

Introduction to Data Manipulation





Workshop Setup:

Wi-Fi

- ▶ Network Name: N/A
- ▶ Password: N/A


Resources

- ▶ R (version 4.0.1) 
- ▶ RStudio (version 1.3.959)  Studio


Packages

- ▶ tidyverse 

What is tidyverse?

Tidyverse is a collection of  packages that are designed for data science tasks, more specifically for data manipulation, transformation, exploration and visualisation.

These packages share a common design philosophy and contain functions that are consistent and uniform in coding style.




 You can read more at <https://www.tidyverse.org/>

Topics

► Workshop aim:

Learn how to do data manipulations using tidyverse packages.

► Topics:

- Learn the “verbs” with 
- Improve your workflow with 
- Simple string manipulation using 

Learn the “verbs” with



One of the most commonly used R packages when dealing with data manipulations is **{dplyr}**. It is very powerful in handling tabular data such as data frames and is easy to use through “verb” functions. You can use **{dplyr}** to:

- ▶ **Select** columns from your data
- ▶ **Filter** your data to keep the rows that meet some conditions
- ▶ **Arrange** your data in some order
- ▶ **Mutate** your data and create new columns
- ▶ **Group** and **summarise** your data

```
library(tidyverse)
```

```
view(starwars)
```

Live Coding Example 1



Use the `starwars` dataset from the `{dplyr}` package to:

1. Select the columns: “name”, “height”, “mass”, “species”.
2. Filter the rows to keep only those characters that are greater than or equal to 175cm.
3. Filter the rows to keep only the “Human” characters.
4. Arrange the rows according to descending “mass” values.
5. Who is the character on the first row?

Live Coding Example 1

selected columns

arranged rows

Who is the character
on the first row?



	name	height	mass	species
1	Darth Vader	202	136.0	Human
2	Owen Lars	178	120.0	Human
3	Jek Tono Porkins	180	110.0	Human
4	Qui-Gon Jinn	193	89.0	Human
5	Gregar Typho	185	85.0	Human
6	Biggs Darklighter	183	84.0	Human
7	Anakin Skywalker	188	84.0	Human
8	Mace Windu	188	84.0	Human
9	Han Solo	180	80.0	Human
10	Dooku	193	80.0	Human
11	Lando Calrissian	177	79.0	Human
12	Lobot	175	79.0	Human
13	Jango Fett	183	79.0	Human
14	Raymus Antilles	188	79.0	Human
15	Boba Fett	183	78.2	Human
16	Obi-Wan Kenobi	182	77.0	Human
17	Wilhuff Tarkin	180	NA	Human
18	Clegg Lars	183	NA	Human
19	Bail Prestor Organa	191	NA	Human

filtered rows

Live Coding Example 1

```
# select columns
df <- select(starwars, name, height, mass, species)

# Filter rows by height condition
df <- filter(df, height >= 175)

# Filter rows by species condition
df <- filter(df, species == "Human")

# Arrange rows by descending mass
df <- arrange(df, desc(mass))
```


Mutate your data

Very often you will want to create new columns from your existing data. The function **mutate()** in the **{dplyr}** package can be used to do exactly this task.

You can actually create multiple columns in a single function call.

```
# Create a column for height in metres  
df <- mutate(starwars, height_m = height/100)
```

Group and summarise your data

Another very common task is to group your data by a column (or more than one column) and then create summarised values for the grouped data. The functions **group_by()** and **summarise()** in the **{dplyr}** package make it very easy to do these transformations.

```
# Find the min/mean/max mass value for each species category
df <- summarise(group_by(starwars, species),
  min_mass = min(mass, na.rm = TRUE),
  mean_mass = mean(mass, na.rm = TRUE),
  max_mass = max(mass, na.rm = TRUE))
```

Live Coding Example 2



Use the `starwars` dataset to:

1. Remove the columns “films”, “vehicles”, “starships” from the data. 💡
2. Remove rows that have missing mass values.
3. Calculate the Body Mass Index (BMI) for each character*.
4. Arrange the rows by descending BMI ... who do you think is at the top?
5. Find the median BMI value for each gender category.

