

Mass Discharge - Outlet Alteck. 2016

PAZ

27 octobre 2016

Purpose

This file computes the discharged mass observed at the outlet. To do that it imports the weekly discharge summary and lab results for isotopes (^{13}C) and s-metolachlor concentrations.

Imports:

- WeeklyHydro_R.csv
- fluxAlteck2016_R.csv
- OutletConc_W0toW17.csv
- MESAlteckWater.csv
- Outlet_Isotopes_W0toW17.csv
- MESAlteck_FilterIsotopes.csv
- AO-Hydrochem.csv

Generates:

- WeeklyHydroContam_R.csv

Required R-packages:

```
library("plyr")
library("dplyr")
```

Working directory

```
# setwd("D:/Documents/these_pablo/Alteckendorf2016/R")
# setwd("/Users/DayTightChunks/Documents/PhD/Routput/Alteck/R")
# setwd("D:/Documents/these_pablo/Alteckendorf2016/00_TransparencyFolder")
getwd()
```

```
## [1] "D:/Documents/these_pablo/Alteckendorf2016/HydrologicalMonitoring"
```

Outlet Data - Alteckendorf 2016

1. Hydrological data on a subweekly basis

```
weeklyhydro = read.csv2("Data/WeeklyHydro_R.csv", header = TRUE)
colnames(wweeklyhydro)[colnames(wweeklyhydro) == "ID"] <- "WeekSubWeek"
head(wweeklyhydro)
```

```
## WeekSubWeek AveDischarge.m3.h Volume.m3 Sampled.Hrs Sampled
## 1 W0-0x 1.204775 14.41714 11.96667 Not Sampled
## 2 W0-1 1.213511 100.15508 82.53333 Sampled
## 3 W0-2x 1.284719 48.34827 37.63333 Not Sampled
## 4 W1-1 14.316647 390.36726 27.26667 Sampled
## 5 W1-2 15.529299 359.24445 23.13333 Sampled
## 6 W1-3x 9.107720 877.37700 96.33333 Not Sampled
```

```
weeklyflux = read.csv2("Data/fluxAlteck2016_R.csv", header = TRUE)
head(weeklyflux)
```

```
## WeekSubWeek ti tf iflux fflux
## 1 W0-0x 2016-03-25 00:04:00 2016-03-25 12:02:00 1.248600 1.129227
## 2 W0-1 2016-03-25 12:04:00 2016-03-28 22:36:00 1.124382 1.313125
## 3 W0-2x 2016-03-28 22:38:00 2016-03-30 12:16:00 1.308100 1.456349
## 4 W1-1 2016-03-30 12:18:00 2016-03-31 15:34:00 1.456080 16.445436
## 5 W1-2 2016-03-31 15:36:00 2016-04-01 14:44:00 16.334349 15.184536
## 6 W1-3x 2016-04-01 14:46:00 2016-04-05 15:06:00 15.203629 5.856380
## changeflux peak valley tdiff chExtreme
## 1 -0.1193728 1.248600 1.118296 11.96667 -0.1303036
## 2 0.1887431 1.380388 1.082199 82.53333 0.2560062
## 3 0.1482496 1.637782 0.929055 37.63333 0.3296817
## 4 14.9893566 38.399790 1.448977 27.26667 36.9437102
## 5 -1.1498131 18.668972 13.201113 23.13333 -3.1332355
## 6 -9.3472489 15.895640 5.471042 96.33333 -9.7325862
```

