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Conductivity of natural and modified DNA measured by scanning tunneling microscopy. The effect of sequence, charge and stacking

Irena Kratochvílová ^{a,*}, Karel Král ^a, Martin Bunček ^b, Alena Víšková ^b, Stanislav Nešpůrek ^c, Anna Kochalska ^c, Tatiana Todorciuc ^c, Martin Weiter ^d, Bohdan Schneider ^e

- ^a Institute of Physics, ASCR, v.v.i., Na Slovance 2, CZ-182 21 Prague, Czech Republic
- ^b GENERI BIOTECH s.r.o., Machkova 587, CZ-500 11 Hradec Kralové, Czech Republic
- ^c Institute of Macromolecular Chemistry ASCR, v.v.i., Heyrovského nám. 2, CZ-162 06 Prague 6, Czech Republic
- ^d Faculty of Chemistry, Brno University of Technology, Purkyňova 118, CZ-612 00 Brno, Czech Republic
- e Institute of Biotechnology AS CR, v.v.i., Vídeňská 1083, CZ-142 20 Prague, Czech Republic

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ABSTRACT

The conductivity of DNA covalently bonded to a gold surface was studied by means of the STM technique. Various single- and double-stranded 32-nucleotide-long DNA sequences were measured under ambient conditions so as to provide a better understanding of the complex process of charge-carrier transport in natural as well as chemically modified DNA molecules. The investigations focused on the role of several features of DNA structure, namely the role of the negative charge at the backbone phosphate group and the related complex effects of counterions, and of the stacking interactions between the bases in Watson-Crick and other types of base pairs. The measurements have indicated that the best conductor is DNA in its biologically most relevant double-stranded form with Watson-Crick base pairs and charged phosphates equilibrated with counterions and water. All the studied modifications, including DNA with non-Watson-Crick base pairs, the abasic form, and especially the form with phosphate charges eliminated by chemical modifications, lower the conductivity of natural DNA.

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1. Introduction

The DNA molecule, containing the genetic code of all living species, has recently become a center of great attention on the part of chemists and physicists, one of the reasons being DNA's potential use in nanoelectronic devices [1–4], both as a template for assembling nanocircuits and as an active element of such circuits. Charge migration along DNA molecules has attracted scientific interest for more than fifty years. The reports on the possible high rates of charge transfer between the donors and acceptors through DNA, obtained from solution chemistry experiments [5], have triggered a series of direct electrical transport measurements on bundles and networks, because a truly conducting form of DNA would have a major impact on the developments in nanotechnologies.

Extended electronic states of DNA could play an important role in biology, e.g. through the processes of DNA-damage sensing or repairing via long-range charge transfer [6,7]. The prevailing DNA architecture, the double helix, has well stacked, nearly parallel bases with overlapping π -electron systems. Such π -electron systems may be

E-mail addresses: krat@fzu.cz (I. Kratochvílová), kral@fzu.cz (K. Král), martin.buncek@generi-biotech.com (M. Bunček), nespurek@imc.cas.cz (S. Nešpůrek), weiter@fch.vutbr.cz (M. Weiter), bohdan@img.cas.cz (B. Schneider).

good candidates for long-distance and one-dimensional (linear) charge transport [8]. Several authors have indicated that DNA conducts electric charge via the hopping mechanism [9–11], but the electronic properties of DNA remain controversial. Charge-transfer reactions and conductivity measurements exhibit a large variety of possible electronic behavior, ranging from Anderson and band-gap insulators to effective molecular wires and induced superconductors. Indeed, understanding the conductance of a complicated polyelectrolyte aperiodic system is in itself a major scientific problem [12–16].

