A Novel Computer Vision Methodology for Intelligent Molecular Modeling and Simulation

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Abstract: Molecular modeling and simulation tools are used to study the structure of the molecules for the purpose of understanding and creating a new generation of technology that works on the nano-scale. The current techniques mainly focus on visualizing the molecule’s structure using many illustrative methods, while they leave the knowledge extraction load on the user that should be aware of many complex sciences. Developing a new innovative method in this perspective becomes crucial to support such fast development in such vital field of sciences. This paper represents a novel computer vision method for molecular modeling and simulation that gives the computer the ability to see and understand the structure of molecules just like the human eyes, and also the ability to analyze its structure without human intervention. The proposed approach is based on using the computer’s memory as a digital representation of the real 3D-physical scaled model of the molecule, and hence accommodates machine learning techniques for an automated analysis job. Moreover, a parallel processing approach has been adopted to speed up the whole process. The realistic case study of a glucose molecule reports the outstanding performance of the proposed approach to model and analyze its structure without human intervention. The proposed methodology makes the developing of an automated molecular expert system a one step away.

1 INTRODUCTION

Today the world turns its eyes on the technologies and phenomena that happens at the Nanoscale, where scientists are studying the cell and the cellular structures such as proteins, their structures, and their functions (Friedrichs et al., 2009; Durrant and McCammon, 2011; Soni et al., 2014). Scientists are learning lessons from nature. They are looking forward to building molecular machines and robots using modified proteins and other nano1-materials to do things that were impossible in the past. Molecular modeling and simulation tools can help scientists in studying and modifying the structure of a molecule by doing the following: visualizing the molecule using computer graphics, simulating the motion of the molecule under different forces and conditions (Dawson et al., 2016; Khatib et al., 2011; Durn-Riveroll et al., 2016; Jallu et al., 2012), simulating the interaction between the molecule and other molecules (Lindert et al., 2013; Friedrichs et al., 2009), analyzing the arrangement and geometrical shapes of the atoms inside the molecule to find the critical points at which the molecule’s structure will change.

The current molecular modeling and simulation tools like Avogadro (Hanwell et al., 2012), VMD (Humphrey et al., 1996), YASARA (Krieger and Vriend, 2014), and RasMol tools (Potterton et al., 2002) pay much attention on rendering the molecule structure and leave the user to study the molecule by himself. They left all the analysis, and knowledge extraction effort to be done by the user who must be an expert in molecular sciences, and theories to take over these tasks. Frequently, the user is even obligated to write a computer program to customize these tools in order to do very simple jobs. This user-dependent approach makes the current methods and tools away of getting the full benefits from the computer sciences’ methods and techniques like machine learning, computer vision, and artificial intelligence. Enhancing the current molecular modeling and simulation tools to overcome their critical limitations be-

11 Nano-meter = 1x10^{-9} meter
come crucial to support the fast development in such vital field of sciences. This will help the scientists to design new molecules and nano-materials that can be used in many applications.

This paper represents a novel computer vision methodology for molecular modeling and simulation which mimics the human eye’s vision and gives the computer the ability to see and understand the molecular structures. It will enable the user to see inner parts of the molecule that may be hidden using the current techniques. It also has the ability to analyze the molecular structures and extract rich knowledge from it without human intervention. Hence, developing molecular expert systems, chemical and physical knowledge bases will become a step away.

The proposed methodology’s approach is based on using the computer’s memory (RAM) as a 3D-digital representation media to model the physical molecular structure (i.e., atoms and bonds) using digits 0 and 1. Each bit represents a cube of 1 picometer$^3$ in the spatial space of the molecule. A parallel processing approach has been adopted to efficiently speed up the process of the whole method. This paper reviews most of the current molecular software tools like RasMol, PyMOL, VMD, Avogadro, GROMACS, and Jmol to discuss all their pros and cons. The extensive simulation studies conducted on a glucose molecule report the outstanding capability of the proposed method to extract knowledge from the molecular structure and analyze it without human intervention, contrarily to current human-dependent approaches.