2. Concentration data (dissolved and suspended solids)

```
outletConc = read.csv2("Data/OutletConc_W0toW17.csv", header = T)
outletConc <- outletConc[outletConc$ID4 != "J+7", ]
outletConc <- outletConc[,c("WeekSubWeek", "Conc.mug.L", "Conc.SD")]
head(outletConc)
```

```
## WeekSubWeek Conc.mug.L Conc.SD
## 1 W0-1 0.2456594 0.01931
## 2 W1-1 6.7882463 0.28942
## 3 W1-2 6.5609982 0.19064
## 4 W2-1 9.4443019 0.33354
## 5 W2-2 1.0421883 0.03904
## 6 W3-1 8.8357358 0.47086
```

```
filters = read.csv2("Data/MESAlteckWater.csv")
filters$M0.mg.L = ifelse(filters$M0.mg.L < 0, 0.0001, filters$M0.mg.L)
head(filters)
```

```
## WeekSubWeek MES.mg.L MES.sd M0.mg.L Conc.Solids.mug.gMES
## 1 W0-1 53.44444 NA 0.0000 0.64472899
## 2 W1-1 62.50000 NA 0.0010 0.12588974
## 3 W1-2 22.50000 NA 0.0001 0.43578716
## 4 W2-1 22.50000 NA 0.0001 0.07935267
## 5 W2-2 5.00000 NA 0.0001 0.05075270
## 6 W3-1 197.50000 NA 0.0058 0.08177487
```

3. Isotope data

```
# Outlet isotope data:
outletIso = read.csv2("Data/Outlet_Isotopes_W0toW17.csv", header = T)
head(outletIso)
```

```
## FileHeader..Filename ID Week Wnum SubWeek WeekSubWeek Repl d.13C.12C
## 1 AO_W0_1-1.dxf AO W0 0 1 W0-1 1 -26.035
## 2 AO_W0_1-2.dxf AO W0 0 1 W0-1 2 -27.740
## 3 AO_W0_1-3_-0001.dxf AO W0 0 1 W0-1 3 -26.219
## 4 AO_W1_1-1_-0001.dxf AO W1 1 1 W1-1 1 -30.591
## 5 AO_W1_1-2_-0001.dxf AO W1 1 1 W1-1 2 -30.411
## 6 AO_W1_1-3_-0001.dxf AO W1 1 1 W1-1 3 -30.404
```

```
# Filter isotope data:
filtersIso = read.csv2("Data/MESAlteck_FilterIsotopes.csv", header = T)
filtersIso$WeekSubWeek = paste(filtersIso$Week, filtersIso$Num, sep = "-")
filtersIso <- filtersIso[filtersIso$Levl != "J+7", ]
head(filtersIso)
```

```
## ID Week Wnum Num Levl Repl d.13C.12C WeekSubWeek
## 1 AFP W2 1 1 1 -25.154 W2-1
## 2 AFP W2 1 1 2 -28.187 W2-1
## 3 AFP W2 1 1 3 -28.283 W2-1
## 4 AFP W2 2 2 1 -30.618 W2-2
## 5 AFP W2 2 2 2 -26.304 W2-2
## 6 AFP W2 2 2 3 -26.024 W2-2
```

4. Hydrochemistry Data

```
hydroChem = read.csv2("Data/A0-Hydrochem.csv", header = T)
hydroChem = hydroChem[, c("WeekSubWeek",
                          "NH4.mM",
                          "TIC.ppm.filt",
                          "Cl.mM",
                          "NO3...mM",
                          "PO4..mM",
                          "NPOC.ppm" ,
                          "TIC.ppm.unfilt",
                          "TOC.ppm.unfilt" )]
head(hydroChem)
```

```
## WeekSubWeek NH4.mM TIC.ppm.filt Cl.mM NO3...mM PO4..mM NPOC.ppm
## 1 W1-1 0.05 51.8 1.48 616.00 NA 4.0
## 2 W1-2 NA 44.8 1574.00 778.00 NA 4.4
## 3 W10-1 NA 60.1 1.17 964.00 NA 2.0
## 4 W10-2 9.00 57.1 1013.00 1174.00 13 5.2
## 5 W10-3 NA 58.2 858.00 1.23 NA 5.0
## 6 W10-4 15.00 26.4 355.00 1409.00 NA 6.4
## TIC.ppm.unfilt TOC.ppm.unfilt
## 1 44.8 4.7
## 2 26.4 5.4
## 3 63.2 2.0
## 4 55.9 4.0
## 5 60.4 4.3
## 6 24.5 6.4
```

