

# pyPolyBuilder user's guide and tutorials

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## Abstract

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# 1 General concepts of the software

PyPolyBuilder is a building block (BB) based automated topology builder. Hence, based on a few BB molecular topology files (MTF), it is able to build a larger and more complex MTF in addition to a suggestion of the geometry, as input for another molecular dynamics (MD) software. Currently the code supports gromacs and gromos format.

PyPolyBuilder has two modules of application. The dendrimer module and the network one .

## 1.1 Dendrimer module

PyPolyBuilder has an optimized and easy-to-use module in order to generate dendrimers (star-like polymers). Due to its symmetry, the user is able to define the core, intermediary and terminal blocks only. The connectivity is pre-assumed according to the desired generations of the dendrimer. The software is prepared to make the linkage list, and generate the final MTF and a initial guess for the dendrimer geometry in vacuum.

### 1.1.1 Input files

As inputs to the dendrimer module, there is required the BBs topologies, the dendrimer generation, and a force field parameters list. In topologies files, there are six available fields to pass information that define the building block structure, similar to gromacs topology files (some example cases will be better discussed further in tutorial descriptions). They are: [ `moleculetype` ], [ `atoms` ], [ `bonds` ], [ `angles` ], [ `dihedrals` ] and [ `branches` ].

The field [ `moleculetype` ] is where the building block name should be written.

```
[ moleculetype ]
; Name nrexcl
C4 2
```

This field defines the name of the BB in pyPolyBuilder's object-oriented workflow. The second column, nrexcl, defines the number of excluded neighbors when used in gromacs. However, pyPolyBuilder does not actually use this value. It was maintained here for completeness and to conserve the similarities to the gromacs .itp files.

In the field [ atoms ], each atom should be listed with their index number, type, residue number, residue name, charge group number, coulombic charge and mass, in that order. The required values are:

```
[ atoms ]  
; nr type resnr resid atom cgnr charge mass total_charge  
1 CH2 1 CORE C 1 0.250 14.02700  
2 CH2 1 CORE C 2 0.250 14.02700  
3 NT 1 CORE N 1 -0.750 14.00670  
4 NT 1 CORE N 2 -0.750 14.00670
```

It's important to notice that, pyPolyBuilder has a feature that modifies this pattern of the inputed gromacs-like topology. Some common force fields (such as gromos), represent an aromatic center only by its improper torsionals. Hence, in order to correctly model aromatic molecules when using a force field from the gromos family, one should exclude interactions between 1-4 neighbors in aromatic center. Those interactions may easily be excluded from the final MTF generated by pyPolyBuilder by adding one column with the flag 'AR' after the mass column, as illustrated below:

```
[ atoms ]  
; nr type resnr resid atom cgnr charge mass  
1 C 1 AROM C 1 -0.1298 12.01100 AR  
2 H 1 AROM H 1 0.1298 1.00800 AR  
3 C 1 AROM C 1 -0.1298 12.01100 AR
```

```
4 H 1 AROM H 1 0.1298 1.00800 AR
```

The ‘AR’ flag should only be used for the atoms which are part of a aromatic center and for those that are directly bonded to aromatic centers.

The [ bonds ] section should have the information about which atoms are forming chemical bonds. We need to pass their index numbers and information about the force field parameters:

```
[ bonds ]
; ai aj funct c0 c1
  1 2 2 gb_21
  1 3 2 gb_21
  2 4 2 gb_21
```

Where  $a_i$  and  $a_j$  are the index number of atom i and j, respectively, involved in the chemical bond, and g\_21 is the force field parameter as the input of the gromacs package.

We strongly suggest that the bonds parameter are defined in each BB file. However, they can also be defined using the atom types in the file that parses the parameters list. This input will be discussed subsequently.

The last field that we need to provide for each BB is the [ branches ]. Here, the atoms are defined which are used to form a bond between two building blocks to propagate the monomer.

```
[ branches ]
; donor acceptor
  1 0
  0 4
```

Usually, we can define the donor as the atom that will be bound in the new BB, since it is donating a connection, and the acceptor as the atom that will be bound in the last BB, since it may accept a connection.

The parameters list file needs to have the force field information about bonds, angles and dihedrals that exist in the topology. Some cases may require that the stereochemistry conformation is specified. In these cases, the conformation field can be used. In the cases where the conformation field is not present, the final structure will have two of them. For example:

```
# BONDS
H N gb_2
H NT gb_2

# ANGLES
H NT CH2 ga_11
H NT H ga_10

# IMPROPERST
CH2 C O N gi_1
C N H CH2 gi_1

# DIHEDRALS
CH2 CH2 C O gd_40
CH2 CH2 C N gd_40

# CONFORMATION
CH2 C N CH2 trans
CH2 C N H cis
```

Notice that, by default, the parameters from this parameters list file will be directly copied to the MTF. By default, pyPolyBuilder uses an internal default set of interaction parameters for simplicity. The “# CONFORMATION” field sets if the dihedral should be cis or trans, and it selects the parameter to be used within this internal pseudo-force field. In some complex cases, it might be interesting to make pyPolyBuilder to use the actual force field parameters (it will be exemplified in some of the following tutorials). This is possible by using the –forcefield option with a path to the force field to be used (in gromacs format). All possible options are discussed in following sections.

### 1.1.2 Command line call and optional flags

To call the software, the following command must be run in the command line:

```
pyPolyBuilder --core=core.itp \
              --inter=inter.itp \
              --ter=ter.itp \
              --params=list_param.itp \
              --ngen=1 \
              --name=NAME \
              --output=output.itp \
              --gro=output.gro \
              --dendrimer
```

All the possible options will be discussed exhaustively below. There are some options which are specific for dendrimer module and may not be used if the dendrimer module is not being used. To make it clear, those options are:

- **--core or -c** should define a relative or absolute path for the BB that will be used as core.
- **--inter or -i** should define a relative or absolute path for the BB that will be used as intermediary.

- **--ter** or **-t** should define a relative or absolute path for the BB that will be used as terminal.
- **--ngen** or **-n** should be an integer defining the generation number of the dendrimer that will be built using pyPolyBuilder.
- **--dendrimer** sets the module that will be called by pyPolyBuilder.

Those are the specific options for the dendrimer module. A complete list with the definition and usage of each option will be provided in a further section.

## 1.2 Network module

Differently from dendrimer module, network module does not assume any symmetry for the molecule. Hence, all connections between BBs need to be defined. Nevertheless, this module is absolutely general and can be used to build any molecule.

Here, the valence or connectivity of the BBs are not specified. Because of that, using the [ **branches** ] field as in dendrimer module could lead to a cumbersome syntax. This issue was avoided by defining a new input file that will be referred to as connectivity file. This file may be considered as a topology of topologies. All valence and connectivity for each BB is defined in this file.

### 1.2.1 Input files

The BB files themselves are organized very similarly to what they are in dendrimer modules (see previous section). For instance, the fields [ **moleculetype** ], [ **atoms** ], [ **bonds** ], [ **angles** ], and [ **dihedrals** ] should be defined exactly the same. While the field [ **branches** ] is not necessary anymore since the connections will be defined in the connectivity file.

In respect to the connectivity file, it is organized as follows. There are two needed fields: “# [BUILDING BLOCKS]” and “# [ CONNECTS ]” as illustrated below:

```
#[ BUILDING BLOCKS ]
```

```
;BBn name
```

```
1 PNIPS
```

```
2 PNIP
```

```
3 PNIP
```

```
4 PNIP
```

```
5 PNIPE
```

```
#[ CONNECTS ]
```

```
;BBi BBj ai aj
```

```
1 2 2 1
```

```
2 3 2 1
```

```
3 4 2 1
```

```
4 5 2 1
```

“# [BUILDING BLOCKS]” field receives a list of all monomers that will be used. The names that should be used are the names of the topology file as defined in the [ moleculetype ] field within the BB file. For instance, the BBs should have a field where their name is defined (see previous section):

```
[ moleculetype ]
```

```
; Name nrexcl
```

```
PNIP 2
```

With the BBs defined, the next field defines how they are connected.

“# [CONNECTS]” field receives 4 values. The two first are the index number of the two BBs that are being connected (as defined into “# [BUILDING BLOCKS]” field) followed by the index number of the atoms within each BB that are being used to make the chemical bound. For instance, by defining the line #11 in the example connectivity file above (“1 2 2 1”), pyPolyBuilder connects the atom index 2 from the first BB to the atom index 1 from

the second BB.

### 1.2.2 Command line call and optional flags

With all the BBs, parameters list, and connectivity files ready, pyPolyBuilder should be called using a command line as follows:

```
pypolybuilder --bbs=bb_PNIP-start.itp,bb_PNIP.itp,bb_PNIP-end.itp \
              --in=connect-4.in \
              --params=list_param.itp \
              --name=PNIPAM \
              --output=PNIPAM.itp \
              --gro=PNIPAM.gro \
              --network
```

All possible options will be described in detail in the next section. For clarity, the specific options for network module will be quickly discussed here as well. These are:

- **--bbs or -bbs** should be a list with the path for all BBs separated by comma.
- **--in or -in** should define a path for the connectivity file.
- **--network** sets the module that will be called by pyPolybuilder.

## 1.3 Command line options

PyPolyBuilder help can be called to check a list of all the possible options to be used and a quick description for them by using the following command line:

```
pypolybuilder -h
```

A complete list can be consulted below:

- **--help or -h:**

Shows pyPolyBuilder help, a quick description of each option, as well as the name of the attribute settled by the option.

- **Mandatory dendrimer options**

- **--core or -c:** path

This option inputs a path to a file that will be used as the core block in dendrimer module.

- **--inter or -i:** path

This option inputs a path to a file that will be used as the intermediary block in dendrimer module.

- **--ter or -t:** path

This option inputs a path to a file that will be used as the terminal block in the dendrimer module.

- **--ngen or -n:** integer

It inputs an integer that will set the number of the generation of the dendrimer that will be built.

Each block should be built as previously described (See Section 1.1.1). These options are only needed if the dendrimer module is being used.

- **Mandatory network options**

- **--bbs or -bbs:** comma-separated list of paths

It receives a list containing the path to each BB used.

- **--in or -in:** path

This option receives the path for the connectivity file dictates the structure of the molecule to be built.

Each block and the connectivity file should be built as described in Section 1.2.1. These options are only needed if the network module is being used.

- **Other mandatory options**

- **--params or -l:** path

This option receives the path for the parameters list file that will be included in the generated MTF.

- **Output control**

- **--output or -o:** string

It is used to define the name of the MTF. The resulting .itp file that will be generated will be named accordingly to this option. Default: default.itp.

- **--gro or -g:** string

It is used to define the name of the coordinates file in which the molecular structure is. The name given to the .gro file is set by this option. Default: default.gro.

- **--name or -name:** string

The topology name is the identifier defined into the [ `moleculename` ] field within the MTF (see Sections 1.1.1 and 1.2.1). This option sets the name for the topology. Notice that this name is what the network module uses for refer to the topology in its connectivity file. Default: NEWTOP.

- **Optimization control**

- **--forcefield or -ff:** path

Since pyPolyBuilder uses an internal pseudo-force field to deal with most of common cases, it might be necessary to include real force field parameters in the optimization steps. This options receives the path to get the parameters values from a force field directory in gromacs format.

- **--ngenga or -ngenga:** integer

Set the number of genetic algorithm (GA) generations used. Default: 20.

- **--npop or -npop:** integer

Set the number of GA population used. Default: 25.

- **--nsteps or -nsteps:** integer  
Set the number of max steps used in geometry optimization. Default: 200.
- **--stepLength or -stepLength:** float  
Length of the step used in the geometry optimization. In pyPolyBuilder, the geometry optimization step is not an actual energy minimization since some terms are omitted and by setting the **--nskipLJ** options, for instance, the objective function may be changed during the run. However, this is an indicative of the distance that atoms are moved at each optimization step. Default: 0.0001.
- **--nskipLJ or -nskipLJ:** integer  
To avoid initial bad contacts resulting from the stochastic process from GA, the evaluation of Lennard-Jonnes (LJ) interactions can be avoided in the few first optimization steps. This option sets the number of iterations for which the LJ interactions will not be computed. Default: 0.