*BMI = weight (kg) / height² (m)



Use minus sign “-” to remove columns

Live Coding Example 2

Who is the character
with the highest BMI?



	name	height	mass	hair_color	skin_color	eye_color	birth_year	gender	homeworld	species	height_m	BMI
1	Jabba Desilijic Tiure	175	1358.0	NA	green-tan, brown	orange	600.0	hermaphrodite	Nal Hutta	Hutt	1.75	443.42857
2	Dud Bolt	94	45.0	none	blue, grey	yellow	NA	male	Vulpter	Vulptereen	0.94	50.92802
3	Yoda	66	17.0	white	green	brown	896.0	male	NA	Yoda's species	0.66	39.02663
4	Owen Lars	178	120.0	brown, grey	light	blue	52.0	male	Tatooine	Human	1.78	37.87401
5	IG-88	200	140.0	none	metal	red	15.0	none	NA	Droid	2.00	35.00000
6	R2-D2	96	32.0	NA	white, blue	red	33.0	NA	Naboo	Droid	0.96	34.72222



	gender	median_BMI
1	female	18.06751
2	hermaphrodite	443.42857
3	male	24.70827
4	none	35.00000
5	NA	34.00999

Investigate data quality

Live Coding Example 2

```
# Select columns
df <- select(starwars, -films, -vehicles, -starships)

# Filter rows that have missing mass
df <- filter(df, !is.na(mass))

# Create columns: height in metres and the Body Mass Index (BMI)
df <- mutate(df, height_m = height/100, BMI = mass / (height_m)^2)

# Arrange rows according to descending "BMI" values
df <- arrange(df, desc(BMI))

# Calculate the median BMI value for each gender
df <- summarise(group_by(df, gender), median_BMI = median(BMI))
```

Summary of {dplyr} “verb” functions

Function	Description
select	Select columns by name
filter	Filter rows that meet a condition
arrange	Arrange rows to some order
mutate	Mutate data to create new columns
group_by	Group data by columns
summarise	Summarise data to values

Improve your workflow with



A package that has changed the way we write R code is called **{magrittr}**. It has significantly improved the readability and workflow of code by introducing the “pipe” operator. It acts as a “then” operation where we can pass data from one function to another function very easily.



Fun fact: The package name is inspired by the famous artist René Magritte.

One of his work, a pipe, has the text “this is not a pipe” as a caption ... this is where the **{magrittr}** package gets its image.

Live Coding Example 3



Repeat Example 1 using the pipe operator from the **{magrittr}** package.

1. Select the columns: "name", "height", "mass", "species" THEN filter the rows to keep only those characters that are greater than or equal to 175cm THEN filter the rows to keep only the human characters THEN arrange the rows according to descending "mass" values.

Live Coding Example 3

```
library(magrittr)

# Pipe each data manipulation operation to the next one
df <- starwars %>%
  select(name, height, mass, species) %>%
  filter(height >= 175) %>%
  filter(species == "Human") %>%
  arrange(desc(mass))
```



Try **CTRL+SHIFT+M** (Windows) **CMD+SHIFT+M** (Mac)
and see what happens

Live Coding Example 4



Repeat Example 2 using the pipe operator from the **{magrittr}** package.

1. Remove the columns “films”, “vehicles”, “starships” from the data **THEN** remove rows that have missing mass values **THEN** calculate the Body Mass Index (BMI) for each character **THEN** arrange the rows by descending BMI **THEN** find the median BMI value for each gender category.

Live Coding Example 4

```
# Pipe each data manipulation operation to the next one
df <- starwars %>%
  select(-films, -vehicles, -starships) %>%
  filter(!is.na(mass)) %>%
  mutate(height_m = height/100,
          BMI = mass / (height_m)^2) %>%
  arrange(desc(BMI)) %>%
  group_by(gender) %>%
  summarise(median_BMI = median(BMI))
```

Simple string manipulation using



The package in the tidyverse collection that helps us do data manipulations involving strings is called **{stringr}**. String manipulation is another common task, especially in data cleaning and pre-processing. Here are some examples:

```
# Make all character names as lower case
string <- str_to_lower(starwars$name)

# Combine the name, hair colour & eye colour of characters in a sentence
string <- str_c(starwars$name, " has ",
               starwars$hair_color, " hair and ",
               starwars$eye_color, " eyes.")

# Create an indicator where the specific pattern matches
ind <- str_detect(string = starwars$name, pattern = "Skywalker")
```

Live Coding Example 5



Use the `starwars` dataset to:

1. Transform the character names to upper case.
2. Combine the “name” and the “homeworld” to create a sentence, for example: “Luke Skywalker is from Tatooine”.
3. Create an indicator for the rows where characters have green skin.

