Adaptation of the base-paired double-helix molecular architecture to extreme pressure

Eric Girard1,*, Thierry Prange2, Anne-Claire Dhaussy3, Evelyne Migianu-Griffoni4, Marc Lecouvey4, Jean-Claude Chervin5, Mohamed Mezouar6, Richard Kahn7 and Roger Fourme1

1Synchrotron-SOLEIL, L’Orme des Merisiers, Saint-Aubin, BP 48, 91192 Gif-sur-Yvette Cedex, 2Laboratoire de Cristallographie et RMN Biologiques (UMR 8015 CNRS) Université Paris-5, 4 Avenue de l’Observatoire, 75006 Paris, 3CRISMAT, ENSICAEN, Boulevard du Maréchal Juin, 14000 Caen, 4BioMoCeTi (UMR CNRS 7033), UFR S.M.B.H., Université Paris-13, 74 rue Marcel Cachin, 93017 Bobigny Cedex, 5PMD, IMPMC, Université Pierre et Marie Curie – CNRS UMR7590, 140 rue de Lourmel, 75015 Paris, 6ESRF, BP 220, 38027 Grenoble Cedex and 7IBS, UMR 5075 CEA-CNRS-UJF, 41 rue Jules Horowitz, 38027 Grenoble Cedex, France

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ABSTRACT

The behaviour of the d(GGTATACC) oligonucleotide has been investigated by X-ray crystallography at 295 K in the range from ambient pressure to 2 GPa (~20 000 atm). Four 3D-structures of the A-DNA form (at ambient pressure, 0.55, 1.09 and 1.39 GPa) were refined at 1.60 or 1.65 Å resolution. In addition to the diffraction pattern of the A-form, the broad meridional streaks previously explained by occluded B-DNA octamers within the channels of the crystalline A-form matrix were observed up to at least 2 GPa. This work highlights an important property of nucleic acids, their capability to withstand very high pressures, while keeping in such conditions a nearly invariant geometry of base pairs that store and carry genetic information. The double-helix base-paired architecture behaves as a molecular spring, which makes it especially adapted to very harsh conditions. These features may have contributed to the emergence of a RNA World at prebiotic stage.

INTRODUCTION

Nucleic acids are more plausible than proteins as the components of a self-contained replicating system in the very first stages of the emergence of life on Earth (1–3). Watson–Crick base pairing provided a very plausible mechanism by which a polynucleotide could direct the synthesis of its complement from mononucleotides or short oligonucleotides, while no equivalent mechanism is known for the replication of a polypeptide. The discovery of ribozymes (4,5) and then the demonstration that ribosomal peptide synthesis is a ribozyme-catalysed reaction (6) strengthened the case for an early RNA World. In this context, the central problem for origin-of-life studies is to understand how this seminal world became established on the primitive Earth. Plausible scenarios for the prebiotic chemistry have been proposed (7,8), but the problem is far from being solved. A recurrent theme is that RNA may have emerged from an earlier world under extreme conditions of pressure and/or temperature and pH (8). In all scenarios, molecules with backbones forming stable double helices held together by Watson–Crick base pairing appear as crucial intermediates or crucial building blocks. This fact led us to undertake a programme on the behaviour of such molecules under high pressure.

Some studies were recently devoted to the analysis of the effects of high hydrostatic pressures on the stability of living cells (9–11), the quaternary structures of proteins and viruses (12), protein–protein and protein–DNA interactions (13), catalytic RNA (14) and tRNA (15). Concerning nucleic acids, they were investigated through their single-strand/duplex equilibrium (16), their B/Z transition (17) and their hydration networks (18) all using spectroscopic techniques in solution. Following Le Chatelier’s principle, pressure tends to favour states with smaller specific volumes. In DNA, structural adaptation to high pressure results from several contributions, including the helical structure integrity (helix-to-coil transition), the stacking of bases, the Watson–Crick association and the hydration network around the duplex. Aromatic-ring stacking is favoured by pressure while Coulombic or hydrophobic interactions are

*To whom correspondence should be addressed. Tel: +33 169359609; Fax: +33 169359456; Email: eric.girard@synchrotron-soleil.fr

