## [1] R code for calculation of uncertainty with MCM

```
# number of trials
M <- 10^6
# Parameters for the pdf of input quantities
uCr <- 0.003
uVp1000 <- 0.00075
uVp500 <- 0.00075
uVp100 <- 0.0001
uVp50 <- 0.0001
uVf10 <- 0.003
alpha <- 0.00149
a <- 25 - 5*alpha*10
b <- 25 + 5*alpha*10
# generate the volume
Vp1 <- rnorm(M, 1, uVp1000)
Vp0.5 <- rnorm(M, 0.5, uVp500)
Vp0.1 <- rnorm(M, 0.1, uVp100)
Vp0.05 <- rnorm(M, 0.05, uVp50)
Temp <- runif(M, 20, 30)
Vf \mathrel{<\!\!\!\!-} rnorm(M,\,10,\,uVf10)*(1+alpha*(Temp-25))\,\#\,temperature\,\,effect
# generate the reference material
Cr <- rnorm(M, 1, uCr)
# initialize concentrations
x0AP \leq rep(0, M)
x04HA \le rep(0, M)
# the peak area ratio of the analyte in the sample
y0.AP < -0.2
y0.4HA <- 0.021
# generate working standard solution
W1 <- Cr*rnorm(M, 1, uVp1000)/(rnorm(M, 10, uVf10)*(1+alpha*(Temp-25)))
```

```
W <- W1*rnorm(M, 1, uVp1000)/(rnorm(M, 10, uVf10)*(1+alpha*(Temp-25)))
# generate calibration standard solutions
C1 <- W*rnorm(M, 0.5, uVp500)/(rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500)+
                                      rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500))
C2 <- C1*rnorm(M, 1, uVp1000)/(rnorm(M, 1, uVp1000)+rnorm(M, 1, uVp1000))
C3 <- C2*(rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500))/(rnorm(M, 0.5, uVp500))+rnorm(M, 0.5,
uVp500)+rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500))
C4 \le C3*rnorm(M, 1, uVp1000)/(rnorm(M, 1, uVp1000)+rnorm(M, 1, uVp1000))
C5 < C4*rnorm(M, 1, uVp1000)/(rnorm(M, 1, uVp1000)+rnorm(M, 1, uVp1000))
C6 \leftarrow C5*(rnorm(M, 0.5, uVp500) + rnorm(M, 0.5, uVp500))/(rnorm(M, 0.5, uVp500) + rnorm(M, 0.5, uVp500))
uVp500)+rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500))
C7 <- C6*rnorm(M, 1, uVp1000)/(rnorm(M, 1, uVp1000)+rnorm(M, 1, uVp1000))
C8 <- C7*(rnorm(M,\, 0.5,\, uVp500) + rnorm(M,\, 0.5,\, uVp500))/(rnorm(M,\, 0.5,\, uVp500) + rnorm(M,\, 0.5,\, uVp500)) + rnorm(M,\, 0.
uVp500)+rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500)+rnorm(M, 0.5, uVp500))
# dilution of calibration standard solutions
X1 <- C1*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X2 < C2*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X3 <- C3*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X4 < C4*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X5 <- C5*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X6 <- C6*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X7 <- C7*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50)+rnorm(M, 0.05, uVp50)+rnorm(M, 0.1, uVp100))
X8 < -C8*rnorm(M, 0.1, uVp100)/(rnorm(M, 0.05, uVp50) + rnorm(M, 0.05, uVp50) + rnorm(M, 0.1, uVp100))
```

<sup>#</sup> generate y based on the data for calibration curves

```
## dat.AP cal: the data for calibration curves
## fit.AP: the fitted calibration curves: yi \sim a + b*xi + ei, var(ei) = (sigma*xi)^2
dat.AP cal <- read excel("~/Dropbox/2021 Research/검찰청/Output/calibration.xlsx",