Live Coding Example 5

```
# Make all character names as upper case
string <- str_to_upper(starwars$name)

# Combine the name, hair colour & eye colour of characters in a sentence
string <- str_c(starwars$name, " is from ",
                starwars$homeworld, ".")

# Create an indicator where the specific pattern matches
ind <- str_detect(string = starwars$skin_color, pattern = "green")
```


Other resources – {dplyr} cheat sheet

Data Transformation with dplyr : CHEAT SHEET

dplyr functions work with pipes and expect tidy data. In tidy data:

- Each **variable** is in its own **column**
- Each **observation**, or **case**, is in its own **row**
- $x \%>\% f(y)$ becomes $f(x, y)$

Summary Cases

These apply **summary functions** to columns to create a new table of summary statistics. Summary functions take vectors as input and return one value (see back).

summary function

```
summarise(data, ...)
# Compute table of summaries
summarise(mtcars, avg = mean(mpg))
```

count() ... wt = NULL, sort = FALSE
Count number of rows in each group defined by the variables in ... Also tally().
count(mtcars, Species)

VARIATIONS

- summarise_all()** - Apply funs to every column.
- summarise_at()** - Apply funs to specific columns.
- summarise_if()** - Apply funs to all cols of one type.

Group Cases

Use **group_by()** to create a "grouped" copy of a table. dplyr functions will manipulate each "group" separately and then combine the results.

```
mtcars %>%
  group_by(cyl) %>%
  summarise(avg = mean(mpg))
```

group_by(data, ... add = FALSE)
Returns copy of table grouped by ...
g_mtcars = group_by(mtcars, Species)

ungroup(x, ...)
Returns ungrouped copy of table.
ungroup(g_mtcars)

Logical and boolean operators to use with filter()

```
< <= to.na() %in% xor()
> >= !is.na() | &
```

See ?base::Logic and ?Comparison for help.

MANIPULATE CASES

Row functions return a subset of rows as a new table.

filter(data, ...) Extract rows that meet logical criteria. filter(mtcars, Sepsol.Length > 7)

distinct(data, ..., keep_all = FALSE) Remove rows with duplicate values.
distinct(mtcars, Species)

sample_frac(tbl, size = 1, replace = FALSE, weight = NULL, env = parent.frame()) Randomly select fraction of rows.
sample_frac(mtcars, 0.5, replace = TRUE)

sample_n(tbl, size, replace = FALSE, weight = NULL, env = parent.frame()) Randomly select size rows. sample_n(mtcars, 10, replace = TRUE)

slice(data, ...) Select rows by position. slice(mtcars, 10:15)

top_n(x, n, wt) Select and order top n entries (by group if grouped data). top_n(mtcars, Sepsol.Length)

MANIPULATE VARIABLES

Column functions return a set of columns as a new vector or table.

pull(data, var = 1) Extract column values as a vector. Choose by name or index.
pull(mtcars, Sepsol.Length)

select(data, ...) Extract columns as a table. Also **select_if()**.
select(mtcars, Sepsol.Length, Species)

Use these helpers with select (),
e.g. **select_if()**, **starts_with()** ("Sepsol")

contains(match) num_range(prefix, range) 1, e.g. mpg:cyl
ends_with(match) one_of(...) -, e.g. Species
matches(match) starts_with(match)

MAKE NEW VARIABLES

These apply **vectorized functions** to columns. Vectorized funs take vectors as input and return vectors of the same length as output (see back).

vectorized function

```
mutate(data, ...)
# Compute new column(s)
mutate(mtcars, gpm = 1/mpg)
```

transmute(data, ...)
Compute new column(s), drop others.
transmute(mtcars, gpm = 1/mpg)

mutate_all(tbl, funs, ...) Apply funs to every column. Use with **funs()**. Also **mutate_if()**.
mutate_all(mtcars, funs(log10))

mutate_if(mtcars, is.numeric, funs(log10))

mutate_at(tbl, cols, funs, ...) Apply funs to specific columns. Use with **funs()**, **vars()** and the helper functions for select().
mutate_at(mtcars, vars(Species), funs(log10))

add_column(data, ..., before = NULL, after = NULL) Add new column(s). Also **add_count()**.
add_count(mtcars, new = 1:2)

add_row(tbl, ..., before = NULL, after = NULL) Add one or more rows to a table.
add_row(mtcars, eruptions = 1, waiting = 1)

rename(data, ...) Rename columns.
rename(mtcars, Length = Sepsol.Length)

