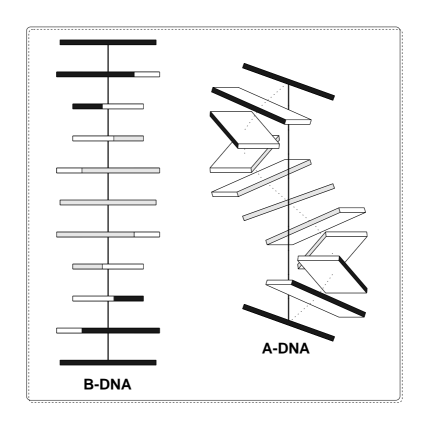
3DNA (v1.5) — A Software Package for the Analysis, Rebuilding and Visualization of 3-<u>D</u>imensional <u>N</u>ucleic <u>A</u>cid Structures



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1 Introduction

Our recent comparison studies (Lu & Olson, 1999; Lu *et al.*, 1999) on seven popular analyzing programs for nucleic acid structures demonstrated that the choice of reference frame rather the mathematical formulation has the greater effect on the calculated parameters. Broadly speaking, there are two classes of reference frames. One is defined by the purine (R) C8 atom and pyrimidine (Y) C6 atom, and the mean base-pair plane as in FreeHelix/NewHelix (Dickerson, 1998). The other uses the average of the two constituent base reference frames in a base-pair as in Curves (Lavery & Sklenar, 1989).

While each has its own advantages and limitations, the base-centered one was recommended at a workshop held on January 13-14, 1999 at the AIST-NIBHT Structural Biology Center in Tsukuba, Japan (Olson *et al.*, 2001).

3DNA is a comprehensive software package for the analysis and rebuilding of nucleic acid structures based on the new recommended reference frame. Mathematically, it follows the CEHS definition (El Hassan & Calladine, 1995) as implemented in SCHNAaP/SCHNArP (Lu et al. 1997a; Lu et al., 1997b) with the global helical parameters replaced by a new set of *local* ones. This matrix based algorithm is rigorous and reversible in that it allows for exact reconstruction of the structure from a set of parameters which can be deduced from the structure. The parameters such defined also have simple geometrical meaning thus the distance between base-pair centers is $\sqrt{\text{Shift}^2 + \text{Slide}^2 + \text{Rise}^2}$, the bending angle between two base-pairs is $\sqrt{\text{Roll}^2 + \text{Tilt}^2}$, and the angle between base-pair normal and local helical axis is $\sqrt{\text{Inclination}^2 + \text{Tip}^2}$.

3DNA is a completely new written software package conforming strictly to the ANSI/ISO C. It is much more robust and efficient than SCHNAaP/SCHNArP and has new and improved features. Its analysis part can automatically classify a dinucleotide step as A-, B-, or TA-like (Guzikevich-Guerstein & Shakked, 1996) based on the positioning of phosphorus atoms with respect to the middle frame. This can be very useful for pinpointing conformational transitions in ligand-bound DNA, especially in a short fragment which is often neglected by other analyzing programs (Lu et al., 2000). High quality "standardized" base-stacking diagrams can be generated by a utility program following the analysis. Authentic CEHS/SCHNAaP parameters are also available, which can be used for comparisons with the 3DNA recommended ones to see how a difference in reference frames can affect the calculated parameters, especially for Rise in heavily deformed DNA structures. Also note that FreeHelix/NewHelix parameters are very similar to the CEHS/SCHNAaP ones due to their similarity in reference frames. For completeness, local parameters using the seven methods (Lu & Olson, 1999; Lu et al., 1999) based on the new recommended reference frame are also calculated. Obviously, they are all quite similar to 3DNA recommended ones.

The rebuilding part of 3DNA can be used for generating sequence-dependent atomic structures, without or with the sugar-phosphate backbone, suitable as starting point for molecular mechanics calculations and molecular dynamics simulations. It is also very convenient to generate publication quality Calladine-Drew style schematic representations of DNA, either in PS, XFIG¹, or Raster3D (Merritt & Bacon, 1997)² format. With XFIG, the picture can be edited and annotated, combined with others, or exported to image formats such as GIF, JPEG and TIFF. With Raster3D, the Calladine-Drew style block representation of bases can be combined with the backbone and protein in either atomic or schematic presentations.

Fifty five different types of fiber DNA and RNA structures based on the work of Chandrasekaran & Arnott (1989) can also be conveniently generated. The atom naming and ordering conventions of the NDB are strictly followed which make it easy for direct comparison with X-ray crystal structures or theoretical models.

¹http://www.xfi g.org/

²http://www.bmsc.washington.edu/raster3d/raster3d.html

2 Installation

The latest version of 3DNA, in binary form for Linux, SGI and Windows, is available from the following URL: http://rutchem.rutgers.edu/~xiangjun/3DNA/. Download the version that fits your system and then do the following:

- gunzip Linux_X3DNA_v1.5.tar.gz, you get file Linux_X3DNA_v1.5.tar
- tar xvf Linux_X3DNA_v1.5.tar, you get directory X3DNA and four subdirectories underneath: BASEPARS, FIBER, bin, and Examples, the contents of which are detailed below.
- If X3DNA is not directly under your home directory, set environment variable X3DNA as follows:

```
in csh | tcsh: setenv X3DNA Your_Directory_Containing_X3DNA
  e.g., setenv X3DNA /usr1/xiangjun/X3DNA
in bash | sh: export X3DNA=Your_Directory_Containing_X3DNA
  e.g., export X3DNA=/usr1/xiangjun/X3DNA
in Windows (suppose 3DNA is installed under C:\X3DNA):
  set X3DNA=C:\X3DNA
  Or do the followings (globally):
    Start
         Settings
             Control Panel
                 System
                     Advanced
                         Environment Variables
                             at "User variables" section
                                Variable: X3DNA
                                Variable Value: C:\X3DNA
```

Then append to your command search path the 3DNA bin subdirectory:

2.1 BASEPARS

This directory contains standard residue geometry files and other parameters controlling various aspects of 3DNA. They are all in text format, so users can view the structures with rasmol or edit the parameters as they see fit.

• Atomic_?.pdb (? = A, C, G, T or U) are the default standard residue geometries used by 3DNA for analyzing and rebuilding full atomic nucleic acid structures in PDB format.

Under its subdirectory ATOMIC, there are four sets of standard residue geometry files:

ADNA_std?.pdb, BDNA_std?.pdb, NDB96_std?.pdb and RNA_std?.pdb.

The base geometries by Clowney $\it et al.$ (1996) are used (downloaded from the NDB archive: http://ndbserver.rutgers.edu/NDB/archives/index.html). The NDB96 set includes base and C1' atoms, without sugar-phosphate backbone. ADNA set uses C3'-endo sugar-backbone conformation as defined by Leslie $\it et al.$ (1980) fiber studies, but with a $\it \chi$ torsion angle of -157° , average of high resolution single crystal X-ray oligonucleotide structures. RNA set is the same as ADNA except for an additional O2' atom for each residue. BDNA set is similarly defined as ADNA except for a C3'-endo sugar fiber sugar-backbone conformation and the $-108^\circ \it \chi$ torsion angle.

The NDB96 set is the default. To use another data set, simply overwrite the corresponding Atomic_?.pdb in BASEPARS or copy them to your current working directory. A utility program, cp_std (see below), can do this automatically for you. You can also use other residue geometries with the help of the utility program std_base.

Note the standard set contains only the five common residues, A, C, G, T and U. Residue I can be got by deleting the N2 atom from G. Their modified counterparts, +A, +C, +G, +I, +T, +U which are changed to lower case by 3DNA, can be approximated by using their normal forms. For +C, for example, 3DNA requires file Atomic_c.pdb, which can be simply a copy of Atomic_C.pdb.

• Block_BP.alc defines the default base-pair rectangular block. It has a size of 10 Å (long) by 4.5 Å (wide) by 0.5 Å (thick) and is in ALCHEMY format. It is used for drawing the Calladine-Drew style schematic presentation of DNA structures. Block_R.alc is for the purine base (R) $(4.5 \times 4.5 \times 0.5)$. Block_Y.alc is for the pyrimidine base (Y) $(3.0 \times 4.5 \times 0.5)$.

Under its subdirectory BLOCK, there six block geometry files. Block_M.alc has half the size of BLOCK_BP.alc, and can be used if you would like the two blocks consisting a base-pair to be of the same size. Block_Ms.alc is slightly smaller in length than BLOCK_M.alc to avoid possible overlaps in a compressed base-pair (*i.e.* with negative Stretch). Furthermore, the blocks do not necessarily to be rectangular, as shown in Block R nr.alc.

- Pxyz.dat contains the xyz coordinates of phosphorus atoms with regard to the middle dinucleotide reference frame. Four sets were defined, corresponding respectively to average values in high resolution A- and B-DNA crystal structures, their intermediate and TA-DNA. New set can be added following the format. This file is used by rebuild for generating DNA structures with only base and phosphorus atoms. PxyzH.dat is the same as above except the coordinates are given in terms of the middle helical frame.
- fig_image.par contains parameters controlling the style of generated XFIG files, which can be edited by users to suit their liking. Similarly, ps_image.par holds parameters defining the drawing style for postscript images. Finally, raster3d.par and my_header.r3d are for Raster3D input.
- misc_3dna.par contains various parameters mainly for analyze and the utility program find_pair.
- baselist.dat contains a comprehensive list of currently known base residues and their standard counterpart. It makes analysis of unusual DNA and RNA structures straightforward.
- trans_pep.pdb & trans_pep.alc are trans peptide unit used for drawing peptide block in protein structures, in the same way as basepair blocks.

2.2 FIBER

This directory contains repeating unit for each type of the 55 fibers DNA and RNA structures. The original data as provided by Chandrasekaran & Arnott (1989) is given in subdirectory Data. Directories Str01–Str55 store the "clean-up" version of each repeating unit in a format suitable for building the structure with utility program fiber.

2.3 bin

This directory contains executables of the 3DNA package. Most of which are utility programs with some in short Perl script. Detailed usage of each program is described in Section 3.

2.4 Examples

Four subdirectories are included to illustrate the various functionalities of the 3DNA package. You are strongly recommended to study these examples carefully in order to use 3DNA more effectively.

• Analysis_Rebuild contains the analysis/rebuilding results of four structures: adh026 (A-DNA), bd1084 (B-DNA), pde0128 and pd0001 (DNA-protein complexes). The *.pdb data files were downloaded from the NDB. The *.inp files are the corresponding input to the analysis routine (analyze) and *.out are the output containing various structural parameters.

Input file multi_str.inp illustrates how to analyze multiple structure from one input file.

README contains detailed information on how to run the analyze/rebuild programs to generate results in this directory, and the RMS deviations between 3DNA rebuilt structures and the experimental ones. For base atoms, the RMS is virtually zero, and with backbone atoms it less than 0.85Å even for the 146bp nucleosomal DNA.

• Calladine_Drew illustrates how to generate the two sets of DNA schematic pictures made popular by Calladine & Drew (1997) (Figures 1 and 2). The README file provides every detail. Note that the plots in these two figures are on the same scale and in the same orientation.

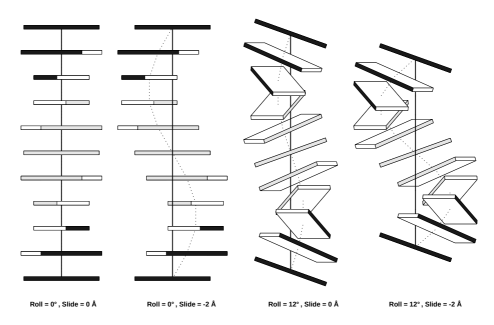


Figure 1: One complete *helical* turn of DNA having twist of 36°, showing the effects of introducing uniform roll and slide at each step (Calladine & Drew, 1997).

- NMR gives an example on how to analyze multiple NMR structures from the PDB.
 Follow the README file there.
- Stacking shows the procedures for generating "standardized" base-stacking diagrams (Figure 3). Check README file there.
- Triplex gives examples on how to analyze triplex and parallel duplex structures. Follow README file there for details.

3 How to run 3DNA

Running 3DNA is simple although it might take a while to use it more effectively. For each of the program, there is a simple on-line help illustrating its usage.

1. Analyzing part

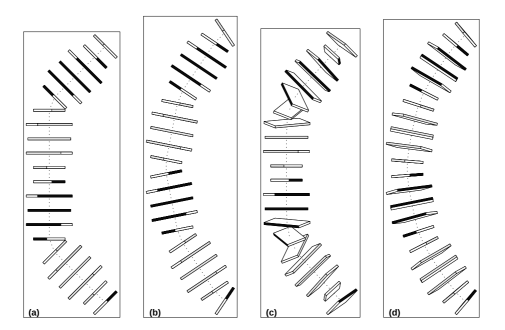


Figure 2: Two complete *helical* turns of DNA, with a curvature of 45° per turn, or 4.5° per step on average. Such tight curvature may be achieved, in principle by any of the distributions of roll angle shown in parts (a) to (d) (Calladine & Drew, 1997).

- analyze [inpfile1 inpfile2 ...]