Historically, two basic experimental approaches have been pursued to investigate conduction through molecules: (i) contacting isolated single molecules or molecules in thin films, and (ii) studying transport in thick films and devices, such as organic thin-film transistors or light-emitting diodes [17,18]. The optimal experimental setup is to position isolated molecules between two electrical contacts, but this is very difficult to implement and especially difficult to verify. Working with the ordered arrays of parallel π -conjugated molecules, where in principle the binding to the substrate can be precisely controlled, offers a possibility to measure the individual molecules by making use of the scanning–tunneling–microscope (STM) tip. STM experiments have already been proven as a very suitable tool for investigating ordered monolayer films, making it possible to contact one or more molecules and obtain images of the structural characteristics of the film [19,20]. The experiments have

^{*} Corresponding author.

also been used to investigate room-temperature electronic properties of twelve base-pair $d(GC)_{12}$ – $d(GC)_{12}$ DNA molecules attached to the gold surface by a thiol link [21]. The STM/STS experiments offer a novel way of probing the electronic properties of biomolecules on surfaces on the atomic level.

In this work, we have used the STM technique in order to study the conductivity of several oligonucleotide sequences in single- and double-stranded forms under ambient conditions. The purpose of these investigations was to understand the complex processes of charge transport through DNA molecules and to dissect the role of the sugar-phosphate backbone, and of its charged phosphate group in particular, as well as of base stacking in the same strand and across strands. To this end, some of the studied oligonucleotide sequences were designed to form non-Watson-Crick base pairs, with the aim of making it possible to estimate the role of these mutation-causing base pairs. The role of the bases and of their stacking interactions in DNA conductivity was also studied at a more fundamental level by removing two or three nitrogenous bases at the central steps of a few sequences forming modified abasic nucleotides. Other sequences contain chemically modified phosphate groups, which eliminate their charge (Fig. 1). Such modifications make it possible to see the roles of the charged backbone and of the alkali-metal and alkaline-earthmetal counterions on conductivity.

2. Experimental

2.1. Material

STM experiments were performed on 32-mer oligonucleotides of various sequences, some of which had been chemically modified. For further reference, all the measured oligonucleotides, including their

Fig. 1. The chemical scheme of the studied nucleotides: (a) a natural nucleotide; b) a part of an abasic oligonucleotide with base removed; c) oligonucleotide with its phosphate neutralized by a *p*-isopropoxy group.

labels, are listed in Table 1. The sequences were designed to investigate the differences: i) between the single-stranded (ss) and doublestranded (ds) DNA of C/G-rich composition, ii) between the singlestranded (ss) and double-stranded (ds) DNA of A/T-rich composition, iii) between dsDNA with only canonical Watson-Crick and containing non-Watson-Crick ("mismatched") base pairs, and iv) between DNA with sequences composed of only natural (non-modified) nucleotides and chemically modified nucleotides. Further, two types of chemical modifications were studied: i) the elimination of nitrogenous bases, thus creating abasic nucleotides, in order to be able to gauge the influence of the stacked bases on the DNA conductivity, and ii) the replacement of the charged phosphate group with its neutral surrogate, which makes it possible to estimate the role of the phosphate charge on the DNA conductivity. The investigated DNA samples (Table 1) were measured in pairs selected in such a way that only one parameter would vary in a pair (see Fig. 1 for the chemical structures of the nucleotides). For instance, the phosphate charge present in the standard oligonucleotide (Fig. 1, label A) is eliminated in DNA E of the identical sequence, the sequences differ between A and B, or between C and its abasic mutant D.

The sequences were selected so that they would not form such special structural features as loops or complex tertiary structures. Guanosine nucleotides were evenly distributed as every fourth base in the mixed sequences. The pure G/C (Table 1, ss: A and ds: 1) and A/T (Table 1, B and 2) oligonucleotides were designed for the evaluation of the effect which the guanine base might have on the overall DNA conductivity. The C/G, T/A and the mixed C/G/T/A sequences were prepared as single-stranded and double-stranded with their respective complementary sequences in order to evaluate the effect of single- versus double-stranded DNA forms. So as to ascertain the influence of the heterocyclic nitrogenous bases on the overall charge transport in DNA, we compared the conductivity of the mixed C/G/T/A sequence (Table 1, C - MIX) containing all four bases in equal proportion with guanine evenly distributed on the one hand with a chain with the same sequences except for three removed bases in the middle (Positions 15–17, Table 1, D – MIX_S) on the other. These abasic sequences produce a gap with three base pairs missing in the respective ds sample (Table 1, ds 3 and 4). Oligo d(T) where two bases in the middle were substituted by either two guanines or by abasic spacer, making two base gaps (Fig. 1b), were also synthesized. To study the impact of charge in the DNA backbone on the charge transport, all the above sequences were also synthesized with a chemically modified phosphate group, which eliminates the phosphate charge. Thus the sequences were prepared as standard phosphodiesters (Fig. 1a), rendering a negative charge on each phosphate, and also as neutral but still polar p-isopropoxy derivatives (Fig. 1c, Table 1, E and NGC). How non-canonical (non-W-C) base pairs affect the conductivity was measured on DNA chains modified by introducing sequences with two non-W-C ("mismatched") pairs (Table 1, ds 6 and 7).

