BioBlender 2.1 Manual

2020-04-08

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What is BioBlender?

BioBlender is a software package based on open source Blender 3D modeling software.

Biology works at the nanoscale, with objects invisible to the human eye. With BioBlender it is possible to show some of the characters that populate our cells, based on scientific data and the highest level of 3D manipulation. Scientists from around the world study proteins at the atomic level and deposit information in the public Protein Data Bank repository, where each molecule is described as the list of its atoms and their 3D coordinates. With BioBlender, users can manage proteins in 3D space, display their surface in a photorealistic way, and construct protein motions based on known conformations.

BioBlender is an implementation of Blender, an open source, freely distributed, cross-platform, interoperable, and compatible 3D animation, visual effects, and video game.

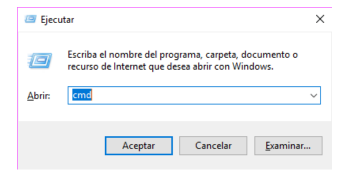
The combination of 3D computer graphics, game engines and scientific programs make BioBlender a complete instrument for elaborating the movement of proteins and displaying their surface characteristics using visual code based on textures and special particle effects. In this way, it is possible to show the physical and chemical properties of a moving molecule.

Requirements

- Install Python 3.5.3
- Install Blender 2.79b
- Install Pymol
- Install the Numpy, Scipy and Prody libraries

To install these libraries:

- 1. Open windows console (Windows $+ \mathbf{R}$)
- 2. Type cmd



3. Then in the Windows console type:

pip install numpy

4. Once finished, proceed to install Prody:

pip install Prody

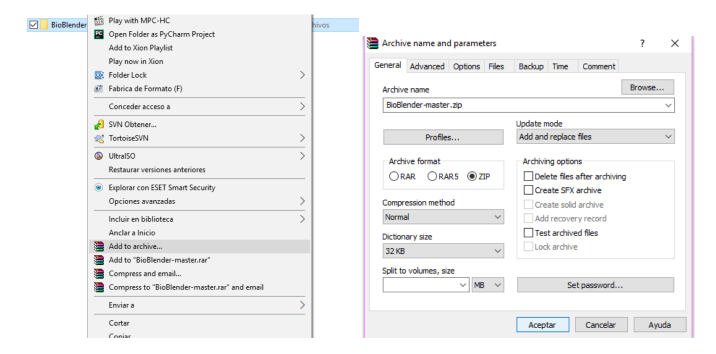
5. By last. Necessary for the proper functioning of the Prody library:

pip install scipy

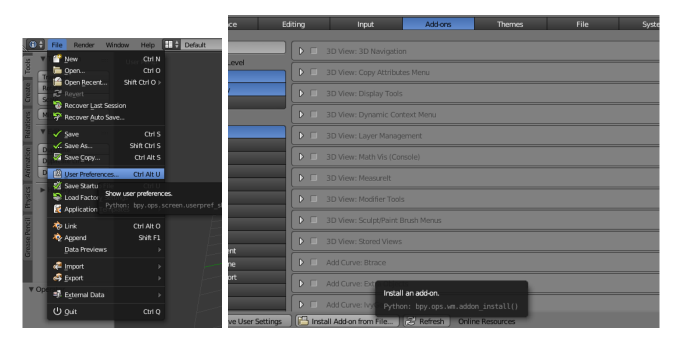
How to install BioBlender?

• Steps:

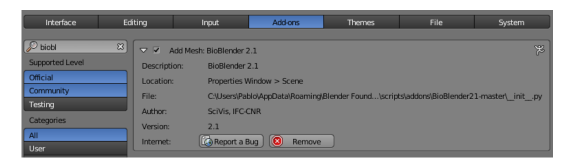
1. Compress the BioBlender-master folder in a .zip file (In case the folder is not compressed)



- 2. Once the folder is compressed, open the Blender and add the BioBlender-master.zip file to Blender. For this you have to open:
 - File > User Preferences > Add-ons > Install Add-ons From File (Here look for the directory, select the .zip file and press Install Add-ons From File)



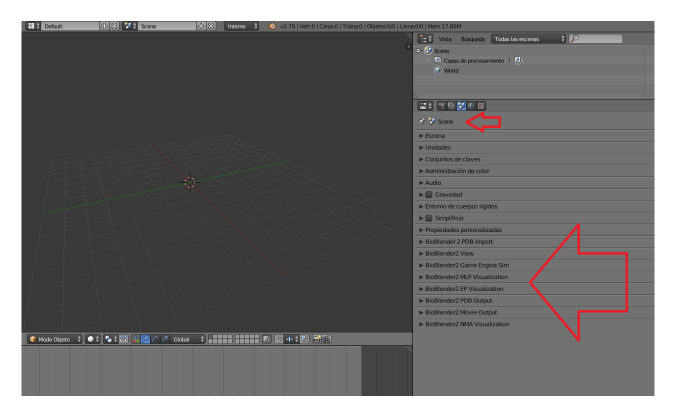
3. Once the BioBlender has been added, it should be activated automatically, but in case it has not been activated, go to: newline File > User Preferences > Add-ons (Here activates BioBlender 2.0) newline Then save the preferences of user.