 Sample input files are given in directory Examples/Analysis.
- cehs [inpfile1 inpfile2 ...]

 cehs gives the original CEHS/SCHNAaP parameters, to which FreeHelix/NewHelix

 ones should be quite similar. cehs is provide for completeness and comparison purpose.
- find_pair [options] pdbfile inpfile find_pair is used to generate an input file for analyze/cehs, starting directly from a PDB file. It can also generate input files for the popular nucleic acid analysis program Curves with option -c.
- manalyze [-cehs] inpfile
 A Perl script for analyzing multiple structures
- nmr_strs [-cehs] inpfile n1 n2
 A Perl script for analyzing multiple NMR structures

2. Rebuilding part

- rebuild [options] [-negx] inpfile outfile
 For rebuilding DNA structures of either atomic model in PDB format or schematic
 representation in ALCHEMY format.
- regular_dna [options] outfile
 A utility program to generate input file for rebuild for the construction of regular DNA structures.
- frame_mol [options] -n1[,n2] reffile inpfile outfile
 A utility program for attaching local helical frames and set the orientation of a

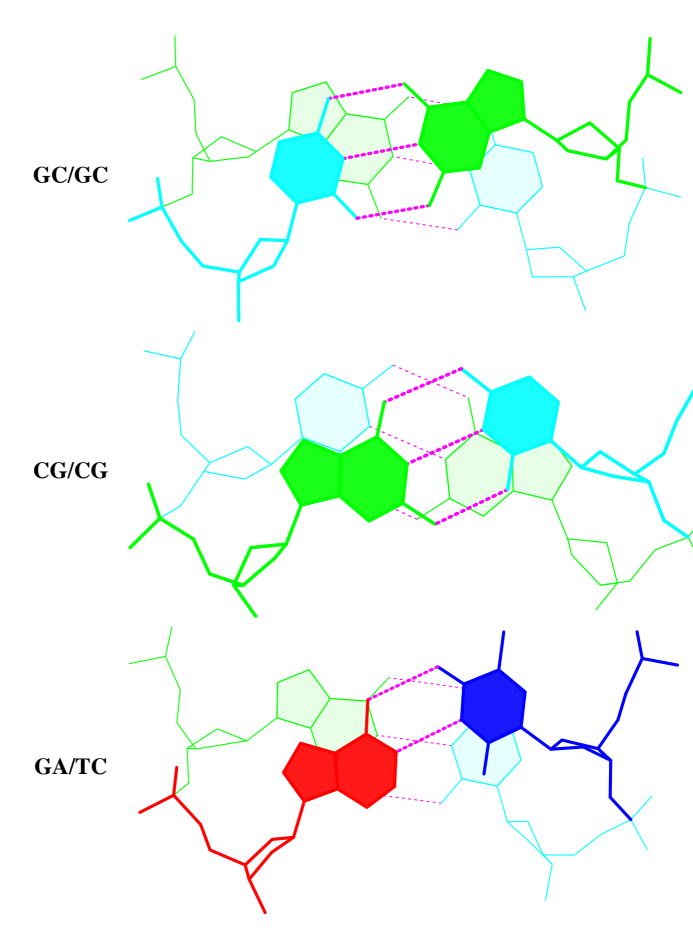


Figure 3: "Standardized" base stacking diagrams of three consecutive dimer steps of the 1.4~Å B-DNA structure d(CGCGAATTCGCG) $_2$ (Shui *et al.*, 1998) (BDL084).

- rotate_mol rotfile inpfile outfile

 A utility program for adjusting the orientation of a structure by rotations in

 MolScript style. In RasMol, a structure can be rotated etc, but the corresponding new coordinates can *not* be saved. rotate_mol is provided simply for this purpose: Rotate your structure in RasMol to the orientation you like, then write molscript rotfile.
- alc2img [options] [-s=factor] inpfile outfile Generating schematic representation of DNA from a ALCHEMY file.
- pdb2img [options] [-s=factor] inpfile outfile Generating schematic block representation of nucleic acid bases or protein peptide bonds directly from a PDB file.

3. Fiber models

• fiber options pdbfile

The options is -num, where num is between 1 and 38. -a|-b|-c|-d|-z

can be used for A-DNA, B-DNA, C-DNA, D-DNA and Z-DNA models.

4. Stacking diagrams

- stack2img [options] [-s=factor] inpfile outfile Program to generate "standardized" base-stacking diagram and atomic representations of nucleic acid structures with filled base rings and hydrogen bonds.
- mstack2img ['options'] stackfile n1 n2 prefix
 A Perl utility calling stack2img to generate stacking diagrams for a set of dinucleotide structures.

5. Set standard bases

• std_base [options] [angle1 distance angle2] inpfile outfile A utility program for setting standard base geometries used by analyze and rebuild.

6. Other utilities

• r3d_atom

A utility program for generating scenes of atomic CPK, ball-and-stick, filled-base-ring, hydrogen bonds between bases, colored by atomic type, base residue or in gray, for input to Raster3D. Some of its functionalities are similar to rods, balls and ring3d of the Raster3D distribution, but with a more general approach for locating base rings.

• cp_std prefix
A Perl script for changing standard base geometry files.

• ex_str -num [-nmr] pdbfile

A utility program for extracting a specific structure unit or an NMR structure from a multiple-structure PDB file.

- get_part [options] inpfile outfile Extracting protein, nucleic acid or other components from a PDB file.
- olp_o2p inpfile [outfile]
 Checking the designation of O1P and O2P atoms in a PDB file. Useful for RMS fittings.
- block_atom

A simple Perl script for generating block plus atomic representation of a nucleic acid structure in ALCHEMY format, which can be displayed using Ras-Mol. For example, with block_atom bdl084 (assuming a PDB data file bdl084.pdb exists), one get a file bdl084.alc, which can be display with rasmol -alchemy -noconnect bdl084.alc. Note the undocumented option -noconnect is needed to avoid RasMol recalculating bond connections.

dcmnfile
 A simple Perl script for clean-up. Various programs in 3DNA generate auxiliary files with fixed names which can be deleted with this utility program.

4 A tutorial introduction

This section shows some detailed examples on how to use 3DNA properly and effectively. Only some of the main features are covered for beginners to get started. As mentioned above, a serious user should go through the Examples directory provided with 3DNA and play with it.

The universal molecular graphics program rasmol (version 2.6+, but not 2.7.1) is indispensable. Also, gv (ghostview), xfig (version 3.2+) and Raster3D (version 2.5+) are highly recommended. Most likely they might be already available on your machine, since they are all extremely valuable freewares.

4.1 Base-pair geometry parameters

Figures 4 and 5 illustrate the definition of various parameters describing the base-pair geometry in a nucleic acid structure. These two figures were themselves generated with the help of 3DNA.

4.2 Analysis of the Tc3 transposon DNA

The crystal structure of Tc3 transposase in complex with transposon DNA was solved by van Pouderoyan *et al.* (1997). This transposon DNA structure is unusual in that it contains an A/B junction: the G+C rich end adopts the A-DNA conformation while the A+T rich part has the B-DNA conformation. As a consequence, this DNA is non-linear, *i.e.*, curved.

The NDB reference code for this structure is pde0128. Download the structure and display it with rasmol, you will find that protein and DNA are together. To analyze the helical DNA structure, the program (analyze) needs to know which base-

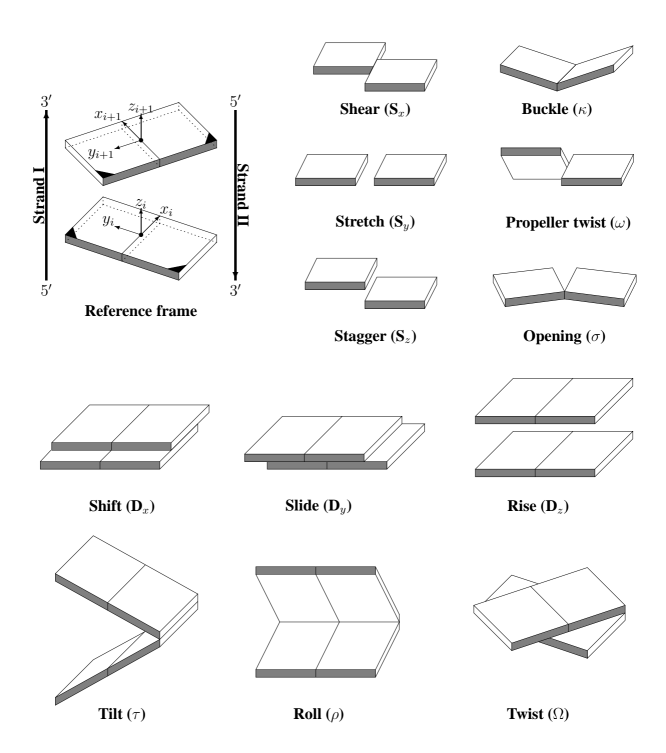


Figure 4: Pictorial definitions of parameters that relate complementary base pairs and sequential base-pair steps. The base-pair reference frame is constructed such that the *x*-axis points away from the (shaded) minor groove edge. Images illustrate positive values of the designated parameters (Dickerson *et al.*, 1989).

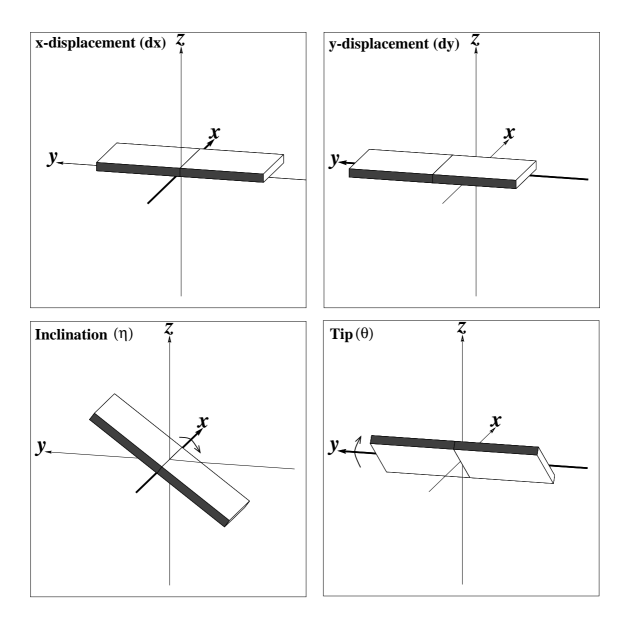


Figure 5: Helical parameters. Pictorial definitions of parameters that relate base pairs to its helical frame.

residue (sequential number in the PDB file) pairs with which. This information (in file pde0128.inp) can be obtained with the utility program find_pair as follows:

```
find_pair pde0128.pdb pde0128.inp
```

where pde0128.pdb is the data file in PDB format downloaded from the NDB and pdeo128.inp is the file required by analyze.

```
analyze pde0128.inp
```

The above two steps can be combined as follows:

```
find_pair pde0128.pdb stdout | analyze
```

Among several other output files, the one named pde0128.out (see page 36) gives a detailed listing of the DNA structural parameters. Each part is briefly summarized as follows:

- RMS deviation ··· gives the rms deviation between each experimental base and the standard reference. Normally the value is less than 0.05 Å. For each base, the corresponding PDB residue information is provided. A base-pair is classified as either Watson-Crick or non-Watson-Crick (denoted with a star).
- H-bonding information gives atom-list and their length in Å of all possible H-bonds in each of the base-pairs.
- ullet Overlap area in \mathring{A}^2 between polygons defined by atoms on successive bases. Polygons projected in the mean plane of the designed base-pair step.
 - Values in parentheses measure the overlap of base ring atoms only. Those outside parentheses include exocyclic atoms on the ring.
- Origin (Ox, Oy, Oz) ··· gives the origins and mean base-pair normal vectors in the coordinate system of the given structure. Dickerson *et al.* has found it very useful by drawing the Ny vs. Nx normal plot to demonstrate the curvature of a DNA molecule.
- Local base-pair parameters, *i.e.*, Shear, Stretch, Stagger, Buckle, Propeller, and Opening (Figure 4). *Generally speaking*, Propeller always has a value of about -10° in A- and B-DNA (Calladine & Drew, 1997), while Buckle can be either positive or negative of up to 20° . The other four show much less variations.
- Local base-pair step parameters gives the familiar Shift, Slide, Rise, Tilt, Roll, and Twist (Figure 4). Slide-Roll-Twist show more variations than the other three. Slide is the best single parameter among the six in discriminating Afrom B-DNA.
- Local base-pair helical parameters, *i.e.*. X-displacement, Y-displacement, Helical rise, Inclination, Tip, and Helical twist, (Figure 5) are also commonly used. For canonical A-DNA, for example, the X-displacement characterizes the "hole" in the top view (Figure 6), and the Inclination means that the bases are non-perpendicular to the helical axis in the side view (see cover image).

- lambda ··· gives the virtual angle between C1'-YN1 or C1'-RN9 glycosidic bonds and the base-pair C1'-C1' line. For mismatched base-pairs, the two λ angles are normally quite different.
- Classification of each dinucleotide step ··· is based on the Zp and ZpH values. It applies to right-handed A-, B- and TA-DNA with Watson-Crick base-pair steps. This classification should be checked against the step/helical parameters and the backbone torsion angles etc for final assignment. Generally speaking, this classification can be taken as a good starting point for pinpointing possible structural transitions in large DNA structures. For pde0128, this section says that the transposon DNA is not simply a deformed B-DNA, but one with an A/B junction.
- Minor and major groove widths are calculated based on the method proposed by El Hassan & Calladine (1998). It uses simple cross-strand P-P distances with and without an angle correction, and assign the groove widths to each dinucleotide step. Please note that the sum of vdW radii (5.8 Å) of the two phosphate groups is **not** subtracted from the values given here.
- Structure classification gives the overall assignment of the structure as right-handed (A-, B-, TA-DNA), left-handed Z-DNA, and other two topologically possible forms.
- Global linear helical axis is defined by equivalent C1' and RN9/YN1 atom pairs as in SCHNAaP. It gives the unit vector along the helical axis and the two end points which can be used to locate the axis by adding these two points as ATOM/HETATM records in your PDB file and draw a line bewteen them. While such a linear-fitting makes no much sense for a strongly curved DNA structure, the deviation from regular linear helix is a good measure of the overall structural deformation.
- Main chain and chi torsion angles.
- Sugar conformational parameters include five internal torsional angles, and amplitude and phase angle of pseudorotation of the sugar ring.
- Same strand P-P and C1'-C1' virtual bond distances have been used in the literature to discriminate A- and B-DNA conformations, although it is now clear that they are not as good as Zp and Slide.
- Helix radius gives the radial displacement of P, O4', and C1' atoms in local helix frame of each dimer.
- Position (Px, Py, Pz) and local helical axis vector (Hx, Hy, Hz) for each dinucleotide step.