The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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disfavoured (19). Hydrogen bonds and associated networks should be, as in protein and protein assemblies (20–22), stabilized by pressure. How these various parameters influence the structure and stability of the DNA duplex is not easy to predict. Clearly, besides spectroscopic data, accurate 3D structures were required in order to describe the behaviour under high pressure at the molecular level and provide a firm starting point for simulations. With respect to pioneering work based on high-pressure beryllium cell (23), our technical developments combining diamond anvil cell and synchrotron radiation of ultrashort wavelength (24) have considerably extended the possibilities of high-pressure macromolecular crystallography (HPMX) in a pressure range increased by one order of magnitude (from 0.2 to 2 GPa). After studies on monomeric (20), then multimeric (25) proteins and a complex protein assembly (21), this article reports the first application of HPMX to nucleic acids.

The d(GGTATACC) oligonucleotide was selected owing to the high stability of its crystalline A-DNA crystalline form, its strong base stacking (2.9 Å compared to the 3.3 Å in B-DNA duplex), and its particularly well-defined hydrogen-bonding network, mostly located in the major groove (26). The last reason for this choice is an interesting feature related to the crystal packing of d(GGTATACC) crystals. Besides Bragg reflections of the A-form crystal, a pattern was observed on diffraction pictures of nucleotide crystals, similar to fibre diagrams that led Watson and Crick to their interpretation of the DNA structure (27). Main features of this pattern include a characteristic diffuse crossed X-ray pattern, a set of streaks perpendicular to the c⁺ direction of the reciprocal lattice and two strong elongated meridian reflections (28). The following interpretation for the origin of this pattern was proposed (28). Oligonucleotide molecules pack in infinite super-helices of duplexes down the 6-fold axis of the P6₁ space group (26). The central channel of the super-helix can trap oriented molecules of DNA. Simulations of streaks allow excluding the A-form for occluded molecules and favour the B-form. The meridional reflections are consistent with the base-pair stacking in B-form molecules. According to these results, we have in hand a system in which both A and B forms of DNA can be simultaneously monitored against external hydrostatic pressure, although information derived on the B-form is obviously quite limited.

Crystals of the nucleotide were compressed up to 2 GPa at 295 K. We describe the high-resolution crystal structures of A-DNA at four pressures from ambient to 1.39 GPa. Meridional streaks attributed to B-DNA are observed up to at least 2 GPa. These results highlight the remarkable adaptation of the base-paired double-helix architecture base-paired architecture to high pressure.

**MATERIALS AND METHODS**

**Preparation and crystallization**

The d(GGTATACC) sequence was synthesized by standard phosphoramidite chemistry on solid support (Applied Biosystems 391 DNA Synthetizer). Crystals were obtained following the batch method (26). Ten milligrams of lyophilized octamer were dissolved in 200 μl of a 15% methyl-pentane-diol (MPD) solution, buffered by sodium cacodylate (5 × 10⁻² M, pH 7) and containing additives: sodium azide, spermine tetrahydrochloride and MgCl₂. To this solution, was added each day 5 μl of the same solution but containing 50% of MPD, followed by a rapid mixing. After 5 days, crystals began to appear as elongated hexagonal rods and grew easily to a size of 0.2 × 0.2 × 1.0 mm³ in a week. They were stabilized by increasing the final MPD concentration to ~35%.

**High-pressure cell**

Crystals were hydrostatically compressed in a diamond anvil cell. The compression chamber (diameter 400 μm, height 200 μm) was drilled in a copper or stainless steel (for pressure above 0.7 GPa) gasket (Figure 1). Gasket preparation and sample loading were performed as described (24). The crystallization solution with 35% MPD was used as compression medium. Pressure monitoring was performed by using the wavelength shift of laser-excited fluorescence from a small ruby sphere loaded in the compression chamber. Two different diamond anvil cells were used for data collection. The first one had a useful aperture of 62° and a standard diamond mount (24). The second one was of a novel design to provide both a larger useful aperture (82°) and a pressure range up to ~2.5 GPa (29).

