## PDB-Hadoop Manual

# Jamie Al-Nasir and Hugh Shanahan February 24, 2015

#### 1 Introduction

PDB-Hadoop is a framework that facilitates the parallel execution of protein structure analysis tools to be carried out on the entire (or large subsets of) the Protein Databank (PDB) using the Apache Hadoop platform. Apache Hadoop is a software platform that allows for the processing of large scale datasets using clusters consisting of commodity hardware and is highly scalable.

PDB-Hadoop is designed so that structural Biologists can use the Hadoop platform without having to write the relatively complex Java code that Hadoop is implemented for. This framework is easily scalable and uses a *mapper* architecture that functions stand-alone or can be extended to include further *mapreduce* operations. This does away with the necessity for one to implement ones own map-reduce applications or re-write existing code for the *map-reduce* formalism, although the way in which PDB-Hadoop is implemented ensures this approach is still available for users wishing to exploit data aggregation properties of the *map-reduce* method.

This manual is organised as follows. We will discuss the scripts developed here and where to download them. We then describe how to download PDB files from the FTP archive and how to uncompress them (users with their own mirror or set of PDB files may skip this section). We then describe how the PDB files of interest can be prepared and placed on the local distributed file systems for Hadoop (called HDFS). We show how to configure, test and run your own jobs to run on PDB-Hadoop and then show how to parse the outputs as required.

## 2 Scripts within this package and their function

PDB-Hadoop is comprised of a series of Bash Shell scripts, listed in table 1. They can be downloaded from the following GitHub repository

blah blah ZENODO DOI. BIT ON HOW TO DOWNLOAD FROM GITHUB.

## 3 Acquisition of the PDB (Protein databank)

As of December 5th 2014 there are 105,383 structures deposited in the protein databank. Downloading the entire or sections of the PDB is commonly achieved

pdb- $gunzip.sh$	Given a folder, recursively extracts ent.gz files.
pdb-to-line.sh	Given a pdb file, converts this into a single line of text (used
	by pdb-build.sh).
pdb-build.sh	Given a folder of pdb files (*.ent), builds a special single,
	large pdb file suitable for Hadoops hdfs.
pdb-hadoop.sh	Hadoop mapper, this is run within a Hadoop job and han-
	dles execution of the user program on the pdb. This file
	requires customisation to set the path of the user program
	and temporary folder.
$Log ext{-}to ext{-}pdb.sh$	Given a single concatenated PDB-Hadoop log file extracts
	individual PDB file output as separate files (these comple-
	ment the input PDB files and are of the same name).

Table 1: List of scripts in PDB-Hadoop and descriptions of what they do.

using the Unix command rsync (which synchronises a local folder with a subfolder of the ftp archive of the PDB (rcsb.org) and can be carried out with the following command:

/usr/bin/rsync --progress -rlpt -v -z --delete --port=873 rsync.ebi.ac.uk::pub/databases/pdb/data/structures/divided/pdb/PDB

This creates a local folder in the current directory named PDB and synchronises it with the entire PDB (in sub-divided folders). rsync should have the same path on any UNIX distribution but this may vary. There is however a well-written script by Thomas Solomon circulating in the public domain which is useful for this purpose and allows further customisation of what is downloaded/synchronised available at

https://github.com/dbolser/PDB-rsync/blob/master/rsyncPDB-v1.0.sh

## 4 Unzipping of the compressed .ent.gz PDB files

The files on the rcsb.org ftp archive are normally in compressed gzip format and require "unzipping" with *gunzip*. Assuming the files are in a folder entitled PDB, this can be achieved using the pre-written script that recursively processes the PDB folder as follows:

./pdb-gunzip.sh ./PDB

# 5 Conversion of a collection of PDB files into a single condensed file for Hadoop

Hadoop distributed file system (hdfs) is not optimised for a large number of small files that comprise the protein databank. This problem can be overcome by packaging the entirety of the PDB into a single file ( $\tilde{\ }$ 80gb for all of the PDB as January 2015). The script is pdb-to-line.sh which is automatically run by pdb-build.sh script. The user should run pdb-build.sh as below, providing the

directory of the PDB files of interest and a target file to build, i.e. PDB-text-full.txt

./pdb-build.sh <PDBfolder> ./PDB-text-full.txt

## 6 Configuration and Testing

Before running your own code on PDB-Hadoop there are some small configuration steps and (optional) testing.