Summarizing IRMS data

```
isoOutSummary = ddply(outletIso, c("WeekSubWeek"), summarise,
                      N = length(d.13C.12C),
```

```

      diss.d13C = mean(d.13C.12C),
      SD.d13C = sd(d.13C.12C),
      se.d13C = SD.d13C / sqrt(N))

head(isoOutSummary)

##   WeekSubWeek N diss.d13C   SD.d13C   se.d13C
## 1      W0-1 3 -26.66467 0.9357993 0.54028398
## 2      W1-1 3 -30.46867 0.1060016 0.06120004
## 3      W1-2 3 -30.61967 0.1513550 0.08738484
## 4     W10-1 2 -29.47350 1.9905056 1.40750000
## 5     W10-2 3 -29.27067 0.6003202 0.34659502
## 6     W10-3 3 -29.76967 0.3411749 0.19697744

isoFiltSummary = ddply(filtersIso, c("WeekSubWeek"), summarise,
      N      = length(d.13C.12C),
      filt.d13C = mean(d.13C.12C),
      filt.SD.d13C = sd(d.13C.12C),
      filt.se.d13C = filt.SD.d13C / sqrt(N))

head(isoFiltSummary)

##   WeekSubWeek N filt.d13C filt.SD.d13C filt.se.d13C
## 1      W2-1 3 -27.20800    1.779464    1.0273738
## 2      W2-2 3 -27.64867    2.575326    1.4868653
## 3      W6-3 3 -28.00667    1.593462    0.9199856
## 4      W9-1 2 -26.79150    1.745847    1.2345000
## 5      W9-2 3 -27.69633    2.013989    1.1627772
## 6      W9-3 3 -26.94633    1.685361    0.9730434

```

Merging and data wrangling steps

1. Merge all data sets by the *WeekSubWeek* column ID, including:

```

# Dissolved
out.CoIs = merge(outletConc, isoOutSummary, by = "WeekSubWeek", all = T)

# Filters (MES, Conc.MES)
out.CoIs = merge(out.CoIs, filters, by = "WeekSubWeek", all = T)
out.CoIs = merge(out.CoIs, isoFiltSummary, by = "WeekSubWeek", all = T)

# Pure and cuve isotope average
d13Co = -31.21

# Lab enrichment:
# epsilon = -1.61

# Lab enrichment:
# Alteck
epsilon_max = -1.5 # +/- 0.3 (@ 20C, 20% vwc)
epsilon_min = -2.0 # +/- 0.2 (@ 20C, 40% vwc)
epsilon_mean = -1.75

# Remaining fraction
out.CoIs$DD13C.diss <- (out.CoIs$diss.d13C - (d13Co))

```

```

out.CoIs$DD13C.filt <- (out.CoIs$filt.d13C - (d13Co))

out.CoIs$f.diss <- (((10**(-3)*out.CoIs$diss.d13C + 1)/(10**(-3)*d13Co + 1))**(1000/(epsilon_mean)))
out.CoIs$f.filt <- (((10**(-3)*out.CoIs$filt.d13C + 1)/(10**(-3)*d13Co + 1))**(1000/(epsilon_mean)))

out.CoIs$B.diss <- (1 - out.CoIs$f.diss)*100
out.CoIs$B.filt <- (1 - out.CoIs$f.filt)*100
#out.CoIs$invf <- 1/out.CoIs$f

# Discharge times
out.CoIs = merge(weeklyhydro, out.CoIs, by = "WeekSubWeek", all = T)

# Discharge summary
out.CoIs = merge(weeklyflux, out.CoIs, by = "WeekSubWeek", all = T)

# Hydrochemistry
out.CoIs = merge(out.CoIs, hydroChem, by= "WeekSubWeek", all = T)

out.CoIs$tf <- as.POSIXct(out.CoIs$tf, "%Y-%m-%d %H:%M", tz = "EST")
out.CoIs$ti <- as.POSIXct(out.CoIs$ti, "%Y-%m-%d %H:%M", tz = "EST")
class(out.CoIs$tf)

