# MASARYK UNIVERSITY FACULTY OF INFORMATICS



# Maya2CellVIEW: 3D Package Integrated Tool for Creating Large and Complex Molecular Scenes

Master's Thesis

David Kouřil

Brno, Fall 2016

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## **Declaration**

Hereby I declare that this paper is my original authorial work, which I have worked out on my own. All sources, references, and literature used or excerpted during elaboration of this work are properly cited and listed in complete reference to the due source.

David Kouřil

Advisor: Ivan Viola

# Acknowledgement

This is the acknowledgement for my thesis, which can span multiple paragraphs.

## **Abstract**

Scientific illustrators communicate the cutting edge of research through their illustrations. There are numerous software tools that assist them with this job. The aim of this thesis is to push abilities of illustrators working on a large scale molecular scenes. This is done by connecting two software packages - Maya and cellVIEW - combining the rendering possibilities of cellVIEW and modeling tools of Maya which results in more effective and efficient workflow.

# Keywords

keyword1, keyword2, ...

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### 1 Introduction

In this age, scientists come to new findings everyday. Unfortunately, not all of these are ever shown to the general public. It could be because of several reasons. One of them might be that the new facts are not easily visualizeable. This can also mean that the things that they found out are a piece of a bigger picture. This is the job of scientific illustrator. These artists are usually experts in their own scientific domain and what they do, is they try to visualize the science in their domain with easily understandable images, animation and other media. There are many tools for this job. Few years back illustrations have been done by hand, drawings and paintings. Then came the age of computer graphics. As graphics tools have become more and more common and sophisticated in a field of computer generated movies (reference to Pixar?). Tools have become more accessible, easier to use and so scientific illustrators adapted and tried to use these tools for their benefit. Tools like Maya, Cinema4D, or XXX are nowadays the industry standards when it comes to authoring 3D content. Inside these programs there usually exist a software called renderer. With this two parts modeler and renderer - the user is able to create images or animations (movies) from the 3d data that he creates in the software. Even with just this, the modern computer graphics brings us the possibility to create amazing imagery and enables illustrators to show ordinary people fenomena from all kinds of science disciplines. But this workflow isn't very effective. One of the obstacles in the work of illustrators is that the software they are using is made for somebody else. Games and movies industry are the leading industries that push computer graphics tool creators forwards and provides most of the revenue for them. This means that these tools, no matter how much they try to be versatile, are being skewed towards the use cases of movies and games industries. That means that people that want to create scientific content might struggle to use the tools sometimes. For this, there are other tools that help them to make the pipeline more effective. This doesn't mean that these tools completely replace Maya or Cinema4D. The domain specific tools are more commonly implemented in a form of a plugin that is developed and loaded into the concrete software packages. In the end - the features that 3D authoring package provides

for modeling 3D scenes are hugely beneficial to the creators. Another things is - people that used these software are used to that and switching from one to another is not a matter of a short time. That's one of the things that we need to take into account. The pipeline of each artist might be a little different and rather than suggesting a solution that completely replaces their pipeline, it's better to make a tools that doesn't break their workflow but rather the artist is able to incorporate the tool, benefit from it but doesn't have to change the way they work a lot. There also exist programs that are completely separated from any of the mentioned authoring software. There is an active research in the domain of visualization, with many conferences every year and thousands (?) of research papers in this domain. Usually, as a biproduct, these paper generate software that showcases the proposed visualization technique or pipeline. Some of these developed into full featured visualization packages and thus provide a way how to illustrate something in a new way. While these programs might hugely benefit scientific illustrators, it's not always the case that they get used. This might be because of several reasons one of which is that the illustrators are simply used to a certain pipeline and incorporating a new software into this pipeline doesn't seems very beneficial to them. Another problem is that because these programs are developed for a certain use cases (showcasing the point of the paper) they might not be easily appliable to more than one purpose.

In this thesis, we made an effort to solve these problems. Our domain of choice is molecular visualization. In this domain we try to visualize living (?) organisms on the smallest possible scale. With this approach, the computer memory and performance requirements are very challenging even for todays available hardware. Still, people have been able to visualize for example model of HIV in blood plasma in it's full detail down do individual atoms of each protein. This however, is achieved by using a very customized rendering method and a custom data format. In 3D graphics, data are usually represented as meshes consisting of triangles (which in turn are made of vertices). If we wanted to represent each atom as a sphere, we would need at least <number> of vertices for one atom and that's only for the roughest level-of-detail. With such representation, a one whole protein could use up to X bytes of memory which already takes up most of the video cards' memory. Thus this representation is not suitable for this task.

Instead, various other techniques have been developed for rendering on molecular data. That being said, these techniques are not usually taken into account when 3D authoring software is developed. As was mentioned, the primary users of these software are video games and movies industry, where mesh representation is the one used. That implies that people that use these programs don't have access to the cutting edge technology of rendering, visualization and illustration of molecular data and there is room for us to improve this situation.

