Introduction to R Programming Factors

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What are factors?

- ▶ A factor is a vector that can contain only predefined values
- ► Factors are used to represent categorical data

Categorical variables

- Categorical variables have a limited and known set of possible outcomes
- Categorical variables typically qualitative

Categorical variables

Examples:

- ▶ flip of a coin (heads or tails)
- ▶ size of a shirt (S, M, L, XL, XXL, XXXL)
- marital status (single, married, divorced, widowed)
- credit rating (very bad, bad, average, good, excellent)
- school grades (A, B, C, D, E, F)

Categorical variables

Person	Sex	Month of birth
Liz Jolene Susan Boris Rochelle Tim Simon Amy	Female Female Male Female Male Male Male Female	April January December September November July July June

Figure 1: Dataset with categorical variables

More about factors

- Factors are built on top of integers
- ► They come with two attributes:
 - Levels, which define the set of allowed values
 - ► Their own class, "factor", which makes them behave differently from regular integers

```
Let's start with a character vector:
```

```
blood <- c("B", "AB", "O", "A", "O", "O", "A")
blood
```

```
## [1] "B" "AB" "O" "A" "O" "O" "A"
```

The factor function encodes a vector as a factor:

```
blood_factor <- factor(blood)
blood_factor</pre>
```

```
## [1] B AB O A O O A
## Levels: A AB B O
```

Note that R sorts the levels alphabetically

```
levels(blood_factor)

## [1] "A" "AB" "B" "O"

typeof(blood_factor)

## [1] "integer"

class(blood_factor)

## [1] "factor"
```

When creating a factor, we can set the ordering of the levels:

```
## [1] B AB O A O O A
## Levels: O A B AB
```

Order levels differently

We can modify the ordering of the levels of an existing factor:

```
blood_factor
## [1] B AB O A O O A
## Levels: A AB B O
factor(blood_factor, levels = c("O", "A", "B", "AB"))
## [1] B AB O A O O A
## Levels: O A B AB
```

Order levels differently

Relevel re-orders the levels of a factor so that the level specified by ref is the first level:

```
blood_factor

## [1] B AB O A O O A
## Levels: A AB B O

relevel(blood_factor, ref = "0")

## [1] B AB O A O O A
## Levels: O A AB B
```

Order levels differently

Levels: O B AB A

```
With rev we can reverse the ordering of the levels of a factor:
blood_factor
## [1] B AB O A O O A
## Levels: A AB B O
factor(blood_factor,
   levels = rev(levels(blood_factor)))
## [1] B AB O A O O A
```

The internal structure of a factor

The str function displays the internal structure of an object:

```
## [1] B AB O A O O A ## Levels: A AB B O
```

```
str(blood_factor)
```

```
## Factor w/ 4 levels "A", "AB", "B", "O": 3 2 4 1 4 4 1
```

- Note that the values are stored as integers!
- ► The levels are just a set of character values to print when the factor is displayed

The internal structure of a factor

##

```
blood_factor2
## [1] B AB O A O O A
## Levels: O A B AB
str(blood_factor2)
```

Factor w/ 4 levels "O", "A", "B", "AB": 3 4 1 2 1 1 2

The internal structure of a factor

To see the underlying integer coding of a factor we can coerce the factor to numeric:

```
blood_factor
```

```
## [1] B AB O A O O A
## Levels: A AB B O
```

```
as.numeric(blood_factor)
```

```
## [1] 3 2 4 1 4 4 1
```

Invalid factor levels

Levels: A AB B O

```
blood_factor

## [1] B AB O A O O A

## Levels: A AB B O

blood_factor[3] <- "C"

blood_factor

## [1] B AB <NA> A O O A
```

Table a factor

##

##

O A B AB

3 2 1 1

```
How many people there are with each type of blood?

table(blood_factor)

## blood_factor

## A AB B O

## 2 1 1 2

table(blood_factor2)

## blood_factor2
```

Renaming factor levels

[1] BT B BT_AB BT_O BT_A BT_O BT_O BT_A

Levels: BT_O BT_A BT_B BT_AB

Levels and labels

- Levels are input (alphabetic order by default)
- ► Labels are associated to levels and control how they are displayed in the output

Levels and labels

```
blood <- c("B", "AB", "O", "A", "O", "O", "A", "B")
factor(blood)
## [1] B AB O A O O A B
## Levels: A AB B O
factor(blood.
        labels = c("BT A", "BT AB", "BT B", "BT O"))
## [1] BT_B BT_AB BT_O BT_A BT_O BT_O BT_A BT_B
## Levels: BT A BT AB BT B BT O
```

Levels and labels

```
factor(blood)
## [1] B AB O A O O A B
## Levels: A AB B O
factor(blood.
        levels = c("0", "A", "B", "AB"),
        labels = c("BT O", "BT A", "BT B", "BT AB"))
## [1] BT_B BT_AB BT_O BT_A BT_O BT_O BT_A BT_B
## Levels: BT O BT A BT B BT AB
```