```

```
## [1] "POSIXct" "POSIXt"
```

```
sum(is.na(out.CoIs$tf))
```

```
## [1] 4
```

```
# Temporarily remove Weeks 16 & 17 (need to get discharge data)
```

```
out.CoIs <- out.CoIs[!is.na(out.CoIs$tf), ]
```

2. Weekly Exported Solids (Kg)

```
# V[m3] * MES [mg/L] * 1000 [L/m3] * [1 Kg/10^6 mg]
```

```
out.CoIs$ExpMES.Kg = out.CoIs$Volume.m3*out.CoIs$MES.mg.L/1000
```

3. Weekly exported S-metolachlor mass (mg)

```
# Dissolved - [mg] S-metolachlor exported per sub-week
```

```
# Conc. [mu.g s-meto/L H2O] * Vol[m3] * [10^3 L/m^3] * [1 mg/10^3 mu.g]
```

```
out.CoIs$DissSmeto.mg = out.CoIs$Conc.mug.L*out.CoIs$Volume.m3
```

```
# Solids - [mg] S-metolachlor in solids exported per sub-week
```

```
# Conc. [mu.g s-meto / g MES] * Kg MES * [10^3 g/Kg] * [1 mg/10^3 mu.g]
```

```
out.CoIs$FiltSmeto.mg = out.CoIs$Conc.Solids.mug.gMES*out.CoIs$ExpMES.Kg
```

```
# Total
```

```
out.CoIs$TotMassOut.mg = out.CoIs$DissSmeto.mg + out.CoIs$FiltSmeto.mg
```

```
# Proportion in dissolved and suspended solids
```

```
out.CoIs$FracDiss = out.CoIs$DissSmeto.mg/out.CoIs$TotMassOut.mg
```

```
out.CoIs$FracFilt = out.CoIs$FiltSmeto.mg/out.CoIs$TotMassOut.mg
```

4. Add the application dates and merge the total mass to the nearest discharge event

The five application dates were:

- 2016-03-20
- 2016-04-05
- 2016-04-13 and 2016-04-14
- 2016-05-26

So the total applied mass mass is merged at the nearest sampling time marker available :

```
ti = c(as.POSIXct('2016-03-25 00:04:00' , tz="EST"),
      as.POSIXct('2016-04-05 15:08:00' , tz="EST"),
      as.POSIXct('2016-04-14 13:52:00' , tz="EST"),
      as.POSIXct('2016-05-24 12:00:00' , tz="EST"))

Appl.Mass.g = c(6369.396, 3128.475, 4744.571, 4982.038)

applics = as.data.frame(ti)
applics$Appl.Mass.g = Appl.Mass.g

out.CoIs = merge(out.CoIs, applics, by = "ti", all = T)
out.CoIs$Appl.Mass.g <- ifelse(is.na(out.CoIs$Appl.Mass.g), 0.0, out.CoIs$Appl.Mass.g)
```