**Data collection**

Diffraction data from crystals compressed in the diamond anvil cell were collected at the ESRF (Grenoble, France), on the ID27 beamline with a MAR CCD 165 mm detector. The wavelength was calibrated and set to 0.3738 Å (Iodine K absorption edge). Four data sets were collected at room temperature and ambient pressure, 0.55, 1.04 and 1.39 GPa. The crystal-to-detector distance,
calibrated using the diffraction rings of a reference silicon powder, was 352.9 mm for the 0.55 GPa data set and 302.9 mm for the others. The X-ray beam was collimated to 50 × 50 μm². Exposure times were 30 or 45 s, depending on the sample, for a rotation step of 1°. As data collection was performed at room temperature, crystals were translated several times, every 10 to 35° of rotation, during exposure in order to irradiate fresh zones (24).

Data processing
Diffraction frames were integrated using XDS (30). All data were independently put on an absolute scale using SCALA (31).

Refinement
The starting molecular model was either the deposited coordinates ref. 115D (26) or 1VJ4 (32) from the Protein Data Bank. In the case of 115D, the isomorphous di-bromo derivative of 1VJ4, thymine residues were reconstructed from original bromo-uracils and all water molecules were removed. A first round of rigid body refinements was done with AMoRe (33), then refinements proceeded at the maximum resolution with individual isotropic thermal factors and bond distance and angle restraints, as used in SHELXL (34). Water molecules were localized during the course of refinement cycles by analysing density peaks in F₀−Fc Fourier difference maps. They were accepted when they met standard criterions like correct bond distances and angles toward polar atoms of the model, and B thermal factors below a given threshold arbitrarily fixed to 65 Å². All analyses of the geometric parameters were done using NEWHEL93 (35). All molecular figures were created using the program PyMol (http://pymol.sourceforge.net/).

RESULTS AND DISCUSSION
Single crystals of d(GGTATACC) were gradually compressed from atmospheric pressure to 2 GPa (Figure 1). Unit cell parameters of the A-DNA crystal were measured at each step of the pressure ramp from diffraction data recorded over a rotation range of 2°. In spite of the elongated (anisotropic) shape of the DNA cylinder, they decrease isotropically up to a pressure value of ~1.5 GPa. The isothermal compressibility of the crystal is defined as χ = −1/V(∂V/∂P)T, where V is the unit-cell volume. The variation of V is shown in Figure 2. Least-squares fit up to 1.5 GPa is V(P) = V(∞) + k₁(exp(−k₂P) with V(∞) = 62290 Å³, k₁ = 11030 Å³ and k₂ = 1.25 GPa⁻¹. The derived isothermal compressibility is χ(P) = V⁻¹, k₁, k₂, exp(−k₂P). The largest value is at ambient pressure (χ₀ = 0.215 GPa⁻¹). We recall that the compressibility of bulk water is 0.35 GPa⁻¹; the compressibility of water in the vicinity of polar atoms is similar to that of an ‘ice-like’ structure, 0.18 GPa⁻¹ (36). The mean value of χ(P) between ambient pressure and 1.5 GPa is 0.088 GPa⁻¹, similar to the compressibility of tetragonal hen egg-white lysozyme crystal (23,24). Above ~1.6 GPa, the cell volume increases (the compressibility is negative) and the crystal quality, as monitored by mosaicity and resolution of diffraction data, gradually deteriorates. The diffraction that extends to 1.6 Å at ambient pressure falls off to 3 Å at ~1.8 GPa and is completely lost at ~2.0 GPa. Four high-resolution structures of the A-DNA form were determined at ambient pressure, 0.55, 1.04 and 1.39 GPa. In each case, one or two crystals were sufficient to acquire high-completeness data (Table 1) in spite of data collection at room temperature. Refinements were performed at 1.60–1.65 Å resolution to R-factors (R-free factors) of 15.2 (19.1), 16.9 (20.1), 19.3 (22.7) and 18.8 (22.6)% respectively (Table 2).

The overall duplex structures shown in Figure 3 evidence axial compression of the helix, which reacts to pressure like a molecular spring. The base-stacking shrinkage is 2.6 Å for the full octamer length (from 23.5 Å at ambient pressure to 20.9 Å at 1.39 GPa, i.e. a relative contraction of 11%). The average base-pair step varies from 2.92 Å down to 2.73 Å (Figure 4). This spectacular plasticity associates only small changes in phosphodiester backbone angles for accounting the denser base stacking. The sugar puckering parameters (Taum and P) describing the sugar conformations were analysed. All of them belong to the canonical C3’endo (or northern) conformation, a behaviour already mentioned earlier (26). The pseudo-rotation P remains remarkably constant over the whole pressure range between standard values of 1 to 30°, including the O3’ end sugar of chain A, which moves from 30.9 to 33.3°. The only exception is the O3’ end sugar of chain B that adopts at ambient pressure a C2’/C1’ twist (nearly a southern conformation) and goes back to 36.8° (northern) at 1.04 GPa and above.

