



ETC4500/ETC5450 Advanced R programming

Week 8: Object-oriented Programming (S3)



- 1 Object oriented programming
- 2 S3
- 3 S3 vctrs
- 4 S4
- 5 R6

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Object oriented programming

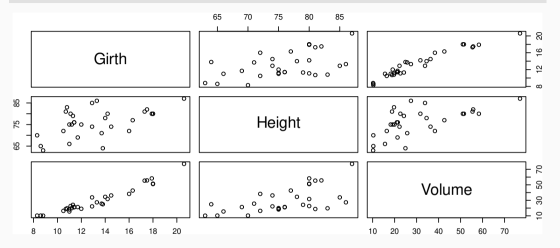
- Encapsulation: objects have secret internals that you don't need to understand
- **Polymorphism**: the same function can do different things to different data as appropriate
- Inheritance: you can take an existing kind of object and make a new, more specialised one

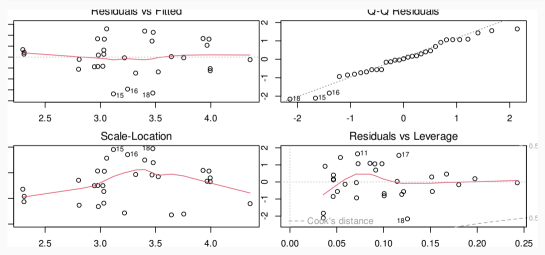
Inheritance turns out to be useful mostly for data infrastructure, but encapsulation and polymorphism are generally valuable

Generic functions and methods

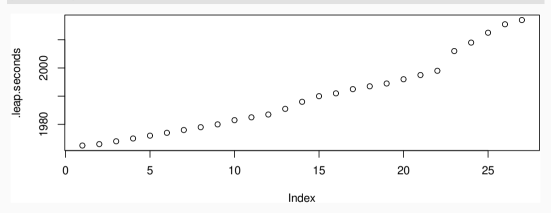
A simple example: plot

plot(trees)

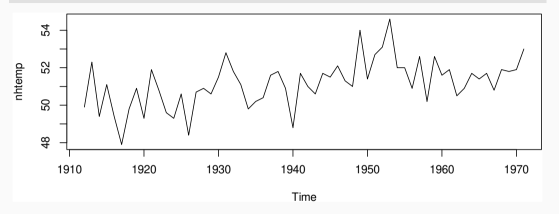








plot(nhtemp)



How?

- Giant switch statement...
- ...that gets updated every time you load a package...
- **???**

Generic functions and methods!

Object systems

R has **a lot** of object systems

- S3
- S3 vctrs
- **S**4
- R6
- R.oo, proto, R7

Main topic for today

- easy to start writing
- no safeguards
- especially good for single-person, small to medium projects
- can be used for large projects with a lot of attention to documentation and communication
- limited use of inheritance
- basis of tidyverse and most of CRAN

S3 vctrs

A tidyverse package for making different sorts of vectors

- handles a lot of formatting and subsetting details
- allows for binary operators
- useful if you want your vectors in a tibble
- enforces some safeguards

S4

- more work to start writing
- objects know their structure
- enforces object structure
- better for large-scale collaborative programming
- better at inheritance
- basis of Bioconductor

R6

- Supports modifiable (mutable) objects
 - database connections, files, etc
 - large data objects we don't want to copy
 - interfacing to eg tensorflow
 - shared state between copies of an object
- not widely used otherwise
- more similar to other languages

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Back to plot

- plot() doesn't do anything
- All the work is done by methods for different types of object
- Methods are just ordinary functions
- When you call plot, R calls the appropriate plot method

Generic functions

- Generic functions don't do anything
- All the work is done by methods for different types of object
- Methods are just ordinary functions
 - with declarations in a package NAMESPACE
 - or R can guess based on function name

When you call the generic function R calls the appropriate method

Try these

```
print
methods("print")
stats:::print.acf
tools:::print.CRAN_package_reverse_dependencies_and_views
plot
methods("plot")
plot.ts
stats:::plot.lm
```

Also, try methods ("plot") after loading another package

What do we notice?

- print functions are all different
- names start with print., then the sort of thing they print
- mostly aren't visible just by name

Generic functions

- methods that actually do the work 'belong to' generic functions
- This is unusual: most popular OOP systems (Java, C++, Python) have methods belonging to data objects
- Important in R because functions are first-class objects (Week 5)
- Useful for functional programming with objects

Classes

- S3 classes are the things that specify which method to use
- Use class to attach a class to an object like a Post-It note
- That's all

```
x<-1
class(x)<-"numbat"</pre>
```

Try it

```
print.numbat <- function(x,...){
  cat(x,"numbats\n")
  invisible(x)
}</pre>
```

Defining classes

- R doesn't care what class you attach to an object
- **You** have to care
- class(x) <- "lm" makes R call lm methods on x</p>
- You are responsible for these methods being appropriate
- Documentation is important
- No real enforcement of encapsulation

Ways to set up classes

- vectors plus attributes (ts, POSIXct, matrix)
- lists plus attributes (lm, data.frame)
- environments plus attributes

Try it

```
unclass(.leap.seconds)
unclass(nhtemp)
unclass(trees)
m<-lm(log(Volume)~log(Girth)+log(Height),data=trees)
str(m)</pre>
```

Defining classes safely

- Document what all is in a valid object of your class
- Have a limited set of places where one is created
- Consider having a pure constructor function (ARP 13.3.1)
- Consider having a validator function (ARP 13.3.1)
- Have a user-friendly function to make valid objects

This is more important if someone else might want to make an object from your class

Constructors

- user-friendly: tibble, lm, acf, svydesign
- pure: new_factor, new_difftime (ARP 13.3.1)

Makes a new object and ensures that it is valid. Stops users creating the object themselves.

