

# Platomic User Guide

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# **Chapter 1**

## **Quick Start Guide**

### **1.1 Quick overview of Platonic**

Platonic is a graphical application built for Plato capable of quantitatively and qualitatively determining the electrical connectivity between atoms in molecules. It uses Plato's tight binding one script and the Hairy Probe method. This project was submitted in partial fulfilment of the requirements for the MSc degree in Computing Science of Imperial College London (2020-2021).

The source code is available here: <https://github.com/aaronlam123/Platomic> and a video playlist walking-through the program is available here: <https://youtube.com/playlist?list=PLCSLQzY0f0zTrq-B2K8wsCT9rXEFVs00y>.

### **1.2 System requirements**

System requirements for the Platonic Binary:

1. Linux

System requirements for Platonic:

1. Linux
2. OSX

Dependencies (tested on):

1. PyQt5 (5.15.4)
2. pyqtgraph (0.11.1)
3. SciPy (1.5.2)
4. pandas (1.2.1)

5. PyAutoGUI (0.9.52)
6. PyOpenGL (3.1.5)
7. NumPy (1.20.2)

Runs purely on Python 3 (tested on 3.8.5). Requires a copy of Plato for full functionality, which you can ask for permission from @horsfielda [here](#) if you are part of his research group. Once given permission to access Plato, you may contact me for a Platonic binary + Plato binary distribution. This distribution runs out of the box with no package / dependency installation required. As such it is most suitable for non-expert users who are unfamiliar with Linux and / or would like to skip over the installation of a number of packages.

## 1.3 Getting started

To run the Platonic binary, navigate to the dist folder, and open a terminal window via right click. Then enter the command “chmod +x setup” to enable execution permissions for the setup bash. Next, type “./setup” then “./main” and it should start. Only ./main is needed thereafter to open Platonic.

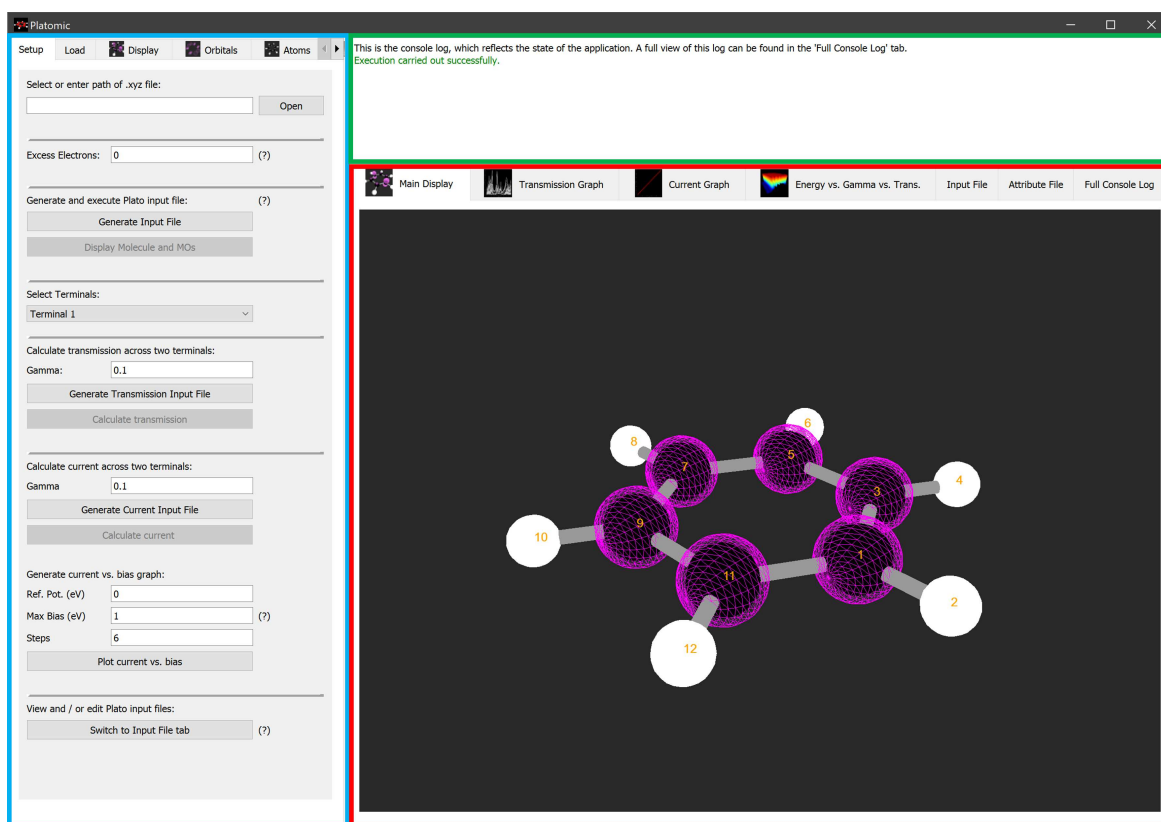
Alternatively, after downloading the repository and the required dependencies you can start Platonic by running main.py.

In both instances a copy of Plato is required to enable full functionality for Platonic. The Plato folder should be located in the same directory as main / main.py.

# Chapter 2

## Using Platomic

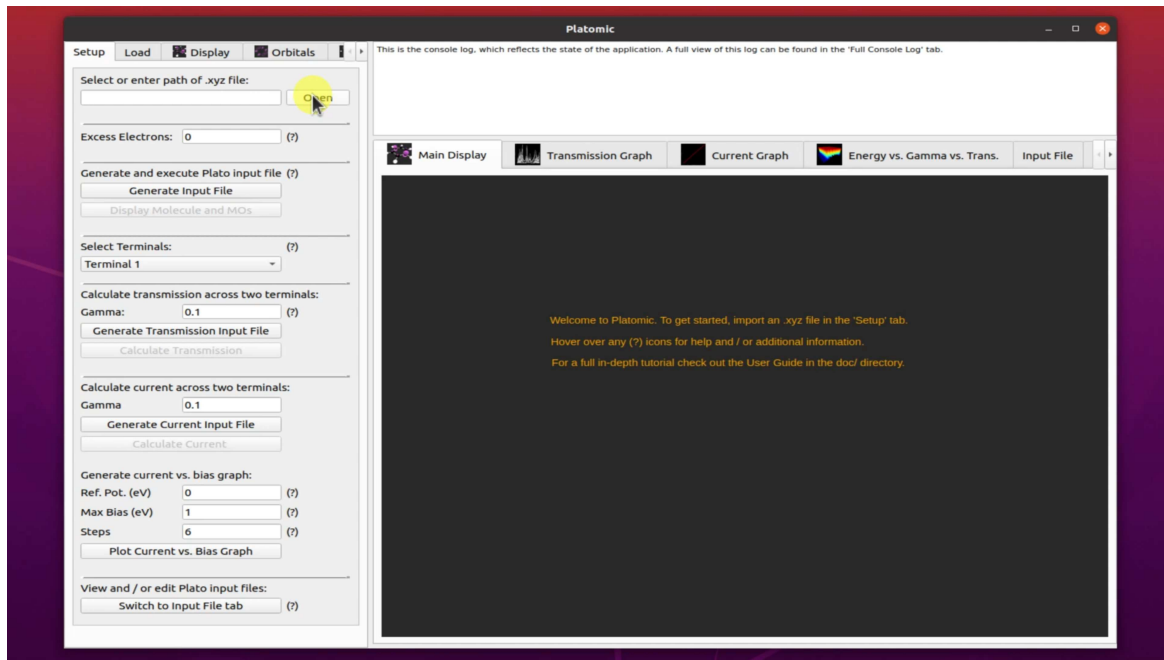
The UI of Platonic consists of three parts, the central window, properties sidebar and console log as seen in Figure 2.1 below. The central window and properties sidebar contain a number of tabs that can be freely switched between. For a brief introduction to each tab, you may refer to the video walk-through linked here: [Section 2.1 - Molecular Orbital Visualisation](#).



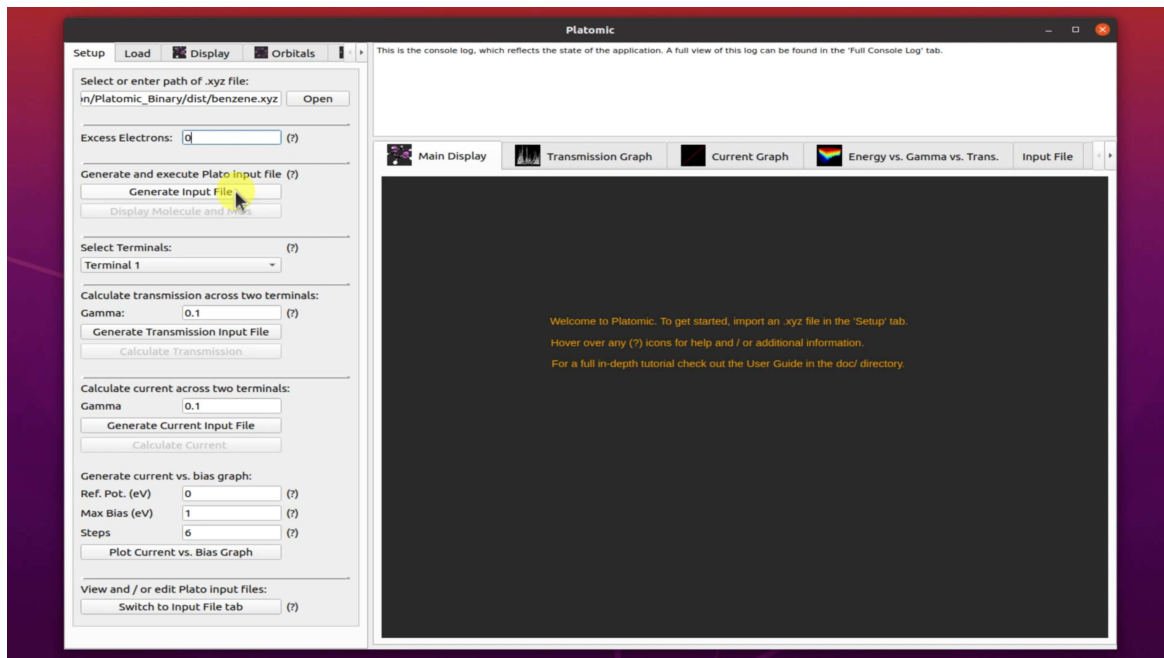
**Figure 2.1:** UI for Platonic. (Red outline) central window, consists of 2D/3D plot and text widgets. (Blue outline) properties sidebar, used to setup the application and change settings. (Green outline) console log, shows console outputs and error messages.

## 2.1 Molecular orbital visualisation

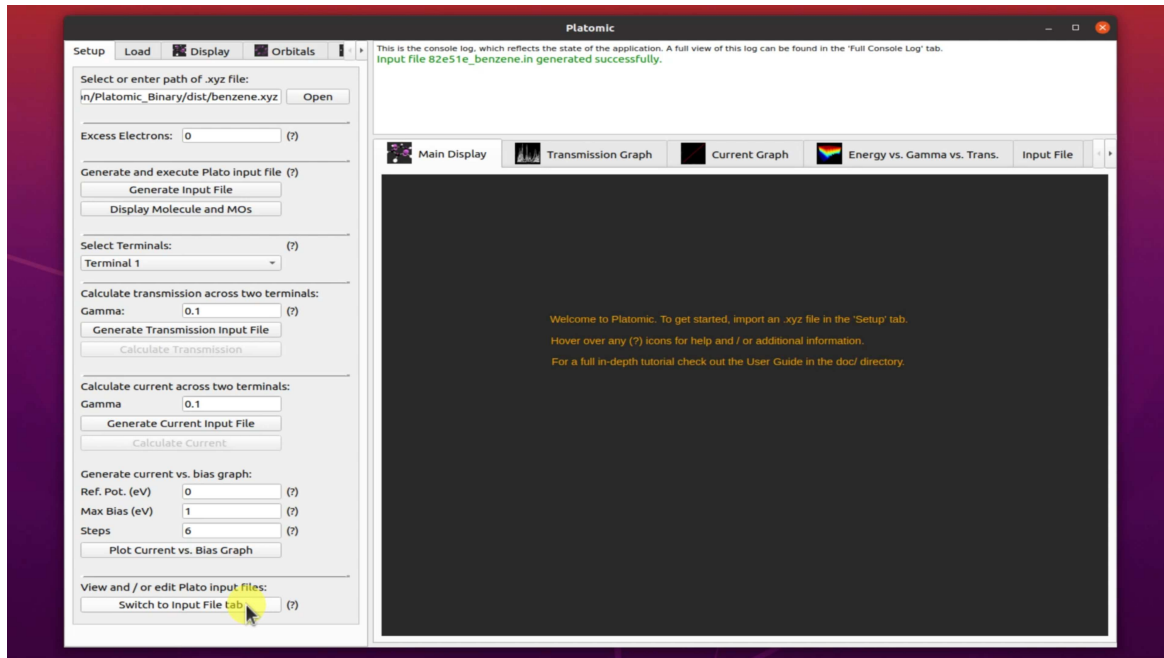
In addition to the walk-through written below, you may refer to the video tutorial linked here: **Section 2.1 - Molecular Orbital Visualisation**.



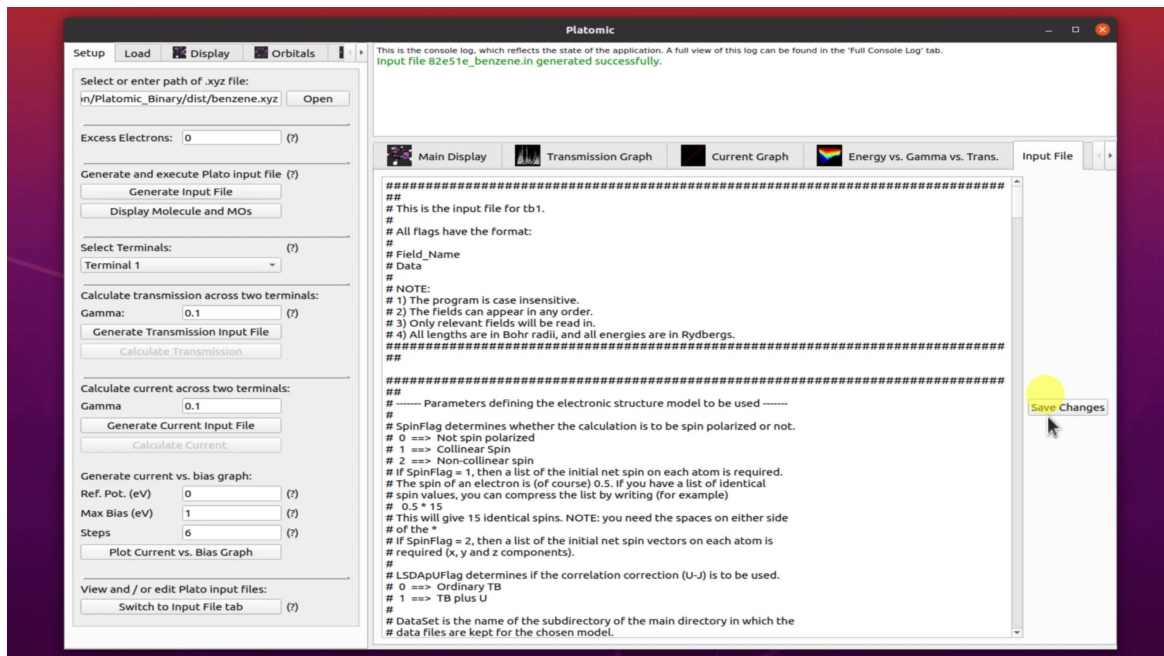
**Figure 2.2:** First load an .xyz file of the desired molecule / molecule system as shown.



