



User's Manual

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Introduction

This document is a work in progress.

What is Tornado?

SimTK Tornado is a dynamic visualization tool for coarse grain (lumped) representations of

RNA and/or protein structure. By shifting the focus from individual atoms to higher order

structures, such as double helical duplexes, the scientist's attention is naturally focuses toward a

larger scale. Biologists will be able to use their intuition to interactively refold RNA structures

and produce morphs from one structure to another.

Purpose

Provides an easy to use application for animating and visualizing RNA and other

macromolecular structures.

Audience

Biomedical and computation researchers interested in RNA, molecular motions, and folding

pathways.

Installing and Running

System Requirements

Platforms: Windows, Mac, and Linux

Java 1.5 or higher required (http://java.sun.com/)

Java Web Start

Tornado uses Java Web Start to load and run directly from the download page at SimTK.org.

The downloaded files are stored in a special area on your computer called the Java Web Start

cache, so Tornado should start more quickly on the second and subsequent runs.

Download Application

- 1. Find the tornado project page at SimTK.org
- 2. Click on Downloads on the left side menu.
- 3. Click on Tornado jnlp file on the right side of the screen under Download Links.
- 4. ToRNAdo will begin to download automatically.
- 5. If a window saying **Warning Security** appears, click the **Run** button.
- 6. Wait a few seconds and the **SimTK ToRNAdo** application will appear.



Figure 1 – Find the Tornado project at SimTK.org



Figure 3 - Select the latest Tornado JNLP file at the right of the Tornado download page



Figure 2 - Find the Tornado download link at the left of the Tornado project page



Figure 4 - Open the JNLP file with Java Web Start. If this does not happen automatically, you may need to install a recent release of Java on your computer.



Warning - Security

The application's digital signature has been verified. Do you want to run the application?

Name: JOGL

Publisher: sun microsystems, inc

From: http://download.java.net

Always trust content from this publisher.

The digital signature has been validated by a trusted source.

More Information...

Figure 5 - Acknowledge the trusted digital signature on the Tornado application

Figure 6 - Acknowledge digital signatures on third party components



Figure 7 - If there is something wrong with a digital signature, please file a problem report at SimTK.org (described in the later section "Submitting Problem Reports and Enhancement Requests"). In this example, the third party component uses a valid certificate that has expired, and thus may pose a minor security risk.



 $\label{eq:Figure 8} \textbf{Figure 8} \text{ - The tornado application is now running on your computer.}$

Troubleshooting Java Web Start

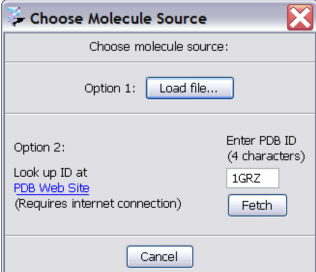
If downloading stalls, try the following:

- 1. Click "Cancel", and reopen the jnlp file.
- 2. If it still fails, clear the Java Web Start cache, and try opening the jnlp file again.

Tutorial

After downloading and running Tornado as described in the previous section of this document, you should see the Tornado application window, as shown below:





Loading Molecules

- Click on Choose Molecule button to choose a molecule structure.
- Option 1: click on the **Load file** button to choose a molecule saved locally.
- Option 2: enter the molecule ID (PDB ID) and click the **Fetch** button to load molecule from the PDB Web Site (requires internet connection).
- Molecule will appear on the screen with default view **Residue Balls**.
- To load a new molecule, go to the **File** menu and click **Load PDB Molecule** and choose option 1 or 2.

Change Molecule Views

- Go to the **View** menu, click on **Molecule Style**.
- A menu of various view styles will come up.
- Choose desired style.

Save Current Molecule View as an Image

- Go to the File menu, click on Save PNG Image.
- Under File Name, choose a desired name followed by .png
- Save the file in the desired location.

Rotating, Zoom In/Out Molecules

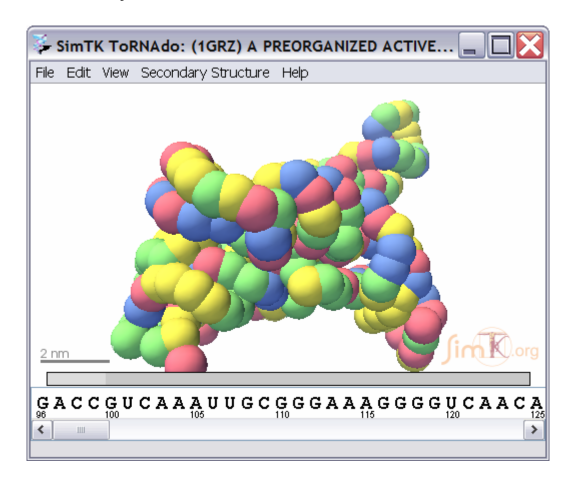
- Go to the **View** menu, click on **Rotation**.
- A menu of various rotation styles will come up.
- Choose desired rotation.

Manual Rotation

- Left click and hold the mouse on current screen.
- Move up, down, left, right to rotate as desired.

Zoom In/Out

- Right click and hold mouse on current screen.
- Move up or down to zoom in and out.



Molecular Visualization

Selecting residues

Select one or more residues to highlight those residues in the display.

The following mouse gestures apply to all molecule displays in Tornado, including the linear sequence display, 3D structure display, and secondary structure display.

