Using C and C++ with R

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Plan

Calling foreign languages

Built-in C interface

The Rcpp package

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Foreign languages

- ▶ C , C++
- ▶ Fortran
- ▶ Java¹.

Other scripting languages

- ► R/Perl² and R/Python³ bidirectional interfaces.
- ► There is also the system() function for direct access to OS functions.

²http://www.omegahat.org/RSPerl/





¹http://www.rforge.net/rJava/

Robert Gentleman, in *R Programming for Bioinformatics*, 2008, about R's built-in C interfaces

Since R is not compiled, in some situations its performance can be substantially improved by writing code in a compiled language. There are also reasons not to write code in other languages, and in particular we caution against premature optimization, prototyping in R is often cost effective. And in our experience very few routines need to be implemented in other languages for efficiency reasons. Another substantial reason not to use an implementation in some other language is increased complexity. The use of another language almost always results in higher maintenance costs and less stability. In addition, any extensions or enhancements of the code will require someone that is proficient in both R and the other language.

Rcpp does make some of the above caution statements slightly less critical.

- ▶ Why? R is getting slow or is not doing well in terms of memory management: for example for loops that can't be vectorised, recursion, . . .
- When? R can't do better and the slow code has been identified → Rprof

- ▶ Why? R is getting slow or is not doing well in terms of memory management: for example for loops that can't be vectorised, recursion, . . .
- When? R can't do better and the slow code has been identified → Rprof

▶ Why? Re-using existing infrastructure

Requirement for C/C++

Working compilers. On Windows, Rtools^{1,2}. On Mac, Xcode^{3,4}.

- 1. http://cran.r-project.org/bin/windows/Rtools/
- 2. http://cran.r-project.org/doc/manuals/R-admin.html#The-Windows-toolset
- 3. http://cran.r-project.org/doc/manuals/R-admin.html#Installing-R-under-_0028Mac_0029-OS-X
- 4. http://cran.r-project.org/doc/manuals/R-admin.html#Mac-OS-X

We will be using the following packages:

- ▶ inline and the cfunction to write inline C code that is compiled on the fly. (There is also a cxxfunction for C++ code).
- ▶ Rcpp, illustrating some of its functionality as well as the cppFunction for inline/on the fly compilation of C++ code.

Example

We have a DNA sequence, represented by a string of A, C, G and T and we want to compute the GC content.

```
x <- "ACCGGGTTTT"
gccountr <- function(x) table(strsplit(x, "")[1])
gccountr(x)
##
## A C G T
## 1 2 3 4</pre>
```

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The R C API

- Very frequent in R but has its quirks.
- ▶ Better know how to program in C.
- ▶ Documentation is not always easy to follow: R-Ext, R Internals as well as R and other package's code.

.C

- Not recommended.
- Arguments and return values must be primitives (vectors of doubles or integers).

.Call

- ► Accepts any R data structures as arguments and return values as SEXP.
- Manual memory management: allocate memory, protect objects to avoid them being garbage collected and subsequently unprotect them.

S (or symbolic) expression

SEXP is a super-type that matches all R data structures. Each data type has its own SEXP sub-type.

- REALSXP and INTSXP for double and integer vectors
- ► LGLSXP and STRSXP for logical and character vectors
- ▶ VECSXP for a list (NB: R list are called vectors at the C level)

Function input and outputs are always SEXP and will have to be coerced to the appropriate SXP sub-type.

