Analysis of Model F

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 Analysis on Model F
suppressMessages(source("ams_initialize_script.R"))
suppressMessages(library(RxODE))
suppressMessages(library(dplyr))
```

Model F is defined below

```
ivsc_4cmtct_shedct = function() {
 model
                  = list()
  model$name
                  = as.character(sys.calls()[[sys.nframe()]])
  #COMPARTMENTS
  model$cmtshort = c('AmtD0','D1','D2','D3','S1','S3','M3','DS1','DS3','DM3','DM1','M1')
  #CALCULATE INITIAL CONDITION WITH NO DRUG PRESENT AND ASSUMING STEADY STATE
  model$init
                  = function(p){
             init = c(AmtD0=0,D1=0,D2=0,D3=0,S1=0,S3=0,M3=0,DS1=0,DS3=0,DM3=0,DM1=0,M1=0)
                  = p %>% t() %>% as.data.frame()
             ksyn = with(p,c(ksynS1,ksynS3,ksynM1,ksynM3))
                                                , -k31S*VS3/VS1
                                                                                                 0,
                  = with(p,matrix(c( keS1+k13S
                                                                          -kshedM1
                                    -k13S*VS1/VS3 , keS3+k31S
                                                                                                -kshedM3
                                                                                                -k31M*VD
                                                    0
                                                                           kshedM1+keM1+k13M
                                     0
                                                     0
                                                                          -k13M*VD1/VD3
                                                                                                 kshedM3
                                  nrow = 4, byrow=TRUE))
                  = solve(K,ksyn)
             init["S1"] = unlist(x[1])
```

```
init["S3"] = unlist(x[2])
            init["M1"] = unlist(x[3])
             init["M3"] = unlist(x[4])
            return(init)
  }
  #PARAMEETRS IN MODEL
                 = c('F','ka','VD1','VD2','VD3','VS1','VS3','VDS1','VDS3',
  model$pin
                      'k12D', 'k21D', 'k13D', 'k31D', 'k13S', 'k31S', 'k13DS', 'k31DS',
                      'ksynS1','ksynS3','ksynM3','keD1','keD3','keS1','keS3','keDS1','keDS3','keM3','ke
                      'kon1', 'koff1', 'kon3', 'koff3',
                      'kshedM3','kshedDM3','ksynM1','kshedM1','kshedDM1','keM1','keDM1','k13M','k31M','
  model$pode
                  = model$pin
                   #INPUT/SYNTHESIS/SHED
                                         DISTRIBUTION (CENTRAL/TUMOR)
                                                                            BINDING
  model$rxode.str = '
                 = AmtD1/VD1;
    d/dt(AmtD0) = -ka *AmtD0;
    d/dt(AmtD1) =(F*ka *AmtD0/VD1 - k13D *D1 + k31D *VD3/VD1*D3 - keD1 *D1 - kon1*D1*S1 + koff1*DS
     d/dt(D2)
                = k12D*VD1/VD2*D1 - k21D*D2;
    d/dt(D3)
                 = k13D *VD1/VD3*D1 - k31D*D3 - keD3 *D3 - kon3*D3*(S3+M3) + koff3*(DS3+DM3);
    d/dt(S1)
                = ksynS1+kshedM1*M1 - k13S *S1 + k31S*VS3/VS1*S3 - keS1 *S1 - kon1*D1*S1
                                                                                                   + ko
    d/dt(S3)
                 = ksynS3 +kshedM3*M3 + k13S *VS1/VS3*S1 - k31S*S3 - keS3 *S3 - kon3*D3*S3 + koff
                 = ksynM3 -kshedM3*M3 -k31M*M3+k13M*VD1/VD3*M1 - keM3 *M3 - kon3*D3*M3 + koff3*DM3;
    d/dt(M3)
    d/dt(DS1)
                 = kshedDM1*DM1 - k13DS*DS1 + k31DS*VDS3/VDS1*DS3 - keDS1*DS1 + kon1*D1*S1 - koff1*DS1
    d/dt(DS3) = kshedDM3*DM3 + k13DS*VDS1/VDS3*DS1 - k31DS*DS3 - keDS3*DS3 + kon3*D3*S3 - koff3*DS
    d/dt(DM3) = -kshedDM3*DM3 - keDM3*DM3 + kon3*D3*M3 - koff3*DM3-k31DM*DM3+k13DM*(VD1/VD3)*DM1;
                = -keDM1*DM1 -kshedDM1*DM1 +kon1*D1*M1 -koff1*DM1-k13DM*DM1+k31DM*(VD3/VD1)*DM3;
    d/dt(DM1)
    d/dt(M1)
                 = ksynM1 - kshedM1*M1 - keM1*M1 + k31M*VD3/VD1*M3 - k13M*M1 - kon1*D1*M1 + koff1*DM1;
  model$rxode
                 = RxODE(model = model$rxode.str, modName = model$name)
  model $rxout
                 = function(result)
   result
                 = as.data.frame(result)
   result = mutate(result,
                   Dtot1 = D1+DS1,
                   Stot1 = S1+DS1,
                   Dtot3 = D3+DS3,
                   Stot3 = S3+DS3,
                   Mtot1 = M1+DM1,
                   Mtot3 = M3+DM3)
  }
 return(model)
}
# Global Variables
model = ivsc_4cmtct_shedct()
tmax = 3*28 # End of the observation time, unit=day
tau = 14 # dosing interval, unit=day
compartment = 2 # compartment to which dosing is applied
# Import parameters
d = readxl::read_excel("../data/ivsc_4cmtct_shedct_param.xlsx",1)
```