All the sequences were analyzed by MALDI-TOF mass spectroscopy in order to prove their identity and evaluate their purity. The sequences with neutralized phosphates charges (*p*-isopropoxy modified) showed a small or undetectable number of Na⁺/K⁺ adduct peaks, and having compared them to MALDI-TOF measured concentration of Na⁺/K⁺ in charged nucleotides, we concluded that the sequences with uncharged phosphates may contain no or only a very small number of Na⁺/K⁺ counterions associated with the oligonucleotide.

Oligonucleotides were synthesized in GENERI BIOTECH, s.r.o., on an ABI394 synthesizer using standard phosphoramidite chemistry. The standard base phosphoramidites, 5'-thiol modifier C6 and dSpacer CE phosphoramidite were purchased from Glen Research, USA, and used according to the manufacturer's recommendations. The *p*-isopropoxy derivatives of base amidites (deoxy adenosine (n-bz) *p*-isopropoxy phosphoramidite, deoxy Cytidine (n-bz) *p*-isopropoxy phosphoramidite, deoxy guanosine (n-ibu) *p*-isopropoxy phosphoramidite and thymidine *p*-isopropoxy phosphoramidite) were purchased from

Table 1The investigated DNA sequences

Code	Name	Sequence	Backbone modification				
A	G/C	Au-s-cgccgccgccgccgccgccgc	CCGCCGCCG-3'	NO			
В	A/T	Au-S-ATAATAATAATAATAATAATAATA	AATAATAAT-3'	NO			
С	MIX	Au-S-GTTAGCACGATAGTCCGATAGTCA	AGTCAGTCC-3'	NO			
D	MIX_dS	Au-S-GTTAGCACGATAGTxxxATAGTCA	AGTCAGTCC-3'	NO			
E	NGC	Au-S-CGCCGCCGCCGCCGCCGCCGC	s-cgccgccgccgccgccgccgccgccgca-3 '				
Code	Sequence		Backbone mod	Backbone modification			
1	Au-S-CGCCGCCC	GCCGCCGCCGCCGCCGCCGCCG-3'	NO				
	1111111						
	3'-GCGGCGGC	CGGCGGCGGCGGCGGCGGC-5'					
2	Au-S-ATAATAA1	CAATAATAATAATAATAATAAT-3'	NO				
	1111111						
	3'-TATTATTA	ATTATTATTATTATTATTATTA-5'					
3	Au-S-GTTAGCAC	CGATAGTCCGATAGTCAGTCC-3'	NO				
	1111111						
	3'-CAATCGTO	GCTATCAGGCTATCAGTCAGTCAGG-5'					
4	Au-S-GTTAGCAC	CGATAGTxxxATAGTCAGTCAGTCC-3'	NO				
	1111111						
	3'-CAATCGTO	GCTATCAxxxTATCAGTCAGTCAGG-5'					
5	Au-S-TTTTTTT	TTTTTTTTGGTTTTTTTTTTTTTT-3'	NO				
	1111111						
	3'-AAAAAAA	AAAAAAACCAAAAAAAAAAAAAAAA-5′					
6	Au-S-TTTTTTT	TTTTTTTGGTTTTTTTTTTTTT-3'	NO				
	1111111						
		AAAAAAAGGAAAAAAAAAAAAAAAA					
7	Au-S-TTTTTTT	TTTTTTTCCTTTTTTTTTTTTT-3'	NO				
	1111111						
	3 ′ - AAAAAAA	AAAAAAACCAAAAAAAAAAAAAAAA-5′					
8	Au-S-TTTTTTT	TTTTTTTCCTTTTTTTTTTTTT-3'	NO				
	1111111						
		AAAAAAAGGAAAAAAAAAAAAAAAA					
9	Au-S-TTTTTTT	TTTTTTTXxTTTTTTTTTTTTTT-3'	NO				
	1111111						
		AAAAAAAxxAAAAAAAAAAAAAAAAAAA					
10	Au-S-CGCCGCCC	GCCGCCGCCGCCGCCGCCG-3'	Uncharged p-	-isopropoxy			
		cgccgccgccgccgccgcc-5'					
11		TAATAATAATAATAATAATAAT-3'	Uncharged p-	-isopropoxy			
		ATTATTATTATTATTATTATTA-5'					
12	Au-S-GTTAGCAC	CGATAGTCCGATAGTCAGTCC-3'	Uncharged p-	-isopropoxy			
	3'-CAATCGTO	GCTATCAGGCTATCAGTCAGTCAGG-5'					

