The Data Validation Cookbook

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Preface

This book is about checking data with the validate package for R.

This version of the book was rendered with validate version 1.0.1. The latest release of validate can be installed from CRAN as follows.

```
install.packages("validate")
```

The purposes of this book include demonstrating the main tools and workflows of the validate package, giving examples of common data validation tasks, and showing how to analyze data validation results.

The book is organized as follows. Chapter 1 discusses the bare necessities to be able to follow the rest of the book. Chapters 2 to 5 form the 'cookbook' part of the book and discuss many different ways to check your data by example. Chapter 6 is devoted to deriving plausibility measures with the validate package. Chapters 7 and 8 treat working with validate in-depth. Chapter 9 discusses how to compare two or more versions of a dataset, possibly automated through the lumberjack package. The section with Biblographical Notes lists some references and points out some literature for further reading.

Prerequisites

Readers of this book are expected to have some knowledge of R. In particular, you should know how to import data into R and know a little about working with data frames and vectors.

Citing this work

To cite the validate package please use the following citation.

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I am greatly indebted by dr Olav ten Bosch for carefully reviewing the manuscript of version 1.0.1. Any mistakes are of course entirely the author's fault.

Contributing

If you find a mistake, or have some suggestions, please file an issue or a pull request on the github page of the package: https://github.com/data-cleaning/validate. If you do not have or want a github account, you can contact the author via the e-mail address that is listed with the package.

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Chapter 1

Introduction to validate

Data Validation is an activity verifying whether or not a combination of values is a member of a set of acceptable combinations (Di Zio et al , 2015).

The validate package is intended to make checking your data easy, maintainable, and reproducible. It does this by allowing you to

- test data against a reusable set of data validation rules:
- investigate, summarise, and visualise data validation results;
- · import and export rule sets from and to various formats;
- · filter, select and otherwise manipulate data validation rules';
- · investigate, summarise, and visualise rule sets.

For advanced rule manipulation there is the validatetools package.

1.1 A quick example

10

2

Here's an example demonstrating the typical workflow. We'll use the built-in cars data set, which contains 50 cases of speed and stopping distances of cars.

```
data(cars)
head(cars, 3)

speed dist
1  4  2
```

Validating data is all about checking whether a data set meets presumptions or expectations you have about it, and the validate package makes it easy for you to define those expectations. Let's do a quick check on variables in the cars data set. We first load the package, and create a list of data quality demands with the validator() function.

```
, speed/dist <= 1.5
, cor(speed, dist)>=0.2)
```

Here, the first three rules are record-wise checks: each record will yield one answer. In the last rule we check whether speed and distance are positively correlated this will yield a single TRUE or FALSE for the whole data set. We now confront the data with those rules and save the output into a variable called out.

```
out <- confront(cars, rules)</pre>
```

The easiest way to check the results is with summary().

```
summary(out)
```

```
name items passes fails nNA error warning
         50
                50
                       A
                            0 FALSE
                                      FALSE
   ٧2
         50
                 50
                       0
                            0 FALSE FALSE
2
                           0 FALSE FALSE
   ٧3
         50
                 48
                       2
3
                           0 FALSE FALSE
   ٧4
                 1
                       A
               expression
  (speed - 0) >= -1e-08
1
2
    (dist - 0) >= -1e-08
       speed/dist <= 1.5
3
4 \text{ cor(speed, dist)} >= 0.2
```

This returns a data frame with one line of information for each rule V1, V2, V3 and V4. To be precise:

- · How many data items were checked against each rule.
- How many items passed, failed or resulted in NA.
- Whether the check resulted in an error (could not be performed) or gave an warning.
- The expression that was actually evaluated to perform the check.

The same information can be summarized graphically as follows (see Figure 1.1).

```
plot(out)
```

In this plot each horizontal bar indicates the percentage of Failing, Passing, and Missing cases. The table in the legend lists the total number of Fails, Passes and Missings, summed over all checks. Here, we have 4 rules. The first three rules yield 50 results each, while the last rule yields a single result. Hence there are 151 validation results in total

Using the function violating we can select the records that violate one or more rules. We select only the first three results because the last rule can not be interpreted record by record.

```
violating(cars, out[1:3])
```

```
speed dist
1 4 2
3 7 4
```

We can extract all individual resuls results using for example as.data.frame.

```
df_out <- as.data.frame(out)
head(df_out, 3)</pre>
```

name value

1.1. A QUICK EXAMPLE 9

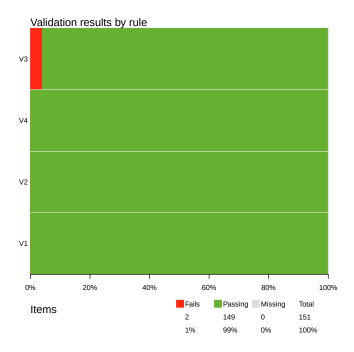


Figure 1.1: Plot of validation output.

```
1 V1 TRUE (speed - \theta) >= -1e-08
2 V1 TRUE (speed - \theta) >= -1e-08
3 V1 TRUE (speed - \theta) >= -1e-08
```

We see that in record 1, rule v1, was satisfied (the result is TRUE), and that validate left a bit of slack when executing the rule, to avoid false negatives caused machine rounding issues.

Summarizing, the basic workflow in validate is to create a rule set, confront a data set with the rules in the rule set, and than analyze or use the results further. To understand which checks you can perform with validate you only need to remember the following.

Any R expression that results in a logical is accepted by validate as a validation rule.

You are now ready to start validating your data, and navigate Chapters 3-5 to learn how to define specific types of checks. Chapter~7, discusses more details about working with validate.

Chapter 2

Variable checks

Variable checks are checks that can be performed on a field-by-field basis. An example is checking that a variable called Age is nonnegative, or of integer type. Variable checks are among the simplest checks.

Data

In this section we will use the SBS2000 dataset, that is included with validate.

```
library(validate)
data(SBS2000)
head(SBS2000, 3)
```

```
id size incl.prob staff turnover other.rev total.rev
1 RET01 sc0
                0.02
                       75
                               NΑ
                                        NA
                                                1130
2 RET02 sc3
                0.14
                        9
                              1607
                                       NA
                                                1607
3 RET03 sc3
                              6886
                                       -33
                0.14
                                                6919
                       NA
 staff.costs total.costs profit vat
                18915 20045 NA
1
         NA
        131
                  1544
                         63 NA
3
        324
                 6493
                          426 NA
```

See ?SBS2000 for a description.

2.1 Variable type

In R, one can test the type of a variable using built-in functions such as is.numeric Or is.character.

```
is.character("hihi")
```

```
[1] TRUE
```

```
is.character(3)
```

[1] FALSE

In validate, any function starting with is. ('is' followed by a dot) is considered a validation function.

```
rules <- validator(
    is.character(size)
    , is.numeric(turnover)
)
out <- confront(SBS2000, rules)
summary(out)</pre>
```

We see that each rule checks a single item, namely one column of data. The first rule is violated (it is in fact a factor variable). The second rule is satisfied.

2.2 Missingness

Use R's standard is.na() to check missin items in individual variables. Negate it to check that values are available.

```
rule <- validator(
   !is.na(turnover)
, !is.na(other.rev)
, !is.na(profit)
)
out <- confront(SBS2000, rule)
summary(out)</pre>
```

```
name items passes fails nNA error warning

1 V1 60 56 4 0 FALSE FALSE

2 V2 60 24 36 0 FALSE FALSE

3 V3 60 55 5 0 FALSE FALSE

expression

1 !is.na(turnover)

2 !is.na(other.rev)

3 !is.na(profit)
```

We see that in 4 cases the variable turnover is missing, while other.rev and profit are missing respectively in 36 and 5 occasions.

To demand that all items must be present or absent for a certain variable, use R's quantifiers: any() or all(), possibly negated.

```
rules <- validator(
  !any(is.na(incl.prob))
  , all(is.na(vat)) )
out <- confront(SBS2000, rules)
summary(out)</pre>
```

2.3. FIELD LENGTH

- · To check whether records or parts thereof are completed, see 4.1.
- To check whether records are available at all, see 3.3.

2.3 Field length

The number of characters in text fields can be tested using either R's standard nchar() function, or with the convenience function field_length.

```
rules <- validator(
   nchar(as.character(size)) >= 2
, field_length(id, n=5)
, field_length(size, min=2, max=3)
)
out <- confront(SBS2000, rules)
summary(out)</pre>
```

```
name items passes fails nNA error warning
   ٧1
         60
               60
                      0
                          0 FALSE
                                  FALSE
2
  V2
         60
               60
                      0
                          0 FALSE
                                   FALSE
  V3 60
3
               60
                      A
                          0 FALSE FALSE
                          expression
       nchar(as.character(size)) >= 2
1
              field_length(id, n = 5)
3 field_length(size, min = 2, max = 3)
```

One advantage of check_field_length is that its argument is converted to character (recall that size is a factor variable). The function field_length can be used to either test for exact field lengths or to check whether the number of characters is within a certain range.

The field length is measured as the number of code points. Use type="width" to measure the printed width (nr of columns) or type="bytes" to count the number of bytes.

2.4 Format of numeric fields

For numbers that are stored in character type, there is a convenience function called number_format() that accepts a variable name and a format specification.

```
dat <- data.frame(x = c("2.54","2.66","8.142","23.53"))
```

To check that the numbers are formatted with one figure before, and two figures after the decimal point, we perform the following check.

```
rule <- validator( number_format(x, format="d.dd"))
values(confront(dat, rule))</pre>
```

```
V1
[1,] TRUE
[2,] TRUE
[3,] FALSE
[4,] FALSE
```

Here, the specification <code>format="d.dd"</code> describes the allowed numeric formats. In this specification the "d" stands for a digit, any other character except the asterisk (*) stands for itself. The asterisk is interpreted as 'zero or more digits'. Here are some examples of how to define number formats.

format	match	non-match
0.dddd	"0.4321"	"0.123","1.4563"
d.ddEdd	"3.14E00"	"31.14E00"
d.*Edd	"0.314E01","3.1415297E00"	"3.1415230"
d.dd*	"1.23", "1.234",	"1.2"

The last example shows how to check for a minimal number of digits behind the decimal point.

There are special arguments to check the number of decimal figures after the decimal separator.

```
x <- c("12.123","123.12345")
number_format(x, min_dig=4)

[1] FALSE TRUE
number_format(x, max_dig=3)

[1] TRUE FALSE
number_format(x, min_dig=2, max_dig=4)

[1] TRUE FALSE
number_format(x, min_dig=2, max_dig=10)

[1] TRUE TRUE
# specify the decimal separator.</pre>
```

```
[1] TRUE
```

The arguments min_dig, max_dig and dec are ignored when format is specified.