In the next chapter, we will describe the state of the art tools that are used for molecular visualization today and we'll mainly focus on showing the gaps in intercompability of the available programs. Then, in chapter 3, we will propose a method of how the problem could be solved.

### 2 State of the art

Popsat: Molecular Flipbook, Molecular workbench, molecular maya, mcell, cellblender, pymol, VMD, ePMV(scripps) In this chapter we will introduce the software that is available for scientific illustrators to use now. Illustrators have gone a long way from doing these works by hand and nowadays there exist a variety of programs that help them do their job communicating the scientific findings. First and foremost, the pipeline of almost all of these illustrators is strongly based on some 3D modeling software. The biggest players right now are Autodesk Maya, Cinema 4D, Autodesk 3Ds Max Design, Blender or even up and comming MODO(https://www.thefoundry.co.uk/case-studies/viscira/), first two named being the dominating choices between illustrators. These programs are complex and aim to provide tool for all kinds of users. Their main functionality revolves around two activities - creation of the 3D scene which is most commonly a mesh models and rendering of this scene. These two stages will be referred to as modeling and rendering. From this brief description we can already see one problem with using a software like this for creating illustrations of molecular models - they make assumptions about the data. While most of the people using this software would actually be using meshes to represent their 3D scenes, in case of molecular models this is not the case. TODO: say something about "scene graph" which is how scene is implemented usually inside these programs. Luckily, it has become a convention that all of these programs provide some kind of scripting interface. This enables its users to extend the program with plugins that use the scripting interface. This way the user can implement functionality that he or she is missing from the basic program. Thanks to that these 3D authoring programs are able to be adapted to more use cases. Some of these plugins have actually already been implemented to help artists in molecular illustration. The most prominent is Molecular Maya which, as the name suggests, extends Maya with the ability load and manipulate models of macromolecules from Protein Database. Molecular Maya plugins gives its users the ability to load macromolecules based on its PDB ID or locally from pdb file. After the model load, user is able to select the display representation (between TODO: vyjmenovat) and also select with how much

detail the model should be rendered. Similar plugin exists for Blender modeling program as well. It's name is cellblender This could be a nice reference: https://pdb101.rcsb.org/learn/resource/molecular-animation-q-and-a-interview

Barely any of these tools is perfect. Some of the tools are very closed and tied to it's creator. The ideal case would be if any new user could approach any of these tools and with a little training be able to use it. This is not the case, unfortunately. From the ones that are fairly simple from the user experience stand point is Molecular Flipbook[ref]. It provides user the ability to create simple molecular models and animate them with a keyframe-like animation workflow.

An important part of any scientific illustration are the data the one is trying to illustrate. In the case of molecular models, there is cellPACK which helps us with this. cellPACK is a software that assembles large molecular scenes from a description (a recipe) of how this scene should look, what it should contain and in what quantity.

If we only focused on the problem that the iteration loop is too long because of the rendering times, we would be able to find solution in "interactive renderers" - something like OTOY's Octane or whatever is in MODO. These are path tracers that show the intermediate result which enables the artists to see the progress of the renderer rendering the scene. The artist can then see how his actions are reflected in the final render. This solution however would not solve everything for us if we consider that we are rendering molecular scenes. These scene are usually very big and we still need to use different representation other then meshes to represent our scene and use this representation to effectively render the scene. The contribution of this thesis is not only the fact that we shorten the iteration loop time of the artist but also to sketch an idea of how interoperability of molecular programs can be done.

## 2.1 Maya

Autodesk Maya is not a tool for molecular visualization per-se.

#### 2.2 cellVIEW

# 3 OLD (integrate into STAR) - Overview of the two programs that I'm going to be bridging -

Here I will talk about in which context are these two 'systems' used and what is each for. The point of this project was to make a bridge between two programs. In this chapter we will look at both of them, see what they bring to the table when used separately, what they lack and how can be benefit from connecting them in this way.

#### 3.1 CellVIEW

Briefly describe what CellVIEW is about. Give links to the papers that are describing CellVIEW and those that are using it.

### 3.1.1 Unity

I don't know if this is needed. I could just go little bit into detail of why is Unity good (which is funny now that we are probably going away from it)

## 3.2 Maya

Point to examples of animations that are produced with Maya. Maya is a 3D content authoring software package developed by Autodesk. It has been introduced in 1998 and since than it has grown into defacto industry standard in many fields that use computer graphics (most notably movies and computer games industry). Because of it's versatility, Maya has also been used for creating scientific illustrations. Although Maya is a very robust piece of software that gives it's users ability to create various types of 3D scenes, it isn't always the best tool for the job. There exist other software packages that might be more specialized and better suited for a task at hand.

3. OLD (integrate into STAR) - Overview of the two programs that  $I^\prime m$  going to be bridging -

### 3.2.1 Extensibility

Fortunately, Maya comes with an enormously useful ability to be extended with so called plugins. Autodesk provides an API that programmers can use to extend Maya's functionality. Several plugins have been developed specifically for the field of molecular visualization and illustration. As an example, we can name Molecular Maya which brings users the ability to import molecular data from online Protein Data Bank. The plugin then allows the user to choose how this data should be represented in a 3D scene and Maya then takes care of rendering such structure.