Import PDB file

Note: Well, first of all, if you want to delete all the objects that load automatically when you open Blender, press the **a** key, then the **x** key and **Enter**. If you want Blender to always open with no objects in the scene, press $\mathbf{Ctrl} + \mathbf{u}$, to save the scene you are in.

The BioBlender functions are found in the properties panel called **Scene**.



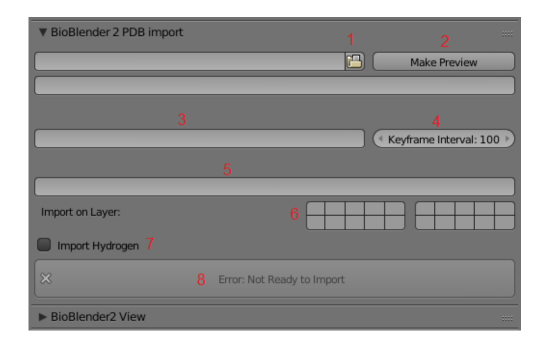
Protein Data Bank (PDB) it is a database of the three-dimensional structure of proteins and nucleic acids. These data, generally obtained by X-ray crystallography or nuclear magnetic resonance, are sent by biologists and biochemists around the world. They are in the public domain and can be freely used. The BioBlender has the possibility to read files in pdb format and obtain information about their atoms to represent them in space.