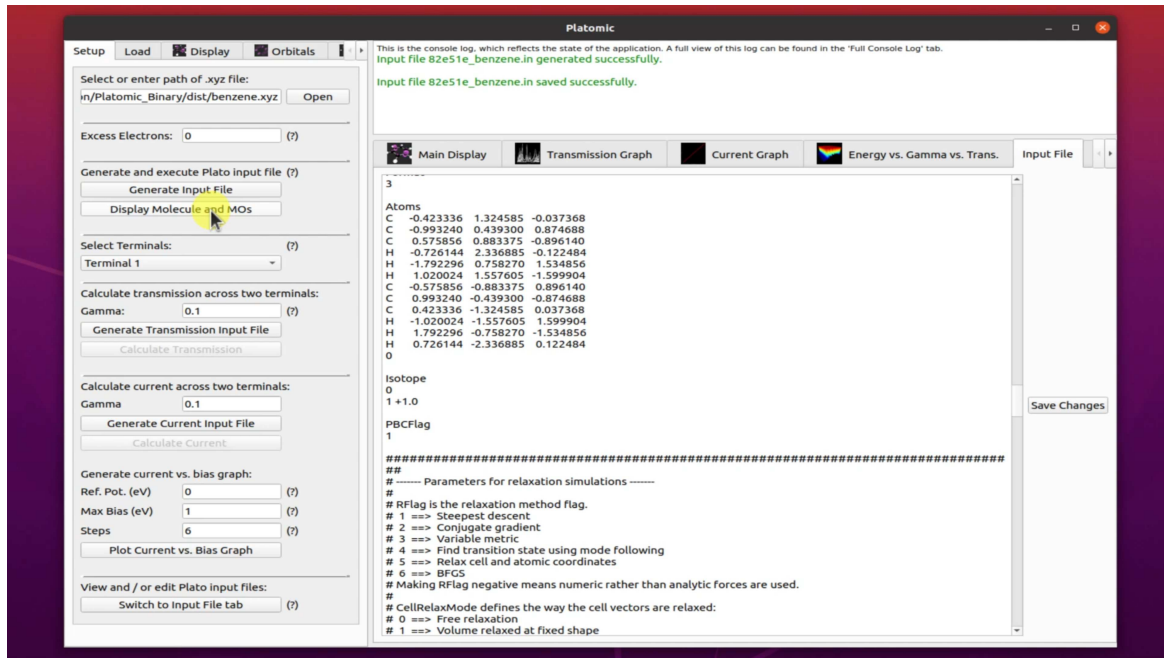
**Figure 2.3:** Next, click the 'Generate Input File' button to generate an input file.



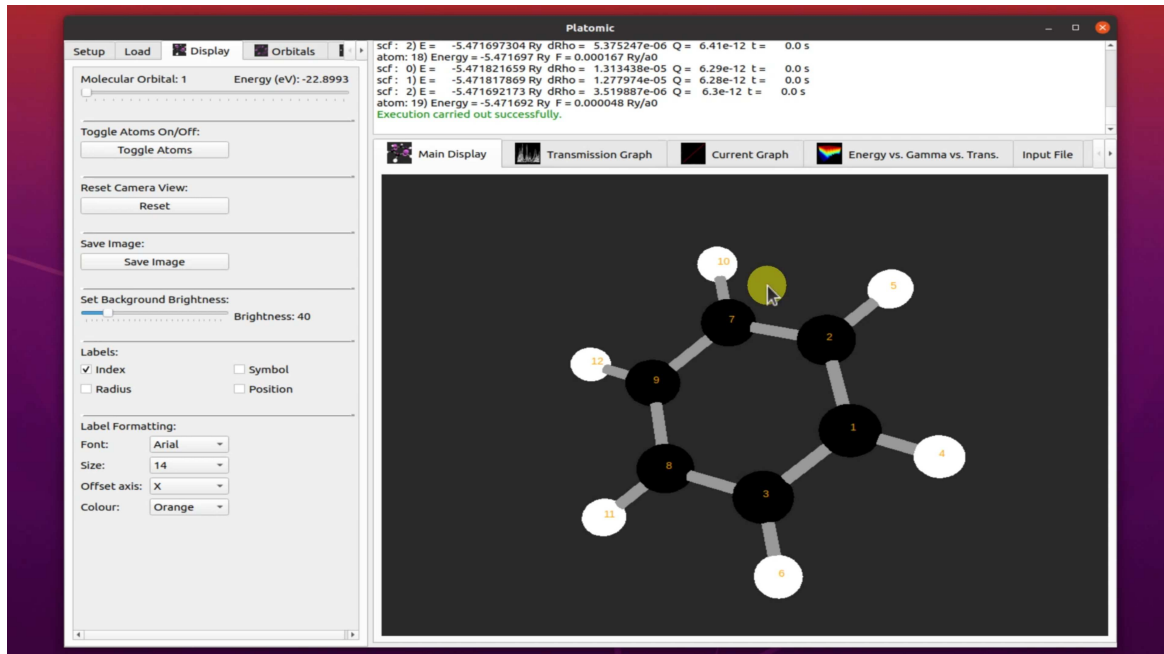
**Figure 2.4:** Optionally edit the input file by switching to the ‘Input File’ tab either by clicking the button or selecting the tab manually on the central window.



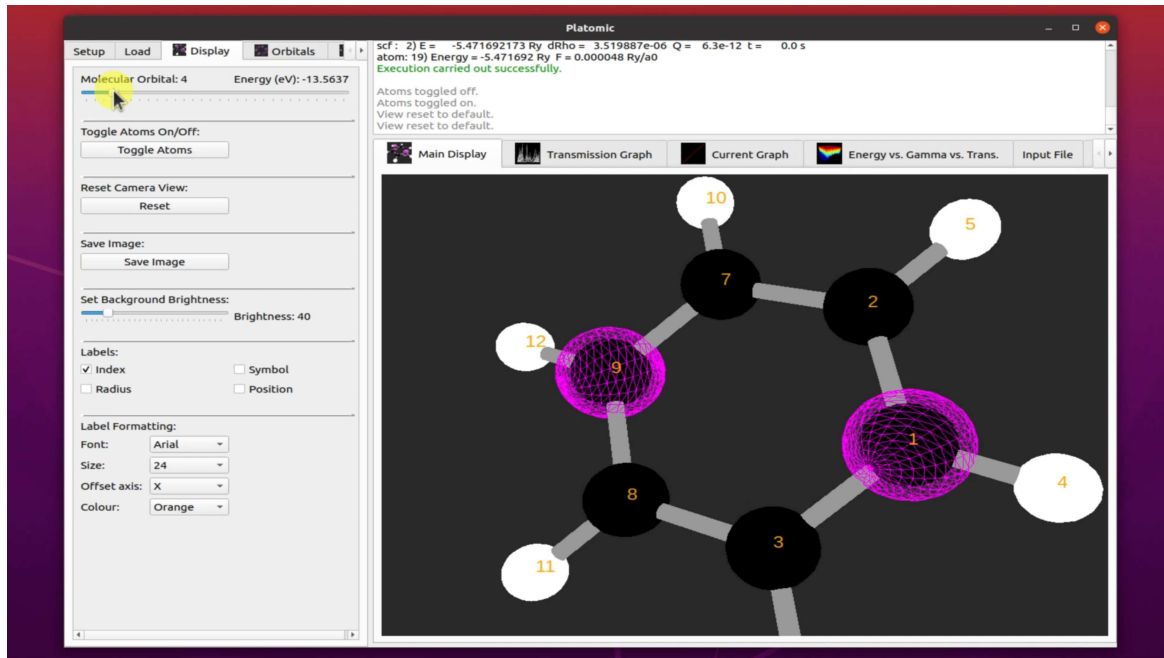
**Figure 2.5:** If you decide to make changes, press the ‘Save Changes’ button. When Plato is executed, it will use the edited input file.



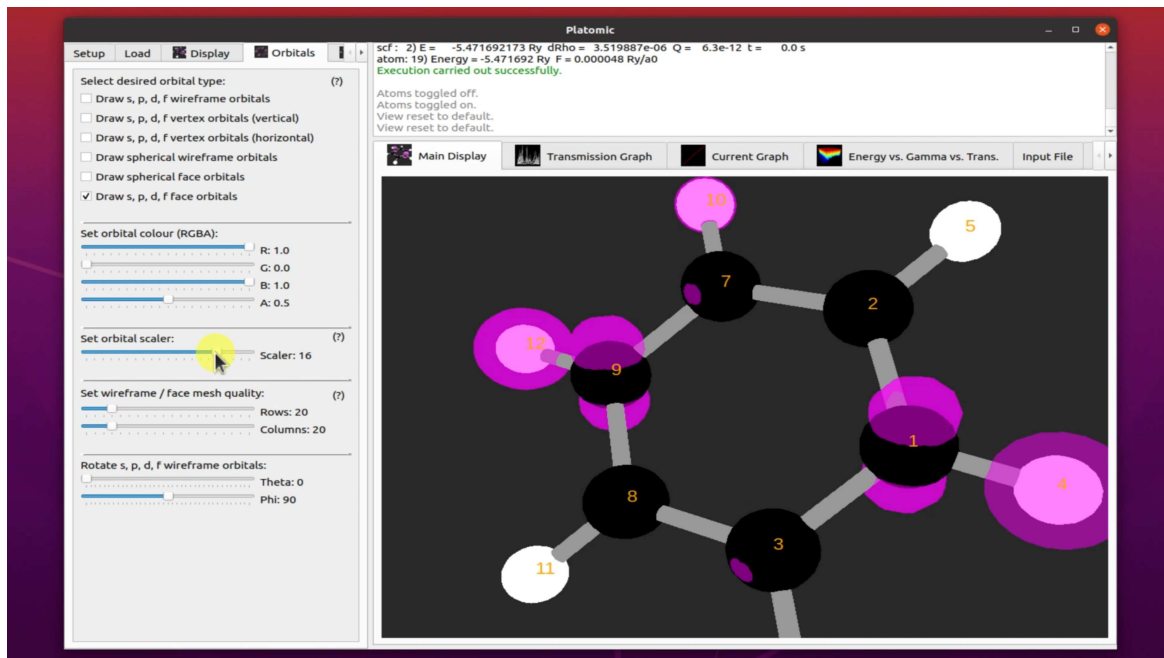
**Figure 2.6:** Finally press the ‘Display Molecule and MOs’ button as shown to run Plato in the back-end on the newly generated input file.



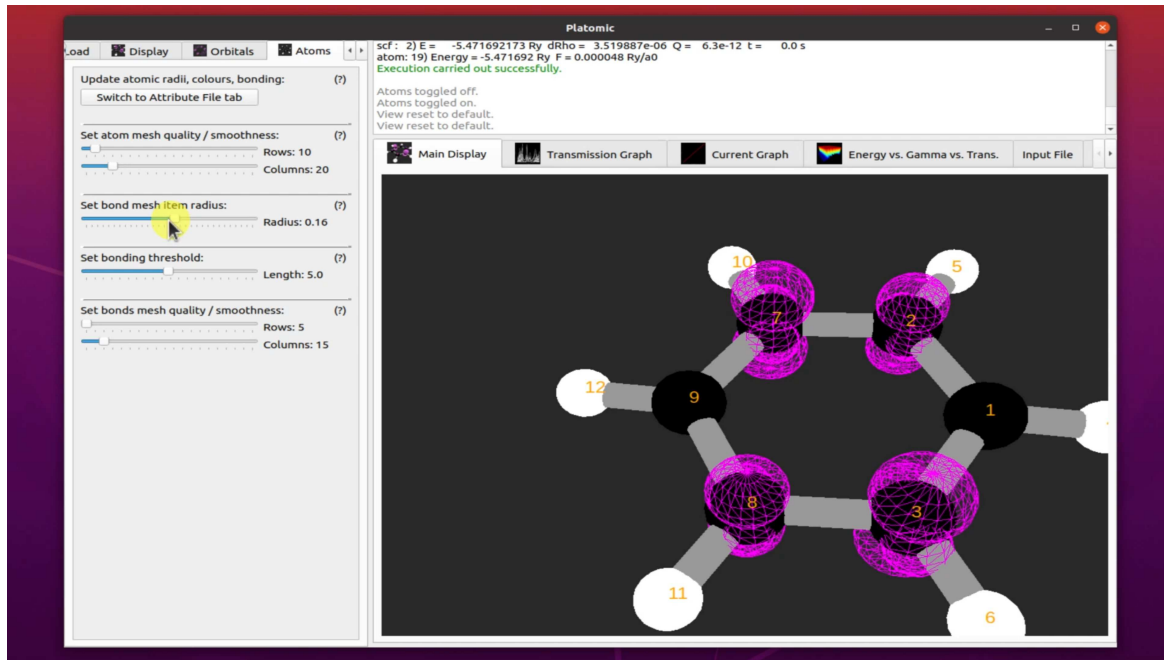
**Figure 2.7:** The central window’s ‘Main Display’ tab should now show the molecule with molecular orbitals. The camera view can be changed by dragging on the widget with the left mouse button. You may also zoom in and out by using the scroll wheel and pan the molecule by dragging while holding the control key.