Vector Functions

TO USE WITH **Mutate ()**

mutate() and **transmute()** apply vectorized functions to columns to create new columns. Vectorized functions take vectors as input and return vectors of the same length as output.

vectorized function

OFFSETS

```
dplyr::lag() - Offset elements by 1
dplyr::lead() - Offset elements by -1
```

CUMULATIVE AGGREGATES

```
dplyr::cumall() - Cumulative all()
dplyr::cumany() - Cumulative any()
dplyr::cummax() - Cumulative max()
dplyr::cummean() - Cumulative mean()
dplyr::cummin() - Cumulative min()
dplyr::cumprod() - Cumulative prod()
dplyr::cumsum() - Cumulative sum()
```

RANKINGS

```
dplyr::cume_dist() - Proportion of all values <=
dplyr::dense_rank() - rank w ties = min, no gaps
dplyr::min_rank() - rank with ties = min
dplyr::ntile() - bins into n bins
dplyr::percent_rank() - min_rank scaled to [0,1]
dplyr::row_number() - rank with ties = "first"
```

MATH

```
+, -, *, /, ^, %%, %%, % - arithmetic ops
log(), log10(), log2() - logs
<, <=, >, >=, == - logical comparisons
dplyr::between() - x >= left & x <= right
dplyr::mean() - safe == for floating point numbers
```

MISC

```
dplyr::case_when() - multi-case if_else()
g_mtcars = mutate(mtcars, color = case_when(
  Species == "versicolor" ~ "yellow",
  Species == "virginica" ~ "pink",
  TRUE ~ "Species"))
```

dplyr::coalesce() - first non-NA values by element across a set of vectors
dplyr::if_else() - element-wise if() & else()
dplyr::na_if() - replace specific values with NA
pmax() - element-wise max()
pmin() - element-wise min()
dplyr::recode() - Vectorized switch()
dplyr::recode_factor() - Vectorized switch() for factors

Also has: rownames(), remove_rownames()

Summary Functions

TO USE WITH **SUMMARISE ()**

summarise() applies summary functions to columns to create a new table. Summary functions take vectors as input and return single values as output.

summary function

COUNTS

```
dplyr::n() - number of values/rows
dplyr::n_distinct() - # of uniques
sum(!is.na()) - # of non-NA's
```

LOCATION

```
mean() - mean, also mean(!is.na())
median() - median
```

LOGICALS

```
mean() - Proportion of TRUE's
sum() - # of TRUE's
```

POSITION/ORDER

```
dplyr::first() - first value
dplyr::last() - last value
dplyr::nth() - value in nth location of vector
```

RANK

```
quantile() - nth quantile
min() - minimum value
max() - maximum value
```

SPREAD

```
spread() - Inter-Quartile Range
mad() - median absolute deviation
sd() - standard deviation
var() - variance
```

Combine Tables

COMBINE VARIABLES

Use **bind_cols()** to paste tables beside each other as they are.

```
bind_cols(x, y)
# Returns tables placed side by side as a single table. BE SURE THAT ROWS ALIGN.
```

Use a **"Mutating Join"** to join one table to columns from another, matching values with the rows that they correspond to. Each join retains a different combination of values from the tables.

```
left_join(x, y, by = NULL, copy = FALSE, suffix = c("x", "y"), ...)
# Join matching values from x to y.
```

```
right_join(x, y, by = NULL, copy = FALSE, suffix = c("x", "y"), ...)
# Join matching values from y to x.
```

```
inner_join(x, y, by = NULL, copy = FALSE, suffix = c("x", "y"), ...)
# Join data. Retain only rows with matches.
```

```
full_join(x, y, by = NULL, copy = FALSE, suffix = c("x", "y"), ...)
# Join data. Retain all values, all rows.
```

COMBINE CASES

Use **bind_rows()** to paste tables below each other as they are.

```
bind_rows(x, y, ...)
# Returns tables one on top of the other as a single table. Set id to a column name to add a column of the original table names (as pictured)
```

intersect(x, y, ...)
Rows that appear in both x and y.

setdiff(x, y, ...)
Rows that appear in x but not y.

union(x, y, ...)
Rows that appear in x or y. (Duplicates removed). union_all() retains duplicates.

Use **setequal()** to test whether two data sets contain the exact same rows (in any order).

