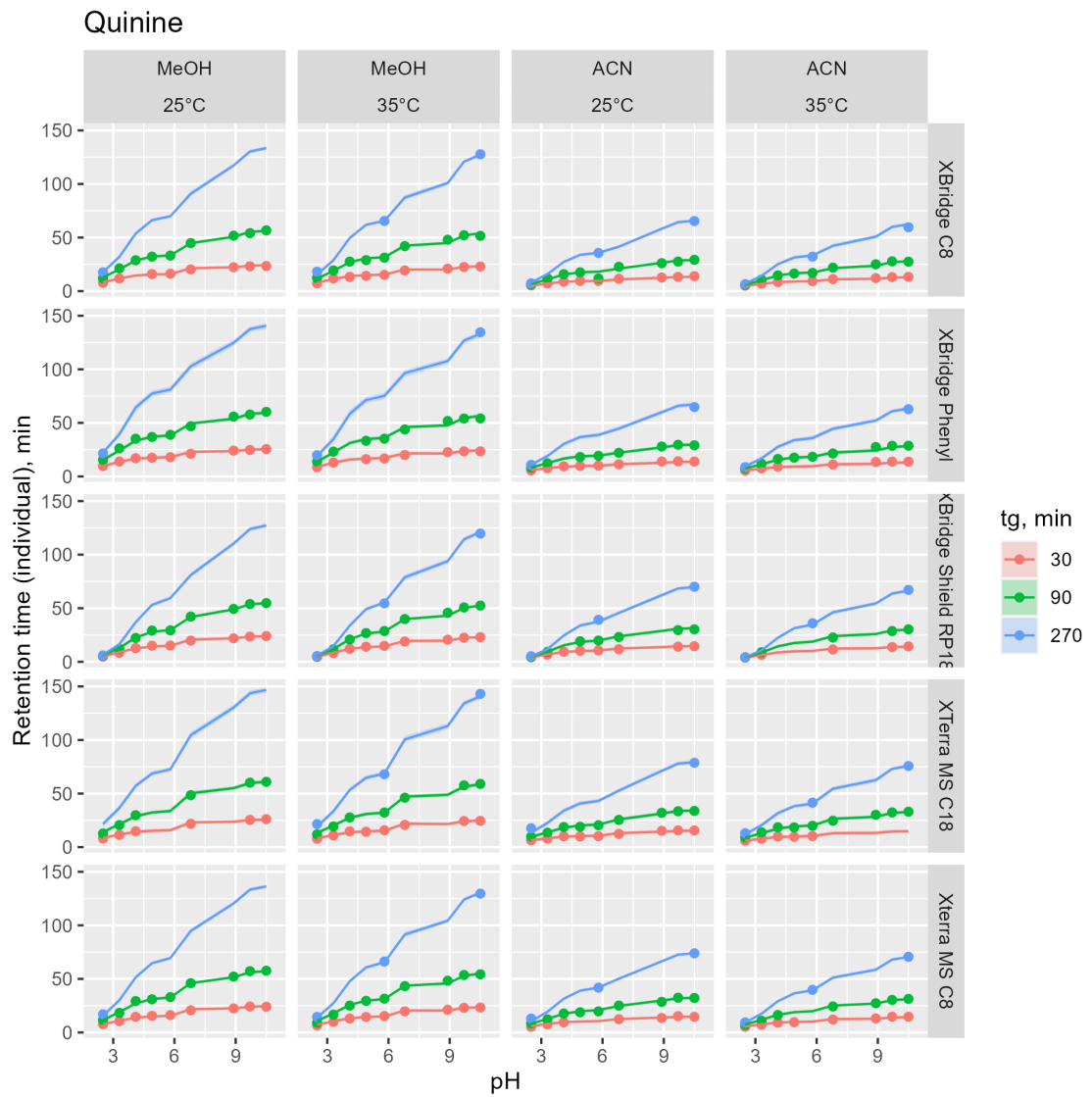


Pioglitazone



Hydrocortisone





7.2 Population predictions:

```
analyte_ID_sample <- c(9,17,33,58,140,180)

for(i in 1:length(analyte_ID_sample)){
  p <- ggplot()+
    geom_point(data=data[which(data$METID %in% analyte_ID_sample[i]),],
               aes(x = pHs, y = RT, color = as.factor(tg)))+
    
```

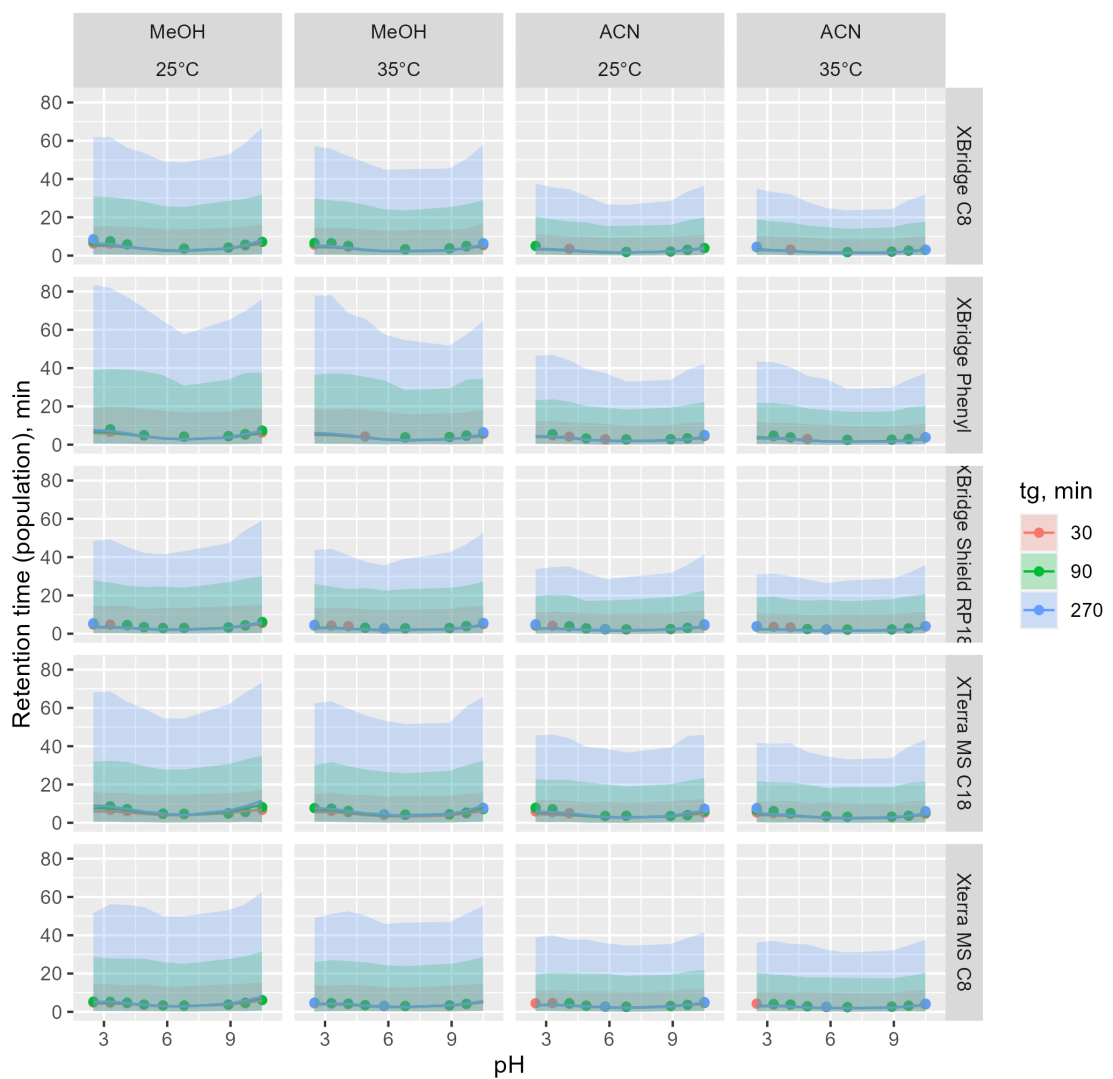
```

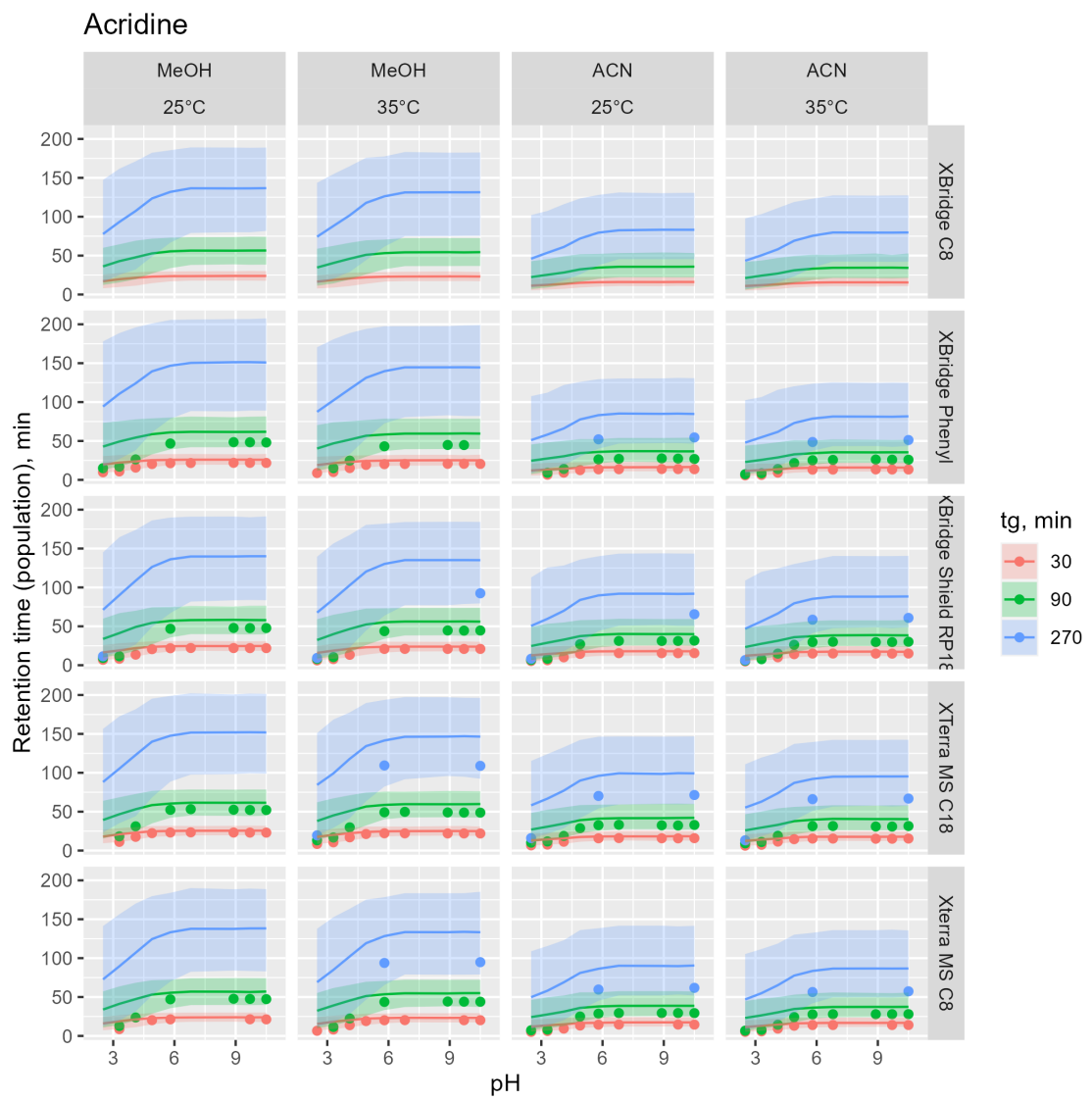
geom_line(data=datasim[which(datasim$METID %in% analyte_ID_sample[i]),],
          aes(x = pHs, y = trPred_m, color = as.factor(tg)))+
  geom_ribbon(data=datasim[which(datasim$METID %in% analyte_ID_sample[i]),],
              aes(x = pHs, ymin = trPred_l, ymax = trPred_h, fill = as.factor(tg)), alpha=0.2)+ 
  facet_grid(ColumnNames~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+ 
  labs(title=paste(dataNames>Name[analyte_ID_sample[i]]),
       x ="pH",
       y = "Retention time (population), min",
       color = "tg, min",
       fill = "tg, min")
print(p)

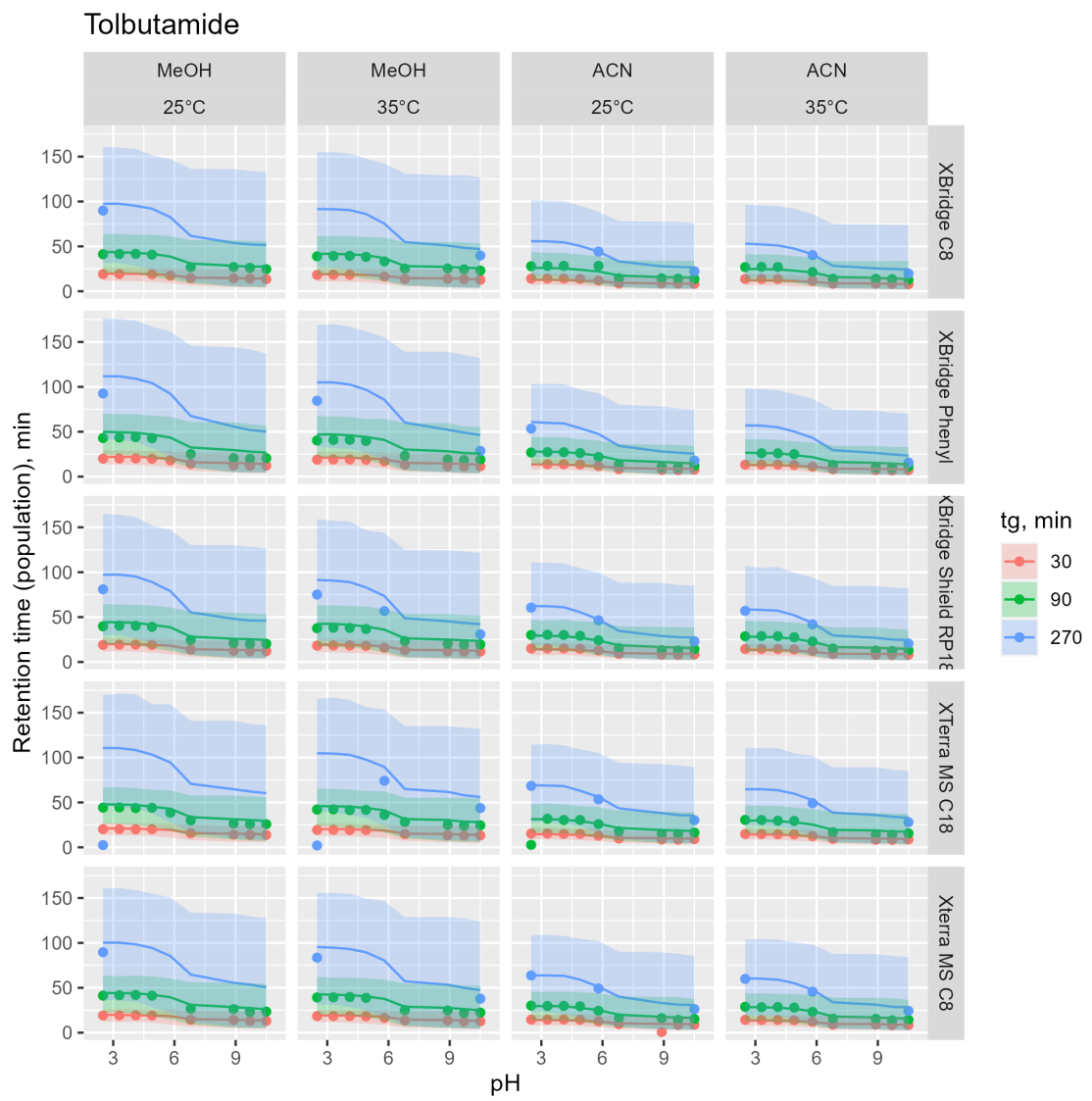
ggsave(paste0("figures\\concordanceplots\\", paste(dataNames>Name[analyte_ID_sample[i]]),
}

```

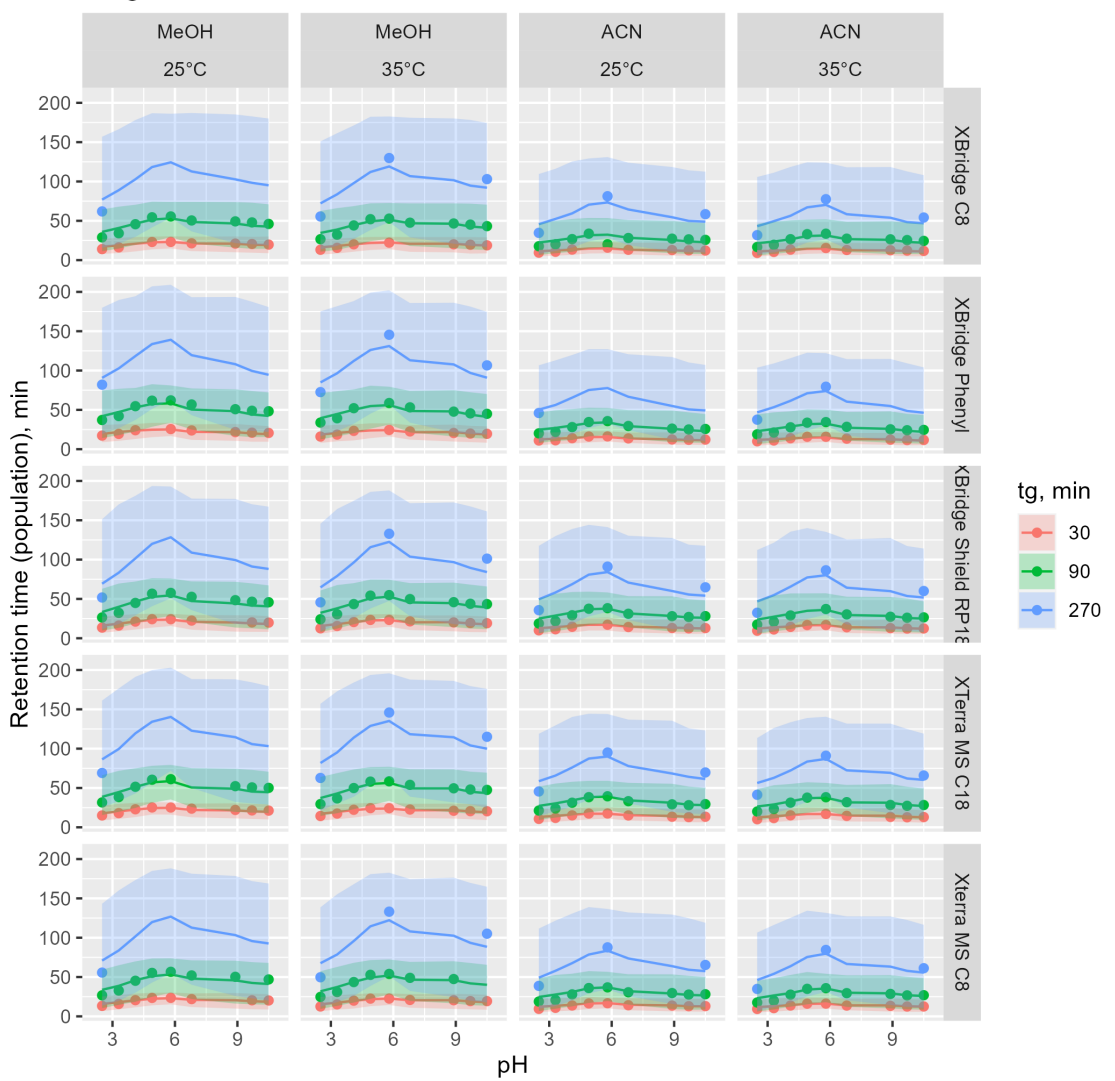
Baclofen

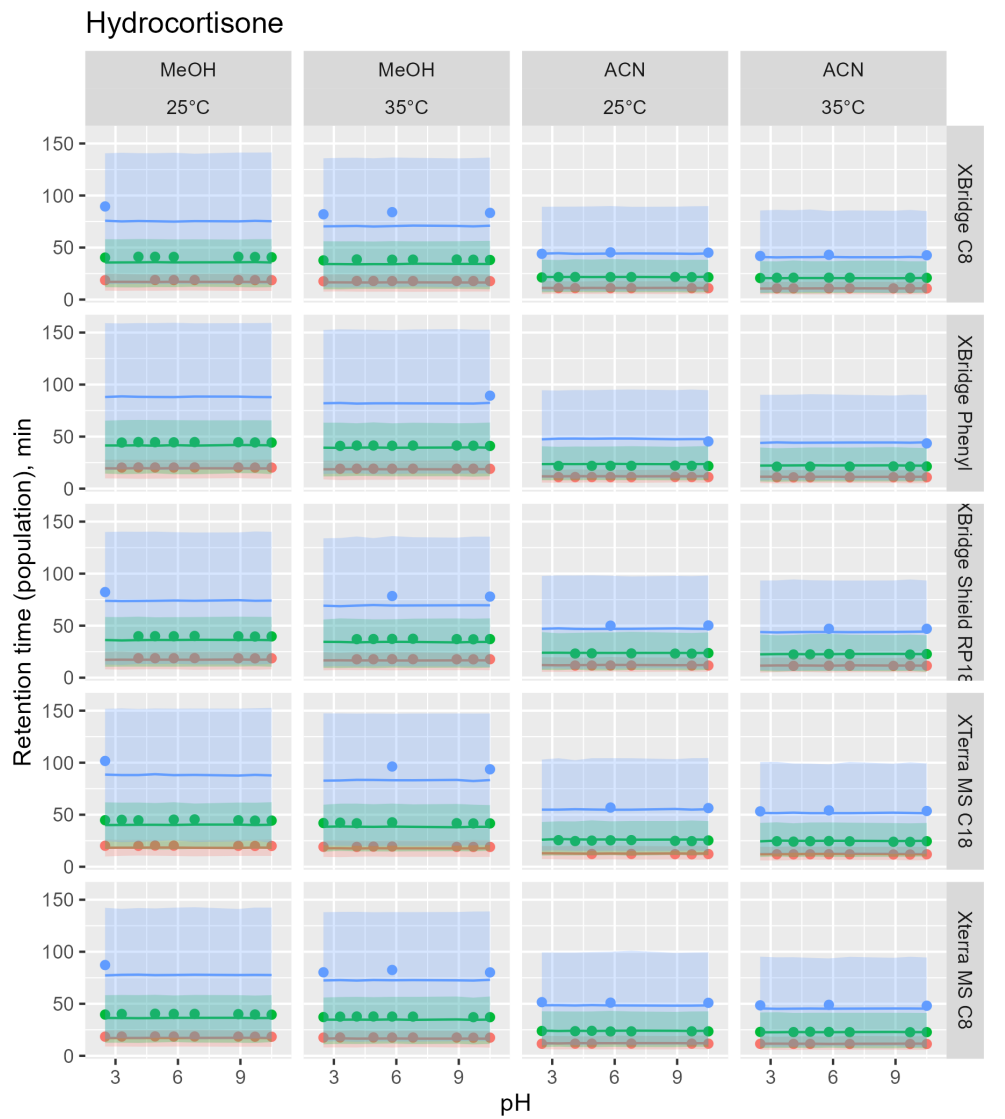


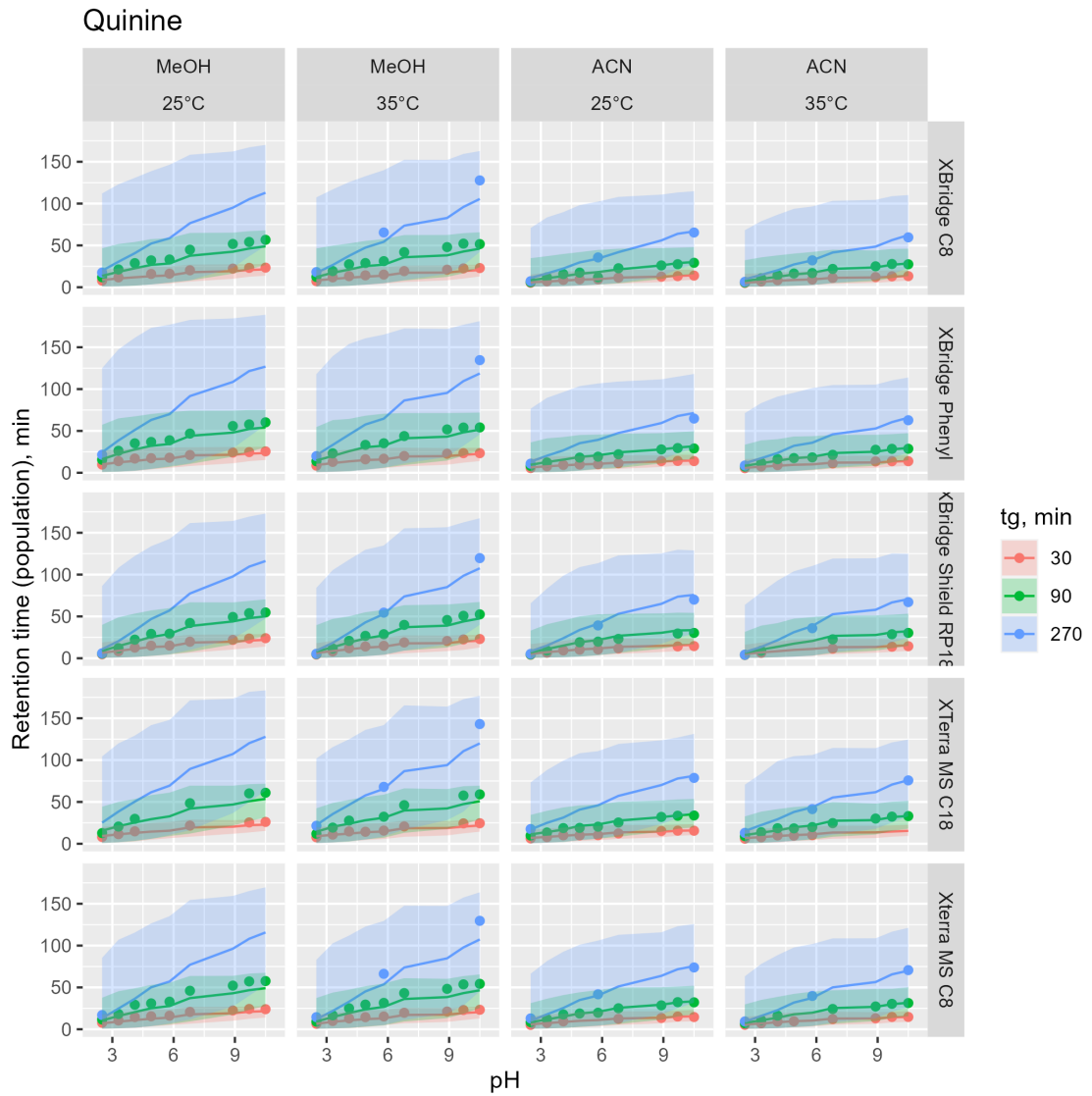




Pioglitazone







8 Uncertainty chromatogram

Below is the example of an uncertainty chromatogram expected for $tg = 90$ min, $pH = 4.9$, $Temp = 25^\circ C$ in MeOH. The individual and population predictions are shown.

```
model_epred <- cmdstan_model("stan/hplc-gra-fivecolumns-epred.stan")
fit_epred <- model_epred$generate_quantities(fitsim,
```

```

data = datastructsim,
seed = 123,
parallel_chains = 4,
output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-epred-202308090811-1-779
))

draws_epred_df <- as_draws_df(x$generated_quantities)

# draws_epred_df <- fit_epred$draws(format = "df")

analyte_ID_sample <-c(9,17,33,58,140,180)

col.labs <- c("XBridge Shield RP18", "XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xter
wpCond= data.frame()
wpPred= data.frame()

for (i in 1:5) {
  idx <- which(datasim$METID %in% analyte_ID_sample &
    datasim$tg==90 & # c(30, 90, 270)
    datasim$pH== 4 & # c(1:9)
    datasim$Column==i & # c(1, 2)
    datasim$Mod2==1 & # c(1, 2)
    datasim$Temp==25) # c(25, 35)

  data_to_plot <- draws_epred_df[,which(colnames(draws_epred_df) %in% paste0("trHatCond[", id
  colnames(data_to_plot) <- paste(dataNames>Name[analyte_ID_sample])
  wpCond1 <- melt(data_to_plot)
  wpCond1$Column <- unname(col.labs[i])
  data_to_plot <- draws_epred_df[,which(colnames(draws_epred_df) %in% paste0("trHatPred[", id
  colnames(data_to_plot) <- paste(dataNames>Name[analyte_ID_sample])
  wpPred1 <- melt(data_to_plot)
  wpPred1$Column <- unname(col.labs[i])
  wpCond= rbind(wpCond,wpCond1)
  wpPred= rbind(wpPred,wpPred1)
}

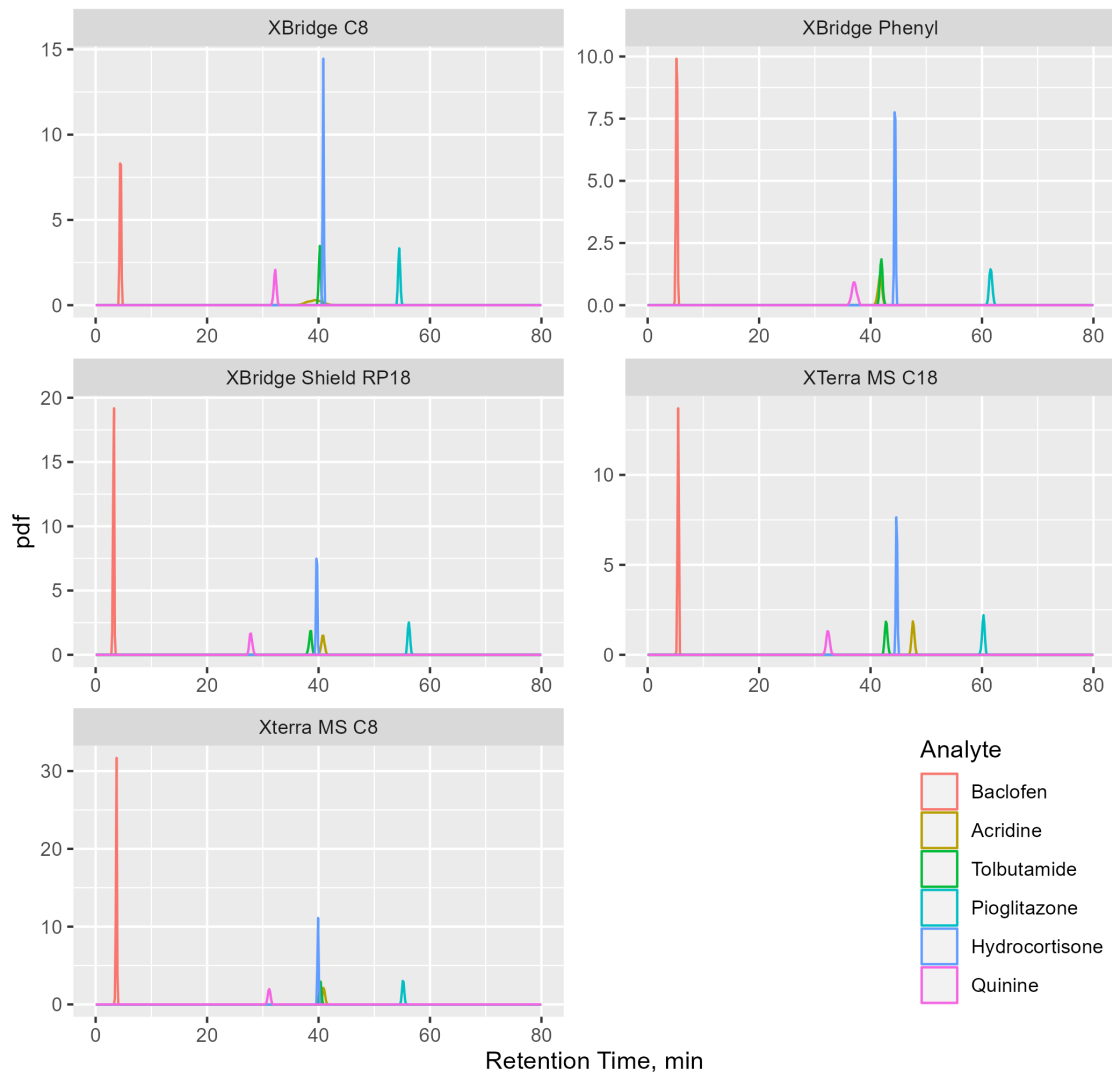
wpCond$Column=as.factor(wpCond$Column)