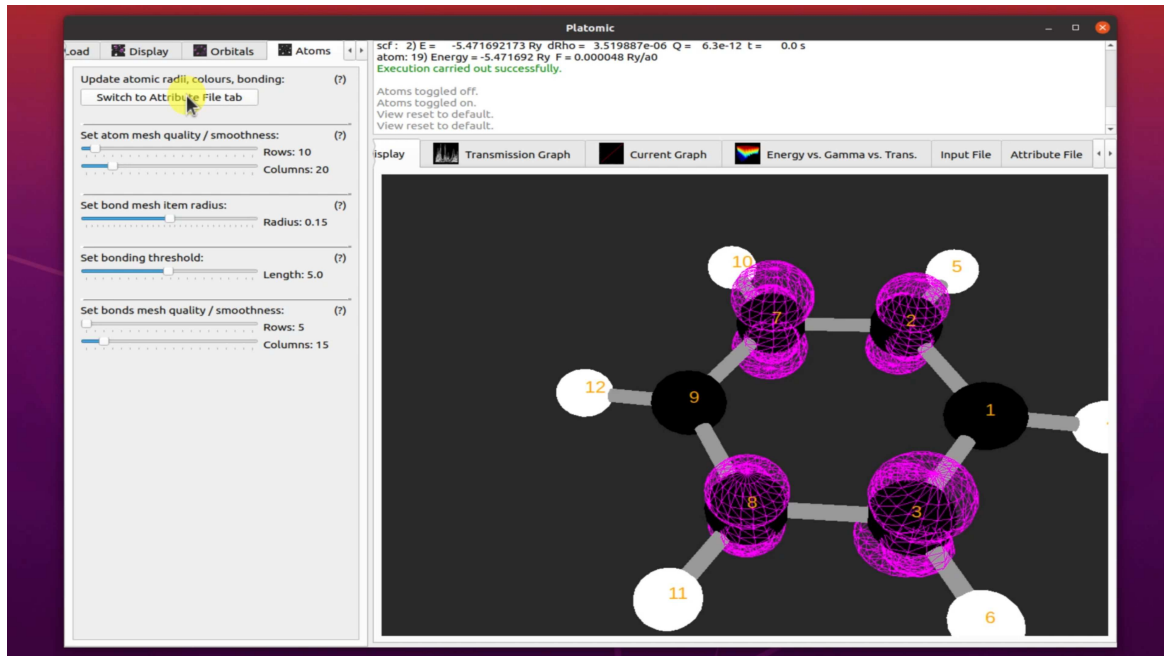
**Figure 2.8:** You may view the different molecular orbitals by dragging the horizontal slider as shown, toggle the atoms on and off, reset the camera view, save the image to disk, change the background brightness and customise labels.



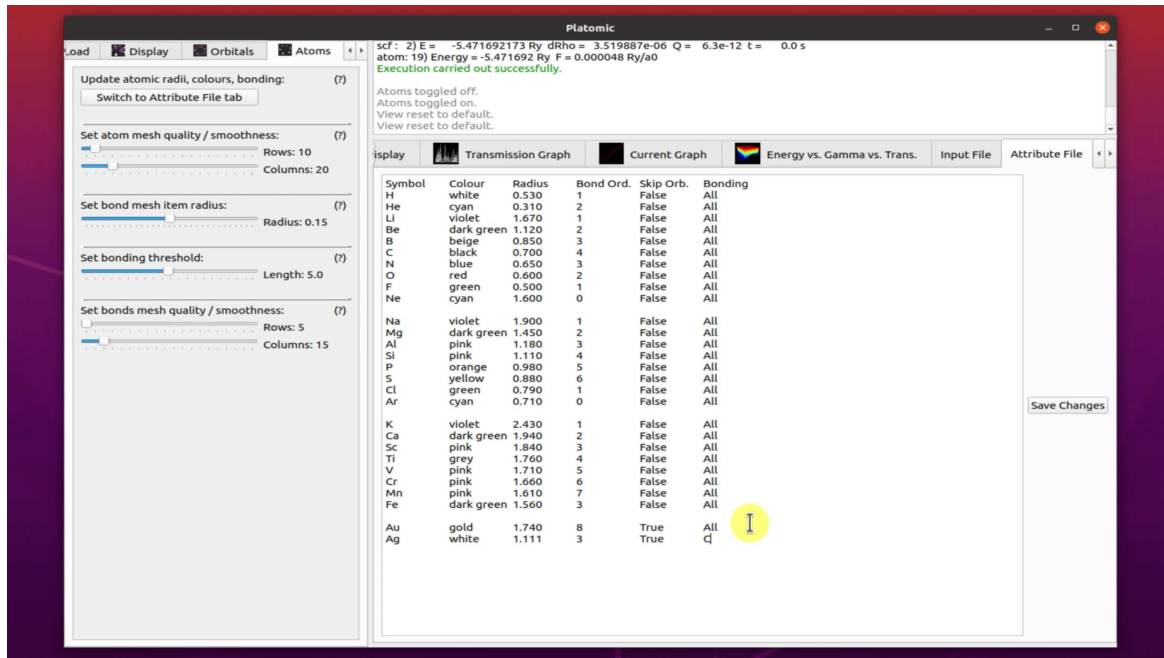
**Figure 2.9:** In the ‘Orbitals’ tab of the properties sidebar, you can change the orbital display type, orbital colour, orbital scaler, orbital mesh item quality and rotate the orbitals.



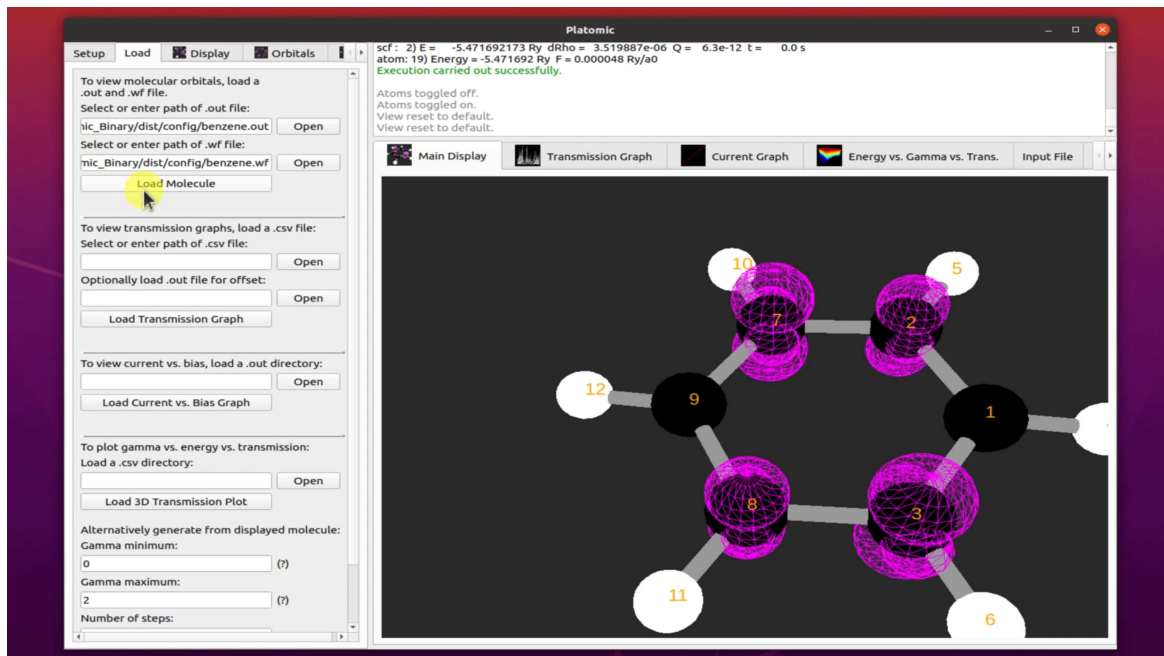
**Figure 2.10:** In the ‘Atoms’ tab of the properties sidebar, you can likewise adjust the atom and bond mesh quality, change the bond mesh item radius and bonding threshold. The bonding threshold allows you to control the upper limit for how long a bond can be before it is skipped.



**Figure 2.11:** You can also edit the attributes file by switching to it manually or clicking the button as shown. These changes affect the default attributes for each element, allowing for properties such as the colour, radius and bonding behaviour to be edited.



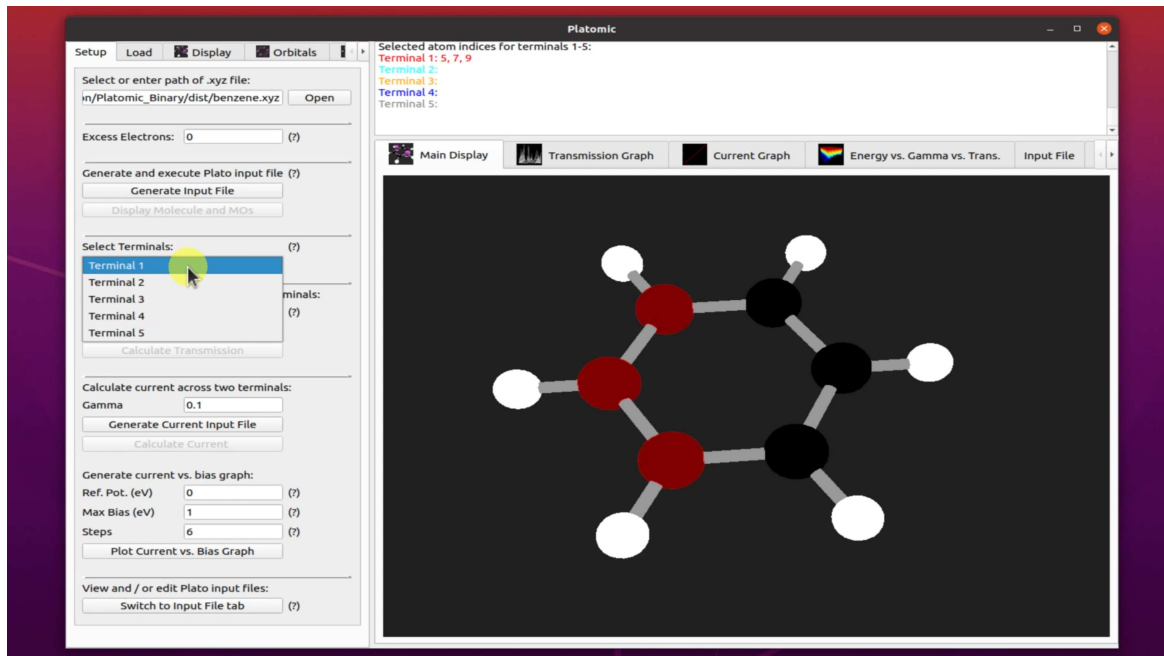
**Figure 2.12:** If an element does not exist, you can add it in a new line as shown for Ag. Changes must be saved by pressing the ‘Save Changes’ button, and will be applied on the next execution.



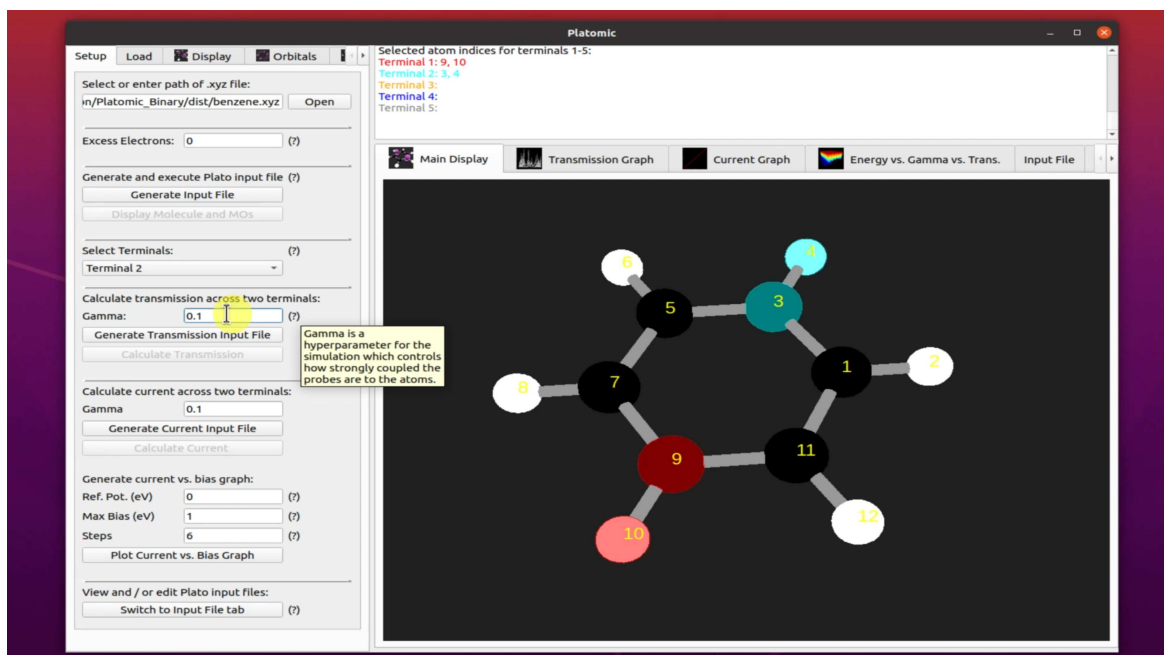
**Figure 2.13:** If you have pre-computed output (.out) and wavefunction files (.wf), you can load them as shown.

## 2.2 2D Transmission graph

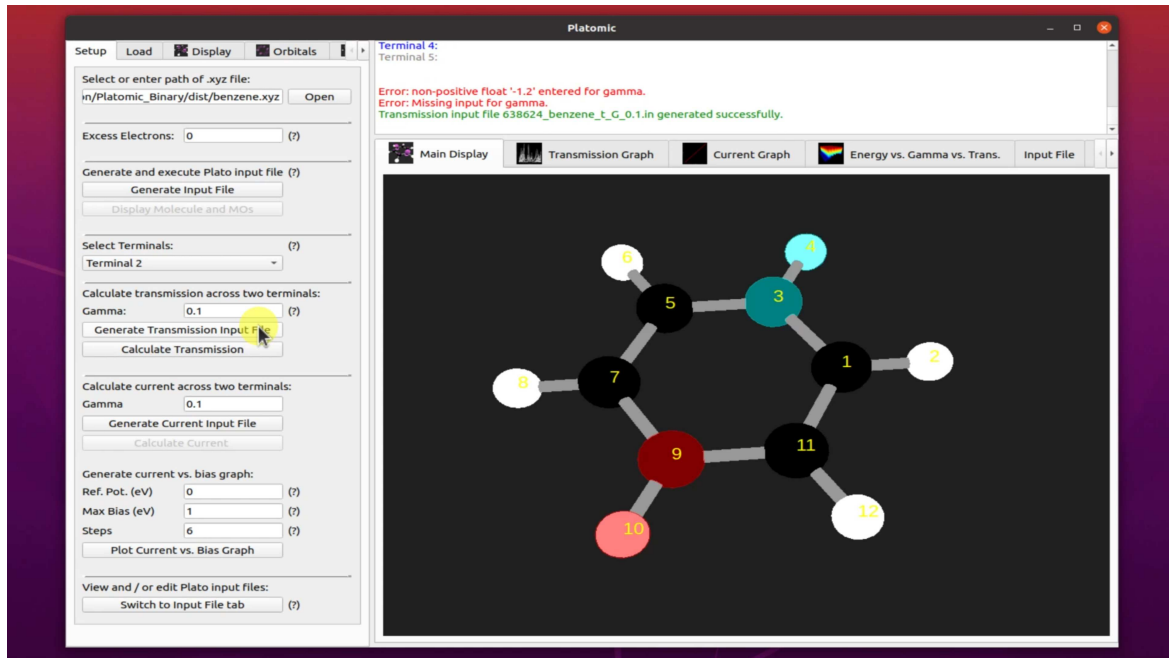
In addition to the walk-through written below, you may refer to the video tutorial linked here: [Section 2.2 - 2D Transmission Graphs](#).