The output file bp_step.par contains information for a rigorous reconstruction of the base-pair geometry. It will be overwritten each time when another analysis is performed unless it is renamed.

```
19 base-pairs
  0 ***local base-pair & step parameters***
      Shear Stretch Stagger Buckle Prop-Tw Opening Shift Slide Rise Tilt Roll Twist
      -0.18 -0.27 0.58 8.57 -9.77 -8.26 0.00 0.00 0.00 0.00 0.00
G-C
                                                                                          0.00
       0.16 -0.26
                    0.13 -5.72 -10.04 -3.52 0.19 -1.13 3.65 5.58 3.88
                                                                                          39.73
G-C
                     0.42 -1.54 -7.63
                                                    0.43 -1.67
                                                                    2.94 -3.14 10.32
      -0.10 -0.25
                                           -0.67
                                                                                          29.99
      0.02 -0.33 0.08 -3.56
                                    -6.03 1.74 0.17 -2.24 3.13 1.31 6.72 31.31
G-C
                                            3.13 0.08 -1.92 3.33 3.67 8.74 27.62
      -0.21 -0.24 -0.13 -3.78 -0.69
G-C
      0.04 \quad -0.22 \quad -0.35 \quad -8.83 \quad -11.08 \quad -3.03 \quad 0.50 \quad -1.91 \quad 3.41 \quad 4.23 \quad 5.60 \quad 31.56
G-C
      -0.32 -0.40 -0.38 -16.06 -17.27 -3.53 0.18 -2.14 3.43 4.68 10.29
                                                                                          27.30
      0.43 \quad -0.40 \quad -0.05 \quad -11.97 \quad -14.80 \quad 12.03 \quad -0.07 \quad -0.42 \quad 3.15 \quad -0.69 \quad 5.91 \quad 35.56
      0.52 \quad -0.43 \quad -0.34 \quad -10.52 \quad 4.23 \quad 3.87 \quad 0.25 \quad 0.76 \quad 3.46 \quad 3.31 \quad 0.88 \quad 37.83
C-G
      0.27 \quad -0.33 \quad 0.52 \quad -15.53 \quad -4.19 \quad -2.52 \quad -1.73 \quad 0.69 \quad 3.77 \quad -7.18 \quad -1.10 \quad 31.39
C-G
      -0.18 -0.23 -0.13 -9.98 -5.91 -1.56 0.35 -0.32 3.19 5.36 6.16
T-A
                                                                                          26.98
      -0.57 -0.49
                     0.47 0.67 -7.27 -0.77 0.41
                                                           1.89 3.06 -6.05 -2.74
A-T
                                                                                         45.60
                    0.33 -3.99 -14.34
      -0.34 -0.35
                                           -5.83 -0.34 -0.18
                                                                  3.44
                                                                           1.73
                                                                                 -3.42
                                                                                          31.69
                    -0.04 -2.98
                                                   0.29
                                           12.91
      -0.69
             -0.02
                                    -0.45
                                                            2.18
                                                                  3.37
                                                                          -2.75
                                                                                  -6.21
                                                                                          49.54
G-C
      -0.51 -0.07 -0.61 3.47
                                    -9.27 12.05 0.33
                                                           1.29 3.25 -0.15 15.17
                                                                                         23.41
      -0.66 -0.15 -0.12 12.32 -13.66 3.93 -0.68 0.30 2.96 -4.96 0.68 35.87
A-T
      -0.41 \quad -0.38 \quad 0.73 \quad 21.10 \quad -23.76 \quad -1.56 \quad -0.06 \quad -0.03 \quad 2.97 \quad -6.65 \quad -0.95 \quad 34.68
A-T
 \texttt{C-G} \qquad 0.40 \quad -0.55 \qquad 0.33 \quad -5.21 \quad -10.24 \quad -3.28 \qquad 0.37 \quad -1.03 \qquad 3.93 \qquad 4.89 \qquad 1.28 \quad 38.27 
T-A 0.56 -0.49 0.46 -21.17 -8.34 -4.72 -0.19 -1.07 3.61 1.91 4.78 37.16
```

Let's rebuild an atomic structure with the following command:

```
rebuild -atomic bp_step.par tc3_base.pdb
```

Depending on your setting of the standard base geometry (*i.e.*. Atomic_?.pdb files), you will get a structure with either only base atoms (the default) or with an approximate sugar-phosphate backbone attached. Use rasmol to have a look.

Your can also rebuild a schematic Calladine-Drew style picture with the following commands:

```
rebuild bp_step.par tc3_bp1.alc[for one block per base-pair]
rebuild -block2 bp_step.par tc3_bp2.alc[for one block
per base]
```

You need to use rasmol with the -alchemy command-line option to display files tc3_bp1.alc and tc3_bp2.alc since they are in ALCHEMY format. *e.g.*,

```
rasmol -alchemy tc3_bp2.alc
```

With this image, the Buckle and Propeller deformations are immediately obvious.

4.3 Fiber models

The 55 types of fiber nucleic acid models by Chandrasekaran & Arnott (1989) can be easily generated with the program fiber. Use fiber -m to get a list of all structures. Here I will use calf thymus A-DNA (number 1 in the list) as an example to illustrate its usage.

```
fiber -a fiber_A.pdb
```

Here -a is the same as -1 and fiber_A.pdb is the output file name. Your will then be asked to input your base sequence. It could either be from a data file (complete sequence) or from keyboard (enter only the repeating sequence, which is the default). Type enter means the default for input from keyboard. You are prompted for repeating unit with a default for polyA. Type atcg (either case is Okay and uncommon bases will be ignored) for a mixed A-T-C-G repeating sequence. Finally you are asked for the number of repeats (default is 10). Type 5 so you get (ATCG)₅, *i.e.*, 20 base-pairs. Display it with rasmol fiber_A.pdb.

High quality postscript picture (Figure 6) of fiber_A.pdb can be generated with the utility stack2img as follows:

```
stack2img -cao fiber_A.pdb fiber_A.ps
```

The option -cao means color-coded (c), atomic-model (a) with filled base-rings (o). If you add the option -f, you will get an image in XFIG format which you can easily edit. This is actually what these *atomic* options are intended for.

4.4 Input for render in Raster3D etc

The Calladine-Drew style base-pair representations are themselves quite useful as shown in Figures 1 and 2. They are, however, even more helpful when combined with atomic and schematic representations of ligands and proteins etc. This can be achieved with the render program of Raster3D as follows:

Use RasMol to find the view you want, and write molscript molfile at the command window. Then use the utility program rotate_mol to get a new PDB file with coordinates corresponding to your chosen view (RasMol does not write back coordinates in a new view). The new PDB coordinates are the *common reference* for scenes generated by different programs (3DNA, MolScript, Raster3D etc) to be properly rendered by Raster3D. Specifically, avoid any further coordinates transformation by MolScript or Raster3D.

You might use the utility get_part to divide a structure into nucleic acid part (default or -n), protein part (-p) and others (-o) to be used with different programs. For example, use 3DNA utilities r3d_atom and pdb2img/stack2img for nucleic acids, and MolScript for the protein part. You might need to delete the header part from MolScript with del_ms -n, a simple Perl script comes with 3DNA. The different scenes can then be concatenated together for render as detailed in Raster3D document.

Most of the images in 3DNA homepage were generated this way.

4.5 Build a DNA structure with sugar-phosphate backbone

The rebuilding of a DNA structure relies on its base sequence and associated base-pair and step parameters, and a set of standard Atomic_?.pdb data file.

The rebuilding algorithm is rigorous as far as the base-pair geometry is concerned. However, approximate sugar-phosphate conformations can be attached to the bases as a nucleic acid structure is built. This way, you get a full atomic structure with the sugar-phosphate backbone.

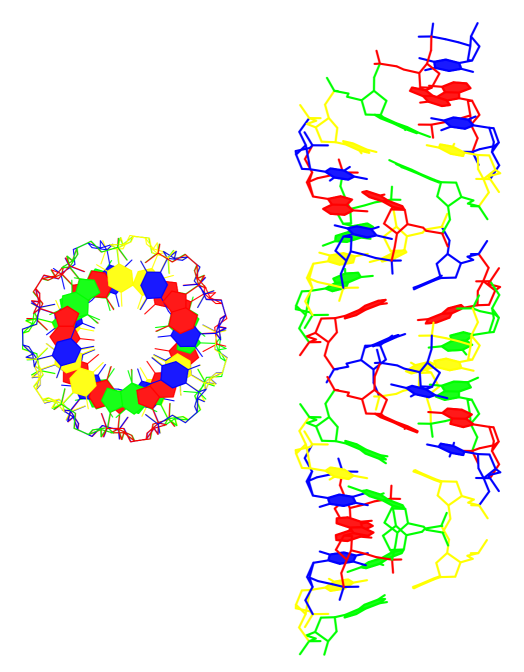


Figure 6: Top and side view images of the 20 base-pair long, (ATCG)₅, fiber A-DNA color coded by residue: A-red, T-blue, G-green, and C-yellow.

In directory BASEPARS/ATOMIC, there are four sets of standard PDB files, corresponding to A-DNA (C3'-endo), B-DNA (C2'-endo), RNA (C3'-endo plus O2') and NDB96 (default, without backbone). The base geometry in each set is identical to ensure the exact same numerical values of base pair parameters no matter which set is used in the analysis procedure.

To use a set other than the default NDB96, simply overwrite these files to their corresponding Atomic_?.pdb counterparts (e.g., BDNA_A.pdb to Atomic_A.pdb). A simple Perl script cp_std helps with this process. For example: cp_std BDNA will put the B-DNA set into your current directory.

3DNA searches for Atomic_?.pdb files in the following order:

- current working directory
- \$X3DNA/BASEPARS (default)
- \$HOME/X3DNA/BASEPARS (if environment variable \$X3DNA is not defined)

As an example, the procedure to rebuild bdl084 with standard B-DNA C2'-endo sugar-phosphate backbone conformation is as follows:

You will get file bp_step.par which contains base-pair sequence and step parameters. (Another file, bp_helical.par, which contains helical parameters, is also suitable for the following rebuilding process.)

Use RasMol to have a look at temp.pdb. It contains explicitly all valence bonds.

To verify that the algorithms used in 3DNA is rigorous and reversible, re-analyze the rebuilt structure temp.pdb, and you will get the same base-pair parameters as those directly from bd1084.pdb. Alternatively, you could superimpose temp.pdb to bd1084.pdb, and you will find that the RMSD (in Å) is virtually zero if only base atoms (i.e., excluding the backbone) are considered. If the sugar-phosphate backbone atoms (full-atom model) are also used, the results for three DNA structures in Examples/Analyze_Rebuild are as follows:

	pd0001	bd1084	adh026
base-atom	0.05	0.02	0.03
full-atom	0.82	0.73	0.52*

In rebuilding full-atomic models, regular sugar-phosphate backbone conformation was used: B-type for pd0001 and bd1084, and A-type for adh026. It should be noted that the labeling of O1P/O2P atoms in adh026 from the NDB is not consistent with the convention. Thus the RMSD values for full-atom model corresponds those after correction with a utility program o1p_o2p.

By matching Atomic_?.pdb with base sequence, it is also possible to generate a nucleic acid structure with mixed A- and B-DNA conformations, as in pde0128.pdb.

5 Some technical details

In our experience, following step-by-step working examples is the best way to understand any nontrivial algorithm. To make things right, every technical detail counts. Over the years, we have received many requests asking for exactly how 3DNA parameters are calculated. Here, we provide details of some key 3DNA components so that interested users could understand them better and possibly apply the same techniques to other related situations.

As an example, we will use the first dinucleotide step GG/CC from A-DNA adh026 (with base sequence GGGCGCC), included in 3DNA distribution, which shows Slide and Roll more clearly.

5.1 Least-squares fitting procedures

3DNA starts with a least-squares procedure to fit a standard base with an embedded reference frame to an observed base structure. It implements a closed-form solution of absolute orientation using unit quaternions first introduced by Horn (1987). This method can be applied when one or both of the structures are perfectly planar. This section is based on the following URL:

```
http://rutchem.rutgers.edu/~olson/jmb/ls_fit.html
```

• Standard reference frame (Olson et al., 2001)

Using base G as an example, its xyz coordinates in standard reference frame in PDB format are as follows (check BASEPARS directory for other cases):

ATOM	1	C1′	G A	1	-2.477	5.399	0.000
ATOM	2	N9	G A	1	-1.289	4.551	0.000
ATOM	3	C8	G A	1	0.023	4.962	0.000
ATOM	4	N7	G A	1	0.870	3.969	0.000
ATOM	5	C5	G A	1	0.071	2.833	0.000
ATOM	6	C6	G A	1	0.424	1.460	0.000
ATOM	7	06	G A	1	1.554	0.955	0.000
ATOM	8	N1	G A	1	-0.700	0.641	0.000
ATOM	9	C2	G A	1	-1.999	1.087	0.000
ATOM	10	N2	G A	1	-2.949	0.139	-0.001
ATOM	11	N3	G A	1	-2.342	2.364	0.001
ATOM	12	C4	G A	1	-1.265	3.177	0.000

• Least-squares fitting procedure

Least-squares fitting in 3DNA uses only (available) ring atoms: nine for purines (' N9 '; ' C8 '; ' N7 '; ' C5 '; ' C6 '; ' N1 '; ' C2 '; ' N3 '; ' C4 '), and six for pyrimidines (' N1 '; ' C2 '; ' N3 '; ' C4 '; ' C5 '; ' C6 ').