The rest of the paper is organized as follows: Section 2 provides a scientific background in molecular modeling and simulation. It also reviews related work. Section 3 provides a detailed explanation of the proposed methodology and the suggested parallel architecture. Section 4 presents the conducted real molecular case study and compare between the proposed methodology and the suggested parallel architecture. The paper is then concluded in Section 5.

2 BACKGROUND

This section explains how the molecular structure is discovered using X-ray. It then reviews the related work in molecular modeling and simulation.

2.1 Scientific Background

As illustrated in figure 1, there is a cycle of steps to discover the structure of any molecule. The first step in Figure 1: The cycle of discovering any molecules structure to extract the target molecule from living organisms. Step two is to get a sufficient amount of such molecule and convert it to a crystal form. Step three is to expose such crystal into an x-ray crystallography device. This device shoots extensive x-ray beams from different angles through the crystal and collects the diffraction of such rays on a light-sensitive sheet. The collected light intensities are then analyzed by a computer program to reveal the position of each atom in the molecule and its chemical type. Here, it is worth to mention that there are other techniques that do the same job like NMR, Mass Spectrometry, and 3D Electron Microscopy, where NMR, for example, use the magnetic field instead of the light diffraction. Nevertheless, they all produce a file called MOL as an abbreviation of the word molecule that reveals the position of each atom in the molecule and its chemical type. Again, there are other types of files different than the MOL like SDF, XYZ, and PDB chemical files. The SDF and XYZ files reveal the same information but in a different file structure, while PDB files are used to describe proteins. Finally, once the MOL file is created, any molecular viewer software takes place to render the molecule is 3Dimensions on any computer’s screen.

2.2 Related Work

In the early 90s, a real improvement in x-ray crystallography and molecular imaging techniques has appeared, since then researchers and scientists have tried to build software tools to study different molecules’ structures. This software can be divided into two categories. The first category includes molecular viewers, molecular editors and molecular designers (Hanwell et al., 2012; Humphrey et al., 1996; Sayle and Milner-White, 1995; Potterton et al., 2002) all of these software tools can visualize molecules. the second category includes molecular dynamics simulation tools that visualize the chemical reactions of the molecules (Humphrey et al., 1996; Emsley and Debreczeni, 2012; Dreher et al., 2013; Phillips et al., 2005). Now, let’s review a group of molecular software with their pros and cons.

The Avogadro tool (Hanwell et al., 2012) visualizes the molecules to the user on a computer’s screen.
using 3D computer graphics. It also enables the user
to choose an atom as the origin and rotate around it using
the mouse buttons and the keyboard buttons. Nevertheless, the user should always memorize the place
and the colors of the atom as well as its chemical types
in order to easily navigate through the molecule without
losing focus.

The VMD tool (Humphrey et al., 1996) visualizes the molecules to the user with different types of
graphical representations. It can do a lot of energy
calculations. It works through a special scripting lan-
guage, that should be used to initiate any complex job.
Learning a new programming language to customize
the molecular graphics software to execute complex
and even simple commands require a lot of time and
effort.

The YASARA tool (Krieger and Vriend, 2014)
renders molecules using 3D graphics with less number
of polygons and in less time, so even smartphones
can render large molecules faster with no hanging or
lagging. The YASARA tool has the advantage of working on molecules anywhere and on any type of
computers from workstations to smartphones. how-
ever, this tool pays much attention to rendering and
rotating the molecules within a tedious workspace
without enough attention to real analysis and knowl-
edge extraction.

(Emsley and Debreczeni, 2012) designs drugs
using molecular graphics tool that can render the
molecules with different complex presentations.
Again, it is the user job to understand these presen-
tations that may take a long time based on his knowl-
edge.

