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# Working with the DICOM and NIfTI Data Standards in R

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### Abstract

Two packages (**oro.dicom** and **oro.nifti**) are provided for the interaction with and manipulation of medical imaging data that conform to the DICOM standard or ANA-LYZE/NIfTI formats. DICOM data, from a single file or single directory or directory tree, may be uploaded into R using basic data structures: a data frame for the header information and a matrix for the image data. A list structure is used to organize multiple DICOM files. The S4 class framework is used to develop basic ANALYZE and NIfTI classes, where NIfTI extensions may be used to extend the fixed-byte NIfTI header. One example of this, that has been implemented, is an XML-based "audit trail" that tracks the history of operations applied to a dataset. The conversion from DICOM to ANALYZE/NIfTI is straightforward using the capabilities of both packages. The S4 classes have been developed to provide a user-friendly interface to the ANALYZE/NIfTI data formats; allowing easy data input, data output, image processing and visualization.

Keywords: export, imaging, import, medical, visualization.

### 1. Introduction

Medical imaging is well established in both the clinical and research areas with numerous equipment manufacturers supplying a wide variety of modalities. The DICOM (Digital Imaging and Communications in Medicine; <a href="http://medical.nema.org">http://medical.nema.org</a>) standard was developed from earlier standards and released in 1993. It is the data format for clinical imaging equipment and a variety of other devices whose complete specification is beyond the scope of this paper. All major manufacturers of medical imaging equipment (e.g., GE, Siemens, Philips)

have so-called DICOM conformance statements that explicitly state how their hardware implements DICOM. The DICOM standard provides interoperability across hardware, but was not designed to facilitate efficient data manipulation and image processing. Hence, additional data formats have been developed over the years to accommodate data analysis and image processing.

The ANALYZE format was developed at the Mayo Clinic (in the 1990s) to store multidimensional biomedical images. It is fundamentally different from the DICOM standard since it groups all images from a single acquisition (typically three- or four-dimensional) into a pair of binary files, one containing header information and one containing the image information. The DICOM standard groups the header and image information, typically a single two-dimensional image, into a single file. Hence, a single acquisition will contain multiple DICOM files but only a pair of ANALYZE files.

The NIfTI format was developed in the early 2000s by the DFWG (Data Format Working Group) in an effort to improve upon the ANALYZE format. The resulting NIfTI-1 format adheres to the basic header/image combination from the ANALYZE format, but allows the pair of files to be combined into a single file and re-defines the header fields. In addition, NIfTI extension allow to store any additional information.

The material presented here provides users with a method of interacting with DICOM, AN-ALYZE and NIfTI files in R (R Development Core Team 2010). Real-world datasets, that are publicly available, are used to illustrate the basic functionality of the two packages: **oro.dicom** and **oro.nifti**. Major features include data input/output, visualization and conversion from DICOM to ANALYZE/NIfTI. These packages should appeal not only to other R package developers, but also to scientists who want to manipulate medical imaging data using R without writing and validating basic data input/output functionality. Packages already available on CRAN that utilize **oro.dicom** and **oro.nifti** include **cudaBayesreg** (Ferreira da Silva 2010a), **dcemriS4** (Whitcher and Schmid 2010) and **dpmixsim** (Ferreira da Silva 2010b).

# 2. oro.dicom: DICOM Data Input/Output in R

The DICOM "standard" for data acquired using a clinical imaging device is very broad and complex. Roughly speaking each DICOM-compliant file is a collection of fields organized into two two-byte sequences (group, element) that are represented as hexadecimal numbers and form a tag. The (group, element) combination establishes what type of information is forthcoming in the file. There is no fixed number of bytes for a DICOM header. The final (group, element) tag should be the "pixel data" tag (7FE0,0010), such that all subsequent information is related to the image(s).

All attributes in the DICOM standard require different data types for correct representation. These are known as value representations (VRs) in DICOM, which may be encoded explicitly or implicitly. There are 27 explicit VRs defined in the DICOM standard, and provided in Table 1. Detailed explanations of these data types are provided in the Section 6.2 (part 5) of the DICOM standard (http://medical.nema.org). The first column provides the two-character string that is present in each entry of the header field and the second column provides a descriptive name for the abbreviated code. The third column provides the maximum length of the data associated with the VR (in bytes), where a length of zero bytes may be interpreted as having an unknown or unlimited number of bytes associated with the VR. The fourth

| Code                | Name                          | Bytes | Fixed |
|---------------------|-------------------------------|-------|-------|
|                     |                               |       |       |
| AE                  | ApplicationEntity             | 16    | 0     |
| AS                  | AgeString                     | 4     | 1     |
| AT                  | AttributeTag                  | 4     | 1     |
| CS                  | CodeString                    | 16    | 0     |
| DA                  | Date                          | 8     | 1     |
| DS                  | DecimalString                 | 16    | 0     |
| $\operatorname{DT}$ | DateTime                      | 26    | 0     |
| $\operatorname{FL}$ | FloatingPointSingle           | 4     | 1     |
| FD                  | ${\bf Floating Point Double}$ | 8     | 1     |
| IS                  | IntegerString                 | 12    | 0     |
| LO                  | LongStrong                    | 64    | 0     |
| $\operatorname{LT}$ | LongText                      | 10240 | 0     |
| OB                  | OtherByteString               | 0     | 0     |
| ow                  | OtherWordString               | 0     | 0     |
| PN                  | PersonName                    | 64    | 0     |
| SH                  | ShortString                   | 16    | 0     |
| $\operatorname{SL}$ | SignedLong                    | 4     | 1     |
| SQ                  | SequenceOfItems               | 0     | 0     |
| SS                  | SignedShort                   | 2     | 1     |
| $\operatorname{ST}$ | ShortText                     | 1024  | 0     |
| TM                  | Time                          | 16    | 0     |
| UI                  | ${\bf Unique Identifier UID}$ | 64    | 0     |
| $\operatorname{UL}$ | UnsignedLong                  | 4     | 1     |
| UN                  | Unknown                       | 0     | 0     |
| US                  | UnsignedShort                 | 2     | 1     |
| UT                  | ${\bf Unlimited Text}$        | 0     | 0     |

Table 1: Value representations in the DICOM standard. A value of zero bytes may be interpreted as having an unknown or unlimited number of bytes associated with the VR.