Using the first G residue in chain A of adh026, denoted thereafter as A_G1, the corresponding 9 ring atoms in standard (sX, sY, sZ) and experimental (eX, eY, eZ) structures are as follows:

		sX	sY	sZ	eX	eY	eZ
1	N9	-1.289	4.551	0.000	11.417	-2.904	-4.880
2	C8	0.023	4.962	0.000	10.759	-1.995	-5.662
3	N7	0.870	3.969	0.000	11.469	-0.913	-5.867
4	C5	0.071	2.833	0.000	12.638	-1.108	-5.156
5	С6	0.424	1.460	0.000	13.759	-0.273	-5.036
6	N1	-0.700	0.641	0.000	14.767	-0.848	-4.249
7	C2	-1.999	1.087	0.000	14.663	-2.116	-3.719
8	N3	-2.342	2.364	0.001	13.625	-2.934	-3.830
9	C4	-1.265	3.177	0.000	12.625	-2.328	-4.545
S	ave:	-0.6897	2.7827	0.0001	e ave:12.8580	-1.7132	-4.7716

Where s_ave and e_ave are the geometric centers of the nine ring atoms in the standard and experimental bases respectively.

We collect the two sets of coordinates in the 9 \times 3 matrices S and E corresponding respectively to the standard and experimental bases. We then construct 3 \times 3 covariance matrix (C) between S and E using the following formula:

-1.2530

-0.7886

0.7419

-0.7886

-0.1167

-1.3526

Here N, the number of atoms in each base, is 9, and i is an N \times 1 column vector consisting of only ones. S' and i' are the transpose of matrix S and column vector i respectively.

From the nine elements of C, we subsequently generate the 4 $\,\times\,$ 4 real symmetric matrix M using the expression:

0.7419

-1.3526

0.1165

3.0312 -0.7417

-0.7889

```
3.0312 \quad -0.7417 \quad -0.7889 \quad 1.2533
```

The largest eigenvalue of matrix M is 3.5896, and its corresponding unit eigenvector (qi, i = 0 -- 3) is:

$$[q0 q1 q2 q3] = [0.5460 -0.2921 0.0524 0.7835]$$

The rotation matrix R is deduced from the qi as:

Following coordinate transformation with matrix R, the translation vector for the least-squares fit between standard and experimental bases is defined as:

```
o = e_ave - s_ave R' = [15.1632 -0.0362 -4.4678]
```

Here o and R are the origin and orientation, i.e., the reference frame, of the experimental base. Moreover, R is guaranteed to be orthogonal.

The least-squares fitted coordinates (F) of the standard base atoms on the experimental structure are then given by:

```
F = S R' + o
 11.4307 -2.9119
                   -4.8818
 10.7607 -1.9933
                   -5.6548
 11.4432 -0.8991
                   -5.8558
 12.6361
         -1.1058
                   -5.1753
 13.7706 -0.2678
                   -5.0319
 14.7583
          -0.8689
                   -4.2591
 14.6659 -2.1180
                   -3.6958
          -2.9091
 13.6138
                   -3.8208
 12.6427
          -2.3449
                   -4.5688
```

Here S is the $(N \times 3)$ matrix of the original coordinates of the standard base.

The difference matrix (D) between F and E, the (N \times 3) matrix of the original coordinates of the experimental base, and the root-mean-square (RMS) deviation between the two structures are found to be:

```
D = E - F
 =
  -0.0137
          0.0079
                     0.0018
  -0.0017
           -0.0017
                     -0.0072
  0.0258
           -0.0139
                    -0.0112
  0.0019
           -0.0022
                     0.0193
  -0.0116
          -0.0052
                    -0.0041
  0.0087
           0.0209
                     0.0101
  -0.0029
           0.0020
                    -0.0232
  0.0112
           -0.0249
                    -0.0092
  -0.0177 0.0169
                    0.0238
```

RMS deviation = $sqrt(sum(d^2)/N) = 0.0236$

5.2 Base reference frames

Following the above least-squares procedure, we get the following for A_G1:

```
RMSD = 0.0236

o = [15.1632 -0.0362 -4.4678]

R = [

-0.2331 -0.8862 -0.4004

0.8249 -0.3983 0.4012

-0.5150 -0.2368 0.8238]
```

where the first column corresponds to x-axis, the 2nd to y-axis, and the 3rd to z-axis respectively.

Similarly for B_C8 (i.e., C8 on chain B) which pairs with A_G1, we have the following:

```
RMSD = 0.0229
  o = [14.9124]
                 0.2803 - 4.7498
  R = [
  -0.2339
             0.9100
                      0.3422 ==>
                                    -0.2339
                                             -0.9100
                                                       -0.3422
   0.7496
             0.3930
                     -0.5326
                                    0.7496
                                             -0.3930
                                                        0.5326
                             ==>
  -0.6191
                     -0.7741
                                 -0.6191
                                             -0.1320
                                                        0.7741]
             0.1320
                            ==>
```

Since their dot product is negative (-0.9884), the z-axis of B_C8 is anti-parallel to that of A_G1. We reverse the y- and z-axes of B_C8 (by a 180-degree rotation around the x-axis) to make its z-axis parallel to that of A_G1, as shown on the right side of R following the arrows.

For A_G2, we have:

```
RMSD = 0.0171

o = [14.8757 	 2.9250 	 -2.4635]

R = [

-0.6807 	 -0.6274 	 -0.3781
```

For B_C7, which pairs with A_G2, we have:

```
RMSD = 0.0213
o = [14.4982 	 3.0313 	 -2.3001]
R = [
-0.5797    0.6905    0.4326 ==> -0.5797    -0.6905    -0.4326
0.3207    0.6814    -0.6579 ==> 0.3207    -0.6814    0.6579
-0.7491    -0.2426    -0.6165 ==> -0.7491    0.2426    0.6165]
```

5.3 General rotation matrix

The general rotation matrix $\mathbf{R}_{\hat{\mathbf{u}}}(\varphi)$ (Eq. [1]) describes a rotation of magnitude φ about an arbitrary unit vector $\hat{\mathbf{u}} = u_1\hat{\mathbf{i}} + u_2\hat{\mathbf{j}} + u_3\hat{\mathbf{k}}$, where $\hat{\mathbf{i}}$, $\hat{\mathbf{j}}$, $\hat{\mathbf{k}}$ are unit vectors along the axes of the local Cartesian frame:

$$\mathbf{R}_{\hat{\mathbf{u}}}(\varphi) = \begin{bmatrix} \cos \varphi + (1 - \cos \varphi)u_1^2 & (1 - \cos \varphi)u_1u_2 - u_3\sin\varphi & (1 - \cos \varphi)u_1u_3 + u_2\sin\varphi \\ (1 - \cos \varphi)u_1u_2 + u_3\sin\varphi & \cos\varphi + (1 - \cos\varphi)u_2^2 & (1 - \cos\varphi)u_2u_3 - u_1\sin\varphi \\ (1 - \cos\varphi)u_1u_3 - u_2\sin\varphi & (1 - \cos\varphi)u_2u_3 + u_1\sin\varphi & \cos\varphi + (1 - \cos\varphi)u_3^2 \end{bmatrix},$$

$$(1)$$

5.4 Base-pair parameters

Given the above base reference frames for A_G1,B_C8 and A_G2,B_C7, we can calculate the base-pair parameters (Shear, Stretch, Stagger, Buckle, Propeller, and Opening). The procedure is exactly the same as for step parameters detailed below. Here we just give the parameters for the two base-pairs, and users who are interested in knowing the details should work them through and should get exactly the same numbers.

The "middle frame" used in calculating the bp parameters becomes the bp reference frame. For base pair A_G1-B_C8, we have:

For base pair A_G2-B_C7, we have:

```
o2 = [14.6869 2.9781 -2.3818]
R2 = [
-0.6319 -0.6594 -0.4072
0.3583 -0.7144 0.6010
-0.6873 0.2339 0.6877]
```

These two reference frames are used in the next two sections to calculate step and helical parameters.

5.5 Step parameters

Given the 2 bp reference frames above, the following procedures are used to calculate step parameters (Shift, Slide, Rise, Tilt, Roll and Twist). It follows CEHS definition, as detailed in SCHNAaP.

1. Hinge axis is the cross product between z1 (3rd column of R1) and z2 (3rd column of R2):

```
[-0.3724 \quad 0.4682 \quad 0.8013] \times [-0.4072 \quad 0.6010 \quad 0.6877]
= [-0.1596 \quad -0.0702 \quad -0.0332]. When normalized, it becomes: [-0.8993 \quad -0.3953 \quad -0.1870]
```

Geometrically, hinge axis is the intersection line between the two base pair planes.

- 2. The RollTilt angle (Γ , i.e., net bending angle) is the magnitude between z1 and z2, which is given by their dot product: $\Gamma = 10.2221$ Å.
- 3. Now we rotate R2 by $(-0.5 * \Gamma = -5.1111)$ degrees around the hinge axis (see Eq. [1]):

```
R_{inge}(-5.1111) = [
 0.9992 - 0.0152
                   0.0359
 0.0181
          0.9966
                   -0.0798
 -0.0345 0.0804 0.9962]
R2' = R_hinge(-5.1111) * R2 = [
 -0.6616 -0.6396
                   -0.3914
 0.4005
         -0.7426
                   0.5368
 -0.6340
         0.1984
                   0.7475]
```

Similarly, we rotate R1 by $(+0.5 * \Gamma)$ degrees along the hinge axis:

```
R1' = R_hinge(+5.1111) * R1 = [
-0.1982   -0.8986   -0.3914
0.7441   -0.3978   0.5368
-0.6381   -0.1848   0.7475]
```

By definition, the z-axes of R1' and R2' (the third column) are the same, i.e., after symmetric rotations, we have perfectly aligned the z-axes of the two bps.

4. The x-, y- and z-axes of the "middle frame" are simply the average between those of R1′ and R2′, and by definition, they are orthogonal. The origin of the bp is the geometric average of o1 and o2. For the above case, we have the "middle frame" as follows:

```
Rm = [ \\ -0.4490 & -0.8033 & -0.3914 \\ 0.5977 & -0.5955 & 0.5368 \\ -0.6642 & 0.0071 & 0.7475] \\ om = [14.8624 & 1.5501 & -3.4953]
```

5. The translational parameters (Shift, Slide, and Rise) are simply the projections of the vector linking from o1 to o2 onto the x-, y-, and z-axes of the "middle frame":

[Shift Slide Rise] =
$$(o2 - o1)$$
 * Rm
= $[-0.3509 2.8561 2.2270]$ * Rm
= $[0.3853 -1.4033 3.3349]$

6. Twist is the angle from y1 to y2 (or from x1 to x2 of the "rotated" R1' and R2' matrices respectively). Its sign is defined with reference to the "middle frame" z-axis (the 3rd column of Rm), following right- handed rule for positive Twist angle. More specifically, the magnitude of the angle between y1 and y2 is: 33.5296°. The signe is determined by:

$$(y1' \times y2')$$
 . $z =$
 $[-0.2162 \ 0.2965 \ 0.4129]$. $[-0.3914 \ 0.5368 \ 0.7475]$
 $= 0.5524$

So Twist is positive.

- 7. The phase angle (ϕ) is the angle from the hinge-axis to the "middle frame" y-axis (2nd column of Rm). By definition, hinge-axis lies in the xy-plane of the "middle-frame" since it is perpendicular to the z-axis of the "middle-frame". The phase angle also has a sign associate with it, determined in the same way as Twist shown above. In this case, $\phi = +16.9598^{\circ}$.
- 8. Roll is defined as:

$$\Gamma * \cos(\phi) = 10.2221 * \cos(16.9598 * \pi/180) = 9.7776^{\circ}.$$

Similarly, Tilt is defined as:

$$\Gamma * \sin(\phi) = 10.2221 * \sin(16.9598 * \pi/180) = 2.9818^{\circ}.$$

9. Overall, the six step parameters are:

```
Shift Slide Rise Tilt Roll Twist 0.3853 -1.4033 3.3349 2.9818 9.7776 33.5296
```

5.6 Local helical parameters

The geometric approach described below gives exactly the same numerical values as those from the RNA (Babcock *et al.*, 1994). The pivot point issue does not apply here: it is only used in defining the base pair reference frame. Given the reference frames of the two base pairs, the procedure used in 3DNA to calculate the local helical parameters (x-displacement, y-displacement, helical rise, inclination, tip, and helical rise) is analogous to the one detailed above for step parameters, by using a tip-inclination combination.

Please note that to define a local helical axis, we need **two** base-pair reference frames. 3DNA finds the single-helical axis (which is actually $dx \times dy$) that brings 1 to coincide with 2 by a Helical Twist angle. The position which this helix passes through is defined by Chasles' theorem as detailed in Figures 12 & 13 of Babcock *et al.* (1994). The calculation of x-displacement, y-displacement, tip and inclination is then exactly as described in SCHNAaP (Lu *et al.*, 1997a).