The variation of base-pair spacing versus pressure (Figure 4) is approximately linear up to 1.0 GPa and becomes steady beyond, which means that the molecule becomes progressively unable to accommodate increasing compression. The gradual degradation of the crystal order beyond ~1.6 GPa may be due, at least partly, to this effect.

Figure 2. A-DNA pressure variation at 295 K of crystal unit-cell volume V versus pressure (rmsd’s on cell dimensions and pressures are estimated as 2 × 10⁻³ Å and 10⁻² GPa, respectively).
Contrary to common B-DNA crystal packing arrangements, where duplexes stack on the top of each other building infinite helices along crystal axes, the hexagonal packing of d(GGTATACC) builds infinite super-helices around the 6-fold crystallographic axis with a close-contact zone between neighbouring helices (26,28). In this region, the wedge effect of hydrostatic pressure produces a hinge point in base stacking between the fifth and the sixth base pairs (Figure 5). This steric effect might also contribute to crystal degradation.

The Watson–Crick type of base-pair association, which represents the foundation of the genetic code transmission, could be another way for the helix to ‘breathe’ transversally. In proteins, in the elastic compression regime, salt bridges and H-bond lengths are usually shortened by \(\Delta \text{C} \approx 0.1 \text{Å.GPa}^{-1}\) (20–22). In the case of A-DNA, evolutions of polar atom distances within G–C and A–T base pairs (Table 3) can be interpreted in terms of small variations of rise, buckle, propeller-twist and other parameters that contribute to the adaptation to high pressure (Table 4). Accordingly, the geometry of Watson–Crick base pairings remains essentially invariant in the pressure domain up to 1.39 GPa. The lengths of vectors \(\text{C} \approx 0.1 \text{Å.GPa}^{-1}\) that may be used to quantify the DNA cylinder transversal squeeze, are all identical within their SDs, i.e. 10.5(2) Å, whatever the applied pressure.

It was suggested from modelling and docking simulations that water hydration under high hydrostatic pressure would tend to build octahedral arrangements (17) around DNA, to account for a negative \(\Delta V\). These arrangements have been invoked in the B/Z transition observed under high pressure (15). The A-form of DNA shows a heavily hydrated major groove and a poorly hydrated minor groove. These features are distinct with respect to other DNA forms, and as such are worth to be investigated under pressure. The hydration scheme of d(GGTATACC)

| Pressure (GPa) | Ambient | 0.55 | 1.04 | 1.39 |
|---------------|---------|------|------|------|
| Cell parameters (Å), \(a = b = 45.034\) \(c = 41.747\) | \(a = b = 43.710\) \(c = 40.710\) | \(a = b = 43.171\) \(c = 40.383\) | \(a = b = 42.830\) \(c = 40.300\) |
| Resolution range (Å) | 18–1.60 | 18–1.65 | 18–1.60 | 18–1.60 |
| Number of crystals used | 2 | 1 | 1 | 1 |
| Number of unique reflections | 6 015 | 5 038 | 5 079 | 5 058 |
| \(R_{\text{merge}}\) (%) | 4.7 (33.0) | 4.3 (18.8) | 5.8 (25.1) | 4.7 (33.1) |
| Completeness (%) | 94.5 (96.2) | 94.1 (94.1) | 89.4 (91.0) | 98.5 (99.3) |
| Redundancy | 4.0 (4.0) | 3.2 (2.6) | 3.2 (3.3) | 3.1 (3.1) |

Table 1. Statistics of data collection at 295 K for the four d(GGTATACC) crystal structures. The space group is P6\(1\). X-ray wavelength was 0.3738 Å.