Defining methods

- A method should have **the same** arguments as the generic (plus maybe more)
- The name of the method is paste(generic,class,sep=".")
- Less ambiguous: use a package and declare the functions
 - S3method(generic, class) in NAMESPACE
 - @method generic class with devtools
- sloop::ftype tells you about the type of a function
- sloop::s3_get_method or getAnywhere finds methods
 even if they're hidden

default methods

Called when there is no specific method for the object (no class, or no matching class)

- mean.default
- summary.default
- head.default

Adding methods to your class

No rules on which methods, but informal standards

- Start with print, [
- summary for more information
- plot or image if possible (ggplot methods take more work)
- coef, vcov, maybe logLik and AIC for models
- resid for models with residuals

Base S3 classes

- The class for method choice isn't just class(x) for base types
- Use sloop::s3_class to be sure

```
> s3_class(1)
[1] "double" "numeric"
> s3_class(matrix(1,1,1))
[1] "matrix" "double" "numeric"
> class(1)
[1] "numeric"
> class(matrix(1,1,1))
[1] "matrix" "array"
```

Ambiguous cases

- t is a generic
- t.test is a generic
- t.test.formula is a method for t.test
- t.data.frame is a method for t
- list is not generic
- list.files isn't a method

Avoid using . as a word separator in function names that aren't methods. Use camelCase or snake_case or some other consistent approach

Defining generics

A typical generic function includes only a call to UseMethod

print

```
function (x, ...)
UseMethod("print")
<bytecode: 0x5f84e6cebfc8>
<environment: namespace:base>
```

Dispatch on another argument

Specify which argument to use for choosing the method (default is the first)

```
> survey:::svymean
function (x, design, na.rm = FALSE, ...)
{
    .svycheck(design)
    UseMethod("svymean", design)
}
```

Inheritance

The class attribute of an object can have multiple elements

- UseMethod() uses the first method that matches, or default
- NextMethod() uses the next method that matches

Polite conduct

- if you define a new generic, you can define methods for new and existing classes
- if you define a new class, you can define methods for new and existing generics
- don't define methods for someone else's class and generic (ask them)
- try not to define a generic with the same name as an existing one

Outline

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S3 vctrs

A package for defining new S3 vectors

- makes it easy to have them as tibble columns
- requires a pure constructor to make new objects, so they will all be valid
- supports double dispatch for binary operators (eg: +,-)
- has a complicated but reliable system for type conversion
- requires a lot of method definitions to get started

vctrs.r-lib.org

An example: S3 and vctrs

The rimu package represents multiple-response questions

remotes::install_github("tslumley/rimu")

```
data(usethnicity)
race<-as.mr(strsplit(as.character(usethnicity$Q5),""))
race<-mr_drop(race," ")
mtable(race)
hispanic<-as.mr(usethnicity$Q4==1,"Hispanic")
ethnicity<-mr_union(race,hispanic)
plot(ethnicity)
e_S3<-ethnicity[101:120]
e_vctrs<-as.vmr(e_S3,na.rm=TRUE)</pre>
```

Inheritance

```
mr_union<-function(x,y,...) UseMethod("mr_union")

mr_union.default<-function(x,y,...){
    x<-as.mr(x)
    y<-as.mr(y)
    if (length(x)!=length(y))
        stop("different numbers of observations in x and y")

.....
}</pre>
```

```
mr_union.vmr<-function(x,y,...) {
    r<-NextMethod()
    as.vmr(r)
}</pre>
```

Encapsulation

- base S3 version is matrix of logical, plus levels attribute
 - works in data.frame
- vctrs S3 version is list of vectors of strings
 - works in tibble

Pure constructor

```
new_vmr <- function(x,levels=unique(do.call(c,x))) {
  vctrs::new_list_of(x, ptype = character(),
  class = "vmr", levs=levels)
}</pre>
```

'helper' functions

```
> methods("as.mr")
[1] as.mr.character* as.mr.data.frame* as.mr.default*
[4] as.mr.factor* as.mr.list* as.mr.logical*
[7] as.mr.mr* as.mr.ms* as.mr.vmr*
> methods("as.vmr")
[1] as.vmr.default* as.vmr.mr*
```

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S4 requires classes and methods to be registered in R code (not just in packages)

- setClass defines the structure of a class
- new creates a new object from a class
- setMethod defines a method

It's possible to ask an object what methods it supports and get a reliable response.

S4 also allows multiple inheritance and multiple dispatch

Bioconductor

- Package system for high-throughput molecular biology
- Large data
- Structured data
- Annotated data
- New data types/structures all the time

It needs consistent infrastructure and large-scale collaboration: S4

bioconductor.org

Multiple dispatch

Choosing a method based on the class of more than one argument

- not very often useful
- important for matrices
- can be useful for plots

Multiple inheritance

AnnDbObjBimap is a class for storing look-up tables between different genomic identifiers (eg from different manufacturers) It is

- (by purpose) a two-way lookup object (BiMap)
- (by construction) an object containing a SQLite database (DbObj)

so it inherits generic functions from both these parents

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R6

Remember function factories from last week?

- Create a function closure with useful variables in its environment
- These variables are visible inside the function

Now do this with

- multiple variables in the shared environment
- multiple functions inheriting this environment
- allow the shared variables to be modified

R6 is good for...

- large data objects, to reduce copying
- external objects that R can't just copy (database connections, files,...)
- shared state such as games

R6 is more similar to C++/Java/Python OOP. It doesn't really support functional programming R6.r-lib.org