- **Log control**

- **--verbose or -v:**  
Be verbose. Tells pyPolyBuilder to explicit, in screen, every action it is making.
- **--debug:**  
Print debug messages.
- **--nogeoem:**  
Do not perform geometry optimization. It can be used in cases that one need a MTF only and not a guess for the geometry. Also useful in saving time in building molecules that require calling pyPolyBuilder more than once by not optimizing intermediary structures. Applications of this option are given in the following tutorials.

- **Building Mode**

- **--dendrimer or -dendrimer:**  
Set building mode to Dendrimer.

- **--network or -network:**  
Set building mode to Network.

- **Output Formats**

- **--gromacs or -gromacs:**  
Set output format to GROMACS. (This is default.)
- **--gromos or -gromos:**  
Set output format to GROMOS.

## 2 Tutorials

This section is divided in three sections:

- **Dendrimer module**, where some examples using only the dendrimer module are provided;
- **Network module**, where the tutorials discussed use only the network module; and
- **Advances examples**, where both modules are combined to ease the construction of the molecule.

Each tutorial directory has a similar structure. The needed BBs are provided in the files with .itp extensions. In particular, BBs for dendrimers are named according to their topology: `core_**.itp`, `inter_**.itp`, and `ter_**.itp` for the core, intermediary, and terminal BBs. In addition, a bash script named `how_to_run_this_example.txt` can be used to automatically run the tutorial.

After using pyPolyBuilder to generate the geometry and the MTF, these files can be tested in the run directory. This directory has the files needed to carry out a MD simulation using gromacs. In order to run the simulations, it is needed to copy the output from pyPolyBuilder in the run directory. That is, if the PAMAM tutorial is being studied, for example, after running the pyPolyBuilder, the following command line needs to be used:

```
cp PAMAM.* run/.
```

In the run directory, there is a bash script with `.sh` extension and named in accordance with the tutorial name. This file can be used to automatically run a workflow for solvate, equilibrate, and a MD simulation steps. The equilibration step is divided into two steps: a nvt equilibration for 100 ps, and also a npt for 100 ps. By default, the MD simulation step will run 100 ps of production too, for time saving purposes. Before executing the bash file, it needs to be edited to use the correct gromacs path on your machine.

## 2.1 Dendrimer module

In the dendrimer module, pyPolyBuilder is automated to build a dendrimer based on three BBs. The force field parameters will be provided in the software through a file defined using the `--params` option. After the building blocks are defined, the only needed input is the dendrimer generation that should be passed through the `--ngen` option. PyPolyBuilder is programmed to calculate the number of monomers needed to generate a dendrimer of the desired generation number and to connect all of them. This process outputs a final topology file and an initial guess for a coordination file that should be minimized afterwards with a proper MD package.

Here we provide some simple tutorials aiming to give the user some prototype files and to show the philosophy of using the dendrimer module of pyPolyBuilder.

### 2.1.1 PAMAM

Poly(amido amine) (PAMAM) dendrimer is defined as the dendrimers built using an ethylene-diamine core, a tertiary amine intermediary and a primary amine terminal. The building blocks are illustrated in Figure 1.

The BBs are provided in the demo directory in pypolybuilder root, whose structure is illustrated below:

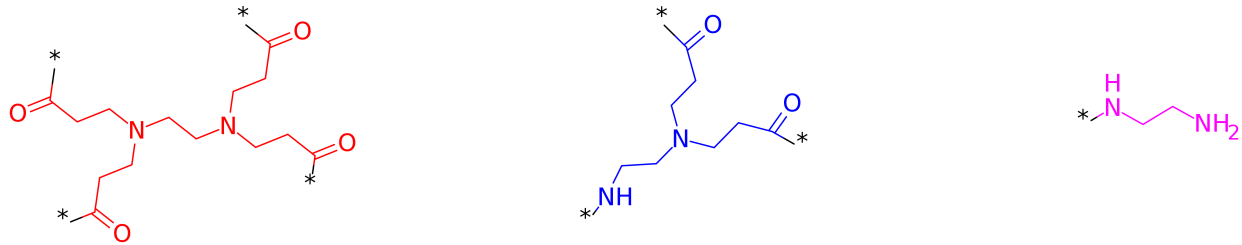


Figure 1: PAMAM dendrimer BBs. The core ethylene diamine block is illustrated in red, the intermediary tertiary amine one in blue, and the terminal primary amine block is displayed in pink.

```
<path/to/pypolybuilder>/demo/gromacs_format/dendrimer/PAMAM
```

```
PAMAM
├── core_PAMAM.itp
├── inter_PAMAM.itp
├── ter_PAMAM.itp
├── list_param.itp
└── run
    ├── PAMAM.sh
    ├── PAMAM.top
    └── mdp
```

The file names are self-explanatory, the core\_PAMAM.itp file is the MTF for the core block, the inter\_PAMAM.itp for the intermediary, and the ter\_PAMAM.itp, the terminal one. Once the BBs and the parameters list are successfully built, one can easily run pypolybuilder by using the code line below (also available in file `how_to_run_this_example.txt` in demo directory) to obtain a generation 2 PAMAM dendrimer (Figure 2):

```
python3 ../../../../../__main__.py \
--core=core_PAMAM.itp \
--inter=inter_PAMAM.itp \
--ter=ter_PAMAM.itp \
--params=list_param.itp \
--ngen=2 \
--name=PAMAM \
--output=PAMAM.itp \
```

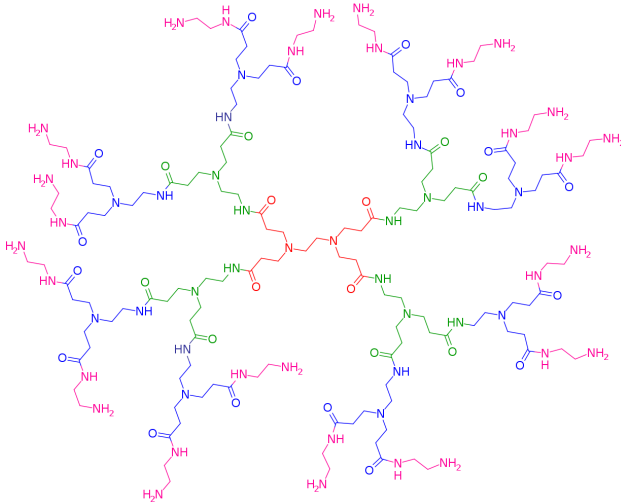


Figure 2: PAMAM G2 dendrimer. The color code is the same as the BBs illustrated in Figure 1. Besides, the first intermediary shell is displayed in green.

```
--gro=PAMAM.gro \
--gromacs \
--dendrimer
```

The used options in this command line were chosen to select each building block (`--core`, `--inter` and `--ter`), the desired dendrimer generation (`--ngen`), the name of the topology, for instance the name that will be placed into [ `moleculename` ] in the MTF (`-name`), the list of force field parameters for pyPolyBuilder (`--params`), as well as to name the coordinates and molecular topology output files (`--gro` and `--output`, respectively). Also, the module and the format for the output were selected using, respectively, `--dendrimer` and `--gromacs`.

Protonated BBs were also provided and the use of them for creating a protonated PAMAM is left as an exercise. They have the same file name as the unprotonated BBs but with the suffix “-protonated”. Protonated BBs were omitted from the directory tree for simplicity.

After pyPolyBuilder finishes the optimization step, one can use any visualization software to check the output geometry. Note that the coordinates are generated considering the molecule in vacuum. Hence, it may not be the expected solvated conformation (see Figure

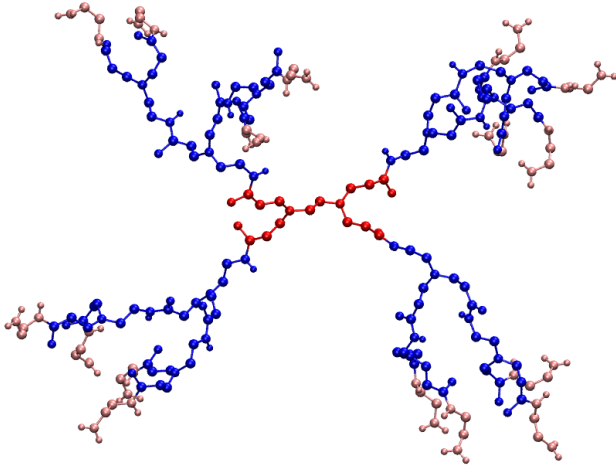


Figure 3: Illustration of PAMAM G2 dendrimer generated with pyPolyBuilder. The snapshot is displayed in accordance with the previously defined color scheme for the BBs. The core is displayed in red, all intermediary shell monomers are displayed in blue,e and the terminal blocks are in pink.

3). Because of that, the run directory has some scripts to run a short MD simulation in order to equilibrate the molecule in water using gromacs.

In order to test the generated MTF and the initial guess for the geometry, we developed an automated script to simulate this tutorial. PAMAM.sh is a script to automatically solvate, equilibrate and simulate the molecule built in this tutorial. Even though the simulated time is not long enough to compute properties consistently, it may be used as basis to future simulations and to evaluate the MTF. PAMAM.top is the topology file for the system and mdp have all required mdp files. However, these scripts were developed for a specific architecture and should be adapted by the user. For instance, the path for gromacs needs to be adapted and the output from pyPolyBuilder (PAMAM.gro and PAMAM.itp) needs to be moved to run directory.

After solvation, an energy minimization cycle, 100 ps of nvt equilibration, and 100 ps of npt equilibration, the system reaches the conformation illustrated in Figure 4.

A powerful feature of the dendrimer module, is that one can easily generate dendrimers of various sizes by only changing one single integer in the input, the `--ngen`. Figure 5 illustrates the procedure of varying `--ngen` from 0 to 5.

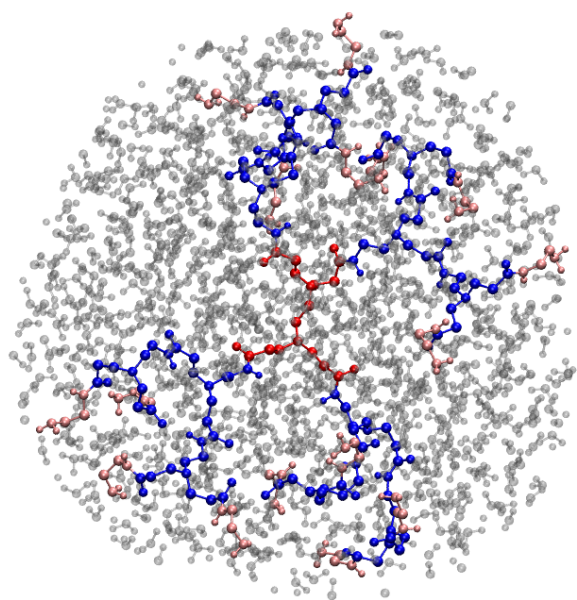


Figure 4: Illustration of PAMAM G2 dendrimer. The core is displayed in red, all intermediary shell monomers are displayed in blue, and the terminal blocks are in pink.