Table 2. Refinement statistics

| Pressure (GPa) | Ambient | 0.55 | 1.04 | 1.39 |
|---------------|---------|------|------|------|
| Number of atoms in the model | 322 | 322 | 322 | 322 |
| Number of water molecules | 57 | 75 | 75 | 76 |
| Number of additional atoms ( spermine) | – | – | 10 | 10 |
| R factor (on observed F)\%, | 14.86 | 16.93 | 18.92 | 18.66 |
| R factor (all F data)\%, | 15.85 | 18.64 | 20.15 | 20.07 |
| R-free factor (8% of F data)\%, | 18.9 | 20.1 | 22.1 | 22.2 |
| Number of parameters/No. restraints | 1523/1585 | 1595/1594 | 1651/1653 | 1691/1729 |
| Solvent fraction in the unit cell (%) | 49.2 | 44.7 | 43.6 | 41.8 |
| Average thermal parameters (Å\(^2\) (Number of atoms in parentheses) | 16.5 | 18.4 | 16.4 | 22.0 |
| Base planes (152) | 19.0 | 22.6 | 20.8 | 27.4 |
| Sugars (96) | 24.3 | 30.5 | 27.6 | 32.3 |
| Phosphate chains (74) | 44.9 | 47.0 | 40.8 | 39.6 |
| All water molecules | 40.6 | 37.1 | 34.6 | 34.1 |
| Common water molecules (46)* | 45.2 | 46.3 | 40.2 | 38.3 |
| 1st shell water molecules | 43.1 | 47.5 | 41.7 | 40.7 |
| 2nd shell water molecules | 0.12 (392) | 0.010 (392) | 0.009 (405) | 0.009 (406) |
| Bond distances (1–2 dist) (Å) | 0.039 (656) | 0.028 (656) | 0.027 (668) | 0.030 (684) |
| Bond angles (1–3 dist) (°) | 0.014 (120) | 0.016 (120) | 0.015 (120) | 0.021 (120) |
| Planarity (Å) | 0.007 (48) | 0.007 (48) | 0.007 (48) | 0.008 (52) |
| Non-zero chiral volumes (Å\(^3\) | –0.23/+0.30 | –0.23/+0.33 | –0.29/+0.39 | –0.26/+0.30 |

Table 2. Refinement statistics

*Water molecules observed in common sites for all structures.
has been extensively analysed because it presents regular pentagonal arrangements (26) in the major groove (Figure 6), a typical motif also observed in some well-ordered proteins at atomic resolution such as crambin (37). The particular conformation of the A-form of DNA allows water molecules to directly bridge phosphate groups along each individual strand. At ambient pressure, 75 direct polar contacts are established through 51 water molecules belonging to the first shell of hydration, including nine over the 12 possible phosphate bridges. As mentioned earlier, the remaining water molecules build an ordered network from rim to rim thus completely filling most of the available space in the major groove. Under pressure, more water sites become apparent. This is illustrated in Figure 7, which shows the same G–C base pair at one end of the duplex superimposed to the $2F_o - F_c$ electron-density maps calculated at different pressures. At 0.55 GPa, the number of observed isolated peaks in the first shell of hydration increases to 69, although some of them were suspected not to correspond to water sites as they fill parts of elongated electron densities that cannot be connected to the model. When pressure is increased above 1.04 GPa, one of the elongated densities resolves as a linear chain, ideally fitting a folded spermine molecule. The stabilization of the hydrogen-bond network is made evident by a slow but continuous decrease of the average normalized ($B_1$) thermal factor of water molecules common to all structures (Table 2) down to a limit of ~34 Å$^2$. The number of direct polar contacts from the first shell of hydration to the oligomer also increases rapidly with pressure up to 0.55 GPa, then more slowly up to 1.39 GPa. Beyond 1 GPa, the first shell of hydration is gradually compressed thus leading to more direct contacts towards the DNA (Table 2). Nevertheless, the pentagonal network located in the major groove remains conserved in the whole pressure range.

The four X-ray data sets were recorded with exposure times appropriate for collecting the A-form diffraction data but too short to get a complete picture of the diffuse scattering pattern described in (28). The series of narrow streaks distributed in the pattern of a cross was barely observable, but the extended meridional streaks associated with the base-pair stacking were clearly observed up to at least 2 GPa. According to the interpretation given in (28), the diffuse scattering pattern is produced by occluded B-DNA molecules in the packing channels of the A-DNA crystal structure. A suitable orientation of crystals in the high-pressure cell allowed us to record these meridional streaks (Figure 8) while ramping pressure, which provided data to determine the average stacking distance. A continuous and smooth shortening is observed, that can be monitored even in the range 1.5–2 GPa where the A-DNA crystal order falls off (Figure 4). The average base-pair step is 3.34 Å at ambient pressure and 3.07 Å at 2 GPa.