```
param.as.double = d$Value
names(param.as.double) = d$Parameter
# Helper function to make the range of a variable when performing sensitivity analysis
lseq = function(from, to, length.out){
    sequence = seq(log(from), log(to), length.out=length.out)
    sequence = exp(sequence)
   return(sequence)
}
simulation = function(dose.nmol, params_file_path, tmax){
  d <- xlsx::read.xlsx(params file path, 1)</pre>
  param.as.double = d$Value
  names(param.as.double) = d$Parameter
  ev = eventTable(amount.units="nmol", time.units="days")
  sample.points = c(seq(-7, tmax, 0.1), 10^{\circ}(-3:0)) # sample time, increment by 0.1
  sample.points = sort(sample.points)
  sample.points = unique(sample.points)
  ev$add.sampling(sample.points)
  ev$add.dosing(dose=dose.nmol, nbr.doses=floor(tmax/tau)+1, dosing.interval=tau,
                dosing.to=2)
  init = model$init(param.as.double)
  out = model$rxode$solve(param.as.double, ev, init)
  out = model$rxout(out)
  out = out %>%
   mutate(Sfree.pct = S1/init["S1"],
             Mfree.pct = M3/init["M3"],
             dose.nmol = dose.nmol)
  return(out)
```

Test the theory and simulation of the lumped parameters: M30, M3tot.ss, B, Davg3

The function below takes a dataset and calculate these lumped parameters from theory

```
lumped.parameters.theory = function(params_file_path, dose.nmol){
    d <- xlsx::read.xlsx(params_file_path, 1)
    param.as.double = d$Value
    names(param.as.double) = d$Parameter
    p = as.data.frame(t(param.as.double))
    Kss = with(p, (koff3 + keDM3 + kshedM3)/kon3)
    Kd = with(p, koff3 / kon3)

# numerators for M3.0 and Mtot3.ss(Mtot3 at steady state, M3 at initial state)
    numerator.DM = with(p, k13DM*(VD1/VD3)*ksynM1+(keDM1+kshedDM1+k13DM)*ksynM3)
    denomenator.DM = with(p, (keDM1+kshedDM1+k13DM)*(keDM3+kshedM3+k31DM)-k31DM*k13DM)
    numerator.M = with(p, k13M *(VD1/VD3)*ksynM1+(keM1 +kshedM1 +k13M) *ksynM3)</pre>
```