Single-stranded samples are labeled with letters A–E and named, double-stranded DNA are numbered 1–12. G, A, C, T are the symbols that designate the deoxynucleotides with either unmodified backbone (NO "backbone modification") or the phosphate charge neutralized by an isopropoxy group (Fig. 1), whereas the symbols "x" indicate abasic nucleotides. Strands chemically attached to the gold plate start with Au–S. For double-stranded samples, "|" labels the Watson–Crick pair while "‡" indicates a non-Watson–Crick pair.

ChemGenes Corporation, USA, and also used according to the manufacturer's recommendations. All oligonucleotide sequences were reverse-phase chromatography purified. The 5'-thiol-modified oligonucleotides were purified according to the manufacturer recommendations, aliquoted immediately after the purification and kept under argon atmosphere before use to prevent oxidative dimerization of the sulfhydryl groups. All the synthesized oligonucleotide sequences were analyzed by HPLC and MALDI-TOF for quality control. According to the MALDI-TOF analysis, the purity of all sequences was >98%.

All the chemicals were purchased from Sigma Aldrich, USA. The blocking solution of 1mM 2-mercaptoethanol in MilliQ water (BS), a hybridization buffer of 100mM Tris.HCl/100mM NaCl in MilliQ water (HB) and a washing buffer of 100mM Tris-HCl/300mM NaCl in MilliQ water (WB) were prepared by dissolving the respective amounts of chemicals in pure MilliQ water, free of DNA/RNA and nucleases (Millipore Inc., USA). The 5'-thiol-modified oligonucleotides were dissolved in MilliQ water to 50µM concentration prior to use. 5µl of 50µM solution of each 5'-thiol-modified oligonucleotide were spotted onto the gold support, forming a drop of ca 3mm in diameter. The spotted gold support slides were kept closed in a cartridge with humidified atmosphere created by water-filled basins. This cartridge

was incubated at 50°C for 16h, after which the gold support was washed twice with MilliQ water, and 5µl of 1mM 2-mercaptoethanol were spotted onto the same places as the oligonucleotides. The 2mercaptoethanol (2-ME) was used to enhance the accessibility of the immobilized probes to the complementary target sequences. The thiol group of 2-ME rapidly displaced the weaker absorptive contacts between the DNA nucleotides and the substrate, leaving the probes tethered primarily through the thiol end groups. The spots were "selfaligned", because the gold support is hydrophobic and any water solution put onto the oligonucleotide-modified gold support concentrates in the oligonucleotide-containing spots. The slides were incubated in the same humidified cartridge at 40°C for 2h, after which the slides were washed 5 times with MilliO water and dried under high-purity nitrogen (9.6). The gold-support slides with singlestranded oligonucleotide spots were kept in a closed box until use. The gold-support slides with the oligonucleotide spots planned to be analyzed as double-stranded were hybridized with the corresponding complementary oligonucleotides. The complementary oligonucleotides were incubated at 1µM concentration in HB at 80°C for 10min. Immediately after the incubation, 5µl of the corresponding complementary oligonucleotides were dropped onto the particular spots and

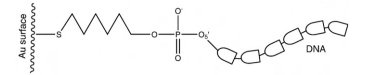


Fig. 2. The scheme of the attachment of the DNA 32-mers to the gold surface.

incubated at 40°C for 1h, after which the spots were washed twice by WB and finally by MilliQ water. Subsequently, the slides were dried under a stream of nitrogen. The whole process was photographed for the easier alignment of the STM tip onto the oligonucleotide spots. All the slides were kept in a closed box in a cool and dry place before use.

