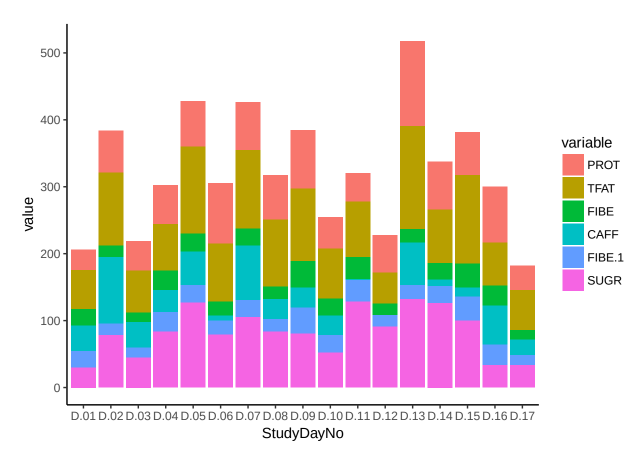
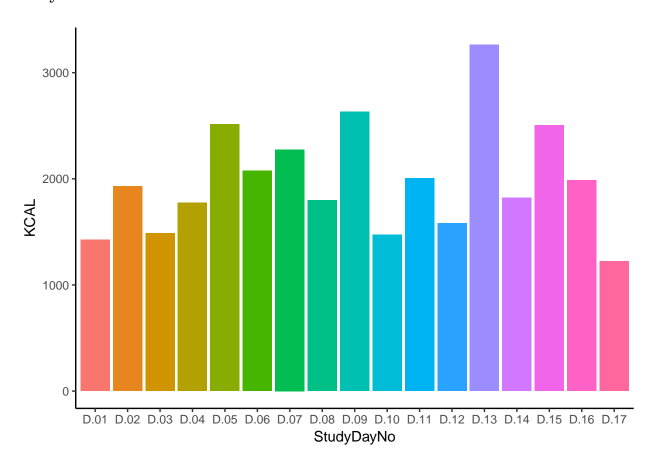
${\bf Subject_22}$

Type	Your Average	Total Average
CALORIES	1986.31	2080.11
PROTEIN	64.85	88.57
TOTAL FAT	90.32	89.97
CARBS	239.28	225.55
FIBER	24.74	21.96

MicroNutrients



Daily Calorie Intake



Microbiome Daily Relative Abundance

'measure' vars in future.

```
# make ggplot bar chart of top 10 most abundant species per day
#melt subtaxasp to get our dataframe in the long format for future usage
  meltdf<- melt(subtaxasp)

## Warning in melt.data.table(subtaxasp): To be consistent with reshape2's
## melt, id.vars and measure.vars are internally guessed when both are 'NULL'.</pre>
```

All non-numeric/integer/logical type columns are conisdered id.vars, which ## in this case are columns [rn]. Consider providing at least one of 'id' or

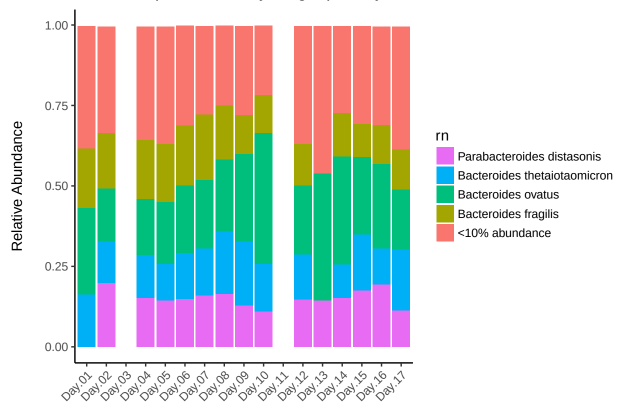
```
#merge to get access to Day var
mergedf<- merge(x=meltdf, y=map, by.x = "variable", by.y= "X.SampleID", all.x=TRUE)

#convert our dataframe species (rn) column to a character
mergedf$rn <- as.character(mergedf$rn)

#series of gsub commands meant to neaten and clarify legend content
mergedf$rn <- gsub(".*s__", "", mergedf$rn)
mergedf$rn <- gsub("\\[", "",mergedf$rn)</pre>
```

```
mergedf$rn <- gsub("\\]", "",mergedf$rn)</pre>
  mergedf$rn <- gsub("_", " ",mergedf$rn)</pre>
  #create <10% abundance category
  mergedf$rn[mergedf$value < 0.1] <- "<10% abundance"</pre>
ggplot(mergedf, aes(x = StudyDayNo, y = value, fill = rn)) +
  geom_bar(stat = "identity") +
  scale_x_discrete(drop = FALSE) +
  theme_classic() +
  theme(strip.text.y = element_text(angle = 0, size = 8, face = "italic"),
        axis.text.x = element_text(angle = 45, hjust = 1),
        axis.title.x = element_blank(),
        plot.title = element_text(hjust = 0.5),
        strip.background = element_rect(color = "grey")) +
  guides(fill = guide_legend(reverse = TRUE,
                              keywidth = 1,
                              keyheight = 1,
                              ncol = 1)) +
  ylab("Relative Abundance\n") +
  ggtitle("Main species within your gut per day")
```

Main species within your gut per day



`geom_smooth()` using method = 'loess'