3 THE PROPOSED COMPUTER VISION METHODOLOGY

3.1 The Main Idea

This section provides a detailed explanation of how
the 3D-physical model of the molecule is represented
in the computers memory, and how the knowledge is
extracted. It also explains the adoption of the paral-
lell architecture to speed up the process of the whole
methodology. The main idea of the proposed methodology is to build a digital model of 0 and 1 into the
computers memory in the form of a 3D-array of bits (i.e., 0 and 1) that
identically simulate the real physical structure of the molecule. First, the proposed methodology reads the input file that describes the structure of the molecule in terms of its atoms that are inter-connected with spe-
cific chemical bonds. Note that the input file de-
cribes any given molecule by listing its atoms and
their spatial distribution in a 3D-dimension. The
current molecular modeling and simulation software tools render this input file using OpenGL library and
use any traditional graphical tools to visualize its
structure. In our approach, the same process is done
but on a 3D-array of bits. As illustrated in figure
2, each atom in this molecule is created in the form
of a sphere of bits of 1, where each bit represent
1 Pico – meter, which is the basic measuring unit
used to represent an atom. The rest of the bits in the
3D-array will be of value 0 to represent the empty
space between the atoms. Note that even the size of
each atom based on its chemical type is identically
reflected inside the new representation.

The result of this process will produce an identi-
cal digital model of the original molecule in the com-
puter’s memory (RAM) as illustrated in Figure 3. The
size of this array can vary depending on the original size of the molecule in Pico-meter unit.

A library has been developed to hold the unique features of all the organic atoms, which help the
computer to recognize each atom later using the pro-
posed computer vision algorithm. To sum up, we can
now claim that this 3D-presentation reflects all major information about the molecule’s structure such as its atoms spatial distribution, size, and the internal distance between atoms.

3.2 The Construction of the 3D Molecular Representation

Here, we explain how we build the 3D-digital model in form of a 3D-array of bits (i.e., 0 and 1) that
identically simulate the real physical structure of the molecule. First, the proposed methodology reads the
input file that describes the structure of the molecule in terms of its atoms that are inter-connected with spec-
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2The creation of this file was explained in section 2, and
it is the same file used as input for any of the existing tools
31 Pico-meter = 1x10^-12 meter
4The organic atoms are: Oxygen, Nitrogen, Hydro-
gen, Sulphur, Phosphorus, Calcium, Magnesium, Potas-
sium, Chlorine, Sodium.
3.3 The Proposed 3D Computer Vision Algorithm

We should remember that the generated 3D-array in the previous step is a collection of zeros and ones that are still in need for interpretation to extract valuable information that can be later transformed to knowledge using machine learning techniques. The proposed computer vision algorithm explained in this subsection will take over this task. Here, the computer vision methodology is composed of two phases, the first phase is scanning the 3D-array, and the second phase is extracting the knowledge from the scanned data.

This algorithm takes either the whole molecule’s 3D-array as input or even a smaller part of it (i.e., a sub 3D-array of bits) to understand and reveal all necessary information about the examined area. It starts from a specific atom in the molecule or even around it within a given space. As illustrated in figure 4, the algorithm starts by reading the given 3D-array of bits by examining the 2D-array in XY plane and then move to the next 2D-array in the Z-direction. It keeps recording the countered spheres in a list that we call the “Hit-List” and recognize each atom through its diameter \( d \). Note that the chemical bonds which exist between atoms are represented in a separate data structure which links each chemical bond with its endpoints atoms that are participating in the bond. This process can be done starting from a specific origin in a certain direction inside a volume with a given depth. The scan works as a transformation and projection from 3D to a 2D-presentation which produce a stream of 2D-images of atoms inside the scanned volume space. The geometric arrangement of the atoms inside the resultant 2D-images such as angles between bonds’ axes and distances between atoms are stored in the Hit-List. The scanning process reveals also the geometric shapes formed by the atoms and the bond’s axes. Note that the produced Hit-List will be the basic input for the next knowledge extraction phase. The steps of the scanning algorithm are illustrated in Figure 5: Algorithm 1: Scanning the 3D-array.

