

NiftiArray: Solving Your Big Nifti Needs

source at https://github.com/muschellij2/niftiarray_talk

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Start of a Package: Plotting a voxel from 800 Subjects in a Shiny App

How can you read in a NIfTI image in R?

Options for reading in NIfTI images

Array like

- oro.nifti::readNIfTI/neurobase::readnii
- AnalyzeFMRI::f.read.volume() can't read gzipped
- neuroim::loadVolume("example.nii")

C++ pointer

- ANTsCore::antsImageRead
- RNifti::readNifti

From RNifti:

https://github.com/jonclayden/RNifti#performance

```
library(microbenchmark)
microbenchmark(AnalyzeFMRI::f.read.volume("example.nii"),
               ANTsRCore::antsImageRead("example.nii"),
               neuroim::loadVolume("example.nii"),
               oro.nifti::readNIfTI("example.nii"),
               RNifti::readNifti("example.nii"),
               tractor.base::readImageFile("example.nii"), unit="ms")
# Unit: milliseconds
                                         expr
                                                    min
                                                                       mean
    AnalyzeFMRI::f.read.volume("example.nii") 26.312881 26.769244 29.685981
      ANTsRCore::antsImageRead("example.nii") 1.150787 1.626918 2.149145
           neuroim::loadVolume("example.nii") 34.596506 37.732245 54.065227
          oro.nifti::readNIfTI("example.nii") 57.059828 61.953430 89.386706
             RNifti::readNifti("example.nii") 0.986773 1.136562 1.683821
   tractor.base::readImageFile("example.nii") 33.380407 34.096574 34.961812
      median
                    ua
                              max neval
   27.301693 28.214908 185.441664
                                    100
    2.210696 2.481675
                        3.376350
                                    100
   40.714061 45.425047 192.978225
                                    100
   65,064212 71,425561 220,709246
                                    100
    1.501528 1.856601
                         7.961566
                                    100
   34.617259 35.257633 42.108584
                                    100
```

What do we want

- 1. Something fast
- 2. Has overloaded functions (e.g. image + image = image)
- 3. Can use most plotting functions on it (e.g. ortho2)
- 4. Is array like but has header
- 5. Keeps memory low
- 6. Has random access

Current issues

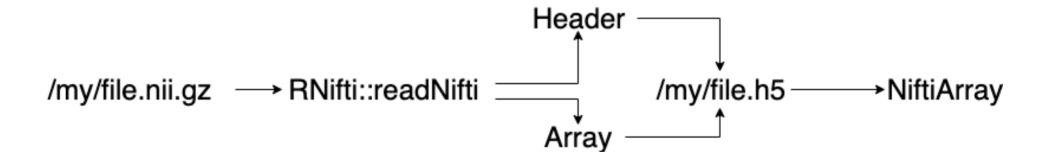
- oro.nifti::readNIfTI slow
- ANTsCore::antsImageRead installation is
- RNifti::readNifti-from
 https://github.com/jonclayden/RNifti/issues/14:

...the "native" data size in the C struct is roughly 4 GiB, while the R array uses 8 GiB because it uses double precision... readNifti(..., internal=FALSE) the object has both the C struct and the R array, for a total of 12 GiB.

None have random access (just read in one slice or 1 timepoint)

Solution: NiftiArray

- Stores image and header in HDF5 (Hierarchical Data Format) format
- Based on HDF5Array from Bioconductor
 (https://bioconductor.org/packages/release/bioc/html/HDF5Array.html)



Takes longer than RNifti (the first time)

```
fname = fslr::mni fname()
system.time({
 rnifti = RNifti::readNifti(fname)
})
##
    user system elapsed
##
    0.124 0.017 0.144
system.time({
 narray = NiftiArray::writeNiftiArray(fname)
 narray@seed@filepath
 narray
})
##
     user system elapsed
    4.658 0.131 4.878
```

Sizes

```
format(object.size(rnifti), units = "Kb")

## [1] "28208.3 Kb"

format(object.size(narray), units = "Kb")

## [1] "8.7 Kb"
```

Header Information

```
print(nifti header(narray))
## NIfTI-1 header
      sizeof hdr: 348
##
        dim info: 0
##
             dim: 3 182 218 182 1 1 1 1
       intent p1: 0
##
       intent p2: 0
##
       intent p3: 0
##
     intent code: 0
        datatype: 8
##
          bitpix: 32
     slice start: 0
         pixdim: -1 1 1 1 0 0 0 0
##
      vox offset: 352
     scl slope: 0
     scl inter: 0
##
       slice end: 0
##
      slice code: 0
    xyzt units: 10
         cal max: 8000
```

cal min: 3000

Accessing voxels

```
narray[18, 5, 14]
## [1] 572
sum(narray > 9990)
## [1] 20
narray[ narray > 9990]
## [1] 9999 9999 9992 9997 9991 9993 9997 9998 9999 9999 9994 9997 9998 9999
## [15] 9998 9991 9993 9991 9996 9994
DelayedArray::extract array(narray, index = list(NULL, NULL, 1))
## , , 1
##
          [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11] [,12]
                                                                                      12/22
   [1,]
                                30
```

What are the benefits?

- Overall most benefits come from saving H5 files on "big" computer
- Random access the DelayedArray package
- Operations on DelayedArray objects, the DelayedMatrixStats package https://bioconductor.org/packages/release/bioc/html/DelayedMatrixStats.html

What are the benefits? Reshaping to a Matrix

```
d = dim(narray); print(d)

## [1] 182 218 182

dim(narray) = c(d, 1)
new_dim = c(prod(d), 1)
newarray = writeNiftiArray(narray)
dim(newarray)
## [1] 182 218 182 1
```

What are the benefits? Reshaping to a Matrix

```
mat = ReshapedNiftiArray(newarray@seed@filepath, dim = new dim)
dim(mat)
## [1] 7221032
head (mat)
## <6 x 1> DelayedMatrix object of type "integer":
##
   [,1]
## [1,]
## [2,]
## [3,] 0
## [4,]
## [5,]
          0
```

[6,]

Why? Voxel-wise operations in low-memory setting

Why? 4D Example

```
## [1] 96 96 60 3
```

Why? 4D Example

Back to an image

```
arr = array(med, dim = dim(from)[1:3])
med_array = writeNiftiArray(arr, header = nifti_header(from))
as(med_array, "niftiImage")

## Image array of mode "double" (4.2 Mb)
## - 96 x 96 x 60 voxels
## - 2.5 x 2.5 x 2.5 mm per voxel
```

Issues

- Need to write H5 file first (do in parallel on cluster)
- Memory tradeoff with speed due to chunking
- Most of this needs things in the same space

Use cases

- Something on Shiny for speed or memory
 - shinyapps.io has very restrictive memory settings for imaging
- Analyzing 1000s of subjects together
- Pass object from R to Matlab: https://www.mathworks.com/help/matlab/ref/hdf5re ad.html

Extensions

- Anything that works with DelayedArrays should work here
- Check out Bioconductor
- Example: mbkmeans mini-batch k-means
 https://bioconductor.org/packages/release/bioc/html/mbkmeans.html