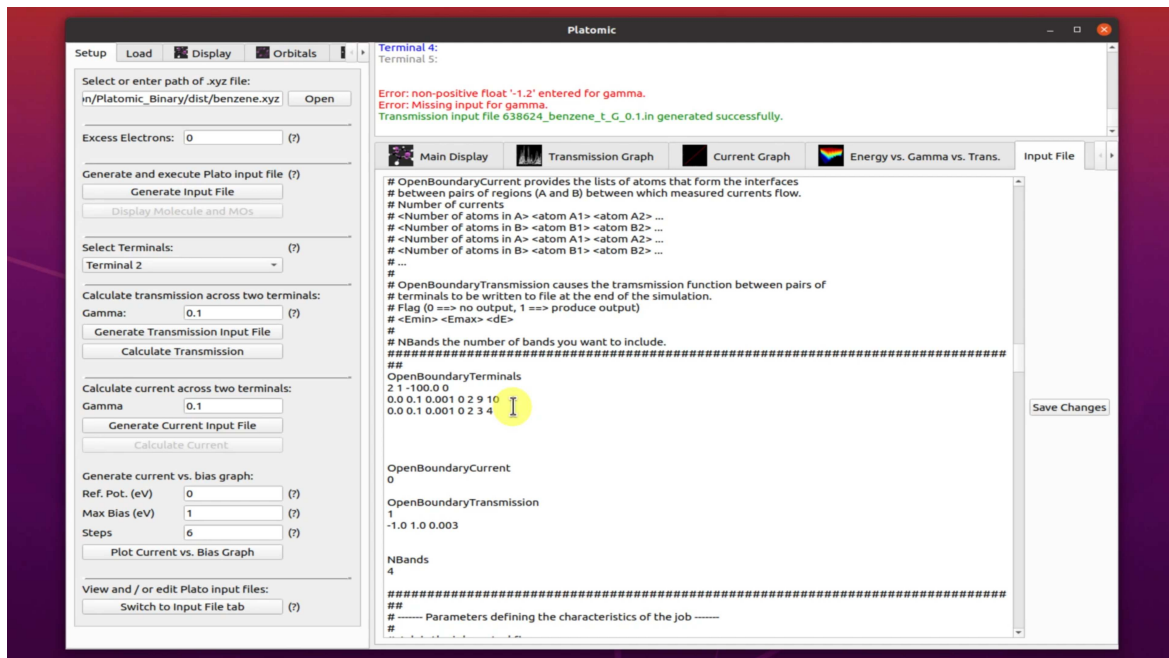
**Figure 2.14:** After a molecule is loaded, you can select / deselect atoms to add / remove them from a terminal by left clicking (terminal changed using the drop down menu).



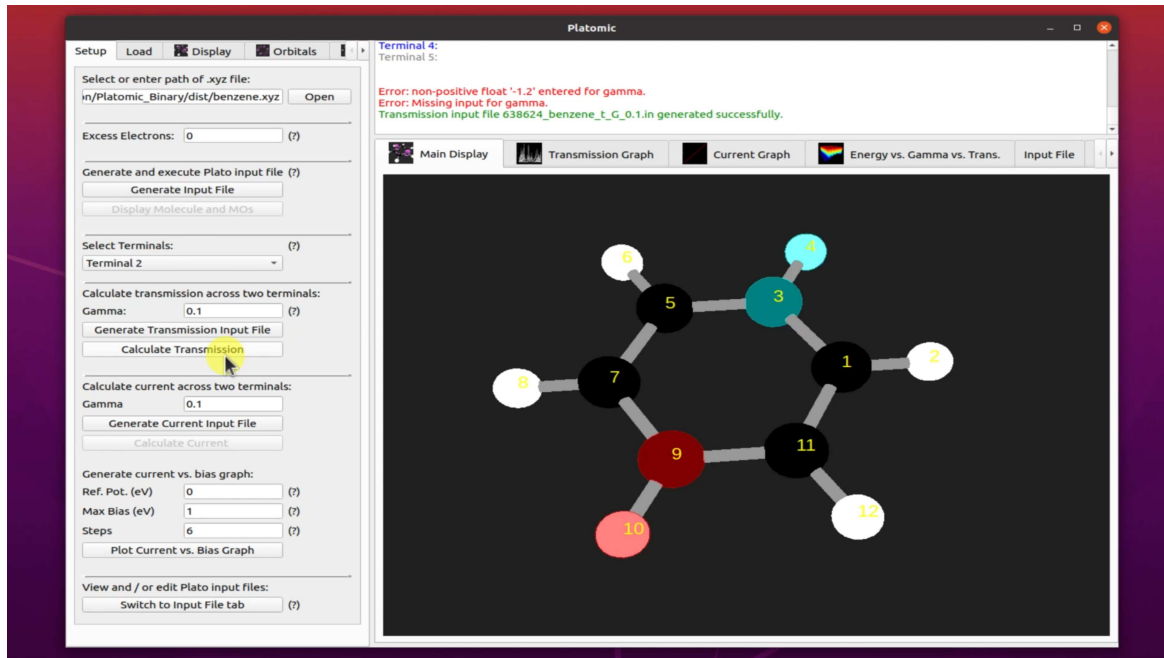
**Figure 2.15:** Here, I have a benzene molecule with two terminals. Terminal one consists of atoms with indices 9 and 10, while terminal two consists of atoms 3 and 4.



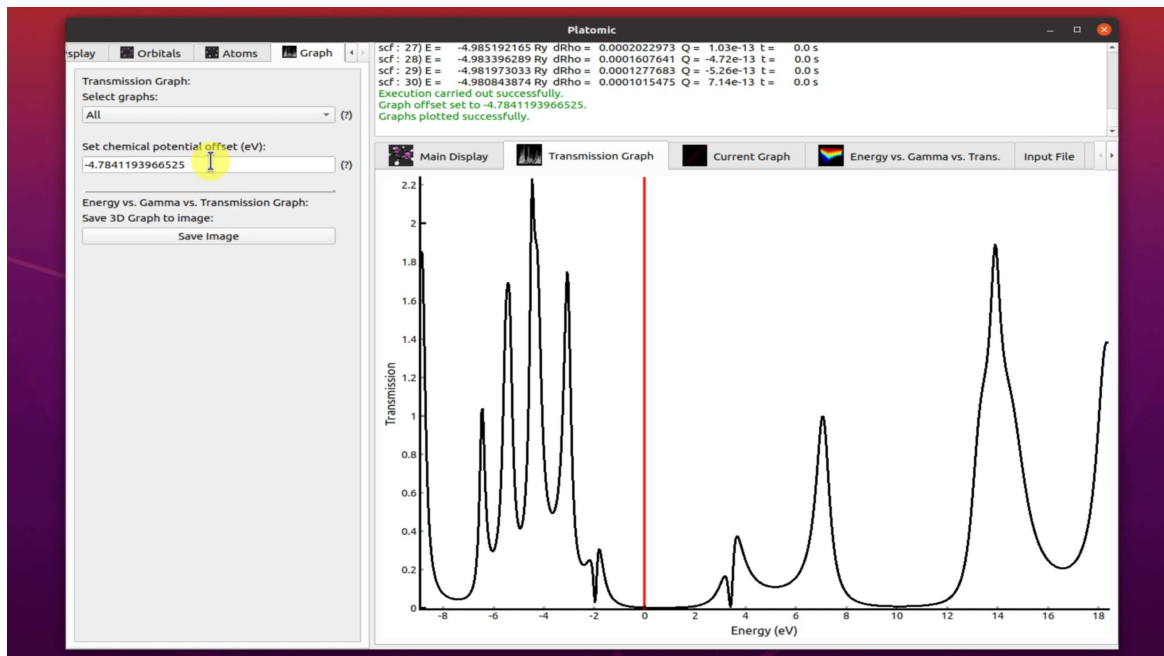
**Figure 2.16:** After specifying the gamma hyperparameter, which controls how strongly coupled probes are to the system, you can generate the transmission input file as shown. Typical gamma values are between zero and one.



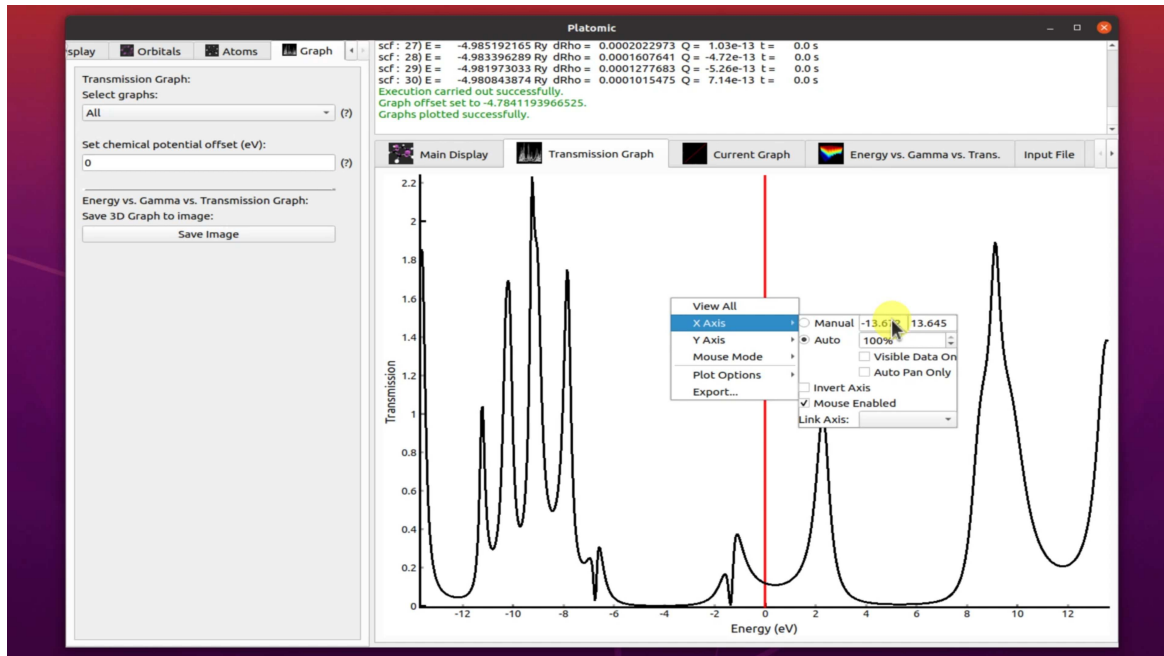
**Figure 2.17:** Like for the molecular orbital visualisation feature in Section 2.1, you can edit the input file in the same way.



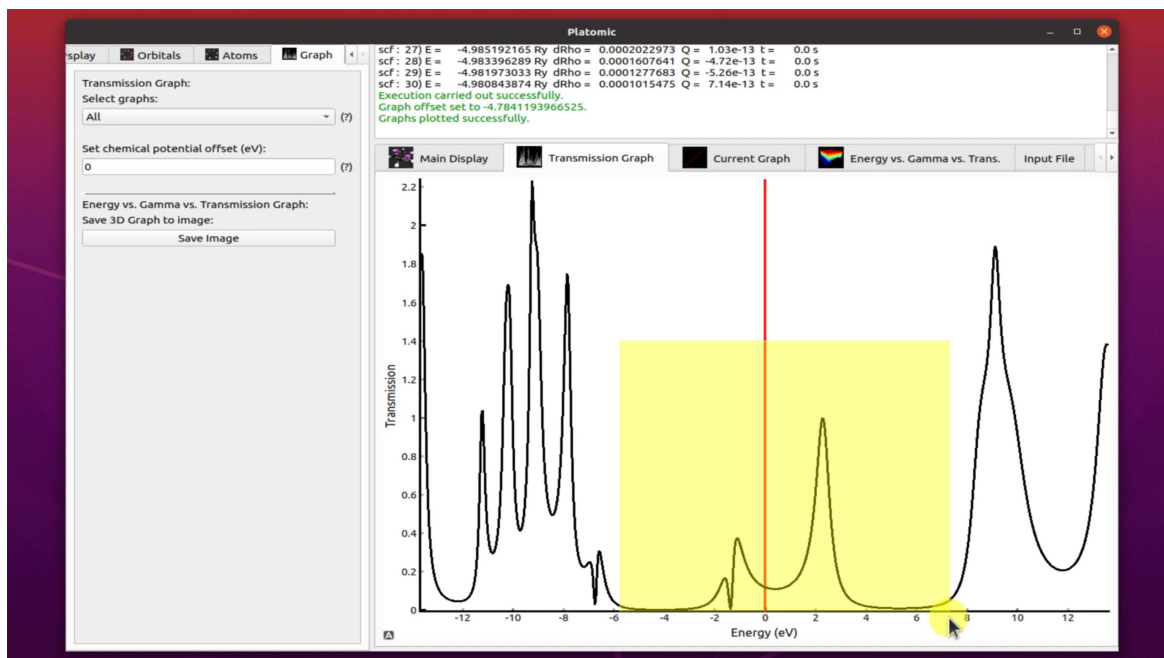
**Figure 2.18:** Finally, to run Plato on the newly generated transmission input file and plot the 2D Transmission graph, you click on the ‘Calculate Transmission’ button as shown.



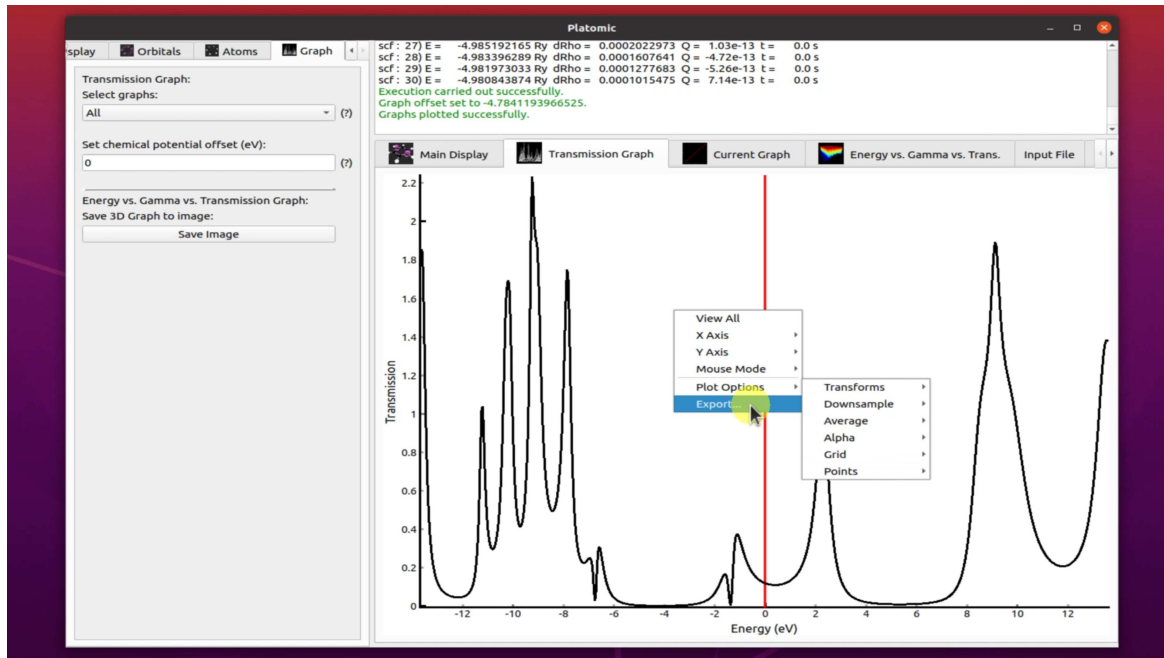
**Figure 2.19:** The average terminal chemical potential is automatically determined from the output file, but you may adjust it manually. Changing this value shifts the position of zero energy.