Rinternals.h defines all C functions, data types and macros.

```
file.path(R.home(), "include", "Rinternals.h")
## [1] "/usr/local/lib/R/include/Rinternals.h"
```

```
library("inline")
## From Hadley Wickham, devtools wiki, adapted from inspect.c
## https://github.com/hadley/devtools/wiki/C-interface
sexp type <- cfunction(c(x = "ANY"), '
  switch (TYPEOF(x)) {
    case NTLSXP:
                      return mkString("NILSXP");
    case SYMSXP:
                      return mkString("SYMSXP");
    case LISTSXP:
                      return mkString("LISTSXP"):
    case CLOSXP:
                      return mkString("CLOSXP");
                      return mkString("ENVSXP");
    case ENVSXP:
    case PROMSXP:
                      return mkString("PROMSXP"):
    case LANGSXP:
                      return mkString("LANGSXP");
    case SPECTALSXP:
                      return mkString("SPECIALSXP");
    case BUILTINSXP.
                      return mkString("BUILTINSXP"):
    case CHARSXP:
                      return mkString("CHARSXP"):
    case LGLSXP:
                      return mkString("LGLSXP");
                      return mkString("INTSXP");
    case INTSXP:
    case REALSXP.
                      return mkString("REALSXP"):
    case CPLXSXP:
                      return mkString("CPLXSXP");
                      return mkString("STRSXP");
    case STRSXP:
    case DOTSXP:
                      return mkString("DOTSXP"):
                      return mkString("ANYSXP");
    case ANYSXP:
    case VECSXP:
                      return mkString("VECSXP");
    case EXPRSXP:
                      return mkString("EXPRSXP"):
    case BCODESXP:
                      return mkString("BCODESXP");
    case EXTPTRSXP:
                      return mkString("EXTPTRSXP");
    case WEAKREESXP.
                      return mkString("WEAKREFSXP"):
    case S4SXP:
                      return mkString("S4SXP"):
    case RAWSXP:
                      return mkString("RAWSXP");
                      return mkString("<unknown>");
    default:
}')
```

```
source("src/sexp.R")
sexp_type(1:3)
## [1] "INTSXP"
sexp type(10L)
## [1] "INTSXP"
sexp_type(TRUE)
## [1] "LGLSXP"
sexp_type(letters)
## [1] "STRSXP"
sexp_type(list(a = 1, b = letters))
## [1] "VECSXP"
sexp type(ls)
## [1] "CLOSXP"
```

Garbage collection

Every R object that is created at the C level (not function arguments, that R is already aware of) must be PROTECTed to avoid being garbage collected. Before the return statement, these must be explicitly UNPROTECTed.

```
SEXP x;
PROTECT(x = ...)
## do stuff
UNPROTECT(1)
return(x)
```

Object creation

- Allocate memory: allocVector, allocMatrix, alloc3DArray
- 2. Initialise objects: memset

```
SEXP x;
PROTECT(x = allocVector(INTSXP, 10) )
memset(INTEGER(x), 0, 10 * sizeof(int))
## do stuff
UNPROTECT(1)
return(x)
```

Accessing/setting SXP elements

- ► REAL(x)[i] if x is a REALSXP
- ► INTEGER(x)[i] if x is a INTSXP
- ▶ LOGICAL(x)[i] if x is a LGLSXP
- **•** . . .
- ► STRING_ELT(x, i) to access individual CHARSXP elements of a STRSXP
- ▶ VECTOR_ELT(x, i) to access individual elements of a VECSXP
- ► SET_STRING_ELT(str, i, x)) to set an element in a string.
- ► SET_VECTOR_ELT(vec, i, x)) to set an element in a list.

```
SEXP gccount(SEXP inseq) {
      int i, 1;
3
      char p;
      SEXP ans, dnaseq;
5
      PROTECT(dnaseq = STRING_ELT(inseq, 0)); // a CHARSXP
7
8
      1 = length(dnaseq);
9
      PROTECT(ans = allocVector(INTSXP, 4)):
10
      memset(INTEGER(ans), 0, 4 * sizeof(int));
11
12
      for (i = 0; i < 1; i++) {
13
        p = CHAR(dnaseq)[i];
14
        if (p == 'A')
15
          INTEGER (ans) [0]++;
16
        else if (p == 'C')
17
         INTEGER (ans)[1]++;
18
        else if (p == 'G')
19
          INTEGER (ans) [2]++:
20
        else if (p == 'T')
21
          INTEGER (ans) [3]++;
22
        else
23
           error("Wrong alphabet");
24
25
      UNPROTECT (2);
26
      return(ans):
27
    }
```

- 1. ingccount: embedding the C directly in R using the inline package.
- 2. gccount: writing the C into its own code file and using .Call.