Data element with explicit VR of OB, OF, OW, SQ, UT or UN:

```
+-----+
| 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 |
+---+---+---+
| <Group-->| <Element>| <VR---->| <0x00000->| <Length------>| <Value->
```

Data element with explicit VR other than as shown above:

```
+-----+
| 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
+---+---+
| <Group-->| <Element>| <VR---->| <Length->| <Value->
```

Data element with implicit VR:

```
+-----+
| 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
+---+---+
| <Group-->| <Element>| <Length----->| <Value->
```

Figure 1: Byte ordering for a single (group, element) tag in the DICOM standard. Explicit VRs store the VR as text characters in two bytes. More information is provided in Section 7, Part 3.5-2009 of the DICOM standard (http://medical.nema.org).

column is not used in the current implementation of the **oro.dicom** package. Internal functions have been written to manipulate each of the value representations and are beyond the scope of this article. The functions **str2date** and **str2time** are useful for converting from the DICOM Datetime and Time value representations to R date and time objects, respectively.

### 2.1. The DICOM Header

Accessing the information stored in a single DICOM file is provided using the dicomInfo function. The basic structure of a DICOM file is summarized in Figure 1, for both explicit and implicit value representations. The first two bytes represent the group tag and the second two bytes represent the element tag, regardless of the type of VR. The third set of two bytes contains the characters of the VR on which a decision about being implicit or explicit is made. Explicit VRs of type (OB, OF, OW, SQ, UT, UN) skip bytes six and seven (counting from zero), convert the next four bytes into an integer length and read length number of objects from the DICOM file. All other explicit VRs follow a slightly different path where bytes six and seven (counting from zero) provide an integer length and all remaining bytes are read in as the value. If the character string in bytes four and five do not correspond to a known VR (Figure 1), then the (group,element) tag is declared to be implicit, the length is taken from bytes four through seven and all remaining bytes contribute to the value.

The basic structure of the resulting object is a list with two elements: the DICOM header (hdr) and the DICOM image (img). The header information is organized in a data frame

with six columns and an unknown number of rows depending on the input parameters.

```
R> fname <- system.file(file.path("dcm", "Abdo.dcm"), package = "oro.dicom")</pre>
R> abdo <- dicomInfo(fname)</pre>
R> names(abdo)
[1] "hdr" "img"
R> abdo$hdr[1:5, ]
  group element
                                        name code length
1 0002
           0000
                                 GroupLength
                                               UL
                                                        4
2 0002
           0001 FileMetaInformationVersion
                                               0B
                                                        2
3 0002
           0002
                    MediaStorageSOPClassUID
                                               UI
                                                       26
  0002
           0003 MediaStorageSOPInstanceUID
                                               UI
                                                       38
   0002
                          TransferSyntaxUID
                                               UI
                                                       20
           0010
                                    value sequence
1
                                      166
2
                                  skipped
3
               1.2.840.10008.5.1.4.1.1.4
4 1.3.46.670589.11.0.4.1996082307380007
5
                     1.2.840.10008.1.2.1
R> abdo$hdr[nrow(abdo$hdr) - 4:0, ]
```

|    | group | element | name                        | code | length | value | sequence |
|----|-------|---------|-----------------------------|------|--------|-------|----------|
| 80 | 0028  | 0102    | HighBit                     | US   | 2      | 11    |          |
| 81 | 0028  | 0103    | ${\tt PixelRepresentation}$ | US   | 2      | 0     |          |
| 82 | 0028  | 1050    | WindowCenter                | DS   | 4      | 530   |          |
| 83 | 0028  | 1051    | ${\tt WindowWidth}$         | DS   | 4      | 1052  |          |
| 84 | 7FE0  | 0010    | PixelData                   | OW   | 131072 |       |          |

The ordering of the rows is identical to the ordering in the original DICOM file. Hence, the first five tags in the DICOM header of Abdo.dcm are: GroupLength, FileMetaInformationVersion, MediaStorageSOPClassUID, MediaStorageSOPInstanceUID and TransferSyntaxUID. The last five tags in the DICOM header are also shown, with the very last tag indicating the start of the image data for that file and the number of bytes (131072) involved. When additional tags in the DICOM header information are queried (via extractHeader)

```
R> extractHeader(abdo$hdr, "BitsAllocated")
[1] 16
R> extractHeader(abdo$hdr, "Rows")
[1] 256
```

```
R> extractHeader(abdo$hdr, "Columns")
```

```
[1] 256
```

it is clear that the data are consistent with the header information in terms of the number of bytes  $(256 \times 256 \times (16/8) = 131072)$ .

The first five columns are taken directly from the DICOM header information (group, element, code, length and value) or inferred from that information (name). Note, the (group,element) values are stored as character strings even though they are hexadecimal numbers. All aspects of the data frame may be interrogated in R in order to extract relevant information from the DICOM header; e.g., "BitsAllocated" as above. The sequence column is used to keep track of tags that are embedded in a fixed-length SequenceItems tag or between a SequenceItem-SequenceDelimitationItem pair.

When multiple DICOM files are located in a single directory, or spread across multiple directories, one may use the function dicomSeparate (applied here to the directory hk-40).

```
R> fname <- system.file("hk-40", package = "oro.dicom")
R> hk40 <- dicomSeparate(fname, verbose = TRUE, counter = 10)

40 files to be processed!
10 files processed...
20 files processed...
30 files processed...
40 files processed...
R> unlist(lapply(hk40, length))
hdr img
40 40
```

The object associated with dicomSeparate is now a nested set of lists, where the hdr element is a list of data frames and the img element is a list of matrices. These two lists are associated in a pairwise sense; i.e., hdr[[1]] is the header information for the image img[[1]]. Default parameters recursive = TRUE and pixelData = TRUE (which is actually an input parameter for dicomInfo) allow the user to search down all possible sub-directories and upload the image in addition to the header information, respectively. Also, by default all files are treated as DICOM files unless the exclude parameter is set to the unwanted file extension; e.g., exclude = "xml".