Once the 3D-array is scanned, the computer will be able to study the relationship between the atoms using molecular geometry functions and extract the following knowledge:

- The distances between the atoms.
- The bonds that exist between the atoms.
- The arrangements of the atoms.
- The electrical charges of each atom.
- The volume of the molecule.
- The volume of empty space inside the molecule.
- The distribution of the atoms’ density.
- The distribution of atoms’ weights.
- The geometric shapes that are formed by the atoms and the bond between them.
- Empty volumes of space inside the molecule and between its atoms.
- Recognition of the molecule’s surfaces and the atoms that compose its surfaces in all directions.
- The dimension of the cuboid that encloses the molecule.

As a matter of fact, the first three extracted points can be mathematically computed from the MOL file directly, however, they are still easily calculated from the new presentation and should be reported as part of the extracted knowledge. Nevertheless, the remaining
Input: Arr3D which is a 3D array of bits with number of 2D matrices M
Data: Circ2D which is a 2D array of bits, DIM which is an integer array of six cells, ChemType which is a char array of two cells
Result: Hitlist which is a list of atoms that lie inside the scanned volume

for $i \leftarrow 0$ to $M$ do
    Read2DMatrix(Arr3D[$i$]);
    if Find(2D circle of bits with value of 1) == true then
        WriteBits(2D circle of bits with value of 1, Circ2D);
        if FindInHitList(Circ2D) == true then
            continue
        else
            DIM = DetermineDimensionsOfAtom(Circ2D);
            ChemType = DetermineChemicalTypeOfAtom(Circ2D);
            AddAtomToHitList(DIM, ChemType, Hitlist);
        end
    end
end

Figure 5: Algorithm 1: Scanning the 3D-array.

3.4 Parallel Processing

The 3D-array can be divided into sub 3D-arrays without losing the spatial arrangements of atoms in the molecule, and hence a strong potential for applying the single instruction multiple data paradigm (SIMD) on these sub 3D-arrays exists in order to accelerate the scanning of the molecule’s representation by harnessing the underlying multi-core hardware. Note that the time consumed by the CPU to build the 3D-digital representation represents only 1% out of the total runtime, so the CPU can build and swap between the cubes very fast.

The proposed algorithm uses the OpenMP C/C++ library for parallel processing of the molecule’s sub 3D-arrays. The proposed parallel implementation divides the molecule’s cube into sub 3D-arrays and assigns each of them to one processing core or thread. The previously mentioned scanning part is then implemented on all the sub 3D-arrays simultaneously, as illustrated in figure 6. After a careful study of both scanning and knowledge extraction phases, we found that a parallel processing architecture can be applied in the scanning phase only while it is not suitable yet for the knowledge extraction one due to the interdependency between sub 3D-arrays at their boundaries. Therefore, all the information that has been collected from the parallel scan are then collected centrally in the Hit-List for the next knowledge extraction phase.

4 EXPERIMENTAL CASE STUDY AND COMPARISON

This section discusses the results of the extensive simulation studies conducted on a realistic glucose molecule using the proposed methodology.

4.1 Simulation Setting

The proposed methodology has been implemented on a Linux operating system using C programming.

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Figure 6: Distributing the sub cubes among the cores of a CPU.
language. OpenGL graphics library for checking the results by rendering the molecules using 3D graphics, and OpenMP for multi-core programming. The studies were implemented on Intel Core i7-3612QM with 6 mega cache CPU and 8 threads each thread works at processing frequency range from 2.10GHz to 3.10GHz and Intel Core i7-4790 with 8 mega cache CPU and 8 threads each thread works at processing frequency range from 3.60 GHz to 4.00 GHz.

The source code is divided into following four main modules:

- The first module is responsible for reading the chemical files.
- The second module is responsible for constructing the 3D-array of bits in the memory and building the molecule’s atoms inside it.
- The third module is responsible for executing the parallel scan of the molecule’s sub 3D-arrays.
- The fourth module is responsible for extracting the knowledge and visualizing the results to the user on the screen using 3D graphics.