HEADER	VIRAL PROTEIN 26-JAN-20 6LU7	SCALEI		υ.	OTO			0.00466		0.00000		
TITLE	THE CRYSTAL STRUCTURE OF COVID-19 MAIN PROTEASE IN COMPLEX WITH AN	SCALE2					0.012582			0.00000		
TITLE	2 INHIBITOR N3	SCALE3		0.	.000			0.021223		0.00000		
COMPND	MOL ID: 1;	ATOM	1	. N	I	SER A	1	-32.073	9.085	33.695	1.00 38.90	N
COMPND	2 MOLECULE: SARS-COV-2 MAIN PROTEASE;	ATOM	2	C	CA	SER A	1	-32.156	8.073	34.741	1.00 37.44	C
COMPND	3 CHAIN: A;	ATOM	3	C		SER A	1	-30.857	8.000	35.536	1.00 34.96	C
COMPND	4 ENGINEERED: YES;	ATOM	4	C)	SER A	1	-30.047	8.926	35.507	1.00 33.29	0
COMPND	5 MOL ID: 2;	ATOM	5	(CB	SER A	1 1	-32.483	6.704	34.140	1.00 44.07	C
COMPND	6 MOLECULE: N-[(5-METHYLISOXAZOL-3-YL)CARBONYL]ALANYL-L-VALYL-N~1~-	ATOM	-)G	SER A	1 1	-31.312	6.067	33.660	1.00 47.56	0
COMPND	7 ((1R,2Z)-4-(BENZYLOXY)-4-OXO-1-{[(3R)-2-OXOFYRROLIDIN-3-	ATOM	1	N	1	GLY A	4 2	-30.665	6.892	36.240	1.00 36.02	N
COMPND	8 YL]METHYL}BUT-2-ENYL)-L-LEUCINAMIDE;	ATOM	8	C	CA	GLY A	4 2	-29.510	6.712	37.092	1.00 34.67	C
COMPND	9 CHAIN: C;	ATOM	9) C	2	GLY A	4 2	-29.828	6.998	38.551	1.00 38.34	C
COMPND	10 ENGINEERED: YES	ATOM	10) ()	GLY A	4 2	-30.810	7.663	38.892	1.00 45.40	0
SOURCE	MOL ID: 1;	ATOM	11	. N	ī	PHE A	7 3	-28.974	6.479	39.430	1.00 38.38	N
SOURCE	2 ORGANISM SCIENTIFIC: SEVERE ACUTE RESPIRATORY SYNDROME CORONAVIRUS	ATOM	12		CA	PHE A	3	-29.155	6.661	40.866	1.00 36.10	C
SOURCE	3 2;	ATOM	13		2	PHE A	3	-27.790	6.744	41.527	1.00 44.18	C
SOURCE	4 ORGANISM COMMON: SARS-COV-2;	ATOM	14	()	PHE A	3	-26.981	5.820	41.399	1.00 40.82	0
SOURCE	5 ORGANISM TAXID: 2697049;	ATOM	15		CB	PHE A	3	-29.978	5.522	41.468	1.00 38.52	C
SOURCE	6 EXPRESSION SYSTEM: ESCHERICHIA COLI BL21(DE3);	ATOM	16		CG	PHE A	3	-30.635	5.875	42.770	1.00 40.78	C
SOURCE	7 EXPRESSION SYSTEM TAXID: 469008;	ATOM	17	(D1	PHE A	3	-31.642	6.824	42.816	1.00 43.38	C
SOURCE	8 EXPRESSION SYSTEM VECTOR TYPE: PLASMID;	ATOM	18		CD2	PHE A	3	-30.247	5.261	43.949	1.00 40.00	C
SOURCE	9 EXPRESSION SYSTEM PLASMID: PGEX-6P-1;	ATOM	19		E1	PHE A	3	-32.251	7.155	44.012	1.00 42.94	C
SOURCE	10 MOL ID: 2;	ATOM	20) (E2	PHE A	3	-30.851	5.586	45.148	1.00 40.35	C
SOURCE	11 SYNTHETIC: YES;	ATOM	21		Z	PHE A	3	-31.854	6.534	45.179	1.00 43.94	C
SOURCE	12 ORGANISM SCIENTIFIC: SYNTHETIC CONSTRUCT;	ATOM	22	N	ī	ARG A	4 4	-27.541	7.844	42.233	1.00 39.42	N
SOURCE	13 ORGANISM TAXID: 32630	ATOM	23		CA	ARG A	4 4	-26.277	8.066	42.915	1.00 38.88	C
KEYWDS	PROTEASE, VIRAL PROTEIN	ATOM	24	C	2	ARG A	4 4	-26.545	8.642	44.296	1.00 40.55	C
EXPDTA	X-RAY DIFFRACTION	ATOM	25)	ARG A	4 4	-27.552	9.320	44.517	1.00 36.11	0
AUTHOR	X.LIU, B.ZHANG, Z.JIN, H.YANG, Z.RAO	ATOM	26		В	ARG A	4 4	-25.367	9.020	42.127	1.00 36.87	C
REVDAT	6 18-MAR-20 6LU7 1 JRNL	ATOM	27		CG	ARG A	4 4	-24.669	8.388	40.936	1.00 42.81	C
REVDAT	5 11-MAR-20 6LU7 1 COMPND SOURCE	ATOM	28		CD	ARG A	4 4	-23.342	7.771	41.340	1.00 42.72	C
REVDAT	4 26-FEB-20 6LU7 1 REMARK	ATOM	29	N	ΙE	ARG A	4 4	-22.460	7.579	40.193	1.00 47.79	N
REVDAT	3 19-FEB-20 6LU7 1 TITLE JRNL	ATOM	30) (CZ	ARG A	4 4	-21.235	7.068	40.270	1.00 53.27	C
REVDAT	2 12-FEB-20 6LU7 1 TITLE COMPND JRNL REMARK	ATOM	31	. N	IH1	ARG A	4 4	-20.744	6.693	41.443	1.00 47.42	N
REVDAT	2 2 1 SHEET LINK SITE ATOM	ATOM	32	N	IH2	ARG A	4 4	-20.502	6.930	39.173	1.00 49.36	N
REVDAT	1 05-FEB-20 6LU7 0	ATOM	33	N	ī	LYS A	₄ 5	-25.636	8.362	45.227	1.00 34.78	N
JRNL	AUTH Z.JIN, X.DU, Y.XU, Y.DENG, M.LIU, Y.ZHAO, B.ZHANG, X.LI, L.ZHANG,	ATOM	34	C	CA	LYS A	₄ 5	-25.667	9.020	46.528	1.00 36.92	C
JRNL	AUTH 2 C.PENG, Y.DUAN, J.YU, L.WANG, K.YANG, F.LIU, R.JIANG, X.YANG, T.YOU,	ATOM	35		2	LYS A	5	-25.399	10.504	46.317	1.00 32.31	C
JRNL	AUTH 3 X.LIU, X.YANG, F.BAI, H.LIU, X.LIU, L.GUDDAT, W.XU, G.XIAO, C.QIN,	ATOM	36		0	LYS A	5	-24.261	10.908	46.053	1.00 37.11	0
JRNL	AUTH 4 Z.SHI, H.JIANG, Z.RAO, H.YANG	ATOM	37		В	LYS A	5	-24.643	8.396	47.471	1.00 39.24	C
JRNL	TITL STRUCTURE OF MPRO FROM COVID-19 VIRUS AND DISCOVERY OF ITS	ATOM	38		CG	LYS A	. 5	-25.062	8.413	48.934	1.00 39.96	C
		ATOM	38	C	G	LYS A	5	-25.062	8.413	48.934	1.00 39.96	

Steps to import the PDB:

- 1. Select the PDB file that you want to import, for this BioBlender has 2 options:
 - Search the local computer for the PDB file and select it.
 - Write only the name of the PDB you want to import. This is based on writing the PDB identifier with only 4 letters, which will be searched in the repository http://www.pdb.org/ (You need the Internet)
- 2. Read the PDB file, clicking Make Preview
- 3. Then a list of numbers appears, these numbers are the models that the PDB contains. This list you have the possibility to modify it and import the model that you want.
- 4. You can modify the interval of the frames between each model where it says **Keyframe Interval**
- 5. Then a list of the strings that the PDB contains appears.