The list of DICOM header information across multiple files may be converted to a single data frame using dicomTable, and written to disc for further analysis; e.g., using write.csv.

```
R> hk40.info <- dicomTable(hk40$hdr)
R> write.csv(hk40.info, file = "hk40_header.csv")
R> sliceloc.col <- which(hk40$hdr[[1]]$name == "SliceLocation")
R> sliceLocation <- as.numeric(hk40.info[, sliceloc.col])
R> sliceLocation[1:5]
```

```
[1] 160.9315 157.8315 154.7315 151.6315 148.5315
R> diff(sliceLocation[1:5])
[1] -3.1 -3.1 -3.1 -3.1
R> unique(extractHeader(hk40$hdr, "SliceThickness"))
[1] 3.125
The tag SliceLocation is extracted from the DICOM header information (at the first element
in the list) and processed using the diff function, and should agree with the SliceThickness
tag. Single DICOM fields may also be extracted from the list of DICOM header information
that contain attributes that are crucial for further image processing; e.g., extracting relevant
MR sequences or acquisition timings.
R> extractHeader(hk40$hdr, "SliceLocation")[1:5]
[1] 160.9315 157.8315 154.7315 151.6315 148.5315
R> modality <- extractHeader(hk40$hdr, "Modality", numeric = FALSE)
R> matchHeader(modality, "mr")[1:5]
[1] TRUE TRUE TRUE TRUE TRUE
R> (seriesTime <- extractHeader(hk40$hdr, "SeriesTime", numeric = FALSE))
 [1] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
 [5] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
 [9] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[13] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[17] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[21] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[25] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[29] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[33] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
[37] "113751.966000" "113751.966000" "113751.966000" "113751.966000"
R> str2time(seriesTime[1:5])
$txt
[1] "11:37:51.96600" "11:37:51.96600" "11:37:51.96600"
[5] "11:37:51.96600"
$time
[1] 41871.97 41871.97 41871.97 41871.97
```

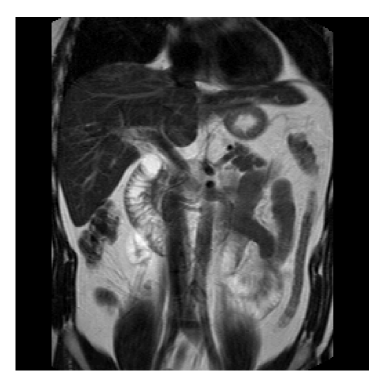


Figure 2: Coronal slice of the abdomen viewed in *neurological* convention (left is right and right is left).

### 2.2. The DICOM Image

Most DICOM files involve a single slice from an acquisition – the image. A notable exception is the Siemens MOSAIC format (addressed in Section 2.2.1). The **oro.dicom** package assumes the image is stored as a flat file of two-byte integers without compression. A variety of additional image formats are possible within the DICOM standard; for example, RGB-colorized, JPEG, JPEG Lossless, JPEG 2000 and run-length encoding (RLE). Going back to the Abdo.dcm example, the image is accessed via

```
R> image(t(abdo$img), col = grey(0:64/64), axes = FALSE, xlab = "", + ylab = "")
```

where the transpose operation is necessary for proper visualization of the image. Figure 2 displays a coronal slice through the abdomen from an MRI acquisition. All information from the original data acquisition should accompany the image through the DICOM header, and this information is utilized as much as possible by **oro.dicom** to simplify the manipulation of DICOM data. As previously shown, this information is easily available to the user by matching DICOM header fields with valid strings. Note, the function <code>extractHeader</code> assumes the output should be coerced via <code>as.numeric</code> but this may be disabled setting the input parameter <code>numeric=FALSE</code>.

```
R> extractHeader(abdo$hdr, "Manufacturer", numeric = FALSE)
```

[1] "Philips"

```
R> extractHeader(abdo$hdr, "RepetitionTime")
[1] 2000
R> extractHeader(abdo$hdr, "EchoTime")
[1] 100
```

The basic DICOM file structure does not encourage the analysis of multi-dimensional imaging data (e.g., 3D or 4D) commonly acquired on clinical scanners. Hence, the **oro.dicom** package has been developed to access DICOM files and facilitate their conversion to the NIfTI or ANALYZE formats. The conversion process requires the **oro.nifti** package and will be outlined in Section 4.

### Siemens MOSAIC Format

Siemens multi-slice EPI (echo planar imaging) data may be collected as a "mosaic" image; i.e., all slices acquired in a single TR (repetition time) frame of a dynamic run are stored in a single DICOM file. The images are stored in an  $M \times N$  array of images. The function create3D will try to guess the number of images embedded within the single DICOM file using the AcquisitionMatrix field. If this doesn't work, one may enter the (M, N) doublet explicitly.

```
R> fname <- system.file(file.path("dcm", "MR-sonata-3D-as-Tile.dcm"),
+    package = "oro.dicom")
R> dcm <- dicomInfo(fname)
R> dim(dcm$img)

[1] 384 384

R> dcmImage <- create3D(dcm, mosaic = TRUE)
R> dim(dcmImage)

[1] 64 64 36
```

Figure 3a is taken from the raw DICOM file, in mosaic format, and displayed with the default margins in R. Figure 3b is displayed after re-organizing the original DICOM file into a three-dimensional array (it was also converted to the NIfTI format for ease of visualization using the overloaded image function in **oro.nifti**).

# 3. oro.nifti: NIfTI-1 Data Input/Output in R

Although the industry standard for medical imaging data is DICOM, another format has come to be heavily used in the image analysis community. The ANALYZE format was originally developed in conjunction with an image processing system (of the same name) at the Mayo Foundation. A common version of the format, although not the most recent, is called

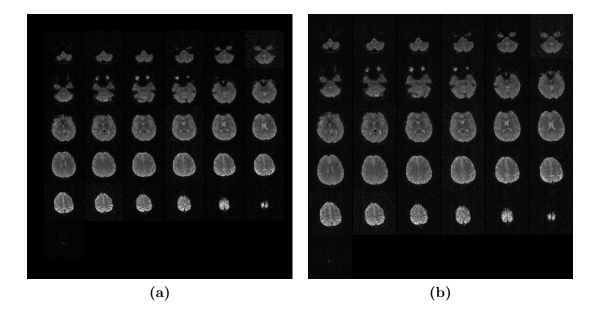


Figure 3: (a) Single MOSAIC image as read in from dicomInfo. (b) Lightbox display of three-dimensional array of images after processing via create3D.

