1 Input file 1

Network extraction code - pnextract

pnextract extracts a conventional pore network from a microCT image. The algorithm is a rewrite of the Dong and Blunt (2009) code. There are major differences though. First, the pore and throat detection algorithm is revised; see Stages 1 and 2 described in Raeini et al. (2017) https://doi.org/10.1103/PhysRevE.96.013312. Raeini et al. (2017) is an extension of this code. The shape of pores in this code are deduced from shape factors, the shape-factor equation is changed compared to the old definition, see Bultreys et al (2018, currently under-review).

1 Input file

The input file for the network extraction code is a mhd header file compatible with Paraview and Fiji (ImageJ with plugins) with additional optional keywords specific to network extraction algorithm. See the file Image.mhd for a sample input.

1.1 Format specifications:

- 1. The order of the first 6 keywords should not be changed for compatibility with third-party software (ImageJ and Paraview)
- 2. Use "#" or "//" for adding comments
- 3. All keyword and its data should be given in a single line

Important keywords:

1 to 3rd keywords (should not be changed):

- 1. ObjectType = Image
- 2. NDims = 3
- 3. ElementType = MET_UCHAR
- keyword: DimSize used to assign the dimensions of the image: Nx, Ny and Nz
- 5. keyword: ElementSize used for assigning voxel size: δx , δy and δz should be equal
- 6. keyword: ElementDataFile specifies the name of binary 8bit data file (.raw), ascii (.dat), .raw.gz, and .tif files are supported too.

1 Input file 2

```
ObjectType = Image
NDims = 3
ElementType = MET_UCHAR

DimSize = 400 400 400
ElementSize = 5.345 5.345 5.345
Offset = 0 0 0
ElementDataFile = Berea.raw
```

Fig. 1: Sample input header file

Medial-surface settings:

The medialSurfaceSettings is an optional technical keyword which can be used for sensitivity analysis, for instance.

```
medialSurfaceSettings 0.1 0.9 0.7 0.5 1.5 1.21 7 0.25 1.6;
```

where the keyword arguments are clipROutx clipROutyz midRFrac RMedSurfNoise lenNf vmvRadRelNf nRSmoothing RCorsf RCors, respectively.

The pnextract code produces few lines showing the settings being used. something like:

The first line is the keyword and its parameters and the rest are short names for each of the parameters and their values. In case you want to do a quick evaluation, you can copy the first line into the pnextract input, the .mhd file, and change the parameters and re-run the code. Here is a short explanation for these parameters:

clipROutx is used to limit the size of maximal-spheres extending outside the rock image in the x direction.

- clipROutyz is used to limit the size of maximal-spheres extending outside the rock image in the y and z directions.
- midRFrac is the relative size of the distance-map of the voxel between two maximal-spheres, for the spheres to be considered part of the same pore.
- RMedSurfNoise is a measure of noise amplitude. Decreasing this will likely increase the number of pores, but it also affects the number of corners per throat.
- lenNf is a relative distance for merging adjacent pores which are too close to each other.
- vmvRadRelNf is the relative size of the throat between the two pore considered for merging, the contraction should be less than this to merge the nearby pores (that are less than lenNf apart), otherwise the pore will not be merged. Decreasing these two will increase the number of pores.
- nRSmoothing applies a small amount of Gaussian-like smoothing on the computed distance map, which in turn affect the rest of the computations. Decreasing this will probably increases the number of pores.
- RCorsf controls the distance between the maximal spheres. This is a sensitive parameter, changing it may need changing other parameters to get good results.
- RCors controls the minimum distance between (small) maximal-spheres. This is a sensitive parameter, changing it may need changing other parameters to get good results.

2 Visualization and optional outputs data

Additional data generated during network extraction can be saved for visualization or further analysis by providing a set of keywords starting with "write_" network extraction code can write additional data The "write_all true" keyword is a short-cut for (and with a higher priority over) the following keywords:

```
write_radius:
                    true; // write distance maps
write_elements:
                    true; // writes pore labels in a _VElems.raw image
    file
write_poreMaxBalls: true; // writes pore maximal balls mapped to image
    (.raw...) format
write_throatMaxBalls: true; // writes throat maximal balls mapped to
    image format
write_throats:
                    true; // writes _throat.vtu file for visualization
    in Paraview -- obsolete
write_hierarchy:
                   true; // writes medial-surface branches in .vtu
write_medialSurface: true; // writes medial-surface branches (partially
    joined to form surfaces) in .vtu format
```

3 The Xmf network format (pnextract version 2021+)

If you are using the public domain version of pnflow please ignore this section. In the pnextract versions 2020+, the classical network data are written in Xmdf format, essentially in the form of a single ASCII (text) xml file which can be opened in any light-weight text editor, or visualized using VTK/Paraview. The file suffix is Net.xmf. This format similar to the format used in gnflow and pnflow versions 2020+ quasi-static (capillary-dominated) two-phase flow codes.