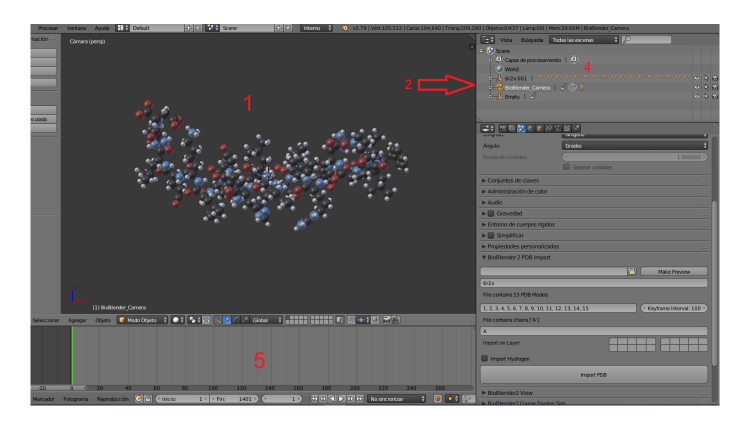
- 6. Where says **Import on Layer**, you select in which layer (layer) of Blender you want to import the molecule.
- 7. Also BioBlender gives you the possibility to import or not, the hydrogens.
- 8. Finally press the click **Import PDB**. The time it takes to import the PDB is proportional to the size of the molecule. If you want to represent a large molecule, the process will be delayed, since the import behaves exponentially.



Once the PDB file is imported, you will be able to see:

- 1. The molecule represented on the scene.
- 2. The list of items that were created during your import.
- 3. In the list of elements you will find a Camera, which is located on layer 20.
 - If you press the key **Num 0**. You will be able to know what the camera is observing.
 - \bullet If you press the key **N** you can modify its location, you can rotate the camera and other options that Blender allows.
- 4. We can also find an element that contains the name of the imported PDB, which has as children (within it) all the imported atoms.

5. You can see the frames, which represent the animation of changing from one model to another in the PDB. In this example, the PDB contains 15 models and the frames reach 1401, this is because the first model is in frame 1, then the second is in frame 101, and so on until the end of the models is reached. (In this example an interval of 100 frames was chosen)

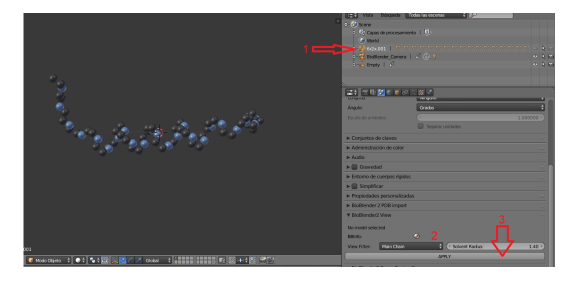


Change the views of the molecule

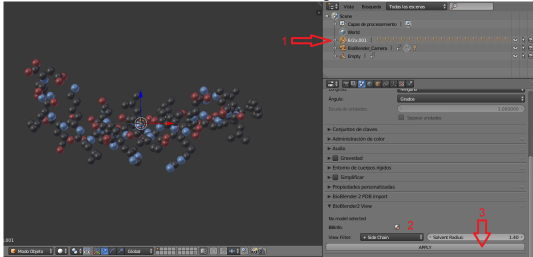
Once the PDB is imported you can change the view of the atoms of the molecule, for this BioBlender has several options:

Note: First you have to select the atoms you want to work with

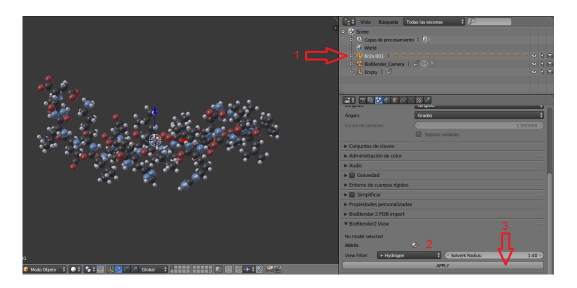
• Main Chain: Show the main chain.



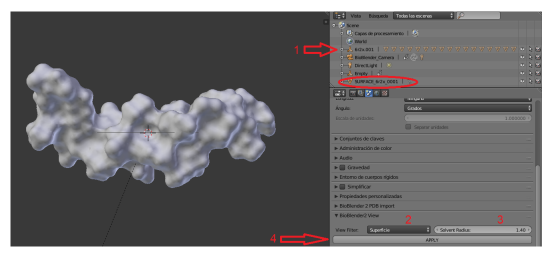
• + Side Chain: Shows the main chain plus the atoms attached to it.



• + **Hydrogen**: Shows all imported atoms plus hydrogens. (This only happens if you imported the hydrogens at the start)



- Surface: Calculate the surface (through the external PyMol program) and show it on the scene. This option counts the property on the right Solvent Radius, which can be modified to calculate the surface of the molecule.
 - The property **Solvent Radius** property represents the radius of the water and defaults to 1.40. If this value is higher, you can calculate a good surface in less time.
 - Once the surface is calculated, an element is automatically added to the scene.
 This has the name: SURFACE, followed by the name of the PDB and then the frame number in which it is located.
 - If you have already calculated a PDB surface in the same number of frames, if you calculate the surface again, it will not calculate it because it will find the element already created.

