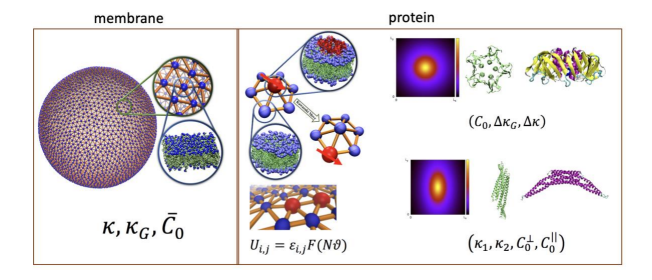
## **User Manual FreeDTS**



### **About FreeDTS**

FreeDTS is a software for performing computational research on biomembranes at the mesoscale. In FreeDTS, a membrane is represented by a dynamically triangulated surface equipped with vertex-based inclusions to integrate the effects of integral and peripheral membrane proteins.

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### **Citations**

Weria Pezeshkian, John H. Ipsen Mesoscale simulation of biomembranes with FreeDTS doi: https://doi.org/10.1101/2023.05.05.539540

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Fluctuations and conformational stability of a membrane patch with curvature inducing inclusions Soft matter **15**, 9974-9981 (2019)

# **Table of Contents**

COMPILING THE SOURCE CODE	3
HOW TO RUN A SIMULATION USING OPENDTS	3
SIMULATION OUTPUT FILES	3
COMMAND LINE OPTIONS	
INPUT FILE OPTION AND FORMAT	
CONSTRAINTS	4
FIXED FRAME TENSION SIMULATIONS	5
Pressure deference or fixed volume simulation	5
COUPLING TO FIXED GLOBAL CURVATURE	5
HARMONIC POTENTIAL BETWEEN TWO GROUPS OF VERTICES	5
APPLYING CONSTANT SURFACE AREA	5
Parallel Tempering	6
MEMBRANES IN CONFINED SPACES	6
OSMOTIC PRESSURE	6
INCLUSION, PROTEIN MODEL	6
TOPOLOGY FILE FORMAT	7
TS FILE	7
"tsi" format files:	
"q" format files:	
GENERATE SCRIPT	
CONVERT SCRIPT	9
TUTORIALS	g
T1: Framed membranes	c
T2: VESICLE SIMULATIONS	
T3: Proteins	11
T4: Protein Sorting	12
T5: PULLING A MEMBRANE NANOTUBE	13
T6: CONFIRMED MEMBRANES	

# Compiling the source code

To compile the OPENDTS source code just execute "compile.sh" script.

./compile.sh

This will create three binary files: **DTS**, **CNV**, and **GEN**. The **DTS** is for running simulations; **GEN** is for generating triangulated files (TS) and **CNV** is for converting different file formats.

# How to run a simulation using OPENDTS

To run a simulation using **OPENDTS**, we need to feed at least two files to the **DTS** binary. An input file (with *dts* extension) that includes all the simulation inputs (see the "Input File" section) and a topology file that contains information about the topology of the TS mesh. The topology file could be in *top* or *tsi* format (see the Topology File sections).

With these two files a simulation can be started as.

\$PATH/DTS -in input.dts -top top.top

Some of the parameters that are defined in the input file could be overwritten in the command line. Note: running a restart simulation is only possible by providing the restart file name in the command line as: -restart restart file name.res

# **Simulation Output files**

There are a number of output files for each DTS simulations that are listed below.

**vtu files:** This is a list of files that shows the evolution of the system and can be opened by paraview. **tsi files:** A list of files that shows the evolution of the system. These files can be converted to different file format using the Convert Script.

restart file: A file to restart a simulation. This file has "res" extension.

**log file:** Stores some information of about the current simulation. This file has "*log*" extension. **energy file:** This file end with "-en.xvg" and contains system the bending energy, box size, volume and etc., at each step.

# **Command line options**

All the option that can be defined in the command line. Also by executing ./DTS -h you can see this list.