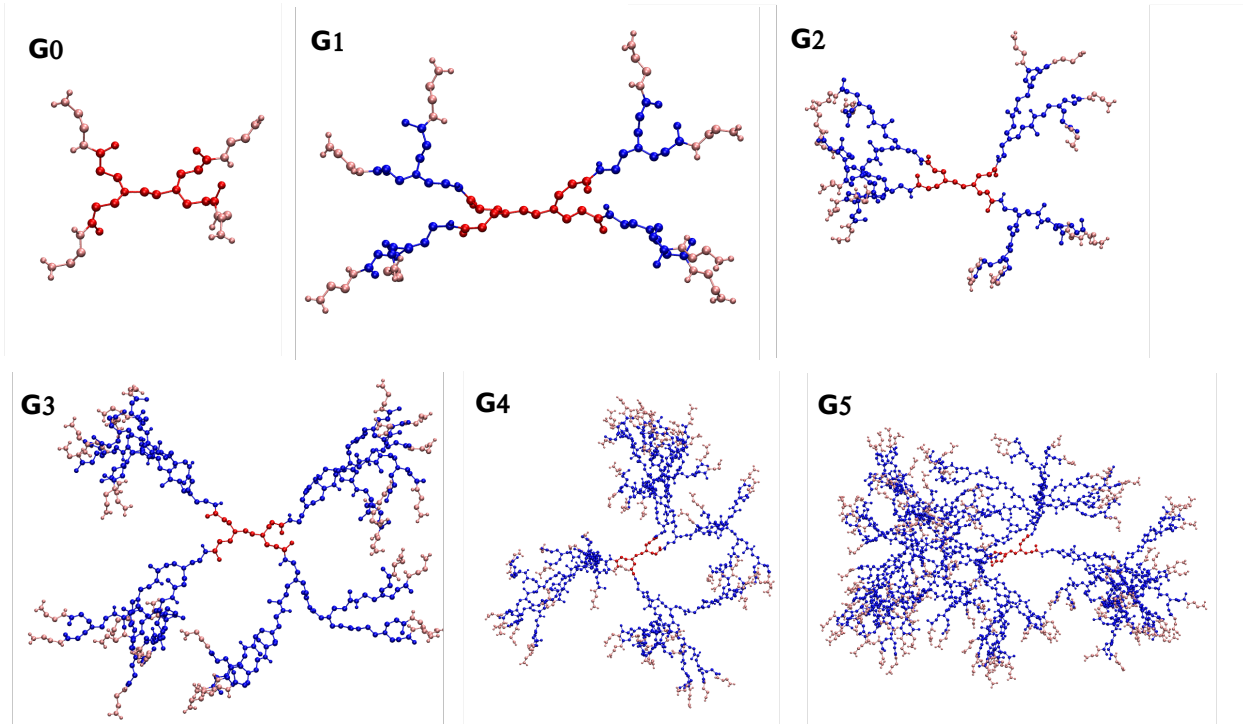


Figure 5: PAMAM dendrimer from generations 0 to 5 built by using pyPolyBuilder and varying the `--ngen` option only between each run. Color scheme is in accordance with Figure 1, that is, the core is displayed in red, all shells of intermediary monomers are in blue, and terminal blocks are in pink.

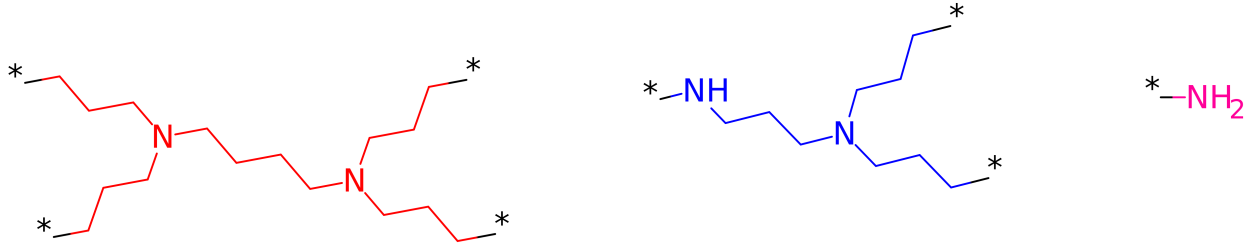


Figure 6: PPI dendrimer BBs. The core diamine butane block is illustrated in red, the intermediary tertiary amine in blue and the terminal primary amine block is displayed in pink.

### 2.1.2 PPI

The BBs for the poly(propyleneimine) (PPI) dendrimer are shown in Figure 6. The core is a diamino butane, intermediary monomer is defined as a tertiary amine in addition to a secondary amine, which is the anchoring point for the core or previous intermediary shell bonding and the terminal is simply a primary amine. It is important to note that PPI dendrimers are commonly defined differently. For instance, this tutorial may be adapted to generate a common used PPI dendrimer by removing the secondary amine from the intermediary monomer and modifying the [ branches ] field in `inter_PPI.itp`.

All PPI BBs, in addition to their protonated versions, are provided in the demo directory in pypolybuilder root, whose structure is illustrated below:

```
<path/to/pypolybuilder>/demo/gromacs_format/dendrimer/PPI
```

```

PPI
├── core_PPI.itp
├── inter_PPI.itp
└── ter_PPI.itp
├── list_param.itp
└── run
    ├── PPI.sh
    ├── PPI.top
    └── mdp

```

The `core_PPI.itp`, `inter_PPI.itp`, and `ter_PPI.itp` files are the MTF for the core, the intermediary, and the terminal blocks, respectively. Once the BBs and the parameters list

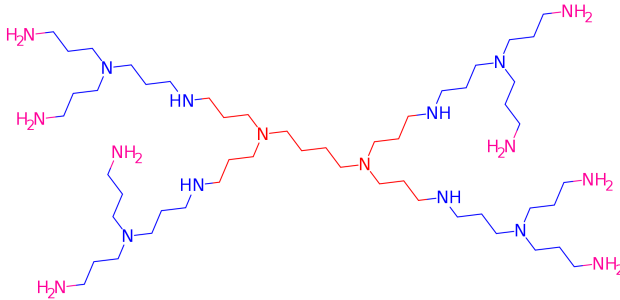


Figure 7: PPI G1 dendrimer. The color code is adopted from Figure 6.

are successfully built, one can easily run pyPolyBuilder by using the code line (also available in `how_to_run_this_example.txt` in demo directory) to obtain a generation 1 PPI dendrimer (Figure 7):

```
python3 ../../../../../__main__.py \
--core=core_PPI.itp \
--inter=inter_PPI.itp \
--ter=ter_PPI.itp \
--params=list_param.itp \
--ngen=1 \
--name=PPI \
--output=PPI.itp \
--gro=PPI.gro \
--dendrimer
```

Each option in this command line was chosen to select each building block (`--core`, `--inter` and `--ter`), the dendrimer generation (`--ngen`), the topology name, for instance the name that will be placed into [ `moleculename` ] in the MTF (`--name`), to parse the list of force field parameters for pyPolyBuilder (`--params`), and to name the coordinates file and MTF output (`--gro` and `--output`, respectively).

It is worth noticing that protonated BBs were also provided. The interested user is welcome to use them. They have the same file name than the unprotonated BBs but with

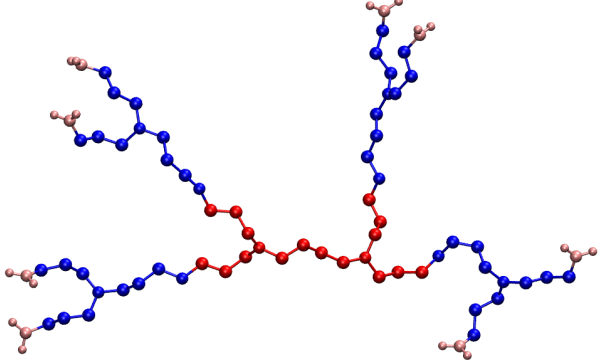


Figure 8: PPI G1 dendrimer built by pyPolyBuilder. The color code is the same as in Figure 6.

the suffix “-protonated”. Protonated BBs were omitted from the directory tree for simplicity.

After pyPolyBuilder finished the optimization step, any visualization software can be used to check the output geometry. Since the coordinates are generated considering the molecule in vacuum, its conformation may not be the expected solvated one (see Figure 8). For that reason, the run directory has some scripts to run a short MD simulation in order to equilibrate the molecule in water using gromacs. However, these scripts were developed for a specific architecture and need to be adapted by each user.

PPI.sh is a script to automatically solvate, minimize energy, equilibrate for 100 ps using nvt and npt ensemble, and run 100 ps of molecular dynamic simulation. The simulation time is actually too small for simulating dendrimers. With this tutorial, we only intend to illustrate how pyPolyBuilder can be used in the creation of a molecular model. PPI.top is the topology file for the system and mdp directory have all required mdp files. However, these scripts were developed for a specific architecture and need to be adapted by each user. For instance, the path for gromacs needs to be adapted and the output from pyPolyBuilder (PPI.gro and PPI.itp) needs to be moved to run directory.

After carrying out the solvation, energy minimization, and 100 ps of both nvt and npt equilibrations, PPI dendrimer is in a more reasonable conformation (Figure 9).

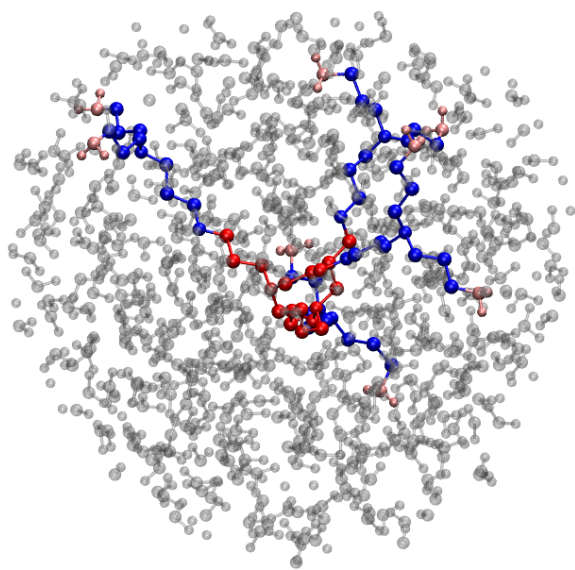


Figure 9: PPI G1 dendrimer in water solution. The color code is the same as in Figure 6. Water molecules are represented as translucent gray molecules.

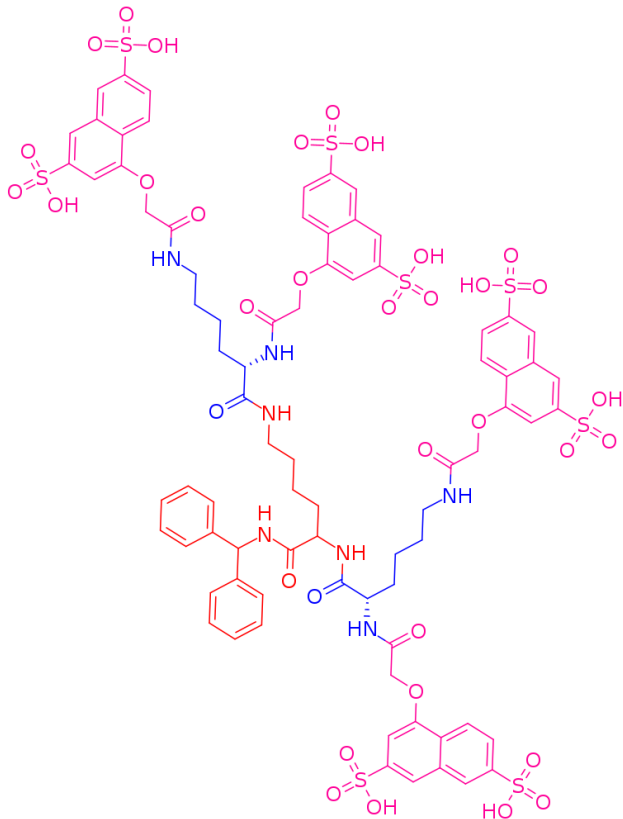


Figure 10: SPL G1 dendrimer. The benzhydrylamine core is illustrated in red, the intermediaries lysine is in blue and the terminals naphthalene disulfonate acid is in pink.

### 2.1.3 SPL7013

Dendrimers of the SPL family are a challenging test case for pyPolyBuilder. Among the examples considered in this tutorial, this is the most complex case for the dendrimer module. Its structure for a generation 1 dendrimer is illustrated in Figure 10.

