



JOINING DATA IN R WITH DATA.TABLE

Complex keys

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Misspecified joins

What happens when you don't use the correct columns for join keys?

- An error is thrown
- The result is a malformed `data.table`

Column type mismatch

Using join key columns with different types will error

```
customers[web_visits, on = .(age = name)]
```

```
Error in bmerge(i, x, leftcols, rightcols, io, xo, roll, rollends,
nomatch,  :
  typeof x.age (double) != typeof i.name (character)
```

customers:

name	gender	age	address
Madeline Martin	F	54	5 Market lane
Madeline Bernard	F	45	4 Jacaranda crescent
George Dimakos	M	39	2a Park square

+

web_visits:

name	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45

Column type mismatch

```
customers[web_visits, on = .(id)]
```

```
Error in bmerge(i, x, leftcols, rightcols, io, xo, roll, rollends,
nomatch,  :
  typeof x.id (integer) != typeof i.id(character)
```

customers:

id	name	gender	age	address
1	"Madeline Martin"	"F"	54	"5 Market lane"
2	"Madeline Bernard"	"F"	45	"4 Jacaranda crescent"
3	"George Dimakos"	"M"	39	"2a Park square"

+

web_visits:

id	name	date	duration
"1"	"Madeline Martin"	2018-05-02	5
"1"	"Madeline Martin"	2018-05-03	32
"2"	"Madeline Bernard"	2018-05-03	12
"3"	"George Dimakos"	2018-04-27	45

Malformed full joins - no common key values

```
merge(customers, web_visits, by.x = "address", by.y = "name", all = TRUE)
```

customers:

name	gender	age	address
Madeline Martin	F	54	5 Market lane
Madeline Bernard	F	45	4 Jacaranda crescent
George Dimakos	M	39	2a Park square

+

web_visits:

name	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45

=

address	name	gender	age	date	duration
2a Park square	George Dimakos	M	39	NA	NA
4 Jacaranda crescent	Madeline Bernard	F	45	NA	NA
5 Market lane	Madeline Martin	F	54	NA	NA
George Dimakos	NA	NA	NA	2018-04-27	45
Madeline Bernard	NA	NA	NA	2018-05-03	12
Madeline Martin	NA	NA	NA	2018-05-02	5
Madeline Martin	NA	NA	NA	2018-05-03	32

Malformed right and left joins - no common key values

```
customers[web_visits, on = .(address = name)]
```

customers:

name	gender	age	address
Madeline Martin	F	54	5 Market lane
Madeline Bernard	F	45	4 Jacaranda crescent
George Dimakos	M	39	2a Park square

+

web_visits:

name	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45

=

name	gender	age	address	date	duration
NA	NA	NA	Madeline Martin	2018-05-02	5
NA	NA	NA	Madeline Martin	2018-05-03	32
NA	NA	NA	Madeline Bernard	2018-05-03	12
NA	NA	NA	George Dimakos	2018-04-27	45

Malformed inner joins - no common key values

```
customers[web_visits, on = .(address = name), nomatch = 0]
```

customers:

name	gender	age	address
Madeline Martin	F	54	5 Market lane
Madeline Bernard	F	45	4 Jacaranda crescent
George Dimakos	M	39	2a Park square

+

web_visits:

name	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45

=

name	gender	age	address	date	duration
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Malformed joins - coincidental common key values

```
customers[web_visits, on = .(age = duration), nomatch = 0]
```

customers:

name	gender	age	address
Madeline Martin	F	54	5 Market lane
Madeline Bernard	F	45	4 Jacaranda crescent
George Dimakos	M	39	2a Park square

+

web_visits:

name	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45

=

name	gender	age	address	i.name	date
Madeline Bernard	F	45	4 Jacaranda crescent	George Dimakos	2018-04-27



Avoiding misspecified joins

Learning what each column represents before joins will help you avoid errors

Keys with different column names

customers:

name	gender	age	address
Madeline Martin	F	54	5 Market lane
Madeline Bernard	F	45	4 Jacaranda crescent
George Dimakos	M	39	2a Park square

web_visits:

person	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45

```
merge(customers, web_visits, by.x = "name", by.y = "person")
```

```
customers[web_visits, on = .(name = person)]
```

```
customers[web_visits, on = c("name" = "person")]
```

```
key <- c("name" = "person")  
customers[web_visits, on = key]
```