```

```
wpPred$Column=as.factor(wpPred$Column)

p1 <- ggplot(data = wpCond)+  
  geom_density(aes(x=value, colour=variable)) +  
  labs(title="uncertainty chromatogram (individual predictions)",  
    x = "Retention Time, min",  
    y = "pdf",  
    colour="Analyte") +  
  
  xlim(c(0,80)) +  
  facet_wrap(.~Column, nrow=3, scales = "free") +  
  theme(legend.position = c(1, 0),  
        legend.justification = c(1, 0))  
print(p1)
```

uncertainty chromatogram (individual predictions)



```

p2 <- ggplot(data = wpPred)+  

  geom_density(aes(x=value, colour=variable)) +  

  labs(title="uncertainty chromatogram (population predictions)",  

       x ="Retention Time, min",  

       y = "pdf",  

       colour="Analyte") +  

  xlim(c(0,80)) +  

  facet_wrap(.~Column, nrow=3,scales = "free") +  

  theme(legend.position = c(1, 0),

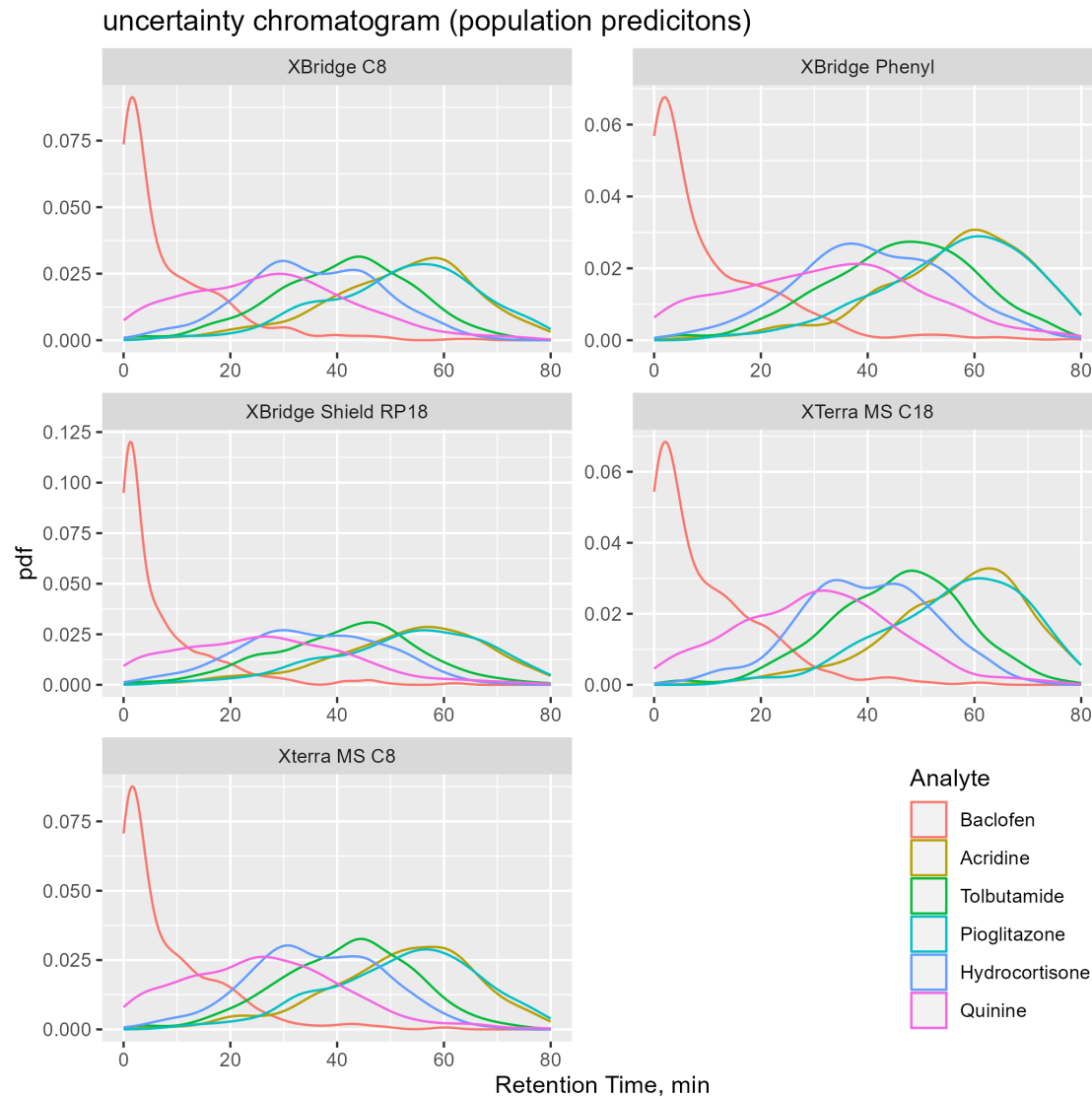
```

```

  legend.justification = c(1, 0))

print(p2)

```



```

ggsave(paste0("figures\\concordanceplots\\", "uncertaintychromatogram.individual", ".png"))

ggsave(paste0("figures\\concordanceplots\\", "uncertaintychromatogram.population", ".png"))

```

8.1 Isocratic predictions (population predictions)

To better asses the impact of parameters on retention we created graphs presenting the isocratic logarithm of retention factor vs. for selected analytes. Separate graphs are shown for each dissociation form (r=1,r=2,r=3). Here the population predictions are shown:

```
analyte_ID_sample <-c(9,17,33,58,140,180)

for(i in 1:length(analyte_ID_sample)){

  idx_analyte = which(unique(data$METID)==analyte_ID_sample[i])
  draws_df_subset <- draws_epred_df[,which(colnames(draws_epred_df) %in% c(
    sprintf("logkwxPred[%s,1,1]",idx_analyte),
    sprintf("logkwxPred[%s,1,2]",idx_analyte),
    sprintf("logkwxPred[%s,1,3]",idx_analyte),
    sprintf("logkwxPred[%s,2,1]",idx_analyte),
    sprintf("logkwxPred[%s,2,2]",idx_analyte),
    sprintf("logkwxPred[%s,2,3]",idx_analyte),
    sprintf("logkwxPred[%s,3,1]",idx_analyte),
    sprintf("logkwxPred[%s,3,2]",idx_analyte),
    sprintf("logkwxPred[%s,3,3]",idx_analyte),
    sprintf("logkwxPred[%s,4,1]",idx_analyte),
    sprintf("logkwxPred[%s,4,2]",idx_analyte),
    sprintf("logkwxPred[%s,4,3]",idx_analyte),
    sprintf("logkwxPred[%s,5,1]",idx_analyte),
    sprintf("logkwxPred[%s,5,2]",idx_analyte),
    sprintf("logkwxPred[%s,5,3]",idx_analyte),
    sprintf("S1xPred[%s,1,1,1]",idx_analyte),
    sprintf("S1xPred[%s,1,1,2]",idx_analyte),
    sprintf("S1xPred[%s,1,1,3]",idx_analyte),
    sprintf("S1xPred[%s,2,1,1]",idx_analyte),
    sprintf("S1xPred[%s,2,1,2]",idx_analyte),
    sprintf("S1xPred[%s,2,1,3]",idx_analyte),
    sprintf("S1xPred[%s,1,2,1]",idx_analyte),
    sprintf("S1xPred[%s,1,2,2]",idx_analyte),
    sprintf("S1xPred[%s,1,2,3]",idx_analyte),
    sprintf("S1xPred[%s,2,2,1]",idx_analyte),
    sprintf("S1xPred[%s,2,2,2]",idx_analyte),
    sprintf("S1xPred[%s,2,2,3]",idx_analyte),
    sprintf("S1xPred[%s,1,3,1]",idx_analyte),
    sprintf("S1xPred[%s,1,3,2]",idx_analyte),
    sprintf("S1xPred[%s,1,3,3]",idx_analyte),
```

```

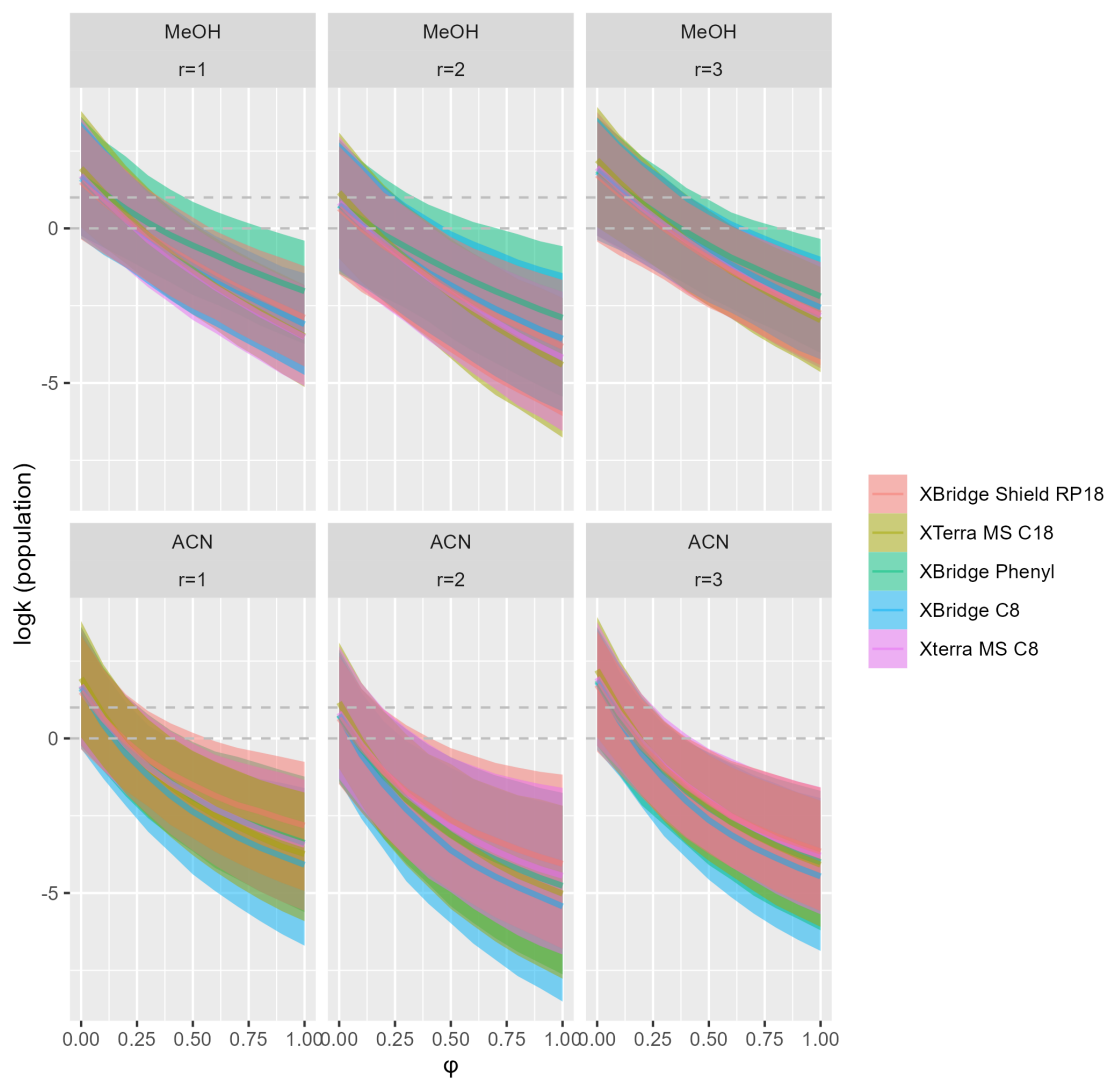
sprintf("S1xPred[%s,2,3,1]",idx_analyte),
sprintf("S1xPred[%s,2,3,2]",idx_analyte),
sprintf("S1xPred[%s,2,3,3]",idx_analyte),
sprintf("S1xPred[%s,1,4,1]",idx_analyte),
sprintf("S1xPred[%s,1,4,2]",idx_analyte),
sprintf("S1xPred[%s,1,4,3]",idx_analyte),
sprintf("S1xPred[%s,2,4,1]",idx_analyte),
sprintf("S1xPred[%s,2,4,2]",idx_analyte),
sprintf("S1xPred[%s,2,4,3]",idx_analyte),
sprintf("S1xPred[%s,1,5,1]",idx_analyte),
sprintf("S1xPred[%s,1,5,2]",idx_analyte),
sprintf("S1xPred[%s,1,5,3]",idx_analyte),
sprintf("S1xPred[%s,2,5,1]",idx_analyte),
sprintf("S1xPred[%s,2,5,2]",idx_analyte),
sprintf("S1xPred[%s,2,5,3]",idx_analyte),
"S2xPred[1,1]",
"S2xPred[2,1]",
".draw",".iteration",".chain"))]

p<-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwxPred[, c, r], S1xPred[, m, c, r], S2xPred[m, ]) %>%
  filter(r<=R[idx_analyte]+1) %>%
  tidyrr::expand_grid(fi = seq(0,1,0.1)) %>%
  mutate(logkPred = logkwxPred-S1xPred*(1+S2xPred)*fi/(1+S2xPred*fi)) %>%
  ggplot(aes(x = fi, y = logkPred, color = as.factor(c), fill = as.factor(c))) +
  ggdist::stat_lineribbon(.width = c(.90), alpha = 1/2) +
  facet_wrap(m~r, nrow = 2, labeller = labeller(m=mod.labs,r=diss.labs))+
  labs(title=paste(dataNames>Name[analyte_ID_sample[i]]),color= " ",fill= " ", y = "logk (",
  scale_fill_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl", "XBridge Phenyl RP18")),
  scale_color_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl", "XBridge Phenyl RP18")),
  geom_hline(yintercept= c(0,1), linetype="dashed",color="gray")
  print(p)

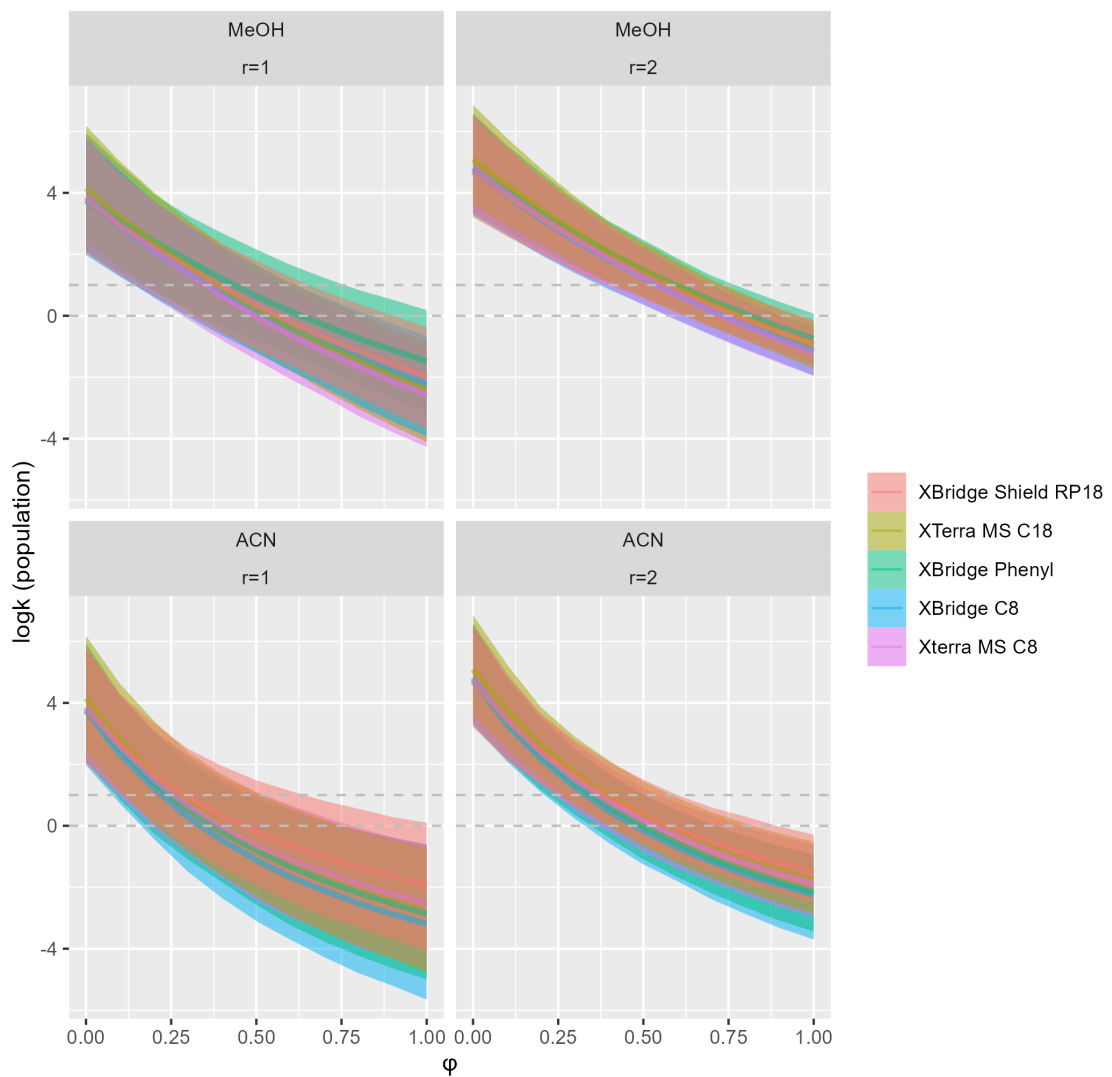
ggsave(paste0("figures\\izoparam\\", paste(dataNames>Name[analyte_ID_sample[i]]), ".isopre")
}

```

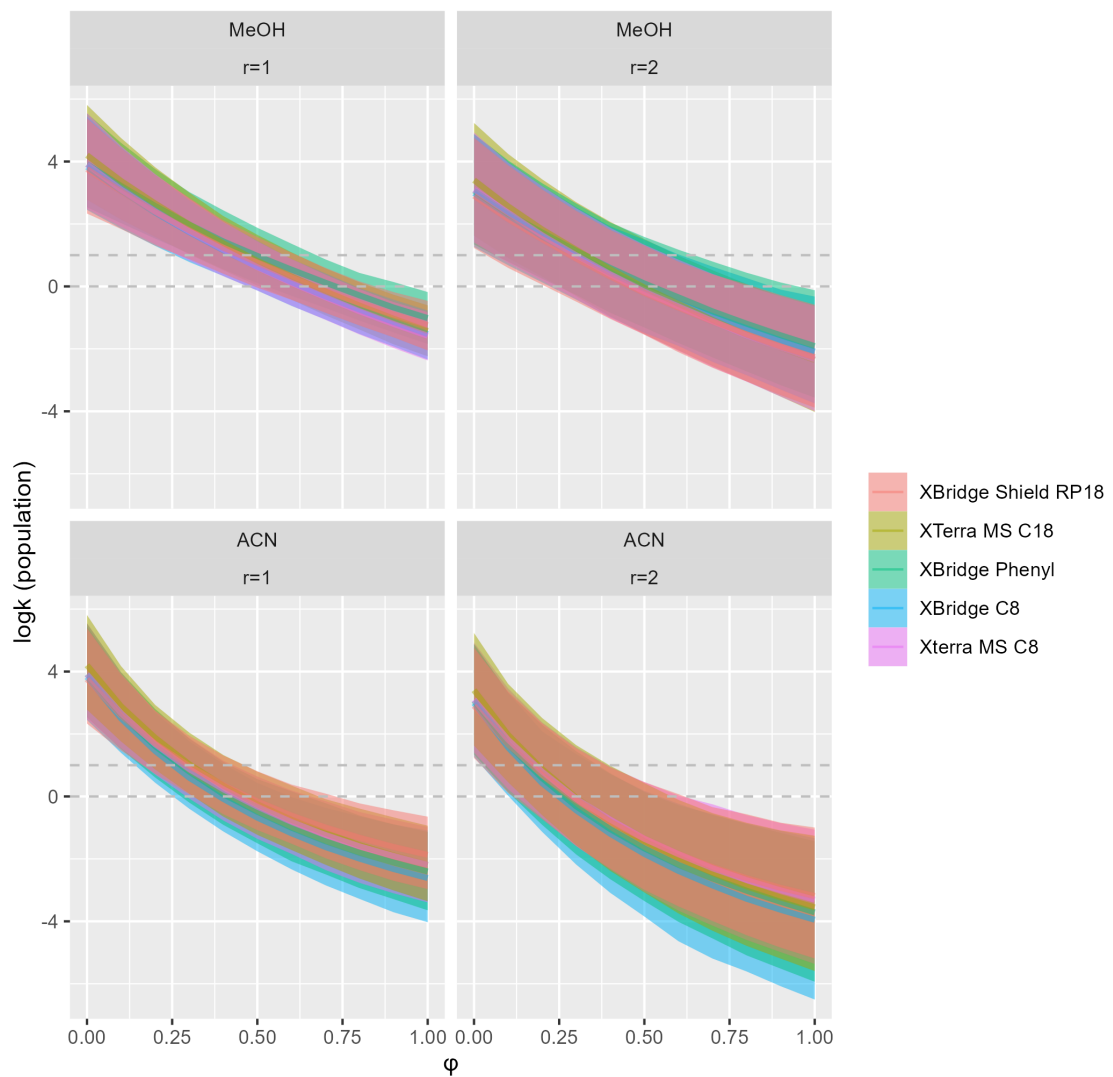
Baclofen



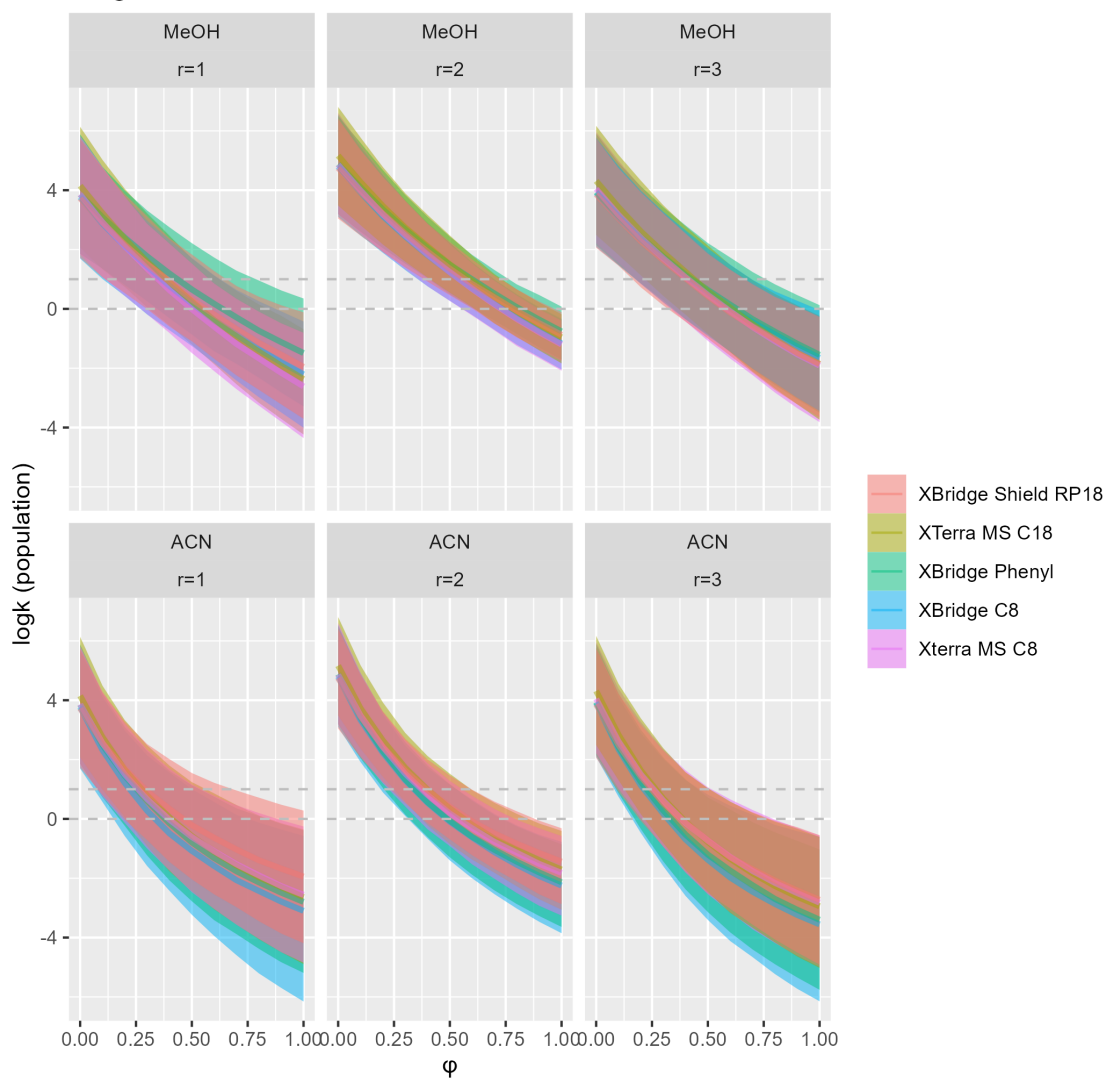
Acridine



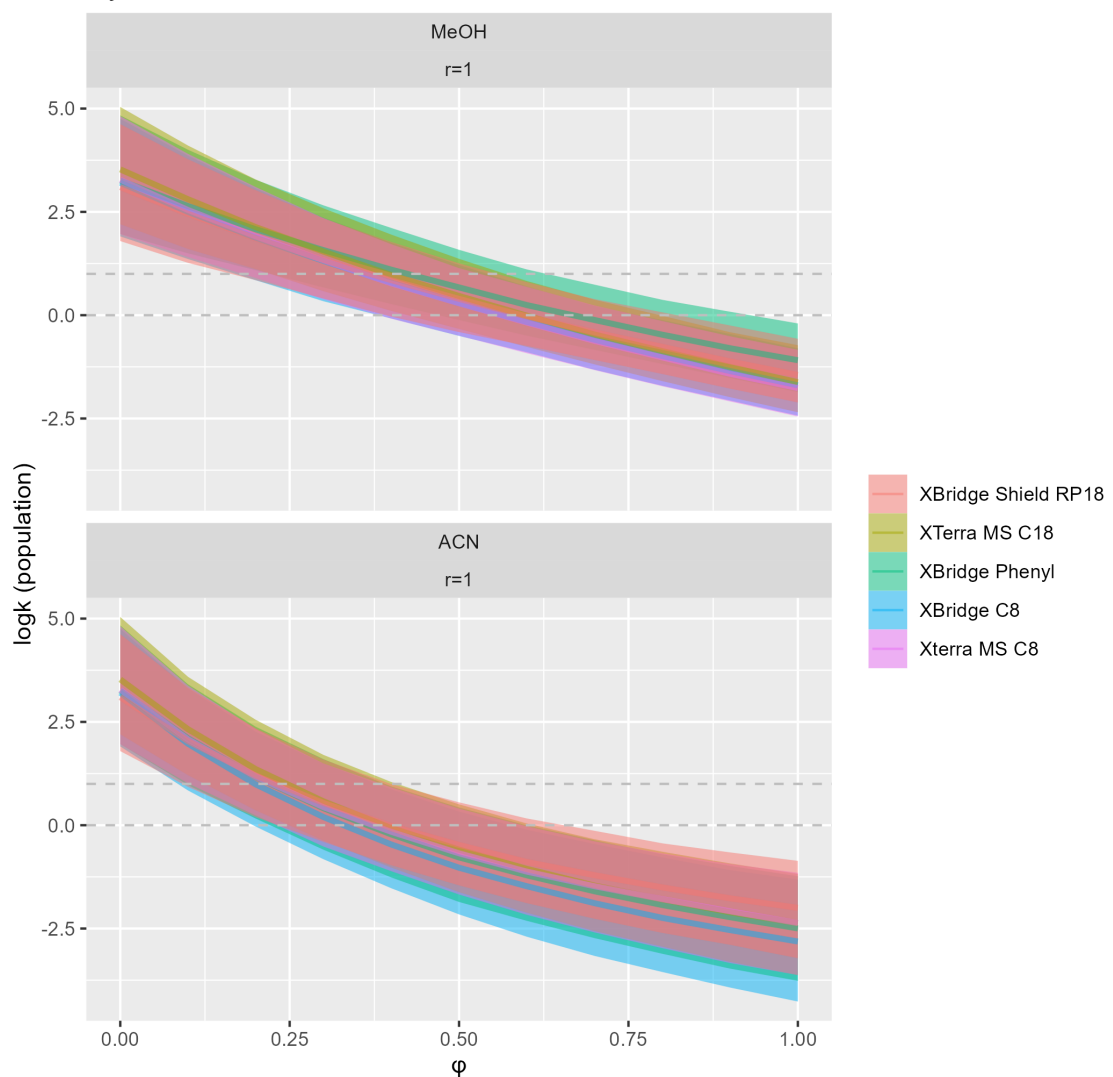
Tolbutamide



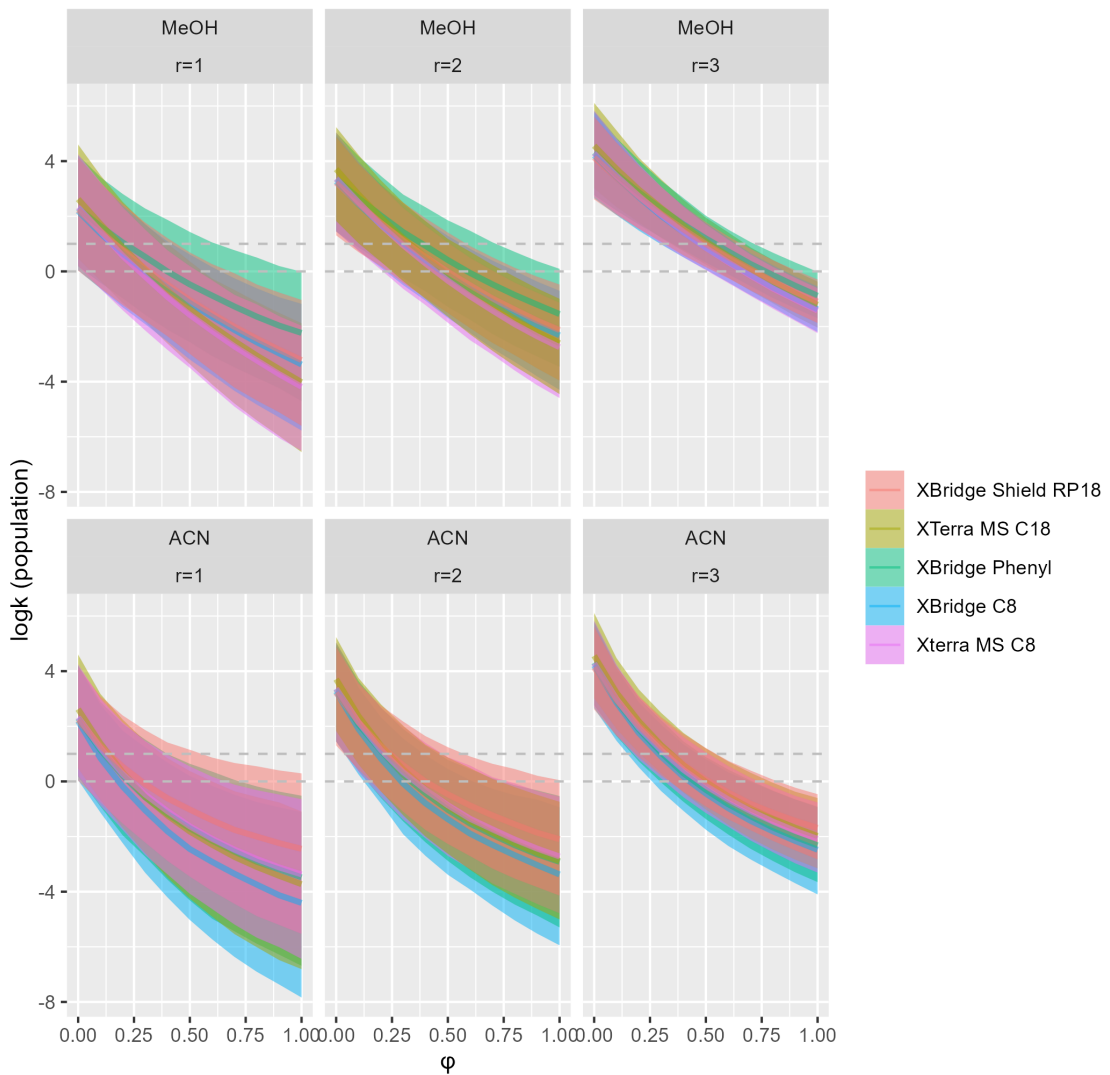
Pioglitazone



Hydrocortisone



Quinine



Similarly we can quantify the column effects (between column differences in logk using XBridge Shield RP18 as a reference column):

```
analyte_ID_sample <- c(9,17,33,58,140,180)

for(i in 1:length(analyte_ID_sample)){
  idx_analyte = which(unique(data$METID)==analyte_ID_sample[i])
```

```

draws_df_subset <- draws_epred_df[,which(colnames(draws_epred_df) %in% c(
  sprintf("logkwxPred[%s,1,1]",idx_analyte),
  sprintf("logkwxPred[%s,1,2]",idx_analyte),
  sprintf("logkwxPred[%s,1,3]",idx_analyte),
  sprintf("logkwxPred[%s,2,1]",idx_analyte),
  sprintf("logkwxPred[%s,2,2]",idx_analyte),
  sprintf("logkwxPred[%s,2,3]",idx_analyte),
  sprintf("logkwxPred[%s,3,1]",idx_analyte),
  sprintf("logkwxPred[%s,3,2]",idx_analyte),
  sprintf("logkwxPred[%s,3,3]",idx_analyte),
  sprintf("logkwxPred[%s,4,1]",idx_analyte),
  sprintf("logkwxPred[%s,4,2]",idx_analyte),
  sprintf("logkwxPred[%s,4,3]",idx_analyte),
  sprintf("logkwxPred[%s,5,1]",idx_analyte),
  sprintf("logkwxPred[%s,5,2]",idx_analyte),
  sprintf("logkwxPred[%s,5,3]",idx_analyte),
  sprintf("S1xPred[%s,1,1,1]",idx_analyte),
  sprintf("S1xPred[%s,1,1,2]",idx_analyte),
  sprintf("S1xPred[%s,1,1,3]",idx_analyte),
  sprintf("S1xPred[%s,2,1,1]",idx_analyte),
  sprintf("S1xPred[%s,2,1,2]",idx_analyte),
  sprintf("S1xPred[%s,2,1,3]",idx_analyte),
  sprintf("S1xPred[%s,1,2,1]",idx_analyte),
  sprintf("S1xPred[%s,1,2,2]",idx_analyte),
  sprintf("S1xPred[%s,1,2,3]",idx_analyte),
  sprintf("S1xPred[%s,2,2,1]",idx_analyte),
  sprintf("S1xPred[%s,2,2,2]",idx_analyte),
  sprintf("S1xPred[%s,2,2,3]",idx_analyte),
  sprintf("S1xPred[%s,1,3,1]",idx_analyte),
  sprintf("S1xPred[%s,1,3,2]",idx_analyte),
  sprintf("S1xPred[%s,1,3,3]",idx_analyte),
  sprintf("S1xPred[%s,2,3,1]",idx_analyte),
  sprintf("S1xPred[%s,2,3,2]",idx_analyte),
  sprintf("S1xPred[%s,2,3,3]",idx_analyte),
  sprintf("S1xPred[%s,1,4,1]",idx_analyte),
  sprintf("S1xPred[%s,1,4,2]",idx_analyte),
  sprintf("S1xPred[%s,1,4,3]",idx_analyte),
  sprintf("S1xPred[%s,2,4,1]",idx_analyte),
  sprintf("S1xPred[%s,2,4,2]",idx_analyte),
  sprintf("S1xPred[%s,2,4,3]",idx_analyte),
  sprintf("S1xPred[%s,1,5,1]",idx_analyte),