```
denomenator.M = with(p, (keM1 +kshedM1 +k13M) *(keM3 +kshedM3+k31M) -k31M *k13M)
    # numerator and denomenator for M3.0 ()
   Mtot3.ss = numerator.DM / denomenator.DM
   M30
            = numerator.M / denomenator.M
    # Target accumulation in the tumor compartment
   Tacc.tum = Mtot3.ss / M30
    # Biodistribution coefficient (reference: ModelF_Appendix)
   B = with(p, (k13D/(keD3 + k31D) * (VD1/VD3)))
    # Clearance
   CL = with(p, (keD1*VD1))
    # Average drug concentration in the central compartment
    \# Cavg1 = with(p, (F * dose.nmol) / (CL * tau))
    # Average drug concentratio in the tumor compartment (I have no idea how to compute it)
    # AFIRT computed with Kss
   AFIRT.theory.Kss = Kss*Tacc.tum*(CL*tau)/(dose.nmol*B)
    # AFIRT computed with Kd
   AFIRT.theory.Kd = Kd*Tacc.tum*(CL*tau)/(dose.nmol*B)
   lumped_parameters_theory = data.frame(type = "theory",
                                          M30 = M30,
                                          Mtot3.ss=Mtot3.ss,
                                          Tacc.tum=Tacc.tum,
                                          B = B,
                                          CL = CL,
                                          AFIRT.Kss = AFIRT.theory.Kss,
                                          AFIRT.Kd = AFIRT.theory.Kd)
   return(lumped_parameters_theory)
}
```

Calculate lumped parameters from theory for Atezolizumab

```
lumped_parameters_theory = lumped.parameters.theory("../data/ModelF_Atezolizumab_Params.xlsx", 80)
print(lumped_parameters_theory)

## type M30 Mtot3.ss Tacc.tum B CL AFIRT.Kss AFIRT.Kd
## 1 theory 2.551332 2.551332 1 0.3333333 0.2 0.525 0.2625
```

Okay, for different inital doses, we do get the same simulated lumped parameters. However, M30 seems to be the same as Mtot3.ss in the simulation which is kind of weird. Also, M30 calculated from theory and M30

calculated from simulation are off by a lot.

- Andy says: the parameetrs are such that M3tot doesn't accumulate much. So this makes sense. We compare the theory and simulation below.
- kable is not working for lumped_parameters_sim or lumped_parameters_theory.
- Hongshan says: AFIRT.sim and AFIRT.theory still differ by a big margin

Sensitivity analysis on lumped parameters calculated from theory with respect to dose.mol

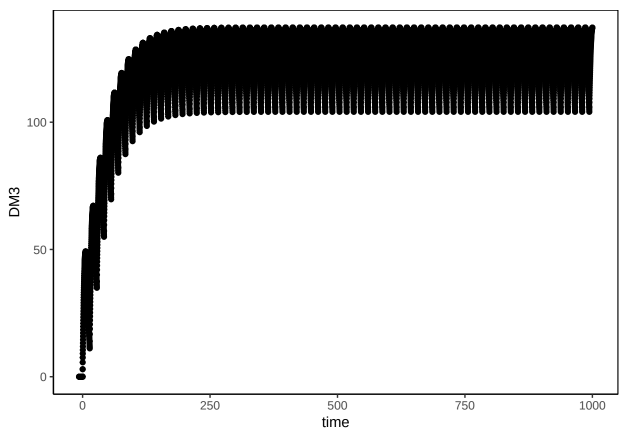
```
dose.nmol.range = lseq(1, 1000, 10)
table = data.frame()
for (dose.nmol in dose.nmol.range){
 row = lumped.parameters.theory("../data/ModelF_Atezolizumab_Params.xlsx", dose.nmol)
 table = rbind(table, row)
print(table)
##
                                                                   AFIRT.Kd
                                              B CL
                                                      AFIRT.Kss
                 M30 Mtot3.ss Tacc.tum
## 1 theory 2.551332 2.551332
                                   1 0.3333333 0.2 42.00000000 21.00000000
## 2 theory 2.551332 2.551332
                                    1 0.3333333 0.2 19.49467310 9.74733655
## 3 theory 2.551332 2.551332
                                   1 0.3333333 0.2 9.04862570 4.52431285
## 4 theory 2.551332 2.551332
                                    1 0.3333333 0.2 4.20000000 2.10000000
## 5 theory 2.551332 2.551332
                                    1 0.3333333 0.2 1.94946731 0.97473366
## 6 theory 2.551332 2.551332
                                    1 0.3333333 0.2 0.90486257
                                                                 0.45243128
## 7 theory 2.551332 2.551332
                                    1 0.3333333 0.2 0.42000000
                                                                 0.21000000
## 8 theory 2.551332 2.551332
                                    1 0.3333333 0.2 0.19494673
                                                                 0.09747337
                                    1 0.3333333 0.2 0.09048626 0.04524313
## 9 theory 2.551332 2.551332
## 10 theory 2.551332 2.551332
                                    1 0.3333333 0.2 0.04200000 0.02100000
```

The function below simulates the lumped parameter