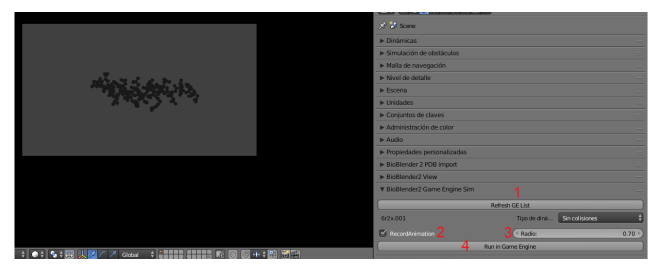


Molecule motion using Blender's physical engine

By importing the molecule into the scene and pressing $\mathbf{Alt} + \mathbf{A}$ the animation of the molecule begins to play. In this case, only the atoms are moving from one location in one conformation to the other, but without taking into account their physical properties, therefore in order to calculate the transition of the protein between two conformations where its properties are taken into account, it is necessary to use the Blender Physics Engine.

Steps to run the Game Engine:

- 1. Click on **Refresh GE List**. This will automatically cause imported molecules to be selected and the name that identifies the molecule will be displayed below this button and the type of dynamic representation to its right (Default is **No Collisions**)
- 2. **Record Animation**: When selected, it allows recording the animation in F-Curves.
- 3. Radius: Represents the radius value for collisions in Blender's physical engine.
- 4. Click on **Run in Game Engine**. This operation once started can be canceled by pressing the key. **Esc**

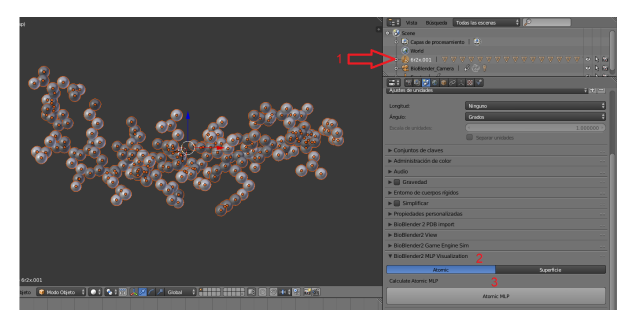


Calculate and visualize the Lipophilic Potential of the molecule (MLP)

BioBlender has the possibility of calculating the MLP, for this it is necessary to use the external Pymol software, which performs the calculations and then they are imported into the Blender.

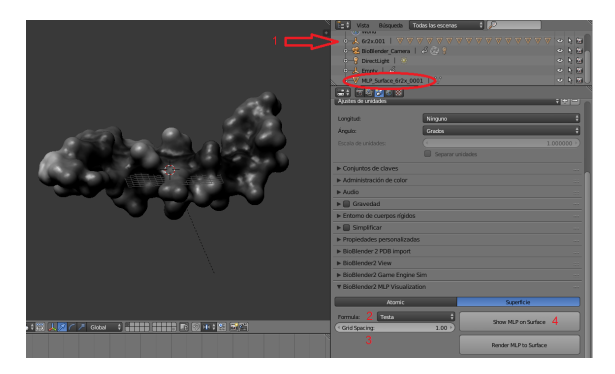
Bioblender has two options:

• Calculate the atomic MLP: To do this you first have to select the atoms in the list of elements. (In this case the use of the external Pymol software is not necessary)

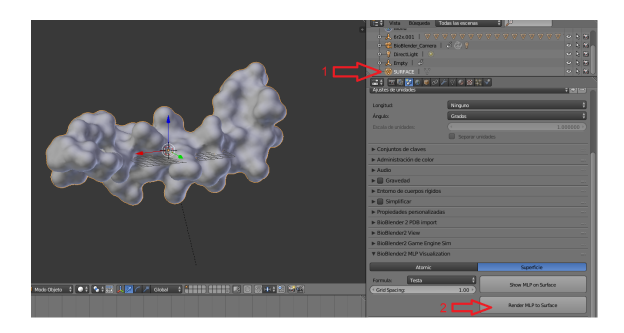


- Calculate the MLP on the surface: PyMLP is used to represent the surface, which obtains the values calculated by PyMol. For its correct operation it is necessary to follow a list of steps.
 - 1. Select the atoms in the item list.
 - 2. If you had previously created a surface through the options of the different views of the molecule, delete it.
 - 3. Select the formula by which to calculate the MLP
 - 4. Select the grid spacing.

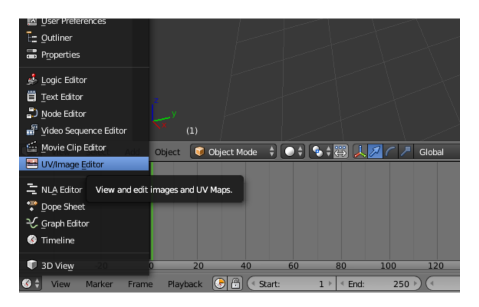
5. Click on Show MLP on Surface.



