

Reading HDF5

Jeffrey Leek Johns Hopkins Bloomberg School of Public Health

HDF5

- Used for storing large data sets
- · Supports storing a range of data types
- · Heirarchical data format
- · groups containing zero or more data sets and metadata
 - Have a *group header* with group name and list of attributes
 - Have a group symbol table with a list of objects in group
- · datasets multidmensional array of data elements with metadata
 - Have a *header* with name, datatype, dataspace, and storage layout
 - Have a data array with the data

http://www.hdfgroup.org/

R HDF5 package

```
source("http://bioconductor.org/biocLite.R")
biocLite("rhdf5")
```

```
library(rhdf5)
created = h5createFile("example.h5")
created
```

```
[1] TRUE
```

- This will install packages from Bioconductor http://bioconductor.org/, primarily used for genomics but also has good "big data" packages
- · Can be used to interface with hdf5 data sets.
- This lecture is modeled very closely on the rhdf5 tutorial that can be found here http://www.bioconductor.org/packages/release/bioc/vignettes/rhdf5/inst/doc/rhdf5.pdf

Create groups

```
created = h5createGroup("example.h5","foo")
created = h5createGroup("example.h5","baa")
created = h5createGroup("example.h5","foo/foobaa")
h5ls("example.h5")
```

```
group name otype dclass dim

0  / baa H5I_GROUP

1  / foo H5I_GROUP

2  /foo foobaa H5I_GROUP
```

Write to groups

```
A = matrix(1:10,nr=5,nc=2)
h5write(A, "example.h5","foo/A")
B = array(seq(0.1,2.0,by=0.1),dim=c(5,2,2))
attr(B, "scale") <- "liter"
h5write(B, "example.h5","foo/foobaa/B")
h5ls("example.h5")</pre>
```

```
group name otype dclass dim

0  / baa H5I_GROUP

1  / foo H5I_GROUP

2  /foo A H5I_DATASET INTEGER 5 x 2

3  /foo foobaa H5I_GROUP

4 /foo/foobaa B H5I_DATASET FLOAT 5 x 2 x 2
```

Write a data set

```
df = data.frame(1L:5L,seq(0,1,length.out=5),
    c("ab","cde","fghi","a","s"), stringsAsFactors=FALSE)
h5write(df, "example.h5","df")
h5ls("example.h5")
```

```
group name otype dclass dim

0  / baa H5I_GROUP

1  / df H5I_DATASET COMPOUND 5

2  / foo H5I_GROUP

3  /foo A H5I_DATASET INTEGER 5 x 2

4  /foo foobaa H5I_GROUP

5 /foo/foobaa B H5I_DATASET FLOAT 5 x 2 x 2
```

Reading data

```
readA = h5read("example.h5","foo/A")
readB = h5read("example.h5","foo/foobaa/B")
readdf= h5read("example.h5","df")
readA
```

```
[,1] [,2]
[1,] 1 6
[2,] 2 7
[3,] 3 8
[4,] 4 9
[5,] 5 10
```

Writing and reading chunks

```
h5write(c(12,13,14), "example.h5", "foo/A", index=list(1:3,1))
h5read("example.h5", "foo/A")
```

```
[,1] [,2]
[1,] 12 6
[2,] 13 7
[3,] 14 8
[4,] 4 9
[5,] 5 10
```

Notes and further resources

- · hdf5 can be used to optimize reading/writing from disc in R
- · The rhdf5 tutorial:
 - http://www.bioconductor.org/packages/release/bioc/vignettes/rhdf5/inst/doc/rhdf5.pdf
- The HDF group has information on HDF5 in general http://www.hdfgroup.org/HDF5/