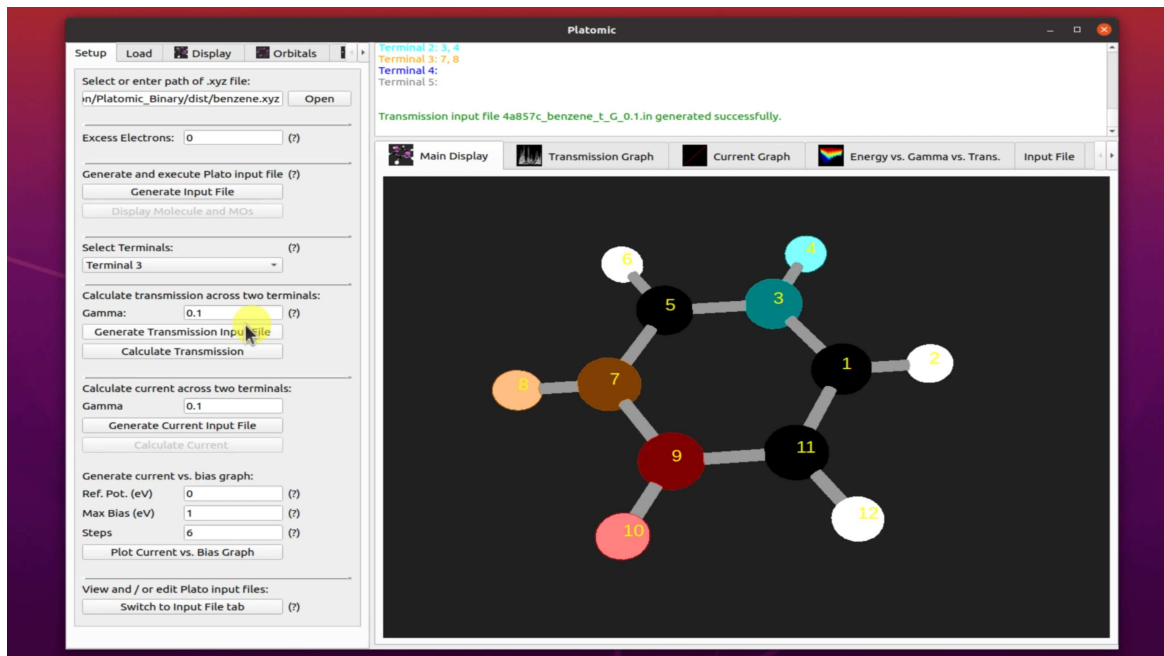
**Figure 2.20:** Right clicking on the graph shows a number of view options, such as changing the x and y axis to specific ranges, enabling grid lines and applying transforms.



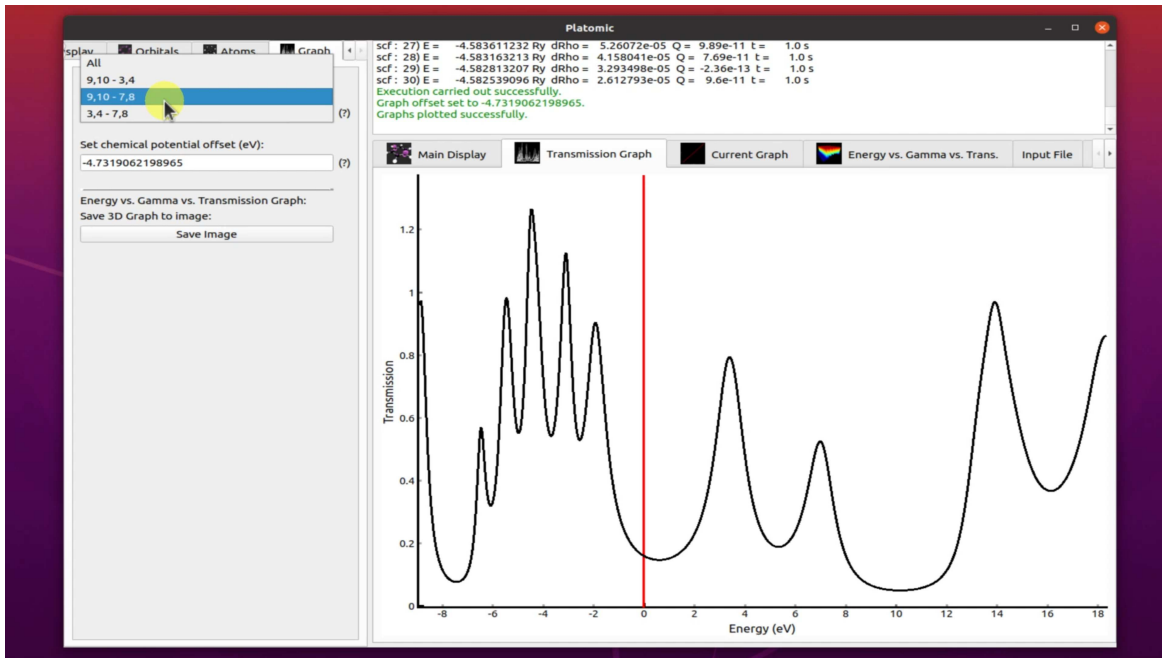
**Figure 2.21:** By changing the mouse mode to 1 button, you can drag a view box around a point of interest and release to automatically resize the graph to the view box.



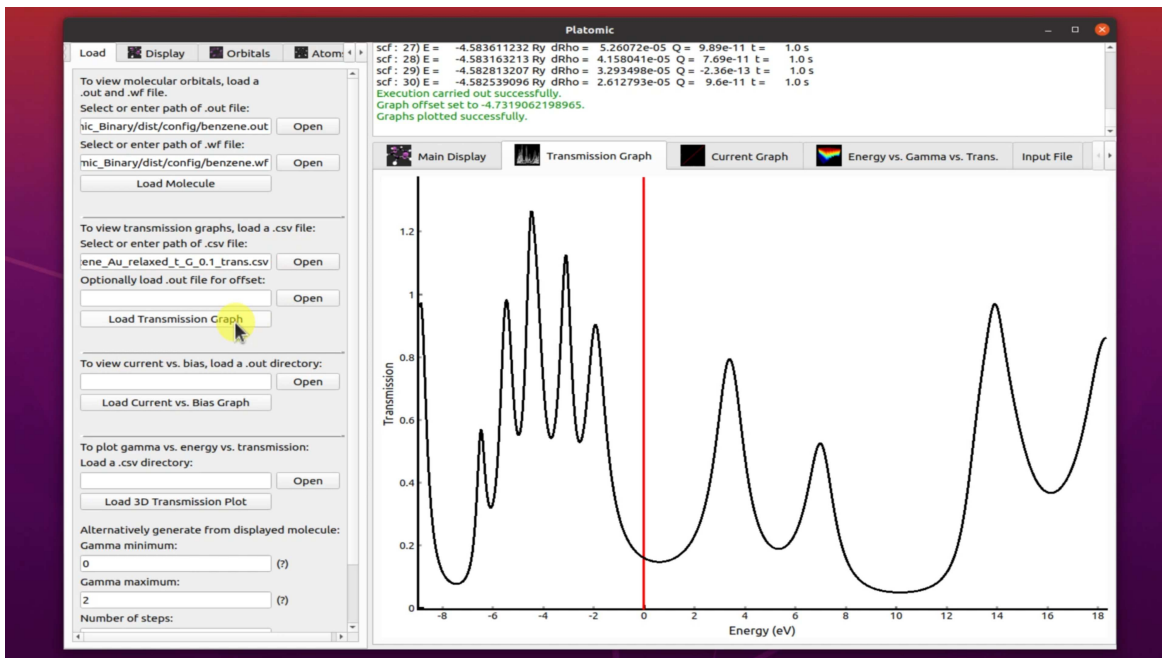
**Figure 2.22:** You can also export the transmission plot to an image file and / or CSV file here.



**Figure 2.23:** When more than two terminals are selected, the ‘Select graphs:’ drop down menu becomes useful (located in the ‘Graph’ tab of the properties sidebar). Here we have a benzene molecule as before with an additional terminal consisting of atoms with indices 7 and 8.



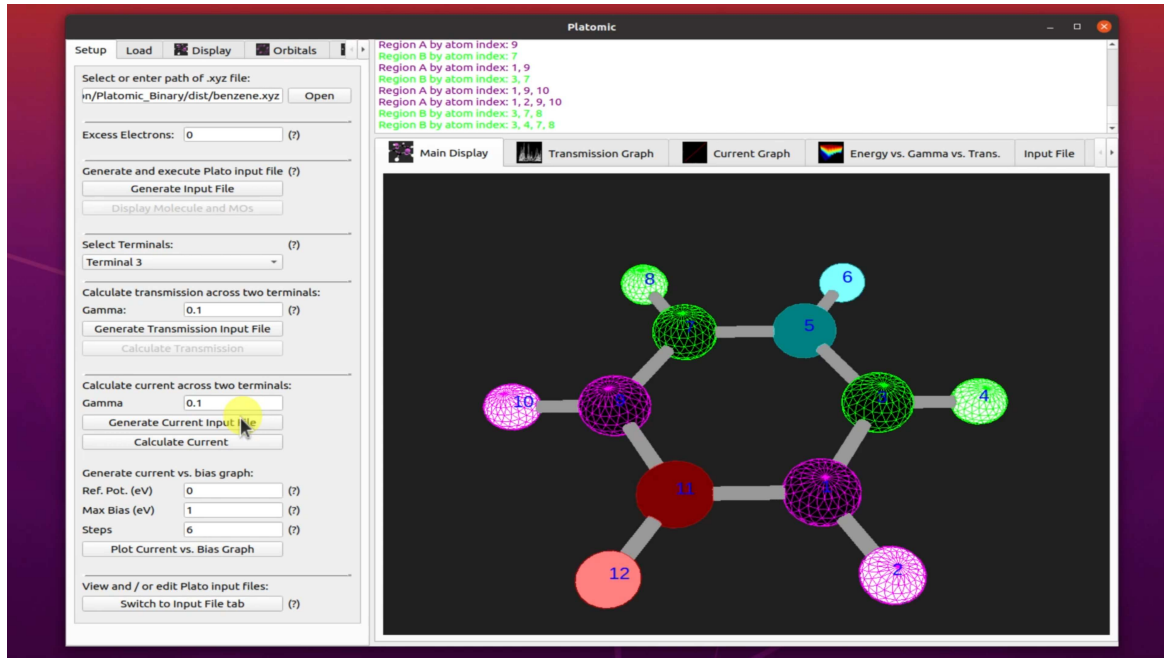
**Figure 2.24:** The ‘Select graphs:’ drop down menu, in addition to showing the indices of atoms which are part of each respective terminal, allows the user to switch between the transmission plots of different terminal pairings.



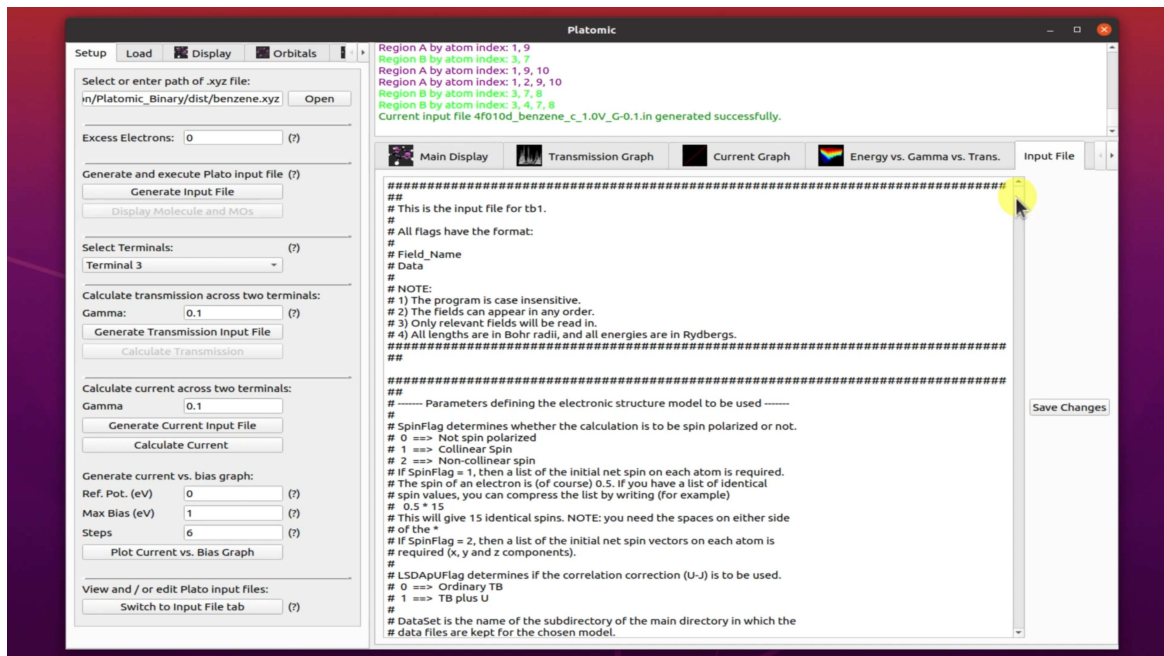
**Figure 2.25:** In the ‘Load’ tab of the properties sidebar, you can load a transmission graph from a pre-computed CSV file and optionally the corresponding .out file to set the terminal chemical potential.