**CONCLUSION**

We have shown that d(GGTATACC) in the crystalline A-form can withstand very high pressures, up to ~1.9 GPa. The gradual loss of long-range order between 1.6 and 1.9 GPa may be related partially or completely to packing effects as mentioned previously and the molecule in solution might be stable even beyond 1.9 GPa. The four high-resolution structures show that geometry of base pairs is well preserved under compression up to at least 1.39 GPa. The B-form molecules occluded in channels of the crystal packing are in a solution-like environment. The information derived on this form from diffuse scattering is limited. Nevertheless, the smooth variation of the period of stacking derived from the evolution of meridional streaks reveals that the B-form is probably stable up to at least 2 GPa. We shall consolidate and extend this preliminary result by performing another single-crystal study on a dodecanucleotide that crystallizes in the B-DNA form.
The remarkable adaptation of d(GGTATACC) to high pressure is clearly associated to the base-paired double-helix topology of the molecule, by which it behaves as a molecular spring. These properties are probably shared by molecules featuring similar topology, with sugar-phosphate or polypeptide backbones. At the prebiotic stage, the base-paired double-helix architecture was crucial in the emergence of molecules with catalytic properties and able to store genetic information. Such architectures could withstand not only pressure in the deepest sea trenches but also much higher pressures found in Earth’s interior or in the context of rare events such as impact of a meteorite. We suggest that this remarkable adaptation to harsh conditions may have played an important role during the sequence of events that led to the seminal RNA World.

Table 3. Main helical parameters in d(GGTATACC) (errors on distances estimated as ±0.035 Å, based on refined coordinates). Temperature is 295 K.

| Pressure (GPa) | Ambient | 0.55 | 1.04 | 1.39 |
|---------------|---------|------|------|------|
| Watson–Crick distances (Å) | | | | |
| G–C base pairs | | | | |
| O6–N4 | 2.90 | 2.95 | 2.96 | 2.90 |
| N1–N3 | 2.92 | 2.88 | 2.89 | 2.83 |
| N2–O2 | 2.84 | 2.80 | 2.81 | 2.76 |
| A–T base pairs | | | | |
| N1–N3 | 2.86 | 2.82 | 2.81 | 2.74 |
| N6–O4 | 2.97 | 2.90 | 2.99 | 2.98 |
| Total number of water molecules located | 57 | 75 | 75 | 76 |
| Number of water molecules in the 1st sphere of hydration (d<3.4 Å) | 51 | 69 | 65 | 71 |
| Number of polar contacts to the 1st sphere of hydration | 75 | 94 | 103 | 115 |
| Distances ( Å) in range | | | | |
| 2.00–2.40 | 2 | 7 | 5 | 7 |
| 2.40–2.80 | 20 | 27 | 29 | 32 |
| 2.80–3.20 | 37 | 44 | 40 | 46 |
| 3.20–3.40 | 16 | 16 | 29 | 30 |
| Common water molecules | | | | |
| Average bond distance (Å) | 2.95 | 2.91 | 2.95 | 2.94 |
| Average isotropic thermal factor B (Å²) | 40.6 | 37.1 | 34.6 | 34.1 |

The remarkable adaptation of d(GGTATACC) to high pressure is clearly associated to the base-paired double-helix topology of the molecule, by which it behaves as a molecular spring. These properties are probably shared by molecules featuring similar topology, with sugar-phosphate or polypeptide backbones. At the prebiotic stage, the base-paired double-helix architecture was crucial in the emergence of molecules with catalytic properties and able to store genetic information. Such architectures could withstand not only pressure in the deepest sea trenches but also much higher pressures found in Earth’s interior or in the context of rare events such as impact of a meteorite. We suggest that this remarkable adaptation to harsh conditions may have played an important role during the sequence of events that led to the seminal RNA World.