4.2 Case Study

Simulation studies have been conducted on different 3D-arrays sizes distributed among a different number of cores to assess the parallelism effect of the proposed methodology and to measure the gain of speeding up the scanning phase. As illustrated in figure 7, in the case of using 1 thread (serial program) the run-time took 32 seconds and the memory consumed is 0.2 Gigabytes. In case of 4 threads (Parallel program) the run-time decreases by 72% and the memory consumption remain the same, after increasing the number of threads to 8 the run-time decreases by 18% again and the memory consumed increased by 50%. Finally, in case of 16 threads the run-time increases up by 2% and the memory consumption increased by 33.3%. It is worth to note that the more threads we apply on the CPU, the more the run-time and memory consumed increase because allocating memory for threads, creating and destroying threads costs overhead processing run-time and memory. Based on the previous run-time degradation, we preferred to implement the following case study using 8 threads in order to get the best performance with the least run-time.

Once the 3D-array is scanned and the Hit-list is created, the proposed methodology displays the glucose molecule in 3D with six carbon atoms colored in dark grey, six oxygen atoms colored in red, and twelve hydrogen atoms colored in light grey, as illustrated in figure 8. The figure displays only 3 shots for the molecule from the XY, XZ, and YZ planes for simplicity as a printed version. The proposed methodology also displays a comprehensive report that reveals the knowledge extracted from the glucose molecule As illustrated in figure 9. The report covers the extracted information previously mentioned before in section 3.3.

To show the capability of the proposed methodology, we conducted a more sophisticated experiment by choosing a specific atom inside the glucose molecule and study only the area behind it for the purpose of going into a deeper level of understanding of each atom in the molecule. The proposed methodology also displays a comprehensive report that reveals the knowledge extracted as illustrated in figure 10. The report covers the following additional information inside the scanned space:
Table 1: Comparison between the available tools and the proposed methodology.

<table>
<thead>
<tr>
<th>Comparison criteria</th>
<th>Available tools</th>
<th>Proposed methodology</th>
</tr>
</thead>
<tbody>
<tr>
<td>User dependent knowledge extraction</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>3D Rendering</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Navigate inside the 3D model</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>View geometric patterns inside the molecule</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Recognition of the molecule surface</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>The volume of empty space inside the molecule</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>The distribution of atoms density of the molecule</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>The distribution of atoms weight of the molecule</td>
<td>No</td>
<td>Yes</td>
</tr>
</tbody>
</table>

- The distances between the chosen atom and the atoms behind it.
- The chemical bonds between the chosen atom and the atoms behind it.
- 3D-images of the atoms spatial distribution.
- 3D-geometric shapes of the spaces between atoms.

Next, the glucose molecule is examined using the current tools like Avogadro and PyMol. As illustrated in figures 11 and 12, the Avogadro and PyMol tools render the glucose efficiently on the screen and wait for the user to choose one atom as the center of rotation. The user will rotate the glucose molecule using the mouse and try to recognize the chemical types of the atoms around the origin atom through their colors using his eyes. The user will try to study the relationships between the origin atom and atoms around it using his knowledge in molecular sciences. The user may have to spend some time in writing a small script in a special programming language or clicking on menus and buttons in order to customize the molecular viewer.

To sum up, we can easily note the outstanding extracted knowledge using the proposed methodology in comparison with the currently available tools that mainly depend on the user, as summarized in Table 1. The proposed solution promises to open a new area of molecular sciences and will significantly enhance the development in this crucial field.

5 CONCLUSION

This paper represents a novel computer vision methodology for molecular modeling and simulation that gives the computer the ability to see, understand, and analyze the molecular structures by itself without human intervention. Its main idea was based on using the computer’s memory (RAM) as a 3D-digital representation of the molecule’s structure. A new algorithm was developed to help the computer to see the new representation, and extract the knowledge about the vital aspects inside the molecule using a parallel architecture to speed up the data processing. This paper reviews most of the current molecular software...
tools like RasMol, PyMOL, VMD, Avogadro, GROMACS, and Jmol to discuss all their pros and cons. The extracted knowledge reports the outstanding capabilities of the proposed methodology in comparison with the current tools.

REFERENCES