```

```

sprintf("S1xPred[%s,1,5,2]",idx_analyte),
sprintf("S1xPred[%s,1,5,3]",idx_analyte),
sprintf("S1xPred[%s,2,5,1]",idx_analyte),
sprintf("S1xPred[%s,2,5,2]",idx_analyte),
sprintf("S1xPred[%s,2,5,3]",idx_analyte),
"S2xPred[1,1]",
"S2xPred[2,1]",
".draw",".iteration",".chain"))]

p<-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwxPred[, c, r], S1xPred[, m, c, r], S2xPred[m, ]) %>%
  filter(r<=R[idx_analyte]+1) %>%
  tidyr::expand_grid(fi = seq(0,1,0.05)) %>%
  mutate(logkPred = logkwxPred-S1xPred*(1+S2xPred)*fi/(1+S2xPred*fi)) %>%
  select(.draw,c,r,m,logkPred,fi) %>%
  tidyr::pivot_wider(names_from = c, values_from = logkPred) %>%
  mutate(cdk2 = `2`-`1` ) %>%
  mutate(cdk3 = `3`-`1` ) %>%
  mutate(cdk4 = `4`-`1` ) %>%
  mutate(cdk5 = `5`-`1` ) %>%
  tidyr::pivot_longer(cdk2:cdk5,names_to = "names_cdk", values_to = "cdk")%>%
  ggplot(aes(x = fi, y = cdk, color = as.factor(names_cdk), fill = as.factor(names_cdk))) +
  ggdist::stat_lineribbon(.width = c(.90), alpha = 1/2) +
  facet_wrap(m~r, nrow = 2,labeller = labeller(m=mod.labs,r=diss.labs)) +
  labs(y = "Between column difference in log (population)", title=paste(dataNames>Name[a
  scale_fill_discrete(labels= c("XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xterra M
  scale_color_discrete(labels= c("XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xterra
  geom_hline(yintercept= c(0), linetype="dashed",color="gray") +
  coord_cartesian(xlim=c(0,1),ylim=c(-1,1))

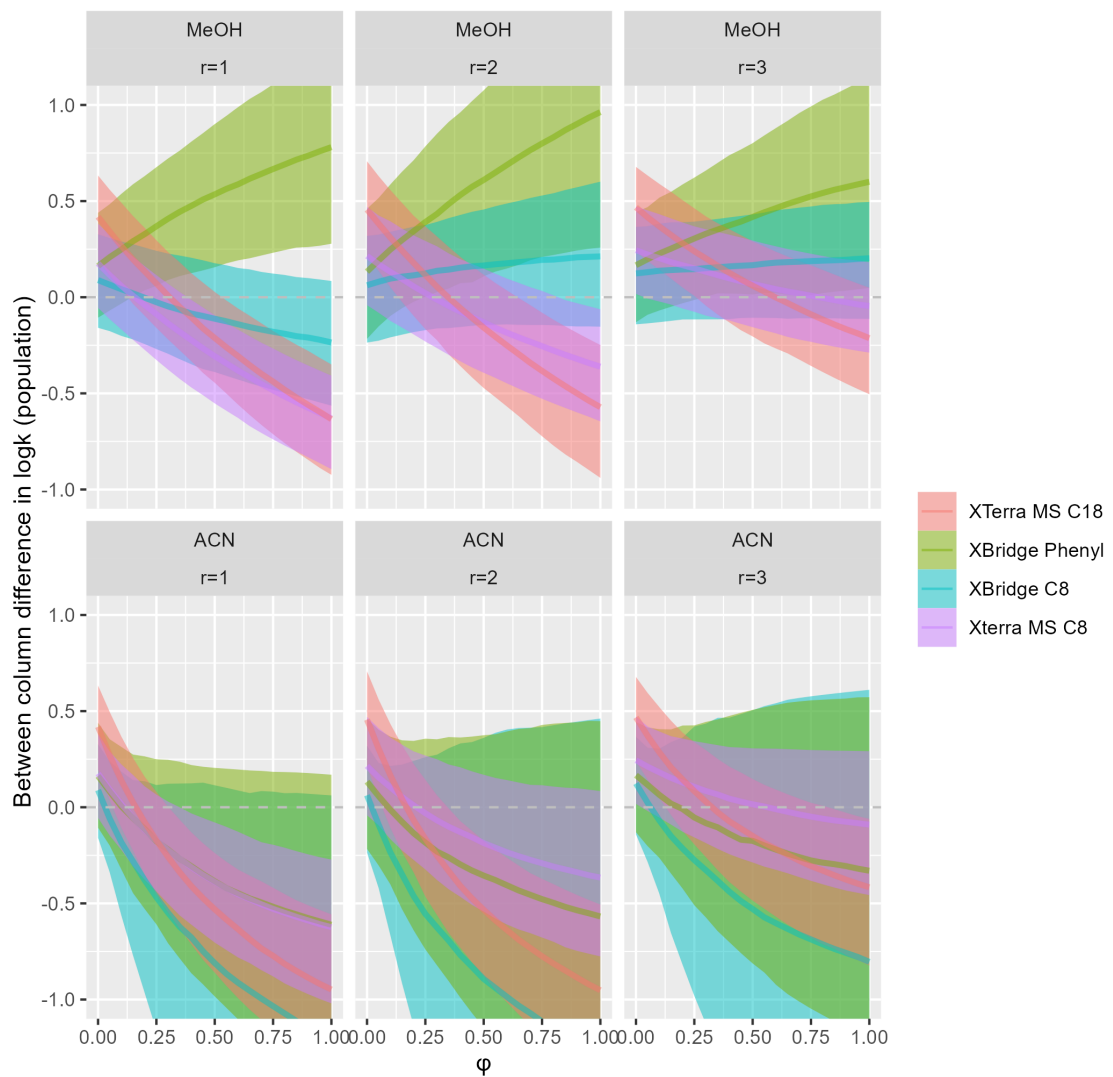
print(p)

ggsave(paste0("figures\\izoparam\\", paste(dataNames>Name[analyte_ID_sample[i]])), "isodif

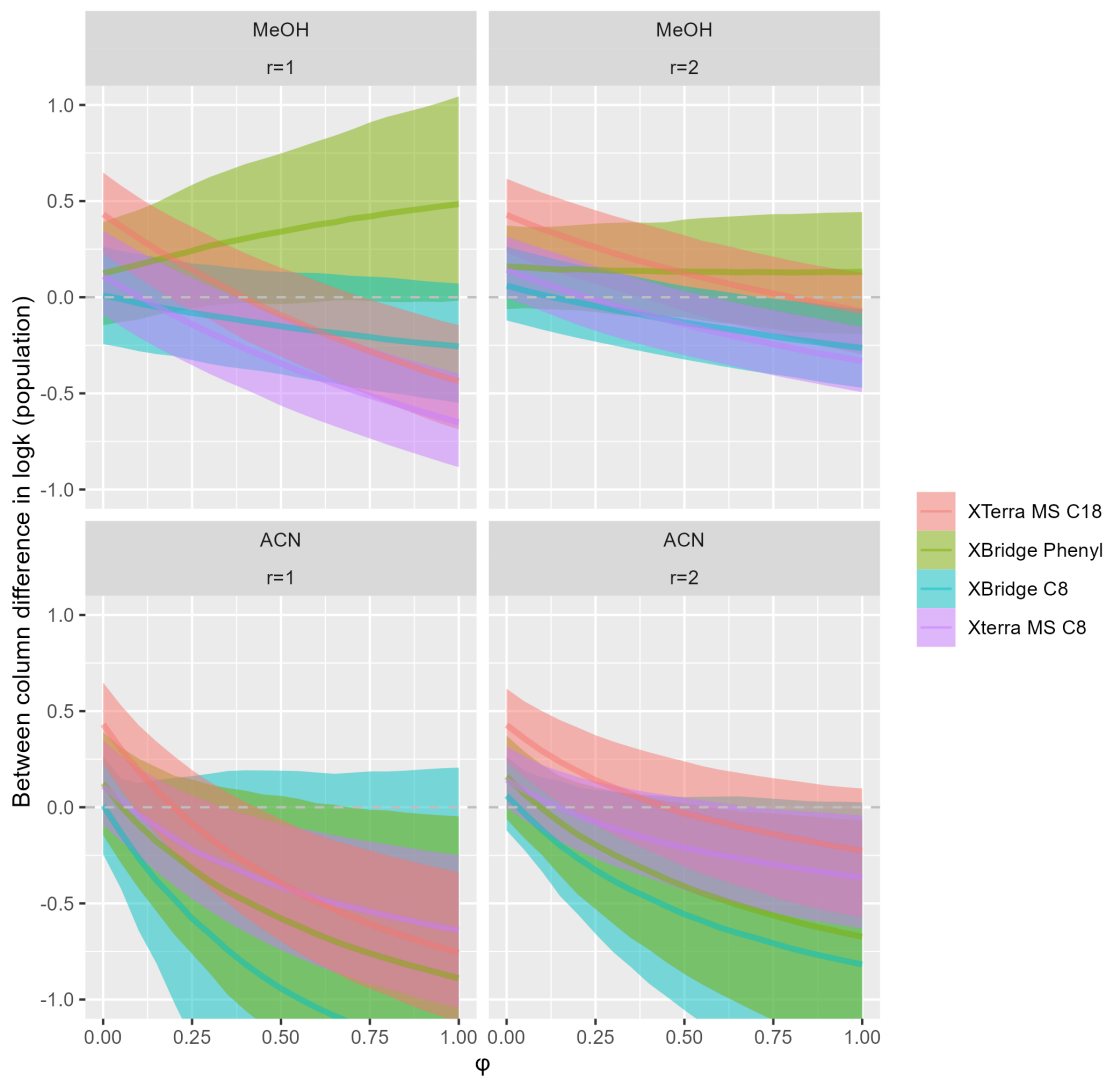
}

```

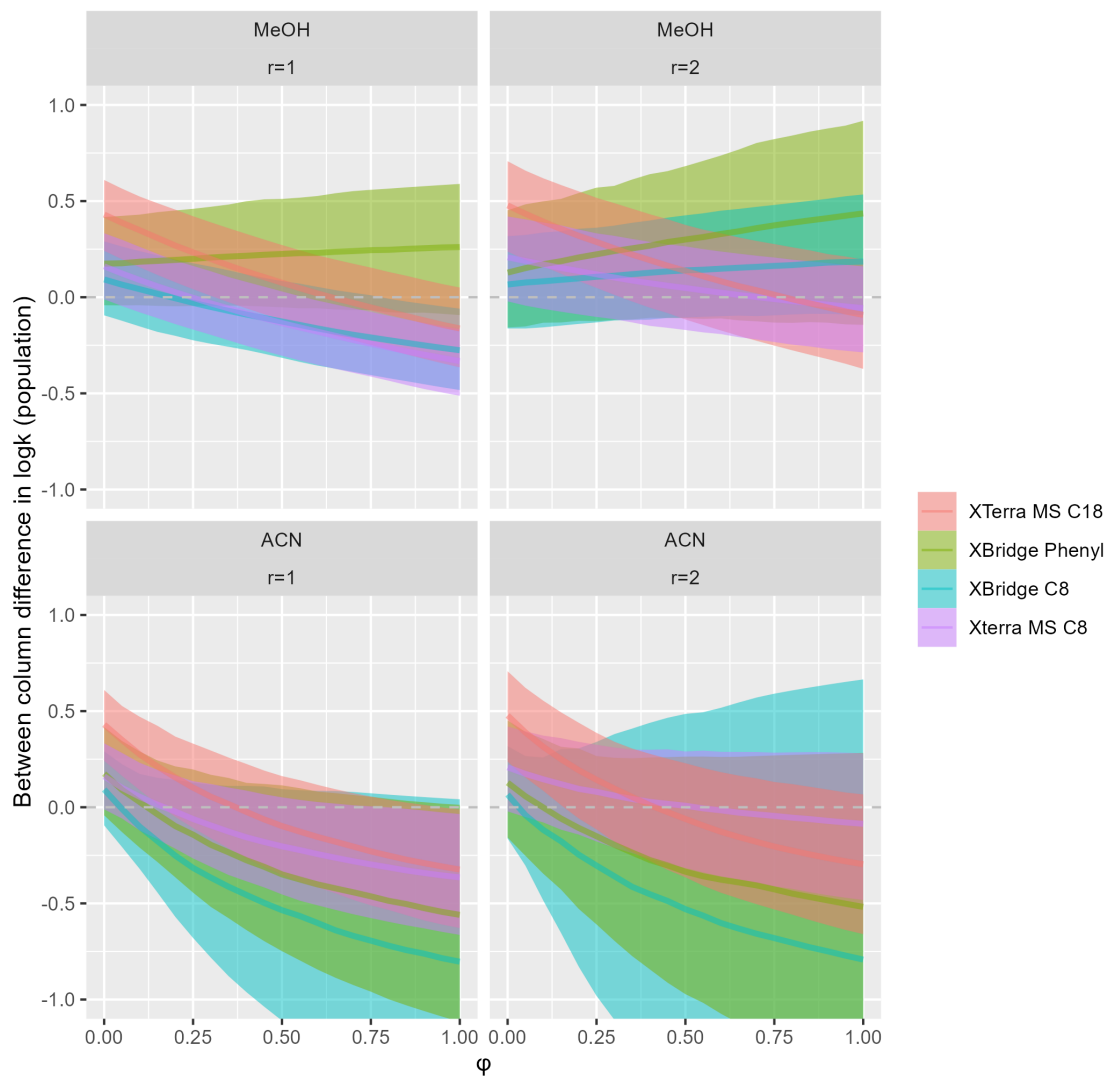
Baclofen



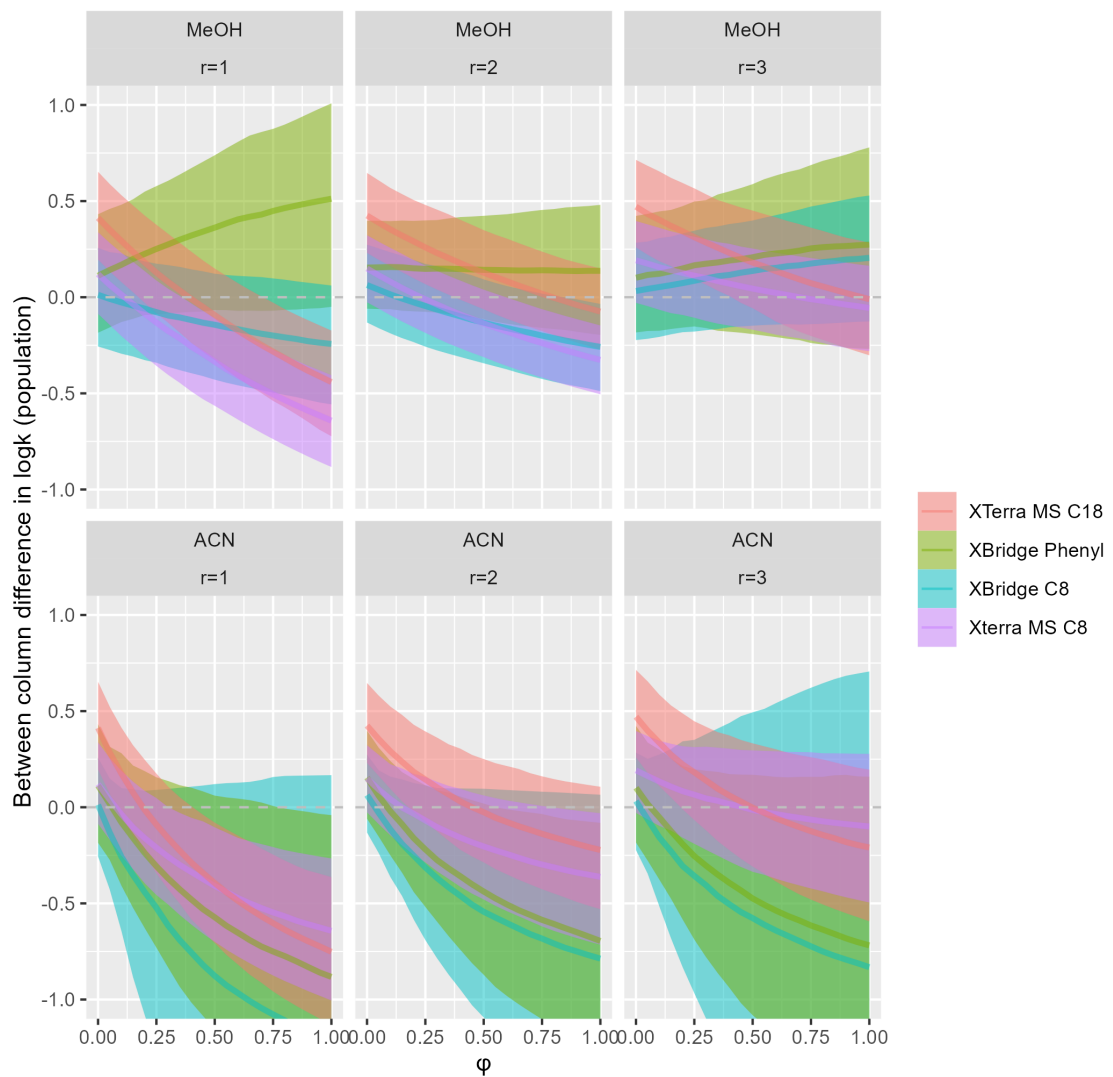
Acridine



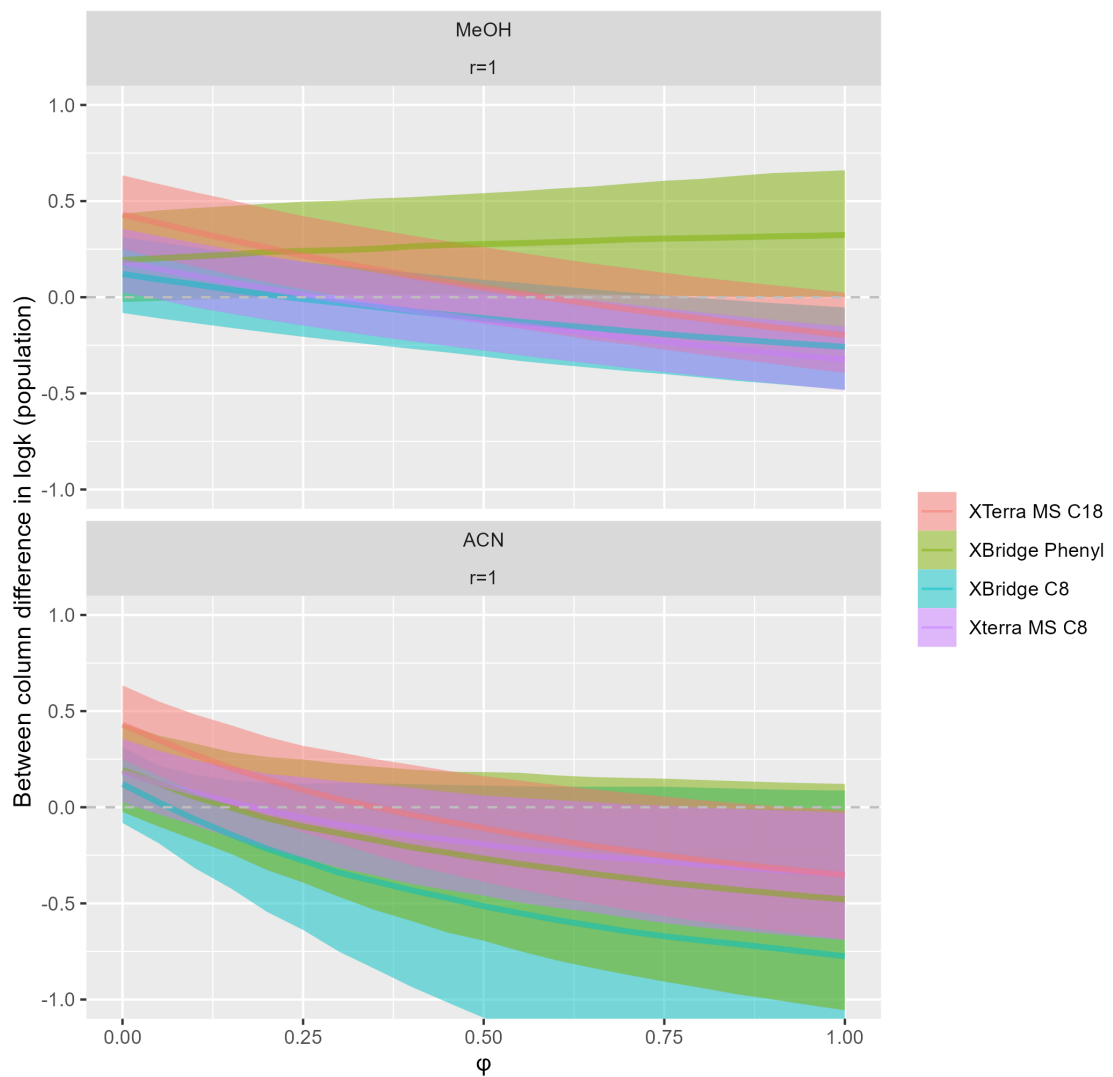
Tolbutamide

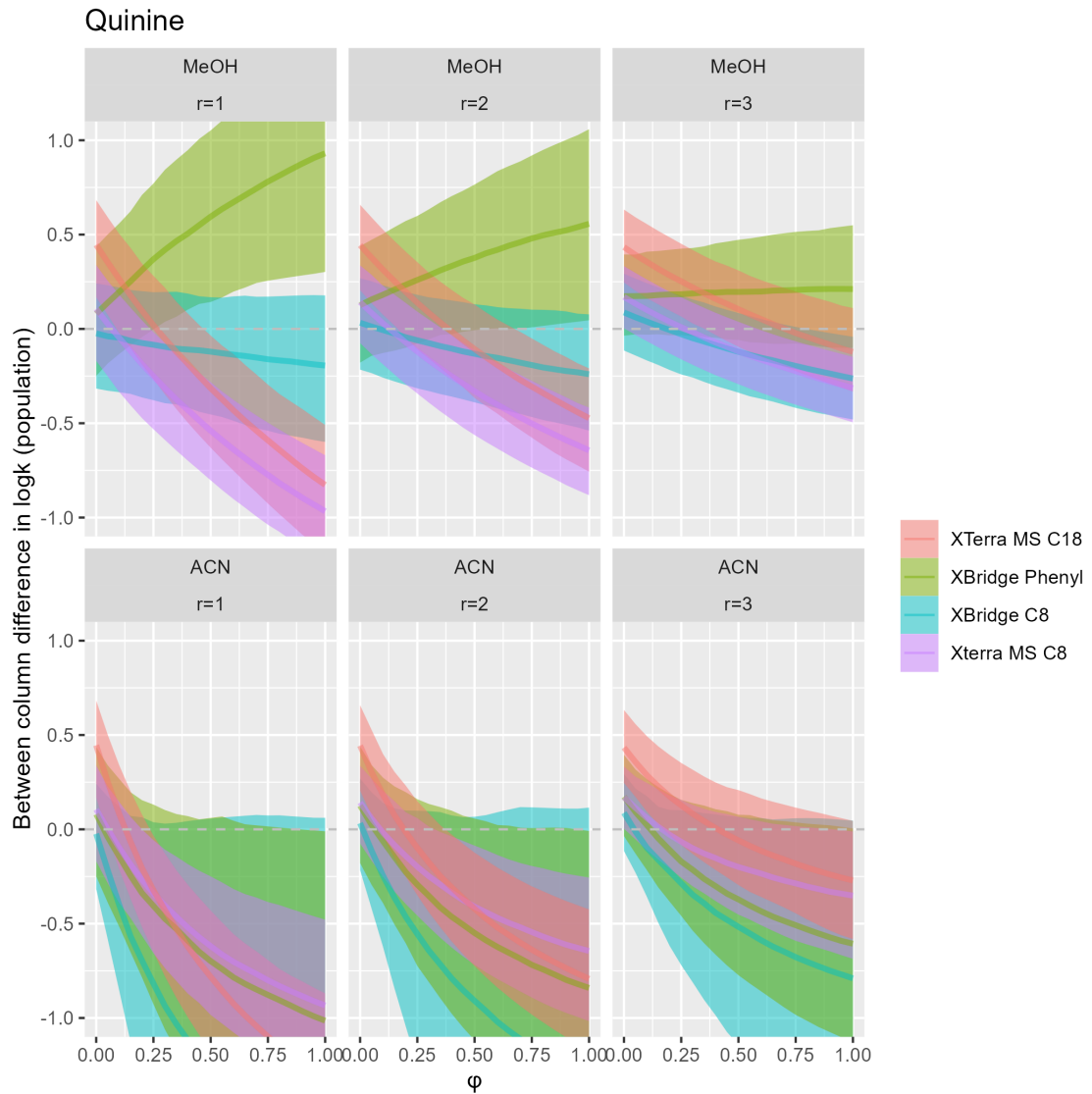


Pioglitazone



Hydrocortisone





or predict the organic modifier content leading to log of 1:

```
analyte_ID_sample <- c(9,17,33,58,140,180)

for(i in 1:length(analyte_ID_sample)){
  idx_analyte = which(unique(data$METID)==analyte_ID_sample[i])

  draws_df_subset <- draws_epred_df[,which(colnames(draws_epred_df) %in% c(
```

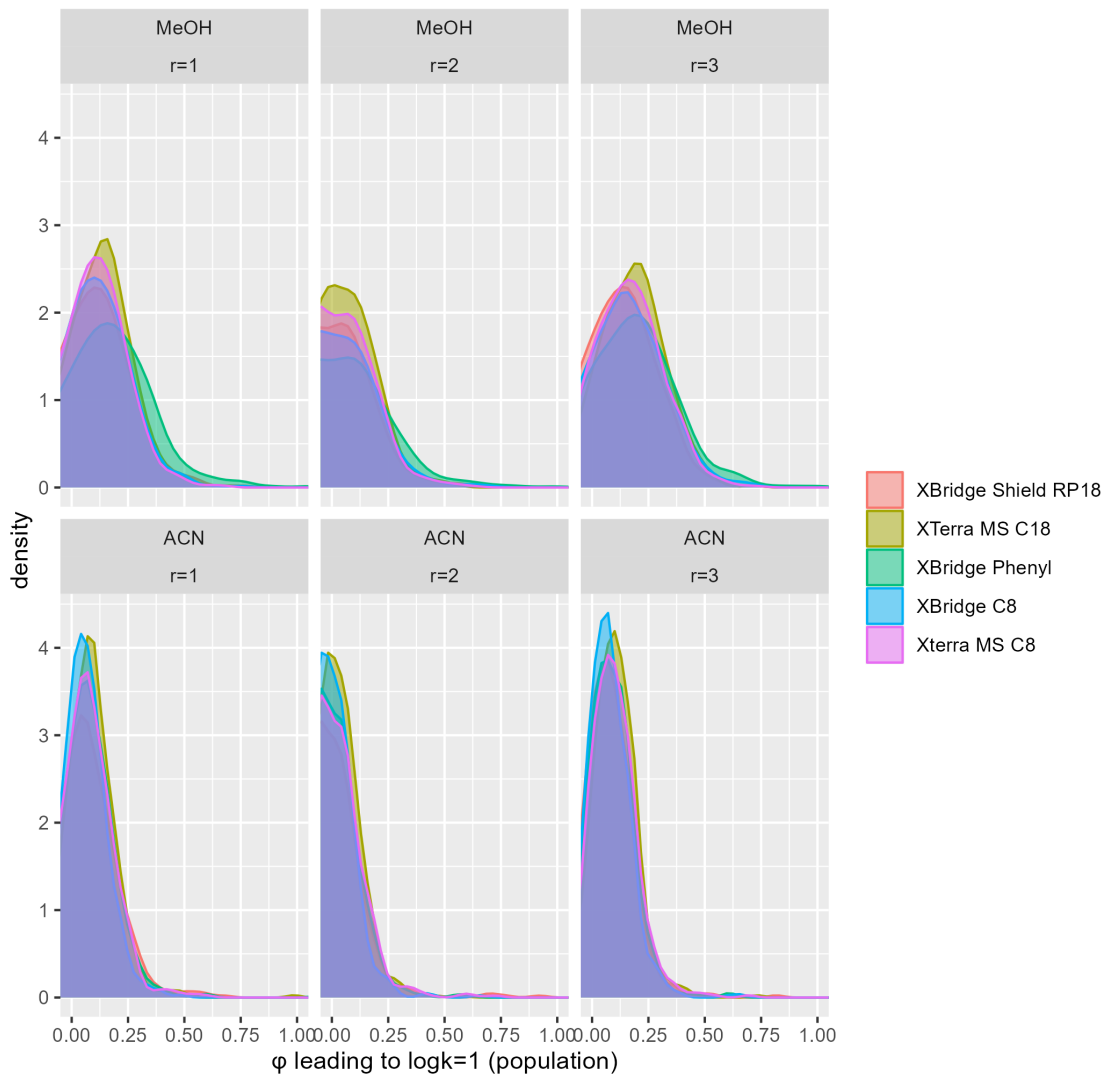
```
    sprintf("logkwxPred[%s,1,1]",idx_analyte),
    sprintf("logkwxPred[%s,1,2]",idx_analyte),
    sprintf("logkwxPred[%s,1,3]",idx_analyte),
    sprintf("logkwxPred[%s,2,1]",idx_analyte),
    sprintf("logkwxPred[%s,2,2]",idx_analyte),
    sprintf("logkwxPred[%s,2,3]",idx_analyte),
    sprintf("logkwxPred[%s,3,1]",idx_analyte),
    sprintf("logkwxPred[%s,3,2]",idx_analyte),
    sprintf("logkwxPred[%s,3,3]",idx_analyte),
    sprintf("logkwxPred[%s,4,1]",idx_analyte),
    sprintf("logkwxPred[%s,4,2]",idx_analyte),
    sprintf("logkwxPred[%s,4,3]",idx_analyte),
    sprintf("logkwxPred[%s,5,1]",idx_analyte),
    sprintf("logkwxPred[%s,5,2]",idx_analyte),
    sprintf("logkwxPred[%s,5,3]",idx_analyte),
    sprintf("S1xPred[%s,1,1,1]",idx_analyte),
    sprintf("S1xPred[%s,1,1,2]",idx_analyte),
    sprintf("S1xPred[%s,1,1,3]",idx_analyte),
    sprintf("S1xPred[%s,2,1,1]",idx_analyte),
    sprintf("S1xPred[%s,2,1,2]",idx_analyte),
    sprintf("S1xPred[%s,2,1,3]",idx_analyte),
    sprintf("S1xPred[%s,1,2,1]",idx_analyte),
    sprintf("S1xPred[%s,1,2,2]",idx_analyte),
    sprintf("S1xPred[%s,1,2,3]",idx_analyte),
    sprintf("S1xPred[%s,2,2,1]",idx_analyte),
    sprintf("S1xPred[%s,2,2,2]",idx_analyte),
    sprintf("S1xPred[%s,2,2,3]",idx_analyte),
    sprintf("S1xPred[%s,1,3,1]",idx_analyte),
    sprintf("S1xPred[%s,1,3,2]",idx_analyte),
    sprintf("S1xPred[%s,1,3,3]",idx_analyte),
    sprintf("S1xPred[%s,2,3,1]",idx_analyte),
    sprintf("S1xPred[%s,2,3,2]",idx_analyte),
    sprintf("S1xPred[%s,2,3,3]",idx_analyte),
    sprintf("S1xPred[%s,1,4,1]",idx_analyte),
    sprintf("S1xPred[%s,1,4,2]",idx_analyte),
    sprintf("S1xPred[%s,1,4,3]",idx_analyte),
    sprintf("S1xPred[%s,2,4,1]",idx_analyte),
    sprintf("S1xPred[%s,2,4,2]",idx_analyte),
    sprintf("S1xPred[%s,2,4,3]",idx_analyte),
    sprintf("S1xPred[%s,1,5,1]",idx_analyte),
    sprintf("S1xPred[%s,1,5,2]",idx_analyte),
```

```

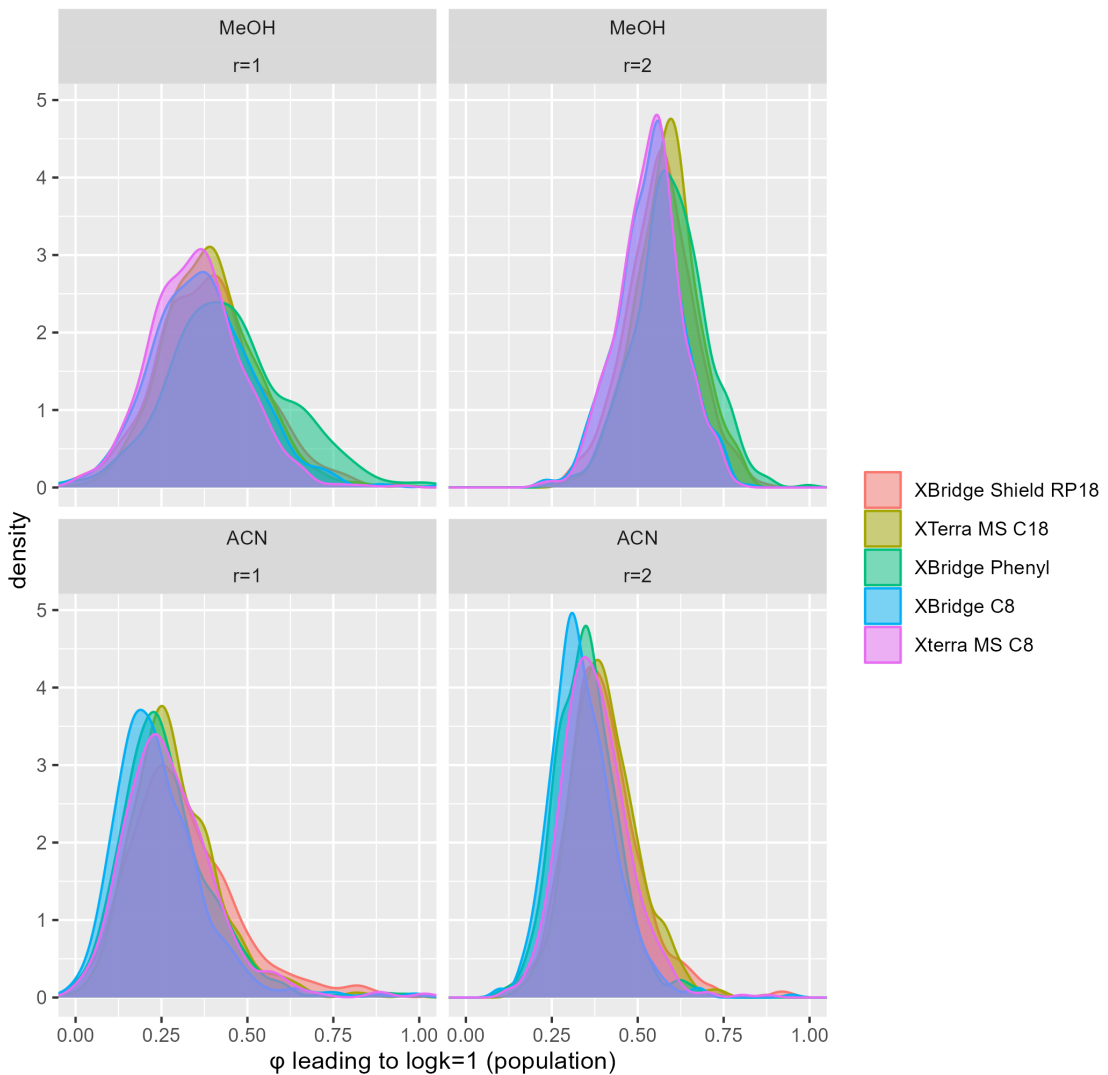
sprintf("S1xPred[%s,1,5,3]",idx_analyte),
sprintf("S1xPred[%s,2,5,1]",idx_analyte),
sprintf("S1xPred[%s,2,5,2]",idx_analyte),
sprintf("S1xPred[%s,2,5,3]",idx_analyte),
"S2xPred[1,1]",
"S2xPred[2,1]",
".draw",".iteration",".chain")]

p<-draws_df_subset %>%
  slice_sample(n=1000) %>%
  tidybayes::spread_draws(logkwxPred[, c, r], S1xPred[, m, c, r], S2xPred[m, ]) %>%
  filter(r<=R[idx_analyte]+1) %>%
  mutate(foo = (logkwxPred-1)/S1xPred/(1+S2xPred)) %>%
  mutate(fix = foo/(1-S2xPred*foo)) %>%
  ggplot(aes(x = fix, color = as.factor(c), fill = as.factor(c))) +
  geom_density(alpha = 1/2) +
  coord_cartesian(xlim=c(0,1))+
  facet_wrap(m~r, nrow = 2,labeller = labeller(m=mod.labs,r=diss.labs))+
  labs(title= paste(dataNames>Name[analyte_ID_sample[i]]),color= " ",fill= " ", x = "\u03c6",
  scale_fill_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl",
  scale_color_discrete(labels= c("XBridge Shield RP18","XTerra MS C18", "XBridge Phenyl",
  facet_wrap(m~r, nrow = 2,labeller = labeller(m=mod.labs,r=diss.labs))
  print(p)
  ggsave(paste0("figures\\izoparam\\", paste(dataNames>Name[analyte_ID_sample[i]]), "filogk"
}
```