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For all pressures, base-plane atoms (152), sugar atoms (96) and phosphate atoms (74) have their (B) parameters in the ranges, [16–20 Å²], [18–26 Å²] and [22–32 Å²], respectively.
Table 4. Base plane geometry versus external hydrostatic pressure, according to NEWHEL93 (35)

| Pressure (GPa) | Step | Rise (Å) | Twist (°) | Roll (°) | Buckle (°) | Propeller (°) | Slide (Å) | Tilt (°) | Inclination (°) |
|---------------|------|----------|-----------|---------|-----------|--------------|----------|---------|----------------|
| Ambient       | 1    | 2.786    | 33.57     | 6.42    | 7.47      | −4.89        | −3.44    | −0.10   | 8.63           |
|               | 2    | 2.817    | 32.09     | −0.08   | 14.03     | −12.11       | −2.80    | 1.63    | 11.82          |
|               | 3    | 3.096    | 29.93     | 12.19   | 6.71      | −9.20        | −2.55    | 0.29    | 14.96          |
|               | 4    | 3.078    | 33.22     | 2.10    | 2.02      | −11.61       | −2.61    | 0.06    | 15.64          |
|               | 5    | 2.994    | 31.38     | 14.86   | −7.64     | −12.51       | −2.85    | −0.92   | 15.32          |
|               | 6    | 2.748    | 31.91     | 3.84    | −7.64     | −14.04       | −3.01    | −0.67   | 13.81          |
|               | 7    | 2.920    | 31.81     | 8.05    | −8.46     | −12.21       | −3.24    | −0.13   | 13.86          |
| Mean          | 8    | 2.92     | 31.99     | 6.77    | −8.78     | −0.49        | −        | −       | 13.36          |
| 0.55          | 1    | 2.754    | 31.95     | 8.07    | 3.67      | −6.12        | −1.81    | −0.11   | 8.57           |
|               | 2    | 2.702    | 36.58     | 0.18    | 11.07     | −15.21       | −1.04    | 1.69    | 10.91          |
|               | 3    | 2.821    | 30.00     | 12.39   | 3.81      | −12.24       | −1.26    | 0.54    | 14.27          |
|               | 4    | 2.913    | 33.98     | −0.04   | 4.04      | −12.01       | −0.96    | −0.06   | 15.20          |
|               | 5    | 3.091    | 30.44     | 18.45   | −8.75     | −15.14       | −1.34    | −0.85   | 14.18          |
|               | 6    | 2.538    | 34.76     | 1.28    | −7.44     | −17.45       | −1.17    | −0.30   | 12.87          |
|               | 7    | 2.867    | 31.76     | 9.39    | −10.07    | −17.50       | −1.65    | −1.46   | 13.96          |
| Mean          | 8    | 2.83     | 32.78     | 7.10    | −4.39     | −12.35       | −1.32    | −0.08   | 12.77          |
| 1.04          | 1    | 2.797    | 31.73     | 7.03    | 6.40      | −6.60        | −1.83    | 0.47    | 7.90           |
|               | 2    | 2.696    | 37.34     | 1.32    | 8.42      | −14.66       | −0.98    | 0.53    | 10.99          |
|               | 3    | 2.712    | 29.51     | 12.25   | 2.30      | −10.26       | −1.28    | 1.04    | 13.29          |
|               | 4    | 2.766    | 35.21     | −0.89   | 4.54      | −13.82       | −1.10    | −0.22   | 15.29          |
|               | 5    | 3.246    | 29.58     | 19.47   | −6.95     | −15.03       | −1.40    | −4.04   | 14.36          |
|               | 6    | 2.352    | 35.94     | −0.79   | −11.82    | −19.16       | −1.15    | 0.28    | 10.35          |
| Mean          | 7    | 2.865    | 31.52     | 8.47    | −9.38     | −19.20       | −1.73    | −1.80   | 12.75          |
| 1.39          | 8    | 2.81     | 32.78     | −4.69   | −12.35    | −1.32        | −0.08    | 12.77   |                |
| Mean          | 1.39  | 2.78     | 32.98     | 6.70    | −1.78     | −12.57       | −1.35    | −0.53   | 11.98          |

Some parameters describing the geometry, in particular RISE and TWIST parameters, are very sensitive to packing hinge effect of one helix over the preceding one in the super-helix built along the c axis as shown in the diagrams in Figure 5.

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Accession codes. Final coordinates and structure factor amplitudes for the structures at ambient pressure, 0.55,
over the degrading A-DNA diffraction pattern.

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