```
# Change the function <simulation> so that it includes more time points
simulation = function(dose.nmol, params_file_path, tmax){
  d <- xlsx::read.xlsx(params_file_path, 1)</pre>
  param.as.double = d$Value
  names(param.as.double) = d$Parameter
  ev = eventTable(amount.units="nmol", time.units="days")
  sample.points = c(seq(-7, tmax, 0.1), 10^{(-3:0)}) # sample time, increment by 0.1
  sample.points = sort(sample.points)
  sample.points = unique(sample.points)
  ev$add.sampling(sample.points)
  ev$add.dosing(dose=dose.nmol, nbr.doses=floor(tmax/tau)+1, dosing.interval=tau,
                dosing.to=2)
  init = model$init(param.as.double)
  out = model$rxode$solve(param.as.double, ev, init)
  out = model$rxout(out)
  out = out %>%
   mutate(Sfree.pct = S1/init["S1"],
```

Run the simulation longer to make sure the system reaches steady state

```
sim = simulation(dose.nmol = 80, params_file_path ="../data/ModelF_Atezolizumab_Params.xlsx", tmax=1000
g = ggplot(sim, aes(x=time, y=DM3)) + geom_point()
print(g)
```



It is quite visible that the system reaches steady state after time > 500.

Simulate lumped parameters during steady state

```
lumped.parameters.simulation = function(params_file_path, dose.nmol, tmax){
    sim = simulation(dose.nmol=dose.nmol, params_file_path = params_file_path, tmax=tmax)
    initial_state = sim %>%
        filter(time==0)
    M30 = initial_state$M3

steady_state = sim %>%
        filter(time>(tmax/2) & time <tmax)
    Mtot3.ss = mean(steady_state$Mtot3)</pre>
```

```
Tacc.tum = Mtot3.ss / M30
  # Average drug concentration in central compartment
  dose applied = sim %>%
   filter(time > 0)
  Cavg1 = mean(dose_applied$D1)
  # Average drug concentration in tumor compartment
  Cavg3 = mean(dose_applied$D3)
  # AFIRT
  AFIRT.sim = mean(steady_state$Mfree.pct)
  lumped_parameters_sim = data.frame(type = "simulation",
                                     M30 = M30,
                                     Mtot3.ss=Mtot3.ss,
                                     Tacc.tum=Tacc.tum,
                                     Cavg1 = Cavg1,
                                     Cavg3 = Cavg3,
                                     AFIRT.sim = AFIRT.sim)
 return(lumped parameters sim)
}
lump_sim = lumped.parameters.simulation(params_file_path="../data/ModelF_Atezolizumab_Params.xlsx", dos
lump_sim
##
                     M30 Mtot3.ss Tacc.tum
                                                        Cavg3 AFIRT.sim
           type
                                              Cavg1
## 1 simulation 2.551332 132.6211 51.98112 10.46519 43.44386 2.484134
```

Sensitivity analysis on lumped parameters calculated from simulation with respect to dose.nmol