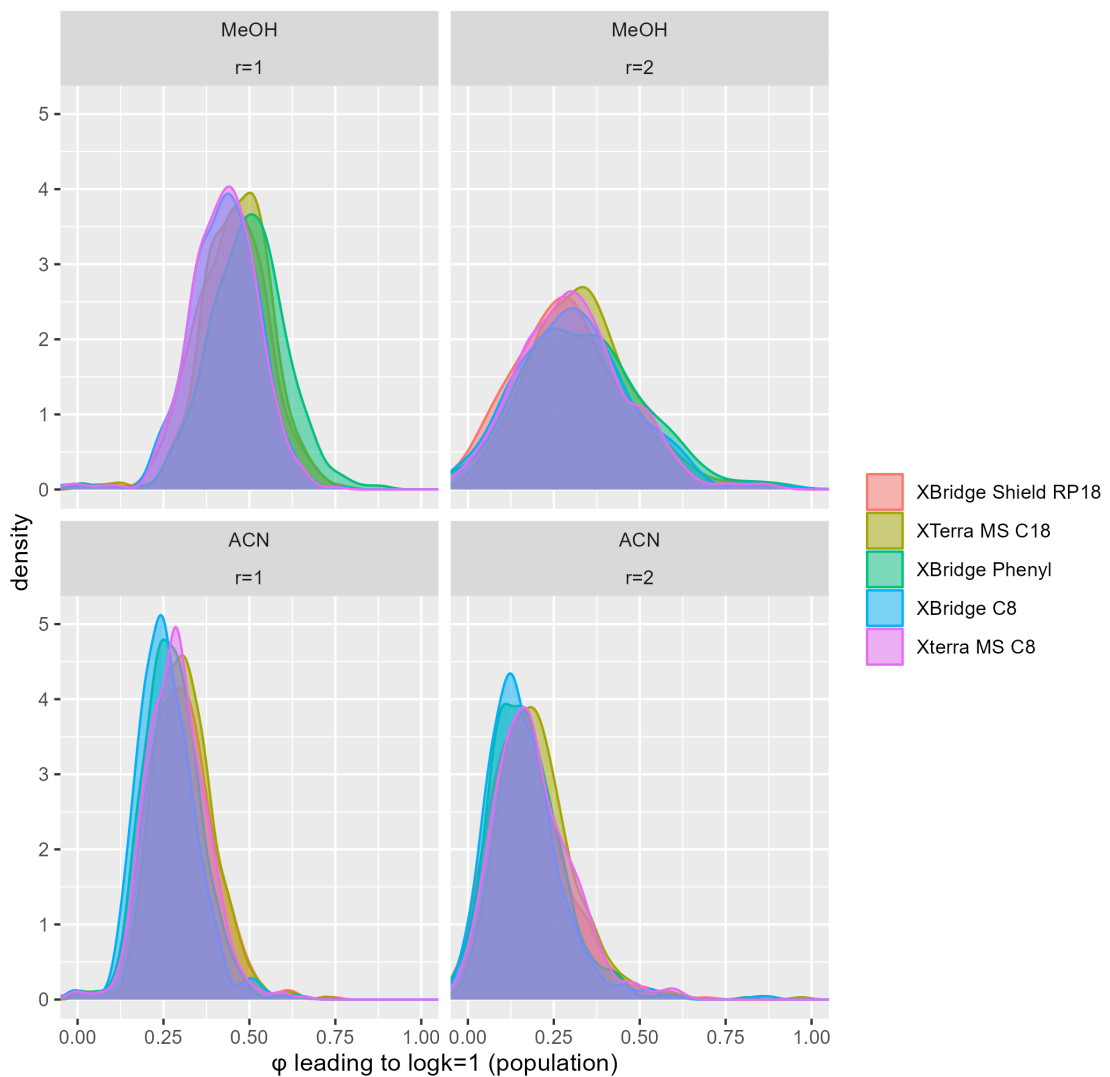
Baclofen



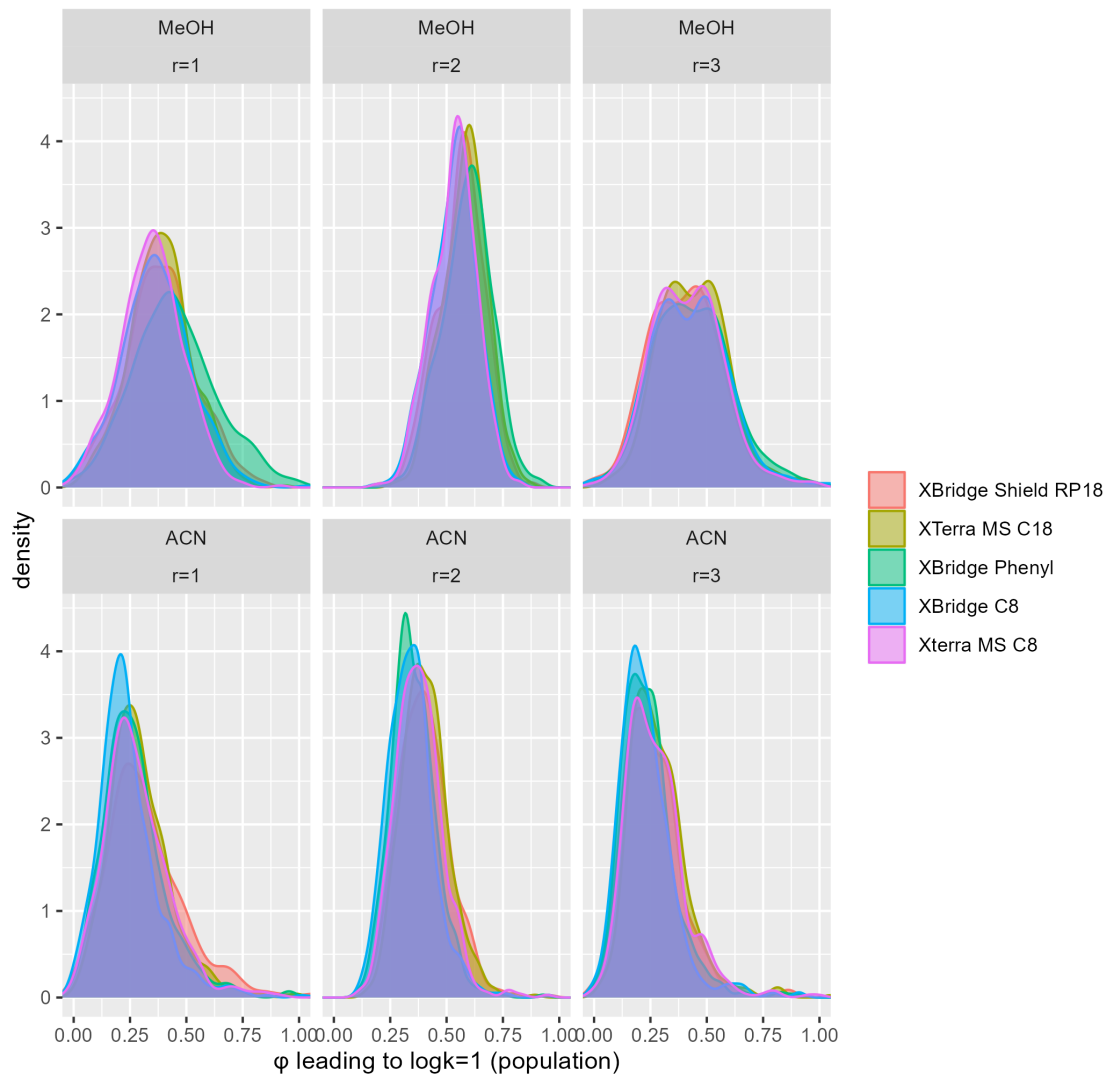
Acridine



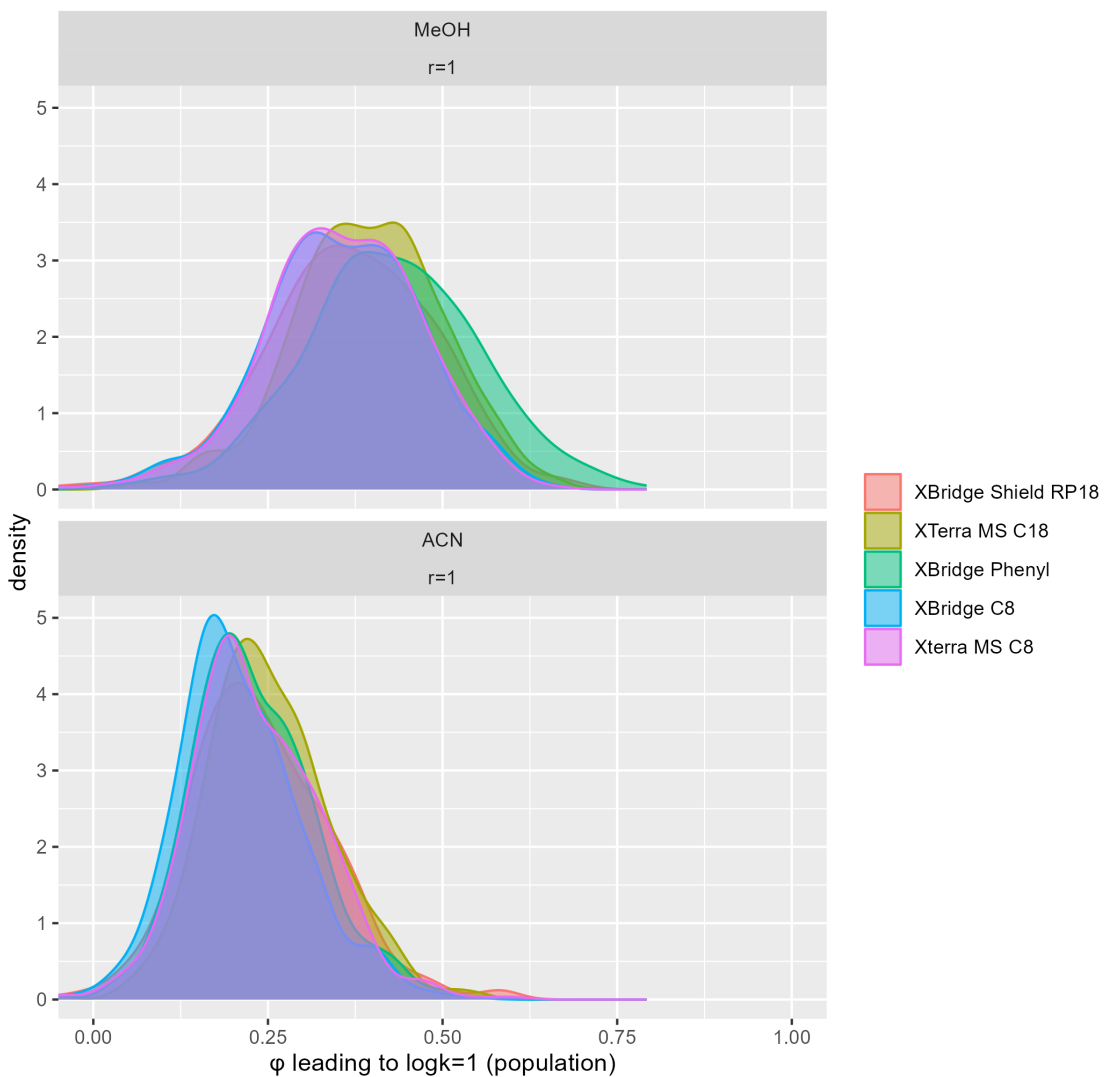
Tolbutamide

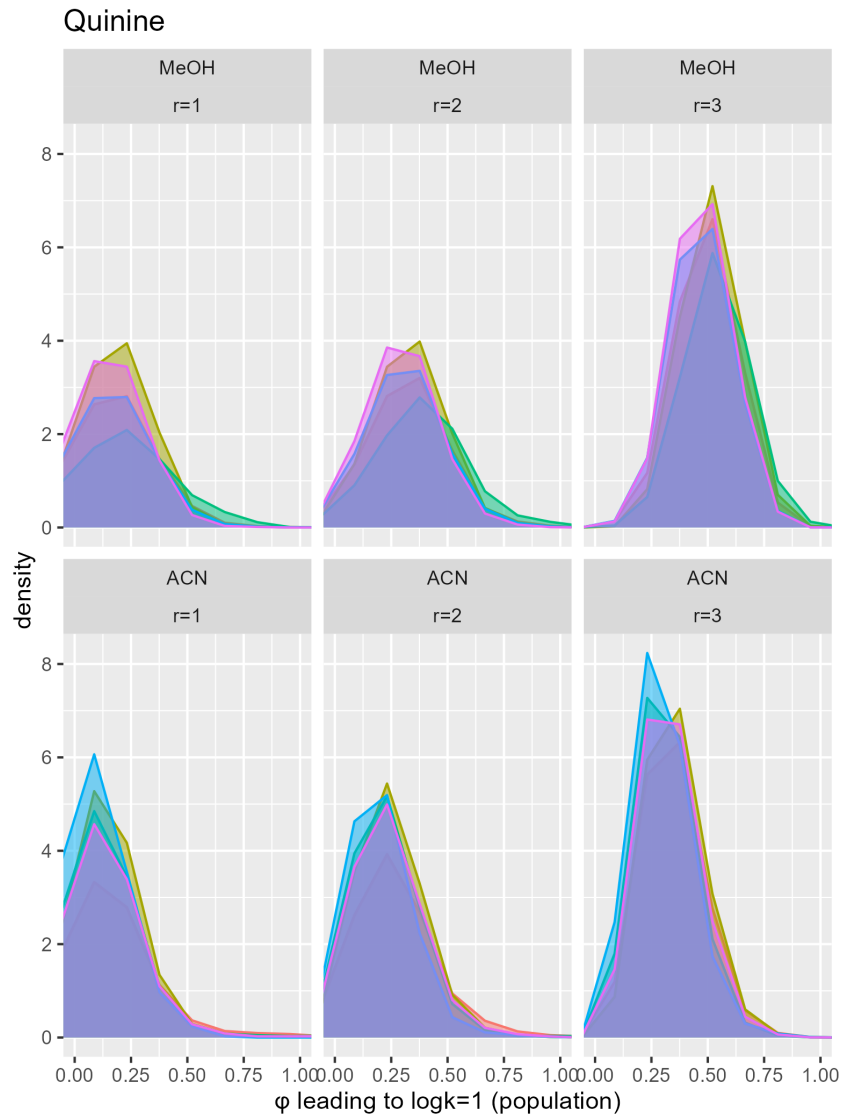


Pioglitazone



Hydrocortisone





9 Effect of logP

The following plot show the effect of logP on isocratic retention times (population predictions) for typical neutral, basic and acidic analyte. The uncertainties are large. They are not shown to improve visibility.

```
model_logP <- cmdstan_model("stan/hplc-gra-fivecolumns-logP.stan")
```



```

        sprintf("logkwxPred[%s,5,3]",idx_analyte),
        sprintf("S1xPred[%s,1,1,1]",idx_analyte),
        sprintf("S1xPred[%s,1,1,2]",idx_analyte),
        sprintf("S1xPred[%s,1,1,3]",idx_analyte),
        sprintf("S1xPred[%s,2,1,1]",idx_analyte),
        sprintf("S1xPred[%s,2,1,2]",idx_analyte),
        sprintf("S1xPred[%s,2,1,3]",idx_analyte),
        sprintf("S1xPred[%s,1,2,1]",idx_analyte),
        sprintf("S1xPred[%s,1,2,2]",idx_analyte),
        sprintf("S1xPred[%s,1,2,3]",idx_analyte),
        sprintf("S1xPred[%s,2,2,1]",idx_analyte),
        sprintf("S1xPred[%s,2,2,2]",idx_analyte),
        sprintf("S1xPred[%s,2,2,3]",idx_analyte),
        sprintf("S1xPred[%s,1,3,1]",idx_analyte),
        sprintf("S1xPred[%s,1,3,2]",idx_analyte),
        sprintf("S1xPred[%s,1,3,3]",idx_analyte),
        sprintf("S1xPred[%s,2,3,1]",idx_analyte),
        sprintf("S1xPred[%s,2,3,2]",idx_analyte),
        sprintf("S1xPred[%s,2,3,3]",idx_analyte),
        sprintf("S1xPred[%s,1,4,1]",idx_analyte),
        sprintf("S1xPred[%s,1,4,2]",idx_analyte),
        sprintf("S1xPred[%s,1,4,3]",idx_analyte),
        sprintf("S1xPred[%s,2,4,1]",idx_analyte),
        sprintf("S1xPred[%s,2,4,2]",idx_analyte),
        sprintf("S1xPred[%s,2,4,3]",idx_analyte),
        sprintf("S1xPred[%s,1,5,1]",idx_analyte),
        sprintf("S1xPred[%s,1,5,2]",idx_analyte),
        sprintf("S1xPred[%s,1,5,3]",idx_analyte),
        sprintf("S1xPred[%s,2,5,1]",idx_analyte),
        sprintf("S1xPred[%s,2,5,2]",idx_analyte),
        sprintf("S1xPred[%s,2,5,3]",idx_analyte),
        "S2xPred[1,1]",
        "S2xPred[2,1]",
        ".draw",".iteration",".chain"))]

}

draws_df_subset<-cbind(mydata[[1]][,1:45],mydata[[2]][,1:45],mydata[[3]])
```

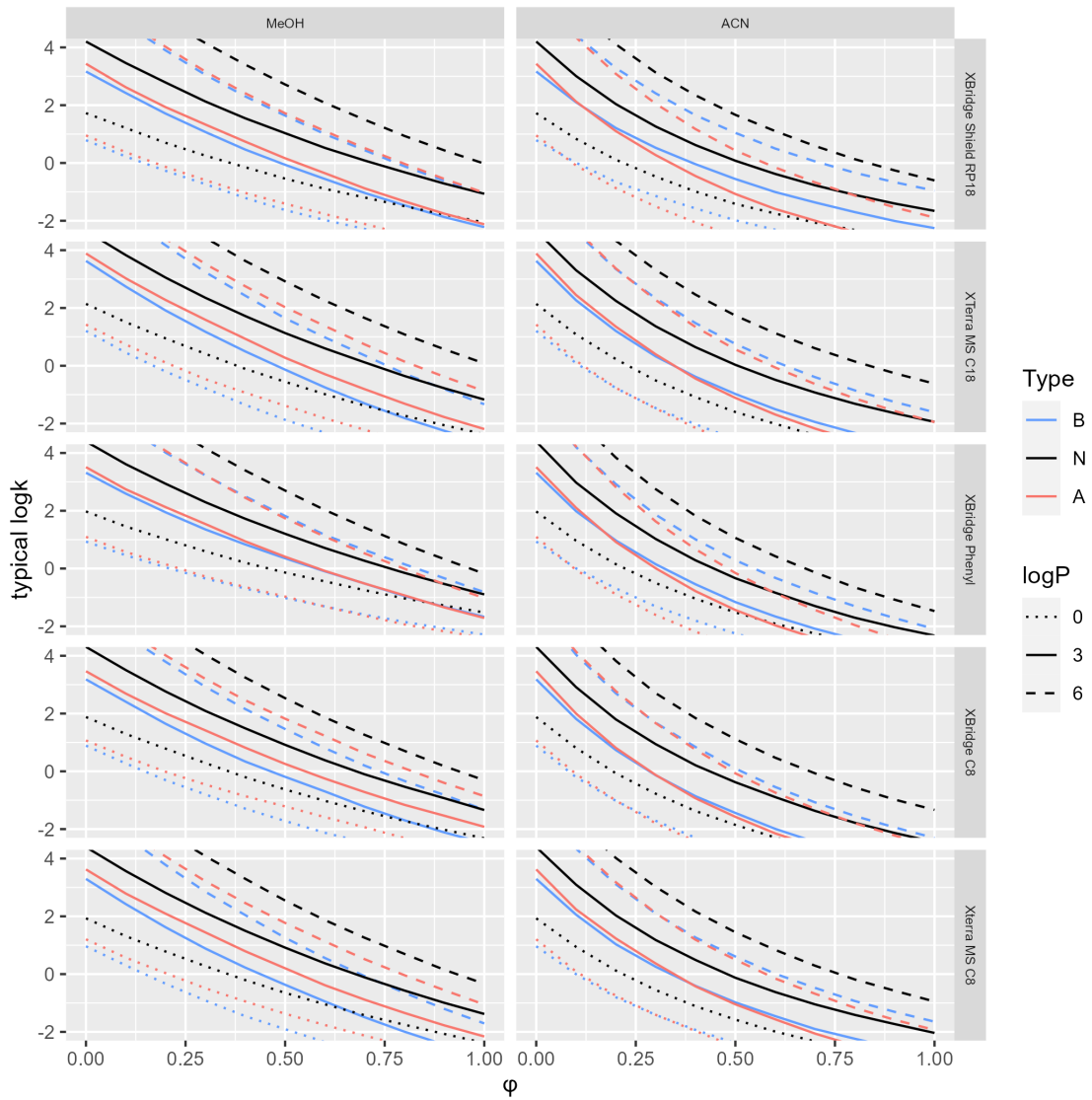
p<-draws_df_subset %>%
 slice_sample(n=500) %>%

```

tidybayes::spread_draws(logkwxPred[p, c, r], S1xPred[p, m, c, r], S2xPred[m, ]) %>%
tidyr::expand_grid(fi = seq(0,1,0.1)) %>%
mutate(logkPred = logkwxPred-S1xPred*(1+S2xPred)*fi/(1+S2xPred*fi)) %>%
ungroup() %>%
group_by(p,m,c,r,fi) %>%
summarise(mlogkPred=median(logkPred)) %>%
ggplot(aes(x = fi, y = mlogkPred, color = as.factor(r), fill = as.factor(r), linetype =
geom_line() +
facet_grid(c~m,labeller = labeller(c=col.labs, m=mod.labs))+
labs(color= "Type", linetype= "logP", y="typical logk",x="\u03c6")+
scale_fill_discrete(labels= c("B", "N", "A")) +
scale_color_discrete(labels= c("B", "N", "A"))+
scale_color_manual(labels = c("B", "N", "A"), values = c("#619cff", "black", "#f8766d"))+
scale_linetype_manual(labels= c("0", "3", "6"), values = c("dotted","solid", "dashed"))+
theme(strip.text.x = element_text(size = 6),strip.text.y = element_text(size = 6))+
coord_cartesian(xlim=c(0,1),ylim=c(-2,4))

print(p)

```



```
ggsave(paste0("figures\\logPeffects\\logPjoined", ".png"), plot=p, width = 20, height = 20)
```

10 Goodness of fit plots

Several goodness of fit plots were used to describe how well the model fits our set of observations.

```

model_restr <- cmdstan_model("stan/hplc-gra-fivecolumns-residuals.stan")

fit_restr <- model_restr$generate_quantities(fitsim,
                                              data = datastruct,
                                              seed = 123,
                                              parallel_chains = 4,
                                              output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-residuals-202308082123-1'
))

draws_restr_df <- as_draws_df(x$generated_quantities)

# draws_restr_df <- fit_restr$draws(format = "df")

trCond <- apply(draws_restr_df[,which(colnames(draws_restr_df) %in% grep("^trHatCond", names(draws_restr_df)))], 1, mean)

trPred <- apply(draws_restr_df[,which(colnames(draws_restr_df) %in% grep("^trHatPred", names(draws_restr_df)))], 1, mean)

restrCond <- apply(draws_restr_df[,which(colnames(draws_restr_df) %in% grep("^restrCond", names(draws_restr_df)))], 1, mean)

restrPred <- apply(draws_restr_df[,which(colnames(draws_restr_df) %in% grep("^restrPred", names(draws_restr_df)))], 1, mean)

data$trCond = trCond
data$trPred = trPred
data$restrCond = restrCond
data$restrPred = restrPred

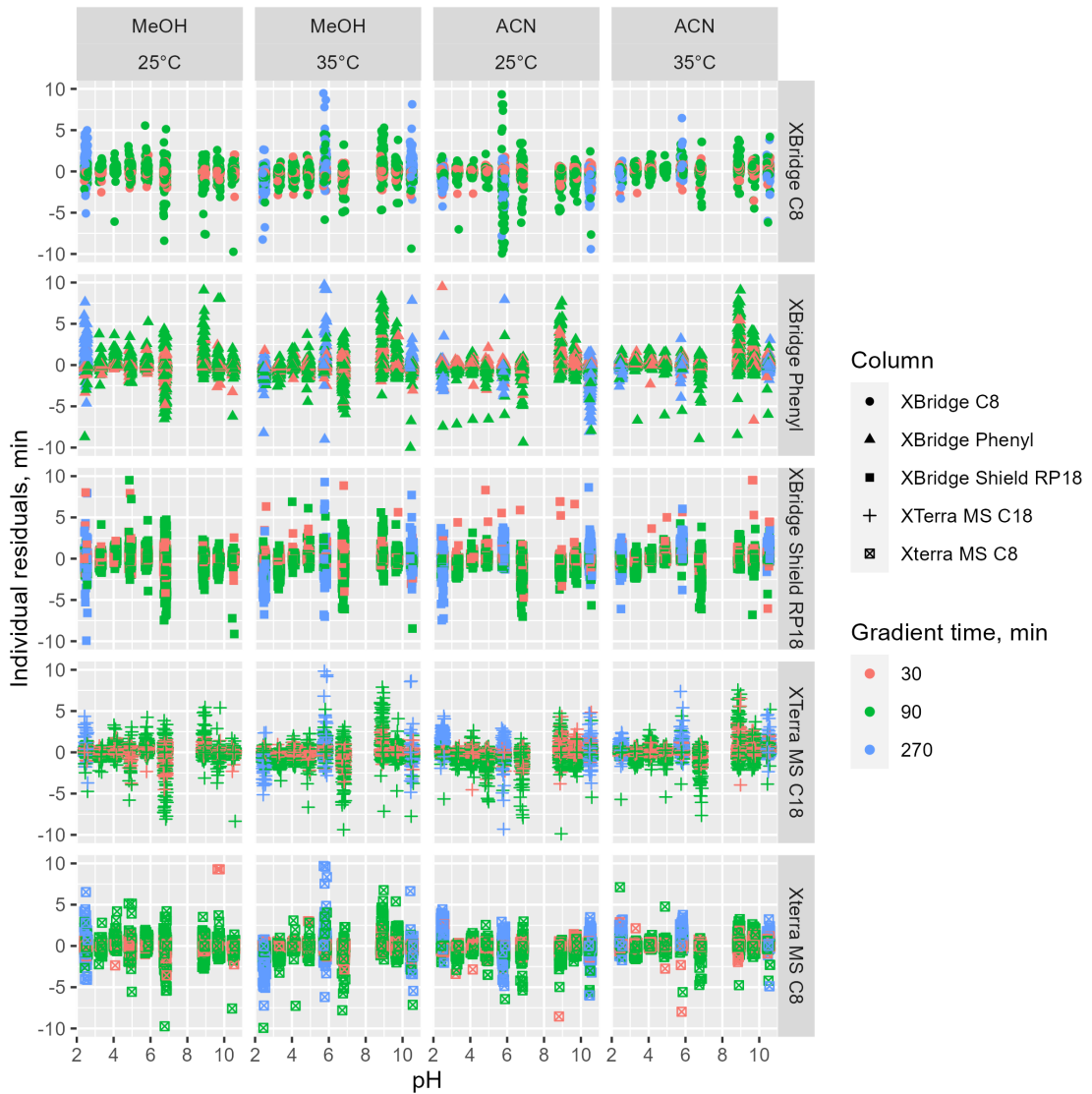
```

Residuals:

```

p1 <- ggplot()+
  geom_jitter(data=data, aes(x = pHs, y = restrCond, color = as.factor(tg), shape=as.factor(Mod2)),
  facet_grid(ColumnName~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+
  labs(x ="pH", y = "Individual residuals, min", color = "Gradient time, min", shape = "Mod2", size = 3)
  print(p1)

```



```
ggsave(paste0("figures\\concordanceplots\\", "residuals", ".png"), plot=p1, width = 20, he
```

Observed vs. population and individual predictions:

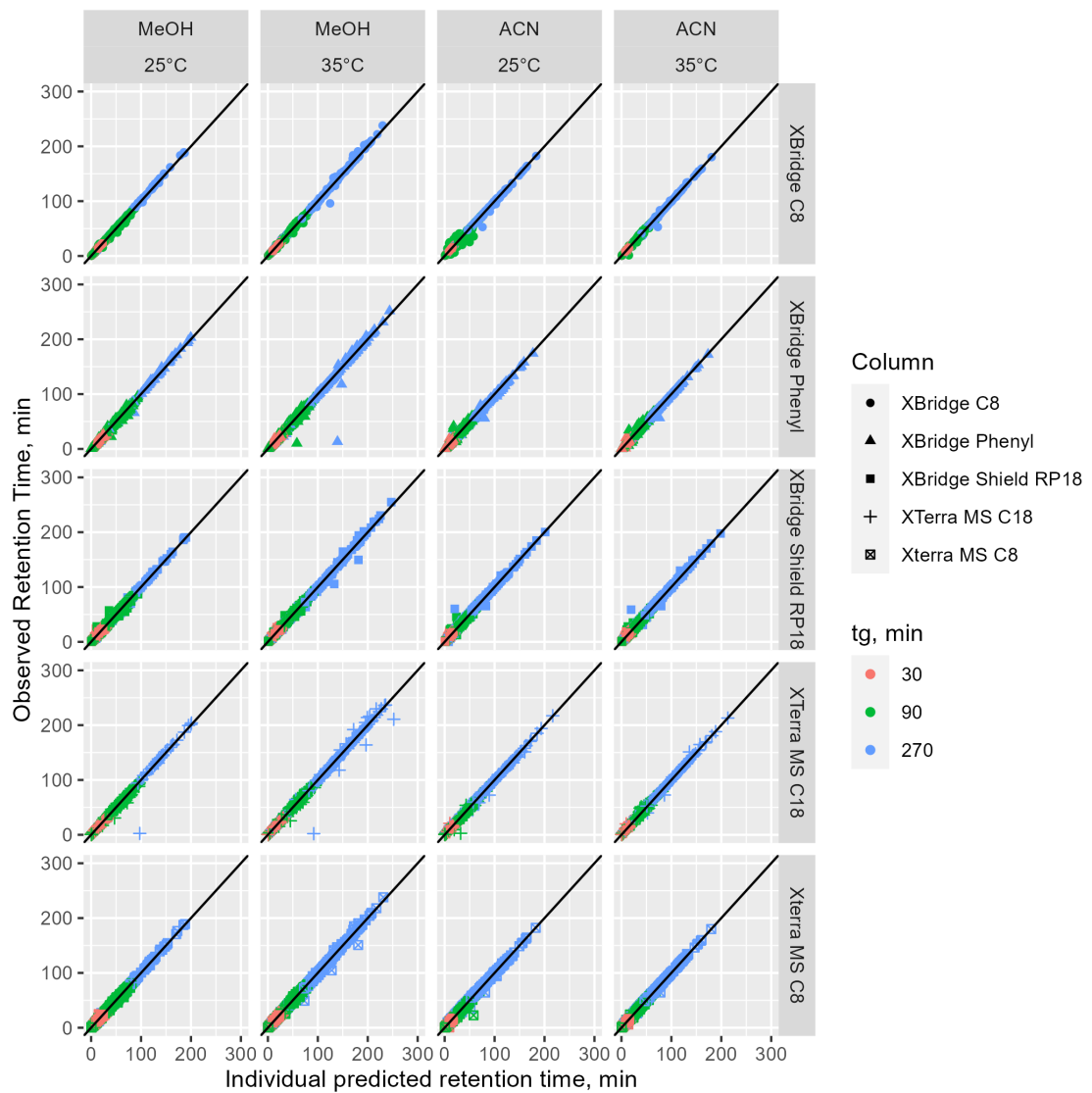
```
p1 <- ggplot()+
  geom_point(data=data, aes(x = trCond, y = RT, color = as.factor(tg), shape=as.factor(
  facet_grid(ColumnName~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+
  labs(y ="Observed Retention Time, min",
  x = "Individual predicted retention time, min",
```

```

      color = "tg, min",
      shape = 'Column')+
  xlim(0,300) +
  ylim(0,300) +
  geom_abline(intercept = 0, slope = 1)

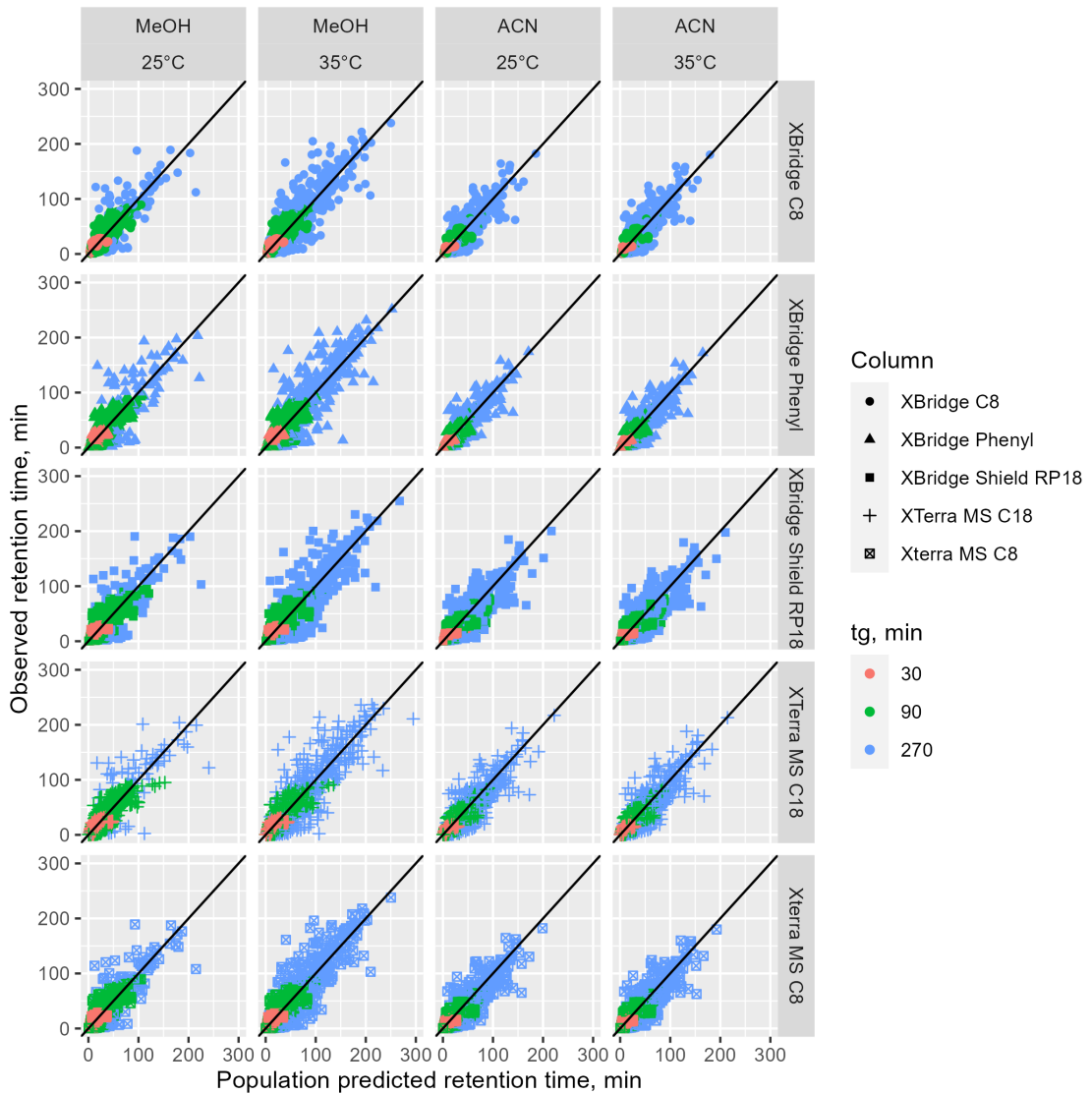
print(p1)

```



```
p2 <- ggplot()+
  geom_point(data=data, aes(x = trPred, y = RT, color = as.factor(tg), shape=as.factor(ColumnN
  facet_grid(ColumnName~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+
  labs(y ="Observed retention time, min",
       x = "Population predicted retention time, min",
       color = "tg, min",
       shape = 'Column')+  
  xlim(0,300) +
  ylim(0,300) +
  geom_abline(intercept = 0, slope = 1)

print(p2)
```



```
ggsave(paste0("figures\\concordanceplots\\", "individual", ".png"), plot=p1, width = 20, height = 15)
ggsave(paste0("figures\\concordanceplots\\", "population", ".png"), plot=p2, width = 20, height = 15)
```

11 Predictions and decision making. Case 1

Predicting best chromatogram using all the available data.

```

fit_red2<- cmdstanr::as_cmdstan_fit(c(
    # 'stanfiles/output_1.csv',
    # 'stanfiles/output_2.csv',
    # 'stanfiles/output_3.csv',
    # 'stanfiles/output_4.csv',
    'stanfiles/output_5.csv',
    'stanfiles/output_6.csv',
    'stanfiles/output_7.csv',
    'stanfiles/output_8.csv'
))

```

11.1 Utility maps

Let's find the best chromatogram that maximize utility using the set of experiments used to build the model.