DNA molecules were attached to the gold surface through a sulfurgold linkage (Fig. 2) [22]. The ssDNA with the 5'-thiol-modifier were used to produce a thiol group at the 5'-terminus of a synthetic oligonucleotide. Recent neutron reflective studies have indicated that ssDNA oligonucleotides form a compact layer on bare gold. Prior to the formation of thiolated-ssDNA, self-assembled 40nm thin Au (99.99% purity) monolayers were vacuum-evaporated on single crystalline silicon substrate <100> wafers, which were cut into 1cm × 1cm pieces. Subsequently, the substrates were blown using pure nitrogen and exposed to a solution of thiolated oligonucleotide probes in a sodium chloride buffer.

2.2. Measurements

The STM measurements were performed with the NTEGRA Prima NT MDT system under ambient conditions. Both topographic and spectroscopic data were obtained using freshly cut Pt/Ir tips. With STM, for a set point of small voltage ~ 10mV and relatively large current ~ 0.5nA and thus quite a small tip-sample distance, we observed the topography of the samples. The topographic images showed a considerable difference between the bare gold substrates and the samples with DNA molecules. In the case of molecular layers, specific surface patterns were observed. The occurrence of these patterns was attributed to an etching of the bottom gold layers arising from the reaction of the thiol-groups with the gold atoms, forming dissolvable complexes [23,24].

The DNA layers thickness have been investigated using variable angle spectroscopic ellipsometry (VASE, J.A.Woollam & co.) working in rotating analyzer mode. The measurements were carried out in the spectral range 300-1100nm at three angles of incidence 65, 70 and 75°. The structural model was constructed to extract the optical functions of the film from the ellipsometry measurements. Structural model consists of: Si substrate/Au layer/DNA layer. The Si-substrate was considered as a semi-infinite medium. The optical constants of Si and Au materials were taken from [25]. The thicknesses of both layers and the optical constants of DNA layer were determined using a direct fitting procedure applied to the experimental ellipsometric data. The DNA molecules were ordered in similar geometric structures with the nearest-neighbor spacing being 1-2nm. The thickness of the DNA layers, determined by ellipsometry was 8.5-9.0nm. For 32-nucleotidelong DNA molecules, this length corresponds to the average distance between the neighboring bases of about 0.3nm, which corresponds to the base-base stacking distance intermediate between the B- and A-DNA forms. The DNA double strand is not likely to be completely straight. The mean inclination of a double helix from the fixed vertical line at the length of 32 base pairs is expected to be about 25° as can be estimated from the generally accepted persistent length P of a dsDNA of "random sequence", P \sim 150 base pairs. A confrontation of the ellipsometry measurement and the persistent length of a DNA double strand leads to a conclusion that DNA molecules are oriented approximately perpendicularly to the Au substrate.

In all cases, we qualitatively compared conductivity of two types of oligonucleotides. In our experimental setting the tip–sample distance was necessary and not easy to control. Firstly, we worked in constant height mode in very small area on both sides of the compared oligonucleotides type border. Secondly, beside current-voltage (I(V)) characteristics we also measured I(h) curves (I — current, h — distance between the tip and the surface) at the same points. I(h) curves have the advantage of being normalized and in [26] new approach for reading sequence of DNA molecule via tunnel-current decay has been demonstrated. The I(h) decay consists of two regions: the area of very short-range chemical interactions between the end of the tip and the terminate of the molecule plus the area of tunneling through the barrier between the tip and sample. The current initially decays slowly and then more rapidly. The I(h) curves are standardly fitted with two exponentials:

$$i = i_0 \exp(-\beta_1 h), 0 < h < h_c$$

 $i = i(h_c) \exp(-\beta_2 (h - h_c)), h_c < h.$

Value of β_2 has been found in [26] as corresponding to decay constant of tunneling current. Thus, it can be supposed that at the breakpoint (h_c) the interactions typical for tip extremely close to the substrate are broken and the tunnel current becomes dominant. The same position of the breakpoint and decay constants (β_2 , β_1) means the same tip-surface interaction conditions. Working with different set points we can arrange the position of the tip in the proper interaction territory so that the distance between the tip and the DNA terminate is under control. Comparing histograms of the decay constants and I(h) curves breakpoints (h_c) measured on various DNA samples we controlled position of the tip above the compared DNA modifications.