## 2.3 Current calculations and current vs. bias graphs

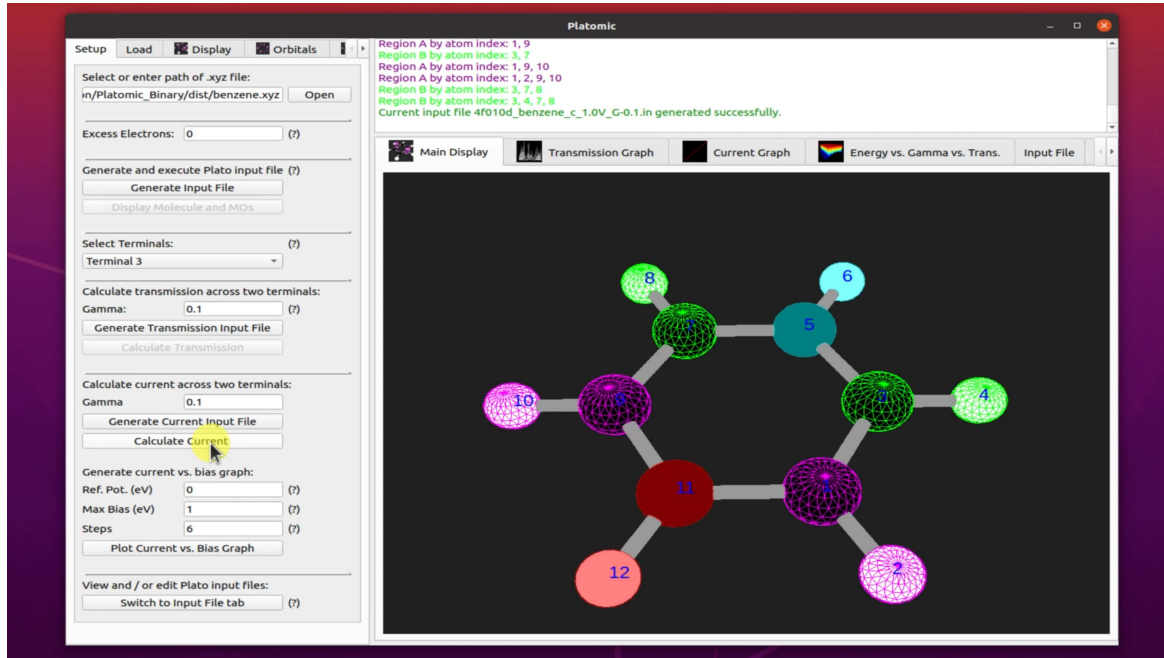
In addition to the walk-through written below, you may refer to the video tutorial linked here: [Section 2.3 - Current Calculations / Current vs. Bias](#).



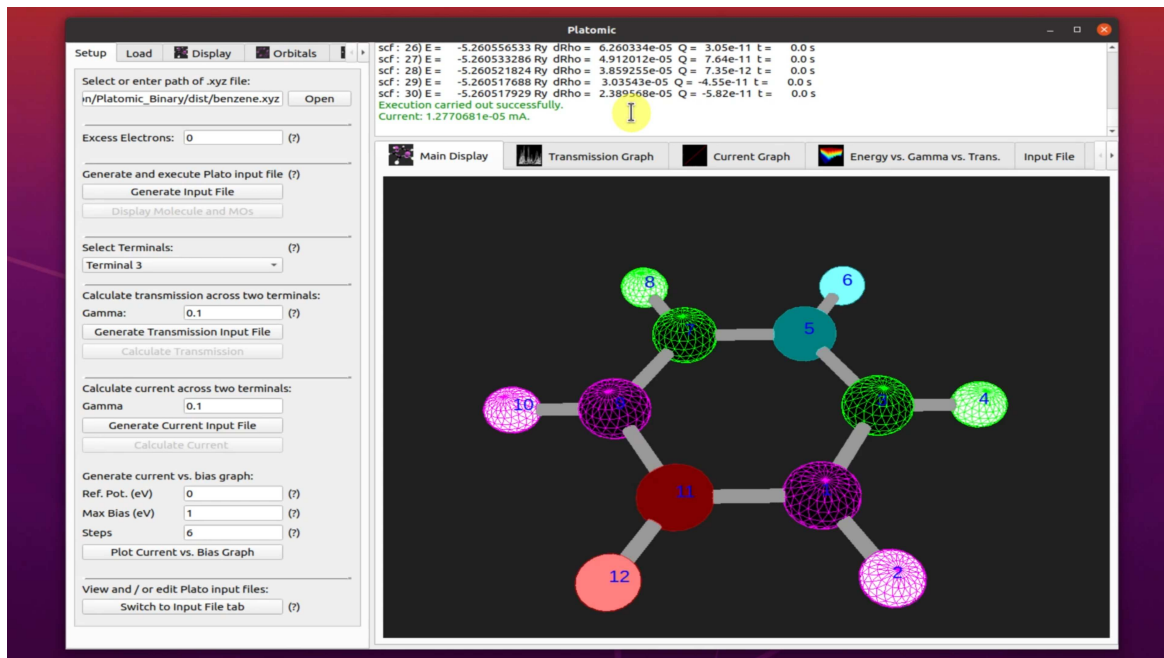
**Figure 2.26:** After loading the molecule, in addition to specifying terminals (see Section 2.2) you must select atoms to add to Regions A (by right clicking) and B (by middle clicking), as current is measured as the flow of change between the two regions.



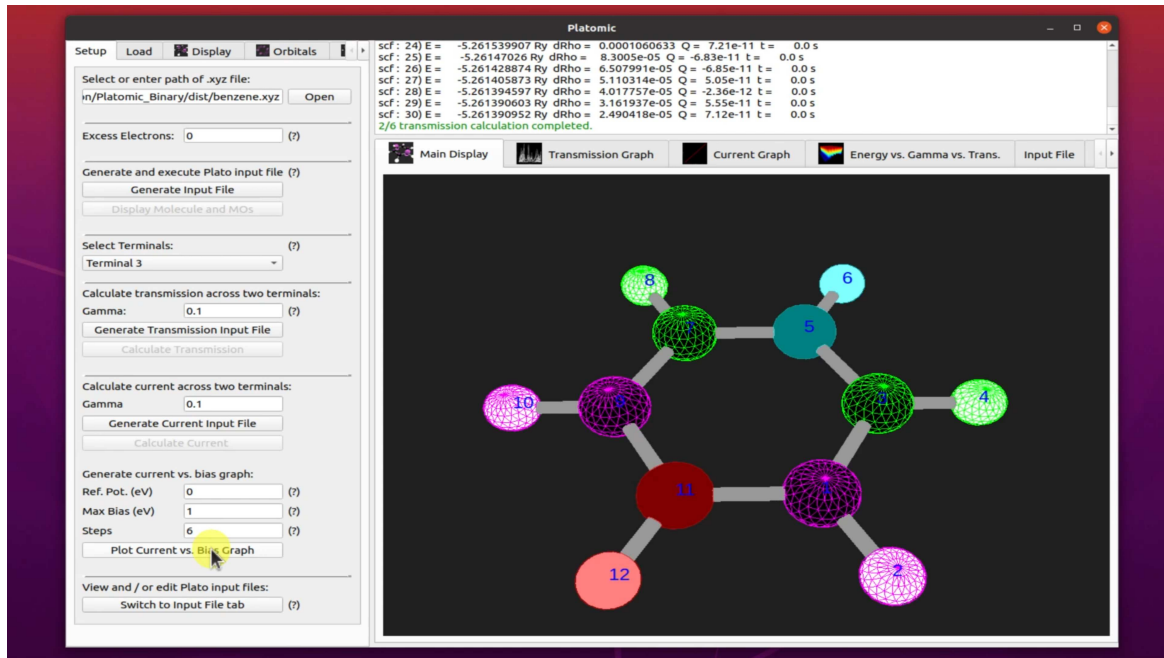
**Figure 2.27:** The current input file can be edited manually as shown if desired.



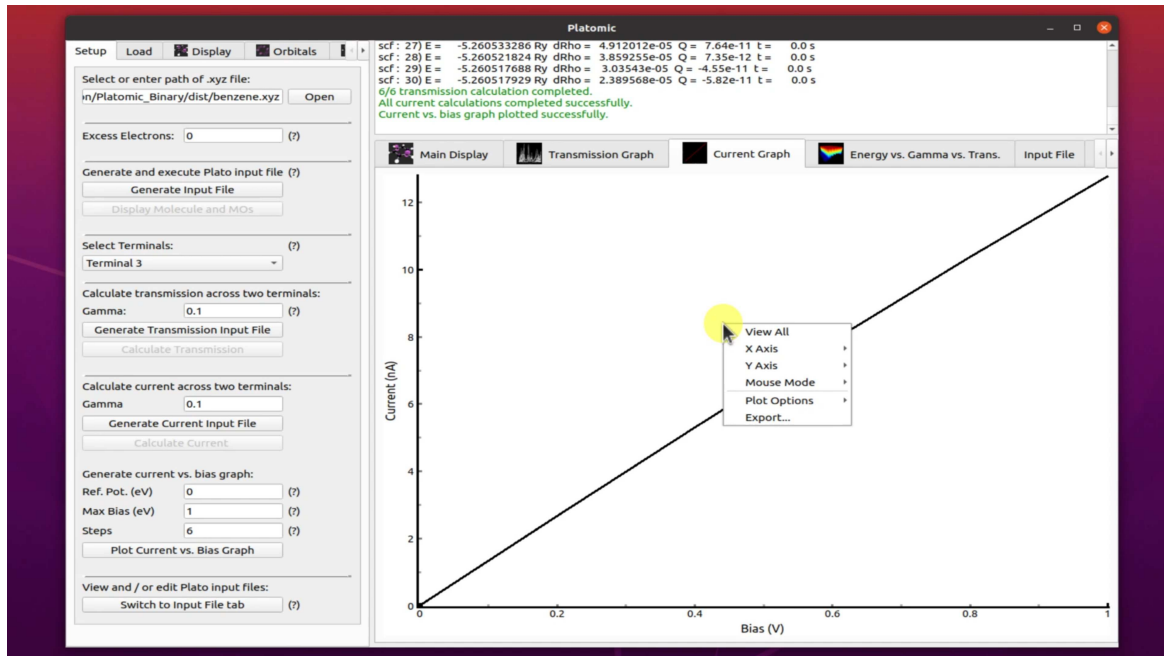
**Figure 2.28:** After a current input file is generated, you can calculate the current between the two terminals by clicking the ‘Calculate Current’ button.



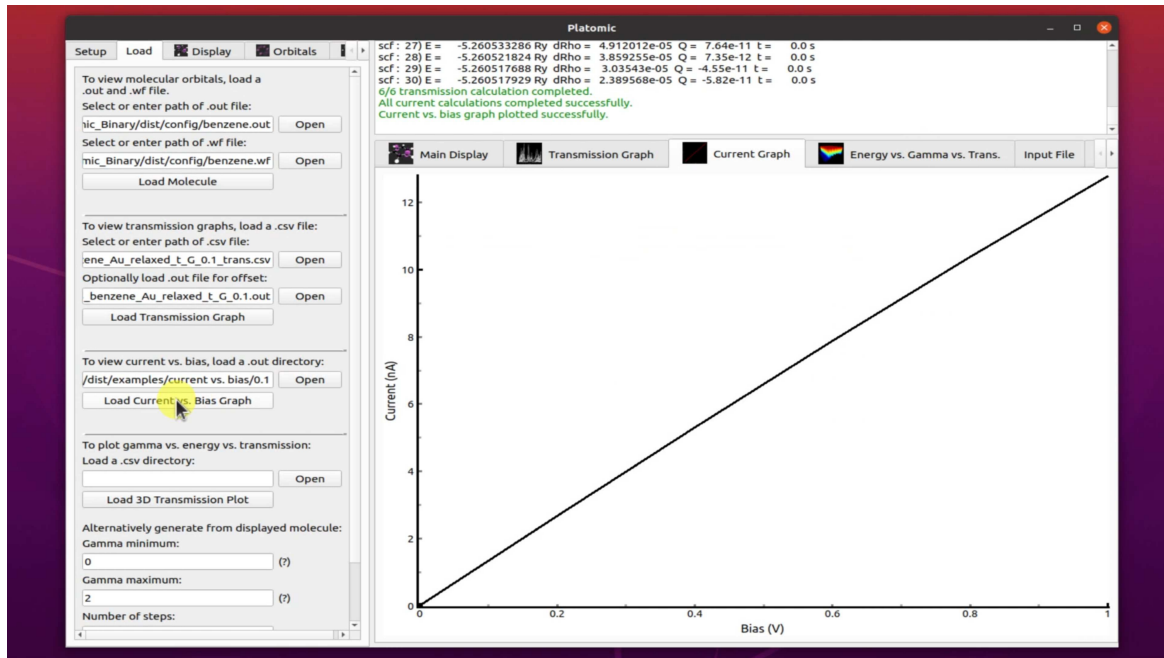
**Figure 2.29:** The result of the current calculation is shown in the console log.



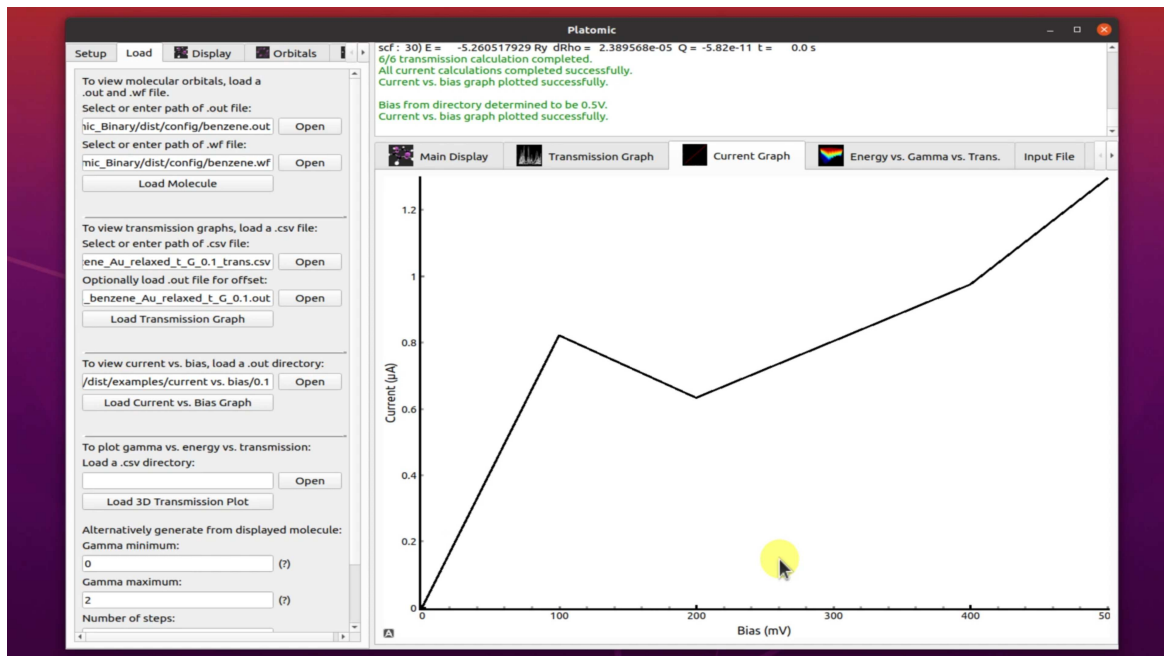
**Figure 2.30:** In addition, you can plot a current vs. bias graph by clicking the corresponding button. The reference potential adjusts the number of electrons in the system. Max bias refers to the maximum value for the graph (it starts from zero and goes to max bias). Steps refer to the number of calculations to make between zero and the max bias, i.e. the number of data points desired for the current vs. bias graph.