Multi-column keys

customers:

first	last	gender	age	address
Madeline	Martin	F	54	5 Market lane
Madeline	Bernard	F	45	4 Jacaranda crescent
George	Dimakos	M	39	2a Park square

web_visits:

first	last	date	duration
Madeline	Martin	2018-05-02	5
Madeline	Martin	2018-05-03	32
Madeline	Bernard	2018-05-03	12
George	Dimakos	2018-04-27	45



Multi-column keys

purchases:

name	date	item	units	price
Madeline Martin	2018-05-03	book	2	\$15.00
Arthur Smith	2018-05-03	shelf	1	\$30.00
Jaqueline Mary	2018-05-03	CD	1	\$12.00
George Dimakos	2018-05-03	plant	3	\$16.00
George Dimakos	2018-04-27	shelf	1	\$30.00

web_visits:

name	date	duration
Madeline Martin	2018-05-02	5
Madeline Martin	2018-05-03	32
Madeline Bernard	2018-05-03	12
George Dimakos	2018-04-27	45



Specifying multiple keys with merge()

```
merge(purchases, web_visits, by = c("name", "date"))
```

```
merge(purchases, web_visits,  
      by.x = c("name", "date"),  
      by.y = c("person", "date"))
```



Specifying multiple keys with the data.table syntax

```
purchases[web_visits, on = .(name, date)]  
purchases[web_visits, on = c("name", "date")]
```

```
purchases[web_visits, on = .(name = person, date)]  
purchases[web_visits, on = c("name" = "person", "date")]
```



JOINING DATA IN R WITH DATA.TABLE

Final Slide



JOINING DATA IN R WITH DATA.TABLE

Problem columns

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Common column names

parents:

name	gender	age
Sarah	F	41
Max	M	43
Qin	F	36

children:

parent	name	gender	age
Sarah	Oliver	M	5
Max	Sebastian	M	8
Qin	Kai-lee	F	7



Common column names

Using the `data.table` syntax

```
parents[children, on = .(name = parent)]
```

	name	gender	age	i.name	i.gender	i.age
1:	Sarah	F	41	Oliver	M	5
2:	Max	M	43	Sebastian	M	8
3:	Qin	F	36	Kai-lee	F	7

Common column names with merge()

Using the `merge()` function

```
merge(x = children, y = parents, by.x = "parent", by.y = "name")
```

	parent	name	gender.x	age.x	gender.y	age.y
1:	Max	Sebastian	M	8	M	43
2:	Qin	Kai-lee	F	7	F	36
3:	Sarah	Oliver	M	5	F	41

Adding context with your own suffixes

The `suffixes` argument can add useful context:

```
merge(children, parents, by.x = "parent", by.y = "name",  
      suffixes = c(".child", ".parent"))
```

	parent	name	gender.child	age.child	gender.parent	age.parent
1:	Max	Sebastian	M	8	M	43
2:	Qin	Kai-lee	F	7	F	36
3:	Sarah	Oliver	M	5	F	41



Renaming columns

Rename all columns using `setnames()`

```
setnames(parents, c("parent", "parent.gender", "parent.age"))
```

```
setnames(parents, old = c("gender", "age"),  
          new = c("parent.gender", "parent.age"))
```

```
parents
```

	parent	parent.gender	parent.age
1:	Sarah	F	41
2:	Max	M	43
3:	Qin	F	36



Joining with data.frames

Join keys for `data.frames` may be in the rownames

```
# A data.frame
parents
```

	gender	age
Sarah	F	41
Max	M	43
Qin	F	36

```
parents <- as.data.table(parents, keep.rownames = "parent")
parents
```

	parent	gender	age
1:	Sarah	F	41
2:	Max	M	43
3:	Qin	F	36



JOINING DATA IN R WITH DATA.TABLE

Let's practice!



JOINING DATA IN R WITH DATA.TABLE

Duplicate matches

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Join key duplicates

```
# Which bacteria could be found at both sites using any method?  
site1_ecology[site2_ecology, on = .(genus)]
```

site1_ecology:

genus	count	method
<i>Nitrosomonas</i>	500	WGS
<i>Nitrosomonas</i>	620	16S
<i>Rhizobium</i>	360	WGS
<i>Rhizobium</i>	300	16S

site2_ecology:

genus	present	method
<i>Nitrosomonas</i>	TRUE	WGS
<i>Nitrosomonas</i>	TRUE	16S
<i>Nitrosomonas</i>	TRUE	Culture
<i>Rhizobium</i>	TRUE	WGS
<i>Rhizobium</i>	TRUE	16S
<i>Rhizobium</i>	FALSE	Culture



Error from multiplicative matches

```
site1_ecology[site2_ecology, on = .(genus)]
```

```
Error in vecseq(f__, len__, if (allow.cartesian || notjoin ||
```

```
!anyDuplicated(f__, :
```

Join results in 12 rows; more than 10 = nrow(x)+nrow(i). Check for duplicate key values in i each of which join to the same group in x over and over again. If that's ok, try by=.EACHI to run j for each group to avoid the large allocation. If you are sure you wish to proceed, rerun with allow.cartesian=TRUE. Otherwise, please search for this error message in the FAQ, Wiki, Stack Overflow and data.table issue tracker for advice.