This function is convenient only for fairly simple number formats. Generic pattern matching in strings is discussed in the next section.

2.5 General field format

number_format("12,123", min_dig=2, dec=",")

A simple way to check for more general format is to use globbing patterns. In such patterns, the asterisk wildcard character (*) is interpreted as 'zero or more characters' and the question mark (?) is interpreted as 'any character'.

For example, to check that the id variable in SBS2000 starts with "RET", and that the size variable has consists of "sc" followed by precisely one character, we can do the following.

```
name items passes fails nNA error warning

1 V1 60 60 0 0 FALSE FALSE

2 V2 60 60 0 0 FALSE FALSE

expression

1 field_format(id, "RET*")

2 field_format(size, "sc?")
```

Here, the globbing pattern "RET*" is understood as 'a string starting with "RET", followed by zero or more characters. The pattern "sc?" means 'a string starting with "sc", followed by a single character.

The most general way to check whether a field conforms to a pattern is to use a regular expression. The treatment of regular expressions is out of scope for this book, but we will give a few examples. A good introduction to regular expressions is given by

J. Friedl (2006) Mastering regular expressions. O'Reilley Media.

In validate one can use grep1 or field_format, with the argument type="regex"

Here, the expression "sc[0-9]\$" is a regular expression that should be read as: the string starts (" *) with "sc", is followed by a number between 0 and 9 ("[0-9]") and then ends ("\$"). The regular expression " * RET\\{d}2" indicates that a string must start (" * ") with "RET", followed by two digits (" * \d{2}\), after which the string must end ("\$").

Globbing patterns are easier to develop and easier to understand than regular expressions, while regular expressions offer far more flexibility but are harder to read. Complex and long regular expressions may have subtle matching behaviour that is not immediately obvious to inexperienced users. It is therefore advisable to test regular expressions with a a small dataset representing realistic cases that contains both matches and non-matches. As a rule of thumb we would advise to use globbing patterns unless those offer insufficient flexibility.

2.6 Numeric ranges

Numerical variables may have natural limits from below and/or above. For one-sided ranges, you can use the standard comparison operators.

If a variable is bounded both from above and below one can use two rules, or use the convenience function in_range.

```
rules <- rules +
  validator(PR = in_range(incl.prob, min=0, max=1))</pre>
```

By default, in_range includes the boundaries of the range, so the above rule is equivalent to incl.prob >= 0 and incl.prob <= 0.

```
out <- confront(SBS2000, rules, lin.ineq.eps=0)</pre>
```

Here we set lin.ineq.eps=0 to keep validate from building in a margin for machine rounding errors.

```
summary(out)
```

```
name items passes fails nNA error warning
   T0
               56
                      0 4 FALSE
                                   FALSE
  TC
2
         60
              55
                      0 5 FALSE FALSE
3
   PR
         60
               60
                      0 0 FALSE FALSE
                           expression
1
                        turnover >= 0
                     total.costs >= 0
2
3 in_range(incl.prob, min = 0, max = 1)
```

For numeric ranges it is often a better idea to work with inclusive inequalities (<, >). Take as an example the strict inequality demand income > 0. This means that *any* income larger than zero is acceptable, including numbers such as 0.01, 0.000001 and 10^{-Googol}. In practice there is almost always a natural minimal acceptable value that is usually dictated by the unit of measurement. For example, if we measure income in whole Euros, a better demand would be income >= 1.

2.7 Ranges for times and periods

For objects of class Date and objects of class POSIXct one can use comparison operators and in_range in the same way as for numerical data. The in_range function has a convenience feature for period data that is coded in character data, as in "201801" for quarterly data.

We first generate some example data.

```
period = sprintf("2018Q%d", 1:4)
period
```

```
[1] "2018Q1" "2018Q2" "2018Q3" "2018Q4"
```

The in_range function is capable of recognizing certain date or period formats.

2.8. CODE LISTS 17

```
in_range(period, min="2017Q2", max = "2018Q2")
```

```
[1] TRUE TRUE FALSE FALSE
```

It is possible to specify your own date-time format using strftime notation. See ?in_range and ?strptime for specifications.

2.8 Code lists

A code list is a set of values that a variable is allowed to assume. For small code lists, one can use the %in% operator.

```
rule <- validator(size %in% c("sc0", "sc1", "sc2", "sc3"))
out <- confront(SBS2000, rule)
summary(out)</pre>
```

```
name items passes fails nNA error warning

1 V1 60 60 0 0 FALSE FALSE
expression

1 size %vin% c("sc0", "sc1", "sc2", "sc3")
```

Notice that validate replaces %in% with %vin%. The reason is that %vin% has more consistent behavior in the case of missing data. In particular,

```
c(1, 3, NA) %in% c(1,2)
```

```
[1] TRUE FALSE FALSE
c(1, 3, NA) %vin% c(1,2)
```

[1] TRUE FALSE NA

For longer code lists it is convenient to refer to an externally provided list. There are two ways of doing this: reading the list in the right-hand-size of %in%, or passing a code list to confront as reference data.

Suppose we have a file called codelist.csv with a column code. We can define a rule as follows.

```
rule <- validator(
   x %in% read.csv("codelist.csv")$code
)
## Or, equivalently
rule <- validator(
   valid_codes := read.csv("codelist.csv")$code
   , x %in% valid_codes
)</pre>
```

The disadvantage is that the rule now depends on a path that may or may not be available at runtime.

The second option is to assume that a variable, say valid_codes exists at runtime, and pass this with confront.

```
codelist <- c("sc0","sc1","sc2","sc3")
rule <- validator(size %in% valid_codes)
# pass the codelist</pre>
```

```
name items passes fails nNA error warning
1 V1 60 60 0 0 FALSE FALSE
expression
```

1 size %vin% valid_codes

This way, (very) large code lists can be used, but note that it does require a 'contract' between variable names used in the rule set and variables passed as reference data.

Chapter 3

Availability and uniqueness

In this Chapter it is demonstrated how to check whether records are available and/or complete with respect to a set of keys, and whether they are unique. The checks described here are typically useful for data in 'long' format, where one column holds a value and all the other columns identify the value.

- To test for missing values in individual variables, see also 2.2.
- To check whether records or parts thereof are completed, see 4.1.

Data

In this Chapter the samplonomy dataset is used that comes with the validate package.

```
library(validate)
data(samplonomy)
head(samplonomy, 3)

region freq period measure value
1 Agria A 2014 gdp 600000
2 Agria A 2014 import 210000
3 Agria A 2014 export 222000
```

3.1 Long data

The samplonomy data set is structured in 'long form'. This means that each record has a single value column, and one or more columns containing character values that together describe what the value means.

```
head(samplonomy,3)

region freq period measure value

1 Agria A 2014 gdp 600000

2 Agria A 2014 import 210000

3 Agria A 2014 export 222000
```

The data set contains several time series for multiple measures of the fictional country 'Samplonia'. There are time series for several subregions of Samplonia.

Long format data is typically used as a transport format: it may be used to bulk-load data into SQL-based data base systems, or to transfer data between organisations in a unambiguous way.

Data in long form is in general much harder to check and process for statistical purpose than data in wide format, where each variable is stored in a separate column. The reason is that in long format relations between different variables are spread out across records, and those records are not necessarily ordered in any particular way prior to processing. This makes interpretation of validation fails intrinsically harder for long-form data then for wide-form data

The samplonomy data set has a particularly nasty structure. It contains both annual and quarterly time series for GDP, Import, Export and the Balance of Trade (export less import). The period column therefore contains both quarterly and annual labels. Furthermore, there are time series for the whole of Samplonia (region Samplonia), for each of its two provinces (regions Agria and Induston) and for each of its districts within Agria (Wheaton and Greenham) and Induston (Smokeley, Mudwater, Newbay and Oakdale).

Naturally, we expect that the key combinations are unique, that all time series are gapless and complete, that the Balance of trade equals Export less Import everywhere, that district values add up to the provinces', and that province values add up to the total of Samplonia. Finally, the quarterly time series must add up to the annual values.

3.2 Uniqueness

The function is_unique() checks whether combinations of variables (usually key variables) uniquely identify a record. It accepts any positive number of variable names and returns FALSE for each record that is duplicated with respect to the designated variariables.

Here, we test whether region, period, and measure uniquely identify a value in the samplonomy data set.

```
rule <- validator(is_unique(region, period, measure))
out <- confront(samplonomy, rule)
# showing 7 columns of output for readability
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 1199 1197 2 0 FALSE FALSE
```

There are 2 fails. After extracting the individual values for each record we can find the duplicated ones using a convenience function from validate.

```
violating(samplonomy, out)
```

```
region freq period measure value
870 Induston Q 2018Q2 export 165900
871 Induston Q 2018Q2 export 170000
```

There are a two subtleties to keep in mind when interpreting uniqueness. The first has to do with missing values, and the second has to do with grouping. To start with the missing value problem, take a look at the following two-record data frame.

```
df \leftarrow data.frame(x = c(1,1), y = c("A", NA))
df
```

```
1 1 A
2 1 <NA>
```

How should we judge whether these two records are unique? A tempting option is to say the the first record is unique, and to return NA for the second record since it contains a missing value: R has the habit of returning NA from calculations when an input value is NA. This choice is not invalid, but it would have consequences for determining whether the first record is unique as well. After all, it is possible to fill in a value in the missing field such that the two records are duplicates. Therefore, if one would return NA for the second record, the correct thing to do is to also return NA for the first record. In R, the choice is made to treat NA as an actual value when checking for duplicates or unique records (see ?duplicated from base R). To see this inspect the following code and output.

```
df \leftarrow data.frame(x=rep(1,3), y = c("A", NA, NA))
is\_unique(df\$x, df\$y)
```

[1] TRUE FALSE FALSE

The second subtlety has to do with grouping. You may want to test whether a column is unique, given one or more other variables. It is tempting to think that this requires a split-apply-combine approach where the dataset is first split according to one or more grouping variables, check for uniqueness of the column in each group, and then combine the results. However, such an approach is not necessary as you can simply add the grouping variables to the list of variables that *together* must be unique.