ANALYZE 7.5. A copy of the file ANALYZE75.pdf has been included in **oro.nifti** (accessed via system.file("doc/ANALYZE75.pdf", package="oro.dicom")) since it does not appear to be available from www.mayo.edu any longer. An ANALYZE 7.5 format image is comprised of two files, the ".hdr" and ".img" files, that contain information about the acquisition and the acquisition itself, respectively. A more recent adaption of this format is known as NIfTI-1 and is a product of the Data Format Working Group (DFWG) from the Neuroimaging Informatics Technology Initiative (NIfTI; http://nifti.nimh.nih.gov). The NIfTI-1 data format is almost identical to the ANALYZE format, but offers a few improvements

- merging of the header and image information into one file (.nii)
- re-organization of the 348-byte fixed header into more relevant categories
- possibility of extending the header information.

### 3.1. The NIfTI Header

The NIfTI header inherits its structure (348 bytes in length) from the ANALYZE data format. The last four bytes in the NIfTI header correspond to the "magic" field and denote whether or not the header and image are contained in a single file (magic = "n+1\0") or two separate files (magic = "ni1\0"), the latter being identical to the structure of the ANALYZE data format. The NIfTI data format added an additional four bytes to allow for "extensions" to the header. By default these four bytes are set to zero.

The first example of reading in, and displaying, medical imaging data in NIfTI format avg152T1\_LR\_nifti.nii.gz was obtained from the NIfTI website (http://nifti.nimh.nih.gov/nifti-1/). Successful execution of the commands

```
R> fname <- system.file(file.path("nifti", "mniLR.nii.gz"),</pre>
      package = "oro.nifti")
R> (mniLR <- readNIfTI(fname))</pre>
NIfTI-1 format
                   : niftiAuditTrail
  Type
  Data Type
                   : 2 (UINT8)
  Bits per Pixel : 8
  Slice Code
                   : 0 (Unknown)
  Intent Code
                   : 0 (None)
  Qform Code
                   : 0 (Unknown)
  Sform Code
                   : 4 (MNI_152)
  Dimension
                   : 91 x 109 x 91
  Pixel Dimension : 2 x 2 x 2
  Voxel Units
                   : mm
  Time Units
                   : sec
R> aux.file(mniLR)
                              11
[1] "none
R> descrip(mniLR)
[1] "FSL3.2beta"
```

produces an S4 "nifti" object (or "niftiAuditTrail" if the audit trail option is set). Two accessor functions are also provided: aux.file and descrip. The former is used to access the original name of the file (if it has been provided) and the latter is the name of a valid NIfTI header field used to hold a "description" (up to 80 characters in length).

### 3.2. The NIfTI Image

Image information begins at the byte position determined by the voxoffset slot. In a single NIfTI file (magic = "n+1\0"), this is by default after the first 352 bytes. Header extensions extend the size of the header and come before the image information leading to a consequent increase of voxoffset for single NIfTI files. Split NIfTI files do not have this problem and voxoffset is set at 0.

The image function has been overloaded so that it behaves differently when dealing with medical image objects (nifti and anlz). The command

### R> image(mniLR)

produces a three-dimensional array of the MNI brain, with the default NIfTI axes, and is displayed on a  $10 \times 10$  grid of images (Figure 4a). The image function for medical image S4 objects is an attempt to balance minimal user input with enough flexibility to customize the display when necessary. For example, single slices may be viewed by using the option plot.type="single" in conjuction with the option z= to specify the slice.

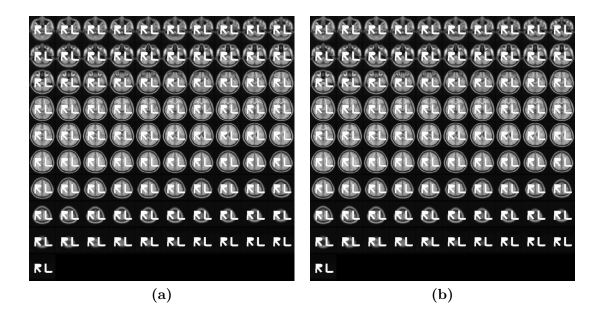


Figure 4: (a) Axial slices of MNI volume mniLR\_nifti stored in the *neurological* convention (right-is-right), but displayed in the *radiological* convention (right-is-left). (b) Axial slices of MNI volume mniRL\_nifti stored and displayed in the *radiological* convention.

The second example of reading in and displaying medical imaging data in the NIfTI format avg152T1\_RL\_nifti.nii.gz was also obtained from the NIfTI website (http://nifti.nimh.nih.gov/nifti-1/). Successful execution of the commands

```
R> fname <- system.file(file.path("nifti", "mniRL.nii.gz"),
      package = "oro.nifti")
   (mniRL <- readNIfTI(fname))</pre>
NIfTI-1 format
  Туре
                   : niftiAuditTrail
                   : 2 (UINT8)
  Data Type
  Bits per Pixel
                   : 8
  Slice Code
                   : 0 (Unknown)
                   : 0 (None)
  Intent Code
  Qform Code
                   : 0 (Unknown)
                   : 4 (MNI_152)
  Sform Code
  Dimension
                   : 91 x 109 x 91
  Pixel Dimension : 2 x 2 x 2
  Voxel Units
                   : mm
  Time Units
                   : sec
```

R> image(mniRL)

produces a three-dimensional array of the MNI brain that is displayed in a  $10 \times 10$  grid of images (Figure 4b). The two sets of data in Figure 4 are stored in two different orientations,

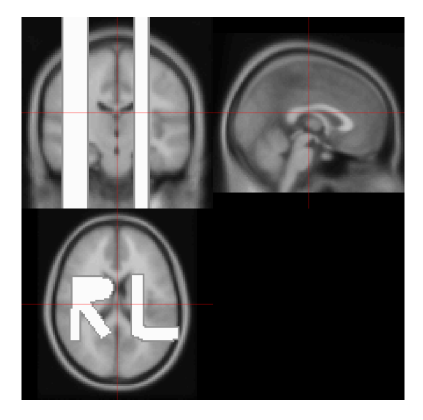


Figure 5: Orthographic display of the MNI volume mniRL\_nifti. By default the mid-axial, mid-saggital and mid-coronal planes are chosen.

commonly referred to as the *radiological* and *neurological* conventions. The neurological convention is where "right is right" and one is essentially looking through the subject. The radiological convention is where "right is left" and one is looking at the subject.