In our experiments, we have measured current passing through the molecules so close to the tip that their contribution is at set voltage higher than the noise. For tip very close to the molecules (defined by the set point value) and very low voltages the number of the molecules contributing to the total current is as small as possible. Current-voltage characteristics were recorded for various feedback voltage and current set points, i.e. for different initial tip-sample distances. In all the experiments, the STM tip acted as the electrical contact on the "top" side of the assembled monolayer of the DNA molecules, whereas the supporting gold substrate acted as the other "bottom" contact. The distance between the top of the molecules and the STM tip was for all the set points 2.5–4Å. The I(V) and I(h) curves were measured at three different set points: 0.1nA, 0.1V; 0.2nA, 0.1V; 0.5nA, 0.1V, corresponding to three different sample–tip separations, using the same strategy for all the samples. The experimental conditions were controlled using I(h) decay analysis and thus we were able to distinguish between the area of tunneling current and current passing through chemically interacted systems. The experimental setup (tip very close to the surface) allows us to expect that the current flows mainly through the molecule nearest to the tip without significant lateral intermolecular contribution to conduction. Two hundred consecutive I(V) sweeps in both voltage directions were

Table 2A comparison of the conductivity of DNA samples measured on the same gold plate in one experiment

DNA samples measured in one experiment	1/2	A/1	3/4	5/9	8/9	1/10	2/11	8/7	5/6	3/12
DNA samples with higher conductivity	1	1	3	5	8	1	2	8	5	3

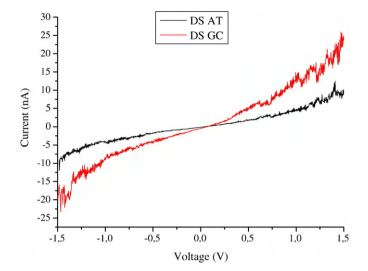


Fig. 3. The typical I(V) curves of DNA GC 32-base double strand (DS GC, see Table 1, ds 1) and DNA AT 32-base double strand (DS AT, see Table 1, ds 2). Set point 0.1 nA, 0.1 V, measured in one voltage direction.

taken on each sample (Table 1). Final results presented in this paper are based on the average of a set from the collected I(V) curves. Only those curves which were not affected much by drift of the STM were included in the statistics.

All the results taken into the consideration showed the same trends, i.e. the symmetry for both biases, linearity and super-linearity for lower and higher voltages, respectively. Due to large statistics we can (from our measurements) set the quantitative proportion of different DNA systems conductivities assuming approximately equal number of molecular densities.

3. Results and discussion

The results of the measurements are summarized in Table 2 and Figs. 3–7. The conductivity measurements for each considered structural and chemical feature and their possible explanations are discussed below. Noise, in the form of height fluctuations, was higher in the case of the topographic images of films than in the case of bare

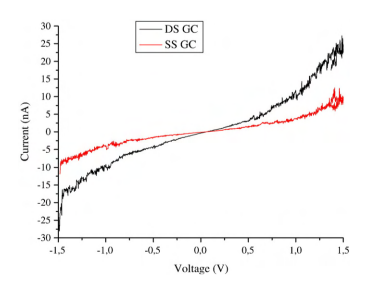


Fig. 4. The typical I(V) curves of DNA GC 32-base single strand (SS GC, see Table 1, SS A) and DNA GC 32-base double strand (DS GC, see Table 1, ds 1). Set point 0.1 nA, 0.1 V, measured in one voltage direction.