Identifier	Туре	Default value	Description
-in	string	Input.dts	input file name
-top	string	topology.top	Topology file name
-b	int	1	initial time step
-е	int	100	final time step
-seed	int	36723	Random number seed
-defout	string	output	A general string for label a specific run
-ndx	string	Index.inx	Index file name
-restart	string	NO	restart file name
-angle	double	-0.5	minimum of cos of the angle between two faces
-minDist	double	1	Square of minimum distance between two vertices

-maxDist double 3	Square of maximum link length
-------------------	-------------------------------

# Input file option and format

The input file must have a dts extension. Below, you can find the most common options defined in this file. However, this file could contain more information, including inclusions and their interactions. Other options are defined in their corresponding sections.

```
Integrator
                                = MC
MC Moves
                                = 1 1 1
                               = 1
Initial Step
                               = 1000000
Final_Step
                               = 1000
Display periodic
OutPutEnergy periodic
                                = 1000
Restart periodic
                                = 10000
                               = 15 0
Kappa
Spont C
                               = 0
                               = 37382
Seed
OutPutTRJ TSI
                               = 1000 10 TrjTSI
GeneralOutputFilename
                               = output
                                = 2.5 2.6 2.5
Cell Size
```

**Integrator:** The algorithm for updating the configuration of the system. Currently only "MC" available. **MC\_Moves:** An option for activating or disactivating the MC moves. First one is vertex move, second is the link flip and the last one is the inclusion moves. 1 1 1 means all the moves are active.

Initial\_Step: Initial step of the simulation, usually it should be one, unless something else is intended.

Final Step: The final step of the simulations

Display\_periodic: Frequency of coordinate file in vtu file format.

**OutPutEnergy\_periodic:** Frequency of energy file. Energy file contains several other information depending on global constrains applied on the system.

**Restart\_periodic**: Frequency of restart file recording.

Kappa: Membrane bending and gaussian rigidity.

**Spont\_C:** spontaneous curvature of the membrane

Seed: Random number seed.

**OutPutTRJ\_TSI:** Frequency for writing trajectory frames in tsi file, the precision of the coordinates and the name of the folder that the files will be stored in.

**GeneralOutputFilename:** A general string to label all the generated files of a specific run with that string.

Cell Size: The size of the unit cells for domain decompositions, should be larger than 1

**Box\_Centering\_F:** To canter the system inside a box after every m steps. The m should be provided as an input.

**FreezingAGroup:** Freezes a group of vertices. Requires an index file in which the name of the group should have been defined in the index file.

**OutPutTRJ\_BTS:** Simulation trajectory output in a binary file format (bts extension). Requires 3 variables. (1) Frequency of saving the coordinates; (2) Precision for file saving (3) the name of the file.

**Min\_Max\_LinkLenghtsSquare:** Square of the maximum and minimum allowed edge size. Two double numbers. Default values are 1 and 3, corresponding to 1 and  $\sqrt{3}$  in length.

**MinfaceAngle:** The minimum allowed dihedral angle between two neighbouring triangles. This number will indicate the minimum value for the cosine of the angle between the two triangle normals.

# **Applying Constraints**

Certain commands can be added to the input file for simulation with constraints such as fixed global curvature or/and fixed volume.

### Fixed frame tension simulations

For doing this, one needs to add below line to the input file. The last two numbers are the frame tension in the unit of  $[k_BT/a^2]$  and inverse frequency of applying the box size check algorithm to maintain the fixed membrane tension respectively. Note: this algorithm only makes sense for predoc membranes.

```
Frame_Tension = on Position_Rescale 0 5
```

### Pressure deference or fixed volume simulation

;Volume_Constraint	= on eqtime	DP	K	gamma	
Volume_Constraint	= on 100	-0.1	0	0	

This command will link the system energy to a potential as

$$E_V = -\Delta PV + \frac{K}{2}(v - \gamma)^2 V_0^2$$

Where 
$$v = \frac{v}{v_0}$$
 and  $V_0 = \frac{\sqrt{A^3}}{6\sqrt{\pi}}$ 

### Coupling to Fixed Global Curvature

To apply a constraint on the global curvature below command can be added to the input file.

```
;CouplingtoFixedGlobalCurvature = state kr m0
CouplingtoFixedGlobalCurvature = on 60 0.1
```