./src/ingccount.R

```
library("inline")
ingccount <- cfunction(
  sig = c(inseq = "character"),
 bodv = "
 int i. 1:
 char p;
 SEXP ans, dnaseq;
 PROTECT(dnaseq = STRING_ELT(inseq, 0)); // a CHARSXP
 1 = length(dnaseq);
 PROTECT(ans = allocVector(INTSXP, 4));
 memset(INTEGER(ans), 0, 4 * sizeof(int));
 for (i = 0: i < 1: i++) {
    p = CHAR(dnaseq)[i];
   if (p == \'A\')
      INTEGER(ans)[0]++:
    else if (p == \'C\')
      INTEGER(ans)[1]++;
    else if (p == \'G\')
      INTEGER(ans)[2]++;
    else if (p == \'T\')
      INTEGER(ans)[3]++:
    else
      error(\"Wrong alphabet\");
 UNPROTECT(2):
 return(ans);
")
```

```
source("./src/ingccount.R")
ingccount(x)
## [1] 1 2 3 4
```

src/gccount.c

```
#include <R.h>
#include <Rdefines.h>
SEXP gccount(SEXP inseq) {
 int i, 1;
 char p;
 SEXP ans, dnaseq;
 PROTECT(dnaseg = STRING_ELT(inseg, 0)); // a CHARSXP
 1 = length(dnaseq);
 PROTECT(ans = allocVector(INTSXP, 4));
 memset(INTEGER(ans), 0, 4 * sizeof(int));
 for (i = 0; i < 1; i++) {
   p = CHAR(dnaseq)[i];
   if (p == 'A')
      INTEGER(ans)[0]++:
    else if (p == 'C')
      INTEGER(ans)[1]++:
    else if (p == 'G')
      INTEGER(ans)[2]++;
    else if (p == 'T')
      INTEGER (ans) [3]++:
    else
      error("Wrong alphabet");
 UNPROTECT(2);
 return(ans);
```

Use directly

- 1. Create a shared library: R CMD SHLIB gccount.c
- Load the shared object: dyn.load("gccount.so")
- 3. Create an R function that uses it:

```
gccountC <-
function(inseq) .Call("gccount", inseq)</pre>
```

4. Use you C code:

```
gccountC(x)
## [1] 1 2 3 4
```

In a package

- ▶ The C code comes in the src directory.
- ▶ The R wrapper will be

```
gccount <- function(inseq)
.Call("gccount", inseq, PACKAGE = "mypackage")</pre>
```

- Document the R function
- Export the R function and useDynLib(mypackge) in the NAMESPACE

```
library(sequences)
gccount
## function (inseq)
## {
       .Call("gccount", inseq, PACKAGE = "sequences")
##
## }
## <environment: namespace:sequences>
gccount(x)
## [1] 1 2 3 4
```

We could check that

```
if (TYPEOF(inseq) != STRSXP)
  error("Need a character vector!");
```

although

```
gccountC(123)
## Error: STRING_ELT() can only be applied to a
'character vector', not a 'double'
```

and type checking could easily be done at the R level. There is also isReal(x), isInteger(x), ... for atomics vectors.

There is of course much more to this ... see references at the end.

Benchmarking

Could we do better in R ? (should be asked first, really)

Could we do better in R? (should be asked first, really)

```
gccountr2 <-
function(x) tabulate(factor(strsplit(x, "")[[1]]))</pre>
```

Could we do better in R ? (should be asked first, really)

```
gccountr2 <-
 function(x) tabulate(factor(strsplit(x, "")[[1]]))
microbenchmark(gccountr(x),
              gccountr2(x),
              ingccount(x),
              gccountC(x),
              times = 1e4)
## Unit: microseconds
##
           expr min lq median uq
                                                 max neval
    gccountr(x) 226.985 233.620 238.439 256.528 4525.72 10000
##
   gccountr2(x) 117.544 122.432 124.807 133.468 4943.36 10000
##
   ingccount(x) 1.118 1.957 3.074 3.423 63.56 10000
##
## gccountC(x) 3.144 4.192 6.915 7.963 49.87 10000
```

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The Rcpp package

The Rcpp package

- Dirk Eddelbuettel and Romain Francois, with contributions by Douglas Bates, John Chambers and JJ Allaire.
- ► A flexible framework that facilitates the integration of R and C/C++ .
- http://www.rcpp.org/
- It comes with loads of documentation and examples: vignette(package = "Rcpp").
- ▶ All basic R types are implemented as C++ classes.
- ▶ No need to worry about garbage collection.