The utility function is based on lowest retention time, highest retention time and the difference of retention times between the critical pair of analytes. This utility is zero if at least one of the analyte has retention higher that 40, less than 2 min or the difference in retention time is less than 2. Otherwise, it favors shorter runs. (.) denotes design variables.

$$U(.) = I((\min tr(.) > 2) \& (\max tr(.) > 2)) \cdot (40 - \max tr(.)) \cdot I(\max tr(.) < 40)$$

$$I(x) = 1 \text{ if condition } x \text{ is true} \\ I(x) = 0 \text{ if condition } x \text{ is false}$$

```

upH <- datasim %>%
  select(1:20) %>%
  distinct(Temp,Mod, pHo, alpha1, alpha2) %>%
  group_by(Temp,Mod) %>%
  tidyr::complete(pHo = seq(min(pHo), max(pHo), len = 17)) %>%
  arrange(Temp,Mod,pHo) %>%
  mutate(alpha1 = zoo::na.approx(alpha1,pHo)) %>%
  mutate(alpha2 = zoo::na.approx(alpha2,pHo)) %>%
  group_by(Temp,Mod) %>%
  mutate(pHid = row_number()) %>%
  group_by(pHid) %>%
  mutate(pHs = round(mean(pHo),2)) %>%
  ungroup()

datasim2 <- datasim %>%

```

```

  select(1:20) %>%
  filter(tg==30, Temp==25) %>%
  select(-tg, -fio, -expid, -pHs, -pHo, -alpha1, -alpha2, -pH) %>%
  distinct() %>%
tidyrr::expand_grid(fio = seq(0.05,0.2,0.05),tg = seq(20,260,20), pHs=unique(upH$pHs))%>%
  left_join(upH, by = join_by(pHs, Mod, Temp), relationship = "many-to-one") %>%
  group_by(tg, fio, Mod, Column, pHs) %>%
  mutate(expid = cur_group_id()) %>%
  ungroup()

datastructsim2 <-datastruct
datastructsim2$nObs=length(datasim2$METID)
datastructsim2$analyte=match(datasim2$METID, unique(datasim2$METID))
datastructsim2$modifier=match(datasim2$Mod2, sort(unique(datasim2$Mod2)))
datastructsim2$steps=4*(2-datasim2$Mod2) + 10*(datasim2$Mod2-1)
datastructsim2$column=match(datasim2$Column, unique(datasim2$Column))
datastructsim2$hplcparam=cbind(datasim2$tg,datasim2$td,datasim2$to,datasim2$te,
                                datasim2$fio,datasim2$fik,datasim2$Mod2-1,datasim2$pHo,
                                datasim2$alpha1,datasim2$alpha2,(datasim2$Temp-25)/10,
                                datasim2$Column-1)
datastructsim2$trobs = rep(0,datastructsim2$nObs)
analyte_ID_sample = c(9,17,33,58,140,180)
idx <- which(datasim2$METID %in% analyte_ID_sample)
datasim_red = datasim2[idx,]
datastructsim_red <-datastructsim2
datastructsim_red$nObs=length(datastructsim_red$analyte[idx])
datastructsim_red$analyte=datastructsim_red$analyte[idx]
datastructsim_red$modifier=datastructsim_red$modifier[idx]
datastructsim_red$steps=datastructsim_red$steps[idx]
datastructsim_red$column=datastructsim_red$column[idx]
datastructsim_red$hplcparam=datastructsim_red$hplcparam[idx,]
datastructsim_red$trobs = rep(0,datastructsim_red$nObs)
datastructsim_red$nexpid=length(unique(datasim_red$expid))
datastructsim_red$expid=match(datasim_red$expid, unique(datasim_red$expid))
datastructsim_red$nAnalytessim=length(unique(datastructsim_red$analyte))
datastructsim_red$analytesim=match(datastructsim_red$analyte, unique(datastructsim_red$analyte))

model_sim_red <- cmdstan_model("stan/hplc-gra-fivecolumns-fixed-sim.stan")

fit_sim_red2 <- model_sim_red$generate_quantities(fit_red2,
                                                    data = datastructsim_red,

```

```

          seed = 123,
          parallel_chains = 4,
          output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-fixed-sim-202308090855-1
))

draws_sim_red_df <- as_draws_df(x$generated_quantities)

#draws_sim_red_df <- fit_sim_red$draws(format = "df")

foo<-datasim_red[!duplicated(datasim_red$expid),]
x<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("^mindifftr", names(draws_
y<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("^maxtr", names(draws_sim_
z<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("^mintr", names(draws_sim_


u=x;

for (i in 1:ncol(x)){u[,i] <- as.numeric(x[,i]>2 & z[,i] > 2) * (40-y[,i]) * as.numeric(y[

pr <- apply(u, MARGIN = 2, FUN = mean)
pr<-as.data.frame(pr)

foo$EUutility=pr$pr

x <- apply(x, MARGIN = 2, FUN = quantile, probs = c(.05,.5,.95))
x<-as.data.frame(t(x))

foo$mindifftr_l=x$`5%
foo$mindifftr_m=x$`50%
foo$mindifftr_h=x$`95%

y <- apply(y, MARGIN = 2, FUN = quantile, probs = c(.05,.5,.95))
y<-as.data.frame(t(y))

foo$maxtr_l=y$`5%
foo$maxtr_m=y$`50%
foo$maxtr_h=y$`95%

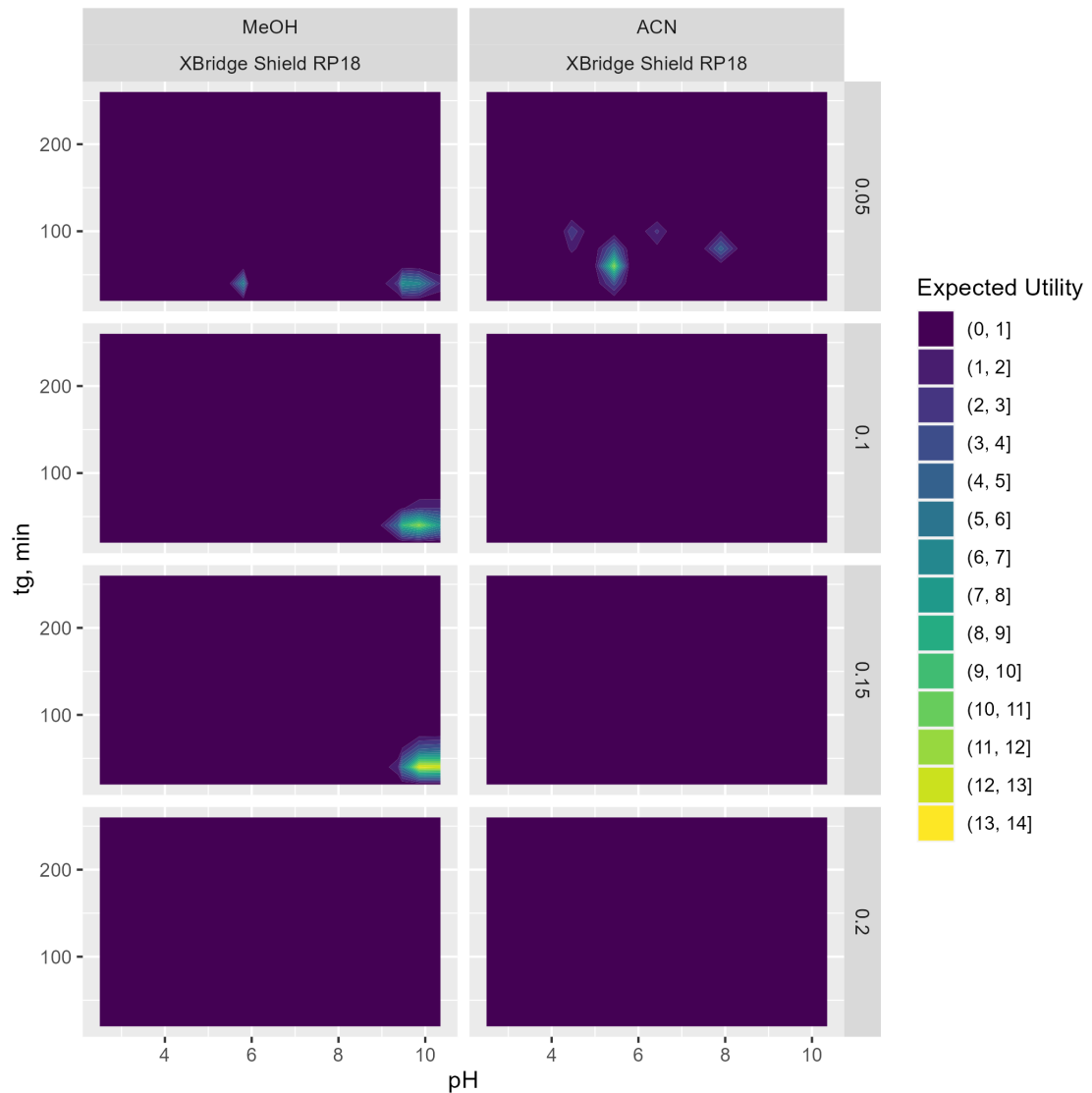
```

```

p11 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==1), aes(x = pHs, y=tg, z = EUtility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p11)

```

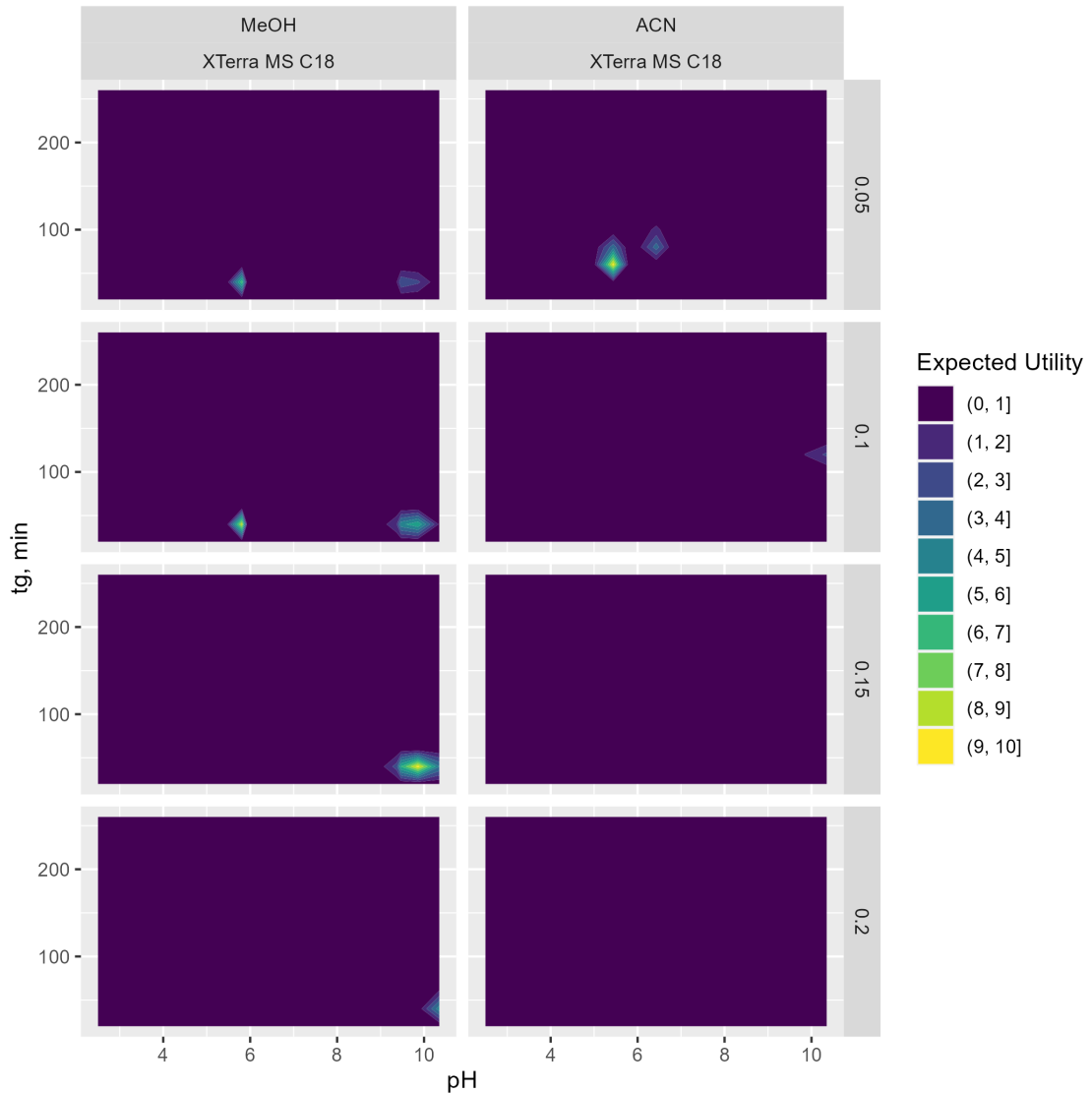


```

p12 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==2), aes(x = pHs, y=tg, z = EUtility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p12)

```

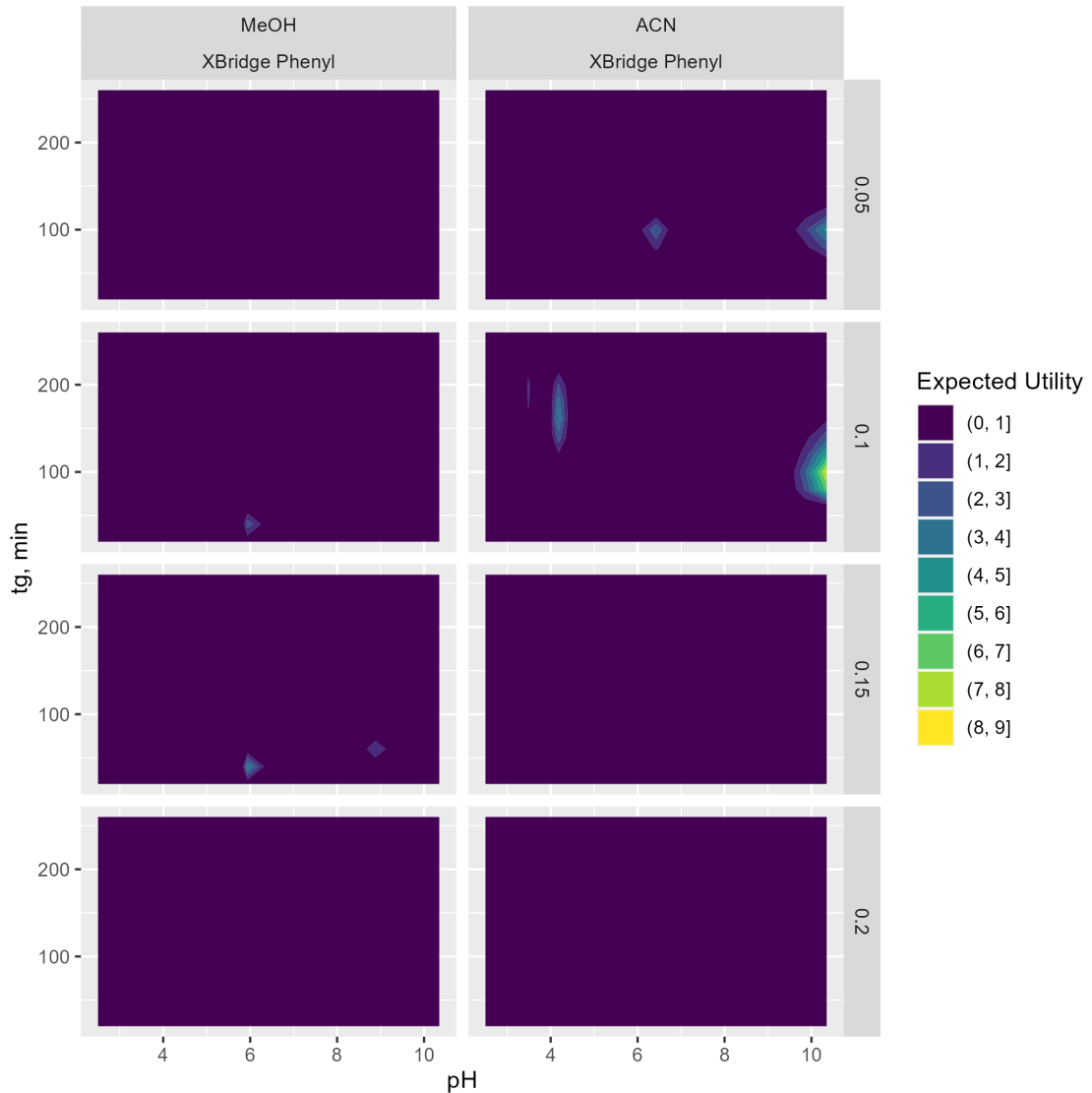


```

p13 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==3), aes(x = pHs, y=tg, z = EUtility))+ 
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p13)

```

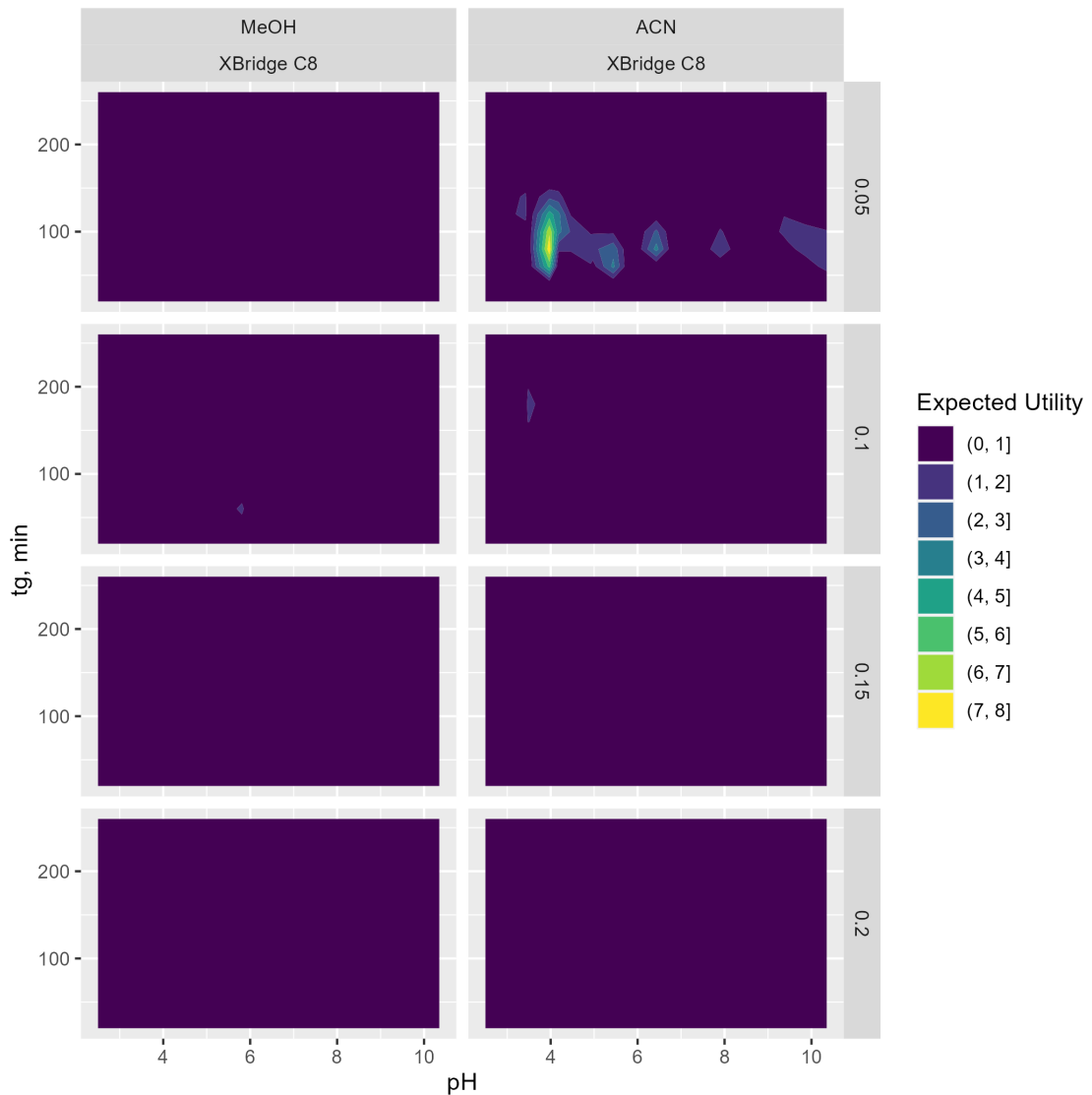


```

p14 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==4), aes(x = pHs, y=tg, z = EUtility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p14)

```

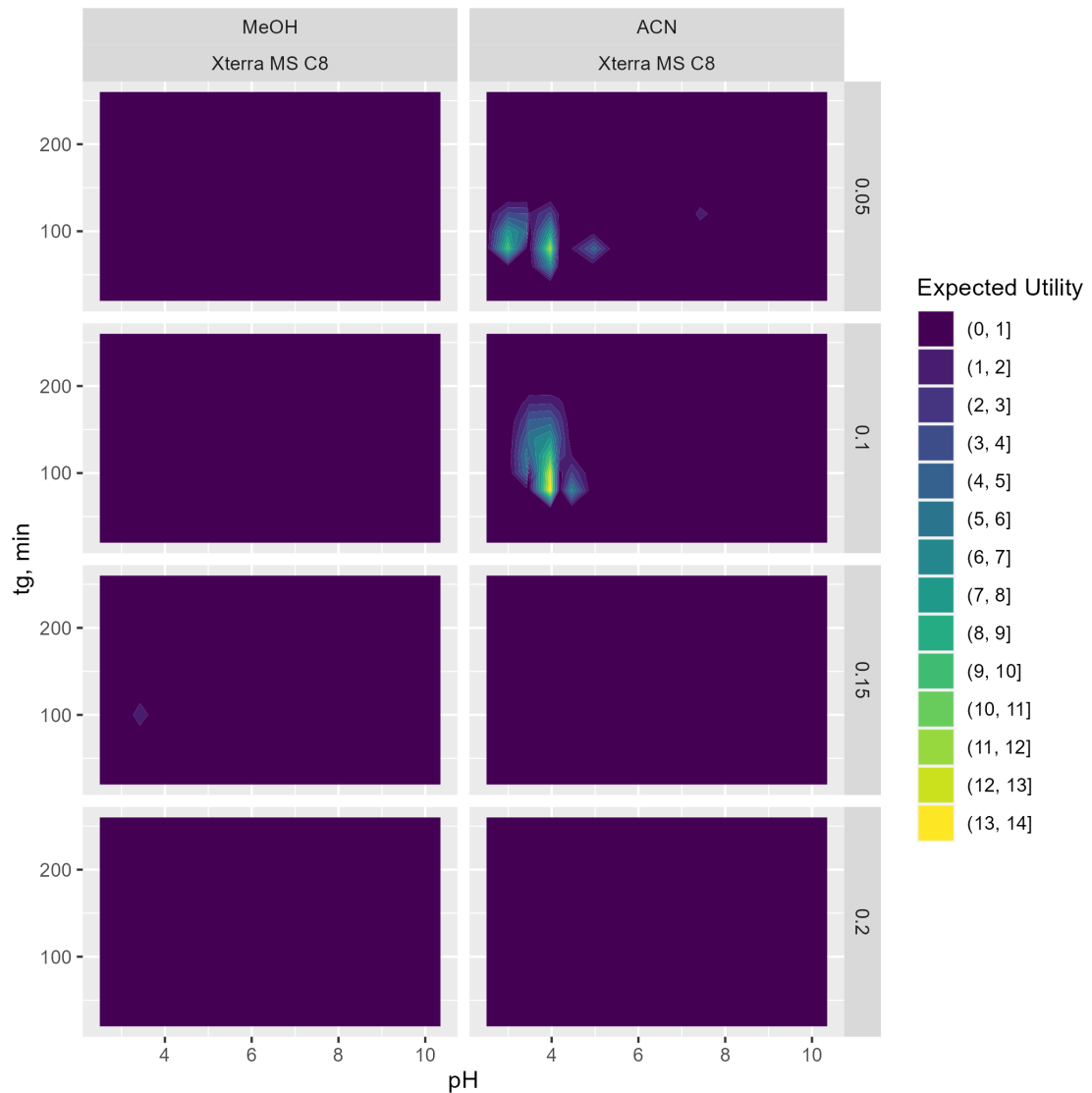


```

p15 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==5), aes(x = pHs, y=tg, z = EUtility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p15)

```



```

ggsave(paste0("figures\\casestudy1\\utilitymap\\", "utilitymap1", ".png"), plot=p11, width=10, height=10)
ggsave(paste0("figures\\casestudy1\\utilitymap\\", "utilitymap2", ".png"), plot=p12, width=10, height=10)
ggsave(paste0("figures\\casestudy1\\utilitymap\\", "utilitymap3", ".png"), plot=p13, width=10, height=10)
ggsave(paste0("figures\\casestudy1\\utilitymap\\", "utilitymap4", ".png"), plot=p14, width=10, height=10)
ggsave(paste0("figures\\casestudy1\\utilitymap\\", "utilitymap5", ".png"), plot=p15, width=10, height=10)

# The best for XBridge Shield RP18

foo1<-foo[foo$Column==1 & foo$pHo<7,];
foo1[which(foo1$EUtility==max(foo1$EUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo','Temp','ColumnName','EUtility')]
# A tibble: 1 x 9
#>   tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUtility
#>   <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
#> 1    60    0.05    0.8 ACN       10    5.50     25 XBridge Shield RP18     11.4

# The best choice for XTerra MS C18

foo2<-foo[foo$Column==2& foo$pHo<7,];
foo2[which(foo2$EUtility==max(foo2$EUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo','Temp','ColumnName','EUtility')]
# A tibble: 1 x 9
#>   tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUtility
#>   <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
#> 1    60    0.05    0.8 ACN       10    5.50     25 XTerra MS C18     9.50

# The best choice for XBridge Phenyl

foo3<-foo[foo$Column==3& foo$pHo<7,];
foo3[which(foo3$EUtility==max(foo3$EUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo','Temp','ColumnName','EUtility')]
# A tibble: 1 x 9
#>   tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUtility
#>   <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
#> 1    40    0.15    0.8 MeOH      12    6.00     25 XBridge Phenyl    4.08

```

```

# The best choice for XBridge C8

foo4<-foo[foo$Column==4& foo$pHo<7,];
foo4[which(foo4$EUtility==max(foo4$EUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo','Temp','ColumnName','EUtility')]

# A tibble: 1 x 9
#>   tg    fio    fik Mod    pHid    pHo  Temp ColumnName EUtility
#>   <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>      <dbl>
1     80    0.05   0.8 ACN       5   4.00    25 XBridge C8      7.73

# The best choice for Xterra MS C8

foo5<-foo[foo$Column==5& foo$pHo<7,];
foo5[which(foo5$EUtility==max(foo5$EUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo','Temp','ColumnName','EUtility')]

# A tibble: 1 x 9
#>   tg    fio    fik Mod    pHid    pHo  Temp ColumnName EUtility
#>   <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>      <dbl>
1     80    0.1    0.8 ACN       5   4.00    25 Xterra MS C8     13.7

```

11.1.1 Uncertainty chromatogram

Let's visualize chromatograms with the highest expected utility

```

analyte_ID_sample <-c(9,17,33,58,140,180)
col.labs <- c("XBridge Shield RP18", "XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xter
datasim_red$fio = as.factor(datasim_red$fio)

wpCond= data.frame()

for (i in 1:5) {
  idx <- which(datasim_red$METID %in% analyte_ID_sample &
    datasim_red$tg == 60 & # c(30, 90, 270)
    datasim_red$pHid == 10 & # c(1:9)
    datasim_red$fio == 0.05 & # ...
    datasim_red$Column == i & # c(1, 2)
    datasim_red$Mod2 == 2 & # c(1, 2)
    datasim_red$Temp == 25) # c(25, 35)

```

```

if (i==3){

  idx <- which(datasim_red$METID %in% analyte_ID_sample &
    datasim_red$tg == 60 & # c(30, 90, 270)
    datasim_red$pHid == 6 & # c(1:9)
    datasim_red$ffio == 0.1 & # ...
    datasim_red$Column == i & # c(1, 2)
    datasim_red$Mod2 == 2 & # c(1, 2)
    datasim_red$Temp == 25) # c(25, 35)

}

if (i==4){

  idx <- which(datasim_red$METID %in% analyte_ID_sample &
    datasim_red$tg == 80 & # c(30, 90, 270)
    datasim_red$pHid == 5 & # c(1:9)
    datasim_red$ffio == 0.05 & # ...
    datasim_red$Column == i & # c(1, 2)
    datasim_red$Mod2 == 2 & # c(1, 2)
    datasim_red$Temp == 25) # c(25, 35)

}

if (i==5){

  idx <- which(datasim_red$METID %in% analyte_ID_sample &
    datasim_red$tg == 80 & # c(30, 90, 270)
    datasim_red$pHid == 5 & # c(1:9)
    datasim_red$ffio == 0.1 & # ...
    datasim_red$Column == i & # c(1, 2)
    datasim_red$Mod2 == 2 & # c(1, 2)
    datasim_red$Temp == 25) # c(25, 35)

}

data_to_plot <- draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% paste0("trHatCond[", 1:10, "]"))]

colnames(data_to_plot) <- paste(dataNames$Name[analyte_ID_sample])
wpCond1 <- melt(data_to_plot)
wpCond1$Column <- unname(col.labs[i])
wpCond= rbind(wpCond,wpCond1)