Allowing multiplicative matches

`allow.cartesian = TRUE` allows the join to proceed:

```
# data.table syntax  
site1_ecology[site2_ecology, on = .(genus), allow.cartesian = TRUE]
```

```
# merge()  
merge(site1_ecology, site2_ecology, by = "genus", allow.cartesian = TRUE)
```

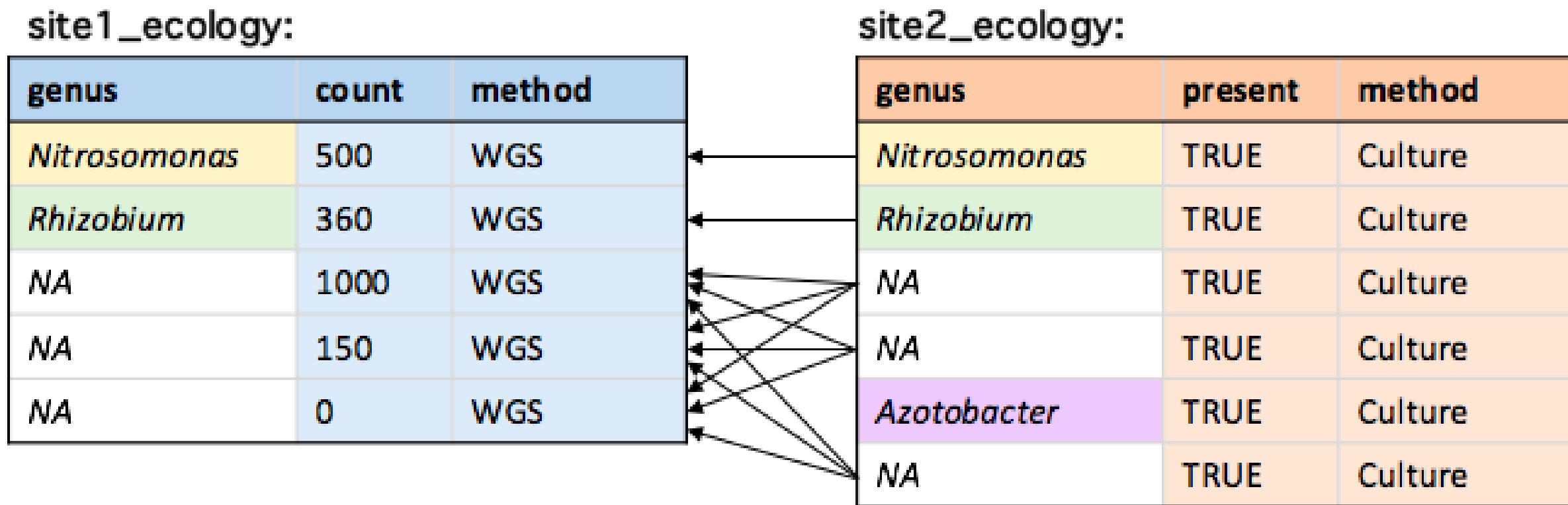
Allowing multiplicative matches

```
site1_ecology[site2_ecology, on = .(genus), allow.cartesian = TRUE]
```

	genus	count	method	present	i.method
1:	Nitrosomonas	500	WGS	TRUE	WGS
2:	Nitrosomonas	620	16S	TRUE	WGS
3:	Nitrosomonas	500	WGS	TRUE	16S
4:	Nitrosomonas	620	16S	TRUE	16S
5:	Nitrosomonas	500	WGS	TRUE	Culture
6:	Nitrosomonas	620	16S	TRUE	Culture
7:	Rhizobium	360	WGS	TRUE	WGS
8:	Rhizobium	300	16S	TRUE	WGS
9:	Rhizobium	360	WGS	TRUE	16S
10:	Rhizobium	300	16S	TRUE	16S
11:	Rhizobium	360	WGS	FALSE	Culture
12:	Rhizobium	300	16S	FALSE	Culture