5. This section converts the observed S-metolachlor concentrations to [g] in dissolved water and suspended solids, assuming 0 for the values where no sample was taken. An approximative model will be tested at a later stage.

```
# Cumulative IN
out.CoIs$CumAppMass.g = cumsum(out.CoIs$Appl.Mass.g)

# First simulate a mass out to deal with missing values
# Option 1, just assume 0.0
out.CoIs$SimOutDiss.g = out.CoIs$DissSmeto.mg/10^3
out.CoIs$SimOutFilt.g = out.CoIs$FiltSmeto.mg/10^3
out.CoIs$SimOutDiss.g = ifelse(is.na(out.CoIs$SimOutDiss.g), 0.0, out.CoIs$SimOutDiss.g)
out.CoIs$SimOutFilt.g = ifelse(is.na(out.CoIs$SimOutFilt.g), 0.0, out.CoIs$SimOutFilt.g)
out.CoIs$SimOutSmeto.g = out.CoIs$SimOutDiss.g + out.CoIs$SimOutFilt.g

# Cumulative OUT
out.CoIs$CumOutDiss.g = cumsum(out.CoIs$SimOutDiss.g)
out.CoIs$CumOutFilt.g = cumsum(out.CoIs$SimOutFilt.g)
out.CoIs$CumOutSmeto.g = out.CoIs$CumOutDiss.g + out.CoIs$CumOutFilt.g

# Balance
out.CoIs$BalMassDisch.g = out.CoIs$CumAppMass.g - out.CoIs$CumOutSmeto.g

# Mass fraction
massOUT = tail(out.CoIs$CumOutSmeto.g, n=1)
out.CoIs$FracMassOut = (out.CoIs$SimOutSmeto.g / massOUT)
out.CoIs$FracDeltaOut = (out.CoIs$SimOutSmeto.g / massOUT)*out.CoIs$diss.d13C
out.CoIs$FracDeltaOut = ifelse(is.na(out.CoIs$FracDeltaOut), 0.0, out.CoIs$FracDeltaOut)

BulkDeltaOut = sum(out.CoIs$FracDeltaOut)
```

The total mass discharged (up to Week 15) and bulk isotope signature (up to week 11) was:

```
# Cumulative S-metolachlor [g] discharged
massOUT
```

```
## [1] 91.10687
```

```
# Bulk isotope signature
```

```
BulkDeltaOut
```

```
## [1] -23.8942
```

6. Testing a regression tree (ommitted for now)

Save files

```
head(out.CoIs)
```

```
##          ti WeekSubWeek          tf      iflux      fflux
## 1 2016-03-25 00:04:00      W0-0x 2016-03-25 12:02:00  1.248600  1.129227
## 2 2016-03-25 12:04:00      W0-1 2016-03-28 22:36:00  1.124382  1.313125
## 3 2016-03-28 22:38:00      W0-2x 2016-03-30 12:16:00  1.308100  1.456349
## 4 2016-03-30 12:18:00      W1-1 2016-03-31 15:34:00  1.456080  16.445436
## 5 2016-03-31 15:36:00      W1-2 2016-04-01 14:44:00  16.334349  15.184536
## 6 2016-04-01 14:46:00      W1-3x 2016-04-05 15:06:00  15.203629  5.856380
##   changeflux      peak      valley      tdiff  chExtreme AveDischarge.m3.h
## 1 -0.1193728  1.248600  1.118296  11.96667 -0.1303036      1.204775
## 2  0.1887431  1.380388  1.082199  82.53333  0.2560062      1.213511
## 3  0.1482496  1.637782  0.929055  37.63333  0.3296817      1.284719
## 4 14.9893566 38.399790  1.448977  27.26667 36.9437102     14.316647
## 5 -1.1498131 18.668972 13.201113  23.13333 -3.1332355     15.529299
## 6 -9.3472489 15.895640  5.471042  96.33333 -9.7325862      9.107720
##   Volume.m3 Sampled.Hrs      Sampled Conc.mug.L Conc.SD N.x diss.d13C
## 1  14.41714    11.96667 Not Sampled      NA      NA  NA      NA
## 2 100.15508    82.53333   Sampled    0.2456594 0.01931  3 -26.66467
## 3  48.34827    37.63333 Not Sampled      NA      NA  NA      NA
## 4 390.36726    27.26667   Sampled    6.7882463 0.28942  3 -30.46867
## 5 359.24445    23.13333   Sampled    6.5609982 0.19064  3 -30.61967
## 6 877.37700    96.33333 Not Sampled      NA      NA  NA      NA
##   SD.d13C      se.d13C MES.mg.L MES.sd MO.mg.L Conc.Solids.mug.gMES N.y
## 1      NA      NA      NA      NA      NA      NA      NA
## 2 0.9357993 0.54028398 53.44444      NA 0e+00      0.6447290  NA
## 3      NA      NA      NA      NA      NA      NA      NA
## 4 0.1060016 0.06120004 62.50000      NA 1e-03      0.1258897  NA
## 5 0.1513550 0.08738484 22.50000      NA 1e-04      0.4357872  NA
## 6      NA      NA      NA      NA      NA      NA      NA
##   filt.d13C filt.SD.d13C filt.se.d13C DD13C.diss DD13C.filt      f.diss
## 1      NA      NA      NA      NA      NA      NA
## 2      NA      NA      NA  4.5453333      NA 0.06892489
## 3      NA      NA      NA      NA      NA      NA
## 4      NA      NA      NA  0.7413333      NA 0.64590754
## 5      NA      NA      NA  0.5903333      NA 0.70603206
## 6      NA      NA      NA      NA      NA      NA
##   f.filt      B.diss B.filt NH4.mM TIC.ppm.filt      Cl.mM NO3...mM PO4...mM
## 1      NA      NA      NA      NA      NA      NA      NA
## 2      NA 93.10751      NA      NA      NA      NA      NA
## 3      NA      NA      NA      NA      NA      NA      NA
## 4      NA 35.40925      NA  0.05      51.8      1.48      616      NA
## 5      NA 29.39679      NA      NA      44.8 1574.00      778      NA
```