An additional graphical display function has been added for nifti and anlz objects that allows a so-called orthographic visualization of the data.

### R> orthographic(mniRL)

As seen in Figure 5 the mid-axial, mid-saggital and mid-coronal planes are displayed by default. The slices used may be set using xyz = c(I,J,K), where (I,J,K) are appropriate indices, and the crosshairs will provide a spatial reference in each plane relative to the other two.

### 3.3. A Note on Axes and Orientation

The NIfTI format contains an implicit generalized spatial transformation from the data coordinate system (i, j, k) into a real-space "right-handed" co-ordinate system. In this real-space system, the (x, y, z) axes are usually set such that x increases from left to right, y increases from posterior to anterior and z increases from inferior to superior.

At this point in time the **oro.nifti** package cannot apply an arbitrary transform to the imaging data into (x, y, z) space – such a transform may require non-integral indices and interpolation

steps. The package does accommodate straightforward transformations of imaging data; e.g., setting the i-axis to increase from right to left (the neurological convention). Future versions of **oro.nifti** will attempt to address more complicated spatial transformations and provide functionality to display the (x, y, z) axes on orthographic plots.

### 3.4. NIfTI and ANALYZE Data in \$4

A major improvement in the **oro.nifti** package is the fact that standard medical imaging formats are stored in unique classes under the \$4 system (Chambers 2008). Essentially, NIfTI and ANALYZE data are stored as multi-dimensional arrays with extra slots created that capture the format-specific header information; e.g., for a nifti object

### R> slotNames(mniRL)

| [1]  | ".Data"       | "trail"       | "extensions"  | "sizeof_hdr"     |
|------|---------------|---------------|---------------|------------------|
| [5]  | "data_type"   | "db_name"     | "extents"     | "session_error"  |
| [9]  | "regular"     | "dim_info"    | "dim_"        | "intent_p1"      |
| [13] | "intent_p2"   | "intent_p3"   | "intent_code" | "datatype"       |
| [17] | "bitpix"      | "slice_start" | "pixdim"      | "vox_offset"     |
| [21] | "scl_slope"   | "scl_inter"   | "slice_end"   | "slice_code"     |
| [25] | "xyzt_units"  | "cal_max"     | "cal_min"     | "slice_duration" |
| [29] | "toffset"     | "glmax"       | "glmin"       | "descrip"        |
| [33] | "aux_file"    | "qform_code"  | "sform_code"  | "quatern_b"      |
| [37] | "quatern_c"   | "quatern_d"   | "qoffset_x"   | "qoffset_y"      |
| [41] | "qoffset_z"   | "srow_x"      | "srow_y"      | "srow_z"         |
| [45] | "intent_name" | "magic"       | "extender"    | "reoriented"     |

R> c(mniRL@cal\_min, mniRL@cal\_max)

[1] 0 255

R> range(mniRL)

[1] 0 255

R> mniRL@datatype

[1] 2

R> convert.datatype(mniRL@datatype)

[1] "UINT8"

Note, an ANALYZE object has a slightly different set of slots. Slots 4–47 are taken verbatim from the definition of the NIfTI format and are read directly from a file. The slot .Data is the multidimensional array (since class nifti inherits from class array) and the slots trail,

extensions and reoriented are used for internal bookkeeping. In the code above we have accessed the min/max values of the imaging data using the "cal\_min" and "cal\_max" slots and matches a direct interrogation of the .Data slot using the range function. Looking at the datatype slot provides a numeric code that may be converted into a value that indicates the type of byte structure used (in this case an 8-bit or 1-byte unsigned integer).

As introduced in Section 3.1 there are currently only two accessor functions to slots in the NIfTI header (aux.file and descrip) – all other slots are either ignored or used inside of functions that operate on ANALYZE/NIfTI objects. The NIfTI class also has the ability to read and write extensions that conform to the NIfTI data format. Customized printing and validity-checking functions are available to the user and every attempt has been made to ensure that the information from the multi-dimensional array is in agreement with the header values.

The constructor function nifti produces valid NIfTI objects, including a consistent header, from an arbitrary array.

```
R> n <- 100
R> (random.image <- nifti(array(runif(n * n), c(n, n, 1))))</pre>
NIfTI-1 format
                   : niftiAuditTrail
  Туре
                   : 2 (UINT8)
  Data Type
                  : 8
  Bits per Pixel
  Slice Code
                   : 0 (Unknown)
                   : 0 (None)
  Intent Code
                   : 0 (Unknown)
  Qform Code
  Sform Code
                   : 0 (Unknown)
  Dimension
                   : 100 x 100 x 1
  Pixel Dimension : 1 x 1 x 1
  Voxel Units
                   : Unknown
  Time Units
                   : Unknown
R> random.image@dim_
[1]
      3 100 100
                   1
                           1
                       1
                               1
                                    1
R> dim(random.image)
[1] 100 100
               1
```

Data types used for NIfTI formats can be achieved from the convert.datatype function.

```
R> cbind(convert.datatype())
```

```
[,1]
UINT8 2
INT16 4
```

```
INT32
            8
FLOAT32
            16
COMPLEX64
            32
FLOAT64
            64
RGB24
            128
INT8
            256
UINT16
            512
UINT32
            768
INT64
            1024
UINT64
            1280
FLOAT128
            1536
COMPLEX128 1792
COMPLEX256 2048
RGBA32
            2304
```