This will couple the system energy to a function as

$$E_s = \frac{k_r}{2A}(M - m_0 A)^2$$

### Harmonic potential between two groups of vertices

To apply a harmonic potential between two groups of vertices, one need to add below command to the input file.

```
; HarmonicPotentialBetweenTwoGroups = on K L0 eqstep Group1 Group2 nx ny nz HarmonicPotentialBetweenTwoGroups = on 10 100 2000000 Group1 Group2 0 0 1
```

You also need to provide an index file for the group definition. This file should have "inx" extension. An example of index file is as below:

```
Group1 1
24
Group2 3
12 16 33
```

### **Applying Constant Surface Area**

Constant total area based on the triangles area

```
;Apply_Constant_Area = state eq_time gamma Ka
Apply_Constant_Area = on eq_time gamma Ka
```

Eq\_time is number of the steps in which the energy of this command reaches its final value Gamma is a number between zero and 1 that fixes the reference area. 1 provides the maximum possible area and zero is the minimum.

$$E_A = \frac{K}{N_T} (A - A_0)^2$$

$$A_0 = N_T (1 + 2\gamma) \sqrt{3} / 4$$

### **Parallel Tempering**

Adding below command allows for additional MC moves, by running simulations at multiple temperatures. The number of the bins will depends on the max and min beta and the provided threads for the openMP

Parallel\_Tempering = on steps minbetta maxbeta

In the command line with -nt the number of the replicas will be defined.

## Membranes in confined spaces

OPENDTS also allows simulating membranes while they are in confined regions. If the initial membrane is not within the defined region, bthe surface is forced to enter this region by defining an equilibrium time. However, if the eq time is short, it may fail to do so. There are four types of confided space defined in OPENDTS: TwoFlatParallelWall, Cuboid, Ellipsoid and EllipsoidalShell. To apply one of these confinements, one of the below commands must be added to the input file.

CoupleToRigidWalls = on TwoFlatParallelWall 10000 X

"X" is a double number which represent half the distance between the two sandwiching walls.

CoupleToRigidWalls	= on Cuboid 10000 10 10 10	
CoupleToRigidWalls	= on Ellipsoid 10000 10 10 10	
CoupleToRigidWalls	= on EllipsoidalShell 10000 10 10 10 0.01	

### Osmotic pressure

Based on the Jacobus van 't Hoff equation  $\Pi = icRT$ .

$$\Delta E(\Delta V) = -P_0 \left( V_0 \ln \left[ 1 + \frac{\Delta V}{V} \right] - \Delta V \right)$$

Add below line to the input.dts file

Osmotic Pressure = Type time gamma PO

$$V_0 = \frac{\gamma}{6} \left(\frac{A_0^3}{\pi}\right)^{1/2}$$

$$V_0 = \frac{1}{6\pi^{1/2}} A^{3/2}$$

Where  $A_0 = 3N_T \frac{3\sqrt{3}}{4}$ 

# **Inclusions (protein model)**

To have inclusion in the simulations, we need to define different type of inclusions. Inclusion type must be defined at the end of the input file as

```
INCLUSION
Define 4 Inclusions
SRotation Type
                        KG
                            ΚP
                                 KL
                                     C0
                                          COP COL
3
           Pro1
                   10
                       0
                            0
                                 0
                                      0
                                          0
                                               \cap
3
           Pro2
                   20
                       0
                            0
                                 0
                                      0
                                          1
                                               0
3
           Pro3
                   20
                       5
                            0
                                 0
                                      0
                                          0
                                               0
2
           Pro4
                   20
                       5
                            0
                                 0
                                      0
                                          0
                                               0
```