Labels

Duplicated values in labels can be used to map different values of the factor to the same level:

```
## [1] BT_B BT_A BT_O BT_A BT_O BT_O BT_A BT_B
## Levels: BT_O BT_A BT_B
```

```
blood <- c("B", "AB", "O", "A", "O", "O", "A", "B")
blood_factor <- factor(blood)
blood_factor[1] < blood_factor[2]</pre>
```

[1] NA

This logical comparison is not meaningful, since the factor is not ordered

Let's build an ordered factor:

```
## [1] M L S S L M L M
## Levels: S < M < L
```

```
tshirt_factor[1] < tshirt_factor[2]

## [1] TRUE

tshirt_factor[1] > tshirt_factor[2]

## [1] FALSE
```

```
Ordered factors differ from factors in their class:
```

```
class(blood_factor)

## [1] "factor"

class(tshirt_factor)

## [1] "ordered" "factor"
```

```
ordered(x) is an alternative to factor(x, ordered = TRUE):
factor(tshirt, ordered = TRUE,
  levels = c("S", "M", "L"))
## [1] M L S S L M L M
## Levels: S < M < L
ordered(tshirt, levels = c("S", "M", "L"))
## [1] M L S S L M L M
## Levels: S < M < L
```

```
The underlying integer coding of the tshirt factor: tshirt factor
```

```
## [1] M L S S L M L M
## Levels: S < M < L
```

as.numeric(tshirt_factor)

```
## [1] 2 3 1 1 3 2 3 2
```

Reordering levels changes the underlying integer coding of the factor:

```
tshirt_factor_2 <- factor(tshirt_factor,
  levels = c("L", "M", "S"))

tshirt_factor_2

## [1] M L S S L M L M
## Levels: L < M < S
as.numeric(tshirt_factor_2)

## [1] 2 1 3 3 1 2 1 2</pre>
```

Testing and coercing

Levels: A < AB < B < O

```
is.factor(blood_factor)

## [1] TRUE
is.ordered(blood_factor)

## [1] FALSE
as.ordered(blood_factor)

## [1] B AB O A O O A B
```

Testing and coercing

[1] B AB O A O O A B ## Levels: A < AB < B < O

```
is.factor(blood)
## [1] FALSE
as.factor(blood)
## [1] B AB O A O O A B
## Levels: A AB B O
as.ordered(blood)
```

Testing and coercing

Curious about the the differences between factor and as.factor?

► See stackoverflow.com/questions/39279238/why-use-as-factor-instead-of-just-factor

Bivariate tables

Bivariate tables

```
table(hair_color, eye_color)
```

```
## eye_color
## hair_color Blue Green
## Black 3 2
## Blond 1 1
## Brown 1 0
```

Three-way tables

```
hair color <- factor(c("Brown", "Black", "Black",
                          "Black", "Blond", "Blond",
                          "Black", "Black"))
eye_color <- factor(c( "Blue", "Blue", "Green",
                          "Green", "Green", "Blue",
                          "Blue", "Blue"))
shirt_size <- factor(c("L", "S", "S", "M", "L",</pre>
                          "L", "S", "S"),
                        ordered = TRUE
```

Three-way table

table(hair_color, eye_color, shirt_size)

Three-way table

```
## , , shirt_size = L
##
##
            eye_color
## hair_color Blue Green
##
       Black
                0
                       0
##
       Blond
                 1
##
       Brown
                       0
##
   , , shirt_size = M
##
##
            eye_color
## hair_color Blue Green
##
       Black
                0
                       1
       Blond
                0
                       0
##
##
       Brown
                0
                       0
##
## , , shirt_size = S
##
##
            eye_color
## hair color Blue Green
##
       Black
       Blond
                       0
##
##
       Brown
                0
                       0
```

Creating factors with cut

- ▶ The cut function transforms numerical vectors into factors
- cut breaks the range of a numerical vector into intervals
- The limits of the intervals are provided as input

Creating factors with cut

```
y <- c(5.4, 1.5, 3.33, 0.01, 2, 4.2, 1.99, 1.01)
limits <- c(0, 2, 4, 6)

y_factor <- cut(y, breaks = limits)

y_factor

## [1] (4,6] (0,2] (2,4] (0,2] (0,2] (4,6] (0,2] (0,2]
## Levels: (0.2] (2,4] (4.6]</pre>
```

Creating factors with cut

```
table(y_factor)
```

```
## y_factor
## (0,2] (2,4] (4,6]
## 5 1 2
```

Open and closed intervals

Intervals closed on the right and open on the left:

```
levels(cut(y, breaks = c(0, 2, 4, 6)))
## [1] "(0,2]" "(2,4]" "(4,6]"
Intervals open on the right and closed on the left:
levels(cut(y, breaks = c(0, 2, 4, 6), right = FALSE))
## [1] "[0,2)" "[2,4)" "[4,6)"
```

Open and closed intervals

Intervals closed on the right and open on the left, but including the lowest value:

```
## [1] "[0,2]" "(2,4]" "(4,6]"
```

Intervals open on the right and closed on the left but including the highest value:

```
## [1] "[0,2)" "[2,4)" "[4,6]"
```