To use the Xmf format the pnflow code, you should assign it to the networkFile: PATH/TO/ROCK/ImageNet.xmf in the pnflow input file. The keyword NETWORK is reserved for reading network files in old (Oren/Statoil) format described in the next section.

4 The Structure of Network Data Files Statoil format

This Section is taken from PhD thesis of Taha Sochi (2007), Appendix I.

The network data are stored in four ASCII files. The format of these files is that of Statoil. The physical data are given in SI unit system.

4.1 Throat Data

The data for the throats are read from the link files. The structure of the link files is as follows:

*_link1.dat file

The first line of the file contains a single entry that is the total number of throats, say N , followed by N data lines. Each of these lines contains six data entries in the following order:

- 1. Throat index
- 2. Pore 1 index
- 3. Pore 2 index
- 4. Throat radius

- 5. Throat shape factor
- 6. Throat total length (pore center to pore center)

Fig. 2: Example of *_link1.dat file

*_link2.dat file

For a network with N throats, the file contains N data lines. Each line has eight data entries in the following order:

- 1. Throat index
- 2. Pore 1 index
- 3. Pore 2 index
- 4. Length of pore 1
- 5. Length of pore 2
- 6. Length of throat
- 7. Throat volume
- 8. Throat clay volume

```
22714 10452 10533 0.178E-04 0.120E-03 0.239E-04 0.218E-13 0.137E-14 22715 10452 10612 0.121E-04 0.747E-04 0.100E-04 0.266E-13 0.355E-14 22716 10453 10534 0.100E-04 0.270E-04 0.139E-04 0.543E-13 0.863E-14 ...
```

Fig. 3: Example of *_link2.dat file

4.2 Pore Data

The data for the pores are read from the node files. The structure of the node files is as follows:

*_node1.dat file

The first line of the file contains four entries: the total number of pores, the length (x-direction), width (y-direction) and height (z-direction) of the network. For a network with M pores, the first line is followed by M data lines each containing the following data entries:

- 1. Pore index
- 2. Pore x-coordinate
- 3. Pore y-coordinate
- 4. Pore z-coordinate
- 5. Pore connection number
- 6. For a pore with a connection number i there are 2(i + 1) entries as follows:
 - (a) The first i entries are the connecting pores indices
 - (b) The (i + 1)st entry is the pore inlet status (0 for false and 1 for true)
 - (c) The (i + 2)nd entry is the pore outlet status (0 for false and 1 for true)
 - (d) The last i entries are the connecting throats indices

Note: the inlet/outlet pores are those pores which are connected to a throat whose other pore is the inlet/outlet reservoir, i.e. the other pore has an index of -1/0. So if the (i+1)st entry is 1, one of the connecting pores indices is -1, and if the (i+2)nd entry is 1, one of the connecting pores indices is 0.

```
12349  0.3000E-02 0.300E-02 0.300E-02

1  0.350E-03 0.000E+00 0.700E-04 3 796 674 2 0 0 522 523 524

2  0.450E-03 0.500E-04 0.000E+00 3 359 31 1 0 0 525 526 524

3  0.880E-03 0.100E-04 0.000E+00 1 392 0 0 527

...
```

Fig. 4: Example of *_node1.dat file

* node2.dat file

For a network with M pores, the file contains M data lines. Each line has five data entries in the following order:

- 1. Pore index
- 2. Pore volume
- 3. Pore radius
- 4. Pore shape factor
- 5. Pore clay volume

```
50 0.3733E-13 0.1957E-04 0.3369E-01 0.7846E-16
51 0.1555E-14 0.8215E-05 0.3262E-01 0.4717E-16
52 0.1711E-13 0.1224E-04 0.3298E-01 0.1485E-15
...
```

Fig. 5: Example of $*_node2.dat$ file

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References:

Publications:

https://www.imperial.ac.uk/earth-science/research/research-groups/pore-scale-modelling/publications/

PhD theses:

https://www.imperial.ac.uk/earth-science/research/research-groups/pore-scale-modelling/phd-theses/