The Inclusion-inclusion interactions should be always placed at the end of the file as

```
Inclusion-Inclusion-Int
1 1 2 0.0 0.0
```

Type 1 n A B

$$e_{i,j} = -A_{i,j} + B_{i,j}(1 + \cos[n\Theta])$$

Type 2 A B n  $\Theta_0$  C  $\gamma_0$ 

$$e_{i,j} = -A_{i,j} - B_{i,j} \cos[n(\Theta - \Theta_0)] - C_{i,j}C_{i,j}(\gamma - \gamma_0)^2$$

Defining inclusions type inside the input file does not means that they are present in the simulation. To create inclusion in the simulations, we need to define them either inside the "*tsi*" file or tell the input file to generate them. Currently we can only generate random distribution of inclusions from the input file. However, some can manually or use a script to add such inclusions to the tsi file.

```
GenerateInclusions
Selection_Type Random
TypeID 1 2 3
Density 0.0 0 0
```

# **Topology file format**

Topology file provides information about the position of all the vertices and how they are linked. There are two types of file that you can provide as a topology file; *top* and *tsi* file format.

**top file:** This file has a "top" extension and contains a list of TS files in q (TS file) file format. An advantage of this topology format is that it allows us to feed multiple TS file into the system. A disadvantage is that it does not include inclusions.

*tsi* file: tsi topology is single file that is also single frames of OPENDTS trajectories. An advantage of this topology file is that it includes the inclusions information.

#### TS file

Triangulated surface files that can be read by FreeDTS either has "q" format or "tsi" format. The Generate script can generate files in both formats (see Generate script section).

### "tsi" format files

The following shows a part of a .tsi file with all necessary keywords highlighted in bold. Every .tsi file starts with a line calling version 1.1. The next line defines the box size (x, y, and z) of the system in nm. The next three sections describe the TS mesh. Each section starts with a keyword (vertex, triangle and inclusion) and their corresponding number. Here, we have 130 vertices (the numbering starts from 0). Each vertex has an index and a position in x, y and z (in nm). The 130 vertices are connected via 256 triangles. Again, every triangle has an index (starting from 0) and is defined by the vertices the triangle connects, i.e. triangle 0 connects vertices 11, 55 and 43. Furthermore, a .tsi file

can have a (protein) inclusion section. Here, there are three inclusions from two different types. Again, each inclusion has an index. The index is followed by the inclusion type (here: type 1 for inclusions 0 and 1, type 2 for inclusion 2) and the corresponding vertex index. The last two (floating point) numbers describe a unit two-dimensional vector (sum of both numbers must be one!) which defines the orientation of the inclusion with respect to the bilayer normal.

the c	menta	tion of tr	ne inci	usion with respec	ct to the bilayer normal.
vers	ion 1.	1			
box	50.0	0000000	00	50.0000000000	50.000000000
verte	ex	13	0		
0	21.1	6062330	83	25.4394806652	25.5960855271
1	27.0	2849954	00	23.2012757654	21.6715285158
2	26.9	9217612	32	25.5136587223	28.0195776981
3	23.3	2732298	96	26.2315165676	28.0075875808
4	26.2	7227731	16	26.3271061222	28.1420707299
5	22.0	3968764	25	23.6080597437	26.8858740866
125	21.5	5562808	60	25.5595098219	26.5363425272
126	23.2	1820253	26	26.8060871266	21.5195141902
127	25.3	1993038	65	24.3519379911	20.6752314764
128	28.0	0932004	58	22.6356946990	23.4685318698
129	21.4	0007412	57	26.5841316766	25.2761757772
tria	ngle	25	6		
0	11	55	43		
1	94	75	14		
2	64	3	91		
3	59	52	40		
253	33	109	44		
254	53	69	47		
255	85	6	74		
incl	usion		3		
0	1	22	0	1	
1	1	5	0	1	
2	2	30	0	1	

## "q" format files

The previous tsi file in q format can be seen below.

Line 1: Box information (3 double numbers).