## 6	NA	NA	NA	NA	NA	NA	NA	NA
##	NPOC.ppm	TIC.ppm.unfilt	TOC.ppm.unfilt	ExpMES.Kg	DissSmeto.mg			
## 1	NA	NA	NA	NA	NA	NA		
## 2	NA	NA	NA	NA	5.352733	24.60403		
## 3	NA	NA	NA	NA	NA	NA		
## 4	4.0	44.8	4.7	24.397953	2649.90908			
## 5	4.4	26.4	5.4	8.083000	2357.00221			
## 6	NA	NA	NA	NA	NA	NA		
##	FiltSmeto.mg	TotMassOut.mg	FracDiss	FracFilt	Appl.Mass.g			
## 1	NA	NA	NA	NA	6369.396			
## 2	3.451062	28.0551	0.8769898	0.123010164	0.000			
## 3	NA	NA	NA	NA	0.000			
## 4	3.071452	2652.9805	0.9988423	0.001157736	0.000			
## 5	3.522468	2360.5247	0.9985078	0.001492239	0.000			
## 6	NA	NA	NA	NA	0.000			
##	CumAppMass.g	SimOutDiss.g	SimOutFilt.g	SimOutSmeto.g	CumOutDiss.g			
## 1	6369.396	0.00000000	0.00000000	0.0000000	0.00000000			
## 2	6369.396	0.02460403	0.003451062	0.0280551	0.02460403			
## 3	6369.396	0.00000000	0.00000000	0.0000000	0.02460403			
## 4	6369.396	2.64990908	0.003071452	2.6529805	2.67451312			
## 5	6369.396	2.35700221	0.003522468	2.3605247	5.03151533			
## 6	6369.396	0.00000000	0.00000000	0.0000000	5.03151533			
##	CumOutFilt.g	CumOutSmeto.g	BalMassDisch.g	FracMassOut	FracDeltaOut			
## 1	0.00000000	0.0000000	6369.396	0.0000000000	0.000000000			
## 2	0.003451062	0.0280551	6369.368	0.0003079361	-0.008211013			
## 3	0.003451062	0.0280551	6369.368	0.0000000000	0.000000000			
## 4	0.006522514	2.6810356	6366.715	0.0291194331	-0.887230300			
## 5	0.010044982	5.0415603	6364.354	0.0259094025	-0.793337267			
## 6	0.010044982	5.0415603	6364.354	0.0000000000	0.000000000			

```

write.csv2(out.CoIs,
            'Data/WeeklyHydroContam_R.csv', row.names = F)

# out.CoIs = read.csv2("Data/WeeklyHydroContam_R.csv")
# out.CoIs$ti = as.POSIXct(out.CoIs$ti, "%Y-%m-%d %H:%M", tz = "EST")

```