As an example, consider the output of the following two approaches.

```
# y is unique, given x. But not by itself

df <- data.frame(x=rep(letters[1:2],each=3), y=rep(1:3,2))

# the split-apply-combine approach
unsplit(tapply(df$y, df$x, is_unique), df$x)</pre>
```

```
[1] TRUE TRUE TRUE TRUE TRUE TRUE
```

```
# the combined approach
is_unique(df$x, df$y)
```

[1] TRUE TRUE TRUE TRUE TRUE TRUE

3.3 Availability of records

This section is on testing for availability of whole records. Testing for individual missing values (NA), is treated in 2.2.

We wish to ensure that for each region, and each variable, the periods 2014, 2015, ..., 2019 are present. Using contains_at_least we can establish this.

```
rule <- validator(
  contains_at_least(
      keys = data.frame(period = as.character(2014:2019))
    , by=list(region, measure) )
)
out <- confront(samplonomy, rule)</pre>
```

```
# showing 7 columns of output for readability
summary(out)[1:7]
```

```
name items passes fails nNA error warning
1 V1 1199 1170 29 0 FALSE FALSE
```

The function contains_at_least splits the samplonomy dataset into blocks according to values of region and measure. Next, it checks that in each block the variable period contains at least the values 2014–2019.

The return value is a logical vector where the number of elements equals the number of rows in the dataset under scrutiny. It is TRUE for each block where all years are present, and FALSE for each block where one of the years is missing. In this case 29 records are labeled as FALSE. These can be found as follows.

```
head(violating(samplonomy, out))
```

```
region freq period measure value
          Α
                        gdp 600000
               2014
1
    Agria
5
    Agria Q 2014Q1
                        gdp 60000
    Agria Q 2014Q2
                        gdp 120000
9
    Agria
          Q 2014Q3
                        gdp 300000
17
    Agria Q 2014Q4
                        gdp 120000
204 Agria
          0 201501
                        adp 58200
```

Inspection of these records shows that in this block, for Agria the GDP for "2015" is missing.

We can perform a stricter check, and test whether for each measure, all quarters "2014Q1" ... "2019Q4" are present for each province (Agria and Induston). First create a key set to test against.

```
years <- as.character(2014:2019)
quarters <- paste0("Q",1:4)

keyset <- expand.grid(
  region = c("Agria", "Induston")
  , period = sapply(years, paste0, quarters))

head(keyset)</pre>
```

```
region period
1 Agria 2014Q1
2 Induston 2014Q1
3 Agria 2014Q2
4 Induston 2014Q2
5 Agria 2014Q3
6 Induston 2014Q3
```

This key set will be referenced in the rule, and passed to confront as reference data.

```
# showing 7 columns of output for readability
summary(out)[1:7]
```

```
name items passes fails nNA error warning
1 V1 1199 899 300 0 FALSE FALSE
```

There are 300 fails. Inspecting the data set as above, we see that for Induston, the export is missing in "201803".

Finally, we do a strict test, to check that for each measure all periods and all regions are reported. We also demand that there are no more and no less records than for each individual measure. For this, the function contains_exactly can be used.

First create a keyset.

```
years <- as.character(2014:2019)
quarters <- paste0("Q",1:4)
keyset <- expand.grid(
 region = c(
   "Agria"
   . "Crowdon"
   . "Greenham"
   , "Induston"
   , "Mudwater"
   , "Newbay"
   , "Oakdale"
   , "Samplonia"
   , "Smokeley"
   , "Wheaton"
 ,period = c(years, sapply(years, paste0, quarters))
)
head(keyset)
```

```
region period
Agria 2014
Crowdon 2014
Greenham 2014
Induston 2014
Mudwater 2014
Newbay 2014
```

The keyset is passed as reference data to the rule using confront.

```
1 V1 1199 0 1199 0 FALSE FALSE
```

To find where the errors reside, we first select the records with an error and then find the unique measures that occur in those records.

```
erroneous_records <- violating(samplonomy, out)
unique(erroneous_records$measure)</pre>
```

```
[1] "gdp" "import" "export" "balance"
```

So here, blocks containing GDP and Export have entire records missing.

3.4 Gaps in (time) series

For time series, or possibly other series it is desirable that there is a constant distance between each two elements of the series. The mathematical term for such a series is called a *linear sequence*. Here are some examples of linear series.

- The natural numbers: 1, 2, 3, ...
- The even natural numbers 2, 4, 6, ...
- · Quarters periods: "2020Q1", "2020Q2", ...
- Years (these are just natural numbers): 2019, 2020, ...

The validate functions is_linear_sequence and in_linear_sequence check whether a variable represents a linear series, possibly in blocks defined by categorical variables. They can be used interactively or as a rule in a validator object. We first demonstrate how these functions work, and then give an example with the samplonomy dataset.

```
is_linear_sequence(c(1,2,3,4))
```

[1] TRUE

```
is_linear_sequence(c(8,6,4,2))
```

[1] TRUE

```
is_linear_sequence(c(2,4,8,16))
```

[1] FALSE

For character data, the function is capable of recognizing certain formats for time periods.

```
is_linear_sequence(c("2020Q1","2020Q2","2020Q3","2020Q4"))
```

[1] TRUE

See ?is_linear_sequence for a full specification of supported date-time formats.

It is not necessary for data to be sorted in order to be recognized as a linear sequence.

```
is_linear_sequence(c("2020Q4","2020Q2","2020Q3","2020Q1"))
```

[1] TRUE

One can force a begin and/or end point for the sequence as well.

```
is_linear_sequence(c("2020Q4","2020Q2","2020Q3","2020Q1")
    , begin = "2020Q2")
```

[1] FALSE

Finally it is possible to split a variable by one or more other columns and check whether each block represents a linear sequence.

```
series <- c(1,2,3,4,1,2,3,3)
blocks <- rep(c("a","b"), each = 4)
is_linear_sequence(series, by = blocks)</pre>
```

[1] FALSE

Now, this result is not very useful since now it is unknown which block is not a linear series. This is where the function in_linear_series comes in.

```
in_linear_sequence(series, by = blocks)
```

[1] TRUE TRUE TRUE TRUE FALSE FALSE FALSE

There are some subtleties. A single element is also a linear sequence (of length 1).

```
is_linear_sequence(5)
```

[1] TRUE

This can yield surprises in cases of blocks of length 1.

```
blocks[8] <- "c"
data.frame(series = series, blocks = blocks)</pre>
```

series blocks

- 1 1 a
- 2 2 a
- 3 3 8
- 4 4 a
- 5 1 b
- 6 2 b
- 7 3 b
- 8 3

in_linear_sequence(series, blocks)

С

[1] TRUE TRUE TRUE TRUE TRUE TRUE TRUE

We now have three linear series, namely

- For "a": 1,2,3,4
- For "b": 1,2,3
- For "c": 3.

We can circumvent this by giving explicit bounds.

```
in_linear_sequence(series, blocks, begin = 1, end = 4)
```

[1] TRUE TRUE TRUE FALSE FALSE FALSE

We now return to the samplonomy dataset. We wish to check that for each measure and each area, the time series are linear series. Since there are time series of different frequencies, we need to split the data by frequency as well.

```
name items passes fails nNA error warning
1 V1 1199 1170 29 0 FALSE FALSE
```

We can find the blocks where records are not in sequence as follows (output not printed here for brevity).

```
violating(samplonomy, out)
```

Inspection of the selected records shows that for Agria the GDP for 2015 is missing, and that for Induston the Export for 2018Q3 is missing while Export for 2018Q2 occurs twice (but with different values)

Chapter 4

Multivariate checks

In this Chapter we treat tests that involve relationships between variables.

Data

In this Chapter we will use the SBS2000 dataset that comes with validate.

```
library(validate)
data(SBS2000)
head(SBS2000, 3)
```

```
id size incl.prob staff turnover other.rev total.rev
1 RET01 sc0 0.02 75 NA
                                NA
                                       1130
2 RET02 sc3
            0.14 9
                        1607
                                NA
                                       1607
3 RET03 sc3
             0.14
                   NA
                         6886
                                -33
                                        6919
 staff.costs total.costs profit vat
      NA
             18915 20045 NA
      131
              1544
2
                     63 NA
3
      324
              6493
                     426 NA
```

4.1 Completeness of records

The functions is_complete() and all_complete() are convenience functions that test for missing values or combinations thereof in records.

```
name items passes fails nNA error warning
1
   ٧1
          60
                 60
                        0
                            0 FALSE
                                      FALSE
2
   ٧2
          60
                 56
                            0 FALSE
                                    FALSE
3
   ٧3
          60
                 52
                            0 FALSE
                                     FALSE
    ٧4
                  1
                        0
                            0 FALSE
                                      FALSE
```

Here, the first rule checks for missing data in the id variable, the second rule checks whether subrecords with id and turnover are complete, and the third rule checks whether subrecords with id, turnover and profit are complete. The output is one logical value (TRUE or FALSE) for each record.

The fourth rule tests whether all values are present in the id column, and it results in a single TRUE or FALSE.

- To test for missing values in individual variables, see also 2.2.
- · To check whether records are available at all, see 3.3.

4.2 Balance equalities and inequalities

Balance restrictions occur for example in economic microdata, where financial balances must be met.

```
rules <- validator(
    total.rev - profit == total.costs
, turnover + other.rev == total.rev
, profit <= 0.6*total.rev
)

out <- confront(SBS2000, rules)
summary(out)</pre>
```

```
name items passes fails nNA error warning
1
   ٧1
                            7 FALSE
                                      FALSE
   ٧2
          60
                 19
                        4 37 FALSE
                                      FALSE
2
    ٧3
                            5 FALSE
                                      FALSE
                        6
                                      expression
1 abs(total.rev - profit - total.costs) < 1e-08
2 abs(turnover + other.rev - total.rev) < 1e-08
            (profit - 0.6 * total.rev) <= 1e-08
```

Here, the first rule checks a balance between income, costs, and profit; the second rule checks a sub-balance, and the third rule is a plausibility check where we do not expect profit to exceed 60 per cent of the total revenue.

Observe that the expressions have been altered by validate to account for possible machine rounding differences. Rather than testing whether variable x equals variable y, validate will check $|x - y| \le \epsilon$, where the default value of ϵ is 10^{-8} . The value of this tolerance can be controlled for linear equalities and inequalities using respectively lin.eq.eps and lin.ineq.eps.

```
out <- confront(SBS2000, rules, lin.ineq.eps=0, lin.eq.eps=0.01)
summary(out)</pre>
```

```
name items passes fails nNA error warning
1
   ٧1
                             7 FALSE
                                       FALSE
                                     FALSE
2
   ٧2
          60
                 19
                         4 37 FALSE
3
    ٧3
          60
                 49
                             5 FALSE
                                       FALSE
```

```
expression

1 abs(total.rev - profit - total.costs) < 0.01

2 abs(turnover + other.rev - total.rev) < 0.01

3 profit <= 0.6 * total.rev
```

See 7.8 for more information on setting and resetting options.