```

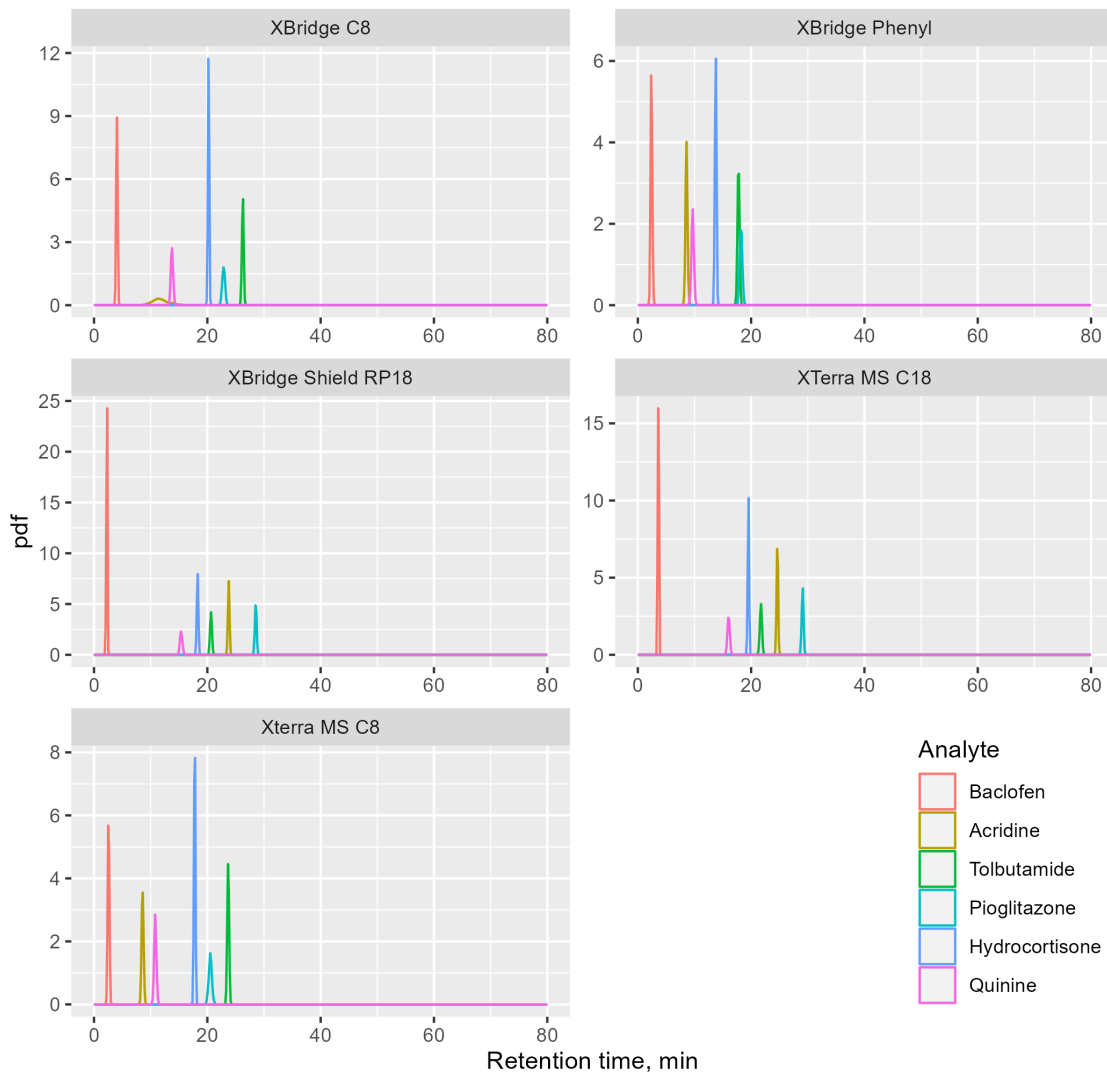
```
}

wpCond$Column=as.factor(wpCond$Column)

p <- ggplot(data = wpCond)+  
  geom_density(aes(x=value, colour=variable)) +  
  labs(title="uncertainty chromatogram (individual predictions)",  
       x ="Retention time, min",  
       y = "pdf",  
       colour="Analyte"  
       )+  
  xlim(c(0,80))+  
  facet_wrap(.~Column, nrow=3,scales = "free") +  
  theme(legend.position = c(1, 0),  
        legend.justification = c(1, 0))

print(p)
```

uncertainty chromatogram (individual predictions)



```
ggsave(paste0("figures\\casestudy1\\utilitymap\\", "chromatogram", ".png"), plot=p, width=10, height=8)
```

12 Predictions and decision making. Case 2

The proposed model can be used for decision making having access to limited data. Here we want to illustrate this by predicting Xterra MS C18, XBridge Phenyl, XBridge C8, Xterra MS C8 retention times based on XBridge Shield RP18 data.

```

analyte_ID_sample = c(9,17,33,58,140,180)

idx <- which(data$METID %in% analyte_ID_sample &
               data$tg %in% c(30, 90, 270) &
               data$pH %in% c(1:9) &
               data$Column %in% c(1) & # c(1, 2)
               data$Mod2 %in% c(1, 2) &
               data$Temp %in% c(25, 35))

datastruct_red = datastruct;
datastruct_red$idx = idx
datastruct_red$nObs2 = length(datastruct_red$idx)

init_red <- function(){
  list(logkwHat=rnorm(1,3.60,0.0793),
       S1mHat=rnorm(1,4.96,0.0832),
       dS1Hat=rnorm(1,0.612,0.0512),
       dlogkwHat=c(rnorm(1,-0.786,0.0718), rnorm(1,-0.970,0.0503)),
       dS1mHat=c(rnorm(1,0.176,0.121), rnorm(1,0.110,0.0744)),
       ddS1Hat=c(rnorm(1,0.279,0.0865), rnorm(1,-0.651,0.0564)),
       logS2mHat=rnorm(1,-0.308,0.0143),
       dlogS2Hat=rnorm(1,0.420,0.00760),
       beta=c(rnorm(1,0.831,0.0407), rnorm(1,0.483,0.0449)),
       alphamHat=c(rnorm(1,2.23,0.155), rnorm(1,-1.36,0.102)),
       dalphaHat=c(rnorm(1,0.212,0.0981), rnorm(1,-0.188,0.0744)),
       dlogkTHat=rnorm(1,-0.0898,0.00292),
       apH=c(rnorm(1,-0.0275,0.00118), rnorm(1,0.0806,0.000800)),
       omega=c(rnorm(1,0.925,0.0549), rnorm(1,0.941,0.0590), rnorm(1,0.565,0.0343)),
       omegaT=rnorm(1,0.0337,0.00212),
       kappa=c(rnorm(1,0.584,0.0350), rnorm(1,0.694,0.0475), rnorm(1,0.57,0.0378)),
       msigma=rnorm(1,0.386,0.0263),
       ssigma=rnorm(1,0.813,0.0496),
       tau=c(rnorm(1,0.882,0.0494), rnorm(1,0.958,0.0584), rnorm(1,0.793,0.0506)),
       rho = matrix(c(1, 0.864, 0.864, 1), nrow=2),
       clogkwHat=c(rnorm(1,0.424,0.0115),rnorm(1,0.173,0.0132),rnorm(1,0.105,0.0118),rnorm(1,0.16
       cS1mHat=c(rnorm(1,0.588,0.0198),rnorm(1,-0.122,0.0218), rnorm(1,0.362,0.0180), rnorm(1,0.4
       cdS1Hat=c(rnorm(1,0.151,0.0134),rnorm(1,0.828,0.0278),rnorm(1,0.552,0.0535),rnorm(1,0.0308
       cdlogkTHat=c(rnorm(1,-0.00595,0.00125),rnorm(1,-0.0208,0.00160),rnorm(1,-0.00605,0.00105),
       cdlogkwHat = matrix(c(rnorm(1,0.0412,0.0154),
       rnorm(1,-0.0515,0.0208),
       rnorm(1,-0.0321,0.0167),

```

```

rnorm(1,0.0426,0.0164),
rnorm(1,0.00532,0.0135),
rnorm(1,-0.0312,0.0154),
rnorm(1,-0.0578,0.0127),
rnorm(1,-0.0205,0.0137)),nrow=(nColumns-1)),
cdS1mHat= matrix(c(rnorm(1,-0.0240,0.0454),
rnorm(1,-0.202,0.0633),
rnorm(1,-0.501,0.0471),
rnorm(1,-0.218,0.0439),
rnorm(1,0.375,0.0352),
rnorm(1,-0.394,0.0392),
rnorm(1,-0.0862,0.0304),
rnorm(1,0.313,0.0333)),nrow=(nColumns-1)),
cddS1Hat= matrix(c(rnorm(1,0.0505,0.0269),
rnorm(1,0.134,0.0531),
rnorm(1,0.457,0.111),
rnorm(1,0.00183,0.0252),
rnorm(1,0.170,0.0154),
rnorm(1,0.573,0.033),
rnorm(1,0.504,0.0743),
rnorm(1,-0.0301,0.0157)),nrow=(nColumns-1)),
cbeta= matrix(c(rnorm(1,0.00719,0.0064),
rnorm(1,-0.00576,0.00755),
rnorm(1,-0.0241,0.00768),
rnorm(1,-0.0102,0.00588),
rnorm(1,-0.0664,0.0109),
rnorm(1,0.113,0.0126),
rnorm(1,-0.0198,0.0111),
rnorm(1,-0.00871,0.00984)),nrow=(nColumns-1)),
capH= matrix(c(rnorm(1,-0.0149,0.00157),
rnorm(1,-0.0219,0.00169),
rnorm(1,0.00482,0.00148),
rnorm(1,-0.0226,0.00146),
rnorm(1,-0.0336,0.00118),
rnorm(1,-0.0440,0.00106),
rnorm(1,-0.0506,0.000901),
rnorm(1,-0.0130,0.00103)),nrow=(nColumns-1)),
comega= abs(matrix(c(rnorm(1,0.107,0.00778),
rnorm(1,0.129,0.00891),
rnorm(1,0.114,0.00751),
rnorm(1,0.0966,0.00638),

```

```

rnorm(1,0.0971,0.0196),
rnorm(1,0.146,0.017),
rnorm(1,0.0901,0.0156),
rnorm(1,0.0713,0.0189),
rnorm(1,0.141,0.011),
rnorm(1,0.309,0.0205),
rnorm(1,0.608,0.0403),
rnorm(1,0.172,0.0123),nrow=(nColumns-1))),
ckappa= abs(matrix(c(rnorm(1,0.0688,0.00622),
rnorm(1,0.116,0.00872),
rnorm(1,0.0855,0.00668),
rnorm(1,0.0873,0.00625),
rnorm(1,0.0714,0.0362),
rnorm(1,0.199,0.0311),
rnorm(1,0.0932,0.0322),
rnorm(1,0.0334,0.0252),
rnorm(1,0.0738,0.0197),
rnorm(1,0.277,0.0253),
rnorm(1,0.717,0.0502),
rnorm(1,0.0861,0.0164),nrow=(nColumns-1))),
corr_L = matrix(c(1,0.568,0.781,0.719,0,0.819,0.129,0.176,0,0,0.604,0.541,0,0,0,0.384), n
comegaT=abs(c(rnorm(1,0.00230,0.00159),rnorm(1,0.0115,0.00123),rnorm(1,0.00179,0.00116),r
paramN = cbind(2+0.75*(logPobs-2.2), 4*matrix(1,nAnalytes,1)+0.5*(logPobs-2.2))
dS1N = matrix(0,nAnalytes,1),
dlogkT = rnorm(nAnalytes,-0.0868,0.0217),
dlogkwA = matrix(-1,nGroupsA,1),
dlogkwB = matrix(-1,nGroupsB,1),
dS1mA = matrix(0,nGroupsA,1),
dS1mB = matrix(0,nGroupsB,1),
dS1A = matrix(0,nGroupsA,1),
dS1B = matrix(0,nGroupsB,1),
clogkwN = matrix(0,nColumns-1,nAnalytes),
cS1mN = matrix(0,nColumns-1,nAnalytes),
cdS1N = matrix(0,nColumns-1,nAnalytes),
etacdlogkT = matrix(0,nColumns-1,nAnalytes),
etacdlogkwA = matrix(0,nColumns-1,nGroupsA),
etacdlogkwB = matrix(0,nColumns-1,nGroupsB),
etacdS1mA = matrix(0,nColumns-1,nGroupsA),
etacdS1mB = matrix(0,nColumns-1,nGroupsB),
etacdS1A = matrix(0,nColumns-1,nGroupsA),
etacdS1B = matrix(0,nColumns-1, nGroupsB),

```

```

pKawA = pKaslitA,
pKawB = pKaslitB,
etaalphamA = matrix(0,nGroupsA,1),
etaalphamB = matrix(0,nGroupsB,1),
etadalphaA = matrix(0,nGroupsA,1),
etadalphaB = matrix(0,nGroupsB,1),
logsigma = rnorm(nAnalytes,log(0.386),0.813),
clogmsigma=c(rnorm(1,0.2,0.0170),
rnorm(1,-0.0170,0.0207),
rnorm(1,-0.260,0.0205),
rnorm(1,-0.100,0.0170)),
cssigma=abs(c(rnorm(1,0.0927,0.0236),
rnorm(1,0.169,0.0200),
rnorm(1,0.161,0.019),
rnorm(1,0.0854,0.0245))),
etaclogsigma = matrix(rnorm(nAnalytes*(nColumns-1),0,0.125),nrow=(nColumns-1))
)
}

mod_fixed <- cmdstan_model("stan/hplc-gra-fivecolumns-fixed.stan",
                           stanc_options = list("O1"))

fit_red <- mod_fixed$sample(
  data = datastruct_red,
  output_dir = "stanfiles",
  init = init_red,
  iter_warmup = 1000,
  iter_sampling = 500,
  chains = 4,
  parallel_chains = 4,
  refresh = 100,
  adapt_delta=0.9
)

fit_red <- cmdstanr::as_cmdstan_fit(c(
  'stanfiles/hplc-gra-fivecolumns-fixed-202308090920-1-544',
  'stanfiles/hplc-gra-fivecolumns-fixed-202308090920-2-544',
  'stanfiles/hplc-gra-fivecolumns-fixed-202308090920-3-544',
  'stanfiles/hplc-gra-fivecolumns-fixed-202308090920-4-544
))

```

12.1 Summary of individual parameters

Here shown for analyte 9 (Baclofen)

```
# which(unique(data$METID)==9) - 3rd compound in stan

fit_red$print(c("logkwx[3,1,1]", "logkwx[3,1,2]", "logkwx[3,1,3]",
               "logkwx[3,2,1]", "logkwx[3,2,2]", "logkwx[3,2,3]",
               "S1x[3,1,1,1]", "S1x[3,1,1,2]", "S1x[3,1,1,3]",
               "S1x[3,2,1,1]", "S1x[3,2,1,2]", "S1x[3,2,1,3]",
               "S1x[3,1,2,1]", "S1x[3,1,2,2]", "S1x[3,1,2,3]",
               "S1x[3,2,2,1]", "S1x[3,2,2,2]", "S1x[3,2,2,3]",
               "apHx[3,1,1]", "apHx[3,1,2]", "apHx[3,1,3]",
               "pKawx[3,1]", "pKawx[3,2]",
               "alphax[3,1,1]", "alphax[3,1,2]",
               "alphax[3,2,1]", "alphax[3,2,2]",
               "S2x[1,1]", "S2x[2,1]",
               "sigmax[3,1]", "sigmax[3,2]"), max_rows = 31)
```

variable	mean	median	sd	mad	q5	q95	rhat	ess_bulk	ess_tail
logkwx[3,1,1]	1.86	1.85	0.93	0.92	0.35	3.37	1.00	2529	1615
logkwx[3,1,2]	1.09	1.11	1.10	1.08	-0.74	2.93	1.00	2787	1492
logkwx[3,1,3]	1.09	1.11	1.10	1.08	-0.74	2.93	1.00	2787	1492
logkwx[3,2,1]	2.28	2.28	0.94	0.93	0.73	3.81	1.00	2566	1563
logkwx[3,2,2]	1.56	1.59	1.11	1.12	-0.29	3.38	1.00	2831	1587
logkwx[3,2,3]	1.56	1.59	1.11	1.12	-0.29	3.38	1.00	2831	1587
S1x[3,1,1,1]	3.88	3.88	0.98	0.99	2.28	5.47	1.00	2252	1400
S1x[3,1,1,2]	4.07	4.08	1.21	1.24	2.11	6.02	1.00	2970	1570
S1x[3,1,1,3]	4.07	4.08	1.21	1.24	2.11	6.02	1.00	2970	1570
S1x[3,2,1,1]	4.50	4.49	1.16	1.15	2.54	6.45	1.00	2274	1222
S1x[3,2,1,2]	4.99	5.03	1.48	1.49	2.54	7.35	1.00	2970	1545
S1x[3,2,1,3]	4.99	5.03	1.48	1.49	2.54	7.35	1.00	2970	1545
S1x[3,1,2,1]	4.61	4.61	0.98	0.98	3.00	6.20	1.00	2245	1382
S1x[3,1,2,2]	4.79	4.80	1.21	1.22	2.84	6.70	1.00	2941	1706
S1x[3,1,2,3]	4.79	4.80	1.21	1.22	2.84	6.70	1.00	2941	1706
S1x[3,2,2,1]	5.39	5.39	1.17	1.16	3.46	7.34	1.00	2331	1324
S1x[3,2,2,2]	5.91	5.92	1.49	1.47	3.46	8.30	1.00	2872	1544
S1x[3,2,2,3]	5.91	5.92	1.49	1.47	3.46	8.30	1.00	2872	1544
apHx[3,1,1]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
apHx[3,1,2]	-0.03	-0.03	0.00	0.00	-0.03	-0.03	1.00	2097	1267
apHx[3,1,3]	-0.03	-0.03	0.00	0.00	-0.03	-0.03	1.00	2097	1267
pKawx[3,1]	7.35	7.35	0.87	0.86	5.97	8.71	1.00	5293	1413

pKawx[3,2]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
alphax[3,1,1]	2.20	2.20	0.99	0.97	0.59	3.84	1.01	4318	1276
alphax[3,1,2]	0.00	0.00	0.00	0.00	0.00	NA	NA	NA	NA
alphax[3,2,1]	2.38	2.41	1.29	1.27	0.17	4.46	1.01	4760	1525
alphax[3,2,2]	0.00	0.00	0.00	0.00	0.00	0.00	NA	NA	NA
S2x[1,1]	0.47	0.47	0.01	0.01	0.44	0.49	1.00	1556	1617
S2x[2,1]	1.20	1.20	0.04	0.04	1.13	1.26	1.00	1020	1252
sigmax[3,1]	0.53	0.39	0.47	0.30	0.11	1.36	1.00	4335	1335
sigmax[3,2]	0.64	0.48	0.58	0.36	0.13	1.66	1.00	4345	1331

12.2 Predicted retention times

```

analyte_ID_sample =  c(9,17,33,58,140,180)
idx <- which(datasim$METID %in% analyte_ID_sample)
datasim_red = datasim[idx,1:20]
datastructsim_red <-datastructsim
datastructsim_red$nObs=length(datastructsim_red$analyte[idx])
datastructsim_red$analyte=datastructsim_red$analyte[idx]
datastructsim_red$modifier=datastructsim_red$modifier[idx]
datastructsim_red$steps=datastructsim_red$steps[idx]
datastructsim_red$column=datastructsim_red$column[idx]
datastructsim_red$hplcparam=datastructsim_red$hplcparam[idx,]
datastructsim_red$trobs = rep(0,datastructsim_red$nObs)
datastructsim_red$nexpid=length(unique(datasim_red$expid))
datastructsim_red$expid=match(datasim_red$expid, unique(datasim_red$expid))
datastructsim_red$nAnalytessim=length(unique(datastructsim_red$analyte))
datastructsim_red$analytesim=match(datastructsim_red$analyte, unique(datastructsim_red$analyte))

model_sim_red <- cmdstan_model("stan/hplc-gra-fivecolumns-fixed-sim.stan")

fit_sim_red <- model_sim_red$generate_quantities(fit_red,
                                                 data = datastructsim_red,
                                                 seed = 123,
                                                 parallel_chains = 4,
                                                 output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-fixed-sim-202308091209-1',
  'stanfiles/hplc-gra-fivecolumns-fixed-sim-202308091209-2',
  'stanfiles/hplc-gra-fivecolumns-fixed-sim-202308091209-3'
)

```

```

  'stanfiles/hplc-gra-fivecolumns-fixed-sim-202308091209-4
  ))}

draws_sim_red_df <- as_draws_df(x$generated_quantities)
#draws_sim_red_df <- fit_sim_red$draws(format = "df")

```

12.2.1 Plot the predicted vs. observed:

```

tr_sim_red_Cond <- apply(draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("tr
tr_sim_Cond<-as.data.frame(t(tr_sim_red_Cond))

datasim_red$trCond_l=tr_sim_Cond$`5%
datasim_red$trCond_m=tr_sim_Cond$`50%
datasim_red$trCond_h=tr_sim_Cond$`95%

idx <- which(data$METID %in% analyte_ID_sample &
  data$tg %in% c(30, 90, 270) &
  data$pH %in% c(1:9) &
  data$Column %in% c(1) & # c(1, 2)
  data$Mod2 %in% c(1, 2) &
  data$Temp %in% c(25, 35))

data_red = data[idx,]

for(i in 1:length(analyte_ID_sample)){
  pi <- ggplot()+
    geom_point(data=data[which(data$METID %in% analyte_ID_sample[i]),], aes(x = pHs, y =
    geom_point(data=data_red[which(data_red$METID %in% analyte_ID_sample[i]),], aes(x =
      geom_line(data=datasim_red[which(datasim_red$METID %in% analyte_ID_sample[i]),], aes(
      geom_ribbon(data=datasim_red[which(datasim_red$METID %in% analyte_ID_sample[i]),], ae
      xlim(2,11)+

  facet_grid(Column~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+

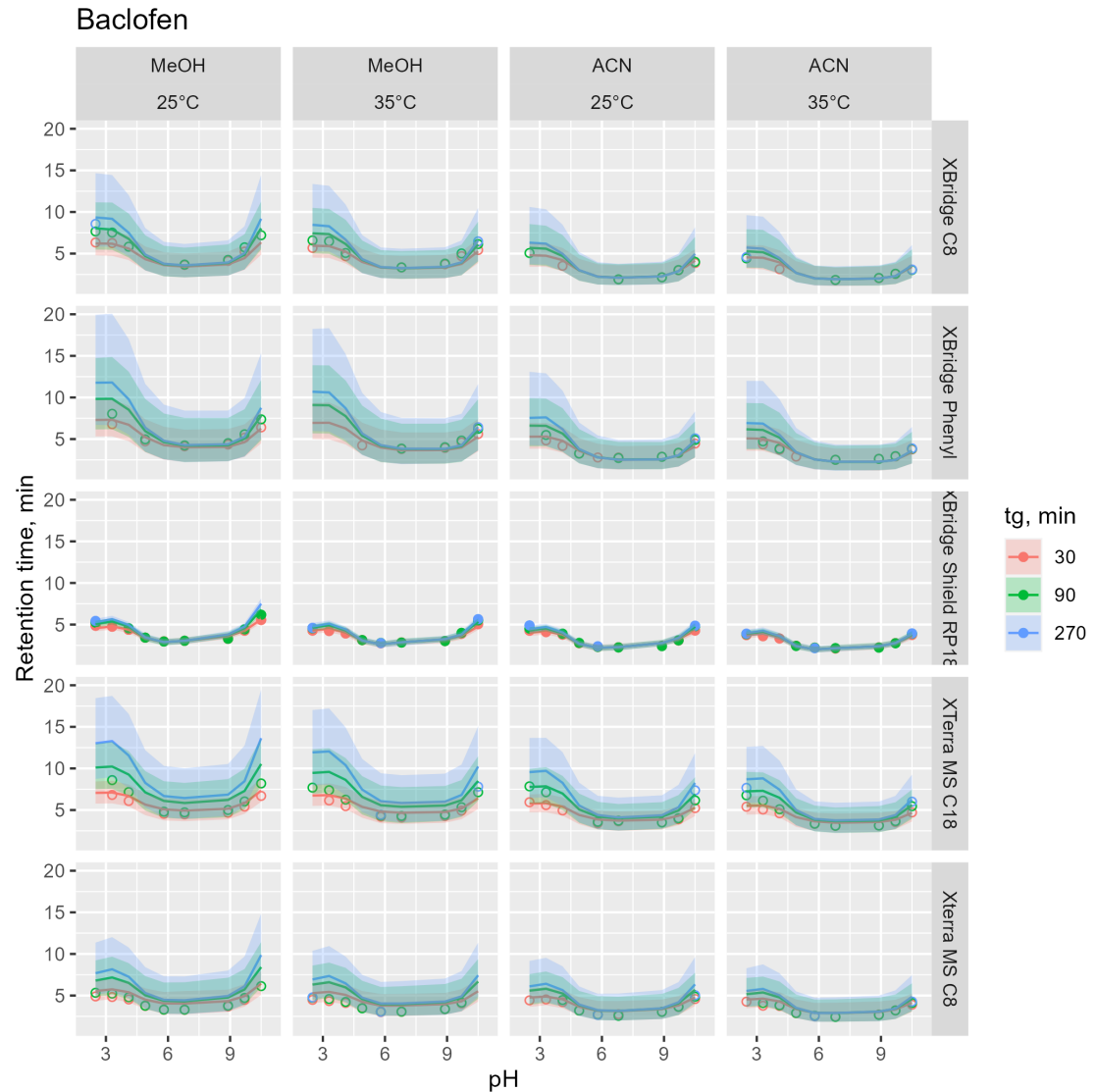
  labs(title=paste(dataNames$name[analyte_ID_sample[i]]), x ="pH", y = "Retention time"

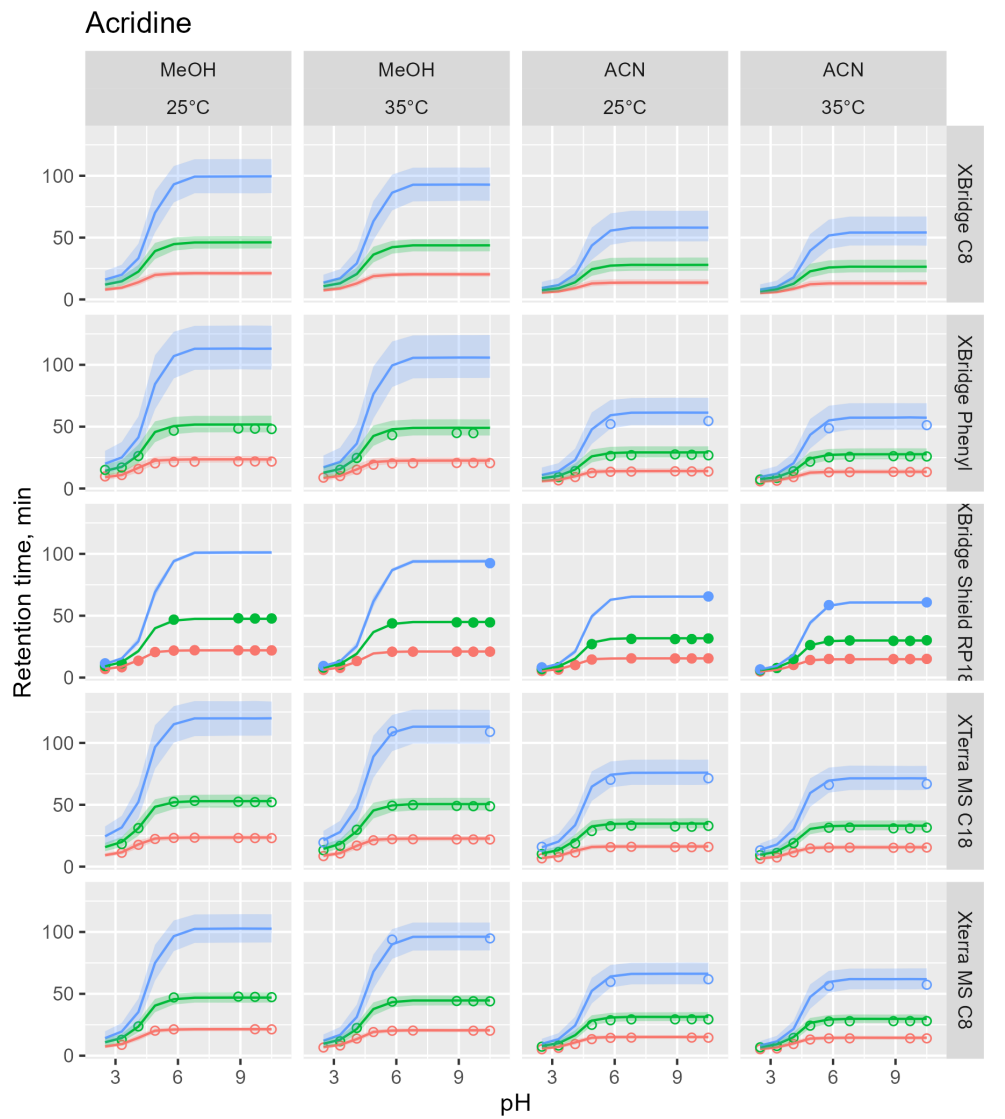
```

```

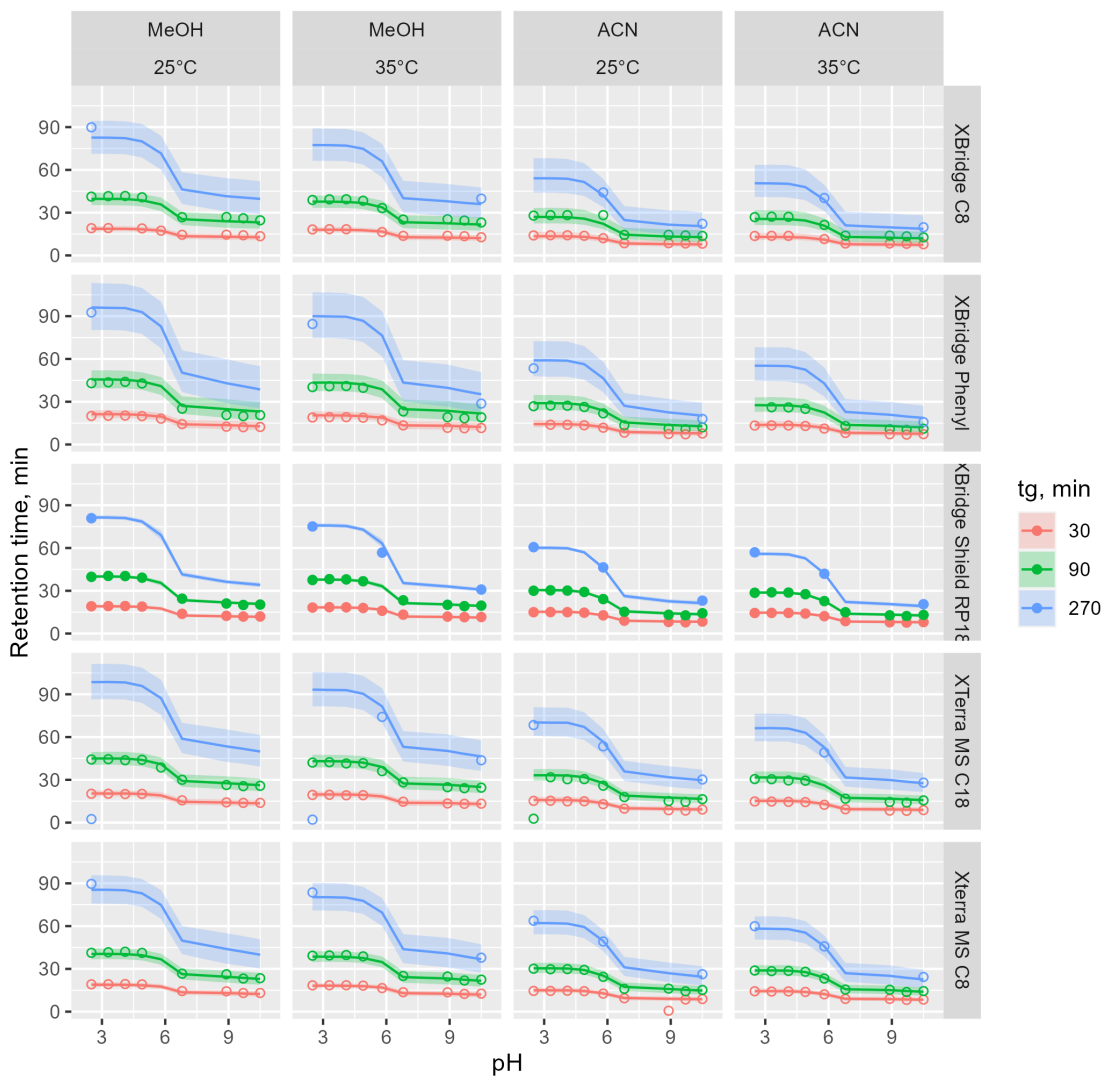
print(pi)

ggsave(paste0("figures\\casestudy2\\concordanceplots\\", paste(dataNames$name[analyte_ID_s
} 
```