To make the above point clear, let's use A1-A2-A3 triplet as an example. First, A1-A2 define a local helical axis and a set of local base-pair helical parameters are calculated. In 3DNA, these parameters are defined in a symmetric manner that bp A1:T1 and bp A2:T2 have exactly the same values. Similarly, step A2-A3 defines another set of local base-pair helical parameters. Thus bp A2:T2 has two sets of helical parameters associated with it depending on its context, i.e., either with bp A1:T1 or with bp A3:T3. Moreover, the local Helical Rise and Helical Twist are directly related to a dinucleotide step. These are the reasons that "Local base-pair helical parameters" as given in 3DNA refer to base-pair steps. As a matter of fact, the schematic diagrams illustrating the local helical parameters (x-displacement, y-displacement, inclination and tip) as given in 3DNA website and this user's manual were based on two perfectly overlapped base pair blocks.

Using the first GG/CC dinucleotide step in adh026 as an example, the detailed precedure is as follows:

1. The local helical axis is defined by the cross product of (x2 - x1) and (y2 - y1):

```
[-0.3997 -0.4307 -0.1183] \times [0.2391 -0.3164 0.4190] = [-0.2179 0.1392 0.2294], which when normalized, gives: h = [-0.6303 0.4026 0.6638]
```

- 2. Local helical frame of base-pair 1:
 - TipInclination angle (Ψ , in magnitude) is the angle between unit vectors h and z1, and is 17.2279°.
 - Hinge axis is defined by a cross product from h to z1, normalized to give:
 [0.0399 0.8707 -0.4902]. It lies in the xy-plane of local helical frame given below.
 - Rotate R1 through the above hinge axis by negative TipInclination angle (i.e., -17.2279°) will align the resultant z-axis with h, which gives us the local helical reference frame:

```
H1 = R_hinge(-17.2279) * R1 = [
-0.1880   -0.7532   -0.6303
   0.7504   -0.5242   0.4026
-0.6337   -0.3973   0.6638]
```

- 3. Similarly, the local helical frame for base pair 2:
 - TipInclination angle (Ψ, in magnitude) is the angle between unit vectors h and z2, and is 17.2279°. Note that by (symmetric) definition, it is the same as for base pair 1.
 - Hinge axis is defined by a cross product from h to z2, normalized to give:
 [-0.4121 0.5511 -0.7256]
 - Rotate R2 through the above hinge axis by negative TipInclination angle (i.e., -17.2279°) will align the resultant z-axis with h, which gives us the local helical reference frame:

4. The "middle helical frame" is the average of H1 and H2, and by definition, it is orthogonal.

- 5. Helical twist is the angle from y(H1) to y(H2) (or from x(H1) to x(H2)) of the local helical reference frames defined above, with reference to the local helical axis h for sign determination: 35.0103° .
- 6. Helical rise is the projection of the vector linking o1 to o2 onto the local helical axis h: 2.8493 Å.
- 7. Use **base pair 1** to calculate tip and inclination (base pair 2 gives the same result):
 - Phase angle ψ is defined from hinge axis ([0.0399 0.8707 -0.4902]) to positive tip axis (2nd column of H1, [-0.7532 -0.5242 -0.3973]) with reference the local helical axis h ([-0.6303 0.4026 0.6638]) for sign control: 106.9598°.
 - Tip is defined as: $\Psi * \cos(\psi) = 17.2279 * \cos(106.9598 * \pi/180) = -5.0254^{\circ}$
 - Inclination is defined as: $\Psi * \sin(\psi) = 17.2279 * \sin(106.9598 * \pi/180) = 16.4787^{\circ}$

- 8. Get the origin of local helical frame of base-pair 1. It is based on Chasles' theorem as used in RNA (Babcock *et al.*, 1994, see Figure 13, page 141 for a detailed illustration). The procedure implemented in 3DNA is as follows:
 - Vector AB is defined by:

```
AB = [o2 - o1] - Helical_Rise * h = [-0.3509 2.8561 2.2270] - 2.8493 * [-0.6303 0.4026 0.6638] = [1.4451    1.7089    0.3357]
```

• Vector AD is defined by an angle of

```
90 - 0.5 * Helical_Twist = 72.4948
```

from vector AB with reference to h:

```
AD = R_h(72.4948) * AB' = [-0.5182   1.6306   -1.4812]'
```

R_h (72.4948) is the rotation matrix along h (Eq. [1]) by an angle of 72.4948°.

AB' changes row vector AB to a column vector to be compatible with the

3-by-3 rotation matrix. When normalized, the AD vector is: [-0.2290 0.7205 -0.654

• The magnitude of AD is defined by:

```
AD_magnitude =

0.5 * AB_magnitude / sin(0.5 * Helical_Twist * pi / 180)

= 0.5 * 2.2631 / 0.3008 = 3.7618
```

• We then get "D", the position vector where the local helical axis passes through. It is the origin of local helical frame of base pair 1.

The origin of the local helical frame of base pair 2 is given by:

```
o2_h = o1_h + Helical_Rise * h = [12.3805 3.9798 -5.1797]
```

The origin of the "middle helical frame" is the average of o1_h and o2_h:

```
[13.2785 3.4062 -6.1253]
```

9. X-displacement and y-displacement are defined as the projections of the vector from o1_h to o1 onto the x- and y-axis of local helical frame of base pair 1 (H1):

```
[01 - o1_h] * H1(:, 1:2) =
([15.0378 \ 0.1221 \ -4.6088] \ - \ [14.1764 \ 2.8326 \ -7.0709]) * H1(:, 1:2)
= [-3.7562 \ -0.2063]
```

So x-displacement = -3.7562 Å and y-displacement = -0.2063 Å.

Using base pair 2 gives exactly the same values due to symmetric definition.

10. Overall, the six local helical parameters are:

```
X-disp Y-disp Rise Incl. Tip Twist -3.7562 -0.2063 2.8493 16.4787 -5.0254 35.0103
```

5.7 Rebuilding based on local step parameters

The local step parameters defined in 3DNA are rigorous and thus reversible. Given a set of step parameters, the relative position and orientation of the two base pairs can be exactly reproduced by a rebuilding procedure as detailed in SCHNAap/SCHNArp (Lu *et al.*, 1997a; Lu *et al.*, 1997b). Here we provide step-by-step working example so users can understand the algorithm better.

Using the six step parameters of the first GG/CC step in adh026 as calculated above:

Shift	Slide	Rise	Tilt	Roll	Twist
0.3853	-1.4033	3.3349	2.9818	9.7776	33.5296

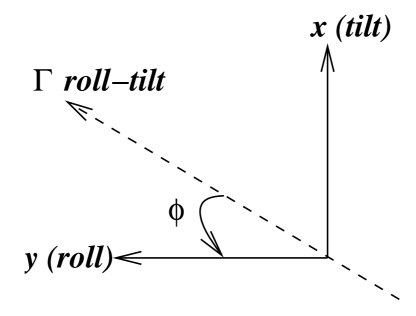


Figure 7: Roll-tilt combination

1. We first Twist base-pairs 2 and 1 about the "middle frame" z-axis (which at this stage is coincident with the z-axes of both base-pair reference frames) through $+\Omega/2$ and $-\Omega/2$ respectively:

$$T2 = \mathbf{R}_z(+\Omega/2), \qquad T1 = \mathbf{R}_z(-\Omega/2)$$
 (2)

Note that to be consistent with previous literatures, the symbol used for Twist here is still Ω (not ω), which is 33.5296°. The rotation matrix \mathbf{R}_z corresponds to the Eq. [1] with unit vector [0 0 1]. Similarly, \mathbf{R}_x corresponds to [1 0 0] and \mathbf{R}_y corresponds to [0 1 0]. Here, the "middle frame" is the "global" reference frame and corresponds to an identity matrix.

2. The RollTilt angle (Γ) is given by:

$$\Gamma = \sqrt{\rho^2 + \tau^2} = \sqrt{9.7776^2 + 2.9818^2} = 10.2221^{\circ}$$
 (3)

3. The Roll-Tilt axis is determined by the values of Roll and Tilt, and it lies in the x-y plane of the "middle frame". For the above case, it is [0.2917 0.9565 0], when normalized.

The angle from the Roll-Tilt axis ([0.2917 0.9565 0]) to the "middle frame" y-axis (Roll) ([0 1 0]) is called ϕ . Its sign is determined with reference to the "middle frame" z-axis ([0 0 1]), with values in range from -180° to 180° . In this case, it is 16.9598° .

4. We now rotate T2 and T1 through $+\Gamma/2$ and $-\Gamma/2$ respectively about the Roll-Tilt axis to give the final orientation of the two base-pairs with respect to the "middle frame". Analytically, the rotation about the Roll-Tilt axis can be expressed in terms of three rotations with reference to "middle frame" axes: First a rotation of angle ϕ around z-axis to align Roll-Tilt axis with y-axis, then a rotation of angle $\pm\Gamma/2$ around y-axis, and finally a rotation of angle $-\phi$ around z-axis to bring the Roll-Tilt axis back. Overall, the required rotations can be written as:

$$T2^{m} = [\mathbf{R}_{z}(-\phi) \mathbf{R}_{y}(+\Gamma/2) \mathbf{R}_{z}(+\phi)] T2$$

$$= \mathbf{R}_{z}(-\phi) \mathbf{R}_{y}(+\Gamma/2) \mathbf{R}_{z}(\phi + \Omega/2)$$

$$T1^{m} = [\mathbf{R}_{z}(-\phi) \mathbf{R}_{y}(-\Gamma/2) \mathbf{R}_{z}(+\phi)] T1$$

$$= \mathbf{R}_{z}(-\phi) \mathbf{R}_{y}(-\Gamma/2) \mathbf{R}_{z}(\phi - \Omega/2)$$
(5)

- 5. To get the correct positions of the origins of the reference frame of base-pair 2 and 1, we apply the translation vectors $(+D_x/2, +D_y/2, +D_z/2)$ and $(-D_x/2, -D_y/2, -D_z/2)$ respectively. Both translations are defined with reference to the "middle frame". Here D_x, D_y, D_z represent Shift, Slide and Rise.
- 6. The set of equations described above for the synthesis of a dinucleotide step defines the positions and orientations of the two base-pairs in a dinucleotide step with respect to the "middle frame". However, by means of a set of simple matrix transformations, we can change the reference frame from the "middle frame" to the reference frame of the base-pair 1. The orientation and position of base-pair 2 and the "middle frame" with respect to base-pair 1 are now given by the following equations:

Rm =
$$[T1^{\mathrm{m}}]^{-1} = \mathbf{R}_z(\Omega/2 - \phi) \mathbf{R}_y(\Gamma/2) \mathbf{R}_z(\phi)$$
 (6)

$$R2 = [\Gamma 1^{\mathrm{m}}]^{-1} \Upsilon 2^{\mathrm{m}} = \mathbf{R}_z(\Omega/2 - \phi) \mathbf{R}_y(\Gamma) \mathbf{R}_z(\Omega/2 + \phi)$$
 (7)

$$o2 = [D_x D_y D_z] Rm'$$
 (8)

Where $R_{\rm m}{}'$ means the transpose of the "middle frame" expressed with reference to base pair 1.

Using the above set of step parameters, we have the following:

```
Rm = [
   0.9537 -0.2873 0.0891
   0.2885
            0.9575
                     -0.0003
                      0.9960]
  -0.0852
           0.0260
R2 = [
   0.8204 -0.5436
                     0.1775
   0.5524 0.8336
                     -0.0006
            0.0985
  -0.1476
                      0.9841]
o2 = [0.3853 -1.4033 3.3349] * Rm' = [1.0677 -1.2336 3.2524]
```

Expressed in terms of the experimental coordinate reference frame of base pair 1, we get the reference frame for base pair 2 which are the same as shown in section 5.4: "Base-pair parameters"

5.8 Rebuilding based on local helical parameters

The local helical parameters defined in 3DNA are rigorous and thus reversible. Given a set of helical parameters, the relative position and orientation of the two base pairs can be exactly reproduced by a rebuilding procedure. Here we provide step-by-step working example so users can understand the algorithm better.

Using the six helical parameters of the first GG/CC step in adh026 as calculated above:

There are four frames for a dinucleotide step consisting of base-pairs 1 and 2: the base-pair frames \mathbf{R}_1 and \mathbf{R}_2 and their corresponding helical frames \mathbf{R}_{1h} and \mathbf{R}_{2h} . \mathbf{R}_{1h} and \mathbf{R}_1 are related by x-displacement, y-displacement, inclination and tip, so are \mathbf{R}_{2h} with \mathbf{R}_2 . \mathbf{R}_{1h} and \mathbf{R}_{2h} are related by *helical* rise and *helical* twist.