4.3 Conditional restrictions

Conditional restrictions are all about demanding certain value combinations. In the following example we check that a business with staff also has staff costs.

```
rule <- validator(if (staff >= 1) staff.costs >= 1)
out <- confront(SBS2000, rule)
summary(out)</pre>
```

```
name items passes fails nNA error warning

1 V1 60 50 0 10 FALSE FALSE
expression

1 !(staff >= 1) | (staff.costs >= 1)
```

Here, combinations where there is a positive number of staff must be accompanied with a positive staff cost.

Validate translates the rule if (P) q to an expression of the form P | q. The reason for this is that the latter can be evaluated faster (vectorised).

The results are to be interpreted as follows. For each record, validate will check that cases where staff>=1 are accompanied by staff.costs >= 1. In cases where this test results in FALSE this means that either the staff number is too high, or the staff costs are too low. To be precise, the results of a conditional restriction match those of an implication in first-order logic as shown in the truth table below.

Ρ	Q	$P \Rightarrow Q$
T	Τ	Т
Τ	F	F
F	Τ	T
F	F	F

4.4 Forbidden value combinations

In some cases it is more convenient to have a list of forbidden (key) value combinations than specifying such combinations individually. The function does_not_contain() supports such situations.

As an example, let's first create some transaction data.

```
transactions <- data.frame(
    sender = c("S21", "X34", "S45","Z22")
, receiver = c("FG0", "FG2", "DF1","KK2")
, value = sample(70:100,4)
)</pre>
```

We assume that it is not possible for senders with codes starting with an "s" to send something to receivers starting with FG. A convenient way to encode such demands is to use globbing patterns. We create a data frame that lists forbidden combinations (here: one combination of two key patterns).

```
forbidden <- data.frame(sender="S*",receiver = "FG*")</pre>
```

Note that the column names of this data frame correspond to the columns in the transactions data frame. We are now ready to check our transactions data frame.

```
rule <- validator(does_not_contain(glob(forbidden_keys)))
out <- confront(transactions, rule, ref=list(forbidden_keys=forbidden))
## Suppress columns for brevity
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 4 3 1 0 FALSE FALSE
```

Observer that we use glob(forbidden_keys) to tell does_not_contain that the key combinations in the forbidden_keys must be interpreted as globbing patterns.

The records containing forbidden keys can be selected as follows.

It is also possible to use regular expression patterns, by labeling the forbidden key set with rx(). If no labeling is used, the key sets are interpreted as string literals.

Chapter 5

Statistical checks

Statistical checks involve group properties such as the means of columns. These characteristics can be checked for whole columns or grouped by one or more categorical variables. It is also possible to use groupwise computed statistics in validation rules. For example if you want to compare individual values with a mean within a group.

For long-form data it is possible to compare aggregate values with underlying details. For example to test whether quarterly time series add up to annual totals. It is also possible to check properties of groups, for example whether in every household (a group of persons) there is exactly one head of household.

Data

In this Chapter we will use the SBS2000 dataset that comes with validate.

```
library(validate)
data(SBS2000)
head(SBS2000, 3)
```

```
id size incl.prob staff turnover other.rev total.rev
1 RET01 sc0
                 0.02
                        75
                                 NA
                                          NA
                                                  1130
2 RET02 sc3
                 0.14
                        9
                               1607
                                          NA
                                                  1607
3 RET03 sc3
                 0.14
                        NA
                               6886
                                         -33
                                                  6919
  staff.costs total.costs profit vat
1
          NA
                  18915 20045
                                NA
                  1544
2
         131
                            63 NA
         324
                  6493 426 NA
```

We shall also use the samplonomy dataset that also comes with validate. See also 3.1.

```
data(samplonomy)
head(samplonomy, 3)
```

```
region freq period measure value
1 Agria A 2014 gdp 600000
2 Agria A 2014 import 210000
3 Agria A 2014 export 222000
```

5.1 Statistical and groupwise characteristics

Any R expression that ultimately is an equality or inequality check is interpreted as a validation rule by validate. This means that any statistical calculation can be input to a rule.

Here we check the mean profit and correlation coefficient between profit and turnover.

```
rule <- validator(
    mean(profit, na.rm=TRUIE) >= 1
   , cor(turnover, staff, use="pairwise.complete.obs") > 0
)
out <- confront(SBS2000, rule)
# suppress some columns for brevity
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 0 0 0 0 TRUE FALSE
2 V2 1 1 0 0 FALSE FALSE
```

There are a few helper functions to compute group-wise statistics, and to make comparing values with group aggregates possible.

For example, here we check whether each turnover is less than ten times the group-wise median.

```
rule <- validator(
  turnover <= 10*do_by(turnover, by=size, fun=median, na.rm=TRUE)
)
out <- confront(SBS2000, rule)
# suppress some columns for brevity
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 60 53 3 4 FALSE FALSE
```

Here, in the right-hand side of the rule the group-wise median of turnover is computed. The function do_by is very similar to functions such as tapply in base R. The difference is that do_by works on vectors only (not on data frames) and always repeats the values of fun so that the length of the output is equal to the length of the input.

```
medians <- with(SBS2000, do_by(turnover, by=size, fun=median, na.rm=TRUE))
head(data.frame(size = SBS2000$size, median=medians))</pre>
```

```
size median
1 sc0 351
2 sc3 2891
3 sc3 2891
4 sc3 2891
5 sc3 2891
6 sc0 351
```

There are also some convenience functions, including sum_by, mean_by, min_by, and max_by.

5.2. GROUP PROPERTIES 33

5.2 Group properties

In this section, we group data by one or more categorical variables and check for each group whether a rule is satisfied. In particular we are going to check whether each household in a small dataset has a unique 'head of household'.

We first create some data with household id (hhid) a person id (person) and that person's role in the household (hhrole).

```
d <- data.frame(
    hhid = c(1, 1, 2, 1, 2, 2, 3)
, person = c(1, 2, 3, 4, 5, 6, 7)
, hhrole = c("h","h","m","m","h","m")
)
d</pre>
```

```
hhid person hhrole
1
          1
2
    1
          2
3
    1
         4
4
                m
5
    2
         5
6
    2
          6
```

With exists_one() we can check that there is exactly one person with the role "h" (head) in each household, by grouping on household id.

```
rule <- validator(exists_one(hhrole == "h", by=hhid))
out <- confront(d, rule)
# suppress some columns for brevity
summary(out)</pre>
```

```
name items passes fails nNA error warning

1 V1 7 3 4 0 FALSE FALSE
expression

1 exists_one(hhrole == "h", by = hhid)
```

m

We can inspect the results by selecting the violating record groups.

```
violating(d, out)
```

We see that household 1 has two heads of household, while household 3 has no head of household.

To test whether at least one head of household exists, one can use exists_any:

```
violating(d, validator(exists_any(hhrole=="h",by=hhid) ))
```

```
hhid person hhrole
7 3 7 m
```

In the following example we check whether there is exactly one region called Samplonia for each period and each measure in the samplonomy dataset.

```
rule <- validator(exists_one(region=="Samplonia", by=list(period, measure)))</pre>
```

The first argument of exists_one() is a rule that has to be checked in every group indicated by the by argument. The output is a logical vector with an element for each record in the dataset under scrutiny. If a group of data fails the test, each record in that group is indicated as wrong (FALSE).

```
out <- confront(samplonomy, rule)
# suppress some columns for brevity
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 1199 1199 0 0 FALSE FALSE
```

Here, there are no groups that violate this assumption.

```
violating(samplonomy, out)
```

```
[1] region freq period measure value
<0 rows> (or 0-length row.names)
```

5.3 Code hierarchies and aggregation

Classifications and ontologies often have a hierarchical structure. A well-known example is the NACE classification of economic activities. In the NACE classification, the economy is divided into 10 basic types of activities such as 'Agriculture' or 'Mining and Quarrying', and each activity is again divided into subclasses, such as 'Growing of rice' and 'Growing of Grapes' under 'Agriculture'. The subdividing can go on for several levels. For statistics that describe an economy according to the NACE classification, it is desirable that the statistics of subclasses add up to their parent classes. This is what the function 'hierarchy' does in 'validate'.

The validate package comes with a version of the NACE classification (Revision 2, 2008) so we will use that as an example.

```
data(nace_rev2)
head(nace_rev2[1:4])
```

```
Order Level Code Parent
1 398481
            1
                  Α
2 398482
            2
                 91
3 398483
            3 01.1
                        Q1
4 398484
           4 01.11
                      01.1
5 398485
           4 01.12 01.1
6 398486
           4 01.13
                      01.1
```

The second and third column contain the necessary information: they list the parent for each NACE code (where each parent is also a NACE code). To demonstrate how hierarchy() works, we first create some example data.

```
nace volume
1 01 100
2 01.1 70
3 01.11 30
4 01.12 40
5 01.2 25
```

We see that the volumes for subclasses "01.11" and "01.12" add up to "01.1" (30 + 40 = 70). However, the volumes for "01.1" and "01.2" do not add up to the volume for "01" ($70 + 25 \neq 100$). The hierarchy() function checks all these relations.

Before using hierarchy in the setting of a validator object, we can examine it directly.

```
dat$check <- hierarchy(dat$volume, dat$nace, nace_rev2[3:4])
dat</pre>
```

```
        nace volume check

        1
        01
        100 FALSE

        2
        01.1
        70 FALSE

        3
        01.11
        30 TRUE

        4
        01.12
        40 TRUE

        5
        01.2
        25 FALSE
```

We see that hierarchy() returns a logical vector with one element for each record in the data. Each record that is involved in one or more aggregation checks that fail is labeled FALSE. Here, this concerns the records with labels "01", "01.1" and "01.2".

We will next look at a more complicated example, but first note the following. The hierarchy() function

- can handle any statistical aggregate, sum() is just the default;
- · supports globbing and regular expressions in the child values;
- has an adjustable tolerance value for comparing observed with computed aggregates;
- has configurable behaviour for cases of missing data;
- can be applied per-group, defined by one or more grouping variables (see next example).

See the help file ?hierarchy for specification and examples.