#### 6.1 Configuration

In the first instance, we suggest setting the permissions of the PDB-Hadoop folder to be accessible to all users for read and execute permissions so that Hadoops' YARN (Yet Another Resource Negotiator) may be able to execute the script within the Hadoop job.

```
chmod -R a+rX ~/pdb-hadoop
```

In order configure the user program for parallel execution on Hadoop, one must edit the main PDB-Hadoop script *pdb-hadoop.sh* file to specify the path to the user program and if necessary a preferred temporary folder for the job. The following lines must be amended:

Specifically, the following PDB-Hadoop variables must/can be set prior to executing a job:

#### 1. \_LEGACY\_PROGRAM\_

[Required] to specify the path to the program to be execute in parallel fashion on Hadoop.

#### 2. \_TEMP\_FOLDER\_

[Optional] to specify the temporary folder (default "/tmp"), which must be writable to YARN (i.e. the Hadoop user).

#### 3. \_POST\_PROC\_PROGRAM\_

[Optional] to specify the path to the users post-processing program that takes output from the execution of user program (defined above) and performs textual processing on the results.

#### 4. \_MAX\_PDB\_SIZE\_

[Optional] If set PDB-Hadoop ignores processing of files greater than specified size (in bytes)

### 6.2 Testing

For testing purposes one can specify a linux command such as "wc -l" as the

```
_LEGACY_PROGRAM_
```

in the main pdb-hadoop.sh script.

In some cases it may be useful to limit the execution time of the user program and this can be achieved by prepending the user job with the Linux *timeout* command, for instance to specify a 4 minute time constraint for the user program /path/myprogram:

```
_LEGACY_PROGRAM_="timeout 4m /path/myprogram"
```

A common reason to do this is to ensure that the user program Job only runs for a shorter period than Hadoops' default container time limit of 5 minutes. If required a time constraint for the container can be set within Hadoop and the Linux timeout command can be prepended to the job path to ensure the timeout set in Hadoop is not exceeded by the user program Job. NB: A user program job that overruns the Hadoop timeout will result in the containiner being terminated by Hadoops' YARN (Yet Another Resource Negotiator). It is also possible to reconfigure Hadoop to have a longer timeout period - please consult with your local hadoop administrator.

## 7 Executing your job on PDB-Hadoop

To execute your job you must first copy the fuile generated in section 5 (generated by pdb-build.sh) to Hadoop's hdfs. This can be achieved as follows:

```
hadoop fs -put /user/hduser/ /local/path/pdb-text-full.txt
```

The job can then be executed by issuing the Hadoop command at the shell prompt:

```
hadoop jar /usr/lib/hadoop-mapreduce/hadoop-streaming.jar \
-Dmapred.sort.avoidance=0 \
-Dmapred.reduce.tasks=0 \
-D stream.non.zero.exit.status.is.failure=false \
-input /user/hduser/pdb-text-full.txt
-output /user/hduser/pdb-legacy-output \
-mapper "/path/to/local/pdb-hadoop/pdb-hadoop.sh" \
```

The specific paths may vary depending on the local configuration of your hadoop cluster.