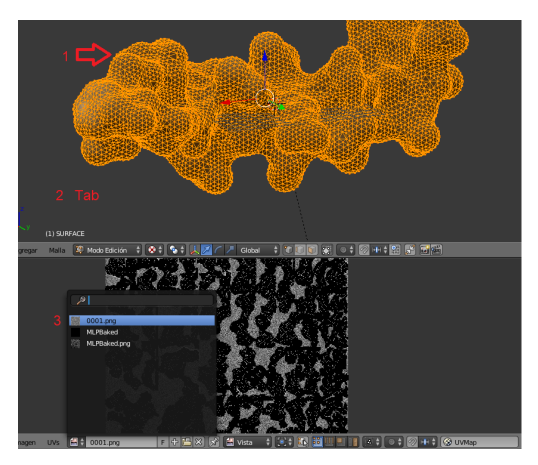
- 6. Once the surface is displayed, an element is automatically added to the scene. This has as name: MLP_Surface, followed by the name that identifies the PDB in the list of elements and then the number where the frame is located.
- 7. If you already calculated a PDB surface in the same number of frames, if you calculate the surface again, it will not calculate it because it will find the element already created.
- 8. Select the created surface.
- 9. Click on Render MLP to Surface



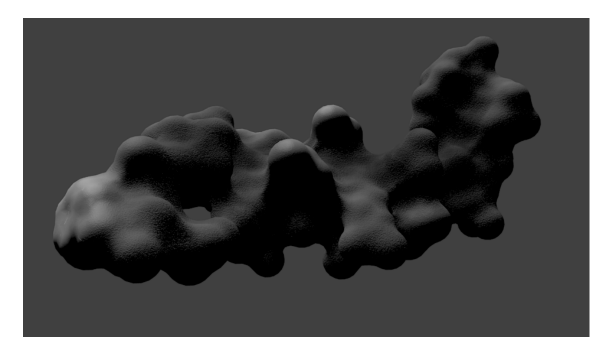
10. Once the texture is created go to **UV** and **Image Editor**



- 11. Select the surface and press the key **Tab**. Automatically all the vertices of the surface were rendered in the UV Editor.
- 12. Select the image 001.png



 $13.\,$ The molecule already presents the surface with the MLP and its texture, ready for rendering.



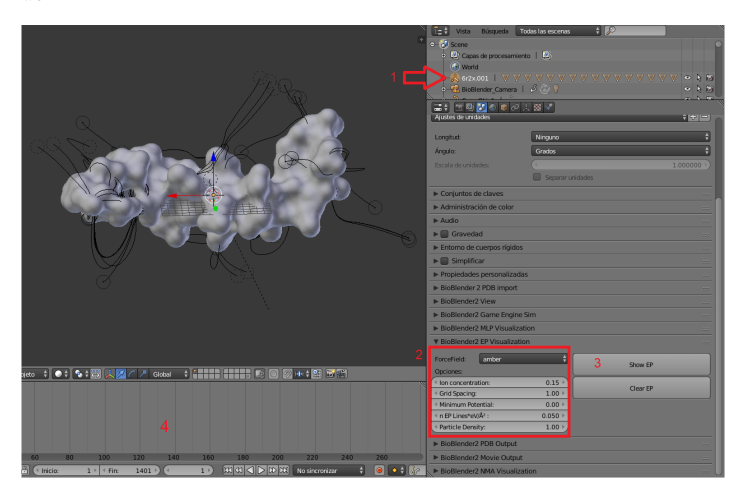
Calculate and visualize the Electrostatic Potential (EP)

BioBlender has the possibility of calculating the electrostatic potential of the molecule and showing its potential lines on the scene. For this BioBlender needs the use of external software **apbs** for carrying out calculations of potential and energy; and the software **SCIVIS** to calculate the lines and export them in a .txt file that BioBlender will later read and make possible their visualization on the scene.

Steps to calculate the EP

- 1. Select the atoms you want to work with.
- 2. This function has various properties which can be modified according to what you want to work with.
 - ForceField: It allows the selection of the force fields with which you want to work.
 - Ion concentration: Allows you to choose the ion concentration of the solvent. Default is 0.15
 - Grid Spacing: Allows you to choose the grid spacing. Smaller is much better, but the calculation is slower.
 - Minimum Potential: It allows choosing the minimum potential from which the lines of the force field start in absolute value.
 - n EP Lines: Represents the concentration of the lines.
 - Particle Density: Represents the density of the particles.
- 3. Click on **Show EP** to calculate the potential.
- 4. Once the EP is calculated, the lines shown in the scene present animation, which you can see since when calculating the EP, the animation automatically plays. (These only go up to approximately 270 frames)

Note: If you want to remove the potential lines from the scene you can click on **Clear EP**, which will remove all the lines not only from the scene, but from the item list as well.