The division for the SPL BBs was made as the colors suggest in Figure 10. The benzhydrylamine bonded to a lysine is chosen to be the core, a lysine molecule alone is the intermediary block and the naphthalene disulfonate is the terminal block. The BBs themselves are illustrated in Figure 11.

MTFs for those BBs are available in the tutorial directory in pyPolyBuilder root, whose structure is illustrated below:

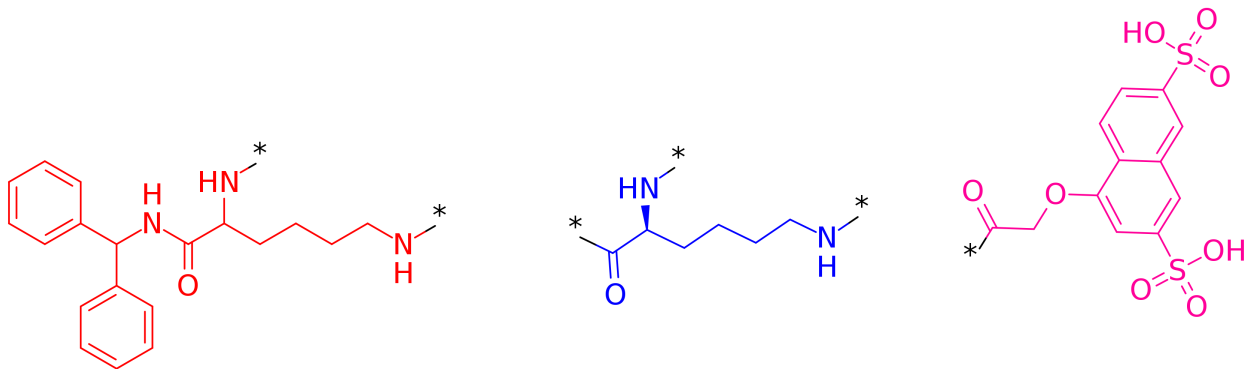


Figure 11: SPL7013 dendrimer BBs. The colors were chosen to match the ones illustrated in Figure 10.

```
<path/to/pypolybuilder>/demo/gromacs_format/dendrimer/SPL7013
```

```
SPL7013
├── core_BHA.itp
├── inter_LYS.itp
├── ter_NDSA.itp
├── list_param.itp
└── run
    ├── SPL7013.sh
    ├── SPL7013.top
    └── mdp
```

The MTFs are named according to its block and molecule name. The core\_BHA.itp is the benzhydrylamine core BB, inter\_LYS.itp the lysine intermediary block, and the ter\_NDSA.itp is the naphthalene disulfonate acid terminal block.

The command line below can be used to build a generation 1 SPL dendrimer (Figure 10):

```
python3 ../../../../../__main__.py \
--core=core_BDA.itp \
--inter=inter_LYS.itp \
--ter=ter_NDSA.itp \
--params=list_param.itp \
--ngen=1 \
```

```
--name=SPL \
--output=SPL.itp \
--gro=SPL.gro \
--nsteps=5000 \
--nskipLJ=100 \
--gromacs \
--stepLength=0.0001 \
--forcefield=../../../../gromos2016h66.ff \
--dendrimer
```

Some of the used option were already used in previous tutorials and explained in Section 1.3. `--core`, `--inter` and `--ter` are used to select the dendrimer BBs and `--ngen` to select its generation number. `--name`, `--output` and `--gro` are used to name the topology, the MTF itself, and the geometry coordinates file, respectively. In turn, `--gromacs` is used to define the format of the output files, in this case, in the gromacs format. Even though `--gromacs` is the default, `--gromos` is also available. `--nsteps` sets how many steps will be made in the geometry optimization. PyPolyBuilder stops the iteration if the energy converges or if the `nstep` is reached. Before doing a local minimization to carry the geometry to a local minimum, pyPolyBuilder carries out a genetic algorithm (GA) optimization only for the torsional dihedrals in order to set an initial quasi-optimum geometry that will be optimized in the local minimization step. Due to the stochastic nature of GA, in very complex geometries it is possible that some atoms are very close at the end of the GA optimization. To avoid these bad interactions, `--nSkipLJ` set how many steps of the geometry optimization will be carried out without evaluating Lennard-Jones (LJ) interactions. Turning LJ interactions off for the first few steps allows the structure to relax only using bonded potentials avoiding bad contacts. `--stepLength` sets the size of the minimization step. Also for complex systems, it is possible that the output of GA is far from optimum. So `--stepLength` can be used to allow a slow minimization in order to avoid overlaps. pyPolyBuilder has default values for bonded and non-bonded parameters. Because of that, the generated structure may not actually be at the minimum of the desired force field. Using the `--forceField` flag, one can pass to pyPolyBuilder the location of a force field (FF) in gromacs format to be used in the

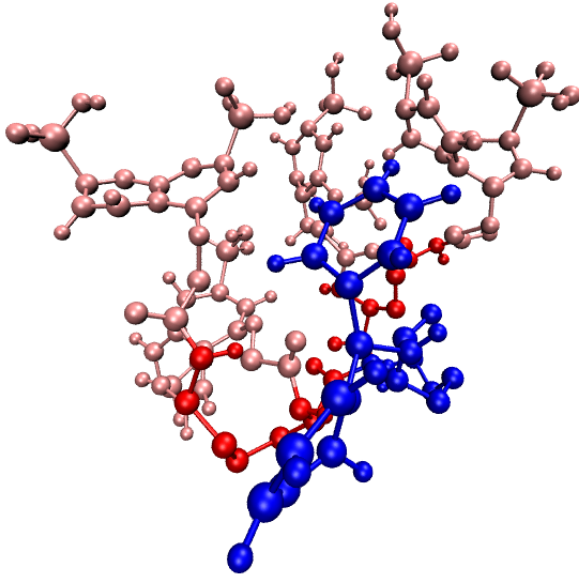


Figure 12: SPL G1 dendrimer built by pyPolyBuilder. The colors were chosen to match Figure 10.

geometry optimization. It may be possible in some cases that the built-in FF is not good enough.

SPL is, for instance, a real challenge due to its very complex structure. In this tutorial, 2016H66 FF was used to optimize its geometry, once the built-in FF leads to an unphysical geometry in which some of the bonds are too long and others are too short (it can be tested by removing the `--forcefield` flag from the previous command-line). Besides, the obtained structure may be equally problematic if the `--nsteps` keyword is not used.

This is a really specific and singular dendrimer. The fact that the pyPolyBuilder is able to build this structure proofs its versatility and robustness.

After pyPolyBuilder finish the optimization stage, one can use any visualization software (such as vmd or pymol) to check the output geometry. Note that the coordinates are generated considering a dendrimer without any partial charge in vaccum. Hence, the obtained conformation is probably not fully realistic. However, if one wants to test the topology of this purely academic case, the run directory has some scripts to run a short molecular dynamics in order to equilibrate the molecule in water using gromacs. These scripts were developed for a specific architecture and should be adapted by the user.

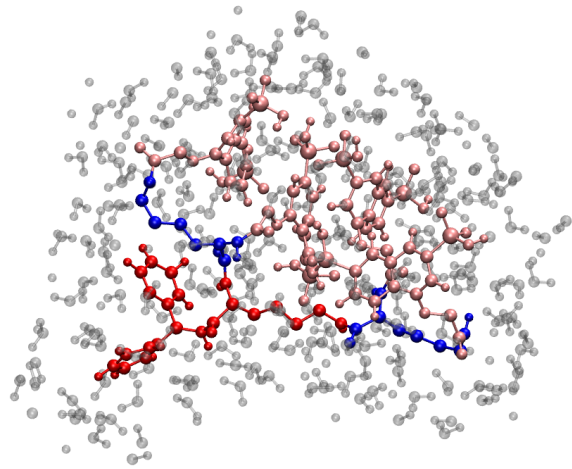


Figure 13: SPL G1 dendrimer equilibrated in water. The colors were chosen to match Figure 10. Water molecules are illustrated as the translucent gray molecules.

By using the available scripts in the run directory to run a small equilibration after solvation and energy minimization in gromacs package, one can see the structure of SPL G1 dendrimer in Figure 13.

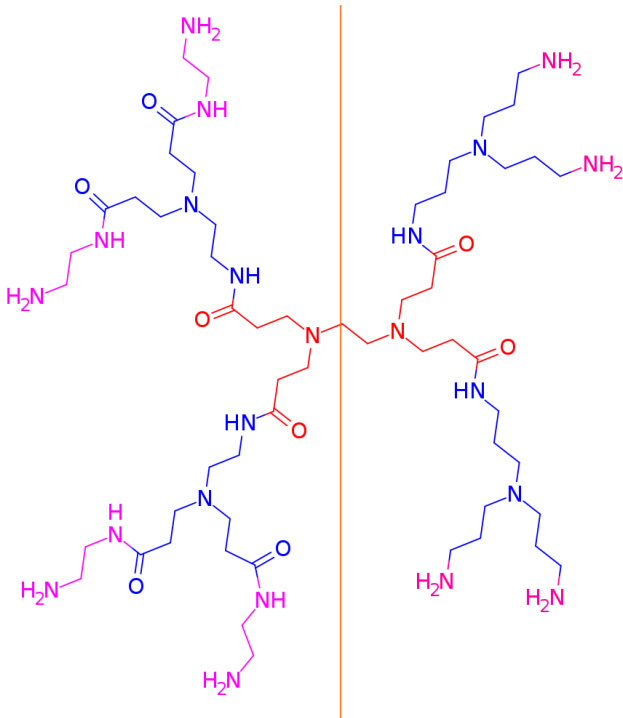


Figure 14: PAMAM/PPI Janus G1 dendrimer.

#### 2.1.4 PAMAM/PPI-Janus

A Janus dendrimer is a amphiphilic dendrimer in which the hydrophobic part is concentrated in a side while the hydrophilic part is concentrated in the opposite side. We illustrated this kind of molecule by concentrating the PAMAM BBs in one region and the PPI BBs at the opposite region of the resulting dendrimer, as depicted in Figure 14. This way, the resulting dendrimer is not amphiphilic due to the similar nature of the PAMAM and PPI chemistry, yet it is a Janus-like dendrimer since it is split into two dendrimeric parts. At the present approach, we used only the dendrimer module for building a method inspired by the divergent synthesis method for creating Janus-like dendrimers. We illustrate this method by using PAMAM and PPI dendrimers BBs.