Associated packages

- RcppArmadillo Armadillo templated C++ library for linear algebra.
- ▶ RcppEigen high-performance Eigen linear algebra library.
- ► RInside use R from inside another C++ by wrapping the existing R embedding API in an easy-to-use C++ class.

C++ classes

Scalar	Vector	Matrix
double	NumericVector	NumericMatrix
int	${\tt IntegerVector}$	${ t Integer Matrix}$
string	${\tt CharacterVector}$	CharacterMatrix
bool	LogicalVector	LogicalMatrix

And Function, List, DataFrame ...

Automatic conversions from R (C) to C++ (R) using as (wrap).

- 1. ingccount2: embedding the C++ directly in R using the Rcpp::cppFunction package.
- 2. gccount2: in a package, writing the C++ into its own code file and using .Call.
- 3. gccountX: using sourceCpp to source the C++ file and export the function to R .

```
IntegerVector ingccount2(CharacterVector inseq) {
      IntegerVector ans(4);
      std::string s = Rcpp::as<std::string>(inseq[0]);
      int n = inseq(0).size();
5
6
7
8
      for (int i = 0; i < n; i++) {
        if (s[i] == 'A')
          ans[0]++;
        else if (s[i] == 'C')
9
          ans[1]++;
10
        else if (s[i] == 'G')
11
          ans[2]++:
12
        else if (s[i] == 'T')
13
          ans[3]++;
14
        else
15
           Rf_error(\"Wrong alphabet\");
16
17
      return(ans);
18
```

```
library("Rcpp")
cppFunction("
IntegerVector ingccount2(CharacterVector inseq) {
  IntegerVector ans(4);
  std::string s = Rcpp::as<std::string>(inseq[0]);
  int n = inseq(0).size();
  for (int i = 0; i < n; i++) {
    if (s[i] == 'A')
     ans[0]++;
    else if (s[i] == 'C')
      ans[1]++;
    else if (s[i] == 'G')
      ans[2]++:
    else if (s[i] == T)
      ans[3]++;
    else
      Rf_error(\"Wrong alphabet\");
  return(ans);
```

```
x <- "ACCGGGTTTT"
source("src/ingccount2.R")
ingccount2(x)
## [1] 1 2 3 4</pre>
```

./src/gccount2.cpp

```
#include <Rcpp.h>
using namespace Rcpp;
RcppExport SEXP gccount2(SEXP inseq)
 Rcpp::IntegerVector ans(4);
 Rcpp::CharacterVector dnaseq(inseq);
 std::string s = Rcpp::as<std::string>(dnaseq[0]);
 for (int i = 0; i < s.size(); i++) {
    char p = s[i];
    if (p=='A')
      ans[0]++;
    else if (p=='C')
      ans[1]++:
    else if (p=='G')
      ans[2]++;
    else if (p=='T')
      ans[3]++:
    else
      Rf_error("Wrong alphabet");
 return(ans);
```

In a package

- 1. You will need a Makevars file in the src directory.
- 2. Modify DESCRIPTION file:

```
Depends: Rcpp (>= 0.10.1)
LinkingTo: Rcpp
```

3. Create an R function that uses it

```
gccount2 <- function(inseq)
.Call("gccount2", inseq, PACKAGE = "mypackage")</pre>
```

- 4. Document the R function
- Export the R function and useDynLib(mypackge) in the NAMESPACE

Check the sequences package. But see Rcpp.package.skeleton below.

Using sourceCpp

- Write the C++ code into a cpp file, including headers and dedicated export statement (see next slide).
- Source it and use the R function.