A more complicated example

Samplonia is divided in two districts, each of which is divided into several provinces. Let us define the hierarchical code list.

```
)
samplonia
   region
              parent
  Agria Samplonia
2 Induston Samplonia
3 Wheaton
              Agria
4 Greenham
               Agria
5 Smokely Induston
6 Mudwater Induston
   Newbay Induston
8 Crowdon Induston
Recall the structure of the samplonomy dataset.
data(samplonomy)
head(samplonomy)
  region freq period measure value
1 Agria
           A 2014
                         gdp 600000
2 Agria
           A 2014 import 210000
         A 2014 export 222000
3 Agria
4 Agria
         A 2014 balance 12000
5 Agria
           Q 2014Q1
                         gdp 60000
6 Agria
            Q 2014Q1 import 21000
We will check whether regions sum to their parent regions, for each period and for each measure.
rule <- validator(
 hierarchy(value, region, hierarchy=ref$codelist, by=list(period, measure))
out <- confront(samplonomy, rule, ref=list(codelist=samplonia))</pre>
summary(out)
  name items passes fails nNA error warning
                237 954 8 FALSE
  V1 1199
                                       TRUE
                                                                            expression
1 hierarchy(value, region, hierarchy = ref[["codelist"]], by = list(period, measure))
We see that some aggregates add up correctly, and some don't. There is also a warning which we should
investigate.
warnings(out)
$V1
[1] "Parent 'Induston' occurs more than once (2 times) in group (2018Q2, export)"
```

If one of the groups contains a parent more than once it is not possible to check whether child values add up to the aggregate. For this reason the duplicated parent and all it's children are marked FALSE. Indeed we find

```
measure == "export")
```

```
region freq period measure value
870 Induston Q 2018Q2 export 165900
871 Induston Q 2018Q2 export 170000
```

Just to see if we can remove the warning, let us remove the duplicate and re-run the check.

```
i <- !duplicated(samplonomy[c("region", "period", "measure")])
samplonomy2 <- samplonomy[i, ]

out <- confront(samplonomy2, rule, ref=list(codelist=samplonia))
# suppress some columns for brevity
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 1198 238 952 8 FALSE FALSE
```

The hierarchy() function marks every record FALSE that is involved in any check. This may make it hard to figure out which check it failed. One can get more detailed information, by checking different parts of the hierarchy in separate rules.

We can now select records involved in violating the highest level rules separately.

```
violating(samplonomy2, out["level0"])
```

```
region freq period measure value
260 Induston A 2015 gdp 1358000
340 Samplonia A 2015 gdp 1940000
814 Agria Q 2018Q3 export 118500
954 Samplonia Q 2018Q3 export 284400
```

From this it appears that in 2015, the GDP for Agria is missing, and in 2018Q3 there is no value for the export of Induston.

5.4 General aggregates in long-form data

Checking aggregations in long-form format is more involved than for data in wide format (as in Section 4.2).

Here, we check in the samplonomy dataset that for each measure and each period, the subregional data adds up to the regional data.

```
rules <- validator(
  part_whole_relation(value
   , labels=region
   , whole="Samplonia"
    , part =c("Agria", "Induston")
    , by=list(measure, period)
)
)</pre>
```

The first argument of part_whole_relation() is the name of the variable containing the values. Here, the column value from the samplonomy dataset. The argument labels indicates the variable that labels parts and wholes. Next, we define the label value that indicates a total. Here, a record with region label "Samplonia" indicates a total. Under argument part we specify the labels that have to add up to Samplonia, here the provinces Agria and Induston. Note that there are more subregions in the dataset, for example the district of Wheaton (a subregion of Agria). Since we do not specify them, these are ignored. In the by argument we specify that the dataset must be split into measure and period prior to checking the regional aggregates.

The output is one boolean value per record. For each block, defined by values of measure and period either all values are TRUE, FALSE, or NA. The latter indicates that the aggregate could not be computed because one of the values is missing, or the computed aggregate could not be compared with the aggregate in the data because it is missing (either the whole record may be missing, or the value may be NA).

```
out <- confront(samplonomy, rules)
# suppress some columns for brevity
summary(out)[1:7]</pre>
```

```
name items passes fails nNA error warning
1 V1 1199 1191 8 0 FALSE FALSE
```

We can extract the truth values and then inspect the blocks with erroneous values using standard R functionality.

```
violating(samplonomy, out)
```

```
region freq period measure
                                    value
260 Induston
                     2015
                              gdp 1358000
340 Samplonia
                     2015
                              gdp 1940000
810
       Agria
                Q 2018Q2 export
                                    47400
814
       Agria
                 Q 2018Q3
                           export
                                   118500
                 Q 2018Q2
                                  165900
870 Induston
                           export
871 Induston
                 Q 2018Q2
                                   170000
                           export
950 Samplonia
                 Q 2018Q2
                                   213300
                           export
954 Samplonia
                 Q 2018Q3
                           export 284400
```

Recall that the rule was executed per block defined by measure and period. Thus, the result indicates three errors: one in the block of records defined by measure=="gdp" and period=="2015", also in the blocks defined

by measure=="export" and period==2018Q2 Or period=="2018Q3".

First, it seems that the 2015 GDP of Agria is missing from the data set. This turns out indeed to be the case.

```
subset(samplonomy, region=="Agria" & period == "2015" & measure == "gdp")
[1] region freq period measure value
<0 rows> (or 0-length row.names)
```

Second, it can be seen that for Induston, there are two export values for "2018Q2" while the export value for "2018Q3" is missing.

Notes

Specifying (group-wise) aggregates is a fairly detailed job in the case of long data. There are a few things to keep in mind when using this function.

- The argument part is optional. If not specified, every record not matching with whole will be considered a detail that is to be used to compute the total. In the current example this was not possible because besides Agria and Induston, we have other subregions.
- In the example we used literal values to specify the keys that define parts and wholes. It is possible to, recognize patterns, for example any years (4 digits) as a whole and a quarter as a part. See also the next example. Supported patterns include regular expressions (shown here) and globbing (see help file).
- It is important that the variables listed in by (if any) uniquely specify a single aggregate. So here, for
 each measure and period, the label "Samplonia" should occur at most once (if it does not occur the
 result will be NA).
- The default way to aggregate is to take the sum. You can specify other ways to aggregate by passing an aggregator argument. For example aggregator=mean.
- By default, the aggregate in the data is compared with the computed aggregate up to a tolerance of 10⁻⁸. This tolerance can be set using the tol argument. E.g. for integer data you may want to set tol=0.

5.5 Aggregates of time series in long format

We are going to check whether quarterly time series add up to the annual time series. This is more complicated because of two subtleties.

First there is not one fixed aggregate key, like "samplonia". Rather, we have a key pattern. Each total is defined by a period label that consists of precisely four digits. So rather than recognizing a specific year we want to recognize that a key represents any year. This can be done using a regular expression of the form "^\\d{4}\s", where the ^ indicates 'start of string', the \\d{4} indicates 'four times a digit' and \\$ indicates 'end of string'.

Second, we wish to check annual totals against the sum over quarters for each region and each measure. However, a value-combination of measure and region does not single out a single value for year. For example, for the Induston export we have the following annual data.

```
subset(samplonomy, region=="Induston" & freq == "A" & measure=="export")
region freq period measure value
```

63 Induston A 2014 export 518000 262 Induston A 2015 export 525000

```
462 Induston A 2016 export 532000
662 Induston A
                 2017 export 560000
862 Induston A
                 2018 export 553000
1062 Induston
                  2019 export 553000
```

So in fact, we need to do the check by year as well as by measure and region. Fortunately, in this case it is easy to derive a variable that indicates the year by selecting the first four characters from period.

```
rules <- validator(part_whole_relation(value
  , labels = period
  , whole = rx("^{\dagger} {4})"
  , by = list(region, substr(period, 1, 4), measure)
out <- confront(samplonomy, rules)
```

We use rx("^\\d{4}") to tell part_whole_relation that this string must be interpreted as a regular expression. Here, we do not indicate part labels explicitly: by default any record not matching whole will be treated as a detail that must be used to compute the total.

```
errors(out)
named list()
# suppress some columns for brevity
summary(out)[1:7]
  name items passes fails nNA error warning
```

FALSE We now get 9 fails and 10 missing values. We can filter out records that have NA (lacking) results.

9 10 FALSE

```
lacking(samplonomy, out)
```

1180

V1 1199

```
region freq period measure value
24
    Crowdon
                  2014 balance 1600
28
    Crowdon
              Q 2014Q1 balance
                                 NA
32
    Crowdon Q 2014Q2 balance
                                 480
36
    Crowdon
              Q 2014Q3 balance
                                 480
40
    Crowdon
              Q 2014Q4 balance
                                 320
1181 Wheaton
                  2019 import 62000
1185 Wheaton Q 2019Q1 import 6200
1189 Wheaton Q 2019Q2 import
1193 Wheaton Q 2019Q3 import 31000
1197 Wheaton
              Q 2019Q4 import 12400
```

There are two blocks where the annual total could not be compared with the sum over quarterly series. The balance value of Crowdon is missing for "2014Q1" as well as the import value of Wheaton for "2019Q2".

Similarly, we can inspect the failing blocks

```
violating(samplonomy, out)
```

```
region freq period measure value
              Q 2015Q1
294
     Agria
                        gdp 58200
208
     Agria
              Q 2015Q2
                        gdp 116400
```

212	Agria	Q	2015Q3	gdp	291000
216	Agria	Q	2015Q4	gdp	116400
862	Induston	Α	2018	export	553000
866	Induston	Q	2018Q1	export	110600
870	Induston	Q	2018Q2	export	165900
871	Induston	Q	2018Q2	export	170000
878	Induston	Q	2018Q4	export	110600

Chapter 6

Indicators

Until now we have discussed various types of data validation rules: decisions that assign True or False values to a data frame. In some cases it is convenient to have a continuous value that can then be used in further assessing the data.

A practical example is the so-called selective editing approach to data cleaning. Here, each record in a data set is assigned a number that expresses the risk a record poses for inferring a faulty conclusion. Records are then ordered from high risk (records that both have suspicious values and large influence on the final result) to low risk (records with unsuspected values and little influence on the final result). Records with the highest risk are then scrutinized by domain experts.

In validate, an indicator is a rule that returns an numerical value. Just like validator objects are lists of validation rules, indicator objects are lists of indicator rules. Indices can be computed by confronting data with an indicator, and using add_indices, the computed indices can be added to the dataset. You can import, export, select, and combine indicator objects in the same way as validator objects.