1. \mathbf{R}_{1h} as reference (1h in superscript)

Starting with \mathbf{R}_{1h} as the reference for the dinucleotide step concerned, then \mathbf{R}_{1h}^{1h} is the identity matrix \mathbf{I} :

$$\mathbf{R}_{1h}^{1h} = \mathbf{I} \tag{9}$$

 \mathbf{R}_{2h}^{1h} is got by the helical twist:

$$\mathbf{R}_{2h}^{1h} = \mathbf{R}_z(\Omega_h) \tag{10}$$

 \mathbf{R}_1^{1h} is got by a combined rotation of magnitude $\Psi (= \sqrt{\eta^2 + \theta^2})$ along tip-inclination axis (Fig. 8 (a)). Thus Ψ is the exact angle between base-pair normal and the local

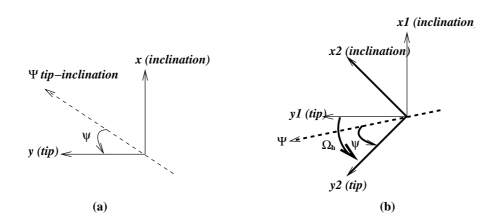


Figure 8: Combination of Tip and Inclination, and calculation of helical Twist.

helical helix. Let the angle from tip-inclination axis to tip axis be ψ , the rotation can be expressed as a rotation about z-axis by angle ψ , followed by a rotation about y-axis by angle Ψ , and then a rotation about z-axis by angle $-\psi$ to bring the axis back:

$$\mathbf{R}_1^{1h} = \mathbf{R}_z(-\psi) \; \mathbf{R}_y(\Psi) \; \mathbf{R}_z(\psi) \tag{11}$$

As shown in Fig. 8 (b), \mathbf{R}_{2}^{1h} is got by a helical twist, and then a rotation about the tip-inclination axis of \mathbf{R}_{2h}^{1h} (which is related to the tip y-axis of \mathbf{R}_{1h} by the helical twist Ω_h):

$$\mathbf{R}_{2}^{1h} = \left[\mathbf{R}_{z}(\Omega_{h} - \psi) \; \mathbf{R}_{y}(\Psi) \; \mathbf{R}_{z}(\psi - \Omega_{h})\right] \; \mathbf{R}_{z}(\Omega_{h}) = \mathbf{R}_{z}(\Omega_{h} - \psi) \; \mathbf{R}_{y}(\Psi) \; \mathbf{R}_{z}(\psi)$$
(12)

2. \mathbf{R}_1 as reference

With Eqs. 9 to 12, we can do some simple matrix transformations to make \mathbf{R}_1 as the reference:

$$\mathbf{R}_1 = \mathbf{I} \tag{13}$$

$$\mathbf{R}_{1h} = [\mathbf{R}_1^{1h}]^{-1} = \mathbf{R}_z(-\psi) \; \mathbf{R}_y(-\Psi) \; \mathbf{R}_z(\psi)$$
 (14)

$$\mathbf{R}_{2h} = [\mathbf{R}_1^{1h}]^{-1} \mathbf{R}_{2h}^{1h} = \mathbf{R}_z(-\psi) \mathbf{R}_y(-\Psi) \mathbf{R}_z(\psi + \Omega_h)$$
 (15)

$$\mathbf{R}_{2} = [\mathbf{R}_{1}^{1h}]^{-1} \mathbf{R}_{2}^{1h}$$

$$= \mathbf{R}_{z}(-\psi) \mathbf{R}_{y}(-\Psi) \mathbf{R}_{z}(\psi) \mathbf{R}_{z}(\Omega_{h} - \psi) \mathbf{R}_{y}(\Psi) \mathbf{R}_{z}(\psi)$$

$$= \mathbf{R}_{z}(-\psi) \mathbf{R}_{y}(-\Psi) \mathbf{R}_{z}(\Omega_{h}) \mathbf{R}_{y}(\Psi) \mathbf{R}_{z}(\psi)$$
(16)

Equ. 16 describes the orientation of base-pair 2 with reference to 1.

- 3. Position vector of base-pair 2 with reference to 1
 - From base-pair 1 origin to base-pair 1 helical origin: $-[dx dy] [x_{1h} y_{1h}]'$
 - From base-pair 1 helical origin to 2 helical origin: $dz \cdot z'_{11} = dz \cdot z'_{21}$

• From base-pair 2 helical origin to base-pair 2 origin: $+[dx dy] [x_{2h} y_{2h}]'$

Combining the above three items, we have the position (base-pair origin) of 2 relative to 1 as follows:

$$\mathbf{o_2} = [dx \ dy] \left\{ [x_{2h} \ y_{2h}]' - [x_{1h} \ y_{1h}]' \right\} + dz \cdot z'_{1h}$$
 (17)

Note that $x_{1h}, y_{1h}, z_{1h}, x_{2h}, y_{2h}, z_{2h}$ are 3×1 column vectors. 'means the transpose of a vector to change it to a row vector.

4. Using the example where inclination = 16.4787° and tip = -5.0254° , we have $\Psi = \sqrt{\eta^2 + \theta^2} = 17.2279^{\circ}$. The tip-inclination axis lies in the x-y plane of local helical frame of base pair 1: [16.4787 -5.0254 0]; which when normalized gives: [0.9565 -0.2917 0]. The angle from the tip-inclination axis ([0.9565 -0.2917 0]) to the base pair 1 helical y-axis (tip) ([0 1 0]) with reference to the helical z-axis ([0 0 1]) is $\psi = 106.9598^{\circ}$.

Using equations 15 to 17, we having the following:

Compared with the numbers based on step parameters, it is clear that Eqs. 16 and 7, Eqs. 17 and 8 are equivalent.

5.9 Relation between local helical and step parameters

To refer the orientation and position of one base-pair relative to the other, 6 parameters (3 rotations and 3 translations) are required. One set of such parameters is (Shift, Slide, Rise, Tilt, Roll and Twist), and the other set is (X-displacement, Y-displacement, Helical Rise, Inclination, Tip and Helical Twist).

Obviously these two sets should be completely reversible/dependent: from any one set you can get the other, rigorously. You can verify this point using step_hel, a utility program in 3DNA. Graphically this is best illustrated by the Calladine-Drew A to B transition model by introducing uniform Roll and Slide values at each dinucleotide step. The key point is that by introducing Roll, you also get Inclination, and with Slide, you get X-displacement.

The rebuild program in 3DNA can construct a DNA structure using either set of these parameters. Examples of such input files (e.g., bp_step.par and bp_helical.par) can be generated by analyze (Examples/Analyze_Rebuild directory.)

We have two sets of simple equations:

$$\theta/\eta = -\tau/\rho$$
$$2\cos\Omega_h = \cos\Omega(1+\cos\Gamma) - (1-\cos\Gamma)$$

6 Citation

Xiang-Jun Lu & Wilma K. Olson (2003). "3DNA: a software package for the analysis, rebuilding and visualization of three-dimensional nucleic acid structures." *Nucl. Acids. Res.* **31(17)**, 5108-5121.

7 Acknowledgments

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8 Appendix: 3DNA output for Tc3 transposon DNA

```
Detailed structural parameters for the analysis of Tc3 transposon DNA (data file pde0128.out)
************************
   3DNA (v1.5, Nov. 2002) by Xiang-Jun Lu at Wilma K. Olson's Lab.
***********************
1. The list of the parameters given below correspond to the 5' to 3' direction
  of strand I and 3' to 5' direction of strand II.
2. All angular parameters, except for the phase angle of sugar pseudo-
  rotation, are measured in degrees in the range of [-180, +180], and all
  displacements are measured in Angstrom units.
*******************
File name: pde0128.pdb
Date and time: Tue Nov 12 00:26:08 2002
Number of base-pairs: 19
Number of atoms: 1287
******************
HEADER
       PROTEIN/DNA
                                            07-JUL-97
                                                     1TC3
       TRANSPOSASE TC3A1-65 FROM CAENORHABDITIS ELEGANS
TITLE
       MOL_ID: 1;
COMPND
COMPND 2 MOLECULE: TC3 TRANSPOSASE;
COMPND 3 CHAIN: C;
COMPND 4 FRAGMENT: SPECIFIC DNA BINDING DOMAIN, RESIDUES 2 - 52;
COMPND 5 ENGINEERED: YES;
COMPND 6 MUTATION: C-TERMINAL 6-HIS TAG;
COMPND 7 MOL_ID: 2;
COMPND 8 MOLECULE: DNA
COMPND 9 (5'-D(*AP*GP*GP*GP*GP*GP*GP*TP*CP*CP*TP*AP*TP*AP*GP*A
COMPND 10 P*AP*CP*TP*T)-3');
COMPND 11 CHAIN: A;
COMPND 12 ENGINEERED: YES;
COMPND 13 MOL_ID: 3;
COMPND 14 MOLECULE: DNA
COMPND 15 (5'-D(*AP*GP*TP*TP*CP*TP*AP*TP*AP*GP*GP*AP*CP*CP*CP*C
COMPND 16 P*CP*CP*T)-3');
COMPND 17 CHAIN: B;
COMPND 18 ENGINEERED: YES
       MOL_ID: 1;
SOURCE
SOURCE 2 ORGANISM_SCIENTIFIC: CAENORHABDITIS ELEGANS;
SOURCE 3 STRAIN: BERGERAC;
SOURCE 4 VARIANT: TR679;
SOURCE 5 ORGANELLE: NUCLEUS;
SOURCE 6 GENE: TC3A;
SOURCE 7 EXPRESSION_SYSTEM: ESCHERICHIA COLI;
SOURCE 8 EXPRESSION_SYSTEM_STRAIN: BL21 (DE3) PLYSS;
SOURCE 9 EXPRESSION_SYSTEM_CELLULAR_LOCATION: CYTOPLASM;
SOURCE 10 EXPRESSION_SYSTEM_VECTOR_TYPE: PET3C;
SOURCE 11 EXPRESSION_SYSTEM_PLASMID: PRP1200;
```

SOURCE 12 EXPRESSION_SYSTEM_GENE: TC3A N1-65;

SOURCE 13 MOL ID: 2;

```
SOURCE 14 SYNTHETIC: YES;
SOURCE 15 MOL_ID: 3;
SOURCE 16 SYNTHETIC: YES
KEYWDS
        TRANSPOSASE, DNA BINDING, HELIX-TURN-HELIX, TC1/MARINER
KEYWDS
        2 FAMILY, COMPLEX (TRANSPOSASE/DNA), PROTEIN/DNA
EXPDTA
        X-RAY DIFFRACTION
AUTHOR
         G. VAN POUDEROYEN, R.F. KETTING, A. PERRAKIS, R.H.A. PLASTERK,
AUTHOR
        2 T.K.SIXMA
REVDAT 1 12-NOV-97
JRNL
           AUTH
                 G. VAN POUDEROYEN, R.F. KETTING, A. PERRAKIS,
JRNL
           AUTH 2 R.H.A.PLASTERK, T.K.SIXMA
                  CRYSTAL STRUCTURE OF THE SPECIFIC DNA-BINDING
JRNL
           TITL 2 DOMAIN OF TC3 TRANSPOSASE OF C. ELEGANS IN COMPLEX
JRNL
           TITL 3 WITH TRANSPOSON DNA
JRNL
JRNL
           REF
                  EMBO J.
                                              V. 16 6044 1997
                  ASTM EMJODG UK ISSN 0261-4189
                                                               0897
JRNL
           REFN
        1 1 ASP C 209 LEU C 220 1
HELIX
HELIX
        2
            2 LEU C 225
                         ILE C 232 1
HELIX
            3 ARG C 236 LYS C 244 1
RMSD of the bases (---- for WC bp, + for isolated bp, x for helix change)
           Strand I
                                      Strand II
                                                        Helix
  1
      (0.025) A:...2_:[..G]G----C[..C]:.119_:B (0.016)
  2
      (0.028) A: ...3_: [..G]G----C[..C]: .118_:B (0.019)
  3
      (0.022) A: ... 4_: [...G]G-----C[...C]: .117_:B (0.014)
  4
      (0.017) A: ...5_: [..G]G----C[..C]: .116_: B (0.017)
  5
      (0.027) A: ...6_: [..G]G----C[..C]: .115_:B (0.020)
  6
      (0.020) A:...7_:[..G]G----C[..C]:.114_:B (0.014)
  7
      (0.034) A: ...8_{:}[..G]G----C[..C]: .113_{:}B (0.019)
      (0.019) A: ...9_{:[..T]T----A[..A]:.112_{:B}} (0.020)
  8
  9
      (0.009) A:..10_:[..C]C----G[..G]:.111_:B (0.017)
  10
      (0.010) A:..11_:[..C]C----G[..G]:.110_:B (0.034)
  11
      (0.023) A:..12_:[..T]T----A[..A]:.109_:B (0.016)
  12
      (0.023) A:..13_{:}[..A]A----T[..T]:.108_{:}B (0.021)
  13
      (0.018) A:..14_:[..T]T----A[..A]:.107_:B (0.024)
  14
      (0.026) A:..15_:[..A]A----T[..T]:.106_:B (0.018)
      (0.017) A:..16_:[..G]G----C[..C]:.105_:B (0.025)
  15
      (0.033) A:..17_{:}[..A]A----T[..T]:.104_{:}B (0.019)
  16
  17
      (0.031) A:..18_{:}[..A]A----T[..T]:.103_{:}B (0.017)
  18
      (0.012) A:..19_:[..C]C----G[..G]:.102_:B (0.026)
       (0.021) A:..20_{:}[..T]T----A[..A]:.101_{:}B (0.023)
*******************
Detailed H-bond information: atom-name pair and length [ON]
  1 G----C [3] 06 - N4 2.53 N1 - N3 2.85 N2 - O2 3.00
  2 G----C [3] O6 - N4 2.61 N1 - N3 2.80 N2 - O2 2.77
   3 G----C [3]
                 06 - N4 2.86 N1 - N3 2.80 N2 - O2 2.62
  4 G----C [3] O6 - N4 2.72 N1 - N3 2.64 N2 - O2 2.40
  5 G----C [3] O6 - N4 2.89 N1 - N3 2.76 N2 - O2 2.59
  6 G----C [3]
                 06 - N4
                          2.64 N1 - N3 2.84 N2 - O2
                                                       2.87
  7 G----C [3]
                 06 - N4 2.57 N1 - N3
                                        2.67
                                             N2 - O2
                                                       2.83
  8 T----A [2] N3 - N1 2.51 O4 - N6 2.90
```

2.64 N4 - O6 2.83

9 C----G [3] O2 - N2 2.38 N3 - N1

```
10 C-----G [3] O2 - N2 2.66 N3 - N1 2.74 N4 - O6 2.78

11 T----A [2] N3 - N1 2.74 O4 - N6 2.77

12 A----T [2] N6 - O4 2.67 N1 - N3 2.61

13 T----A [2] N3 - N1 2.60 O4 - N6 2.60

14 A----T [2] N6 - O4 3.52 N1 - N3 2.86

15 G----C [3] O6 - N4 3.43 N1 - N3 2.94 N2 - O2 2.60

16 A----T [2] N6 - O4 3.13 N1 - N3 2.91

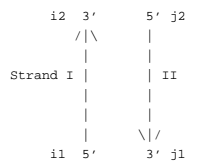
17 A----T [2] N6 - O4 2.91 N1 - N3 2.63

18 C----G [3] O2 - N2 2.47 N3 - N1 2.56 N4 - O6 2.57

19 T----A [2] N3 - N1 2.65 O4 - N6 2.66
```

Overlap area in Angstrom² between polygons defined by atoms on successive bases. Polygons projected in the mean plane of the designed base-pair step.