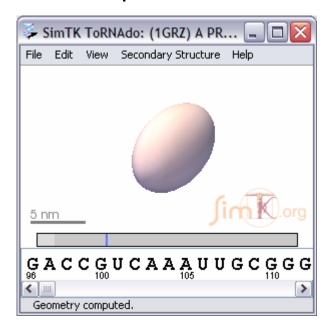
- Click on a residue to select that residue
- Ctrl-click on a residue to add that residue to the list of selected residues.
- Double-click on a residue to center the display on that residue.
- Shift-click to add all residues between the most recently selected residue and the current residue to the list of selected residues.

Representations

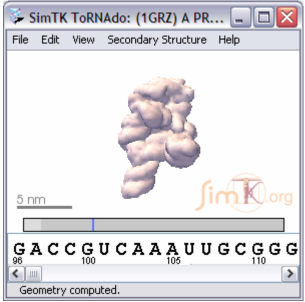
	RNA/DNA	Protein	Ligands	Water
Molecule	ellipsoid	ellipsoid	ellipsoid	not shown
Ellipsoids				
Molecule Blobs	blob	blob	blob	not shown
Secondary	rope and	backbone	space filling	not shown
Structure	cylinder			
Residue Spheres	residue spheres	residue spheres	ball and stick	not shown
Ribbons	ribbon	ribbon	ball and stick	not shown
Bond Lines	lines	lines	lines	not shown
Atom Balls	atom balls	atom balls	atom balls	not shown
Ball and Stick	ball and stick	ball and stick	ball and stick	ball and stick

Coarseness (Fineness) of representation

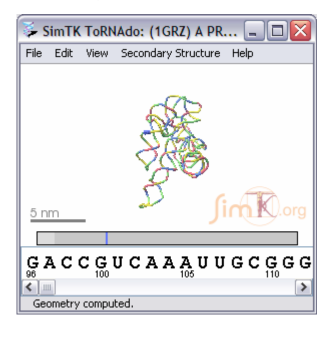
Molecule Ellipsoids



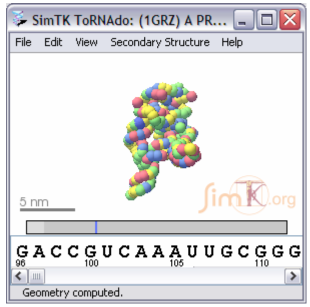
Molecule Blobs



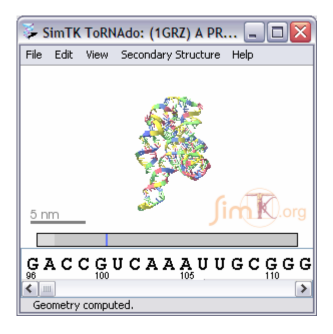
Secondary Structure



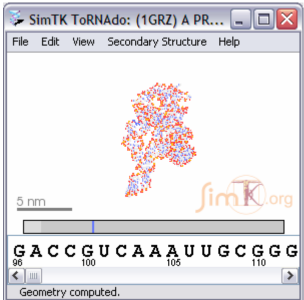
Residue Balls



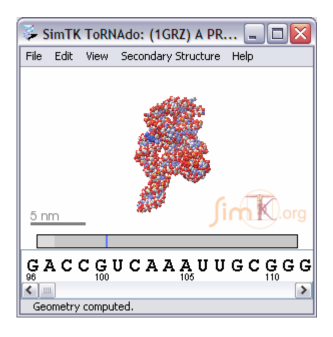
Ribbons



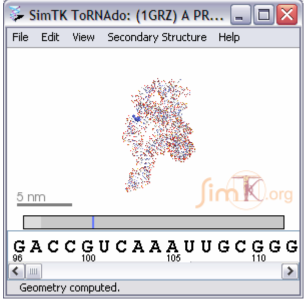
Bond Lines



Atom Balls



Ball and Stick



Nucleic Acid Secondary Structure – RNAML files

RNAML is an xml file format for descriptions of RNA structure. Examples of other applications that use RNAML format are RNAVIEW and MFOLD.

Saving An Image File of the 3D structure

From the menu bar, select "File"->"Save PNG Image".

At the right side of the file save dialog, you can adjust the image magnification, which permits you to save images at a higher resolution than you monitor can display.

Submitting Problem Reports and Enhancement Requests

If you observe a problem with the Tornado program, or if you would like to request a new behavior in future versions of Tornado, please file a bug or feature report at SimTK.org. From the tornado project page, select Advanced->"Features & Bugs"->"Bugs" or "Features"->"Submit New", then fill in the report and click "Submit". Thank you for helping us to improve Tornado.

References

[RNAVIEW] Yang, H., Jossinet, F., Leontis, N., Chen, L., Westbrook, J., Berman, H.M., Westhof, E. (2003). Tools for the automatic identification and classification of RNA base pairs. *Nucleic Acids Research* **31** (13): 3450-3460.

[MFOLD] M. Zuker, D.H. Mathews and D.H. Turner Algorithms and Thermodynamics for RNA Secondary Structure Prediction: A Practical Guide in RNA Biochemistry and Biotechnology, J. Barciszewski and B.F.C. Clark, eds., NATO ASI Series, Kluwer Academic Publishers, (1999)