  sheet = "AP")
dat.4HA_cal <- read_excel("~/Dropbox/2021_Research/검찰청/Output/calibration.xlsx",
  sheet = "4HA")
cal AP y <- read.csv("~/Dropbox/2021 Research/검찰청/Output/cal AP y.txt")
cal 4HA y <- read.csv("~/Dropbox/2021 Research/검찰청/Output/cal 4HA y.txt")
y1 <- rnorm(M, dat.AP cal$y[1], sqrt(cal AP y$var[1]))
y2 <- rnorm(M, dat.AP_cal$y[2], sqrt(cal_AP_y$var[2]))
y3 <- rnorm(M, dat.AP_cal$y[3], sqrt(cal_AP_y$var[3]))
y4 <- rnorm(M, dat.AP_cal$y[4], sqrt(cal_AP_y$var[4]))
y5 <- rnorm(M, dat.AP_cal$y[5], sqrt(cal_AP_y$var[5]))
y6 <- rnorm(M, dat.AP_cal$y[6], sqrt(cal_AP_y$var[6]))
y7 <- rnorm(M, dat.AP cal$y[7], sqrt(cal AP y$var[7]))
for(i in 1:M){
 AP < -data.frame(x = 1000000*rev(c(X1[i], X2[i], X3[i], X4[i], X5[i], X6[i], X7[i])),
           y= c(y1[i], y2[i], y3[i], y4[i], y5[i], y6[i], y7[i]))
 fit.AP <- lm(y\simx, weight=1/x^2, data=AP)
 x0AP[i] <- (y0.AP - fit.AP$coefficients[1])/fit.AP$coefficients[2]
}
y1 <- rnorm(M, dat.4HA_cal$y[1], sqrt(cal_4HA_y$var[1]))
y2 <- rnorm(M, dat.4HA_cal$y[2], sqrt(cal_4HA_y$var[2]))
y3 <- rnorm(M, dat.4HA_cal$y[3], sqrt(cal_4HA_y$var[3]))
y4 \le rnorm(M, dat.4HA\_cal\$y[4], sqrt(cal\_4HA\_y\$var[4]))
y5 <- rnorm(M, dat.4HA_cal$y[5], sqrt(cal_4HA_y$var[5]))
y6 \le rnorm(M, dat.4HA\_cal\$y[6], sqrt(cal\_4HA\_y\$var[6]))
y7 <- rnorm(M, dat.4HA_cal$y[7], sqrt(cal_4HA_y$var[7]))
for(i in 1:M){
 HA \leftarrow data.frame(x = 1000000*rev(c(X2[i], X3[i], X4[i], X5[i], X6[i], X7[i], X8[i])),
```

```
y=c(y1[i], y2[i], y3[i], y4[i], y5[i], y6[i], y7[i]))
 fit.HA <- lm(y\simx, weight=1/x^2, data=HA)
 x04HA[i] <- (y0.4HA - fit.HA$coefficients[1])/fit.HA$coefficients[2]
# relative uncertainty from the GUM
urD <- 0.000866
urR.AP <- 0.0193553
urR.4HA <- 0.0285052
# AP accounting for the uncertainty from method repeatability and urine sample dilution
x0AP.t <- x0AP + rnorm(M, 0, urR.AP*x0.AP) + rnorm(M, 0, urD*x0.AP)
#mean,median
mean(x0AP.t)
median(x0AP.t)
# standard deviation
sd(x0AP.t)
# 95% CI
quantile(x0AP.t, c(0.025, 0.975))
# 4HA accounting for the uncertainty from repeatability and dilution
x0HA.t <- x04HA + rnorm(M, 0, urR.4HA*x0.4HA) + rnorm(M, 0, urD*x0.4HA)
#mean,median
mean(x0HA.t)
median(x0HA.t)
# standard deviation
sd(x0HA.t)
# 95% CI
quantile(x0HA.t, c(0.025, 0.975))
```

## [2] R code for Figure 2

```
library(gridExtra)
library(tidyverse)
library(ggpubr)
library(cowplot)
res <- data.frame(AP = x0AP.t, HA = x0HA.t)

p1 <- ggplot(res, aes(x = AP,
y = ..density.., )) + labs(x="Concentration (ng/mL)", y = "Density", size=1) +
geom_histogram(alpha = 0.5, fill="#00AFBB") +
geom_density(alpha=0.2, size=1.2)

p2 <- ggplot(res, aes(x = HA,
y = ..density..)) + labs(x="Concentration (ng/mL)", y = "Density") +
geom_histogram(alpha = 0.5, fill="#E7B800") +
geom_histogram(alpha=0.2, size=1.2)

p3 <- ggarrange(p1, p2, nrow=1, labels=c("AP", "4HA"), label.x=0.8)
p3
```