Line 2: Number of vertices (1 integer number; Let's call it NV).

Line 3 to NV+2: Vertex ID and coordinate (1 integer number and 3 double numbers).

Line NV+3: Number of triangles (1 integer number; Let's call it NT).

Line NV+4 to NV+NT+3: Triangle ID and ID of its vertices (4 integer number).

50.00	000000	000	50.00	00000000	50.0	00000000	 _
130							
0	21.16	50623308	3	25.439480	6652	25.5960855271	
1	27.02	28499540	0	23.201275	7654	21.6715285158	
2	26.99	92176123	2	25.513658	7223	28.0195776981	
3	23.32	27322989	16	26.231516	5676	28.0075875808	
4	26.27	72277311	.6	26.327106	1222	28.1420707299	
5	22.03	39687642	5	23.608059	7437	26.8858740866	
125		55628086		25.559509		26.5363425272	
126		18202532	-	26.806087		21.5195141902	
127		L9930386	-	24.351937		20.6752314764	
128		09320045		22.635694		23.4685318698	
129	21.40	00074125	'/	26.584131	6766	25.2761757772	
256							
0	11	55	43				
1	94	75	14				
2	64	3	91				
3	59	52	40				
•							
•							
253	33	109	44				
233	33	109	44				

```
254 53 69 47
255 85 6 74
```

## **Generate Script**

Generate (**GEN**) bindery allows you to create triangulated surface files with different topologies in two different file formats, "q" or "tsi". Three options exist, flat bilayer periodic in x and y directions, cylinder periodic in the x-direction and a closed sphere. To see all the options, execute \$path/GEN -h Some example:

### To generate flat bilayers

\$path/GEN -box 50 50 30 -type flat -o topol.q

#### Generating closed membranes

```
$PATH/GEN -box 50 50 50 -type tetrahedron -N 20 -o topol.q
```

Note: You can force this structure to become sphere by using Ellipspoidalshell command and a short simulation.

```
CoupleToRigidWalls = on EllipsoidalShell 10000 10 10 10 0.01
```

# **Convert Script**

The convert (**CNV**) bindery allows for converting "tsi" and "q" files to each other and several other different file format such as "vtu" and "gro". To see all the options, execute \$path/GEN -h

## **Tutorials**

#### T1: Framed membranes

The simplest simulation that can be performed by FreeDTS is a flat membrane (a fluid elastic surface) in a PBC box. To do this, we have to first generate a flat TS file. This can be done by using **GEN** binary. Using this you can generate a TS file in a q file format using the below command line.

```
$path/GEN -box 30 30 100 -type flat -o topol.q
```

This command creates a TS file in q file format with a box size of 50\*50\*100. Next, this file name should be added to a topology file name with top extension (here it is named top.top), and we should add the below line in the top file.

```
topol.q 10
```

The number in front of the file gives an *id* to the entire mesh (the value is not important but needs to be unique if multiple files are included). To run a simulation, an input file (with dts extension) is required to define the simulation parameters (see below box and the input file format section).

```
Integrator
                                 = MC
MC Moves
                                 = 1 1 1
Initial Step
                                 = 1
Final Step
                                 = 100000
Display periodic
                                 = 1000
OutPutEnergy periodic
                                 = 100
Restart periodic
                                 = 10000
                                 = 20 0
Kappa
```

= output

Using these two files we are now able to run DTS simulations as: \$PATH/DTS -in input.dts -top top.top -seed 76532

Everything above could be done by running the ./run.sh script in the T1/dts\_tutorials/T1-FramedMembrane a folder in the source code.

To make a membrane tensionless, we need to add below command to the input file (such the example in the input file section).

Frame Tension

= on Position Rescale 0 5

In the dts\_tutorials/T1/T1-FramedMembrane\_b folder run the ./run.sh script to perform this section.