The function writeNIfTI outputs valid NIfTI class files, which can be opened in other medical imaging software. Files can either be stored as standard .nii files or compressed with gnuzip (default).

```
R> writeNIfTI(random.image, "random")
R> list.files(pattern = "random")
[1] "random.nii.gz"
```

### 3.5. Audit Trail

Following on from the S4 implementation of both the NIfTI and ANALYZE data formats, the ability to extend the NIfTI data format header is utilized in the **oro.nifti** package. First, extensions are properly handled when reading and writing NIfTI data. Second, users are allowed to add extensions to newly-created NIfTI objects by casting them as niftiExtension objects and adding niftiExtensionSection objects to the extensions slot. Third, by default all operations that are performed on a NIfTI object will generate what we call an *audit trail* that consists of an XML-based log. Each log entry contains information not only about the function applied to the NIfTI object, but also various system-level information; e.g., version of R, user name, date, time, etc. When writing NIfTI-class objects to disk, the XML-based NIfTI extension is converted into plain text and saved using ecode = 6 to denote plain ASCII text only. The user may control the tracking of data manipulation via the audit trail using a global option. For example, please use the command

```
R> options(niftiAuditTrail = FALSE)
```

to turn off the "audit trail" option in **oro.nifti**. Table 2 displays output from the accessor function audit.trail(mnilR), the XML-based audit trail that is stored as a NIfTI header extension.

# Table 2: XML-based audit trail obtained via audit.trail(mniLR).

```
<filename>/home/brandon/R/x86_64-unknown-linux-gnu-library/2.12/oro.nifti/nifti/mniLR.nii.gz</filename>
                                                                                                                                                                                                     <workingDirectory>/home/brandon/Software/rigorous/papers/jss_dicom_nifti</workingDirectory>
                                                                                                                                                                                                                                                                                                                                                                                                             <r-version.version.string>R version 2.12.1 (2010-12-16)</r-version.version.string>
                                                                                             <audit-trail xmlns="http://www.dcemri.org/namespaces/audit-trail/1.0">
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         <package-version.Version>0.2.5</package-version.Version>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   <date>Tue Mar 01 02:30:55 PM 2011 GMT</date>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      <user.LOGNAME>brandon</user.LOGNAME>
                                                                                                                                                                                                                                                                                                           <call>readNIfTI(fname = fname)</call>
R> audit.trail(mniLR)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    </audit-trail>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              </system>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 </created>
                                                                                                                                                                                                                                                                                                                                                                <system>
                                                                                                                                                     <created>
```

### 3.6. Interactive visualization

Basic visualization of nifti and anlz class images can be achieved with any visualization for arrays in R. For example, the **EBImage** package provides functions display and animate for visualization (Sklyar *et al.* 2010). Please note that functions in **EBImage** expect greyscale values in the range [0, 1], hence the display of nifti data may be performed using

```
R> mniLR.range <- range(mniLR)
R> display((mniLR - min(mniLR))/diff(mniLR.range))
```

Interactive visualization of multi-dimensional arrays, stored in NIfTI or ANALYZE format, is however best performed outside of R at this point in time. Popular viewers, especially for neuroimaging data, are

- FSLView (http://www.fmrib.ox.ac.uk/fsl/fslview/),
- MRIcron (http://cabiatl.com/mricron/).

The **mritc** package provides basic interactive visualization of ANALYZE/NIfTI data using a Tcl/Tk interface (Feng and Tierney 2010).

### 3.7. Functional MRI Example

Simple Time-Series or Multi-Volume Image

This is an example of reading in, and displaying, a four-dimensional medical imaging data set in NIfTI format filtered\_func\_data obtained from the FSL evaluation and example data suite (http://www.fmrib.ox.ac.uk/fsl/fseds.html). Successful execution of the commands

```
R> (ffd <- readNIfTI("filtered_func_data"))</pre>
```

```
NIfTI-1 format
```

Type : niftiAuditTrail
Data Type : 16 (FLOAT32)

Bits per Pixel : 32

Slice Code : 0 (Unknown)
Intent Code : 0 (None)
Qform Code : 0 (Unknown)
Sform Code : 0 (Unknown)

Dimension :  $64 \times 64 \times 21 \times 180$ 

Pixel Dimension: 4 x 4 x 6 x 3

Voxel Units : mm Time Units : sec

R> image(ffd)

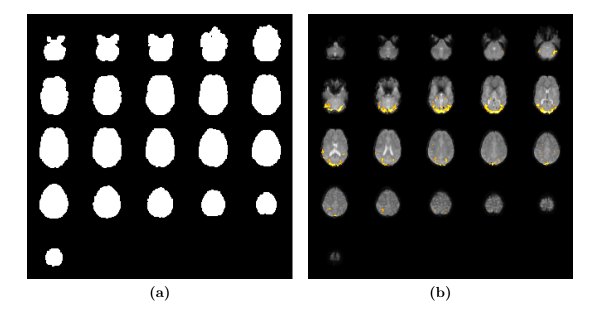


Figure 6: (a) Axial slices of the functional MRI "volume" filtered\_func\_data from the first acquisition. (b) Axial slices of the functional MRI data with the statistical image overlayed. The test statistics were thresholded at  $|Z| \geq 5$  for all voxel.

produces a four-dimensional (4D) array of imaging data that may be displayed in a  $5 \times 5$  grid of images (Figure 6a). The first three dimensions are spatial locations of the voxel (volume element) and the fourth dimension is time for this functional MRI (fMRI) acquisition. As seen from the summary of object, there are 21 axial slices of fairly coarse resolution ( $4 \times 4 \times 6$  mm) and reasonable temporal resolution (3 s). Figure 8 depicts the orthographic display of the filtered\_func\_data using the axial plane containing the left-and-right thalamus to approximately center the crosshair vertically.