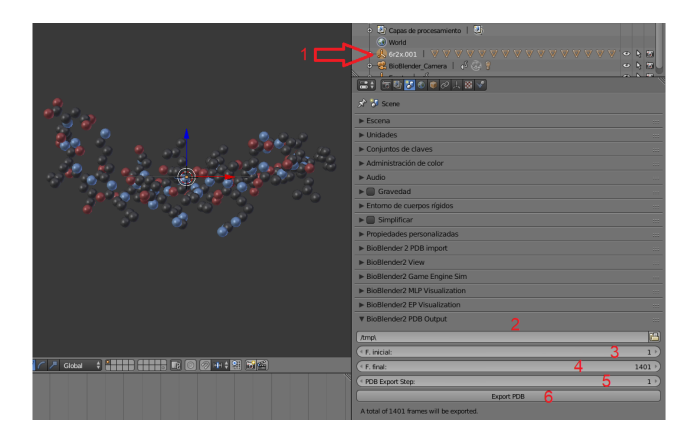


Export PDB file

BioBlender gives the user the possibility of exporting a PDB file according to the location of the molecule in the scene, that is, when exporting the PDB file, it will take the position of all the atoms for each frame of the animation, therefore Keep in mind that if you have 1000 as the final frame, the PDB file generated by the BioBlender will contain 1000 models, since one model is created for each frame.

Steps to export a new PDB file:

- 1. Select the atoms you want to work with.
- 2. Search local storage for the address where you want to save the PDB file.
- 3. Select the starting frame from which you want to start obtaining the position of the atoms to generate your PDB file.
- 4. Select the final frame where you want to stop getting the position of the atoms to generate your PDB file.
- 5. Select the jumps between the frames, by default it is 1. This means that it will increase the frames by 1 until reaching the end of the selected frames.
- 6. Click on Export PDB

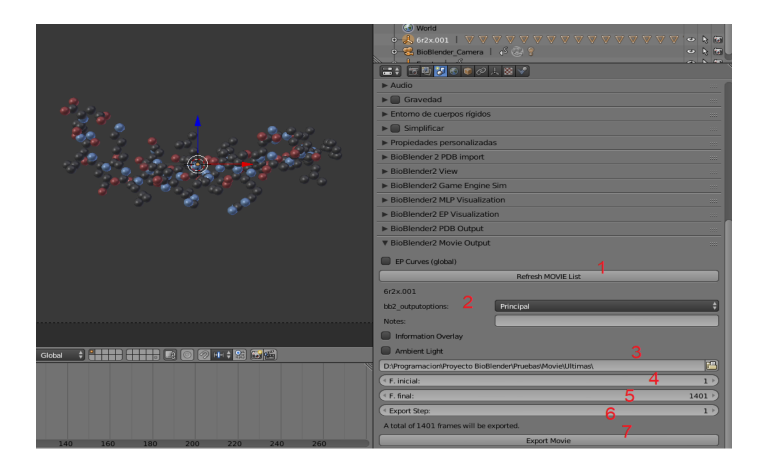


Export Movie

BioBlender offers the possibility of exporting all the frames you want, as well as choosing how you want to observe the molecule in these images. This works through the Blender Render.

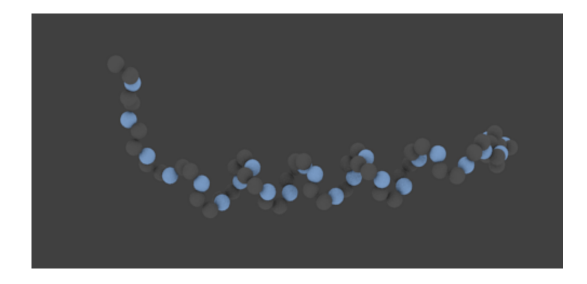
Steps to export the movie:

- 1. Click on Refresh MOVIE List
- 2. Configure the various options it provides:
 - EP Curves (global): When selected, when exporting the molecule in the frame, it also introduces the Electrostatic Potential (EP) into the image.
 - **bb2_outputoptions**: It allows the selection of the view of the molecule that you want to export in the image.
 - Notes: Allows you to add custom text to the image.
 - Information Overlay: Provides detailed information for each frame in the image.
 - Ambient Light: Add light from the environment.
- 3. Select in the local storage the address where you want to save the movie.
- 4. Select the starting frame from which you want to start generating the movie.
- 5. Select the final frame up to where you want to generate the movie.
- 6. Select the jumps between the frames, by default it is 1. This means that it will increase the frames by 1 until reaching the end of the selected frames.
- 7. Click on Export Movie

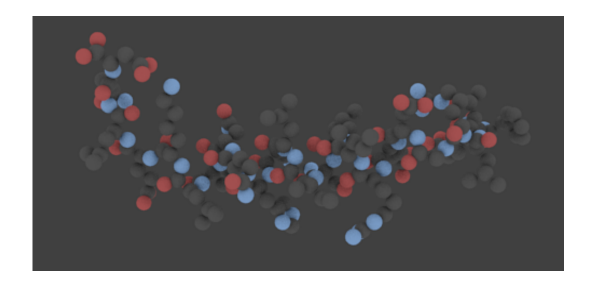


Example of different views of the molecule:

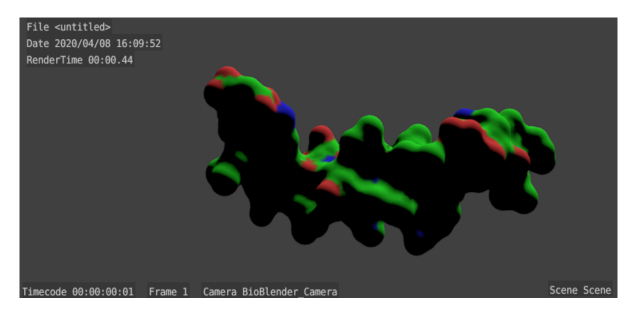
1. View selection **Main** only.