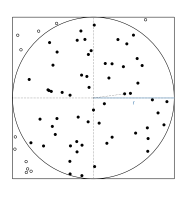
```
sourceCpp("src/gccountX.cpp")
gccountX(x)
## [1] 1 2 3 4
```

```
#include <Rcpp.h>
using namespace Rcpp;
// [[Rcpp::export]]
IntegerVector gccountX(CharacterVector inseq) {
 IntegerVector ans(4);
 std::string s = Rcpp::as<std::string>(inseq[0]);
 for (int i = 0; i < s.size(); i++) {
    if (s[i] == 'A')
      ans[0]++;
    else if (s[i] == 'C')
      ans[1]++:
    else if (s[i] == 'G')
      ans[2]++;
    else if (s[i] == T)
      ans[3]++:
    else
      Rf_error("Wrong alphabet");
 return(ans);
```

sugar

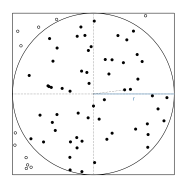
sugar (for syntactic sugar) is a set of C++ functions that (mostly) work and look like their R couterparts. Allows for example compact vectorised expression. Looks like R with the C++ efficiency. (Rcpp-sugar vignette/paper)

- ▶ Vectorised arithmetic and logical operators: +, >, !, ...
- ► Functions: seq_len, all, any, sapply, rnorm, abs, ...



$$\frac{d}{n} = \frac{\pi r^2}{4r^2} = \pi/4$$

$$\pi \approx \frac{4d}{n}$$



```
\frac{\frac{d}{n} = \frac{\pi r^2}{4r^2} = \pi/4
\pi \approx \frac{4d}{n}
```

```
piR <- function(N) {
    x <- runif(N)
    y <- runif(N)
    d <- sqrt(x^2 + y^2)
    4 * sum(d < 1.0) / N
}</pre>
```

```
piR <- function(N) {
    x <- runif(N)
    y <- runif(N)
    d <- sqrt(x^2 + y^2)
    return(4 * sum(d < 1.0) / N)
}</pre>
```

```
piR <- function(N) {
    x <- runif(N)
    y <- runif(N)
    d <- sqrt(x^2 + y^2)
    return(4 * sum(d < 1.0) / N)
}</pre>
```

```
#include <Rcpp.h>
using namespace Rcpp;
// [[Rcpp::export]]
double piSugar(const int N) {
    RNGScope scope; // ensure RNG gets set/reset
    NumericVector x = runif(N);
    NumericVector y = runif(N);
    NumericVector d = sqrt(x*x + y*y);
    return 4.0 * sum(d < 1.0) / N;
}</pre>
```

```
source("src/pi.R")
library("Rcpp")
sourceCpp("src/pi.cpp")
N < -1e + 06
set.seed(42)
resR <- piR(N)
set.seed(42)
resCpp <- piSugar(N)
stopifnot(identical(resR, resCpp))
```

Exercise 1: Translate these R functions into C or C++

```
sum
rowSums
pdistR <- function(x, ys)
   sqrt( (x - ys) ^ 2 )
sety <- function(x, y) {
      x[x > 0] <- y
      x[x < 0] <- -y
      x
}
grY <- function(x, y) x[x > y]
```

Rcpp modules

Using S4 Reference Classes to reflect C++ classes and methods (see the Rcpp-modules vignette).

Interface to the Standard template library

C++ library with data structure and algorithms: vectors, arrays, stacks, iterators, accumulators, . . .

Using Rcpp in packages

```
Rcpp.package.skeleton("mypackage")
mypackage
|-- DESCRIPTION
l-- man
   |-- mypackage-package.Rd
     -- rcpp_hello_world.Rd
 -- NAMESPACE
I -- R.
     -- rcpp_hello_world.R
|-- Read-and-delete-me
 -- src
    I-- Makeyars
    |-- Makevars.win
    |-- rcpp_hello_world.cpp
```

-- rcpp_hello_world.h

References

Further reading

- Writing R Extensions, R Core team.
- ▶ Robert Gentleman, R Programming for Bioinformatics, 2008.
- Rcpp documentation and Rcpp.org.
- ▶ Dirk Eddelbuettel, Seamless R and C++ Integration with Rcpp, Springer, 2013.
- ▶ Dirk Eddelbuettel and Romain Francois, Rcpp: Seamless R and C++ Integration, Journal of Statistical Software, Vol. 40, Issue 8, Apr 2011, http://www.jstatsoft.org/v40/i08/.
- Relevant devtools sections: C interface and Rcpp.

- ▶ This work is licensed under a CC BY-SA 3.0 License.
- Course web page and more material: https://github.com/lgatto/TeachingMaterial

Thank you for you attention