Assignment week 3

Detailed analysis of the glucocorticoid receptor dynamica model $\,$



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1 Assignment 1: Assessing Model Validity

To assess the validity of the model, the experimental data will be compared to the simulation data from previous week. The comparison will be done by plotting the experimental data and the simulation data in the same graph.

1.1 Loading in the data

```
# Load in data from file
data <- read.csv("MPL.csv", na.strings = "NA")
# Rename 'mRNA' and 'Free_receptor' columns to use the same name scheme
names(data)[4:5] <- c("Rm", "R")</pre>
```

The actual experiment data contains multiple values per time, if a plot is made with those raw values the plot below will result.

```
# Create plot with raw values to demonstrate the need of using medians
ggplot(data, mapping = aes(x = time, y = R, width = 2)) + geom_line() + geom_point()
```

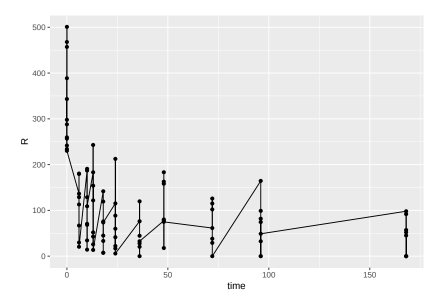


Figure 1: Demonstration of plotting raw data

As can be seen the line goes through each single point and thus is the line not that useful. So to be able to get a meaningful plot the medians are used to create a useful line of the medians that can then be used to compare the values.

1.2 Implementing GRD model with given values

Table 1: Initial Values for MPL

Parameter	Value	Unit	Explanation
Rm	4.74	$fmol/g\ liver$	concentration of receptor mRNA
R	267	$fmol/mg\ protein$	concentration of free receptor in cytosol
DR	0	$fmol/mg\ protein$	concentration of receptor complex in cytosol
DR_N	0	$fmol/mg\ protein$	concentration of receptor complex in nucleus

Table 2: Parameter Values for MPL

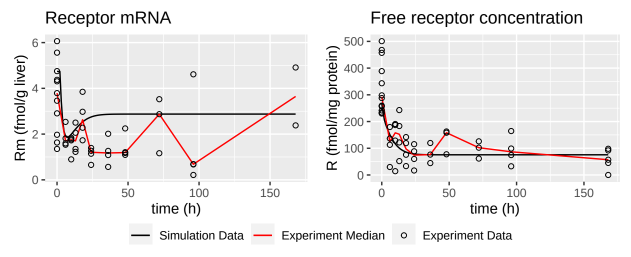
Parameter	Value	Unit	Explanation
k.s_Rm	2.9	fmol/g liver/h	zero-order rate constant of receptor mRNA synthesis
$k.d$ _Rm	0.612	-	first-order rate constant receptor mRNA degradation
$IC.50_Rm$	26.2	$fmol/mg\ protein$	concentration of DR_N where receptor mRNA
			synthesis drops to 50% of base value
k.on	0.00329	L/nmol/h	second-order rate constant of receptor complex
			formation
k.T	0.63	1/h	first-order rate constant of translocation of receptor
			complex to nucleus
k.re	0.57	1/h	first-order rate constant of receptor 'recovery' from
			nucleus to cytosol
R.f	0.49	-	fraction of receptor being recycled from complexes
$k.s_R$	3.22	-	first-order rate constant of receptor synthesis
$k.d_R$	0.0572	1/h	first-order rate constant of receptor degradation
D	-	nmol/L	plasma concentration of corticosteroid

```
# Create function of GRD model
model <- function(time, cur.state, parameters) {</pre>
     # Unpack the current state and the parameters for instant access
     with(as.list(c(cur.state, parameters)), {
            # Calculate delta for each equation and return them in a list
           \label{eq:local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_local_
           delta.R \leftarrow k.s_R * Rm + R.f * k.re * DR_N - k.on * D * R - k.d_R * R
           delta.DR \leftarrow k.on * D * R - k.T * DR
           delta.DR_N <- k.T * DR - k.re * DR_N
           return(list(c(delta.Rm, delta.R, delta.DR, delta.DR_N)))
           })
}
# Initial values and time frame
initial.values <- c(Rm = 4.74, R = 267, DR = 0, DR_N = 0)
times <- seq(0, 168, by = 1)
# Create function that converts concentration of MPL from ng/mL to nmol/L
get.D <- function(ng.ml.concentration) { return(ng.ml.concentration * 1000 / 374.471) }</pre>
# Define parameters determined for methylprednisolone (MPL)
parameters <- c(k.s_Rm = 2.9, k.d_Rm = 0.612, IC.50_Rm = 26.2, k.on = 0.00329, k.T = 0.63,
                                              k.re = 0.57, R.f = 0.49, k.s_R = 3.22, k.d_R = 0.0572, D = get.D(20)
```