**Figure 2.31:** Like the 2D Transmission graph feature in Section 2.2, the same view and export options are available.



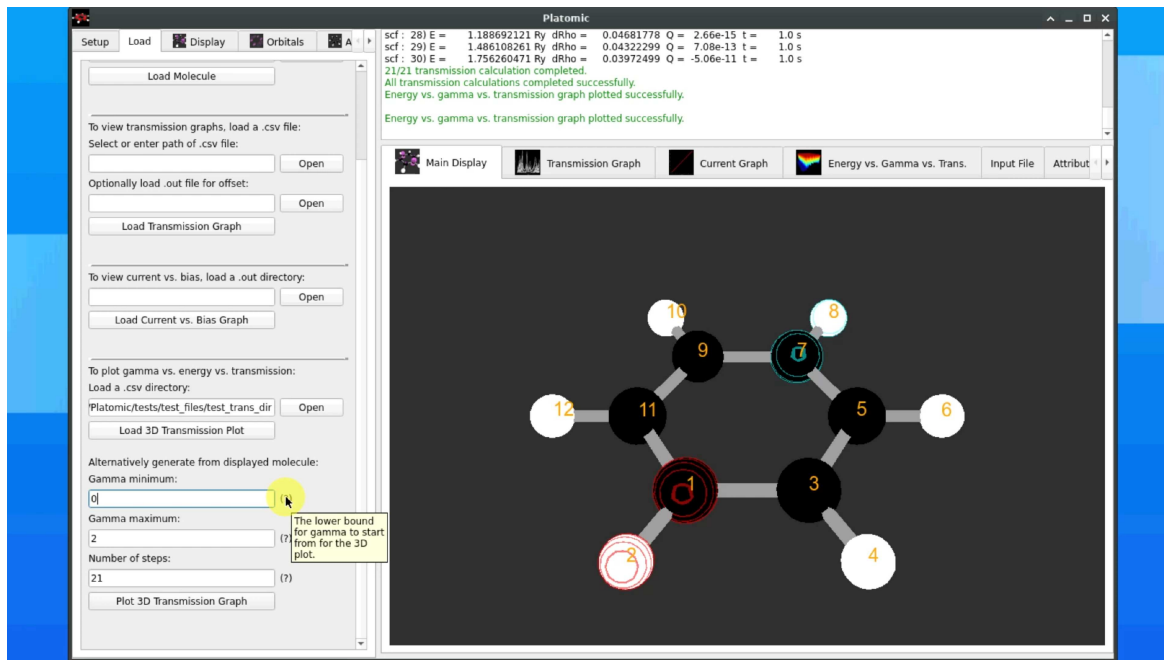
**Figure 2.32:** A current vs. bias graph can be loaded into Platonic by opening a directory of pre-computed Plato .out files.



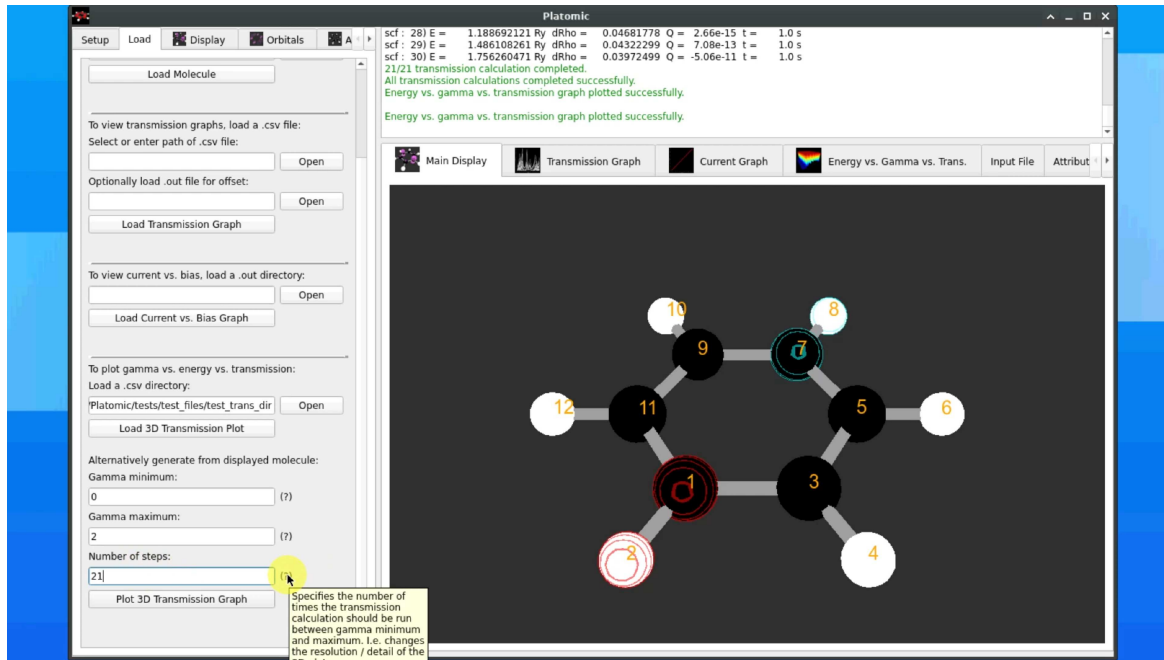
**Figure 2.33:** An example of a current vs. bias graph loaded from a directory of pre-computed .out files.

## 2.4 3D Transmission plots (energy vs. gamma vs transmission graphs)

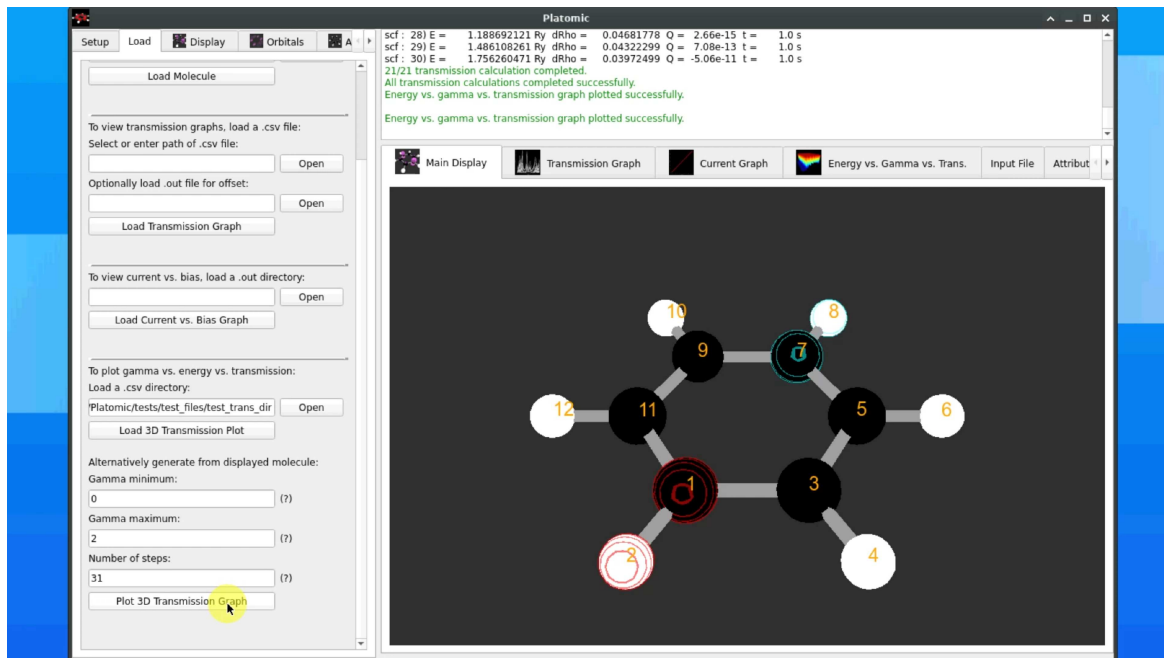
In addition to the walk-through written below, you may refer to the video tutorial linked here: [Section 2.4 - 3D Transmission Plots](#).



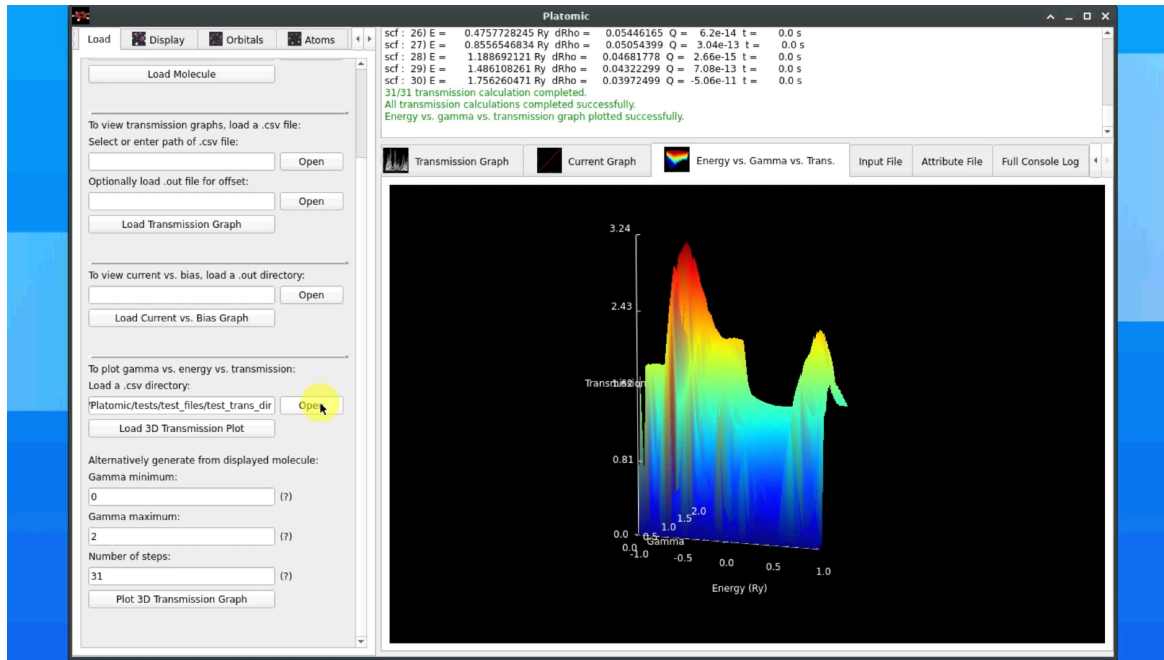
**Figure 2.34:** 3D Transmission plots, which are essentially the 2D Transmission graph with an extra dimension, shows the effect of gamma on the simulation results. Gamma as explained previously in Section 2.2 controls how strongly coupled probes are to the system. After a molecule is loaded and terminals are selected, you can adjust the minimum and maximum gamma values, which defines the range of the gamma axis. Here, we have a benzene molecule with two terminals, where terminal one consists of atoms with indices 1 and 2 and terminal two consists of atoms with indices 7 and 8.



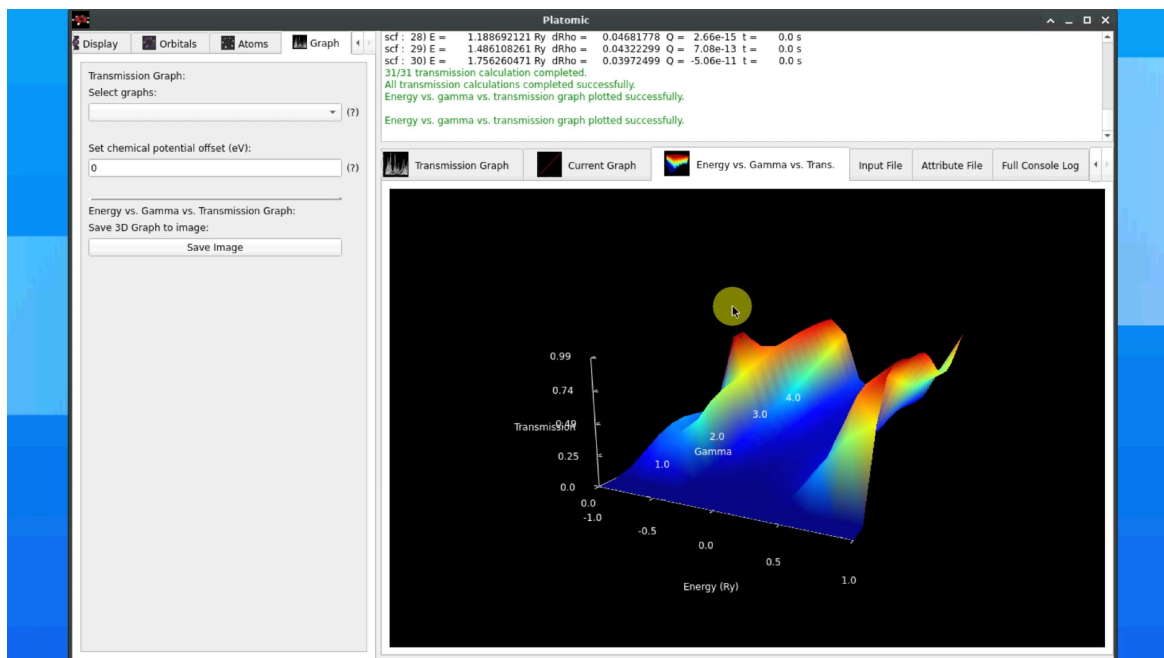
**Figure 2.35:** Next, you can adjust the number of steps, which refers to the number of times a transmission calculation should be carried out from min to max gamma. I.e. changes the resolution of the 3D Transmission plot.



**Figure 2.36:** Then to start the executions and subsequently plot the 3D Transmission graph, press the ‘Plot 3D Transmission Graph’ button.



**Figure 2.37:** The camera view of the 3D Transmission graph can be adjusted like the molecular orbital visualisation feature in Section 2.1. The 3D Transmission graph can also be loaded from a directory of pre-computed CSV files.



**Figure 2.38:** Another example of a 3D Transmission graph for benzene, which was loaded from a directory of pre-computed transmission CSV files.

# Chapter 3

## Developing Platonic

### 3.1 Programs and scripts

#### 3.1.1 atom.py

`_init_`: initialises the Atom object.

`decrement_bond_count`: decrements bondCount member variable when bond is plot.

`get_bondCount`: getter for the remaining number of times bonds can be plot.

`get_bonding`: getter for bonding behaviour as read from attributes.txt.

`get_colour`: getter for atom colour.

`get_eigenenergies`: getter for the eigenenergies of the system.

`get_eigenenergy`: getter for a specific eigenenergy.

`get_eigenvector`: getter for a specific eigenvector.

`get_index`: getter for the atom index.

`get_isSelectedCurrA`: getter for whether the atom is selected for region A.

`get_isSelectedCurrB`: getter for whether the atom is selected for region B.

`get_isSelectedTrans`: getter for whether the atom belongs to a terminal.