```
dose.nmol.range = lseq(1, 1000, 10)
table = data.frame()
for (dose.nmol in dose.nmol.range){
 row = lumped.parameters.simulation("../data/ModelF_Atezolizumab_Params.xlsx", dose.nmol, tmax=1000)
 table = rbind(table, row)
}
print(table)
##
           type
                     M30
                          Mtot3.ss
                                      Tacc.tum
                                                     Cavg1
                                                                  Cavg3
## 1 simulation 2.551332
                          2.583974
                                      1.012794 8.116794e-03 8.999798e-03
## 2 simulation 2.551332
                          2.624257
                                      1.028583 1.815418e-02 2.030361e-02
## 3 simulation 2.551332 2.722778 1.067199 4.266990e-02 4.881300e-02
## 4 simulation 2.551332 3.019877
                                      1.183648 1.146346e-01 1.410843e-01
## 5 simulation 2.551332 4.643720
                                      1.820116 4.480822e-01 7.420969e-01
## 6 simulation 2.551332 19.584284
                                      7.676103 2.557543e+00 7.256225e+00
```

```
simulation 2.551332 216.348104 84.798112 1.712243e+01 6.227119e+01
## 8 simulation 2.551332 432.293678 169.438451 5.710637e+01 1.036092e+02
## 9 simulation 2.551332 607.736635 238.203701 1.437300e+02 1.401636e+02
## 10 simulation 2.551332 853.647835 334.589132 3.304390e+02 2.007661e+02
##
      AFIRT.sim
## 1
      1.002147
## 2
      1.004751
      1.010925
## 3
## 4
      1.028222
## 5
      1.103235
## 6
      1.466487
## 7
      2.935337
## 8
      3.682950
## 9
      3.897053
## 10 3.880476
lumped.parameters.theory("../data/ModelF_Atezolizumab_Params.xlsx", 80)
                M30 Mtot3.ss Tacc.tum
                                              B CL AFIRT.Kss AFIRT.Kd
       type
## 1 theory 2.551332 2.551332
                                    1 0.3333333 0.2
                                                         0.525
                                                                 0.2625
```

under high dose, AFIRT.thy should agree with AFIRT.sim

Make a plot to demonstrate that

Make data frames for lumped parameters(thy and sim) at different doses

```
dose.nmol.range = lseq(80, 8000, 50)

df_sim = data.frame() # put all simulations for different dose into one data frame
for (dose.nmol in dose.nmol.range){
   row = lumped.parameters.simulation("../data/ModelF_Atezolizumab_Params.xlsx", dose.nmol,
   df_sim = rbind(df_sim, row)
}

df_thy = data.frame() # put all theoretical calculations of lumped parameters at different
for (dose.nmol in dose.nmol.range){
   row = lumped.parameters.theory("../data/ModelF_Atezolizumab_Params.xlsx", dose.nmol)
   df_thy = rbind(df_thy, row)
}
```

Get AFIRT from both df_sim and df_thy

```
Dose = as.data.frame(dose.nmol.range)
AFIRT_sim = as.data.frame(df_sim$AFIRT.sim)
AFIRT_Kd= as.data.frame(df_thy$AFIRT.Kd)
AFIRT_Kss = as.data.frame(df_thy$AFIRT.Kss)
AFIRT = data.frame(Dose, AFIRT_Kd, AFIRT_Kss, AFIRT_sim)
names(AFIRT) = c("Dose", "AFIRT_Kd", "AFIRT_Kss", "AFIRT_sim")
AFIRT
```

