# apply-prep.Rmd

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## Reusing code - Apply functions

Introduction to repeating things in R

```
est_mass <- function(volume) {
   mass <- 2.65 * volume^0.9
   return(mass)
}

est_mass(1.6)

## [1] 4.045329

est_mass(5.6)

## [1] 12.49151</pre>
```

### Using vectorized functions

```
est_mass <- function(volume) {
   mass <- 2.65 * volume^0.9
   return(mass)
}

est_mass(1.6)

## [1] 4.045329

est_mass(5.6)

## [1] 12.49151</pre>
```

```
c(1, 2, 3) * 2
## [1] 2 4 6
volumes <-c(1.6, 5.6, 3.1)
est mass(volumes)
## [1] 4.045329 12.491515 7.336204
# install.packages("stringr")
library(stringr)
str_to_sentence(c("dipodomys", "chaetodipus"))
## [1] "Dipodomys"
                      "Chaetodipus"
genus <- c("dipodomys", "chaetodipus", "dipodomys")</pre>
species <- c("ordii", "baileyi", "spectabilis")</pre>
combine_genus_species <- function(genus, species) {</pre>
  genus_cap <- str_to_sentence(genus)</pre>
  genus_species <- paste(genus_cap, species)</pre>
  return(genus_species)
combine_genus_species(genus, species)
## [1] "Dipodomys ordii"
                                 "Chaetodipus baileyi"
                                                          "Dipodomys spectabilis"
data <- data.frame(genus, species)</pre>
combine_genus_species(data$genus, data$species)
## [1] "Dipodomys ordii"
                                 "Chaetodipus baileyi"
                                                          "Dipodomys spectabilis"
```

### Apply functions

```
library(stringr)
```

```
est_mass <- function(volume) {
  if(volume > 5) {
  mass <- 2.65 * volume^0.9
  } else {
    mass <- NA
  }
  return(mass)
}</pre>
```

```
volumes <- c(1.6, 5.6, 3.1)
# est_mass(volumes)
sapply(volumes, est_mass)

## [1] NA 12.49151 NA

c(est_mass(volumes[1]), est_mass(volumes[2]), est_mass(volumes[3]))

## [1] NA 12.49151 NA</pre>
```

#### The mapply function

```
est_mass <- function(volume, veg_type) {
  if(veg_type == "tree") {
  mass <- 2.65 * volume^0.9
  } else {
    mass <- NA
  }
  return(mass)
}

volumes <- c(1.6, 5.6, 3.1)
veg_type <- c("shrub", "tree", "tree")</pre>
```

```
mapply(FUN = est_mass, volume = volumes, veg_type = veg_type)
```

## [1] NA 12.491515 7.336204

### Combining functions with dplyr

```
est_mass <- function(volume, veg_type) {
  if(veg_type == "tree") {
  mass <- 2.65 * volume^0.9
  } else {
    mass <- NA
  }
  return(mass)
}</pre>
```

```
est_mass_vectorized <- function(volume) {
  mass <- 2.65 * volume^0.9
  return(mass)
}</pre>
```

```
volumes <- c(1.6, 5.6, 3.1)
veg_type <- c("shrub", "tree", "tree")
plant_data <- data.frame(volumes, veg_type)</pre>
```

```
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
       intersect, setdiff, setequal, union
plant_data %>%
  rowwise() %>%
  mutate(masses = est_mass(volumes, veg_type))
## # A tibble: 3 x 3
## # Rowwise:
   volumes veg_type masses
                     <dbl>
##
      <dbl> <chr>
## 1
       1.6 shrub
                     NA
## 2
        5.6 tree
                      12.5
## 3
        3.1 tree
                      7.34
get_biomass <- function(volumes) {</pre>
  masses <- est_mass_vectorized(volumes)</pre>
  biomass <- sum(masses)</pre>
 return(biomass)
get_biomass(volumes)
## [1] 23.87305
plant_data %>%
  group_by(veg_type) %>%
  summarize(biomass = get_biomass(volumes))
## # A tibble: 2 x 2
## veg_type biomass
## <chr>
              <dbl>
## 1 shrub
               4.05
## 2 tree
              19.8
```