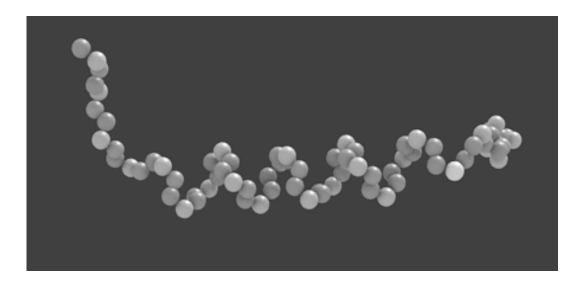
2. Selecting the view + **Side** only.



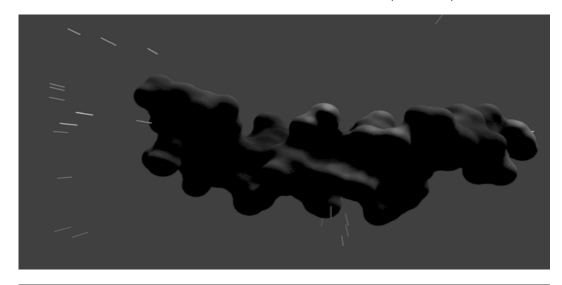
3. Selecting the view **Superficie** and of **Information Overlay**

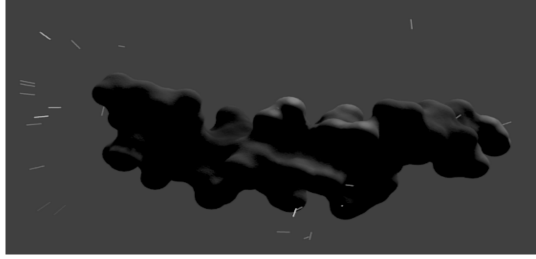


4. Selecting the view MLP Main:



5. Selecting the view MLP Surface and EP Curves (global)





Calculate the NMA

BioBlender has the possibility of calculating the path of the **Normal Mode Analysis**, for this the use of the **Prody** library is required, which allows the analysis to be carried out. The NMA is only calculated for the first model in the PDB file that was imported at startup, so if you only want to calculate the NMA when working with BioBlender, you will not need to import all other models from the PDB.

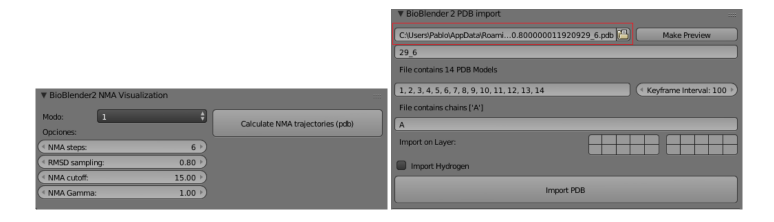
To calculate the NMA, it is not necessary to select the atoms with which you want to work.

Options provided by the calculation of the NMA:

- Modo: Select a normal mode analysis to show.
- NMA Steps: Represents the Number of conformations to be calculated in each direction.
- RMSD sampling: RMSD between the given and the farthest conformation.
- NMA cutoff: NMA cutoff distance (A) for pairwise interactions.
- NMA Gamma: Represents the NMA spring constant

Note: When calculating the NMA a PDB file is exported in the folder

This file contains the calculated path and the address where it is located is automatically entered in the Import function, ready for import.



Important notes for your knowledge

The vast majority of processes in BioBlender export files of vital importance to their operation, which can be very useful for personal use apart from BioBlender. These files are located in the folder:

Functions that generate files in this folder:

ullet BioBlender2 View	: When ca	lculating t	he Surface .:
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- original.pdb
- surface.pml
- tmp.pdb
- tmp.wrl

• BioBlender2 MLP Visualization: When calculating the MLP on the Surface:

- original.pdb
- surface.pml
- tmp.pdb
- tmp.wrl
- tmp.dx

• BioBlender2 MLP Visualization: When calculating the texture of the MLP on the Surface, it generates the necessary images to carry out the texture:

- -0001.png
- MLPBaked.png
- noise.png

- composite.blend This file is typical of Blender, which keeps the composition of the texture with it.

• BioBlender2 EP Visualization: When calculating the EP:

- scenewide.obj
- scenewide.in
- scenewide.pqr
- scenewide.wrl
- scenewide.pdb
- scenewide-input.p
- surface.pml
- -io.mc
- tmp.txt
- pot.dx
- apbs.exe