Each run will give multiple outputs. Folder VTU\_Frames contains *paraview* readable files to visualize the evolution of the system. TrjTSI folder provides files in *tsi* format for analysis of the simulation. *output-en.xvg* file contains information about system energy, box size (for constant tensions simulations) membrane volume for constant volume simulations (see the vesicle tutorials) .... as a function of the simulation steps.

### T2: Vesicle simulations

### Perpetrating a Vesicle

For a simulation of a vesicle, we first need to create a vesicle structure. For this, we use **GEN** script to generate a tetrahedron and run a short simulation with *EllipsoidalShell* confinement to shape the tetrahedron into a vesicle. The below command creates a *TS* file in the shape of tetrahedron. The size of the tetrahedron can be changed by -N option

\$PATH/GEN -box 50 50 50 -type tetrahedron -N 20 -o topol.q

Next, we add the generated TS file to the topology file and run a normal simulation while having below line in the input file (for more see "Membranes in confined spaces" section in the manual).

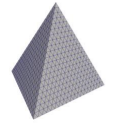
CoupleToRigidWalls

= on EllipsoidalShell 10000 10 10 10 0.01

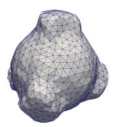
After about 20K steps, you should get a spherical TS file (the run steps may differ for different system)

In the <a href="https://dec.archevideo.com/dec.archevideo">dts\_tutorials/T2-Vesicle</a> folder run the <a href="https://dec.archevideo.com/dec.arc

**Note**: one can easily create a TS file with a spherical shape. However, we prefer this method because it allow us to create a TS file with regular edge size (between range required for DTS simulations) on many different shapes.











#### **Constant Volume Vesicle**

Using the obtain spherical TS file, we can obtain a shape of a vesicle with a constant volume. For this use the **CNV** script and convert last frame of the previous simulation into a q file for a new simulation.

```
$PATH/CNV -in output20.tsi -o vesicle.q
```

Remove below line from your input file

```
CoupleToRigidWalls = on EllipsoidalShell 10000 10 10 10 0.01
```

And add below line to the input file (see "Osmotic pressure or fixed volume simulation" section)

```
Volume_Constraint = on 1000000 0 0.1 0.4
```

In the dts tutorials/T2-Vesicle folder run the ./run 2.sh script to perform this part.

#### T3: Proteins

In this tutorial, we want to make a flat membrane covered by 20% curvature inducing proteins.

**Inclusions**: Proteins are modeled as inclusions. The type and the model parameters of the corresponding inclusion must be defined in the input file (\*.dts file). Below is an example of defining two inclusion types in the input files.

```
INCLUSION
Define 2 Inclusions
SRotation Type
                                  C0
                                      COP COL
                     KG KP
                              KL
          Pro1
                                  0.4
                                            0
                                                 0
                 10 0
                 20
0
          Pro2
                                  -0.4
                                             0
                                                 0
```

**Interactions:** Inclusion-inclusion interactions are also defined in the input file. For interaction type 1 (the tested version)  $E_{ij} = -A + B \cos N\Theta$  it will be as:

```
Inclusion-Inclusion-Int
i j ftype N A B
```

In this dts file, this will be defined as below. Please note: this section MUST always be the last section of the input file (\*.dts file).

```
Inclusion-Inclusion-Int
      1
           1
                2
                        2
                               0.0
1
      2
                 2
                        0
1
            1
                               0.0
2
      2
                 2
                        2
            1
                               0.0
```

**Defining inclusions:** The number of the inclusions in a simulation can be defined in two different ways. If the simulation starts from a q file, a random distribution of inclusions must be defined in the input file.

```
GenerateInclusions
Selection_Type Random
TypeID 1 2 3
Density 0.3 0.1 0
```

However, if the simulation started from a tsi file, inclusions will be read from the inclusion section of this file.

In this first step of the tutorial, we will generate the inclusions from an input file and force them to cluster to enhance the aggregation of the inclusions (to accelerate the process). For this, we will turn off the vertex, edge flip and box change moves.

```
MC Moves = 0 0 1
```

In the dts\_tutorials/T3-Membranes\_Inclusions folder run the ./run.sh script to perform this.