`get_maxBonds`: returns maximum number of bonds permitted for this atom as read from attributes.txt.

`get_mi`: returns mesh item object associated with the atom.

`get_quantum_dict`: returns orbital quantum numbers for the atom.

`get_radius`: returns radius for the atom as read from attributes.txt.

`get_skipOrbital`: returns flag to skip orbital plotting in Platonic for optimisation reasons.

`get_symbol`: returns the atomic symbol.

`get_total_orbitals`: returns total number of orbitals in the system.

`get_xyz`: returns coordinates of the atom as a list.

`reset_bonds`: helper function to reset bondCount to maxBonds.

`set_bonding`: setter for bonding behaviour as read from attributes.txt.

`set_colour`: setter for atom colour.

`set_eigenenergies`: setter for the eigenenergies of the system.

**set\_eigenvector:** setter for the 2D array of eigenvectors.  
**set\_isSelectedCurrA:** sets atom to be part of region A.  
**set\_isSelectedCurrB:** sets atom to be part of region B.  
**set\_isSelectedTrans:** sets atom to a specific terminal.  
**set\_maxBonds:** sets the maximum number of bonds the atom can form as read from attributes.txt.  
**set\_mi:** sets mesh item object to Atom instance.  
**set\_quantum\_dict:** sets orbital quantum dictionary to Atom instance.  
**set\_radius:** sets radius to Atom instance as read from attributes.txt.  
**set\_skipOrbital:** sets skip orbital plotting flag for Atom instance as read from attributes.txt.  
**set\_total\_orbitals:** sets the total number of orbitals in the system.

### 3.1.2 glaxis.py

#### Label class

**\_\_init\_\_:** inherits from gl.GLGraphicsItem.GLGraphicsItem parent class.  
**setGLViewWidget:** sets Label instance to a GLViewWidget.  
**paint:** renders Label onto a GLViewWidget.

#### GLAxis class

**\_\_init\_\_:** inherits from GLAxisItem parent class.  
**add\_labels:** adds Labels to GLAxis mesh item.  
**add\_tick\_values:** adds tick values to GLAxis mesh item.  
**paint:** renders GLAxis axes and tick marks.

### 3.1.3 glview.py

#### GLView class

**\_\_init\_\_:** inherits from GLViewWidget parent class.  
**mousePressEvent:** custom mouse click interaction for terminals and regions.  
**itemsAt:** returns list of mesh items in a region.  
**paintGL:** used to render coordinate based labels.  
**readQImage:** captures widget view as an image.

### 3.1.4 glview3D.py

#### GLView3D class

`_init_`: inherits from GLViewWidget parent class.

`readQImage`: captures widget view as an image.

### 3.1.5 input.py

`correct_quantum`: converts Plato magnetic quantum numbers which are whole numbers back to the traditional integer representation. E.g. 0, 1, 2 to -1, 0, 1.

`create_all_atoms`: creates the Atom array for the molecule.

`create_orbital_dict`: creates a dictionary with orbital and quantum number information for each element in the molecule.

`curr_plato_input`: generates Plato current input file.

`eig_arr_from_wf`: returns 2D array with all eigenvectors split by energy.

`find_current_in_file`: parses output file for current.

`get_last_two_chars`: returns last two characters in file which matches a string.

`get_line_number`: returns line number in file which matches a string.

`get_lines_between`: returns lines between start and end strings with options.

`input_file_setup`: generates molecular orbital input file.

`isfloat`: boolean checks if a string is a float.

`isnatnumber`: boolean checks if a string is a natural number.

`isposfloat`: boolean checks if a string is a positive float.

`lines_that_contain`: returns the line from file which matches a string.

`lines_that_start_with`: returns the line from file if the line's start matches a string.

`process_current_out`: processes directory of current output files into arrays.

`process_energy_gamma_trans_csv`: processes directory of .csv files into arrays.

`return_occupied_keys`: returns the number of selected terminals.

`return_occupied_keys_list`: returns the terminals selected.

`set_attr_from_file`: sets attributes from attributes.txt to Atom array.

`set_eig_to_atoms`: sets 2D array of eigenvalues, orbital count and quantum dictionary to Atom array.

`total_orbitals`: returns the total number of orbitals in the system.

`trans_plato_input`: generates Plato transmission input file.

`transmission_headers`: parses transmission .csv file for column headers and maps headers to Platonic selections.

`xyz_to_plato_input`: generates Plato molecular orbital input file.

### 3.1.6 main.py

`_init_`: inherits from QtWidgets.QMainWindow class.

`draw`: updates main display by plotting atoms, bonds and orbitals.

`execute`: interacts with Plato back-end.

**onBiasLineEditChanged:** checks bias line edit for invalid inputs for current vs. bias graphs.

**onCurrExecuteLoadedButtonClicked:** loads current vs. bias graph from pre-computed directory of .out files.

**onCurrentSelectionA:** called when the user right clicks on an atom, writes to console log user selection and updates member variable for region A.

**onCurrentSelectionB:** called when the user middle clicks on an atom, writes to console log user selection and updates member variable for region B.

**onExcessLineEditChanged:** checks excess electrons line edit for invalid inputs.

**onExecute3DGraphButtonClicked:** executes Plato back-end and plots energy vs. gamma vs. transmission 3D graph.

**onExecuteButtonClicked:** executes Plato back-end and plots the molecule with molecular orbitals.

**onExecuteCurrButtonClicked:** executes Plato back-end and prints current to console log.

**onExecuteCurrGraphButtonClicked:** executes Plato back-end and graphs current vs. bias graph.

**onExecuteLoadedButtonClicked:** loads molecule and molecular orbitals from pre-computed .out and .wf files.

**onGammaEndLineEditChanged:** checks gamma end line edit for invalid inputs.

**onGammaExecuteLoadedButtonClicked:** loads pre-computed directory of .csv files for energy vs. gamma vs. transmission 3D graph.

**onGammaLineEditChanged:** checks gamma line edit for invalid inputs.

**onGammaLineEditChanged2:** checks gamma line edit for invalid inputs.

**onGammaStartLineEditChanged:** checks gamma start line edit for invalid inputs.

**onGammaStepsLineEditChanged:** checks gamma steps line edit for invalid inputs.

**onGenerateCurrInputFileButtonClicked:** generates current input file.

**onGenerateInputFileButtonClicked:** generates molecular orbital input file.

**onGenerateTransInputFileButtonClicked:** generates transmission input file.

**onOpenCsvFileButtonClicked:** opens QFileDialog for pre-computed transmission .csv files.

**onOpenDirButtonClicked:** opens QFileDialog for a pre-computed directory of .out files (for current vs. bias graphs).

**onOpenDirGammaButtonClicked:** opens QFileDialog for a pre-computed directory of .csv files for energy vs. gamma vs. transmission graphs.

**onOpenFileButtonClicked:** opens QFileDialog for a .xyz file.

**onOpenOutFileButtonClicked:** opens QFileDialog for a .out file.

**onOpenWfFileButtonClicked:** opens QFileDialog for a .wf file.

**onReferenceLineEditChanged:** checks reference potential line edit for invalid inputs.

**onResetViewButtonClicked:** resets view of the main display.

**onSave3DImageButtonClicked:** saves energy vs. gamma vs. transmission plot as a PNG.

**onSaveAttributeFileButtonClicked:** saves changes to edits made to the attribute file.

**onSaveImageButtonClicked:** saves main display view as PNG.

**onSaveInputFileButtonClicked:** saves changes to edits made to the input file.

**onStepsLineEditChanged:** checks steps (current vs. bias graph) line edit for invalid inputs.

**onSwitchToAttrFileDialogClicked:** switches central window tab to attribute file tab.

**onSwitchToInputFileDialogClicked:** switches central window tab to input file tab.

**onToggleAtomsButtonClicked:** toggles atoms on / off in the main display.

**onTransExecuteButtonClicked:** executes Plato back-end and plots 2D transmission graph.

**onTransExecuteLoadedButtonClicked:** loads 2D transmission graph from pre-computed .csv file.

**onTransSelection:** called when the user left clicks on an atom, writes to console log user selection and updates member variable for terminal selection.

**replaceTextEdit:** helper function to update the text edit widget for input files.

**setAtomColSliderLabel:** called when atom column mesh slider is moved.

**setAtomRowSliderLabel:** called when atom row mesh slider is moved.

**setBondColSliderLabel:** called when bond column mesh slider is moved.

**setBondRadiusSliderLabel:** called when bond radius slider is moved.

**setBondRowSliderLabel:** called when bond row mesh slider is moved.

**setBondThresholdSliderLabel:** called when bond threshold slider is moved.

**setBrightnessSliderLabel:** called when main display brightness slider is moved.

**setCheckBoxIndex:** called when index coordinate label checkbox is changed.

**setCheckBoxPosition:** called when position coordinate label checkbox is changed.

**setCheckBoxRadius:** called when radius coordinate label checkbox is changed.

**setCheckBoxSymbol:** called when symbol coordinate label checkbox is changed.

**setColourASliderLabel:** called when orbital alpha colour slider is moved.

**setColourBSliderLabel:** called when orbital blue colour slider is moved.

**setColourComboBox:** called when colour for coordinate labels is changed.

**setColourGSliderLabel:** called when orbital green colour slider is moved.

**setColourRSliderLabel:** called when orbital red colour slider is moved.

**setFontComboBox:** called when font for coordinate labels is changed.

**setGraphComboBox:** called when graph header for 2D transmission graphs is changed, so the transmission plot widget is updated.

**setHorizontalSliderLabel:** called when the molecular orbital slider is moved.

**setOffsetComboBox:** called when the offset direction for coordinate labels is changed.

**setOrbColSliderLabel:** called when the orbital column mesh slider is moved.

**setOrbRowSliderLabel:** called when the orbital row mesh slider is moved.

**setPhiSliderLabel:** called when the phi rotation slider is moved.

**setScalerSliderLabel:** called when the orbital scaler slider is moved.

**setSizeComboBox:** called when the font size for coordinate labels is changed.

**setTerminalComboBox:** called when the terminal index is changed.

**setThetaSliderLabel:** called when the theta rotation slider is moved.

**updateAtomColSliderLabel:** called when atom column mesh slider is changed.  
**updateAtomRowSliderLabel:** called when atom row mesh slider is changed.  
**updateBondColSliderLabel:** called when bond column mesh slider is changed.  
**updateBondRadiusSliderLabel:** called when bond radius slider is changed.  
**updateBondRowSliderLabel:** called when bond row mesh slider is changed.  
**updateBondThresholdSliderLabel:** called when bond threshold slider is changed.  
**updateBrightnessSliderLabel:** called when main display brightness slider is changed.  
**updateColourASliderLabel:** called when orbital alpha colour slider is changed.  
**updateColourBSliderLabel:** called when orbital blue colour slider is changed.  
**updateColourGSliderLabel:** called when orbital green colour slider is changed.  
**updateColourRSliderLabel:** called when orbital red colour slider is changed.  
**updateHorizontalSliderLabel:** called when the molecular orbital slider is changed.  
**updateOrbColSliderLabel:** called when the orbital column mesh slider is changed.  
**updateOrbRowSliderLabel:** called when the orbital row mesh slider is changed.  
**updatePhiSliderLabel:** called when the phi rotation slider is changed.  
**updateScalerSliderLabel:** called when the orbital scaler slider is changed.  
**updateThetaSliderLabel:** called when the theta rotation slider is changed.  
**writeErrorToLogs:** helper function to write errors to both console logs.  
**writeToLogs:** helper function to write messages to both console logs.

### 3.1.7 orbital.py

**advanced\_orbital:** generates s, p, d, f orbital mesh data.

### 3.1.8 plot.py

**colours:** returns either the colour as a string or RGBA tuple.  
**current\_graph:** plots the current vs. bias graph on a plot widget.  
**draw\_advOrbFaces:** draws s, p, d, f orbitals with face meshes.  
**draw\_advOrbHorz:** draws s, p, d, f vertex orbitals horizontally.  
**draw\_advOrbVert:** draws s, p, d, f vertex orbitals vertically.  
**draw\_advOrbWf:** draws s, p, d, f orbitals as a wireframe.  
**draw\_atoms:** draws atoms using passed in Atom array.  
**draw\_bonds:** draws bonds using passed in Atom array.  
**draw\_selection:** draws terminal selection highlights.  
**draw\_sphOrbFaces:** draws spherical orbitals with face meshes.  
**draw\_sphOrbWf:** draws spherical orbitals as a wireframe.  
**energy\_gamma\_trans\_graph:** plots the energy vs. gamma vs. transmission graph.  
**transmission\_graph:** plots the 2D transmission graph.

## 3.2 Directory map

### 3.2.1 Platonic Binary

```

Platonic
└── build/ (stores metadata, is useful for debugging)
└── dist/
    ├── LICENSE
    ├── benzene.xyz (example .xyz file)
    ├── benzene_Au_relaxed.out (example Plato .out file)
    ├── benzene_Au_relaxed.wf (example Plato .wf file)
    ├── benzene_Au_relaxed.xyz (example .xyz file)
    ├── main (main executable)
    ├── setup (simple bash file to setup Platonic)
    └── config/
        ├── attributes.txt (configure atom and bond plotting attributes)
        ├── benzene.out (example Plato .out file)
        ├── benzene.wf (example Plato .wf file)
        ├── default.in (template for molecular orbital input file gen)
        ├── default_curr.in (template for current input file gen)
        ├── default_trans.in (template for transmission input file gen)
        ├── icon.png
        ├── icon1.png
        ├── icon2.png
        ├── icon3.png
        ├── icon4.png
        ├── icon5.png
        ├──mainwindow5.ui (PyQt5 UI file for Platonic)
        └── platonic.png (Platonic application icon)
    └── doc/
        └── User_Guide.pdf
    └── Plato/ (Plato repository or binary)

```

### 3.2.2 Platonic

```

Platonic
├── LICENSE
├── README.md
├── atom.py (Atom class)
├── glaxis.py (custom GLAxis class for 3D graphs)
├── gview.py (custom GLWidget class for main display)
├── gview3D.py (custom GLWidget class for 3D graphs)
├── input.py (processing input and output files)
├── main.py (main program)
├── orbital.py (mesh data generation for s, p, d, f orbitals)
├── plot.py (plotting of atoms, bonds, orbitals and graphs)
└── benzene.xyz

```

```
benzene_Au_relaxed.out
benzene_Au_relaxed.wf
benzene_Au_relaxed.xyz
config/
    attributes.txt
    benzene.out
    benzene.wf
    default.in
    default_curr.in
    default_trans.in
    icon.png
    icon1.png
    icon2.png
    icon3.png
    icon4.png
    icon5.png
   mainwindow5.ui
    platonic.png
doc/
    User_Guide.pdf
Plato/
tests/ (directory used to store unit tests)
    config
    test_files
    coverage.py (used to generate html coverage reports)
    glaxis_test.py
    glview3D_test.py
    glview_test.py
    input_test.py
    main_test.py
    orbital_test.py
    plot_test.py
```