R > orthographic(ffd, xyz = c(34, 29, 10))

### Statistical analysis

The R programming environment provides a wide variety of statistical methodology for the quantitative analysis of medical imaging data. For example, function MRI (fMRI) data are typically analysed by applying a multiple linear regression model, commonly referred to in the literature as a general linear model (GLM), that utilizes the stimulus experiment to construct the design matrix. Estimation of the regression coefficients in the GLM produces a statistical image; e.g., Z-statistics for a voxel-wise hypothesis test on activation in fMRI experiments (Friston et al. 1994, 1995).

The 4D volume of imaging data in filtered\_func\_data was acquired in an experiment with a TR=3 s, using both visual and auditory stimuli. The visual stimulus was a on/off with a duration of 60 seconds and the auditory stimulus was on/off with a duration of 90 seconds. We use a parametric haemodynamic respone function (HRF), with mean 6 and standard deviation 3, similar to the default values in FSL. Figure 8 depicts the visual and auditory stimuli convolved with the HRF.

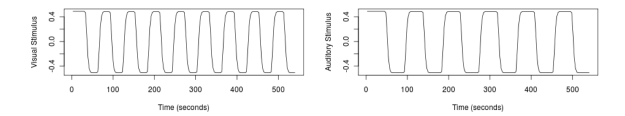


Figure 7: Visual (30 seconds on/off) and auditory (45 seconds on/off) stimulus convolved with a Gamma haemodynamic response function, used in the GLM fMRI analysis.

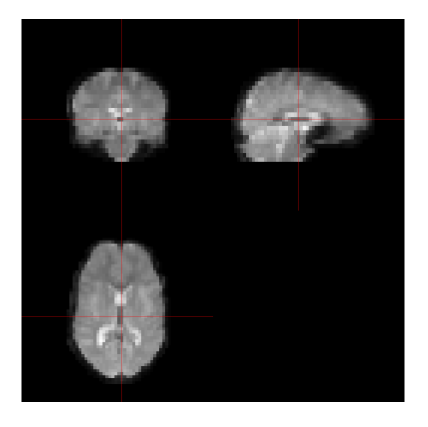


Figure 8: Orthographic display of the first volume from the functional MRI dataset filtered\_func\_data.

```
R > visual <- rep(c(rep(-0.5, 30), rep(0.5, 30)), 12)
R> auditory <- rep(c(rep(-0.5, 45), rep(0.5, 45)), 12)
R > hrf <- c(dgamma(1:15, 4, scale = 1.5))
R> visual.hrf <- rep(NA, 540)
R> for (k in 60:599) {
      visual.hrf[k - 59] \leftarrow sum(visual[k - (1:15)] * hrf)
+ }
R> auditory.hrf <- rep(NA, 540)
R> for (k in 90:629) {
      auditory.hrf[k - 89] \leftarrow sum(auditory[k - (1:15)] * hrf)
+ }
R> visual.hrf <- c(rep(0.5, 30), visual.hrf)
R> auditory.hrf <- c(rep(0.5, 45), auditory.hrf)
R > index <- seq(3, 540, by = 3)
R> visual.hrf <- visual.hrf[index]</pre>
R> auditory.hrf <- auditory.hrf[index]</pre>
R> visual.hrf <- visual.hrf - mean(visual.hrf)</pre>
R> auditory.hrf <- auditory.hrf - mean(auditory.hrf)</pre>
```

The design matrix is than used in a voxel-wise GLM using the the standard 1m function in R for parameter estimation in linear regression. From this, for each voxel, t-statistics and associasted p-values for a hypothesis test on no effect of each stimulus along with an F-statistic for a hypothesis test on no effect of any stimuli can be computed.

For anatomical context, statistical images are displayed as an overlay on top of a reference image. Successful execution of the command allows one to display the statistical image (of voxel-wise activations) overlayed on the original EPI (echo planar imaging) data taken from the functional MRI experiment.

```
R> overlay(ffd, ifelse(abs(Z.visual) > 5, Z.visual, NA), zlim.x = range(ffd),
+ zlim.y = range(Z.visual, na.rm = TRUE))

R> overlay(ffd, ifelse(abs(Z.auditory) > 5, Z.auditory, NA),
+ zlim.x = range(ffd), zlim.y = range(Z.auditory, na.rm = TRUE))
```

The 4D array of parameter estimates are overlayed on the original data for anatomical reference in Figure 6b. The function overlay extends the capabilities of displaying "images" by allowing one to add a statistical image to a structural image of the same dimension.

## 4. Converting DICOM to NIfTI

The **oro.dicom** and **oro.nifti** packages have been specifically designed to use as much information as possible from the metadata-rich DICOM format and use that information in the construction of the NIfTI data volume. The function dicom2nifti converts a list of DICOM images into an nifti object, and likewise dicom2analyze converts them into an anlz object. Historically, data conversion from DICOM to NIfTI (or ANALYZE) has been provided outside of R using one of several standalone software packages:

- Xmedcon (Nolf 2003),
- FreeSurfer (http://surfer.nmr.mgh.harvard.edu),
- MRIConvert (http://lnci.oregon.edu/~jolinda/MRIConvert).

This is by no means an exhaustive list of software packages available for DICOM conversion. In addition there are several other R packages with the ability to process DICOM data

- fmri (Polzehl and Tabelow 2007),
- tractor.base (Clayden 2010) (part of the tractor project http://code.google.com/p/tractor).