EXTRACT ROWS

```
x
y
# x %>% filter(y) ==
```

Use a **"Filtering Join"** to filter one table against the rows of another.

```
semi_join(x, y, by = NULL, ...)
# Return rows of x that have a match in y. USEFUL TO SEE WHAT WILL BE JOINED.
```

```
anti_join(x, y, by = NULL, ...)
# Return rows of x that do not have a match in y. USEFUL TO SEE WHAT WILL NOT BE JOINED.
```



Get the cheat sheet at: <https://rstudio.com/resources/cheatsheets/>

Other resources – {stringr} cheat sheet

String manipulation with stringr : : CHEAT SHEET

The **stringr** package provides a set of internally consistent tools for working with character strings, i.e. sequences of characters surrounded by quotation marks.

Detect Matches

- str_detect(string, pattern)** Detect the presence of a pattern match in a string. `str_detect(fruit, "a")`
- str_which(string, pattern)** Find the indexes of strings that contain a pattern match. `str_which(fruit, "a")`
- str_count(string, pattern)** Count the number of matches in a string. Also `str_count_all(fruit, "a")`
- str_locate(string, pattern)** Locate the positions of pattern matches in a string. Also `str_locate_all(fruit, "a")`

Subset Strings

- str_sub(string, start = 1L, end = -1L)** Extract substrings from a character vector. `str_sub(fruit, 1, 3); str_sub(fruit, -3)`
- str_subset(string, pattern)** Return only the strings that contain a pattern match. `str_subset(fruit, "b")`
- str_extract(string, pattern)** Return the first pattern match found in each string, as a vector. Also `str_extract_all(fruit, "oaiou")`
- str_match(string, pattern)** Return the first pattern match found in each string, as a matrix with a column for each (1) group in pattern. Also `str_match(sentences, "(a|the) (l|t) ")`

Manage Lengths

- str_length(string)** The width of strings (i.e. number of code points, which generally equals the number of characters). `str_length(fruit)`
- str_pad(string, width, side = c("left", "right", "both"), pad = " ")** Pad strings to constant width. `str_pad(fruit, 17)`
- str_truncate(string, width, side = c("right", "left", "center"), ellipsis = "...")** Truncate the width of strings, replacing content with ellipsis. `str_truncate(fruit, 3)`
- str_trim(string, side = c("both", "left", "right"))** Trim whitespace from the start and/or end of a string. `str_trim(fruit)`

Mutate Strings

- str_sub<v>(string, value)** Replace substrings by identifying the substrings with `str_sub()` and assigning into the results. `str_sub(fruit, 1, 3) <- "a"`
- str_replace(string, pattern, replacement)** Replace the first matched pattern in each string. `str_replace(fruit, "a", "-")`
- str_replace_all(string, pattern, replacement)** Replace all matched patterns in each string. `str_replace_all(fruit, "a", "-")`
- str_to_lower(string, locale = "en")** Convert strings to lower case. `str_to_lower(sentences)`
- str_to_upper(string, locale = "en")** Convert strings to upper case. `str_to_upper(sentences)`
- str_to_title(string, locale = "en")** Convert strings to title case. `str_to_title(sentences)`

Join and Split

- str_c<v>(string, sep = "", collapse = NULL)** Join multiple strings into a single string. `str_c(letters, LETTERS)`
- str_collapse<v>(string, sep = "", collapse = "")** Collapse a vector of strings into a single string. `str_collapse(LETTERS)`
- str_dup(string, times)** Repeat strings times times. `str_dup(fruit, times = 2)`
- str_split_fixed(string, pattern, n)** Split a vector of strings into a matrix of substrings (splitting at occurrences of a pattern match). Also `str_split` to return a list of substrings. `str_split(fruit, " ", n = 2)`
- str_glue<v>(string, sep = "", env = parent.frame())** Create a string from strings and (expressions) to evaluate. `str_glue("PI is {pi}")`
- str_glue_data<v>(string, sep = "", env = parent.frame(), na = "NA")** Use a data frame, list, or environment to create a string from strings and (expressions) to evaluate. `str_glue_data(mtcars, "rownames(mtcars) has {hp} hp")`

Order Strings

- str_order(x, decreasing = FALSE, na_last = TRUE, locale = "en", numeric = FALSE, ...)** Return the vector of indexes that sorts a character vector. `str_order(x)`
- str_sort(x, decreasing = FALSE, na_last = TRUE, locale = "en", numeric = FALSE, ...)** Sort a character vector. `str_sort(x)`