The dendrimer module was designed for building homogeneous perfect dendrimers. Hence, it is not straightforward to build a dendrimer using BBs of different types. The BBs that are going to be used in this tutorial are exactly the same ones used in previous tutorials (see Figure 15). These files are provided in the demo directory in pypolybuilder root, whose

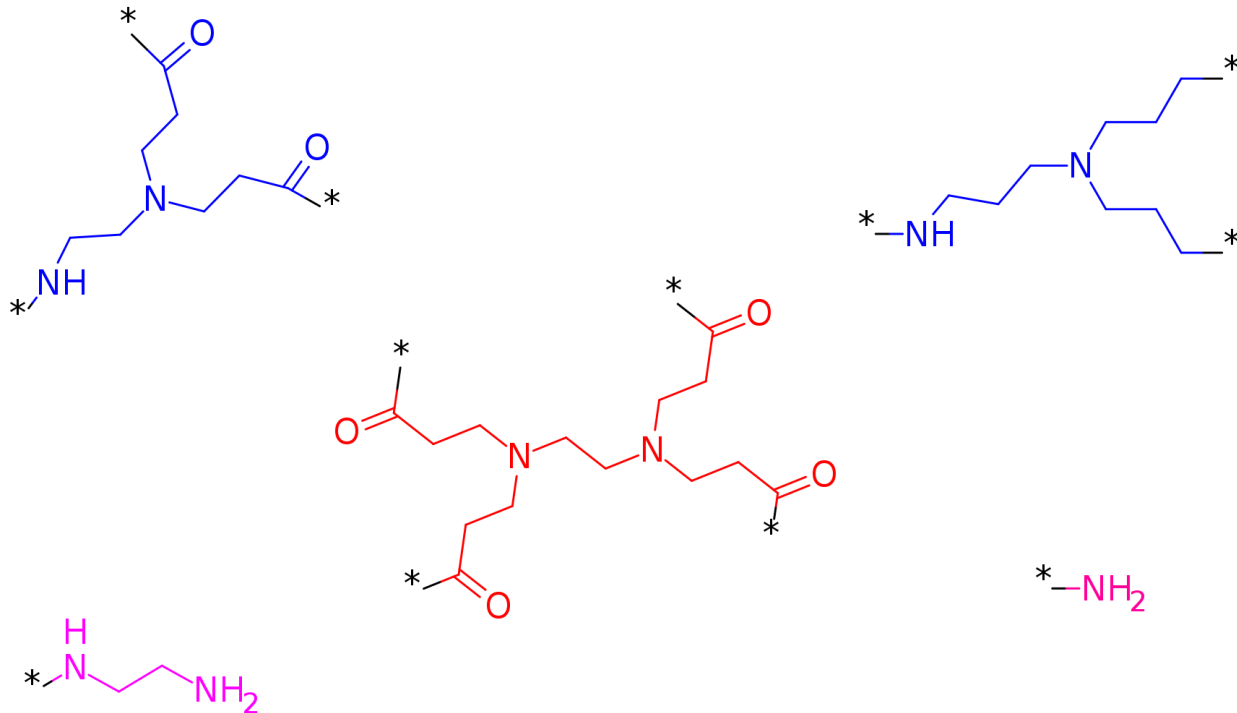


Figure 15: PAMAM/PPI Janus dendrimer BBs. The PAMAM core block is illustrated in red, intermediary and core blocks are placed on the left in blue and red, respectively. On the right, the BBs for PPI intermediary and core, also in blue and red, respectively.

structure is illustrated below:

```
<path/to/pypolybuilder>/demo/gromacs_format/dendrimer/PAMAM_PPI
```

```
PAMAM_PPI
├── core_PAMAM.itp
├── inter_PAMAM.itp
├── ter_PAMAM.itp
├── inter_PPI.itp
├── ter_PPI.itp
├── list_param.itp
└── run
    ├── PAMAM_PPI.sh
    ├── PAMAM_PPI.top
    └── mdp
```

In order to build a Janus dendrimer, we need to make it in two steps. The whole process is automated in `how_to_run_this_example.txt`. Therefore, this file may be executed

with bash in order to get the final geometry. Nevertheless, the process is described in the following. First, we need to grow one half of the dendrimer without growing the opposite half. Afterwards, we use the pyPolyBuilder to grow the second half. The pyPolyBuilder connects the branches using the dendrimer module according to the [ branches ] field. The atom indexes in the “acceptor” column will receive a new bond from the atom in the “donor” column in the incoming new intermediary block. Therefore, we can edit this field in order to select which branches will be created in each step. The `core_PAMAM.itp` file in this tutorial has its [ branches ] field edited as shown below:

```
[ branches ]
; donor acceptor
; 0 10
; 0 13
    0 16
    0 19
```

Lines that begin with “;” are comments and will not be interpreted by pyPolyBuilder. With this [ branches ], the intermediary blocks will only be connected at atoms 16 and 19, respectively. First, the PAMAM branches are included by using the command line code:

```
python3 ../../../../__main__.py \
--core=core_PAMAM.itp \
--inter=inter_PAMAM.itp \
--ter=ter_PAMAM.itp \
--params=list_param.itp \
--ngen=1 \
--name=PAMH \
--output=PAMAMhalf.itp \
```

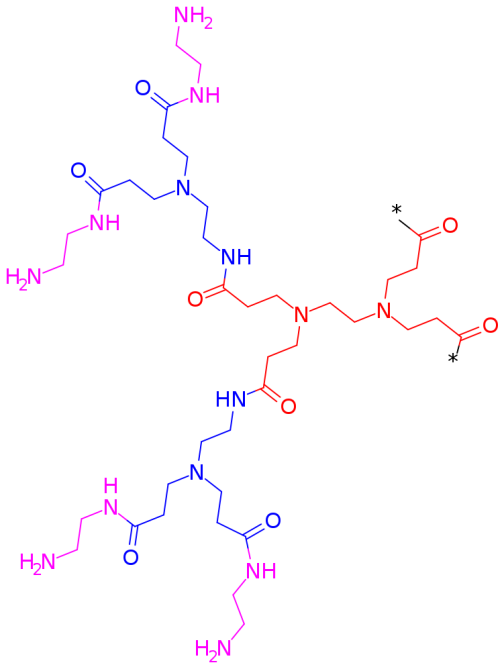


Figure 16: The dendrimer with only the PAMAM BBs attached to the core BB.

```
--nogeom \
--dendrimer
```

In this command line, `--core`, `--inter` and `--ter` were used for selecting the building blocks, `--params` for parsing force fields parameter files, `--ngen` to select the generation of the PAMAM that will be grown, `--name` for naming the topology, and `--output` for naming the generated MTF. Notice that the option `--gro` was not used here since the geometry will not be optimized and an initial guess for the geometry will not be produced. Accordingly, we use the `--nogeom` flag. As this is just an intermediary step in the process of creating the molecule, only the MTF is needed. See the Section 1.3. After the build is done, a new `PAMAMhalf.itp` MTF with the PAMAM half of the Janus dendrimer (Figure 16) will be produced.

The PAMAM core has 4 branch points, at the atoms 10, 13, 16 and 19. We have already used the branch points 16 and 19 for growing the part in which we used the PAMAM BBS. However, the atoms 10 and 13 are available for growing the second half of the dendrimer using the PPI BBS. We need to edit the `PAMAMhalf.itp` file to include the desired [ `branches` ]

shown below in the final of the file:

```
[ branches ]  
;  
; donor acceptor  
0 10  
0 13  
; 0 16  
; 0 19
```

After editing the `PAMAMhalf.itp` file, the second part of the dendrimer may be grown by running the following command line:

```
python3 ../../../../__main__.py \  
--core=PAMAMhalf.itp \  
--inter=inter_PPI.itp \  
--ter=ter_PPI.itp \  
--params=list_param.itp \  
--ngen=1 \  
--name=PAMPPPI \  
--output=PAMAM_PPI.itp \  
--gro=PAMAM_PPI.gro \  
--dendrimer
```

Here, as the geometry is wanted, the `--nogeom` flag was not used. Instead, `--gro` option was used to name the gro file that will be generated.

After pyPolyBuilder has finished the optimization step, one can use any visualization software to check the output geometry. Note that the coordinates are generated considering the molecule in vaccum. Hence, it may not be the expected conformation in solvent (Figure 17). Because of that, the run directory has some scripts to run a short MD simulation

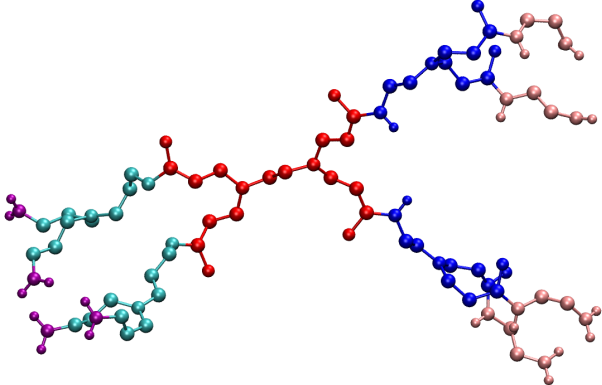


Figure 17: The Janus-like dendrimer generated by pyPolyBuilder using PAMAM and PPI BBs. The PAMAM core is displayed in red, PAMAM and PPI intermediaries are displayed, respectively, in blue and cyan, and the terminal PAMAM and PPI blocks are in pink and purple, respectively. For instance, in this snapshot, the PAMAM half is at the right side of the structure, and PPI half at the left side.

in order to equilibrate the molecule in water using gromacs. However, these scripts were developed for a specific architecture and needs to be adapted by the user.

Using the topology generated by pyPolyBuilder, the Janus-like dendrimer was solvated, its energy was minimized, and equilibrations in nvt and npt ensembles were carried out for 100 ps each (Figure 18).

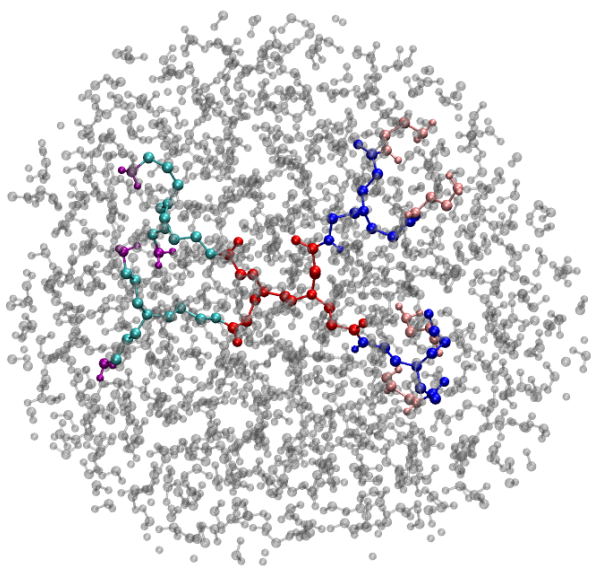


Figure 18: The created Janus-like structure after equilibration in water solvent. Similarly to the Figure 17, PAMAM monomers are at the right side of the molecule while PPI monomers are at the left side, at this snapshot. The color scheme was chosen to be the same as in Figure 17.

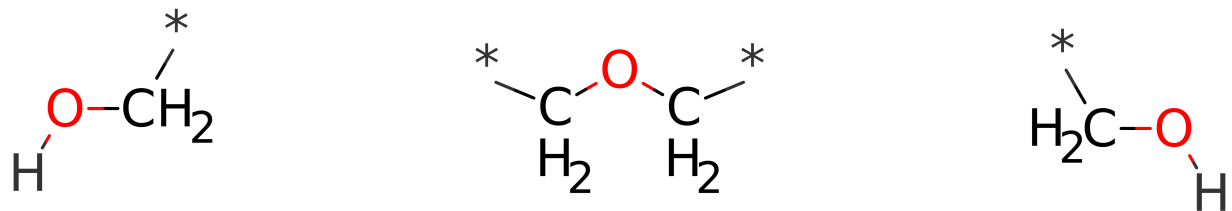


Figure 19: PEG BBs. The left and the right monomers are the initial and final monomers, respectively. The middle one is the repeating monomer unit.

## 2.2 Network module

Differently from the dendrimer module, there is no pre-assumed polymer topology. Hence, the user needs to completely define how every BB is connected. In order to make all information about the molecule connectivity available in one single file, the network module does not use the [ `branches` ] field. Besides, all the connections should be passed in the connectivity file through the `--in` option. Its content was already discussed in Section 1.2.1. Also, when using this module, all the building blocks should be passed using the `--bbs` option (differently from dendrimer module that uses the options `--core`, `--inter` and `--ter`).

Here we provide some simple tutorials aiming to give the user some prototype files and to show the philosophy of using the network module of pyPolyBuilder.

### 2.2.1 PEG

Poly ethyleneglycol (PEG) polymer is made by multiple ethyleneglycol monomers. Each monomer for building PEG in this tutorial was chosen to be the oxygen atom plus its two neighbor atoms in accordance to the GROMOS united atoms FF (Figure 19).

The monomers are pretty simple to be built due to the small number of atoms that need to be defined. The building blocks as defined in Figure 19 are available in the tutorial directory as `bb_PNIP-start.itp`, `bb_PNIP.itp` and `bb_PNIP-end.itp`.