### 4.1. Single-Series Example

Using the 40 images from the hk40 object (previously defined in Section 2.1) it is straightforward to perform DICOM-to-NIfTI conversion using only default settings and plot the results in either lightbox or orthographic displays.

```
R> dput(formals(dicom2nifti))
list(dcm = , datatype = 4, units = c("mm", "sec"), rescale = FALSE, reslice = TRUE, DIM =
R> (hk40n <- dicom2nifti(hk40))</pre>
```

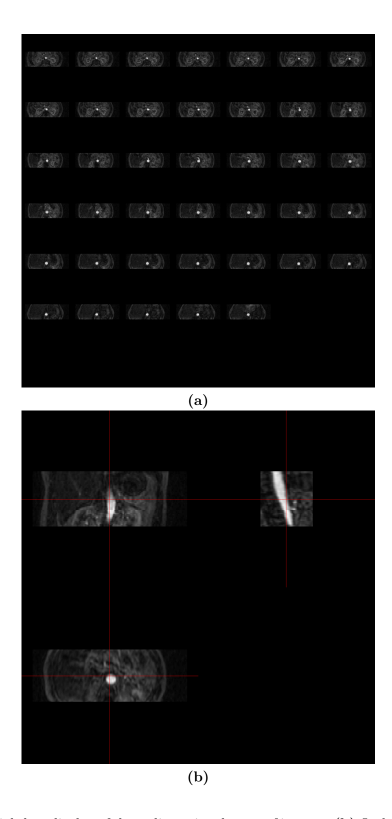


Figure 9: (a) Lightbox display of three-dimensional array of images. (b) Orthographic display of the same three-dimensional array (using the default settings for orthographic).

NIfTI-1 format

Type : niftiAuditTrail

Data Type : 4 (INT16)

Bits per Pixel : 16

Slice Code : 0 (Unknown)
Intent Code : 0 (None)
Qform Code : 0 (Unknown)
Sform Code : 0 (Unknown)
Dimension : 256 x 256 x 40
Pixel Dimension : 1.56 x 1.56 x 3.12

Voxel Units : mm Time Units : sec

R> image(hk40n)

R> orthographic(hk40n)

At default dicom2nifti takes all image data from the DICOM list and creates a 3D image. Four-dimensional image volumes (three in space plus one in time) are also converted automatically by specifying DIM=4, where slice positions are taken from the ImagePositionPatient DICOM header field. For example, using DIM=4 on the hk40 DICOM data,

R> (hk40n <- dicom2nifti(hk40, DIM = 4))

NIfTI-1 format

Type : niftiAuditTrail

Data Type : 4 (INT16)

Bits per Pixel : 16

Slice Code : 0 (Unknown)
Intent Code : 0 (None)
Qform Code : 0 (Unknown)
Sform Code : 0 (Unknown)
Dimension : 256 x 256 x 40
Pixel Dimension : 1.56 x 1.56 x 3.12

Voxel Units : mm Time Units : sec

will also produce a three-dimensional volume of images, since the ImagePositionPatient field is unique for each single slice of the volume.

The functions dicom2nifti and dicom2analyze will fail when the dimensions of the individual images in the DICOM list do not match. However, they do not check for different series numbers or patient IDs so caution should be exercised when scripting automated workflows for DICOM-to-NIfTI conversion. In cases where a DICOM file includes images from more than one series, the corresponding slices have to be chosen before conversion, using dicomTable, extractHeader, and matchHeader.

### 4.2. Multiple-Series Example

The National Biomedical Imaging Archive (NBIA) is a searchable, national repository integrating *in vivo* cancer images with clinical and genomic data. The NBIA provides the scientific community with public access to DICOM images, image markup, annotations, and rich metadata.<sup>1</sup> The multiple MRI sequences processed here were downloaded from the "RIDER Neuro MRI" collection.<sup>2</sup> A small for loop has been written to operate on a subset of the DICOM directory structure, where the SeriesInstanceUID DICOM header field is assumed to be 100% accurate in series differentiation.

```
R> subject <- "1086100996"
R> DCM <- dicomSeparate(subject, verbose = TRUE, counter = 100)
   564 files to be processed!
   100 files processed...
   200 files processed...
   300 files processed...
   400 files processed...
   500 files processed...
R> seriesInstanceUID <- extractHeader(DCM$hdr, "SeriesInstanceUID",
R> for (uid in unique(seriesInstanceUID)) {
      index <- which(unlist(lapply(DCM$hdr, function(x) uid %in%</pre>
          x$value)))
      uid.dcm <- list(hdr = DCM$hdr[index], img = DCM$img[index])</pre>
      patientsName <- extractHeader(uid.dcm$hdr, "PatientsName",</pre>
          FALSE)
      studyDate <- extractHeader(uid.dcm$hdr, "StudyDate",</pre>
          FALSE)
      seriesDescription <- extractHeader(uid.dcm$hdr, "SeriesDescription",
      fname <- paste(gsub("[^0-9A-Za-z]", "", unique(c(patientsName,
          studyDate, seriesDescription))), collapse = "_")
      cat("## ", fname, fill = TRUE)
      if (gsub("[^0-9A-Za-z]", "", unique(seriesDescription)) ==
          "axtensor") {
          D < -4
          reslice <- FALSE
      }
      else {
          D <- 3
          reslice <- TRUE
      uid.nifti <- dicom2nifti(uid.dcm, DIM = D, reslice = reslice,</pre>
```

http://cabig.nci.nih.gov/tools/NCIA

<sup>&</sup>lt;sup>2</sup>http://wiki.nci.nih.gov/display/CIP/RIDER

```
descrip = c("PatientID", "SeriesDescription"))
      writeNIfTI(uid.nifti, fname)
+ }
##
     281949_19040720_axtensor
##
     281949_19040720_ax30flip
##
     281949_19040720_ax15flip
##
     281949_19040720_ax25flip
##
     281949_19040720_ax20flip
##
     281949_19040720_ax10flip
##
     281949_19040720_ax5flip
```

Note, the diffusion tensor imaging (DTI) data axtensor is assumed to be four dimensional and all other series (the multiple flip-angle acquisitions) are assumed to be three dimensional. There is always a balance between what information should be pre-specified versus what can easily be extracted from the DICOM headers or images.

### 5. Conclusion

Medical image analysis depends on the efficient manipulation and conversion of DICOM data. The **oro.dicom** and **oro.nifti** packages have been developed to provide the user with a set of functions that mask as many of the background details as possible while still providing flexible and robust performance.

The future of medical image analysis in R will benefit from a unified view of the imaging data standards: DICOM, NIfTI and ANALYZE. The existence of a single package for handling imaging data formats would facilitate interoperability between the ever increasing number of R packages devoted to medical image analysis. We do not assume that the data structures in **oro.dicom** or **oro.nifti** are best-suited for this purpose and we welcome an open discussion around how best to provide this standardization to the end user.

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