Values in parentheses measure the overlap of base ring atoms only. Those outside parentheses include exocyclic atoms on the ring. Intra- and inter-strand overlap is designated according to the following diagram:



	step	i1-i2	i1-j2	j1-i2	j1-j2	sum
1	GG/CC	4.01(2.20)	0.00(0.00)	0.03(0.00)	0.81(0.00)	4.85(2.20)
2	GG/CC	3.97(2.70)	0.00(0.00)	0.19(0.00)	0.00(0.00)	4.16(2.70)
3	GG/CC	3.18(1.65)	0.00(0.00)	1.16(0.00)	0.00(0.00)	4.34(1.65)
4	GG/CC	3.36(1.96)	0.00(0.00)	0.56(0.00)	0.00(0.00)	3.93(1.96)
5	GG/CC	4.26(2.90)	0.00(0.00)	0.43(0.00)	0.00(0.00)	4.69(2.90)
6	GG/CC	3.30(1.83)	0.00(0.00)	0.61(0.00)	0.00(0.00)	3.91(1.83)
7	GT/AC	6.08(1.72)	0.00(0.00)	0.00(0.00)	4.79(3.35)	10.87(5.06)
8	TC/GA	5.10(0.61)	0.00(0.00)	0.00(0.00)	4.69(1.63)	9.78(2.25)
9	CC/GG	1.18(0.00)	0.00(0.00)	0.00(0.00)	8.23(5.73)	9.41(5.73)
10	CT/AG	7.29(1.31)	0.00(0.00)	0.00(0.00)	2.86(1.70)	10.14(3.01)
11	TA/TA	2.56(0.00)	0.00(0.00)	0.00(0.00)	1.53(0.00)	4.08(0.00)
12	AT/AT	5.73(1.71)	0.00(0.00)	0.00(0.00)	6.31(2.41)	12.05(4.12)
13	TA/TA	1.76(0.00)	0.00(0.00)	0.00(0.00)	2.27(0.00)	4.03(0.00)
14	AG/CT	6.78(4.02)	0.00(0.00)	0.00(0.00)	4.40(0.27)	11.18(4.29)
15	GA/TC	3.83(0.87)	0.00(0.00)	0.00(0.00)	4.69(0.53)	8.52(1.40)
16	AA/TT	4.06(2.70)	0.00(0.00)	0.00(0.00)	5.91(0.32)	9.97(3.02)
17	AC/GT	5.32(4.25)	0.00(0.00)	0.00(0.00)	7.39(2.57)	12.70(6.81)
18	CT/AG	4.16(0.08)	0.00(0.00)	0.18(0.00)	3.04(2.60)	7.39(2.68)
* * * :	*****	*****	*****	*****	*****	*****

Origin (Ox, Oy, Oz) and mean normal vector (Nx, Ny, Nz) of each base-pair in the coordinate system of the given structure

bp	Ox	Oy	Oz	Nx	Ny	Nz
1 G-C	-0.32	155.89	29.45	0.62	-0.76	-0.20
2 G-C	2.94	154.00	28.78	0.70	-0.69	-0.15