Helpers

- str_conv(string, encoding)** Override the encoding of a string. `str_conv(fruit, "ISO-8859-1")`
- str_view(string, pattern, match = NA)** View HTML rendering of first regex match in each string. `str_view(fruit, "[aeiou]")`
- str_view_all(string, pattern, match = NA)** View HTML rendering of all regex matches. `str_view_all(fruit, "[aeiou]")`
- str_wrap(string, width = 80, indent = 0, exdent = 0)** Wrap strings into nicely formatted paragraphs. `str_wrap(sentences, 20)`

Need to Know

Pattern arguments in **stringr** are interpreted as regular expressions (i.e. sequences of characters surrounded by quotes ("") or single quotes ('')).

In R, you write regular expressions as strings, sequences of characters surrounded by quotes ("") or single quotes ('').

Some characters cannot be represented directly in an R string. These must be represented as special characters, sequences of characters that have a specific meaning, e.g.

Special Character	Represents
\	backslash
\\	double backslash
\n	new line (return)
\t	tab
\s	any whitespace (S for non-whitespaces)
\d	any digit (D for non-digits)
\w	any word character (W for non-word chars)
\b	word boundaries
[]	characters in brackets
{ }	quantifiers
	alternation
()	grouping
^	start of string
\$	end of string

Run `??` to see a complete list.

Because of this, whenever a \ appears in a regular expression, you must write it as \\ in the string that represents the regular expression.

Use `writeLines()` to see how R views your string after all special characters have been parsed.

```
writeLines("a\\b")
# [1] "a\b"

writeLines("a\\b\\n")
# [1] "a\b\n"
```

Regular Expressions -

Regular expressions, or regexps, are a concise language for describing patterns in strings.

```
see <- function(x) str_view_all("abcABC123456789", x)
```

MATCH CHARACTERS	regex	matches	example
string (type: chr)	regex (a etc.)	a (etc.)	abc ABC 123 .7890
\\	backslash	backslash	abc ABC 123 .7890
\\n	new line (return)	new line	abc ABC 123 .7890
\\t	tab	tab	abc ABC 123 .7890
\\s	any whitespace (S for non-whitespaces)	any digit (D for non-digits)	abc ABC 123 .7890
\\d	any digit (D for non-digits)	any word character (W for non-word chars)	abc ABC 123 .7890
\\w	any word character (W for non-word chars)	word boundaries	abc ABC 123 .7890
\\b	word boundaries	digits	abc ABC 123 .7890
[]	characters in brackets	lowercase letters	abc ABC 123 .7890
{ }	quantifiers	uppercase letters	abc ABC 123 .7890
	alternation	letters and numbers	abc ABC 123 .7890
()	grouping	punctuation	abc ABC 123 .7890
^	start of string	letters, numbers, and punctuation	abc ABC 123 .7890
\$	end of string	space characters (i.e. \s)	abc ABC 123 .7890
.	any character except a new line	space and tab (but not new line)	abc ABC 123 .7890
.	any character except a new line	every character except a new line	abc ABC 123 .7890

Many base R functions require classes to be wrapped in a second set of [], e.g. `[digit]`.

ALTERNATES

regex	matches	example
[a b]	a or b	abcde
[a b c]	a or b or c	abcde
[a b c d]	a or b or c or d	abcde

ANCHORS

regex	matches	example
^	start of string	aaa
\$	end of string	aaa

LOOK AROUND

regex	matches	example
(?<=)	looked at	ba cad
(?<=)	not followed by	ba cad
(?<=)	preceded by	ba cad
(?<=)	not preceded by	ba cad

QUANTIFIERS

regex	matches	example
?	zero or one	quant("a")
*	zero or more	quant("a")
+	one or more	quant("a")
{n}	exactly n	quant("a", 2)
{n, m}	between n and m	quant("a", 2, 4)

GROUPS

regex	matches	example
(a b c)	a or b or c	abcde
(a b c)	a or b or c	abcde

Use parentheses to set precedence (order of evaluation) and create groups.

Use an escaped number to refer to and duplicate parentheses groups that occur earlier in a pattern. Refer to each group by its order of appearance.

string	regex	matches	example
ba cad	(?<=)	ba cad	ba cad
ba cad	(?<=)	ba cad	ba cad
ba cad	(?<=)	ba cad	ba cad



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