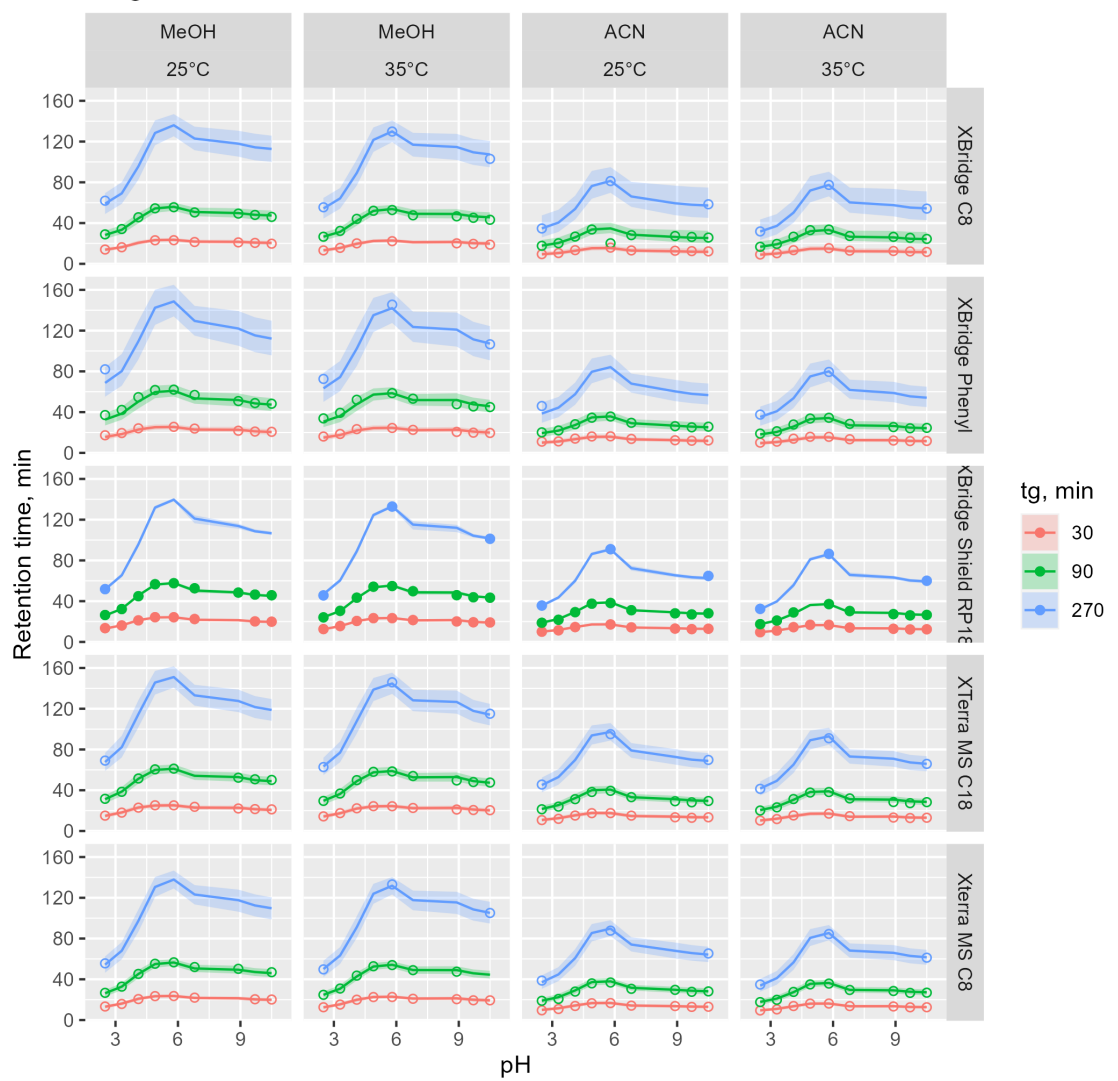




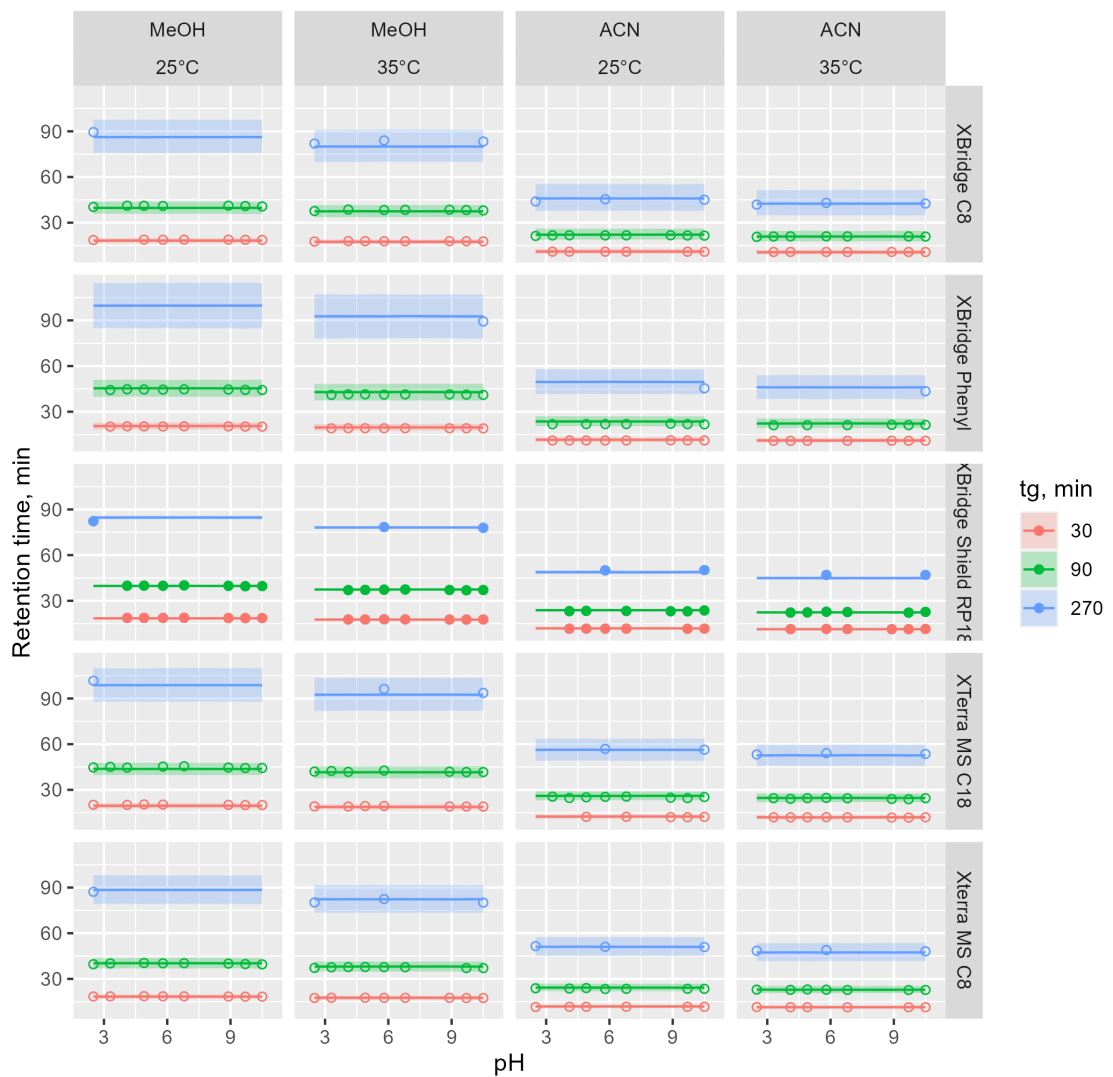
Tolbutamide

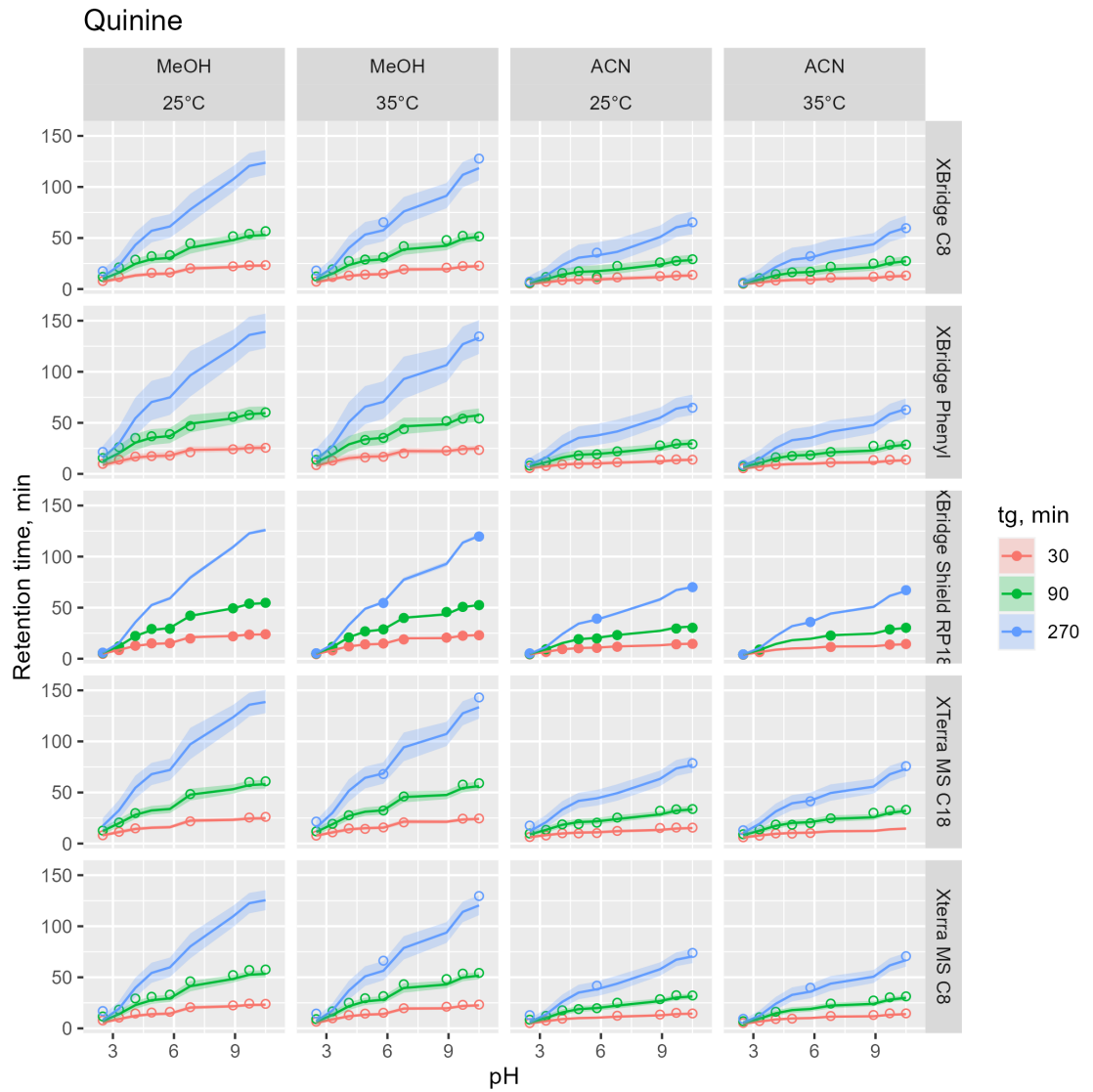


Pioglitazone



Hydrocortisone





12.3 Decision making

The same utility function was used:

```
foo<-datasim_red[!duplicated(datasim_red$expid),]

x<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("mindifftr", names(draws_)

y<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("maxtr", names(draws_sim_
```

```

z<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("mintr", names(draws_sim_))

u=x;

for (i in 1:ncol(x)){u[,i] <- as.numeric(x[,i]>2 & z[,i] > 2) * (40-y[,i]) * as.numeric(y[

u <- apply(u, MARGIN = 2, FUN = mean)
u<-as.data.frame(u)

foo$EUutility=u$u
x <- apply(x, MARGIN = 2, FUN = quantile, probs = c(.05,.5,.95))
x<-as.data.frame(t(x))

foo$mindifftr_l=x$`5%
foo$mindifftr_m=x$`50%
foo$mindifftr_h=x$`95%

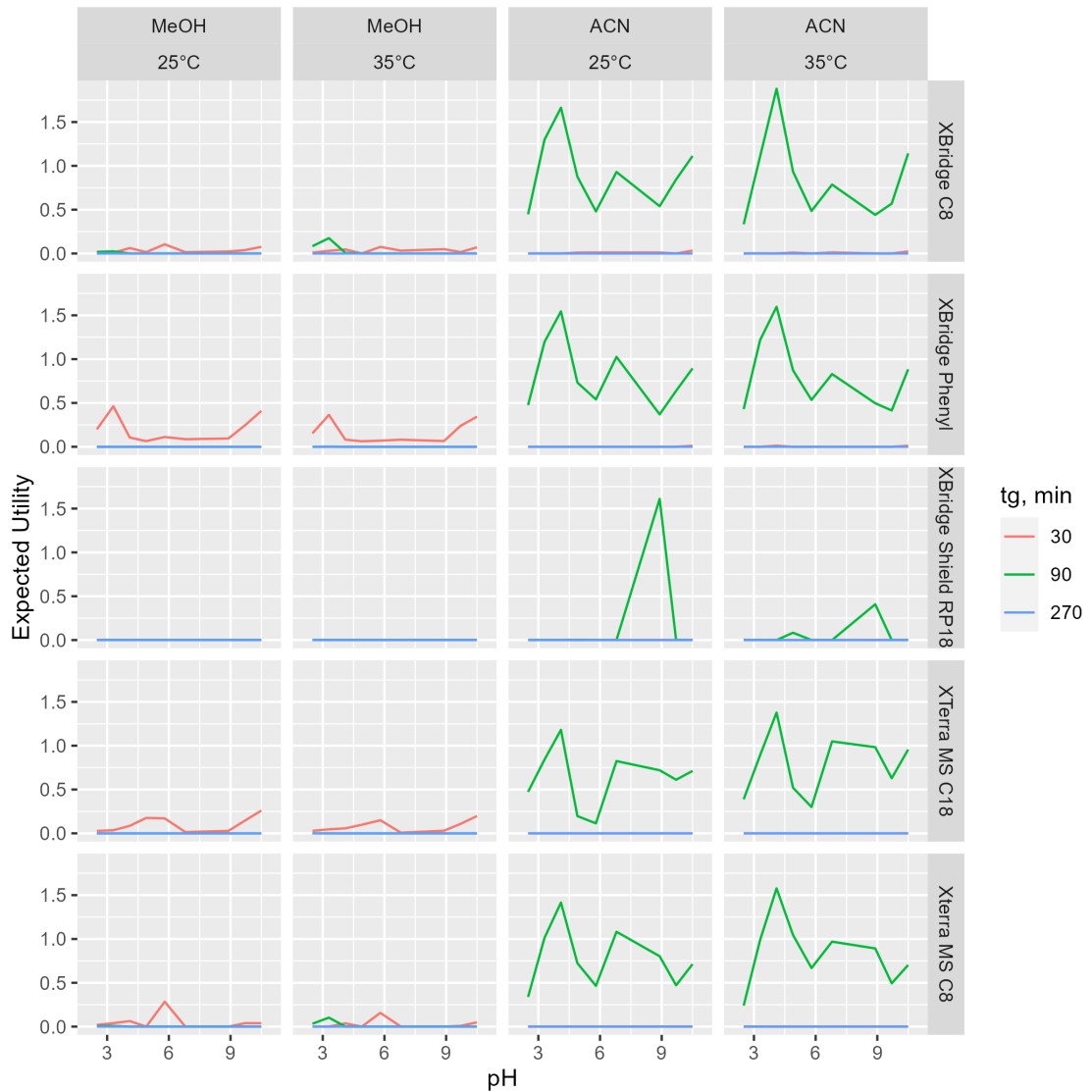
y <- apply(y, MARGIN = 2, FUN = quantile, probs = c(.05,.5,.95))

y<-as.data.frame(t(y))
foo$maxtr_l=y$`5%
foo$maxtr_m=y$`50%
foo$maxtr_h=y$`95%

p1 <- ggplot()+
  geom_line(data=foo, aes(x = pHs, y = EUutility, color = as.factor(tg)))+
  xlim(2,11)+
  facet_grid(ColumnNames~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+
  labs(x = "pH",
       y = "Expected Utility",
       color = "tg, min")

print(p1)

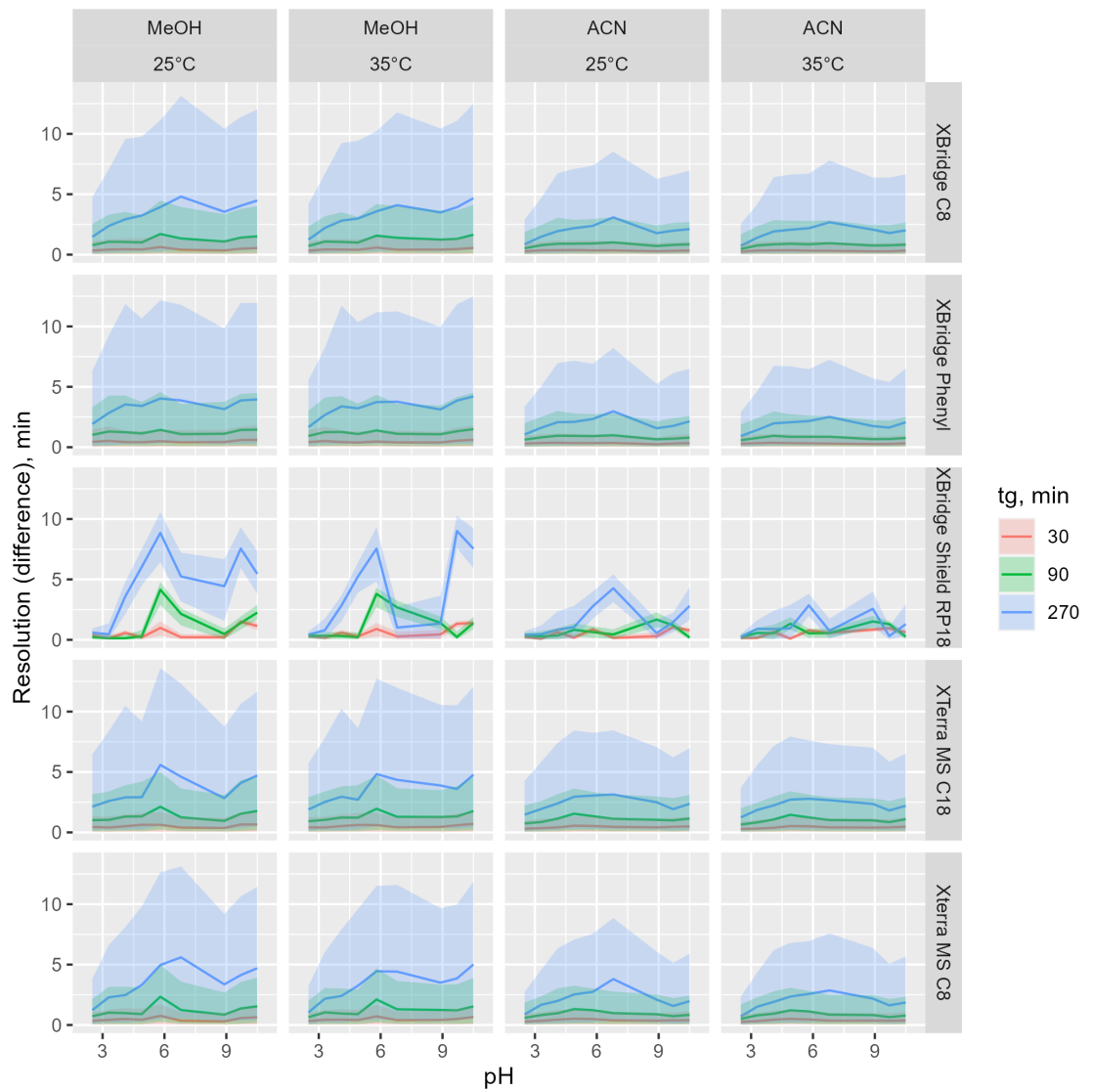
```



```

p2 <- ggplot()+
  geom_line(data=foo, aes(x = pHs, y = mindifftr_m, color = as.factor(tg)))+
  geom_ribbon(data=foo, aes(x = pHs, ymin = mindifftr_l, ymax = mindifftr_h, fill = as.factor(tg)))+
  facet_grid(ColumnName~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+
  labs(x ="pH",
       y = "Resolution (difference), min",
       color = "tg, min",
       fill = "tg, min")
  
```

```
print(p2)
```



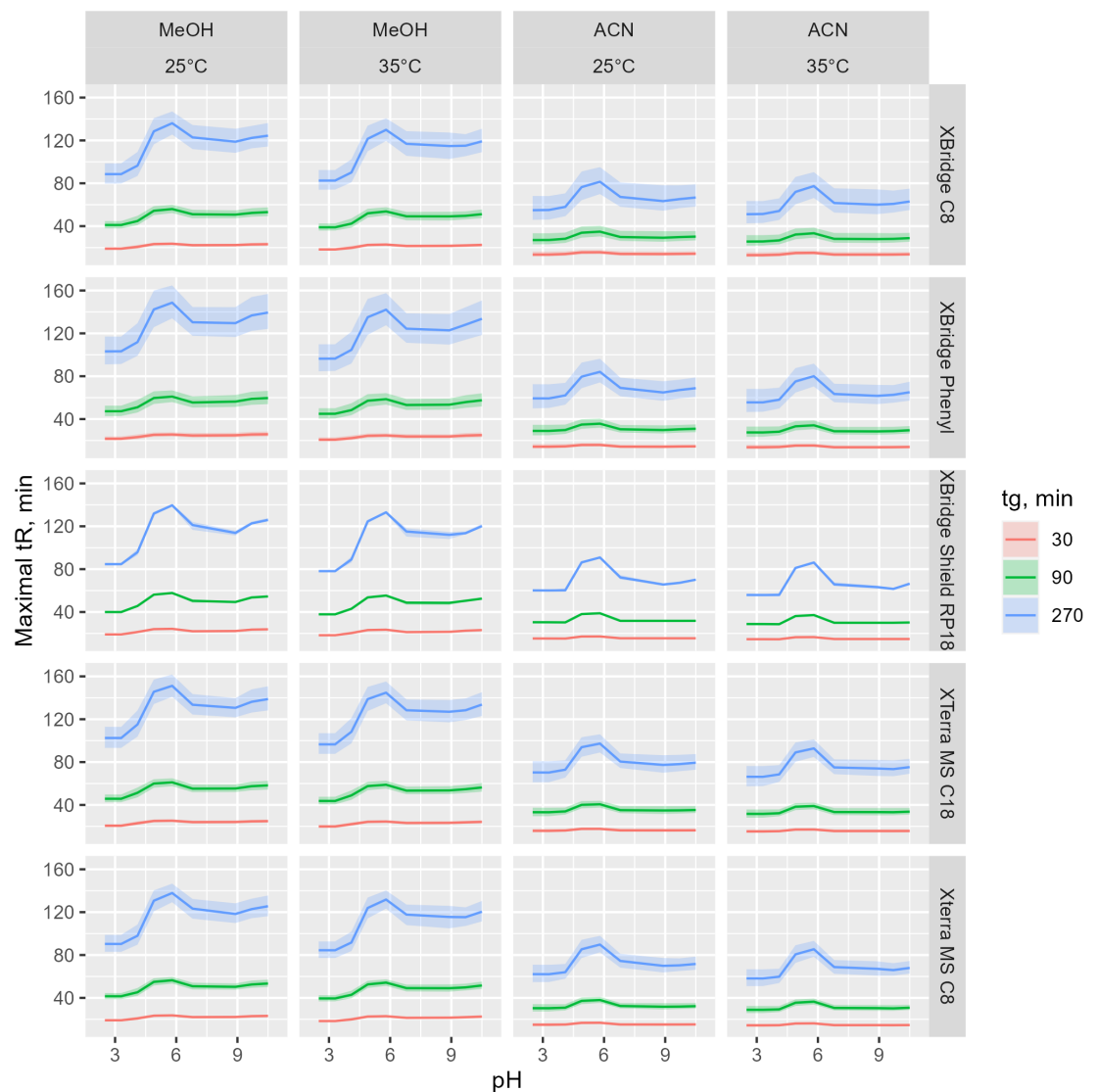
```
p3 <- ggplot()+
  geom_line(data=foo, aes(x = pHs, y = maxtr_m, color = as.factor(tg)))+
  geom_ribbon(data=foo, aes(x = pHs, ymin = maxtr_l, ymax = maxtr_h, fill = as.factor(tg)))
  xlim(2,11)+
  facet_grid(ColumnName~Mod2+Temp, labeller = labeller(Temp=temp.labs,Mod2=mod.labs))+
  labs(x ="pH",
```

```

y = "Maximal tR, min",
color = "tg, min",
fill = "tg, min")

print(p3)

```



```

ggsave(paste0("figures\\casestudy2\\decision\\", "ExpectedUtility", ".png"), plot=p1, width = 10)
ggsave(paste0("figures\\casestudy2\\decision\\", "Resolution", ".png"), plot=p2, width = 10)

```

```

ggsave(paste0("figures\\casestudy2\\decision\\", "maxtr", ".png"), plot=p3, width = 20, height = 10)

# The best for XBridge Shield RP18
foo1<-foo[foo$Column==1,];
foo1[which(foo1$EUtility==max(foo1$EUtility)),c('tg','fio', 'fik','Mod', 'pH','pHo','Temp')]

tg  fio fik Mod pH      pHo Temp      ColumnName EUtility
158 90 0.05 0.8 ACN  7 8.87588  25 XBridge Shield RP18 1.611051

# The best choice for XTerra MS C18
foo2<-foo[foo$Column==2,];
foo2[which(foo2$EUtility==max(foo2$EUtility)),c('tg','fio', 'fik','Mod', 'pH','pHo','Temp')]

tg  fio fik Mod pH      pHo Temp      ColumnName EUtility
15404 90 0.05 0.8 ACN  3 4.202902  35 XTerra MS C18 1.377401

# The best choice for XBridge Phenyl
foo3<-foo[foo$Column==3,];
foo3[which(foo3$EUtility==max(foo3$EUtility)),c('tg','fio', 'fik','Mod', 'pH','pHo','Temp')]

tg  fio fik Mod pH      pHo Temp      ColumnName EUtility
30632 90 0.05 0.8 ACN  3 4.202902  35 XBridge Phenyl 1.596929

# The best choice for XBridge C8
foo4<-foo[foo$Column==4,];
foo4[which(foo4$EUtility==max(foo4$EUtility)),c('tg','fio', 'fik','Mod', 'pH','pHo','Temp')]

tg  fio fik Mod pH      pHo Temp ColumnName EUtility
45860 90 0.05 0.8 ACN  3 4.202902  35 XBridge C8 1.880014

# The best choice for Xterra MS C8
foo5<-foo[foo$Column==5,];
foo5[which(foo5$EUtility==max(foo5$EUtility)),c('tg','fio', 'fik','Mod', 'pH','pHo','Temp')]

tg  fio fik Mod pH      pHo Temp      ColumnName EUtility
61088 90 0.05 0.8 ACN  3 4.202902  35 Xterra MS C8 1.577515

```

12.3.1 Uncertainty chromatogram

Below is the example of a chromatogram (along with uncertainty) expected for $tg = 90$ min, pH = 8.9, Temp = 25°C in ACN.

```
analyte_ID_sample <- c(9, 17, 33, 58, 140, 180)

col.labs <- c("XBridge Shield RP18", "XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xter

wpCond= data.frame()

for (i in 1:5) {

  idx <- which(datasim_red$METID %in% analyte_ID_sample &
                datasim_red$tg==90 & # c(30, 90, 270)
                datasim_red$pH== 7 & # c(1:9)
                datasim_red$Column==i & # c(1, 2)
                datasim_red$Mod2==2 & # c(1, 2)
                datasim_red$Temp==25) # c(25, 35)

  data_to_plot <- draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% paste0("trHatCond[

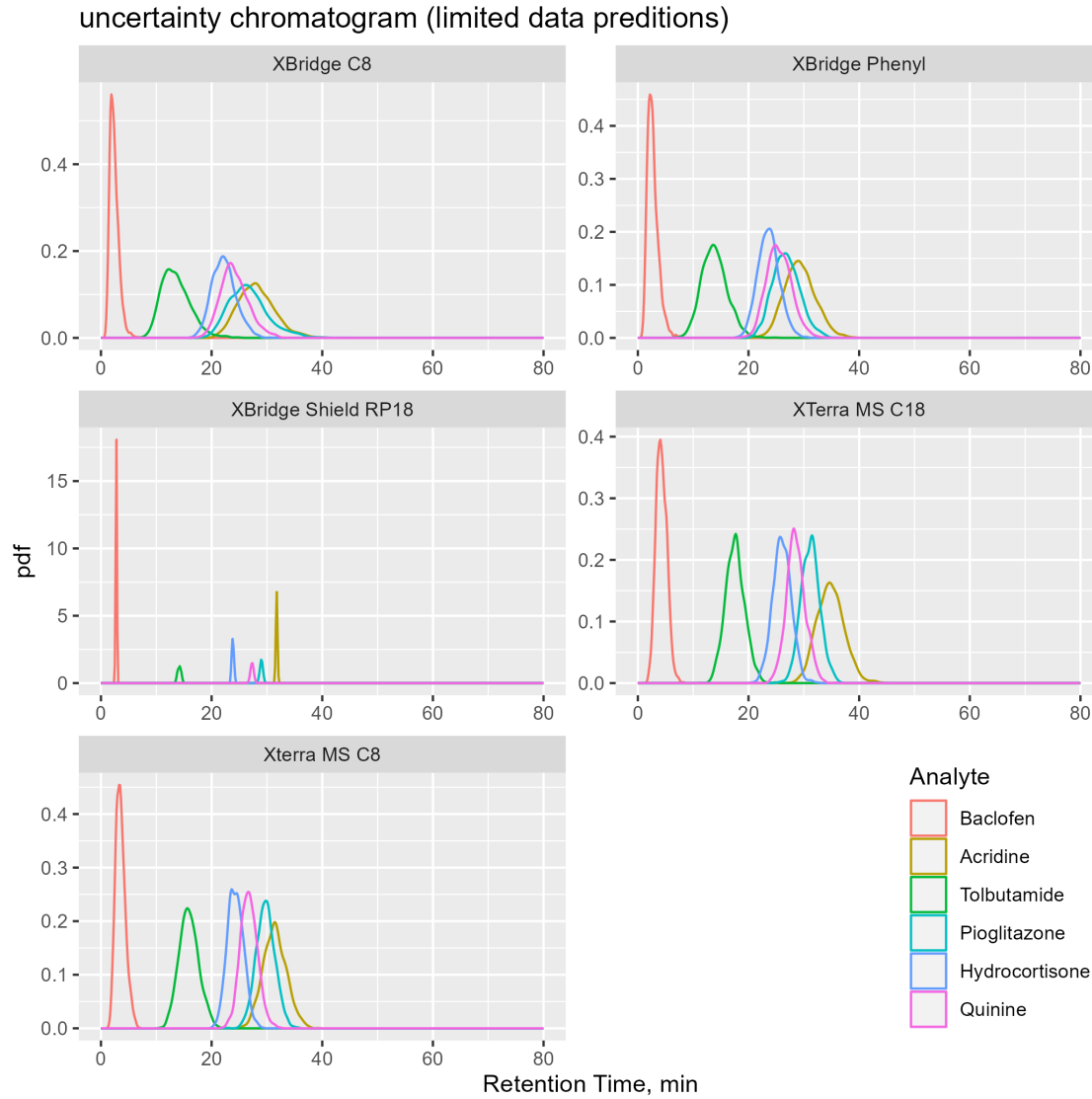
  colnames(data_to_plot) <- paste(dataNames>Name[analyte_ID_sample])

  wpCond1 <- melt(data_to_plot)
  wpCond1$Column <- unname(col.labs[i])
  wpCond= rbind(wpCond,wpCond1)

}

wpCond$Column=as.factor(wpCond$Column)
p <- ggplot(data = wpCond)+
  geom_density(aes(x=value, colour=variable)) +
  labs(title="uncertainty chromatogram (limited data predictions)",
       x ="Retention Time, min",
       y = "pdf",
       colour="Analyte")+
  xlim(c(0,80))+
  facet_wrap(.~Column, nrow=3,scales = "free")+
  theme(legend.position = c(1, 0),
        legend.justification = c(1, 0))
```

```
print(p)
```



```
ggsave(paste0("figures\\casestudy2\\decision\\", "Chromatogram", ".png"), plot=p, width =
```

12.4 Utility maps

We can also determine more detailed graph presenting relationship between the expected utility and design variables:

```

upH <- datasim %>%
  select(1:20) %>%
  distinct(Temp,Mod, pHo, alpha1, alpha2) %>%
  group_by(Temp,Mod) %>%
  tidyr::complete(pHo = seq(min(pHo), max(pHo), len = 17)) %>%
  arrange(Temp,Mod,pHo) %>%
  mutate(alpha1 = zoo::na.approx(alpha1,pHo)) %>%
  mutate(alpha2 = zoo::na.approx(alpha2,pHo)) %>%
  group_by(Temp,Mod) %>%
  mutate(pHid = row_number()) %>%
  group_by(pHid) %>%
  mutate(pHs = round(mean(pHo),2)) %>%
  ungroup()

datasim2 <- datasim %>%
  select(1:20) %>%
  filter(tg==30, Temp==25) %>%
  select(-tg, -fio, -expid, -pHs, -pHo, -alpha1, -alpha2, -pH) %>%
  distinct() %>%
  tidyr::expand_grid(fio = seq(0.05,0.2,0.05),tg = seq(20,260,20), pHs=unique(upH$pHs))%>%
  left_join(upH, by = join_by(pHs, Mod, Temp), relationship = "many-to-one") %>%
  group_by(tg, fio, Mod, Column, pHs) %>%
  mutate(expid = cur_group_id()) %>%
  ungroup()
datastructsim2 <-datastruct
datastructsim2$nObs=length(datasim2$METID)
datastructsim2$analyte=match(datasim2$METID, unique(datasim2$METID))
datastructsim2$modifier=match(datasim2$Mod2, sort(unique(datasim2$Mod2)))
datastructsim2$steps=4*(2-datasim2$Mod2) + 10*(datasim2$Mod2-1)
datastructsim2$column=match(datasim2$Column, unique(datasim2$Column))
datastructsim2$hplcparam=cbind(datasim2$tg,datasim2$td,datasim2$to,datasim2$te,
                               datasim2$fio,datasim2$fik,datasim2$Mod2-1,datasim2$pHo,
                               datasim2$alpha1,datasim2$alpha2,(datasim2$Temp-25)/10,
                               datasim2$Column-1)
datastructsim2$trobs = rep(0,datastructsim2$nObs)
analyte_ID_sample = c(9,17,33,58,140,180)
idx <- which(datasim2$METID %in% analyte_ID_sample)
datasim_red = datasim2[idx,]
datastructsim_red <-datastructsim2
datastructsim_red$nObs=length(datastructsim_red$analyte[idx])
datastructsim_red$analyte=datastructsim_red$analyte[idx]

```

```

datastructsim_red$modifier=datastructsim_red$modifier[idx]
datastructsim_red$steps=datastructsim_red$steps[idx]
datastructsim_red$column=datastructsim_red$column[idx]
datastructsim_red$hplcparam=datastructsim_red$hplcparam[idx,]
datastructsim_red$trobs = rep(0,datastructsim_red$nObs)
datastructsim_red$nexpid=length(unique(datasim_red$expid))
datastructsim_red$expid=match(datasim_red$expid, unique(datasim_red$expid))
datastructsim_red$nAnalytessim=length(unique(datastructsim_red$analyte))
datastructsim_red$analytesim=match(datastructsim_red$analyte, unique(datastructsim_red$ana
model_sim_red <- cmdstan_model("stan/hplc-gra-fivecolumns-fixed-sim.stan")
fit_sim_red <- model_sim_red$generate_quantities(fit_red,
                                                 data = datastructsim_red,
                                                 seed = 123,
                                                 parallel_chains = 4,
                                                 output_dir = "stanfiles")

x<- cmdstanr::read_cmdstan_csv(c(
  'stanfiles/hplc-gra-fivecolumns-fixed-sim-202308091217-1
))

draws_sim_red_df <- as_draws_df(x$generated_quantities)
#draws_sim_red_df <- fit_sim_red$draws(format = "df")

```

The best chromatogram can be sought based on utility map:

```

foo<-datasim_red[!duplicated(datasim_red$expid),]

x<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("^mindifftr", names(draws_
y<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("^maxtr", names(draws_sim_
z<-draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% grep("^mintr", names(draws_sim_
u=x;

for (i in 1:ncol(x)){u[,i] <- as.numeric(x[,i]>2 & z[,i] > 2) * (40-y[,i]) * as.numeric(y[

pr <- apply(u, MARGIN = 2, FUN = mean)

pr<-as.data.frame(pr)