6.1 A first example

Here is a simple example of the workflow.

```
library(validate)
ii <- indicator(
    BMI = (weight/2.2046)/(height*0.0254)^2
, mh = mean(height)
, mw = mean(weight))
out <- confront(women, ii)</pre>
```

In the first statement we define an indicator object storing indicator expressions. Next, we confront a dataset with these indicators. The result is an object of class indication. It prints as follows.

```
Object of class 'indication'
```

```
Call:

confront(dat = women, x = ii)
```

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Confrontations: 3
Warnings : 0
Errors : 0

To study the results, the object can be summarized.

```
summary(out)
```

```
name items
                  min
                           mean
                                      max nNA error warning
   BMI
          15 22.0967 22.72691
                                 24.03503
                                            0 FALSE
                                                       FALSE
2
   mh
           1 65.0000 65.00000 65.00000
                                            0 FALSE
                                                       FALSE
3
           1 136.7333 136.73333 136.73333
                                            0 FALSE
                                                       FALSE
                           expression
1 (weight/2.2046)/(height * 0.0254)^2
2
                         mean(height)
3
                         mean(weight)
```

Observe that the first indicator results in one value per record while the second and third indicators (mh, mw) each return a single value. The single values are repeated when indicator values are added to the data.

```
head(add_indicators(women, out), 3)
```

```
height weight BMI mh mw
1 58 115 24.03503 65.0000 136.7333
2 59 117 23.63114 136.7333 65.0000
3 60 120 23.43589 65.0000 136.7333
```

The result is a data frame with indicators attached.

The columns error and warning indicate whether calculation of the indicators was problematic. For example because the output of an indicator rule is not numeric, or when it uses variables that do not occur in the data. Use warnings(out) or errors(out) to obtain the warning and error messages per rule.

6.2 Getting indicator values

Values can be obtained with the values function, or by converting to a data.frame. In this example we add a unique identifier (this is optional) to make it easier to identify the results with data afterwards.

```
women$id <- letters[1:15]</pre>
```

Compute indicators and convert to data.frame.

```
out <- confront(women, ii,key="id")
tail( as.data.frame(out) )</pre>
```

```
id name
                 value
                                                 expression
         BMI 22.15113 (weight/2.2046)/(height * 0.0254)^2
12
              22.09670 (weight/2.2046)/(height * 0.0254)^2
13
         BMI
            22.17600 (weight/2.2046)/(height * 0.0254)^2
14
         BMI
15
         BMI 22.24240 (weight/2.2046)/(height * 0.0254)^2
             65.00000
                                               mean(height)
16 < NA>
          mh
17 <NA>
         mw 136.73333
                                               mean(weight)
```

45

Observe that there is no key for indicators ${\tt mh}$ and ${\tt mw}$ since these are constructed from multiple records.

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Chapter 7

Working with validate

7.1 Reading rules from file

It is a very good idea to store and maintain rule sets outside of your R script. Validate supports two file formats: simple text files and yaml files. Here we only discuss simple text files, yaml files are treated in 7.4.

To try this, copy the following rules into a new text file and store it in a file called myrules.R, in the current working directory of your R session.

```
# basic range checks
speed >= 0
dist >= 0

# ratio check
speed / dist <= 1.5</pre>
```

Note that you are allowed to annotate the rules as you would with regular R code. Reading these rules can be done as follows.

```
rules <- validator(.file="myrules.R")</pre>
```

7.2 Manipulating rule sets

Validate stores rulesets into something called a validator object. The validator() function creates such an object.

```
v <- validator(speed >= 0, dist>=0, speed/dist <= 1.5)
v</pre>
```

```
Object of class 'validator' with 3 elements:

V1: speed >= 0

V2: dist >= 0

V3: speed/dist <= 1.5
```

Validator objects behave a lot like lists. For example, you can select items to get a new validator. Here, we select the first and third element.

```
w \leftarrow v[c(1,3)]
```

Here w is a new validator object holding only the first and third rule from v. If not specified by the user, rules are given the default names "v1", "v2", and so on. Those names can also be used for selecting rules.

```
w <- v[c("V1","V3")]
```

Validator objects are reference objects. This means that if you do

```
w <- v
```

then w is not a copy of v. It is just another name for the same physical object as v. To make an actual copy, you can select everything.

```
w <- v[]
```

It is also possible to concatenate two validator objects. For example when you read two rule sets from two files (See 7.1). This is done by adding them together with +.

```
rules1 <- validator(speed>=0)
rules2 <- validator(dist >= 0)
all_rules <- rules1 + rules2</pre>
```

An empty validator object is created with validator().

If you select a single element of a validator object, an object of class 'rule' is returned. This is the validating expression entered by the user, plus some (optional) metadata.

```
v[[3]]
```

Object of class rule.

expr : speed/dist <= 1.5

name : V3
label :
description:

origin : command-line

created : 2020-12-09 12:18:03

meta : language<chr>, severity<chr>

Users never need to manipulate rule objects, but it can be convenient to inspect them. As you see, the rules have some automatically created metadata. In the next section we demonstrate how to retrieve and set the metadata.

7.3 Rule metadata

Validator objects behave a lot like lists. The only metadata in an R list are the names of its elements. You can get and set names of a list using the names<- function. Similarly, there are getter/setter functions for rule metadata.

- origin(): Where was a rule defined?
- · names(): The name per rule

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```
• created(): when were the rules created?
```

- · label(): Short description of the rule
- description(): Long description of the rule
- meta(): Set or get generic metadata

Names can be set on the command line, just like how you would do it for an R list.

```
rules <- validator(positive_speed = speed >= 0, ratio = speed/dist <= 1.5)
rules</pre>
```

```
Object of class 'validator' with 2 elements:

positive_speed: speed >= 0

ratio : speed/dist <= 1.5
```

Getting and setting names works the same as for lists.

```
names(rules)
```

```
[1] "positive_speed" "ratio"
names(rules)[1] <- "nonnegative_speed"</pre>
```

The functions origin(), created(), label(), and description() work in the same way. It is also possible to add generic key-value pairs as metadata. Getting and setting follows the usual recycling rules of R.

```
# add 'foo' to the first rule:
meta(rules[1], "foo") <- 1
# Add 'bar' to all rules
meta(rules, "bar") <- "baz"</pre>
```

Metadata can be made visible by selecting a single rule:

```
v[[1]]
```

```
Object of class rule.

expr : speed >= 0
name : V1
label :
```

description:
origin : command-line

created : 2020-12-09 12:18:03

meta : language<chr>, severity<chr>

Or by extracting it to a data.frame

```
meta(v)
```

```
3 validate 1.0.1 error
```

Some general information is obtained with summary,

```
summary(v)
```

```
block nvar rules linear
```

Here, some properties per *block* of rules is given. Two rules occur in the same block if when they share a variable. In this case, all rules occur in the same block.

The number of rules can be requested with length

```
length(v)
```

[1] 3

With variables, the variables occurring per rule, or over all the rules can be requested.

```
variables(v)
```

```
[1] "speed" "dist"
variables(v,as="matrix")
```

```
variable
rule speed dist
V1 TRUE FALSE
V2 FALSE TRUE
V3 TRUE TRUE
```

7.4 Metadata in text files: YAML

YAML is a data format that aims to be easy to learn and human-readable. The name 'YAML' is a recursive acronym that stands for

YAML Ain't Markup Language.

Validate can read and write rule sets from and to YAML files. For example, paste the following code into a file called myrules.yaml.

```
rules:
```

```
- expr: speed >= 0
  name: 'speed'
  label: 'speed positivity'
  description: |
    speed can not be negative
  created: 2020-11-02 11:15:11
  meta:
    language: validate 0.9.3.36
    severity: error
- expr: dist >= 0
  name: 'dist'
  label: 'distance positivity'
```

```
description: |
    distance cannot be negative.
  created: 2020-11-02 11:15:11
 meta:
    language: validate 0.9.3.36
    severity: error
- expr: speed/dist <= 1.5
  name: 'ratio'
  label: 'ratio limit'
  description: |
   The speed to distance ratio can
    not exceed 1.5.
  created: 2020-11-02 11:15:11
 meta:
    language: validate 0.9.3.36
    severity: error
```

We can read this file using validator(.file=) as before.

```
rules <- validator(.file="myrules.yaml")
rules</pre>
```

```
Object of class 'validator' with 3 elements:

speed [speed positivity] : speed >= 0

dist [distance positivity]: dist >= 0

ratio [ratio limit] : speed/dist <= 1.5
```

Observe that the labels are printed between brackets. There are a few things to note about these YAML files.

- rules: starts a list of rules.
- 2. Each new rule starts with a dash (-)
- Each element of a rule is denoted name: <content>. The only obligated element is expr: the rule expression.
- Spaces matter. Each element of a rule must be preceded by a newline and two spaces. Subelements
 (as in meta) are indented again.

A full tutorial on YAML can be found at W3Cschools.io.

To export a rule set to yaml, use the export_yaml() function.

```
rules1 <- rules[c(1,3)]
export_yaml(rules1, file="myrules2.yaml")</pre>
```

We will return extensively to reading rules from YAML or other text files in Chapter 8.

7.5 Rules in data frames

You can read and write rules and their metadata from and to data frames. This is convenient, for example in cases where rules are retrieved from a central rule repository in a data base.

Exporting rules and their metadata can be done with as.data.frame.

```
rules <- validator(speed >= 0, dist >= 0, speed/dist <= 1.5)
df <- as.data.frame(rules)</pre>
```

Reading from a data frame is done through the .data argument.

```
rules <- validator(.data=df)</pre>
```

It is not necessary to define all possible metadata in the data frame. It is sufficient to have three character columns, named rule, name and description in any order.

7.6 Validation rule syntax

Conceptually, any R statement that will evaluate to a logical is considered a validating statement. The validate package checks this when the user defines a rule set, so for example calling validator(mean(height)) will result in a warning since just computing mean(x) does not validate anything.

You will find a concise description of the syntax in the syntax help file.

?syntax

In short, you can use

- · Type checks: any function starting with is.
- Binary comparisons: <, <=, ==, !=, >=, > and %in%
- Unary logical operators: !, all(), any()
- Binary logical operators: &, &&, |, || and logical implication, e.g. if (staff > 0) staff.costs > 0
- Pattern matching grep1
- Functional dependency: $X \rightarrow Y + Z$ is represented by $X \sim Y + Z$.

There are some extra syntax elements that help in defining complex rules.

- Inspect the whole data set using ., e.g. validator(nrow(.) > 10).
- Reuse a variable using :=, e.g. validator(m := mean(x), x < 2*m).
- Apply the same rule to multiple groups with var_group. For example validator(G:=var_group(x,y),
 G > 0) is equivalent to validator(x>0, y>0).

A few helper functions are available to compute groupwise values on variables (vectors). They differ from functions like aggregate or tapply in that their result is always of the same length as the input.