Next, we use the last frame (tsi file) of the previous simulation to start a new simulation with all the required moves.

In the dts\_tutorials/T3-Membranes\_Inclusions folder run the ./run\_2.sh script to perform this.

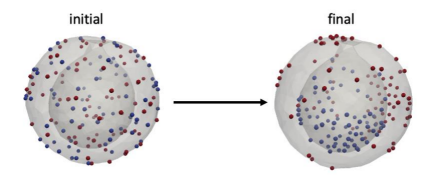
### T4: Protein Sorting

With FreeDTS, we can look at the protein sorting without allowing for membrane shape change. In this way, we can talk for example the shape of membrane structure from Cryo-ET and obtain protein organizations on the membrane. For this, we need to turn off the vertex, edge flip moves and only allow for inclusion moves.

```
MC\_Moves = 0 0 1
```

We will take the stomotosyte shape in the tutorial 2 and add two different protein type.

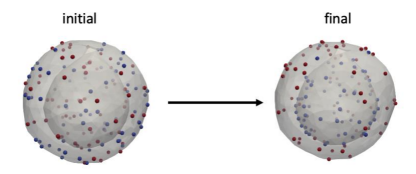
```
INCLUSION
Define 2 Inclusions
SRotation Type K
                                       COP
                                            COL
                     KG KP KL C0
                20 0 0 0 0.3
1
         Pro1
                                        0
                                            0
                      0
                20 0
                         0 -0.3
1
                                            0
         Pro2
                                        0
                20 0 10
                             0 1
1
                          0
         Pro2
GenerateInclusions
Selection Type Random
TypeID
          1
Density
          0.1
               0.1 0.0
Inclusion-Inclusion-Int
         1
             0
                   1
                        0.0
2
    2
         1
             0
                   1
                        0.0
    2
1
         1
             0
                   1
                        0.0
```



In the dts\_tutorials/ T4-protein\_sorting folder run the ./run.sh script to perform this.

The same simulation can be done a system that inclusion one prefers inclusion 2 but still the get sorted by the membrane curvature. For this

Inc	lusio	n-In	clusio	on-Int	
1	1	1	0	0	0.0
2	2	1	0	0	0.0
1	2	1	0	1	0.0



In the dts\_tutorials/ T4-protein\_sorting folder run the ./run\_2.sh script to perform this.

### T5: Pulling a membrane nanotube

To pull a nanotube from a membrane, we need to have a membrane under tension (see previous tutorials for such a simulations). Then we can apply a harmonic potential between two groups, one group can be the single vertex while the other group can be all the other vertices. Using a corresponding index file (See the index file section) and adding below command to the input file we have everything we need to pull a nanotube from a flat membrane.

HarmonicPotentialBetweenTwoGroups = on 10 100 1000000 Group1 Group2 0 0 1

Next, just run below command and wait for the nanotube to form.

\$PATH/DTS -in input.dts -top top.top -seed 76532 -ndx index.inx

In the dts\_tutorials/T4-tether\_pulling folder run the ./run.sh script to perform this tutorial.

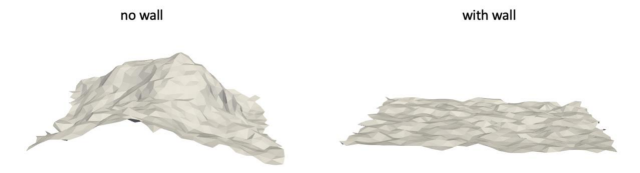
Note: this tutorial might take long.

#### T6: Confirmed membranes

How membranes behave in a confined space. This can be studied using FreeDTS. In this tutorial we will look how two sandwiching hard walls. For this, we need add below command to the inpot file

CoupleToRigidWalls = on TwoFlatParallelWall 10000 0.5

We also set the kappa to 4 to make the



In the dts\_tutorials/T6\_confined\_membranes folder run the ./run.sh script to perform this tutorial.