All needed files are provided in the demo directory in pyPolyBuilder root.

```
<path/to/pypolybuilder>/demo/gromacs_format/polymer/PolyEthylene_glycol
```

```

PEG
├── polyeyhylene-start.itp
├── polyeyhylene.itp
├── polyeyhylene-end.itp
├── list_param.itp
├── connect.in
├── connect-4.in
└── run
    ├── PEG.sh
    ├── PEG.top
    └── mdp

```

The command for building a PEG polymer with 5 monomers is automated in a bash file within tutorial directory called `how_to_run_this_example.txt`.

In order to have a PEG polymer with 5 monomers, with the files provided in the tutorial directory, one can use the following command line:

```

python3 ../../../../__main__.py \
--bbs=etyleneglycol-start.itp,etyleneglycol.itp,etyleneglycol-end.itp \
--in=connect.in \
--params=list_param.itp \
--name=PEG \
--output=PEG5x.itp \
--gro=PEG5x.gro \
--network

```

Here, `--bb` option receives the three used BBs: the beginning of the polymer, than the repetition monomer, and a final BB. The connectivity file (named “connect.in”) is input through `--in` option. FF parameters are defined in `list_param.itp` used in `--params`. Also, the output is named according to `--name`, `--output` and `--gro`. At this tutorial, the `--network` flag needs to be used so the network module will be called.

After pyPolyBuilder has finished the optimization step, one can check the molecule geometry in vacuum (Figure 20). This initial geometry and the MTFs can be tested by running a MD simulation using the created files. A script automating a workflow to solvate the

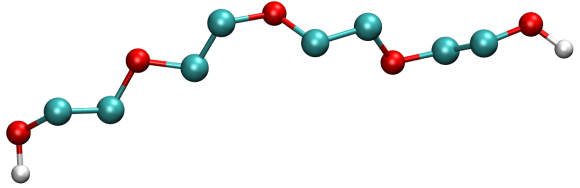


Figure 20: The PEG polymer in the way it is built in this tutorial.

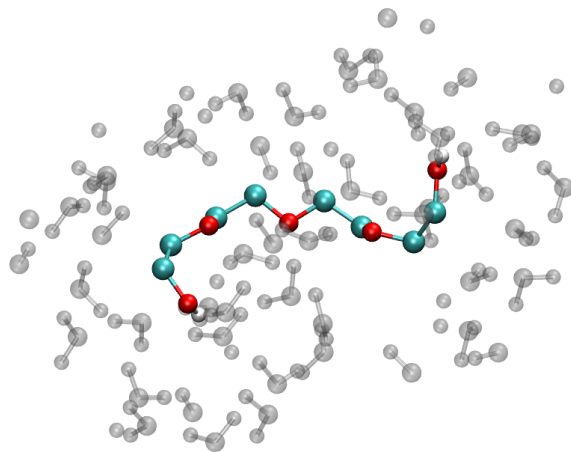


Figure 21: The PEG polymer in the way it is built in this tutorial. Water molecules are displayed as translucent gray molecules.

polymer and run a small MD simulation is included in the tutorial directory.

PEG.sh has the workflow automated but PEG5x.gro and PEG5x.itp need to be moved to the run directory within the tutorial directory, and the script needs to be edited in order to use a actual gromacs path on your computer.

Once equilibrated, PEG polymer conformation can be seen in Figure 21.

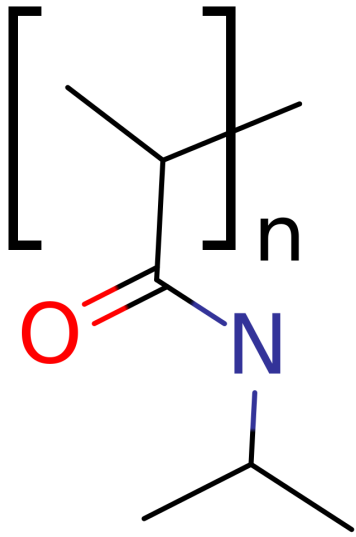


Figure 22: The PNIPAM polymer in the way it is built in this tutorial.

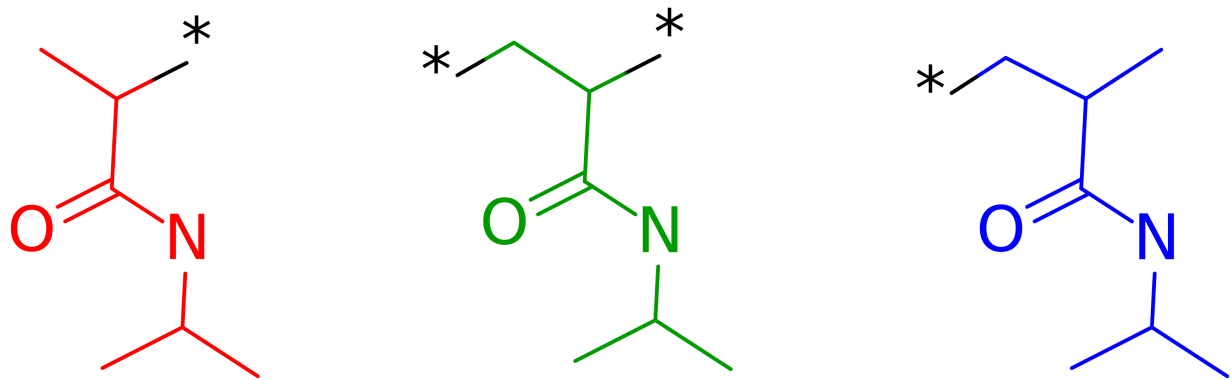


Figure 23: The BBs for building PNIPAM molecules that were used in this tutorial.

### 2.2.2 PNIPAM

The PNIPAM polymer, as it is going to be built in this tutorial, has its monomers as illustrated in Figure 22. Since the connectivity of the first and last monomers are slightly different (shown in Figure 23 by the asterisks), three very similar BBs were created for building PNIPAM. The BBs are provided in the demo directory in pypolybuilder root, whose structure is illustrated in Figure 23: Note that the only difference between each BB is the number of carbon atoms in the backbone.

Where `bb_PNIP-start.itp`, `bb_PNIP.itp`, and `bb_PNIP-end.itp` are the first, the repeated monomer and the last BBs, respectively. The `list_param.itp` is the parameters file,

and it should be written according to the selected force field. The `connect.in` file is the connectivity file. In this file, one should define the used BBs and how they are connected. Using a smaller polymer connectivity file as an example, the `connect-5.in` file is provided with the following content:

```
# Connectivity file to generate a PNIPAM with 5 monomers

#[ BUILDING BLOCKS ]

#BBNUM BBNAME
1  PNIPS
2  PNIP
3  PNIP
4  PNIP
5  PNIPE

#[ CONNECTS ]

#BBI BBJ IAT JAT
1 2 2 1
2 3 2 1
3 4 2 1
4 5 2 1
```

In the field [ BUILDING BLOCKS ], the user should define the names of the BBs that will be used. Those names should agree with the [ moleculetype ] within each BB. Once the BBs are defined, the field [ CONNECTS ] has the information about which BBs are connected (using their index numbers defined in the previous field) and which atom in each BB is used to form the connection. For example, according to line 13: 1 2 2 1, pyPolyBuilder will create a covalent bond between BBs 1 and 2 (the PNIPS and the first PNIP, respectively) using the atom 2 of the first BB, i.e., the PNIPS, and the atom 1 of the second BB, the first PNIP. Consistently, this file contains the informations about what bonds to create in order

to get the polymer built.

To build a PNIPAM with 5 monomer units, one can run the following code:

```
python3 ../../../../__main__.py \
--bbs=bb_PNIP-start.itp,bb_PNIP.itp,bb_PNIP-end.itp \
--in=connect-5.in \
--params=list_param.itp \
--name=PNIPAM \
--output=PNIPAM.itp \
--gro=PNIPAM.gro \
--network
```

Note that the `--network` was explicitly invoked and that all the BBs are passed at once using the `--bbs` option, to be used by the `connect.in` file.

Once the optimization step is gone, the PNIPAM MTF will be ready and an initial guess for the coordinates file will be provided. It is important to note that this guess is evaluated in vacuum and a further energy minimization stage is highly advisable to be carried out in a proper molecular dynamics package.

Since pyPolyBuilder optmization steps are made in vacuum, the geometry (Figure 24) may not be in the same conformation than in solution. To carry a small simulation in water, the run directory is provided. Copy the output from pyPolyBuilder `PNIPAM.*` to `run` and edit the `PNIPAM.sh` script to use the correct gromacs path on your machine. By running `PNIPAM.sh`, the molecule generated by pyPolyBuilder (`PNIPAM.gro`) will be solvated in water, equilibrated, and simulated for 100 ps. After equilibrated, PNIPAM conformation can be seen in Figure 25.

Changing the size of the polymer is not as easy as using dendrimer module. However it is still easily doable. Due to pyPolyBuilder philosophy, all connections are defined in a single file, hence, in order to make more connections, one only needs to edit the connectivity file. Within this tutorial directory, two connectivity files are provided: `connect-5.in` and

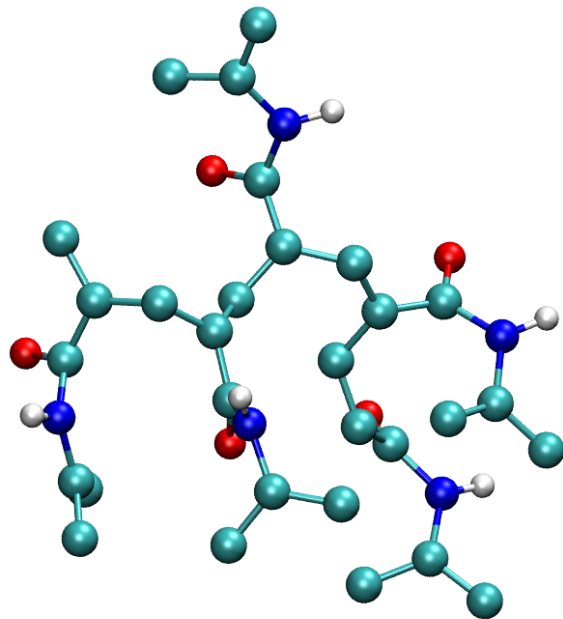


Figure 24: PNIPAM polymer with 5 monomers created by pyPolyBuilder.

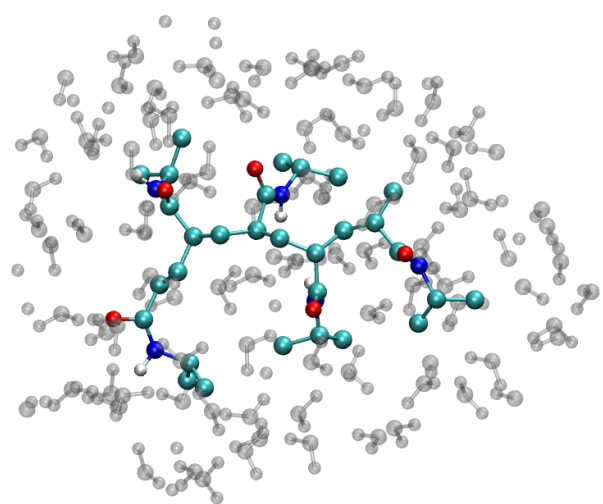


Figure 25: PNIPAM polymer with 5 monomers in water.

`connect.in`. The former connects 5 monomers of the PNIPAM polymer, while the latter uses the same three BBs to build a 30-mer. Using the `connect.in` as the connectivity file, one can simply run the code:

```
python3 ../../../../__main__.py \
--bbs=bb_PNIP-start.itp,bb_PNIP.itp,bb_PNIP-end.itp \
--in=connect.in \
--params=list_param.itp \
--name=PNIPAM \
--output=PNIPAM.itp \
--gro=PNIPAM.gro \
--network
```

That is very similar to the previous one. In fact, only the `--in` option was changed. Both built polymers can be seen in Figure 26.