```

```

foo$EUtility=pr$pr

x <- apply(x, MARGIN = 2, FUN = quantile, probs = c(.05,.5,.95))

x<-as.data.frame(t(x))

foo$mindifftr_l=x$`5%
foo$mindifftr_m=x$`50%
foo$mindifftr_h=x$`95%

y <- apply(y, MARGIN = 2, FUN = quantile, probs = c(.05,.5,.95))

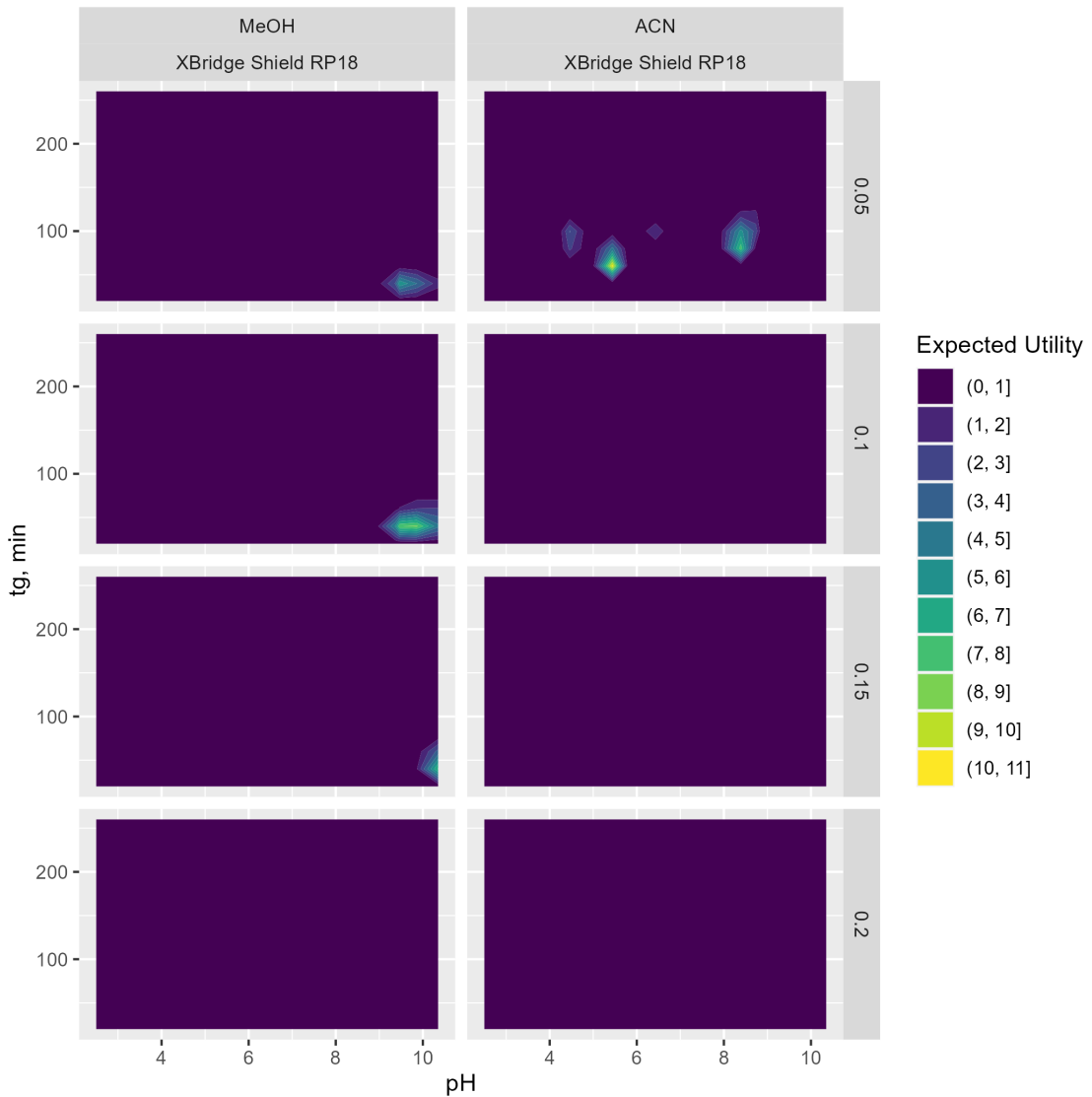
y<-as.data.frame(t(y))

foo$maxtr_l=y$`5%
foo$maxtr_m=y$`50%
foo$maxtr_h=y$`95%

p11 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==1), aes(x = pHs, y=tg, z = EUtility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p11)

```

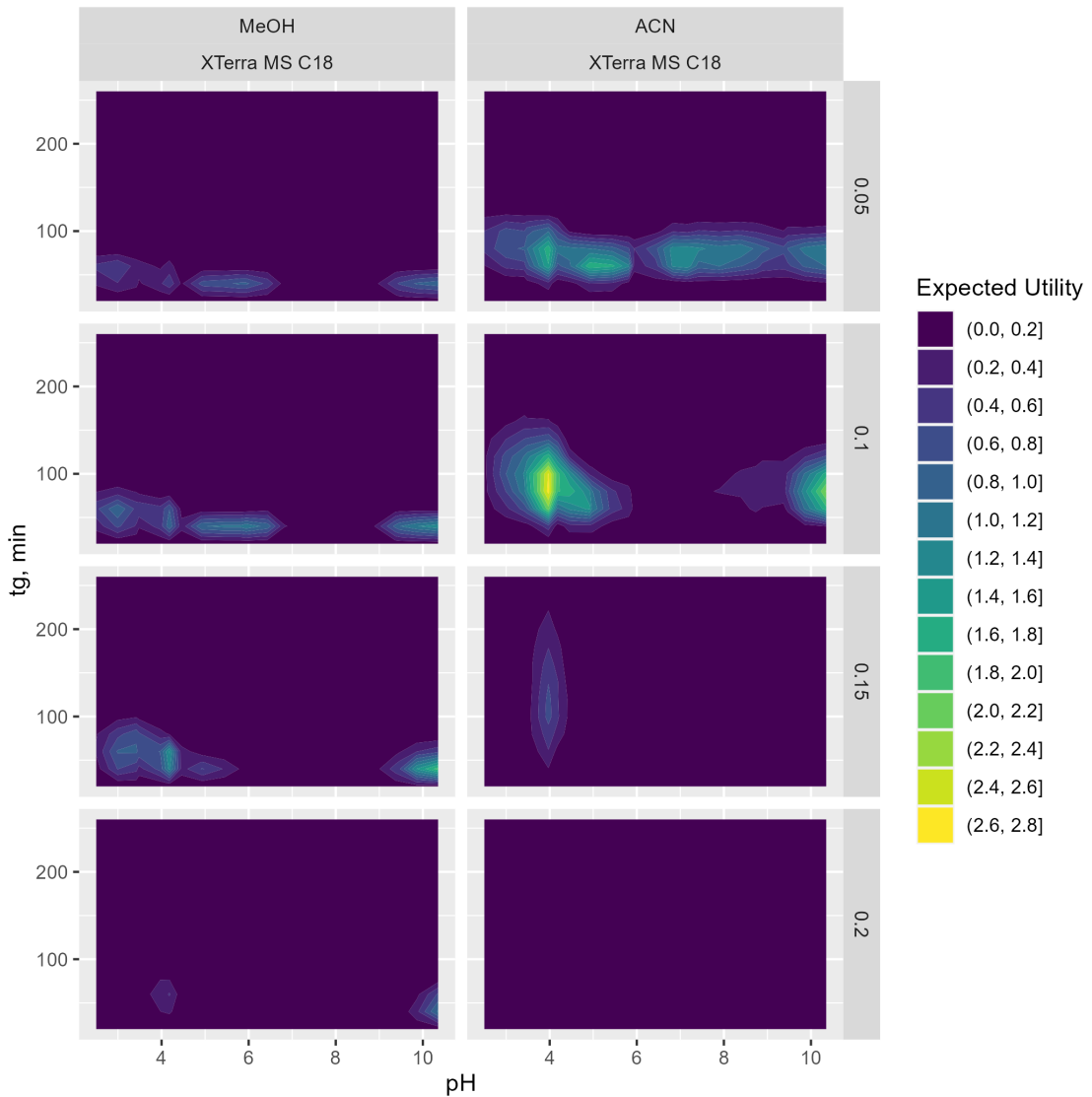


```

p12 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==2), aes(x = pHs, y=tg, z = EUutility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x ="pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p12)

```

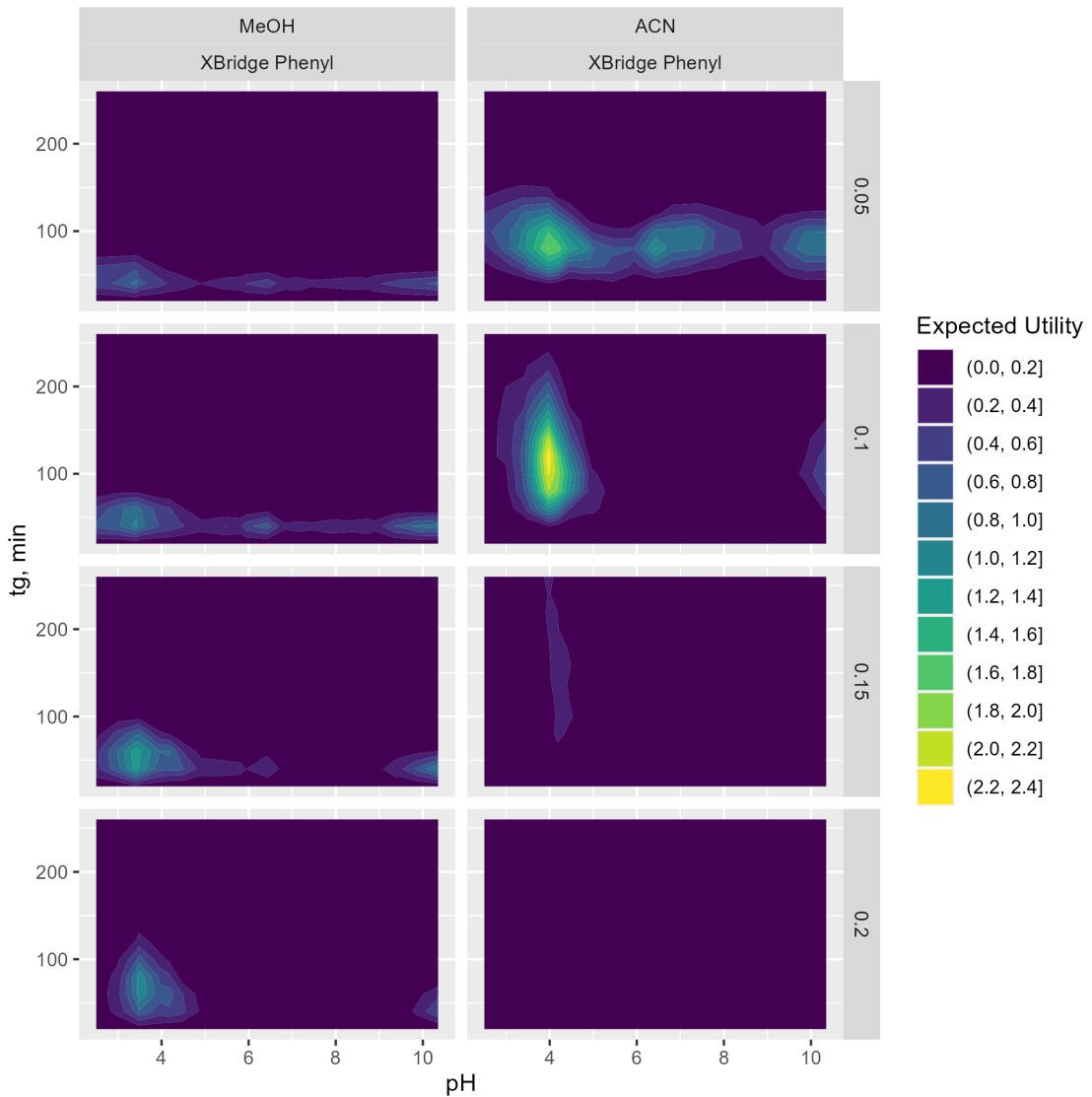


```

p13 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==3), aes(x = pHs, y=tg, z = EUutility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x ="pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p13)

```

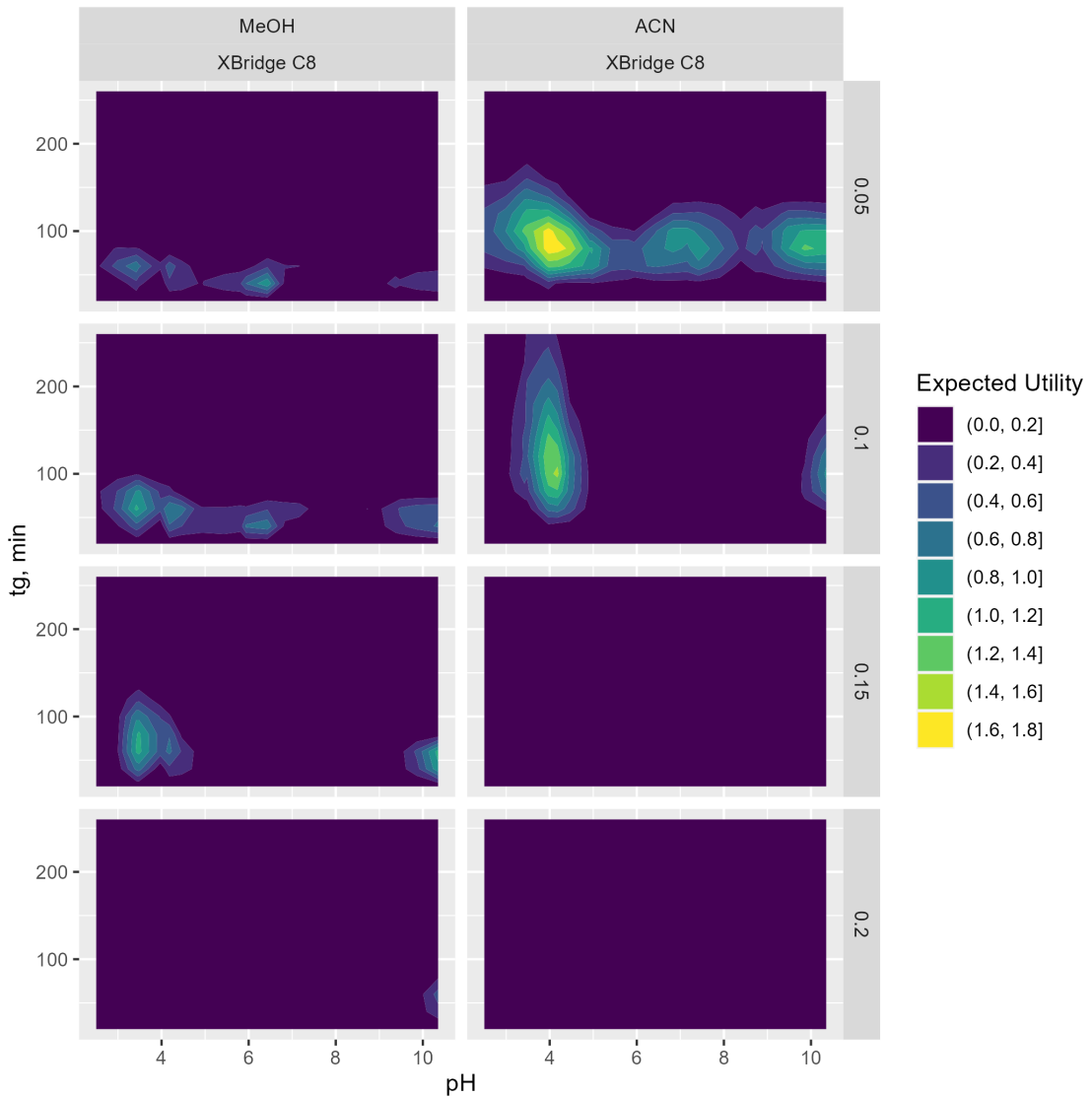


```

p14 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==4), aes(x = pHs, y=tg, z = EUutility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x ="pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p14)

```

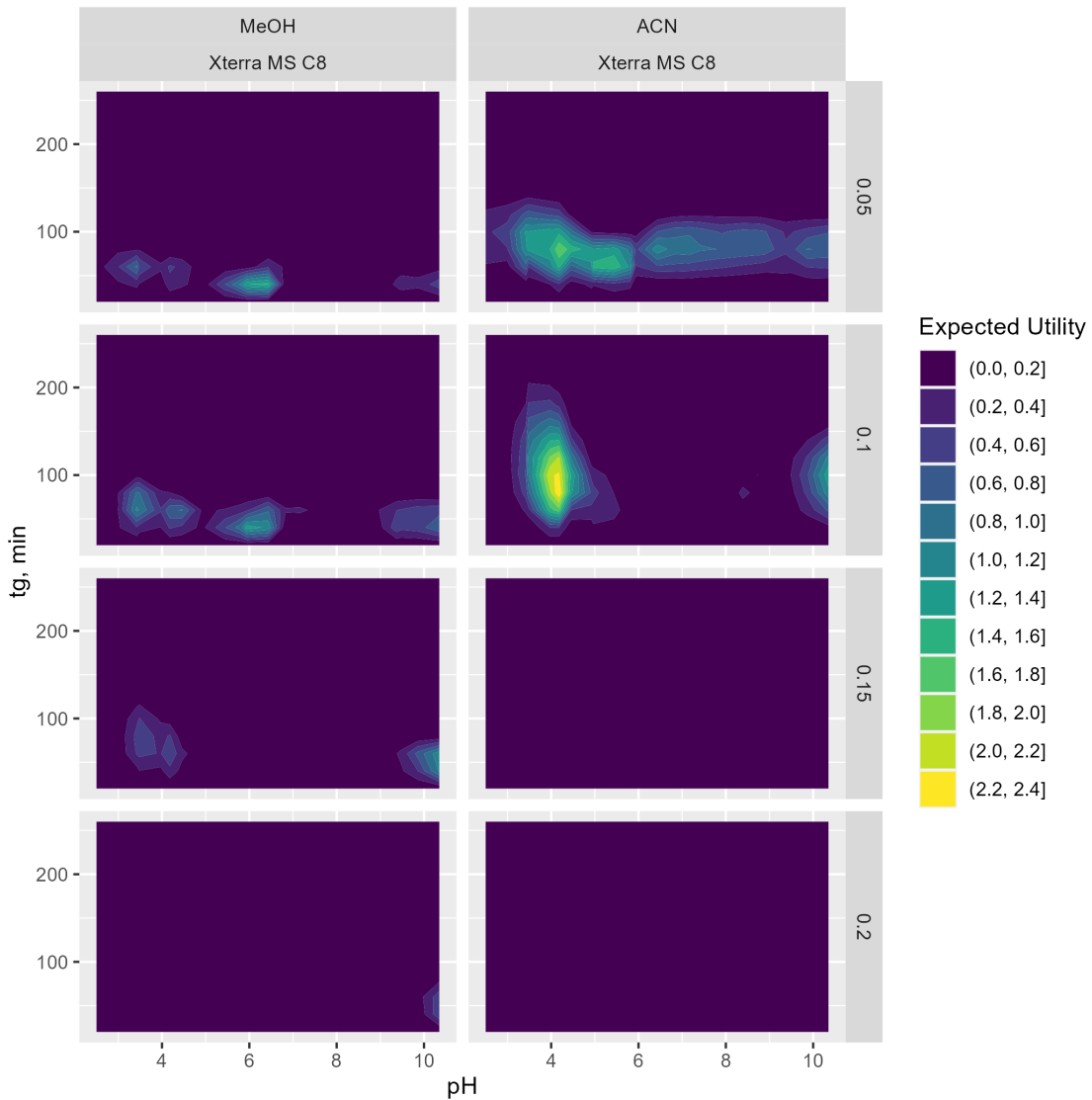


```

p15 <- ggplot()+
  geom_contour_filled(data=subset(foo,Column==5), aes(x = pHs, y=tg, z = EUutility))+
  facet_grid(fio~Mod2+ColumnName, labeller = labeller(Mod2=mod.labs))+ 
  labs(x = "pH",
       y = "tg, min",
       fill = "Expected Utility")

print(p15)

```



```

ggsave(paste0("figures\\casestudy2\\utilitymap\\", "utilitymap1", ".png"), plot=p11, width=10, height=8)
ggsave(paste0("figures\\casestudy2\\utilitymap\\", "utilitymap2", ".png"), plot=p12, width=10, height=8)
ggsave(paste0("figures\\casestudy2\\utilitymap\\", "utilitymap3", ".png"), plot=p13, width=10, height=8)
ggsave(paste0("figures\\casestudy2\\utilitymap\\", "utilitymap4", ".png"), plot=p14, width=10, height=8)
ggsave(paste0("figures\\casestudy2\\utilitymap\\", "utilitymap5", ".png"), plot=p15, width=10, height=8)

# The best for XBridge Shield RP18
foo1<-foo[foo$Column==1,];
foo1[which(foo1$EUtility==max(foo1$EUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo','Te

```

```

# A tibble: 1 x 9
  tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUUtility
  <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
1    60    0.05   0.8 ACN      10   5.50    25 XBridge Shield RP18     10.4

  # The best choice for XTerra MS C18
  foo2<-foo[foo$Column==2,];
  foo2[which(foo2$EUUtility==max(foo2$EUUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo', 'Te

# A tibble: 1 x 9
  tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUUtility
  <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
1    80    0.1    0.8 ACN      5    4.00    25 XTerra MS C18     2.69

  # The best choice for XBridge Phenyl
  foo3<-foo[foo$Column==3,];
  foo3[which(foo3$EUUtility==max(foo3$EUUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo', 'Te

# A tibble: 1 x 9
  tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUUtility
  <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
1   120    0.1    0.8 ACN      5    4.00    25 XBridge Phenyl    2.36

  # The best choice for XBridge C8
  foo4<-foo[foo$Column==4,];
  foo4[which(foo4$EUUtility==max(foo4$EUUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo', 'Te

# A tibble: 1 x 9
  tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUUtility
  <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
1    80    0.05   0.8 ACN      5    4.00    25 XBridge C8      1.74

  # The best choice for Xterra MS C8
  foo5<-foo[foo$Column==5,];
  foo5[which(foo5$EUUtility==max(foo5$EUUtility)),c('tg','fio', 'fik','Mod', 'pHid','pHo', 'Te

# A tibble: 1 x 9
  tg    fio    fik Mod    pHid    pHo  Temp ColumnName      EUUtility
  <dbl> <dbl> <dbl> <chr> <int> <dbl> <int> <chr>           <dbl>
1    80    0.1    0.8 ACN      6    4.17    25 Xterra MS C8     2.39

```

12.4.1 Uncertainty chromatogram

Let's visualize chromatograms with the highest expected utility given the experimental data.

```
analyte_ID_sample <- c(9,17,33,58,140,180)

col.labs <- c("XBridge Shield RP18", "XTerra MS C18", "XBridge Phenyl", "XBridge C8", "Xter

datasim_red$fio = as.factor(datasim_red$fio)

wpCond= data.frame()

for (i in 1:5) {

  idx <- which(datasim_red$METID %in% analyte_ID_sample &
                datasim_red$tg == 60 & # c(30, 90, 270)
                datasim_red$pHid == 10 & # c(1:9)
                datasim_red$fio == 0.05 & # ...
                datasim_red$Column == i & # c(1, 2)
                datasim_red$Mod2 == 2 & # c(1, 2)
                datasim_red$Temp == 25) # c(25, 35)

  if (i==2){

    idx <- which(datasim_red$METID %in% analyte_ID_sample &
                  datasim_red$tg == 80 & # c(30, 90, 270)
                  datasim_red$pHid == 5 & # c(1:9)
                  datasim_red$fio == 0.1 & # ...
                  datasim_red$Column == i & # c(1, 2)
                  datasim_red$Mod2 == 2 & # c(1, 2)
                  datasim_red$Temp == 25) # c(25, 35)

  }

  if (i==3){

    idx <- which(datasim_red$METID %in% analyte_ID_sample &
                  datasim_red$tg == 120 & # c(30, 90, 270)
                  datasim_red$pHid == 5 & # c(1:9)
                  datasim_red$fio == 0.1 & # ...
                  datasim_red$Column == i & # c(1, 2)
                  datasim_red$Mod2 == 2 & # c(1, 2)
```

```

datasim_red$Temp == 25) # c(25, 35)

}

if (i==4){

idx <- which(datasim_red$METID %in% analyte_ID_sample &
  datasim_red$tg == 100 & # c(30, 90, 270)
  datasim_red$pHid == 5 & # c(1:9)
  datasim_red$fio == 0.05 & # ...
  datasim_red$Column == i & # c(1, 2)
  datasim_red$Mod2 == 2 & # c(1, 2)
  datasim_red$Temp == 25) # c(25, 35)

}

if (i==5){

idx <- which(datasim_red$METID %in% analyte_ID_sample &
  datasim_red$tg == 100 & # c(30, 90, 270)
  datasim_red$pHid == 6 & # c(1:9)
  datasim_red$fio == 0.1 & # ...
  datasim_red$Column == i & # c(1, 2)
  datasim_red$Mod2 == 2 & # c(1, 2)
  datasim_red$Temp == 25) # c(25, 35)

}

data_to_plot <- draws_sim_red_df[,which(colnames(draws_sim_red_df) %in% paste0("trHatCond", colnames(data_to_plot)) <- paste(dataNames>Name[analyte_ID_sample])]

wpCond1 <- melt(data_to_plot)
wpCond1$Column <- unname(col.labs[i])
wpCond= rbind(wpCond,wpCond1)

}

wpCond$Column=as.factor(wpCond$Column)

p <- ggplot(data = wpCond)+  

  geom_density(aes(x=value, colour=variable)) +  

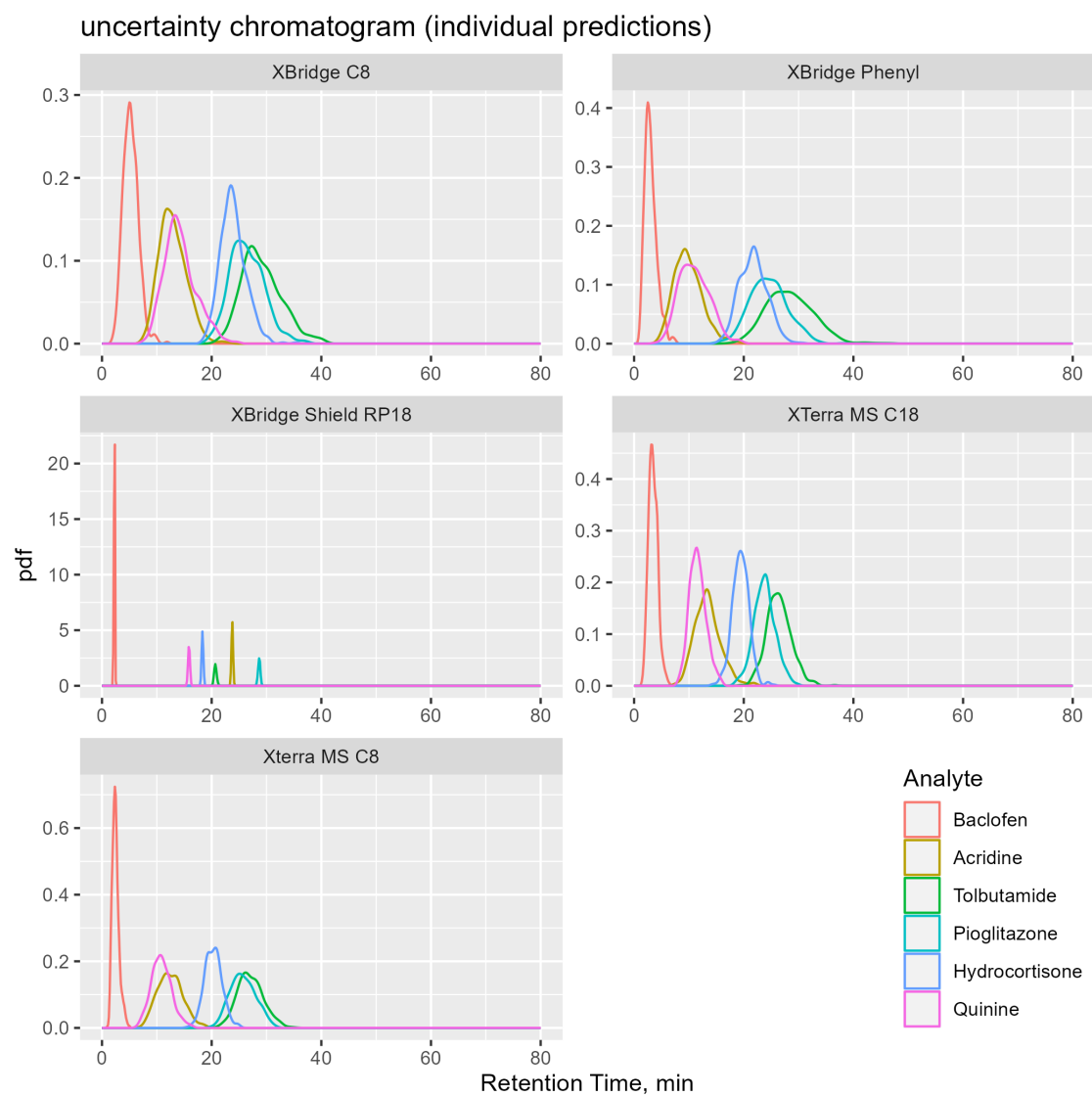
  labs(title="uncertainty chromatogram (individual predictions)",
```

```

x ="Retention Time, min",
y = "pdf",
colour="Analyte")+
xlim(c(0,80))+
facet_wrap(.~Column, nrow=3,scales = "free")+
theme(legend.position = c(1, 0),
legend.justification = c(1, 0))

print(p)

```



```
ggsave(paste0("figures\\casestudy2\\utilitymap\\", "chromatogram", ".png"), plot=p, width=10, height=10)
```

13 Conclusions

The large gradient retention time data and multilevel modeling allow to characterize separation of RP HPLC stationary phases.

References

- Kamedulska, Agnieszka, Łukasz Kubik, Julia Jacyna, Wiktoria Struck-Lewicka, Michał J. Markuszewski, and Paweł Wiczling. 2022. “Toward the General Mechanistic Model of Liquid Chromatographic Retention.” *Analytical Chemistry* 94 (31): 11070–80. <https://doi.org/10.1021/acs.analchem.2c02034>.
- Kamedulska, Agnieszka, Łukasz Kubik, and Paweł Wiczling. 2022. “Statistical Analysis of Isocratic Chromatographic Data Using Bayesian Modeling.” *Analytical and Bioanalytical Chemistry* 414 (11): 3471–3348. <https://doi.org/10.1007/s00216-022-03968-x>.
- Kubik, Łukasz, Julia Jacyna, Wiktoria Struck-Lewicka, Michał J. Markuszewski, and Paweł Wiczling. 2022a. “LC-TOF-MS Data Collected for 300 Small Molecules. XBridge Shield RP18 Column.” *Osf.io/1* (1): 1. <https://doi.org/10.17605/OSF.IO/ZQTJ7>.
- . 2022b. “LC-TOF-MS Data Collected for 300 Small Molecules. XBridge-C8 Column.” *Osf.io/1* (1): 1. <https://doi.org/10.17605/OSF.IO/Y6S8P>.
- . 2022c. “LC-TOF-MS Data Collected for 300 Small Molecules. XTerra MS C18 Column.” *Osf.io/1* (1): 1. <https://doi.org/10.17605/OSF.IO/QBV7J>.
- . 2022d. “LC-TOF-MS Data Collected for 300 Small Molecules. XTerra-C8 Column.” *Osf.io/1* (1): 1. <https://doi.org/10.17605/OSF.IO/2MCNW>.
- . 2022e. “LC-TOF-MS Data Collected for 300 Small Molecules. XBridge Phenyl Column.” *Osf.io/1* (1): 1. <https://doi.org/10.17605/OSF.IO/EVUJ9>.
- Kubik, Łukasz, Roman Kalisz, and Paweł Wiczling. 2018. “Analysis of Isocratic-Chromatographic-Retention Data Using Bayesian Multilevel Modeling.” *Analytical Chemistry* 90 (22): 13670–79. <https://doi.org/10.1021/acs.analchem.8b04033>.
- Nikitas, P., and A. Pappa-Louisi. 2002. “New Equations Describing the Combined Effect of pH and Organic Modifier Concentration on the Retention in Reversed-Phase Liquid Chromatography.” *Journal of Chromatography. A* 971 (1-2): 47–60. [https://doi.org/10.1016/s0021-9673\(02\)00965-2](https://doi.org/10.1016/s0021-9673(02)00965-2).
- Wiczling, Paweł, Agnieszka Kamedulska, and Łukasz Kubik. 2021. “Application of Bayesian Multilevel Modeling in the Quantitative Structure–Retention Relationship Studies of Heterogeneous Compounds.” *Analytical Chemistry* 93 (18): 6961–71. <https://doi.org/10.1021/acs.analchem.0c05227>.

Licenses

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Original Computing Environment

```
sessionInfo()

R version 4.1.3 (2022-03-10)
Platform: x86_64-w64-mingw32/x64 (64-bit)
Running under: Windows 10 x64 (build 22621)

Matrix products: default

locale:
[1] LC_COLLATE=Polish_Poland.1250  LC_CTYPE=Polish_Poland.1250
[3] LC_MONETARY=Polish_Poland.1250 LC_NUMERIC=C
[5] LC_TIME=Polish_Poland.1250

attached base packages:
[1] stats      graphics   grDevices utils      datasets   methods    base

other attached packages:
[1] kableExtra_1.3.4  GGally_2.1.2      posterior_1.3.1   bayesplot_1.9.0
[5] reshape2_1.4.4   knitr_1.40      cmdstanr_0.5.3   gridExtra_2.3
[9] ggplot2_3.4.2   dplyr_1.1.2      pracma_2.3.8

loaded via a namespace (and not attached):
[1] Rcpp_1.0.9          lattice_0.20-45    svglite_2.1.1
[4] tidyverse_1.3.0     zoo_1.8-11        digest_0.6.29
[7] utf8_1.2.2         R6_2.5.1          plyr_1.8.7
[10] ggridges_0.5.3     backports_1.4.1    coda_0.19-4
[13] evaluate_0.16      httr_1.4.5        pillar_1.9.0
[16] tidybayes_3.0.2    rlang_1.1.0        data.table_1.14.2
[19] rstudioapi_0.14    checkmate_2.1.0    rmarkdown_2.16
[22] textshaping_0.3.6  labeling_0.4.2     webshot_0.5.4
[25] stringr_1.5.0      munsell_0.5.0     compiler_4.1.3
[28] xfun_0.32          pkgconfig_2.0.3    systemfonts_1.0.4
[31] htmltools_0.5.3    tidyselect_1.2.0   tibble_3.2.1
```

```
[34] tensorA_0.36.2      arrayhelpers_1.1-0    matrixStats_0.62.0
[37] codetools_0.2-18    reshape_0.8.9       fansi_1.0.3
[40] viridisLite_0.4.1   withr_2.5.0        ggdist_3.2.0
[43] grid_4.1.3          distributional_0.3.1 jsonlite_1.8.4
[46] gtable_0.3.1        lifecycle_1.0.3    magrittr_2.0.3
[49] scales_1.2.1        cli_3.4.0         stringi_1.7.8
[52] farver_2.1.1        xml2_1.3.3        ragg_1.2.5
[55] generics_0.1.3      vctrs_0.6.2        RColorBrewer_1.1-3
[58] tools_4.1.3          svUnit_1.0.6       glue_1.6.2
[61] purrrr_1.0.1        abind_1.4-5        fastmap_1.1.0
[64] yaml_2.3.5          colorspace_2.0-3   isoband_0.2.5
[67] rvest_1.0.3
```