# 8 Format of PDB-Hadoop job output - user program results

The pdb-hadoop job (executed by mapper: pdb-hadoop.sh) will execute the user program in a parallelised fashion on Hadoop. This is an example of the output:

```
Extracted/writing file /tmp/pdb3zz1.ent
pdb3zz1.ent-000000001 PDB Dihedral angle calculation program
pdb3zz1.ent-0000000002
                        By Jamie Al-Nasir
pdb3zz1.ent-0000000003 Royal Holloway dept. of Computer Science
pdb3zz1.ent-0000000004 http://jamie.al-nasir.com
pdb3zz1.ent-
pdb3zz1.ent-0000000005 Using PDB file: /tmp/pdb3zz1.ent
pdb3zz1.ent-
pdb3zz1.ent-0000000006 Computation for 712 residues(s) in the structure
pdb3zz1.ent-
pdb3zz1.ent-000000007 Phi
                                                Ch. Residue
                                Psi
                                        Omega
pdb3zz1.ent-0000000008 0.00
                                -103.12 168.38
                                                 Α
                                                    VAL1
pdb3zz1.ent-0000000009
                        -123.59 147.71
                                        -178.71
                                                    VAL2
                        -66.19 143.08
pdb3zz1.ent-0000000010
                                        -179.97
                                                    PRO3
pdb3zz1.ent-0000000011
                       -67.74
                               146.86
                                       176.65
                                                    PRO4
pdb3zz1.ent-0000000012
                        -64.85
                               151.53
                                        -179.54
                                                 Α
                                                    ALA5
pdb3zz1.ent-0000000013 80.26
                                        176.39
                                                    GLY6
                                11.45
pdb3zz1.ent-000000014
                        -119.22 158.85
                                        179.83
                                                    THR7
pdb3zz1.ent-000000015
                        -51.76
                               -44.05
                                                    PR08
                                        178.34
                                                 Α
pdb3zz1.ent-0000000016
                       -68.62
                                -34.23
                                        179.35
                                                 Α
                                                    TRP9
pdb3zz1.ent-0000000017
                        -63.71
                                -45.73
                                        178.06
                                                    GLY10
pdb3zz1.ent-000000018
                        -68.99
                                -38.64
                                        175.79
                                                    THR11
                                                 Α
pdb3zz1.ent-0000000019
                        -61.31
                                -38.36
                                        177.48
                                                 Α
                                                    ALA12
Extracted/writing file /tmp/pdb3zzs.ent
pdb3zzs.ent-0000000001
                       PDB Dihedral angle calculation program
pdb3zzs.ent-0000000002
                       By Jamie Al-Nasir
pdb3zzs.ent-0000000003
                       Royal Holloway dept. of Computer Science
pdb3zzs.ent-0000000004 http://jamie.al-nasir.com
pdb3zzs.ent-
pdb3zzs.ent-000000005 Using PDB file: /tmp/pdb3zzs.ent
pdb3zzs.ent-
pdb3zzs.ent-000000006 Computation for 585 residues(s) in the structure
pdb3zzs.ent-
pdb3zzs.ent-0000000007
                       Phi
                                Psi
                                                Ch. Residue
                                        Omega
pdb3zzs.ent-0000000008
                       0.00
                                150.04
                                       176.62
                                                 Α
                                                    SER7
pdb3zzs.ent-0000000009
                        -60.85 157.36
                                        172.78
                                                    ASP8
pdb3zzs.ent-0000000010
                        -139.54 163.05
                                                    PHE9
                                        177.34
                                                 Α
pdb3zzs.ent-0000000011
                        -123.13 149.23
                                        177.06
                                                 Α
                                                    VAL10
pdb3zzs.ent-0000000012
                       -112.04 123.65
                                        -179.08
                                                 Α
                                                    VAL11
```

Each pdb file that is extracted from pdb-full.txt by the Hadoop mapper will have a line stating "Extracted/writing file /tmp/pdb3zz1.ent". followed by the results from the user program. Lines from the user program are prefixed with the original pdb filename and a line-number to facilitate sorting of the output by Hadoop. In this case the user program parallelised was dihedrals-64 a Linux command-line tool for computation of dihedral/torsional angles in peptide molecules.

## 9 Extracting user program results

Hadoop stores the output of the job on hdfs as a collection of text files of size dependent on the Hadoop cluster configuration. It is often most convenient to extract these concatenated as a single text file saved to the local file system. This can be achieved using Hadoops *getmerge* command as follows:

hadoop fs -getmerge /users/job-folder/ PDB-Hadoop-log.txt

## 10 Post-processing of user program results

PDB-Hadoop facilitates the processing of output data from the user program prior to deposition on HDFS. For example the initial job may create an extensive log file which then needs to be parsed to extract a subset of the results. This allows the user to extract and distil information from the user job within the parallelised map process that is running on Hadoop. As noted previously, this can be done by setting

\_POST\_PROC\_PROGRAM\_

to the path of the users post-processing program.

# 11 Extracting individual files from the PDB-Hadoop generated concatenated log file

As previously explained the Hadoops getmerge command can be used to obtain a single concatenated log file. In some cases it is required to extract the output of a PDB-Hadoop job as individual files. For instance some legacy programs merely output a report for the input PDB file, while others output a new PDB file. Extraction of the newly generated PDB files (or user program reports) is achieved with the PDB-Hadoop log-to-pdb.sh script. This takes the single concatenated log-file input and splits this into individual PDB files/reports. In each case this output can be captured in individual files using the PDB-Hadoop log-to-pdb.sh script as follows:

./log-to-pdb.sh ./PDB-Hadoop-log.txt