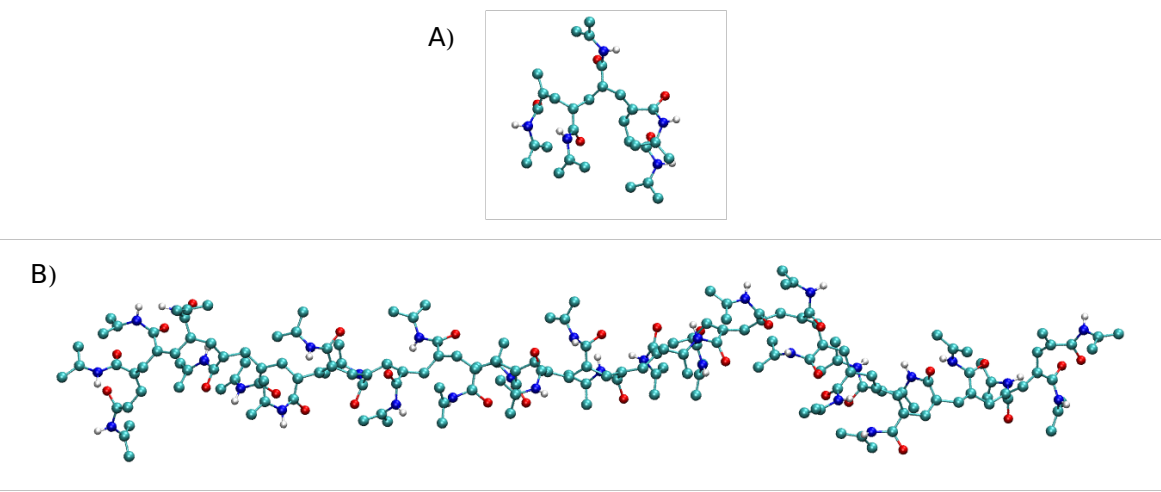


Figure 26: PNIPAM polymers created by pyPolyBuilder. These two polymers were used to illustrate the simplicity of using pyPolyBuilder for varying the size of the created molecule. A) is a PNIPAM with only 5 monomers while B) is a PNIPAM with 30 monomers. The same BBs were used for both polymers.

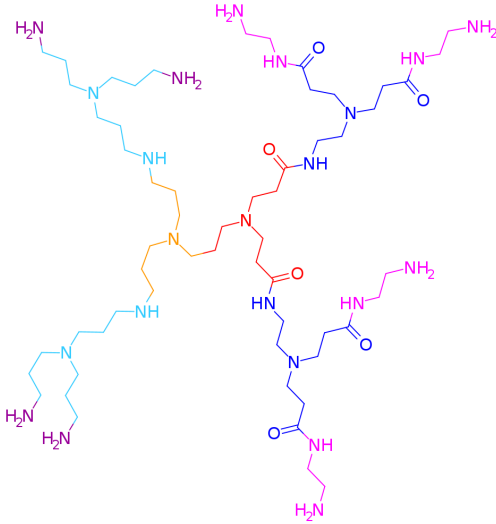


Figure 27: PAMAM/PPI dendrimer in which each side is composed by either PAMAM or PPI dendrimer. The PAMAM core block is illustrated in red, intermediary and core blocks are displayed in blue and pink, respectively. PPI part is displayed in orange, cyan and purple for core, intermediary, and terminal BBs, respectively.

## 2.3 Advanced examples

For some complex cases, it may be necessary to combine both modules to build different parts of the molecule separately. In the following tutorials, parts of the molecule were built using the dendrimer module, and the network module was used to connect them. Generally, these tasks may also be realized with the network module alone since it is completely generic, however, the dendrimer module is specialized in building any dendritic structure, and it is desirable to use it to ease the process.

### 2.3.1 PAMAM/PPI-half

Similarly to the PAMAM/PPI-Janus (see Section 2.1.4), this tutorial aims to build a dendrimer in which each side of the dendrimer is composed by BBs of a different dendrimer. However, differently to the PAMAM/PPI-Janus tutorial, here the molecule is supposed to be completely divided between PAMAM and PPI side. That is, even the core block is split (Figure 27).

All needed files are provided in the demo directory:

```
<path/to/pypolybuilder>/demo/gromacs_format/polymer/PAMAM_PPI_Half
```

```
PAMAM_PPI_Half
├── coreHalf_PAMAM.itp
├── coreHalf_PPI.itp
├── inter_PAMAM.itp
├── inter_PPI.itp
├── ter_PAMAM.itp
└── ter_PPI.itp
├── list_param.itp
├── connect.in
└── run
    ├── PAMAM_PPI_Half.sh
    ├── PAMAMPPITop.top
    └── mdp
```

All the following procedure for building the molecule with a side using only PAMAM BBs, and the other side using only PPI BBs, is automated in an available bash file called `how_to_run_this_example.txt`.

In order to do that, we created separated topology files of the core block for each side of the dendrimer: `coreHalf_PAMAM.itp` and `coreHalf_PPI.itp`. Intermediary and terminal blocks are the same as used in PAMAM and PPI tutorials (See Figure 28).

Therefore, firstly each half will be built using the dendrimer module by using the following command lines to build the first side using PAMAM BBs:

```
python3 ../../../../__main__.py \
--core=coreHalf_PAMAM.itp \
--inter=inter_PAMAM.itp \
--ter=ter_PAMAM.itp \
--params=list_param.itp \
--ngen=1 \
--name=PAMH \
```

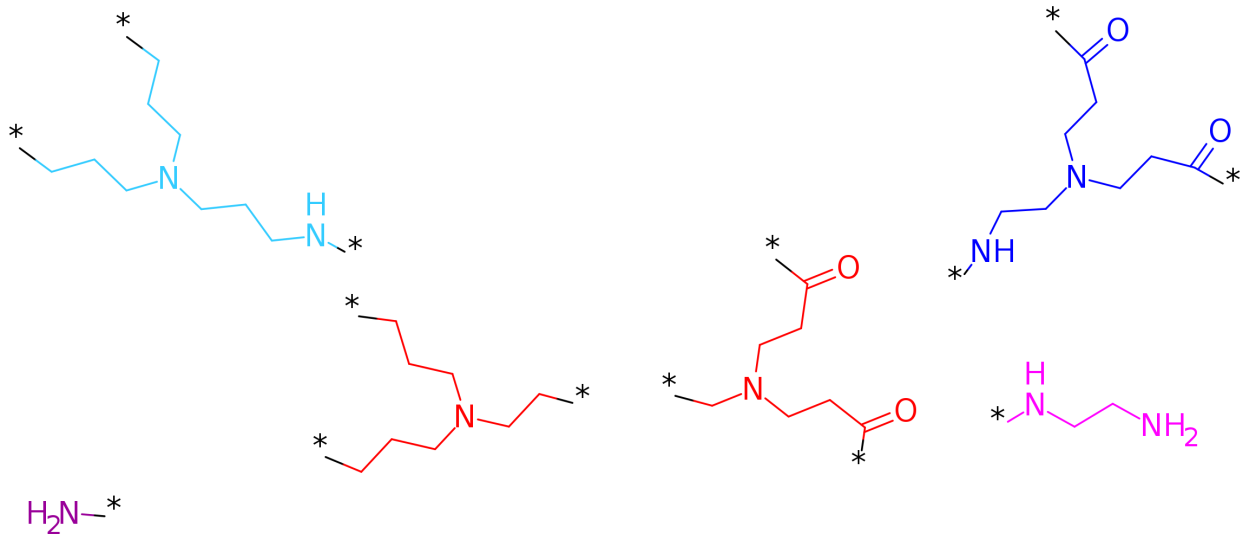


Figure 28: PAMAM/PPI dendrimer in which each side is composed of PAMAM or PPI. PAMAM BBs are illustrated on the right, where the core block is illustrated in red, and intermediary and core blocks are displayed in blue and pink, respectively. PPI part is on the left and displayed in orange, cyan, and purple for core, intermediary, and terminal BBs, respectively.

```
--output=PAMAMhalf.itp \
--nogeom \
--dendrimer
```

Here the usage of dendrimer module is exactly the same as when generating a whole dendrimer. The only difference is that at the core BB, the [ branches ] field tells pyPolyBuilder that only the two available branching points will be used.

```
[ branches ]
; donor acceptor
 0 6
 0 9
```

Secondly, PPI half is built. The procedure is similar to the one for building the PAMAM

half:

```
python3 ../../../../__main__.py \
--core=coreHalf_PPI.itp \
--inter=inter_PPI.itp \
--ter=ter_PPI.itp \
--params=list_param.itp \
--ngen=1 \
--name=PPIH \
--output=PPIhalf.itp \
--nogeo \
--dendrimer
```

In both calls of pyPolyBuilder, the option `--nogeo` was used since we're not interested in optimizing the geometry for an intermediary molecule. Hence, this options can be used to save time skipping the optimization stage and the generation of a coordinates file for the intermediary molecules.

Once both halves are done, one will have the MTFs for each side of the dendrimer (see Figure 29). Now these two parts need to be connected by using the network module. In order to do that, the `connect.in` file simply uses both sides and connect the fist atom of each topology.

```
#[ BUILDING BLOCKS ]
1 PAMH
2 PPIH

#[ CONNECTS ]
1 2 1 1
```

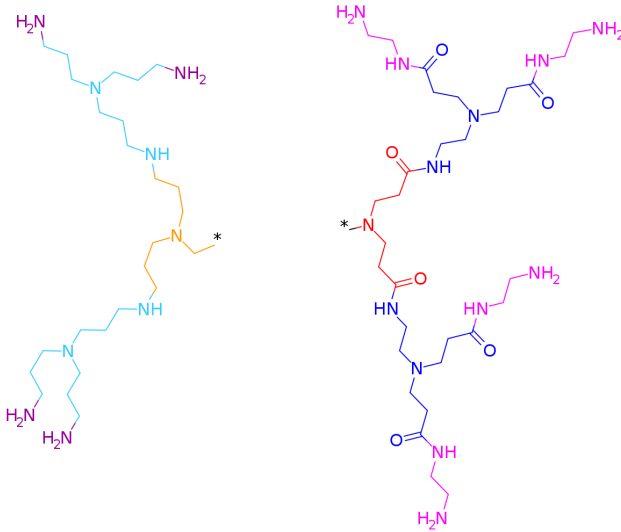


Figure 29: PAMAM/PPI dendrimer in which each side is composed by either PAMAM or PPI dendrimer. The PAMAM core block is illustrated in red, intermediary and core blocks are displayed in blue and pink, respectively. PPI part is displayed in orange, cyan and purple for core, intermediary and terminal BBs, respectively.

Connecting both sides, the final molecule will be generated (Figure 29). The command line to connect the dendrimer is the following one:

```
python3 ../../../../../__main__.py \
--bbs=PAMAMhalf.itp,PPIhalf.itp \
--in=connect.in \
--params=list_param.itp \
--name=PAMPPI \
--output=PAMAMPPI.itp \
--gro=PAMAMPPI.gro \
--nsteps 500 \
--network
```

Once this last command line is executed, the dendrimer will be readily prepared, including its geometry optimized in vacuum (Figure 30). In this tutorial, the internal pseudo-force

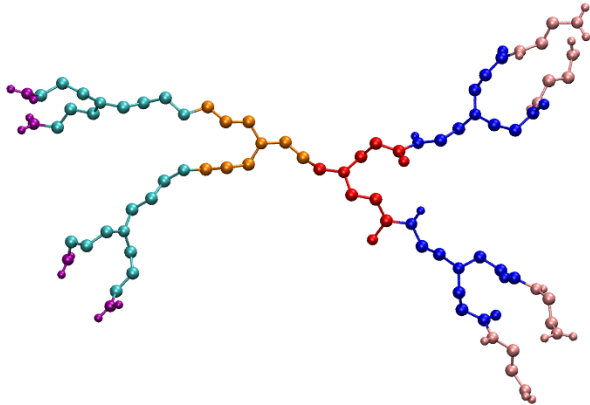


Figure 30: PAMAM/PPI dendrimer in which each side is composed by either PAMAM or PPI dendrimer. The PAMAM core block is illustrated in red, and intermediary and core blocks are displayed in blue and pink, respectively. PPI part is displayed in orange, cyan, and purple for core, intermediary and terminal BBs, respectively.

field of pyPolyBuilder was used, so it is strongly suggested that, after building any molecule in pyPolyBuilder, a MD simulation package should be used to minimize the energy using an actual force field (if the option `--forcefield` was not used in pyPolyBuilder) and to equilibrate the molecule in the desired solvent.