```
##
            Dose
                    AFIRT_Kd
                                AFIRT Kss AFIRT sim
                                           2.340834
## 1
        80.00000 0.262500000 0.525000000
##
  2
        87.88329 0.238953272 0.477906543
                                           2.524922
        96.54341 0.217518728 0.435037456
##
  3
                                           2.691186
##
  4
       106.05691 0.198006902 0.396013803
                                           2.838241
       116.50788 0.180245322 0.360490644
##
  5
                                           2.967006
##
  6
       127.98870 0.164076988 0.328153976
                                           3.079226
## 7
       140.60085 0.149358983 0.298717967
                                           3.176797
## 8
       154.45582 0.135961210 0.271922421
                                            3.261495
##
  9
       169.67607 0.123765242 0.247530484
                                           3.334883
##
       186.39614 0.112663274 0.225326549
  10
                                            3.398303
##
       204.76383 0.102557173 0.205114347
                                            3.452893
##
       224.94150 0.093357608 0.186715216
   12
                                           3.499609
##
   13
       247.10749 0.084983260 0.169966521
                                            3.539253
##
  14
       271.45774 0.077360107 0.154720214
                                            3.572495
##
       298.20750 0.070420765 0.140841529
                                            3.599893
##
  16
       327.59320 0.064103894 0.128207787
                                           3.621905
       359.87461 0.058353658 0.116707315
##
  17
                                            3.638911
##
       395.33707 0.053119228 0.106238457
  18
                                           3.651216
##
   19
       434.29404 0.048354337 0.096708673
                                           3.659065
##
  20
       477.08987 0.044016865 0.088033729
                                           3.662648
##
  21
       524.10285 0.040068472 0.080136943
                                           3.662104
## 22
       575.74854 0.036474257 0.072948513
                                           3.657535
##
  23
       632.48346 0.033202449 0.066404899
                                            3.649000
##
  24
       694.80910 0.030224129 0.060448258
                                           3.636526
##
  25
       763.27638 0.027512970 0.055025940
                                            3.620111
##
  26
       838.49051 0.025045006 0.050090013
                                            3.599727
##
   27
       921.11632 0.022798424 0.045596847
                                            3.575321
##
  28 1011.88417 0.020753363 0.041506727
                                            3.546821
  29 1111.59640 0.018891749 0.037783498
                                            3.514143
  30 1221.13437 0.017197125 0.034394249
                                            3.477186
  31 1341.46635 0.015654511 0.031309022
                                            3.435848
   32 1473.65598 0.014250273 0.028500546
                                            3.390021
  33 1618.87172 0.012971998 0.025943995
                                            3.339604
   34 1778.39719 0.011808386 0.023616772
                                            3.284504
##
  35 1953.64248 0.010749152 0.021498304
                                           3.224647
  36 2146.15664 0.009784934 0.019569867
                                            3.159990
     2357.64136 0.008907207 0.017814414
                                           3.090516
     2589.96603 0.008108214 0.016216429
                                            3.016256
##
  39 2845.18424 0.007380893 0.014761786
                                           2.937292
  40 3125.55195 0.006718813 0.013437627
                                           2.853766
  41 3433.54741 0.006116124 0.012232247
                                           2.765884
##
  42 3771.89309 0.005567496 0.011134992
                                           2.673925
  43 4143.57974 0.005068082 0.010136163
                                           2.578243
  44 4551.89282 0.004613465 0.009226931
                                           2.479269
## 45 5000.44154 0.004199629 0.008399258
                                            2.377496
  46 5493.19076 0.003822915 0.007645830
                                           2.273490
  47 6034.49605 0.003479992 0.006959985
                                           2.167863
  48 6629.14218 0.003167831 0.006335661
                                           2.061270
  49 7282.38542 0.002883670 0.005767341
                                            1.954387
## 50 8000.00000 0.002625000 0.005250000
                                           1.847895
```

Hongshan says: We don't actually need a plot to realize that the theory and simulation differ by a big margin But I will make a plot anyway, just to see which one is linear

```
names = names(AFIRT)
data = AFIRT %>% gather(key, value, -c(get(names[1])))
g = ggplot(data, aes(Dose, value, color=key)) +
  scale.x.log10() +
  scale.y.log10() +
  geom_point()
g
                                                                            key
                                                                             AFIRT_Kd
value
   0.1
                                                                                AFIRT_Kss
                                                                                AFIRT_sim
  0.01
           100
                                         1000
                                                                      10000
```

Dose

Hongshan says: Deeply confused