Missing values

Missing values (`NA`) will match all other missing values:



Filtering missing values

`!is.na()` can be used to filter rows with missing values

```
site1_ecology <- site1_ecology[!is.na(genus)]  
site1_ecology
```

	genus	count	method
1:	Nitrosomonas	500	WGS
2:	Rhizobium	360	WGS

```
site2_ecology <- site2_ecology[!is.na(genus)]  
site2_ecology
```

	genus	present	method
1:	Nitrosomonas	TRUE	Culture
2:	Rhizobium	TRUE	Culture
3:	Azotobacter	TRUE	Culture



Keeping only the first match

```
site1_ecology[site2_ecology, on = .(genus), mult = "first"]
```

site1_ecology:

genus	Year	count	method
Nitrosomonas	2018	620	16S
Nitrosomonas	2017	603	16S
Nitrosomonas	2016	591	16S
Rhizobium	2018	290	16S
Rhizobium	2017	300	16S
Rhizobium	2016	280	16S
Azotobacter	2018	1230	16S
Azotobacter	2017	0	16S
Azotobacter	2016	0	16S

site2_ecology:

genus	present	method
Nitrosomonas	TRUE	WGS
Rhizobium	TRUE	WGS
Azotobacter	FALSE	WGS



Keeping only the last match

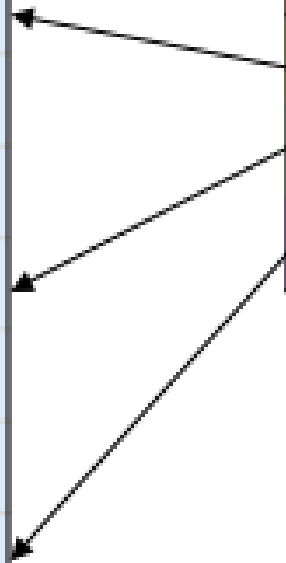
```
children[parents, on = .(parent = name), mult = "last"]
```

site1_ecology:

genus	Year	count	method
<i>Nitrosomonas</i>	2018	620	16S
<i>Nitrosomonas</i>	2017	603	16S
<i>Nitrosomonas</i>	2016	591	16S
<i>Rhizobium</i>	2018	290	16S
<i>Rhizobium</i>	2017	300	16S
<i>Rhizobium</i>	2016	280	16S
<i>Azotobacter</i>	2018	1230	16S
<i>Azotobacter</i>	2017	0	16S
<i>Azotobacter</i>	2016	0	16S

site2_ecology:

genus	present	method
<i>Nitrosomonas</i>	TRUE	WGS
<i>Rhizobium</i>	TRUE	WGS
<i>Azotobacter</i>	FALSE	WGS





Identifying and removing duplicates

`duplicated()` : what rows are duplicates?

`unique()` : **filter** a `data.table` to just unique rows

The duplicated() function

Using values in all columns:

```
duplicated(site1_ecology)
[1] FALSE FALSE FALSE FALSE
```

Using values in a subset of columns:

```
duplicated(site1_ecology,
           by = "genus")
[1] FALSE TRUE FALSE TRUE
```

site1_ecology:

genus	count	method
<i>Nitrosomonas</i>	500	WGS
<i>Nitrosomonas</i>	620	16S
<i>Rhizobium</i>	360	WGS
<i>Rhizobium</i>	300	16S

The unique() function

```
unique(site1_ecology, by = "genus")
```

site1_ecology:

genus	count	method
<i>Nitrosomonas</i>	500	WGS
<i>Nitrosomonas</i>	620	16S
<i>Rhizobium</i>	360	WGS
<i>Rhizobium</i>	300	16S

X

X

genus	count	method
<i>Nitrosomonas</i>	500	WGS
<i>Rhizobium</i>	360	WGS

Changing the search order

`fromLast = TRUE` changes the direction of the search to start from the last row

```
 duplicated(site1_ecology, by = "genus", fromLast = TRUE)
[1] TRUE FALSE TRUE FALSE
 unique(site1_ecology, by = "genus", fromLast = TRUE)
```

site1_ecology:

genus	count	method		genus	count	method
Nitrosomonas	500	WGS	X	Nitrosomonas	620	16S
Nitrosomonas	620	16S		Rhizobium	300	16S
Rhizobium	360	WGS	X			
Rhizobium	300	16S				



JOINING DATA IN R WITH DATA.TABLE

Let's practice!