To mimic the procedure of energy minimization, equilibration and simulation, the run directory was provided where the script `PAMAM_PPI_half.sh` can be used to automate the whole process. The outputs from pyPolyBuilder `PAMAMPPI.*` needs to be copied to the run directory and `PAMAM_PPI_half.sh` need to be changed to include an actual path for a gromacs package. By running `PAMAM_PPI_half.sh`, it will automatically solvate the molecule, minimize the energy of the system, equilibrate it for 100 ps in nvt and npt ensembles, and carry out 100 ps of molecular dynamics simulation. An equilibrated structure is shown in Figure 31.

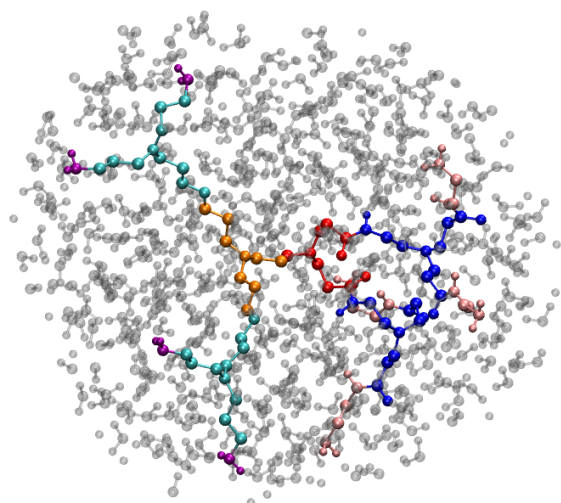


Figure 31: PAMAM/PPI dendrimer in which each side is composed of either PAMAM or PPI in water. The color scheme for the dendrimer is in accordance with Figure 30 and water molecules are translucent in gray.

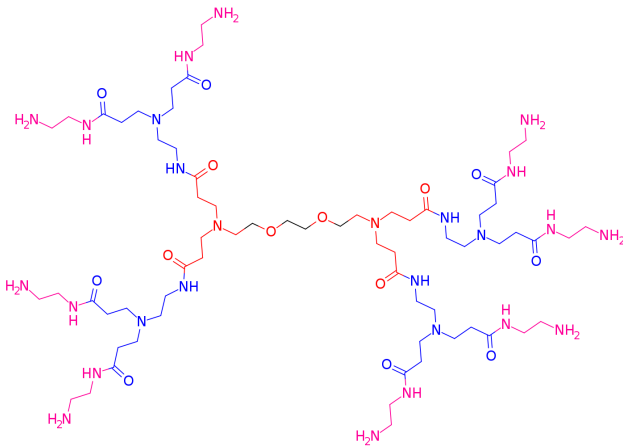


Figure 32: PAMAM-PEG dendrimer in which each side is connected by a PEG polymer. The PAMAM core block is illustrated in red, intermediary and core blocks are displayed in blue and pink, respectively. The PEG moiety of the molecule is placed between the two PAMAM halves.

### 2.3.2 PEG connected PAMAM dendrimer

This tutorial aims to generate a molecule in which two halves of a PAMAM dendrimer are connected by a Poly-ethyleneglycol (PEG) linker. Similarly to the PAMAM/PPI-Half tutorial, this tutorial needs to use both modules in order to prepare the molecule conveniently (Figure 32).

All needed files are provided in demo directory:

```
<path/to/pypolybuilder>/demo/gromacs_format/polymer/PAMAM_PolyEtleneglycol
```

```
PAMAM_PolyEtleneglycol
├── coreHalf_PAMAM.itp
├── inter_PAMAM.itp
├── ter_PAMAM.itp
├── ethleneglycol.itp
├── list_param.itp
├── connect.in
└── run
    ├── PAMAM_PEG.sh
    └── PAMAM_PEG.top
```

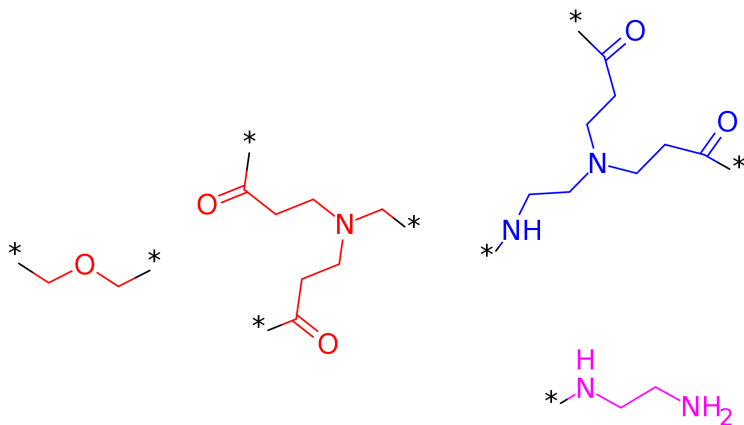


Figure 33: PAMAM-PEG dendrimer BBs. The PAMAM core, intermediary and terminal blocks are displayed in red, blue and pink, respectively. The PEG monomer is placed on the left of the figure.

```
|_ mdp
```

All the following procedure for building the PAMAM dendrimer, in which its core is split into two halves connected by a PEG polymer, is automated in the available bash script `how_to_run_this_example.txt`.

For building this molecule, we will make the dendritic part of the molecule first and then connect two of these, using the network module to get all BBs together. The core BB for the dendrimer module is a PAMAM core split in the middle. Intermediary and terminal blocks are the same as used in PAMAM tutorial (See Figure 33).

The dendrimer module can be called to generate the dentritic part of the molecule by using the following command line to build the first part of the molecule using PAMAM BBs:

```
python3 ../../../../../__main__.py \
--core=coreHalf_PAMAM.itp \
--inter=inter_PAMAM.itp \
--ter=ter_PAMAM.itp \
--params=list_param.itp \
--ngen=1 \
```

```
--name=PAMH \
--output=PAMAMhalf.itp \
--nogeo \
--dendrimer
```

Here the usage of dendrimer module is exactly the same as when generating a whole dendrimer. However, the [ branches ] field is made in a way that pyPolyBuilder uses only the two available branching points.

```
[ branches ]
; donor acceptor
  0 6
  0 9
```

In the first pyPolyBuilder call, the option --nogeo was used because we are not interested in optimizing the geometry of a molecule, which is only an intermediate step of building the final MTF. Hence, this options can be used to save time skipping the optimization stage and the generation of a coordinates file for the intermediary molecules.

Once we have built the dendritic part of the molecule (Figure 34), the network module can be called to connect every BB we need. For instance, we will use the blocks in Figure 33 to build the molecule in Figure 32. In order to do that, the `connect.in` file defines how the BBs are connected. To make it simple, only two monomers of PEG will be used.

```
#[ BUILDING BLOCKS ]
1 PAMH
2 EG
3 EG
4 PAMH
```

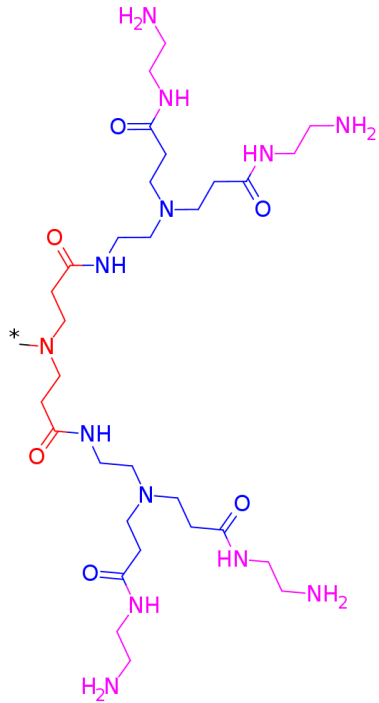


Figure 34: PAMAM part of the PAMAM-PEG dendrimer. The color scheme is adopted in accordance with the Figure 33.

```
# [ CONNECTS ]
1 2 1 1
2 3 3 1
3 4 3 1
```

After building all the BBs, the final molecule will be done and can be visualized in any software (Figure 34). The command line to gather all BBs together is the following one:

```
python3 ../../../../../__main__.py \
--bbs=PAMAMhalf.itp,etleneglycol.itp \
--in=connect.in --params=list_param.itp \
--name=PAMPEG --output=PAMAM_PEG.itp \
--gro=PAMAM_PEG.gro \
--network
```

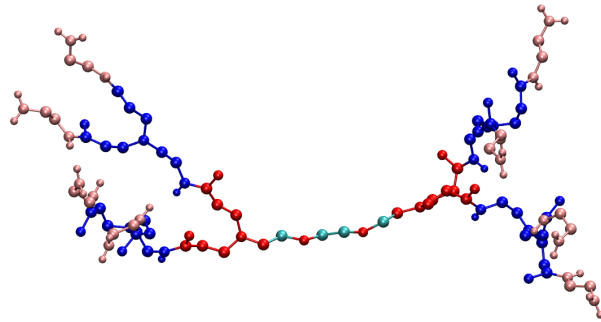


Figure 35: PAMAM\_PEG dendrimer in which each side is connected by a PEG linker. The PAMAM core block is illustrated in red, intermediary and core blocks are displayed in blue and pink, respectively. The PEG moiety of the molecule is placed between the two PAMAM halves.

Once this last command line is done, the dendrimer will be completed, including its geometry optimized in vacuum (Figure 35). It is strongly suggested that the geometry generated by pyPolyBuilder be energy-minimized in a simulation package. The procedure of energy minimization, equilibration, and simulation, is automated in a script file called **PAMAM\_PEG.sh** in the run directory.

The outputs from pyPolyBuilder **PAMAM\_PEG.\*** needs to be copied to the run directory and the input path for the gromacs package in the **PAMAM\_PEG.sh** needs to be changed. **PAMAM\_PEG.sh** can be run in order to automatically solvate the molecule, minimize the energy of the system, equilibrate it for 100 ps in nvt and npt ensembles, and carry out 100 ps of molecular dynamics simulation. Figure 36 shows a snapshot of an equilibrated conformation of the PAMAM-PEG molecule.

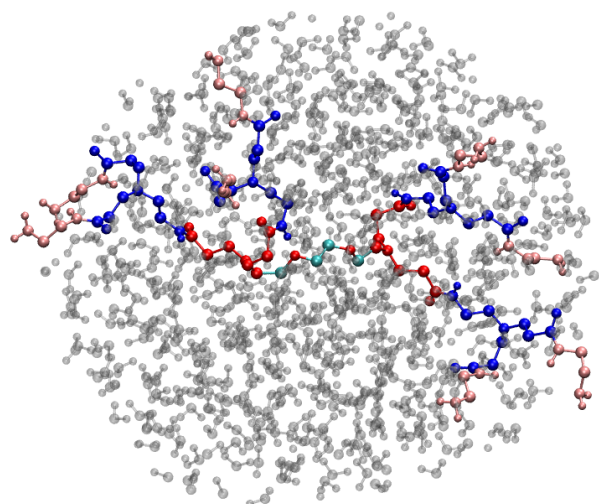


Figure 36: PAMAM\_PEG dendrimer in water. The color scheme is in accordance with Figure 35. The water molecules are displayed as translucent silver molecules.