```
sum_by(1:10, by = rep(c("a", "b"), each=5))
```

[1] 15 15 15 15 15 40 40 40 40 40

This is useful for rules where you want to compare individual values with group aggregates.

function	computes		
do_by	generic groupwise calculation		
sum_by	groupwise sum		
min_by, max_by	groupwise min, max		
mean_by	groupwise mean		
median_by	groupwise median		

See also Section 5.1.

There are a number of functions that perform a particular validation task that would be hard to express with basic syntax. These are treated extensively in Chapters 2 to 5, but here is a quick overview.

function	checks
in_range	Numeric variable range
is_unique	Uniqueness of variable combinations
all_unique	Equivalent to all(is_unique())
is_complete	Completeness of records
all_complete	Equivalent to all(is_complete())
exists_any	For each group, check if any record satisfies a rule
exists_one	For each group, check if exactly one record satisfies a rule
is_linear_sequence	Linearity of numeric or date/time/period series
in_linear_sequence	Linearity of numeric of date/time/period series
hierarchy	Hierarchical aggregations
part_whole_relation	Generic part-whole relations
field_length	Field length
number_format	Numeric format in text fields
field_format	Field format
contains_exactly	Availability of records
contains_at_least	Availability of records
contains_at_most	Availability of records
does_not_contain	Correctness of key combinations

7.7 Confrontation objects

The outcome of confronting a validator object with a data set is an object of class confrontation. There are several ways to extract information from a confrontation object.

- summary: summarize output; returns a data.frame
- aggregate: aggregate validation in several ways
- sort : aggregate and sort in several ways
- · values: Get the values in an array, or a list of arrays if rules have different output dimension structure
- errors: Retrieve error messages caught during the confrontation
- warnings: Retrieve warning messages caught during the confrontation.

By default aggregates are produced by rule.

```
v <- validator(height>0, weight>0, height/weight < 0.5)
cf <- confront(women, rules)
aggregate(cf)</pre>
```

NULL

To aggregate by record, use by='record'

```
head(aggregate(cf,by='record'))
```

Aggregated results can be automatically sorted, so records with the most violations or rules that are violated most sort higher.

```
# rules with most violations sorting first:
sort(cf)
```

NULL

Confrontation objects can be subsetted with single bracket operators (like vectors), to obtain a sub-object pertaining only to the selected rules.

```
summary(cf[c(1,3)])
```

7.8 Confrontation options

By default, all errors and warnings are caught when validation rules are confronted with data. This can be switched off by setting the raise option to "errors" or "all". The following example contains a specification error: hite should be height and therefore the rule errors on the women data.frame because it does not contain a column hite. The error is caught (not resulting in a R error) and shown in the summary,

```
v <- validator(hite > 0, weight>0)
summary(confront(women, v))
```

```
name items passes fails nNA error warning expression

1 V1 0 0 0 0 TRUE FALSE hite > 0

2 V2 15 15 0 0 FALSE FALSE weight > 0
```

Setting raise to all results in a R error:

```
# this gives an error
confront(women, v, raise='all')
```

```
Error in fun(...): object 'hite' not found
```

Linear equalities form an important class of validation rules. To prevent equalities to be strictly tested, there is an option called lin.eq.eps (with default value 10⁻⁸) that allows one to add some slack to these tests. The amount of slack is intended to prevent false negatives (unnecessary failures) caused by machine rounding. If you want to check whether a sum-rule is satisfied to within one or two units of measurement, it is cleaner to define two inequalities for that.

7.9 Using reference data

For some checks it is convenient to compare the data under scrutiny with other data artifacts. Two common examples include:

- Data is checked against an earlier version of the same dataset.
- We wish to check the contents of a column against a code list, and we do not want to put the code list hard-coded into the rule set

For this, we can use the ref option in confront. Here is how to compare columns from two data frames row-by-row. The user has to make sure that the rows of the data set under scrutiny (women) matches row-wise with the reference data set (women1).

```
women1 <- women
rules <- validator(height == women_reference$height)</pre>
cf <- confront(women, rules, ref = list(women_reference = women1))</pre>
summary(cf)
 name items passes fails nNA error warning
1 V1 15 15 0 0 FALSE FALSE
                             expression
1 height == women_reference[["height"]]
Here is how to make a code list available.
rules <- validator( fruit %in% codelist )</pre>
fruits <- c("apple", "banana", "orange")</pre>
dat <- data.frame(fruit = c("apple", "broccoli", "orange", "banana"))</pre>
cf <- confront(dat, rules, ref = list(codelist = fruits))</pre>
summary(cf)
 name items passes fails nNA error warning
         4 3 1 0 FALSE FALSE
           expression
1 fruit %vin% codelist
```

Chapter 8

Rules in text files

This Chapter is about importing and exporting rules from and to file, both in free-form text and in YAML. We also discuss some more advanced features like how to have one rule file include another file.

- Importing from free-form text files was discussed in @ref{sect:readfromfile}
- Importing from simple YAML files was discussed in @ref{sect:yamlfiles}

8.1 Setting options

Both free-form and YAML files can optionally start with a header section where options or file inclusions can be set. The header section is enclosed by lines that contain three dashes (---) at the beginning of the line.

For example, in the following rule file we make sure that errors are not caught but raised to run-time level, and we set the tolerance for checking linear equalities and inequalities to zero.

```
options:
    raise: errors
    lin.eq.eps: 0
    lin.ineq.eps: 0
---

turnover >= 0

staff >= 0

total.rev - profit == total.costs
```

The options you set here will be part of the validator object, that is created once you read in the file. The options are valid for every confrontation you use this validator for, unless they are overwritten during the call to confront().

The header section is interpreted as a block of YAML, so options and file inclusions must be specified in that format

8.2 Including other rule files

In validate, rule files can include each other recursively. So file A can include file B, which may inclide file C. This is useful for example in surveys where the first part of the questionnaire goes to all respondents, and for the second part, the contents of the questionnaire (and hence its variables) depend on the respondent type. One could create a files with specific rules for the second part: one for each respondent group, and have each specific rule file include the general rules that must hold for every respondent. It can also be useful when different persons are responsible for different rule sets.

File inclusion can be set through the include option in the YAML header.

```
include:
    - petes_rules.yaml
    - nancys_rules.yaml
options:
    raise: errors
---
# start rule definitions here
```

8.3 Exporting validator objects

There are three ways to do that. You can either write to a yaml file immediately as follows

```
v <- validator(height>0, weight> 0)
export_yaml(v, file="my_rules.yaml")

or you can get the yaml text string using as_yaml
cat(as_yaml(v))

Finally, you can convert a rule set to data frame and then export it to a database.
```

```
df <- as.data.frame(v)
```

Chapter 9

Comparing data sets

When processing data step by step, it is useful to gather information on the contribution of each step to the final result. This way the whole process can be monitored and the contribution of each step can be evaluated. Schematically, a data processing step can be visualised as follows.

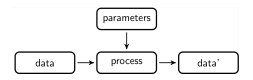


Figure 9.1: Data, process, changed data

Here, some input data is processed by some procedure that is parameterized, usually by domain experts. The output data is again input for a next step.

In the following two sections we discuss two methods to compare two or more versions of a data set. In the last section we demonstrate how validate can be combined with the lumberjack package to automate monitoring in an R script.

9.1 Cell counts

One of the simplest ways to compare different versions of a data set is to count how many cells have changed. In this setting it can be useful to distinguish between changes from available to missing data (and *vice versa*) and changes between data where the values change. When comparing two data sets, say the input and the output data, the total number of cells can be decomposed according to the following schema.

The total number of cells (fields) in the output data can be decomposed into those cells that are filled (available) and those that are empty (missing). The missing ones are decomposed into those that were already missing in the input data and those that are still missing. Similarly, the available values can be decomposed into those that were missing before and have been imputed. And those that already were available can be decomposed in those that are the same as before (unadapted) and those that ave been changed (adapted).

With the validate package, these numbers can be computed for two or more datasets using cells(). As an example, we first create three versions of the sbs2000 dataset. The first version is just the unaltered data. In

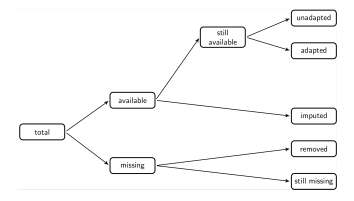


Figure 9.2: decomposition of output fields

the second version we replace a revenue column with it's absolute value to 'repair' cases with negative revenues. In the third version, we impute cases where turnover is missing with the vat (value added tax) value, when available.

```
library(validate)
data(SBS2000)
original <- SBS2000
version2 <- original
version2$other.rev <- abs(version2$other.rev)
version3 <- version2
version3$turnover[is.na(version3$turnover)] <- version3$vat[is.na(version3$turnover)]</pre>
```

We can now compare version2 and version3 to the original data set as follows.

```
cells(input = original, cleaned = version2, imputed = version3)
```

Object of class cellComparison:

```
cells(input = original, cleaned = version2, imputed = version3)
```

	input	cleaned	imputed
cells	660	660	660
available	580	580	581
still_available	580	580	580
unadapted	580	579	579
adapted	0	1	1
imputed	0	0	1
missing	80	80	79
still_missing	80	80	79
removed	0	0	0

The cells function accepts an arbitrary number of name=data frame arguments. The names provided by the user are used as column names in the output. From the output we see that the cleaned data set (version2) and in the imputed data set (version3) have one adapted value compared to the original data. Similarly, no imputations took place in preparing the cleaned data set, but a single value was imputed in the imputed

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dataset.

Since each data frame is compared to the first data frame, the last column can be considered a 'cumulative' record of all changes that took place from beginning to end. It is also possible to print differential changes, where each data set is compared with the previous one.