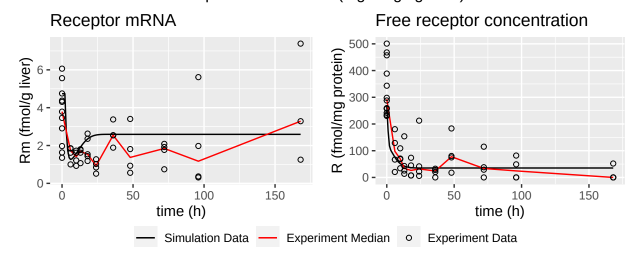
1.3 Creating plots to compare experiment data with simulation data

```
# Get all unique doses present in the data except 0.0
all.doses <- unique(data$dose)</pre>
all.doses <- all.doses[!all.doses %in% (0.0)]
# Create function that calculates median MPL concentration for a dose
get.median.mpl_conc <- function(i) { median(data$MPL_conc[data$dose==i], na.rm=T) }</pre>
# Create data frame with mpl concentration for each dose
median.mpl_conc <- data.frame("dose" = all.doses,</pre>
                               "mpl conc" = unlist(lapply(all.doses, get.median.mpl conc)))
get legend<-function(myggplot){</pre>
  tmp <- ggplot_gtable(ggplot_build(myggplot))</pre>
  leg <- which(sapply(tmp$grobs, function(x) x$name) == "guide-box")</pre>
 legend <- tmp$grobs[[leg]]</pre>
 return(legend)
for (current.dose in all.doses) {
  # Change to correct D parameter for current dose
  parameters$D <- get.D(median.mpl conc$mpl conc[median.mpl conc$dose == current.dose])</pre>
  # Get subset of experiment data for current dose
  cur.data <- subset(data, dose == 0.0 | dose == current.dose)</pre>
  # Get subset of experiment median data for current dose
  cur.data.medians <- subset(data.medians, dose == 0.0 | dose == current.dose)</pre>
  # Perform ODE function using the model with the given values for MPL
  cur.data.simulated <- as.data.frame(ode(times = times, y = initial.values,</pre>
                                        parms = parameters, func = model, method = "euler"))
  # Create plot for receptor mRNA
  plot.Rm <- ggplot(cur.data.simulated, aes(x = time, y = Rm)) +
    geom_line(aes(color = "Simulation Data")) +
    geom line(aes(color = "Experiment Median"), data = cur.data.medians) +
    geom_point(aes(color = "Experiment Data"), data = cur.data, shape = 1) +
    ggtitle("Receptor mRNA") + ylab("Rm (fmol/g liver)") + xlab("time (h)") +
    theme(legend.position = "bottom", plot.margin = margin(0.25, 0.25, 0.25, 0.25, "cm")) +
    scale color manual("",
                       limits=c("Simulation Data", "Experiment Median", "Experiment Data"),
                       values = c("black","red","black")) +
    guides(color = guide_legend(override.aes = list(linetype = c(1, 1, NA), shape = c(NA, NA, 1))))
  # Create plot for free receptor concentration
  plot.R \leftarrow ggplot(cur.data.simulated, aes(x = time, y = R)) +
    geom_line(aes(color = "Simulation Data")) +
    geom_line(aes(color = "Experiment Median"), data = cur.data.medians) +
    geom_point(aes(color = "Experiment Data"), data = cur.data, shape = 1) +
    ggtitle("Free receptor concentration") + ylab("R (fmol/mg protein)") + xlab("time (h)") +
    theme(legend.position = "bottom", plot.margin = margin(0.25, 0.25, 0.25, 0.25, "cm")) +
    scale_color_manual("", limits=c("Simulation Data", "Experiment Median", "Experiment Data"),
                       values = c("black","red","black")) +
```

Graphs for dose = 0.1 (mg drug/kg rat/h)



Graphs for dose = 0.3 (mg drug/kg rat/h)



1.4 Results of plot comparison

[2] How do the results of the simulations depend on the dose and concentration of the drug? Compare the model variables mRNA, R with the experimental data

[3] Are the results of the model in line with experimental data? If not, what could be the reason? Think of at least one explanation.

Try to test it with simulations (you will get bonus points for that, your explanation does not need to be correct, but should be logical).

References

[1] Barnes, P.J. (2011), Glucocorticosteroids: current and future directions, British Journal of Pharmacology, 163: 29-43, https://doi.org/10.1111/j.1476-5381.2010.01199.x (accessed May 11, 2022).