### 4 Method

I have introduced the field. I have sketched the problem that we are trying to solve (improving the workflow of animators/illustrators). Now it's time to describe the suggested solution. We talked about improving the workflow of scientific illustrator that uses Maya as an authoring software. The way that we are going to accomplish this is by taking advantage of extensibility of both Maya and Unity. There are basically two stages of authoring a scientific visualization. These are modeling stage and rendering stage. TODO: image of the pipeline with time estimates of how long each stage takes, compared to our new approach What we basically did is use the fact that modern GPUs are able to render what we want in a decent visual quality and that they can do it really fast (real time). This way we can shorten the time it takes to the illustrator to create his artwork. From Maya we aimed to use it's various modeling tools - tools that enable translating, rotating, scaling of objects in scene. With that we wanted to use particle system to generate randomly molecular scene (not scientifically correct but close enough for illustration purposes). The vision was that there would be cellVIEW which renders the scene on one side and Maya on the other. In Maya the scene would be appropriately simplified. At the beginning we had an idea that there would just be cubes with maybe textures that would give the user at least some visual information about the protein type. We ended up using very rough meshes that represent the proteins in the scene in Maya. As this project wasn't meant to be software intended for certain user group, we work with very little input requirements and so the design and user experience side of things have been explored over the course of development. We however had initial input from one of the most respected domain experts in scientific illustrator - Drew Berry. From this persona came the initial idea to use Maya as the modeling package as that's the software that Drew Berry uses. He expressed his interest in rendering style that cellVIEW provides and suggested that if we were to make cellVIEW usable in his workflow with Maya, it would greatly improve his productivity and shorten the production of his movies. His input sparked our interest in making cellVIEW usable as a part of a pipeline rather than a standalone tool. This then lead to research of which

resulted in the following proposed method. This general method is presented as an idea of an approach that tools like cellVIEW could take in making themselves usable in art authoring pipeline. In this chapter we present this idea. Further in implementation chapter we focus on our use case - using Maya and cellVIEW but we believe that this method should be appliable on any two modeling and rendering tools (with possibility to be used in other applications as well). In computer science there are not a lot of ways to translate information from one program to another. The majority of this communication is done via files that are exported on one side and imported on another. This lead to a need for standardized format in which this data are formatted so that both of these programs understand each other. This simple approach is fine for most of the application where it's used. In our case however this isn't sufficient. There is one big problem. If user needs to perform the actual export and import step, this greatly reduces his productivity. Another problem that is found in our domain has already been mentioned - rendering times. Our method tries to solve these two problems. The problem of import/export is solved by using shared memory instead of files managed by the user. The rendering times problem is solved by using modern rendering techniques which enable us to render what we want in an interactive framerate. The reason why it's possible to take this approach of using shared memory comes from the character of data that we work with. Molecular scenes consist of molecules. Molecules are made of atoms which can be, and usually are, represented as spheres. Atoms of different elements are visualized by having different radii and colors. So for every molecule we need to keep track of what atoms it consists of, positions of those atoms and type of each of those atoms. This is even more leveraged by Protein Data Bank, which stores data about various proteins that have been found through out the years. For us, this means that we only need to keep track of the type of molecule and by fetching data from Protein Data Bank we get information about the protein cataloged under the protein identifier. Our scenes are (simplified) made of macromolecules. That means that in the end for rendering the scene we need a list of molecules, where for each molecule instance we need to save it's position, rotation and type. That's all the info that we need to be able to render our large molecular scenes. Even though this is still a lot of data, we are slowly getting to being able to render this on commonly available hardware. Because of the fact that this data size is "manageable" by modern computers we are able to store the data in shared memory. This method is mostly valuable as an example of how interconnection of software can be done and what improvements to the users this can bring. There is some prerequisities - both programs need to be extensible to the extend of the programmers must be able to develop extension that are able to read from and write to the shared memory using operating system api calls.

## 5 Implementation

The key element of the implementation has been how to use all the different parts of the ecosystem, all the extensibility possibilities, and connect them together in a way that would show our method. The architecture of the system basically consists of three parts - a plugin for Maya, plugin for Unity and a Unity script.

#### 5.1 WinAPI

### 5.2 Maya side

Topics - Architecture of the plugin, what parts/classes of API are used, memory layout, what I tried and didn't work.

### 5.3 Unity side

Topics - Architecture of the plugin, very generally about plugins (it's just a basic C++ dll plugin), interface between C++ and C#

## 6 Demonstration

In this chapter we present two use cases of how the user can approach illustration with this new proposed system.

## 7 Discussion, future work

Limitations (of the tool) - what was done just for the use case and should be worked on for final production. The project has been presented to Drew Berry (I think). Also, it was mention in a talk in Utah, presentation by Peter Mindek.

# A An appendix

Here you can insert the appendices of your thesis.