```
cells(input = original, cleaned = version2, imputed = version3
, compare="sequential")
```

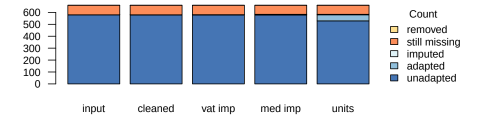
Object of class cellComparison:

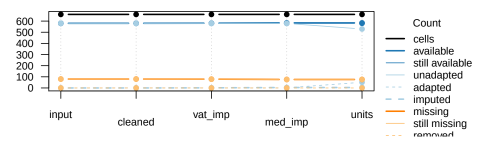
```
cells(input = original, cleaned = version2, imputed = version3, compare = "sequential")
```

input	cleaned	imputed
660	660	660
580	580	581
580	580	580
580	579	580
0	1	0
0	0	1
80	80	79
80	80	79
0	0	0
	580 580 580 580 0 0 80	580 580 580 580 580 579 0 1 0 0 80 80

plot(out)

The output of cells() is an array of class cellcomparison. The most interesting about this is that validate comes with two plot methods for such objects. To demonstrate this, we will create two more versions of the SBS2000 dataset.





The bar plot and line plot convey the same information. The line plot is better when the data sets are instances resulting from a sequential process. The bar plot can be used more generally since it does not suggest a particular order.

9.2 Comparing rule violations

When processing data it is interesting to compare how many data validations are violated before and after a processing step. Comparing output data with input data, we can decompose the total number of validation results of the output data as follows.

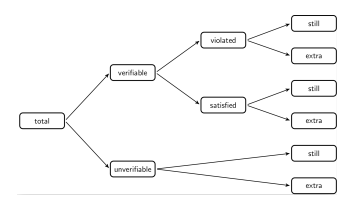


Figure 9.3: decomposition of validation output

The total number of validation results in the output data van be split into those that are verifiable (TRUE or FALSE) and those that are unverifiable (NA). The unverifiable cases can be split into those that were also unverifiable in the input data (still) and those that were verifiable in the input data but can now not be verified, because certain fields have been emptied. The verifiable cases can be split into those that yielded FALSE (violated) and

those that yielded TRUE (satisfied). Each can be split into cases that stayed the same or changed with respect to the input data.

With validate the complete decomposition can be computed with compare(). It takes as first argument a validator object and two or more data sets to compare. We will use the data sets developed in the previous paragraph.

Object of class validatorComparison:

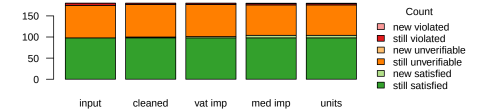
```
compare(x = rules, input = original, cleaned = version2, vat_imp = version3, med_imp = version4, units
```

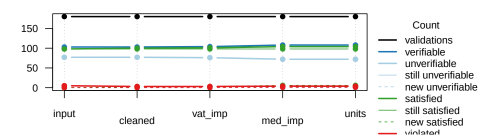
,	Version	า			
Status	input	cleaned	vat_imp	${\sf med_imp}$	units
validations	180	180	180	180	180
verifiable	103	103	104	108	108
unverifiable	77	77	76	72	72
still_unverifiable	77	77	76	72	72
new_unverifiable	0	0	0	0	0
satisfied	98	100	101	104	104
still_satisfied	98	98	98	98	98
new_satisfied	0	2	3	6	6
violated	5	3	3	4	4
still_violated	5	3	3	3	3
new_violated	0	0	0	1	1

By default each data set is compared to the first dataset (input=original). Hence the last column represents the cumulative change of all processing steps since the first data set. It is possible to investigate local differences by setting how='sequential'.

It is possible to plot the output for a graphical overview in two different ways: a bar plot and a line plot.

```
par(mfrow=c(2,1))
barplot(comparison)
plot(comparison)
```





9.3 validate and lumberjack

The lumberjack package makes it easy to track changes in data in a user-defined way. The following example is slightly adapted from the JSS paper.

We create a script that reads data, performs a few data cleaning steps and then writes the output. The script is stored in clean_supermarkets.R and has the following code.

```
## Contents of clean_supermarkets.R
library(validate)

# 1. simulate reading data
data(SBS2000)
spm <- SBS2000[c("id", "staff", "turnover", "other.rev", "total.rev")]

# 2. add a logger from 'validate'
start_log(spm, logger=lbj_cells())

# 3. assume empty values should be filled with 0
spm <- transform(spm, other.rev = ifelse(is.na(other.rev),0,other.rev))

# 4. assume that negative amounts have only a sign error
spm <- transform(spm, other.rev = abs(other.rev))

# 5a. ratio estimator for staff conditional on turnover
Rhat <- with(spm, mean(staff,na.rm=TRUE)/mean(turnover,na.rm=TRUE))

# 5b. impute 'staff' variable where possible using ratio estimator
spm <- transform(spm, staff = ifelse(is.na(staff), Rhat * turnover, staff))</pre>
```

```
# 6. write output
write.csv(spm, "supermarkets_treated.csv", row.names = FALSE)
```

In the first section we do not actually read data from a data source but take a few columns from the SBS2000 data set that comes with the validate package. The data to be processed is stored in a variable called spm. Next, in section two, we use the lumberjack function start_log() to attach a logging object of type lbj_cells() to the data under scrutiny. Two things are of note here:

- The call to library(validate) is necessary to be able to use lbj_cells(). Alternatively you can use validate::lbj_cells().
- 2. It is not necessary to load the lumberjack package in this script (although it is no problem if you do).

In sections three and four, values for other revenue are imputed and then forced to be nonnegative. In section 5 a ratio model is used to impute missing staff numbers. In section 7 the output is written.

The purpose of the lbh_cells() logger is to record the output of cells() after each step. To make sure this happens, run this file using run_file() from the lumberjack package.

```
library(lumberjack)
run_file('clean_supermarkets.R')
```

Dumped a log at /home/mark/projects/validate/cookbook/spm_lbj_cells.csv

This command executed all code in clean_supermarkets.R, but run_file() also ensured that all changes in the spm variable were recorded and logged using lbj_cells(). The output is written to a csv file which we can read.

```
logfile <- read.csv("spm_lbj_cells.csv")</pre>
```

The logfile variable has quite a lot of columns, so here show just two rows.

```
logfile[3:4,]
```

```
time
  step
     2 2020-12-09 12:18:05
3
     3 2020-12-09 12:18:05
                                                                   expression
3 spm <- transform(spm, other.rev = ifelse(is.na(other.rev),0,other.rev))</pre>
                         spm <- transform(spm, other.rev = abs(other.rev))</pre>
4
  cells available still_available unadapted adapted imputed
3
    300
               288
                                252
                                          252
               288
                                288
                                          287
    300
                                                     1
  missing still_missing removed
3
       12
                      12
                                0
4
       12
                      12
                                0
```

Each row in the output lists the step number, a time stamp, the expression used to alter the contents of the variable under scrutiny, and all columns computed by cells(). Since the logger always compares two consecutive steps, these numbers are comparable to using cells(comapare='sequential'). For example, we see that after step four, one value was adapted compared to the state after step three. And in step three, 36 values were imputed compared to the state created by step 2. In step four, no values were imputed.

It is also interesting to follow the progression of rule violations as the spm dataset gets processed. This can

be done with the <code>lbj_rules()</code> logger that is exported by <code>validate</code>. Since <code>lumberjack</code> allows for multiple loggers to be attached to an R object, we alter the first part of the above script as follows, and store it in <code>clean_supermarkets2.R</code>

Running the file again using lumberjack, we now get two log files.

```
run_file("clean_supermarkets2.R")
```

Dumped a log at /home/mark/projects/validate/cookbook/spm_lbj_cells.csv

Dumped a log at /home/mark/projects/validate/cookbook/spm_lbj_rules.csv

Let's read the log file from spm_lbj_rules.csv and print row three and four.

```
read.csv("spm_lbj_rules.csv")[3:4,]
  sten
3
     2 2020-12-09 12:18:05
     3 2020-12-09 12:18:05
3 spm <- transform(spm, other.rev = ifelse(is.na(other.rev),0,other.rev))</pre>
4
                         spm <- transform(spm, other.rev = abs(other.rev))</pre>
  validations verifiable unverifiable still_unverifiable
3
          240
                      225
                                     15
4
          240
                      225
                                     15
                                                         15
  new_unverifiable satisfied still_satisfied new_satisfied
3
                 A
                          218
                                          152
                                                           66
                  0
                          220
                                           218
                                                            2
  violated still_violated new_violated
3
         7
                         5
         5
                         5
                                       0
```

We get the full output created by validate::compare(). For example we see that after step three, 66 new cases satisfy one of the checks while two new violations were introduced. The fourth step adds two new

satisfied cases and no new violations. The total number of violations after four steps equals five.

Until now the logging data was written to files that were determined automatically by lumberjack. This is because lumberjack automatically dumps logging data after processing executing the file when the user has not done so explicitly. You can determine where to write the logging data by adding a stop_log() statement anywhere in your code (but at the end would usually make most sense).

For example, add the following line of code at the end of clean_supermarkets2.R to write the output of the lbj_rules logger to my_output.csv.

```
stop_log(spm, logger="lbj_rules",file="my_output.csv")
```

The format and way in which logging data is exported is fixed by the logger. So <code>lbj_rules()</code> and <code>lbj_cells()</code> can only export to csv, and only the data we've seen so far. The good news is that the <code>lumberjack</code> package itself contains other loggers that may be of interest, and it is also possible to develop your own logger. So it is possible to develop loggers that export data to a database. See the lumberjack paper for a short tutorial on how to write your own logger.

Bibliographical notes

More background on the validate package can be found in the paper for the R Journal.

MPJ van der Loo and E de Jonge (2020). Data Validation Infrastructure for R. *Journal of Statistical Software* (Accepted).

The theory of data validation is described in the following paper.

MPJ van der Loo, and E de Jonge (2020). Data Validation Infrastructure for R. *In Wiley StatsRef: Statistics Reference Online (eds N. Balakrishnan, T. Colton, B. Everitt, W. Piegorsch, F. Ruggeri and J.L. Teugels*).

Data validation is described in the wider context of data cleaning, in Chapter 6 of the following book.

MPJ van der Loo and E de Jonge (2018) Statistical Data Cleaning With Applications in R. *John Wiley & Sons, NY*.

The following document describes data validation in the context of European Official Statistics. It includes issues such as lifecycle management, complexity analyses and examples from practice.

M. Zio, N. Fursova, T. Gelsema, S. Giessing, U Guarnera, J. Ptrauskiene, Q. L. Kalben, M. Scanu, K. ten Bosch, M. van der Loo, and K. Walsdorfe (2015) Methodology for data validation. *Deliverable of the ESSNet on validation*.

The lumberjack package discussed in Chapter 9 is described in the following paper.

MPJ van der Loo (2020). Monitoring Data in R with the lumberjack package. *Journal of Statistical Software*, Accepted for publication.