3	G-C	5.94	152.64	29.63	0.62	-0.79	-0.01	
4	G-C	8.66	150.82	31.66	0.55	-0.83	0.07	
5	G-C	10.24	148.16	33.94	0.42	-0.90	0.15	
6	G-C	10.19	144.79	35.97	0.30	-0.94	0.17	
7	G-C	9.09	141.17	37.41	0.14	-0.99	0.07	
8	T-A	9.15	137.99	37.55	0.15	-0.99	-0.03	
9	C-G	10.37	134.67	37.58	0.12	-0.99	-0.08	
10	C-G	9.49	130.73	38.76	0.14	-0.99	0.04	
11	T-A	10.52	127.69	38.52	0.26	-0.97	-0.03	
12	A-T	10.13	124.46	40.11	0.15	-0.99	0.01	
13	T-A	10.83	121.09	39.68	0.18	-0.98	-0.05	
	A-T	9.50	117.55	38.32	0.19	-0.97	-0.17	
15	G-C	8.98	114.42	36.82	-0.05	-1.00	-0.08	
	A-T	9.55	111.47	36.25	-0.05	-0.98	-0.17	
	A-T	9.56	108.56	35.71	0.04	-0.97	-0.24	
	C-G	8.48	104.79	34.55	-0.05	-0.97	-0.25	
	T-A	7.31	101.48	33.18	-0.03	-0.94	-0.33	
		********						***
		-pair parame						
	р	Shear	Stretch	Stagger	Buckle	Propeller	Opening	
	G-C	-0.18	-0.27	0.58	8.57	-9.77	-8.26	
	G-C	0.16	-0.26	0.13	-5.72	-10.04	-3.52	
	G-C	-0.10	-0.25	0.42	-1.54	-7.63	-0.67	
	G-C	0.02	-0.33	0.08	-3.56	-6.03	1.74	
	G-C	-0.21	-0.24	-0.13	-3.78	-0.69	3.13	
	G-C	0.21	-0.22	-0.35	-8.83	-11.08	-3.03	
	G-C G-C	-0.32	-0.22	-0.38	-16.06			
						-17.27	-3.53	
	T-A	0.43	-0.40	-0.05		-14.80	12.03	
	C-G	0.52	-0.43	-0.34	-10.52	4.23	3.87	
	C-G	0.27	-0.33	0.52	-15.53	-4.19	-2.52	
	T-A	-0.18	-0.23	-0.13	-9.98	-5.91	-1.56	
	A-T					-7.27		
	T-A	-0.34	-0.35	0.33			-5.83	
	A-T	-0.69				-0.45		
	G-C	-0.51	-0.07	-0.61	3.47			
	A-T		-0.15					
	A-T	-0.41		0.73				
	C-G					-10.24		
19	T-A		-0.49			-8.34		
		~~~~~~~~~						
	ave.	-0.09						
	s.d.					6.42		
*****	****	*****	****	*****	*****	*****	*****	***
Local	base-	-pair step p	arameters					
st	ep	Shift			Tilt	Roll	Twist	
	G/CC		-1.13	3.65	5.58	3.88	39.73	
2 G	G/CC	0.43	-1.67	2.94	-3.14	10.32	29.99	
3 G	G/CC	0.17	-2.24	3.13	1.31	6.72	31.31	
4 G	G/CC	0.08	-1.92	3.33	3.67	8.74	27.62	
5 G	G/CC	0.50	-1.91	3.41	4.23	5.60	31.56	
6 G	G/CC	0.18	-2.14	3.43	4.68	10.29	27.30	
7 G	ST/AC	-0.07	-0.42	3.15	-0.69	5.91	35.56	

8 TC/GA 0.25 0.76 3.46 3.31 0.88 37.83

_	CC/GG	-1.73	0.69	3.77	-7.18	-1.10	31.39				
10	CT/AG	0.35	-0.32	3.19	5.36	6.16	26.98				
11	TA/TA	0.41	1.89	3.06	-6.05	-2.74	45.60				
12	AT/AT	-0.34	-0.18	3.44	1.73	-3.42	31.69				
13	TA/TA	0.29	2.18	3.37	-2.75	-6.21	49.54				
14	AG/CT	0.33	1.29	3.25	-0.15	15.17	23.41				
15	GA/TC	-0.68	0.30	2.96	-4.96	0.68	35.87				
16	AA/TT	-0.06	-0.03	2.97	-6.65	-0.95	34.68				
17	AC/GT	0.37	-1.03	3.93	4.89	1.28	38.27				
18		-0.19									
		0.03									
		0.53									
	********************										
		air helical			_						
		X-disp									
		-2.13									
		-4.56				5.83					
		-5.13				-2.38					
		-5.58		2.60							
		-4.42									
	GG/CC			2.47							
7	GT/AC	-1.48									
	TC/GA			3.48							
9		1.48		4.03		13.05					
10		-2.07				-11.16	28.17				
11		2.64				7.75					
12		0.32				-3.15					
		3.03				3.26					
		-1.36				0.33					
15	GA/TC	0.40	0.46	3.03	1.09	8.01	36.20				
16	AA/TT	0.09	-0.80	2.93	-1.57	11.03	35.30				
17	AC/GT	-1.75	0.16	3.92	1.95	-7.42	38.59				
	OTT / 7. O	-2.34									
18											
18	~ ~ .	-1 56									
18	ave.	-1.56 2.81	0.07	3.12	7.36	-0.80	35.16				

Structure classification:

This is a right-handed nucleic acid structure

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lambda: virtual angle between Cl'-YN1 or Cl'-RN9 glycosidic bonds and the base-pair Cl'-Cl' line

C1'-C1': distance between C1' atoms for each base-pair RN9-YN1: distance between RN9-YN1 atoms for each base-pair RC8-YC6: distance between RC8-YC6 atoms for each base-pair

bp	lambda(I)	lambda(II)	C1'-C1'	RN9-YN1	RC8-YC6
1 G-C	51.0	58.7	10.7	9.0	9.8
2 G-C	58.2	48.2	10.6	8.9	9.7
3 G-C	62.1	54.4	10.3	8.8	9.8

4	G-C	51.7	52.9	10.5	8.7	9.7
5	G-C	51.6	55.9	10.4	8.7	9.8
6	G-C	54.2	60.3	10.5	8.9	9.8
7	G-C	62.5	57.0	10.2	8.7	9.5
8	T-A	57.3	53.0	9.9	8.3	9.5
9	C-G	52.6	54.7	10.3	8.6	9.6
10	C-G	49.3	52.7	10.6	8.8	9.6
11	T-A	56.0	57.1	10.4	8.8	9.7
12	A-T	54.9	59.9	10.1	8.6	9.5
13	T-A	55.0	50.7	10.5	8.8	9.7
14	A-T	49.2	62.3	10.4	8.7	9.9
15	G-C	59.0	60.5	10.2	8.8	9.9
16	A-T	53.9	62.2	10.3	8.8	9.8
17	A-T	45.7	51.3	10.5	8.5	9.5
18	C-G	57.3	57.8	10.2	8.7	9.5
19	T-A	46.6	45.9	10.4	8.5	9.3
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Classification of each dinucleotide step in a right-handed nucleic acid structure: A-like; B-like; TA-like; intermediate of A and B, or other cases

ste	р Хр	Үр	Zp	ХрН	ҮрН	ZpH	Form	
1 GG	/CC -2.03	8.22	1.96	-4.14	8.00	2.77	A	
2 GG	/CC -2.00	8.35	2.44	-6.36	7.15	4.91	A	
3 GG	/CC -1.96	8.21	2.78	-6.89	7.47	4.41	A	
4 GG	/CC -1.61	8.45	2.50	-6.98	7.34	4.87	A	
5 GG	/CC -1.44	8.13	2.57	-5.65	7.57	3.94	A	
6 GG	/CC -2.30	8.54	1.72	-8.31	7.41	4.67	A	
7 GT	/AC -3.10	8.76	0.52	-4.51	8.57	1.89		
8 TC	/GA -3.37	8.71	-0.37	-2.26	8.72	-0.19	В	
9 CC	/GG -2.98	8.85	-0.08	-1.58	8.84	-0.56	В	
10 CT	/AG -3.21	8.87	-0.51	-4.98	8.79	1.29	В	
11 TA	/TA -2.15	8.08	-0.04	0.28	8.06	-0.45	В	
12 AT	/AT -3.22	8.85	-0.34	-2.89	8.76	-1.29	В	
13 TA	/TA -2.31	8.32	-0.37	0.44	8.22	-1.32	В	
14 AG	/CT -3.69	8.49	-0.96	-5.02	7.68	3.76	В	
15 GA	/TC -3.87	8.83	-1.10	-3.34	8.85	-0.88	В	
16 AA	/TT -3.52	8.81	0.03	-3.19	8.81	-0.12	В	
17 AC	/GT -3.62	9.14	0.44	-5.27	9.12	0.70	В	
18 CT	/AG -2.81	9.09	0.83	-5.00	8.91	1.98		
*****	******	*****	* * * * * * * *	*****	*****	*****	*****	*****

Minor and major groove widths: direct P-P distances and refined P-P distances which take into account the directions of the sugar-phosphate backbones

(Subtract 5.8 Angstrom from the values to take account of the vdw radii of the phosphate groups, and for comparison with FreeHelix and Curves.)

Ref: M. A. El Hassan and C. R. Calladine (1998). "Two Distinct Modes of Protein-induced Bending in DNA." J. Mol. Biol., v282, pp331-343.

	Minor	Groove	Major Groove		
	P-P	Refined	P-P	Refined	
1 GG/CC					
2 GG/CC					

3	GG/CC	16.3		17.1	
4	GG/CC	17.0	16.0	20.4	18.6
5	GG/CC	16.0	15.3	20.8	18.5
6	GG/CC	14.7	13.8	18.6	18.0
7	GT/AC	13.9	13.2	17.9	17.9
8	TC/GA	12.8	12.5	16.4	16.2
9	CC/GG	13.0	13.0	15.9	15.9
10	CT/AG	13.1	13.1	17.3	17.2
11	TA/TA	11.6	11.6	15.8	15.5
12	AT/AT	10.5	10.5	17.6	16.9
13	TA/TA	11.8	11.8	17.2	16.6
14	AG/CT	14.0	14.0	16.8	16.7
15	GA/TC	13.7	13.6	16.1	15.7
16	AA/TT	12.2		19.2	
17	AC/GT				
18	CT/AG				

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Global linear helical axis defined by equivalent C1' and RN9/YN1 atom pairs Deviation from regular linear helix: 2.85(0.76)

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Main chain and chi torsion angles:

Note: alpha: 03'(i-1)-P-05'-C5'

beta: P-O5'-C5'-C4'
gamma: O5'-C5'-C4'-C3'
delta: C5'-C4'-C3'-O3'
epsilon: C4'-C3'-O3'-P(i+1)
zeta: C3'-O3'-P(i+1)-O5'(i+1)

chi for pyrimidines(Y): 04'-C1'-N1-C2
 chi for purines(R): 04'-C1'-N9-C4

## Strand I

base	alpha	beta	gamma	delta	epsilon	zeta	chi
1 G		-170.2	73.1	82.0	-152.8	-70.0	-172.9
2 G	-79.5	-178.4	49.9	90.6	-149.4	-67.0	-162.6
3 G	-78.2	169.8	57.7	73.5	-163.3	-64.3	-172.9
4 G	-92.9	-177.5	71.5	76.5	-140.6	-93.3	-176.2
5 G	-59.6	158.4	55.7	67.7	-155.6	-61.5	-168.2
6 G	-71.0	161.4	62.0	73.1	-156.0	-88.8	-177.2
7 G	123.5	-165.1	-137.7	125.7	-147.4	-82.7	-164.8
8 T	-176.7	-142.6	149.5	101.5	-148.3	-61.2	-153.5
9 C	-58.8	-174.1	21.1	159.4	-110.7	171.2	-79.4
10 C	-92.0	143.7	68.4	154.1	-156.8	-107.9	-122.7
11 T	-35.9	-174.3	-6.0	169.1	-117.1	167.1	-69.6
12 A	-93.0	148.2	66.1	161.4	-120.5	-151.0	-109.0
13 T	-33.2	120.8	38.5	116.9	-123.5	-172.6	-123.3
14 A	-57.0	149.4	48.3	158.9	-164.6	-84.7	-94.1
15 G	-82.2	-176.7	40.0	154.1	-96.9	159.0	-69.5
16 A	-75.0	135.8	50.2	121.8	-147.3	-154.9	-112.8
17 A	6.5	98.0	45.3	80.0	177.5	-81.0	-136.9
18 C	105.5	-173.8	-165.4	81.5	83.6	54.5	-153.6
19 T	-142.8	170.5	85.0	80.9			-167.3

Strand	II						
base	alpha	beta	gamma	delta	epsilon	zeta	chi
1 C	-72.4	-176.5	67.2	128.1			-129.0
2 C	-43.5	172.3	43.4	148.7	-173.4	-112.6	-122.5
3 C	142.3	-163.6	-168.5	97.5	-136.0	-81.8	-170.0
4 C	-60.4	174.5	48.5	83.2	-179.6	-85.3	-166.2
5 C	-43.3	172.4	34.5	79.4	-151.6	-66.2	-155.8
6 C	107.3	-161.6	-159.1	89.0	-141.1	-88.0	-165.4
7 C	161.3	-116.0	148.8	88.1	-155.2	-103.4	-151.3
8 A	-78.6	126.0	59.1	86.8	175.4	-81.2	-135.3
9 G	-69.2	179.3	44.6	141.5	-109.5	-178.9	-92.7
10 G	-99.0	-157.7	76.8	111.6	-174.7	-74.4	-108.6
11 A	-64.6	168.0	47.6	139.1	153.4	-72.2	-88.7
12 T	-82.1	-174.6	63.1	146.1	-132.6	-163.4	-112.7
13 A	-70.6	130.0	55.8	140.4	175.2	-94.7	-97.2
14 T	-82.3	-167.0	69.2	156.5	-102.6	171.9	-88.1
15 C	-92.1	153.6	77.9	78.5	-170.8	-83.1	-129.7
16 T	38.9	90.6	38.3	104.6	-156.8	-86.5	-119.6
17 T	154.2	-128.5	165.8	80.1	-145.5	-126.3	-179.1
18 G	98.2	-172.9	-121.4	136.6	176.7	-104.8	-131.4
19 A			-157.3	94.7	-173.6	-100.2	-137.4

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## Sugar conformational parameters:

Note: v0: C4'-O4'-C1'-C2'
v1: O4'-C1'-C2'-C3'
v2: C1'-C2'-C3'-C4'
v3: C2'-C3'-C4'-O4'
v4: C3'-C4'-O4'-C1'

tm: amplitude of pseudorotation of the sugar ring
P: phase angle of pseudorotation of the sugar ring

Strand	d I							
base	v0	v1	v2	v3	v4	tm	Р	Puckering
1 0	1.7	-26.8	40.6	-40.2	24.6	42.3	16.2	C3'-endo
2 0	11.7	-33.7	42.2	-36.2	15.7	42.2	2.8	C3'-endo
3 0	-5.5	-25.6	45.2	-50.0	35.1	49.9	25.1	C3'-endo
4 0	-22.3	-11.0	38.7	-52.3	46.7	52.7	42.8	C4′-exo
5 G	-5.4	-28.9	50.6	-55.1	38.0	55.4	24.1	C3'-endo
6 0	-4.0	-27.0	47.5	-51.1	34.3	51.6	23.1	C3'-endo
7 0	39.7	-32.4	14.6	7.4	-28.7	38.0	292.5	C1'-endo
r 8	-21.3	7.8	7.4	-19.8	25.5	25.3	73.1	04'-endo
9 (	-22.0	44.1	-47.8	36.3	-9.4	48.3	172.1	C2'-endo
10 0	-32.4	50.5	-49.1	31.8	0.3	51.9	161.2	C2'-endo
11 7	-18.0	43.1	-48.9	41.5	-15.1	48.9	178.3	C2'-endo
12 A	-15.5	35.7	-41.9	34.2	-11.8	41.9	177.7	C2'-endo
13 1	-51.6	50.3	-29.7	0.8	31.5	52.3	124.6	C1'-exo
14 A	-7.3	30.1	-40.2	36.9	-19.3	40.6	188.6	C3'-exo
15 0	-23.0	41.8	-42.1	30.7	-5.1	43.2	167.4	C2'-endo
16 A	-36.4	38.4	-26.2	6.3	18.6	38.6	132.8	C1'-exo
17 A	-55.8	34.1	-0.4	-32.8	55.7	58.0	90.3	04'-endo

18 C	-6.0	-19.6	36.1	-40.6	29.0	40.4	26.8	C3'-endo	
19 T	-41.2	10.7	21.0	-45.1	52.9	53.0	66.6	C4′-exo	
Strand II									
base	v0	v1	v2	v3	v4	tm	P	Puckering	
1 C	-21.4	29.0	-26.2	14.7	4.0	29.2	153.8	C2'-endo	
2 C	-35.3	47.2	-41.3	22.0	8.1	46.9	151.6	C2'-endo	
3 C	8.7	-24.7	30.5	-26.9	11.4	30.5	3.0	C3'-endo	
4 C	-18.5	-9.5	31.4	-43.1	39.0	43.1	43.3	C4'-exo	
5 C	-1.5	-21.6	36.9	-38.7	24.3	39.4	20.7	C3'-endo	
6 C	4.2	-27.5	39.4	-38.2	21.3	40.4	12.9	C3'-endo	
7 C	-44.0	21.3	6.9	-32.2	47.4	47.6	81.7	04'-endo	
8 A	-18.2	-4.7	24.3	-35.3	33.2	36.0	47.6	C4'-exo	
9 G	-39.6	49.4	-40.7	19.0	12.6	48.7	146.6	C2'-endo	
10 G	-33.8	28.7	-14.6	-4.5	23.9	32.9	116.3	C1'-exo	
11 A	13.5	4.9	-19.9	28.4	-27.3	28.9	226.4	C4'-endo	
12 T	-39.1	48.1	-38.2	17.2	13.5	46.9	144.6	C2'-endo	
13 A	-27.6	39.7	-37.0	22.0	3.3	40.2	156.9	C2'-endo	
14 T	-10.5	30.5	-37.5	33.0	-14.4	37.5	183.2	C3'-exo	
15 C	-29.8	3.6	22.2	-40.0	44.0	44.2	59.8	C4′-exo	
16 T	-37.5	34.3	-18.7	-2.4	25.0	37.3	120.2	C1'-exo	
17 T	-28.6	1.7	24.4	-41.8	43.4	44.7	57.0	C4′-exo	
18 G	-3.0	15.4	-21.2	20.1	-11.0	21.6	190.9	C3'-exo	
19 A	-44.0	27.3	-1.9	-23.3	41.5	44.3	92.5	04'-endo	
*****************									

Same strand P--P and Cl'--Cl' virtual bond distances

Strand I				Strand II				
k	oase	PP	C1'C1'	k	oase I	PP	C1'C1'	
1	G/G	5.8	5.6	1	C/C	6.6	5.3	
2	G/G	5.7	5.3	2	C/C	6.3	5.6	
3	G/G	5.8	5.4	3	C/C	6.3	5.5	
4	G/G	5.4	5.7	4	C/C	5.4	5.4	
5	G/G	5.9	5.1	5	C/C	6.6	5.6	
6	G/G	5.8	5.7	6	C/C	6.7	5.4	
7	G/T	6.7	5.3	7	C/A	6.7	4.6	
8	T/C	6.9	5.2	8	A/G	6.1	4.8	
9	C/C	6.4	5.4	9	G/G	6.7	4.4	
10	C/T	7.2	4.9	10	G/A	7.0	4.7	
11	T/A	6.2	5.0	11	A/T	6.5	5.9	
12	A/T	7.1	4.3	12	T/A	7.0	4.5	
13	T/A	6.3	5.7	13	A/T	6.7	5.1	
14	A/G	7.1	4.4	14	T/C	6.6	5.6	
15	G/A	6.6	4.9	15	C/T	6.3	4.7	
16	A/A	6.7	4.4	16	T/T	6.6	5.2	
17	A/C	6.5	4.9	17	T/G	7.1	4.5	
18	C/T	6.7	4.9	18	G/A		5.0	

Helix radius (radial displacement of P, O4', and C1' atoms in local helix frame of each dimer)

Strand I				Strand II			
step	P	04′	C1'	P	04′	C1′	

1	GG/CC	8.9	8.1	7.2	9.1	7.2	6.5		
2	GG/CC	8.0	8.3	7.8	11.1	10.1	9.4		
3	GG/CC	9.9	9.6	9.1	10.4	9.6	9.1		
4	GG/CC	10.7	10.4	10.0	9.5	9.4	9.0		
5	GG/CC	9.4	9.1	8.6	9.5	9.2	8.6		
6	GG/CC	10.9	10.9	10.4	11.4	9.8	9.4		
7	GT/AC	9.4	7.4	6.5	10.0	7.3	6.5		
8	TC/GA	9.3	6.0	5.3	8.7	6.1	5.3		
9	CC/GG	10.3	7.3	6.9	7.8	4.3	3.8		
10	CT/AG	10.8	7.7	7.3	9.4	6.9	6.1		
11	TA/TA	7.2	4.3	4.1	9.0	6.6	6.1		
12	AT/AT	10.3	6.9	6.4	8.1	5.0	4.6		
13	TA/TA	7.4	5.0	4.6	9.0	6.0	5.8		
14	AG/CT	9.5	6.8	6.0	9.0	7.6	6.6		
15	GA/TC	10.0	6.8	6.2	9.0	5.7	5.1		
16	AA/TT	9.0	6.0	5.1	9.8	7.0	6.2		
17	AC/GT	11.0	7.5	6.8	10.1	7.3	6.4		
18	CT/AG	10.5	7.9	7.2	10.0	7.2	6.6		
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Position (Px, Py, Pz) and local helical axis vector (Hx, Hy, Hz) for each dinucleotide step

	bp	Px	Py	Pz	Hx	Ну	Hz
1	GG/CC	1.94	155.38	31.00	0.62	-0.78	-0.03
2	GG/CC	2.47	151.13	32.67	0.43	-0.85	-0.30
3	GG/CC	3.81	149.58	33.38	0.45	-0.88	-0.11
4	GG/CC	4.52	147.45	33.83	0.35	-0.92	-0.17
5	GG/CC	6.63	145.33	32.97	0.33	-0.94	-0.05
6	GG/CC	5.82	142.28	32.01	0.41	-0.89	-0.20
7	GT/AC	9.07	139.52	36.07	0.30	-0.95	0.03
8	TC/GA	9.17	136.12	38.33	0.20	-0.97	-0.11
9	CC/GG	8.08	132.84	37.25	-0.09	-1.00	0.03
10	CT/AG	11.95	129.88	38.43	0.34	-0.91	0.25
11	TA/TA	7.96	125.84	38.27	0.16	-0.98	-0.14
12	AT/AT	11.32	123.00	39.49	0.27	-0.96	0.02
13	TA/TA	11.22	119.89	36.48	0.31	-0.95	-0.09
14	AG/CT	7.77	116.25	37.52	-0.12	-0.78	-0.61
15	GA/TC	9.57	112.91	37.03	0.08	-0.99	-0.12
16	AA/TT	10.14	110.11	35.51	0.11	-0.99	-0.06
17	AC/GT	8.61	107.25	33.63	0.01	-0.93	-0.36
18	CT/AG	8.50	103.79	31.77	0.09	-0.96	-0.27

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