

# Using 3DNA: A Short Introduction

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In the following guide we'll show a step by step introduction to 3DNA from installation of the software to practical examples.

## 1 Installation

To install 3DNA you will first have to download the software from:

`http://adna.rutgers.edu`

The webpage will ask you to register before allowing you to download the program.

Once registered you'll be able to download the file **Linux\_X3DNA\_v2.0.tar.gz** if you selected the Linux version.

To uncompress issue the following command in a terminal

```
tar -xvzf Linux_X3DNA_v2.0.tar.gz
```

This will create a new folder named **X3DNA** which you will have to include in your environment variables by changing your shell environment configuration file named ".bashrc" in bash, and ".chsrc" in csh and tcsh.

If you are using .bashrc the following lines will need to be added:

```
export X3DNA=/home/yourusername/X3DNA
export PATH=$PATH:$X3DNA/bin
```

A detailed explanation of how to configure X3DNA can be found in the pdf file **x3dna\_v1.5.pdf** found in the X3DNA/doc folder. If you're stuck in the installation process feel free to drop me at [mauricio.esguerra@gmail.com](mailto:mauricio.esguerra@gmail.com)

## 2 fiber

The **fiber** program included in the 3DNA package allows you to easily generate protein data bank (pdb) files of fiber models of common nucleic acid conformations, for example A-DNA, B-DNA, and Z-DNA. To get a list of all possible fiber models which you can reconstruct using **fiber**, type:

```
fiber -l
```

If you type **fiber** in the terminal you will get a usage message.

Stop and read the usage message carefully.

Now, you know that the crystal structures of A-RNA and B-DNA must have a total of 11 and 10 residues per turn respectively, so, go ahead and use **fiber** to make an A-RNA with 12 residues and a B-DNA with 11 residues. Using a molecular visualization program such as pymol, vmd, or chimera check the generated structure to corroborate that you have the correct number of residues per turn.

## 3 find\_pair

The **find\_pair** program of 3DNA allows you to find pairs in a given pdb file. To read the usage message open a terminal and type:

```
find_pair
```

Now go to the protein data bank website at <http://www.pdb.org> and download the structure with PDB\_ID: 1ehz. This is the structure for yeast phenylalanine tRNA. Run it through **find\_pair**:

```
find_pair 1ehz.pdb 1ehz.inp
```

You will get a file named 1ehz.inp. Go ahead and open the resulting file in a text editor and you will see that the `find_pair` program has found the canonical Watson-Crick base-pairs G·C and A·U, as well as non-canonical base-pairs such as G·U and A·G.

## 4 analyze

The `analyze` program of the 3DNA software package allows you to compute the base-pair and base-pair step parameters associated with a given nucleic acid structure, among other analysis features, such as computing the overlap areas between stacked nucleic acid bases, sugar conformations, and sugar-phosphate backbone torsion angles.

A requirement of the `analyze` program is that you have to run the `find_pair` program beforehand. Go ahead and read the usage message of the `analyze` program by typing:

```
analyze --help
```

in your terminal.

Now run the `analyze` program on the file previously produced using the `find_pair` program:

```
analyze 1ehz.inp
```

Notice that you can run the `find_pair` and `analyze` program in one line by issuing the command:

```
find_pair 1ehz.pdb stdout | analyze
```

This command will produce a wealth of files with structural information, but the main one will just have the name of the input file followed by the `.out` extension. That is, you should now have a 1ehz.out file in your folder. Go ahead and take a look at the 1ehz.out file in your favorite text editor.

## 5 rebuild

The very useful 3DNA program called `rebuild` can do just that, rebuild nucleic acid structures. It uses as input either a set of base step, or base-pair step parameters, or a set of helical parameters to create a `pdb` file. To test the `rebuild` program first run `find_pair` and `analyze` on the A-RNA structure you previously created:

```
find_pair A-RNA.pdb stdout | analyze
```

Issuing the previous command must have created a file called *bp\_step.par*. Open this file in a text editor and change the Twist values from 31.5 degrees to a smaller value, say 25.0 degrees, and save the new file with the name *undertwisted.par*.

Rebuild the undertwisted RNA by typing the following command in your good-ole terminal:

```
rebuild -atomic undertwisted.par undertwisted.pdb
```

This will generate a `pdb` structure with no sugar-phosphate backbone. To add a sugar-phosphate backbone you will need to issue the `cp_std` command before you perform the `rebuild` command, in the following way:

```
cp_std RNA
```

```
rebuild -atomic undertwisted.par undertwisted.pdb
```

Now rebuild a `pdb` file using the unmodified base-pair step parameters, that is, using the original *bp\_step.par* file. Open both structures in a molecular visualization program and confirm that one helix is undertwisted with respect to the original.

## 6 Exercises

1. Run `find_pair` and `analyze` on your A-RNA structure and modify the resulting `bp_step.par` file so that the sequence is AAAGGGUUUCCC instead of AAAAAAAAAAAAAA. Use `rebuild` to generate a `pdb` file with the AAAGGGUUUCCC sequence.
2. Download the file with PDB\_ID: 1aoi, that is, chromatin's nucleosome core particle, from the protein data bank website. Run it through `find_pair` and `analyze`.
3. Produce a “publication quality” plot of Slide vs. base-pair step number for the chromatin parameters you've computed in the previous exercise. Ask around (e.g. drop me a line) for which solutions are good for such a task.  
I recommend `xmgrace`, the `matplotlib` library of `python`, and `igor-pro`.
4. Repeat the recipe for Twist vs. base-pair step number (`bpsn`), and Roll vs. `bpsn`.
5. Compare your graphs to Figure 2 of the journal article “DNA Sequence-Directed Organization of Chromatin: Structure-Based Computational Analysis of Nucleosome-Binding Sequences” *Biophysical Journal*, **2009**, 96, 2245-2260.  
What regularities do you see?  
How would you describe the DNA wrapping around the nucleosome in terms of Sliding, Twisting, and Rolling?