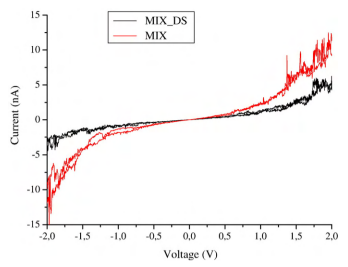


Fig. 5. *I*(*V*) curves of double-helical DNA 32-mers with mixed G, A, C, T sequences (Table 1, ds sample 3) and double-helical DNA 32-mers with mixed G, A, C, T sequences with three bases missing in the middle (MIX_DS, see Table 1, ds 4). Set point 0.1 nA, 0.1 V, measured in both voltage directions.

gold. All the organic molecules that have been studied by STM until now show symmetry of the (I(V)) characteristics. The symmetry of the I(V) characteristics has been explained on a simple model [27].

The conductivity of all 32-nucleotide-long DNA molecules including the sulfur spacer was comparable in the STM experiments. We measured the conductivity of the DNA molecules using a scanning tunneling microscope under ambient conditions, i.e. in a humid air (40% humidity) environment and room temperature. Xu et al. [21], who measured DNA samples using ultrahigh vacuum scanning spectroscopy, concluded that DNA was a wide-band-gap insulator. From this comparison, we can see that the absence or at least substantially lower content of water in the ultrahigh vacuum plays a significant role in the charge pathway. In agreement with what Enders et al. pointed out in their summary [28], we have concluded that water molecules supporting the molecular structure improve the conditions for the electrical current passing through the whole DNA molecule.

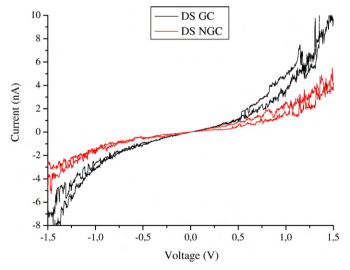


Fig. 6. I(V) curves of double-helical DNA 32-mers with G-C bases without charge-bearing groups on the phosphates – DS NGC (see Table 1, ds 10) and standard double-helical DNA 32-mers with G-C bases – DS GC (see Table 1, ds 1). Set point 0.1 nA, 0.1 V, measured in both voltage directions.

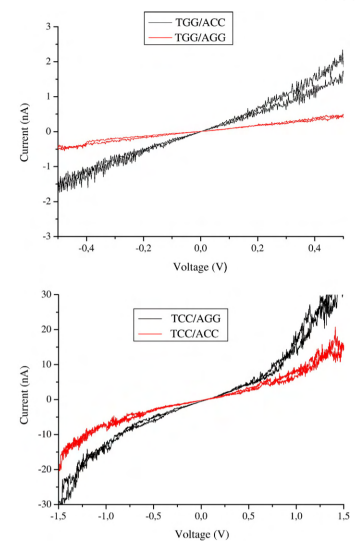


Fig. 7. Typical *I(V)* curves of DNA molecules containing 4 non-Watson-Crickbase pairs in the middle (TGG vs. AGG and TCC vs. ACC, see Table 1, ds 6 and ds 7, respectively) and standard dsDNA (TGG vs. ACC and TCC vs. AGG, see Table 1, ds 5 and ds 8, respectively). Set point 0.1 nA, 0.1 V, measured in both voltage directions.

3.1. G/C chain vs. A/T chain

The Guanine/Cytosine-rich and Adenine/Thymine-rich oligonucleotides (Table 1, ds 1 and 2) were selected for the evaluation of the guanine effect on the overall DNA conductivity.

We found that a double-helical sample with G/C bases is a better conductor than the double strand made of A/T bases (Fig. 3, Table 2). The guanine base is known to be easily oxidized, generating a charge carrier (hole). Once charges, and especially holes, are created in the uniform DNA chain, the hopping charge transport can apparently occur between discrete guanine sites or delocalized (e.g. polaron) domains [29]. Furthermore, the stacking of the adjacent base pairs is also likely to affect the conductivity via the π -electron overlap. The G-C pairs with more compact stacking compared to that in the A-T pairs have higher conductivity.

3.2. Double-stranded vs. single-stranded DNA

Typical I(V) curves of both ss and ds GC sequences (A and 1 in Table 1) are shown in Fig. 4. The conductivity of the double-stranded form is higher than that of the single strand when complementary sequences are compared, which is in agreement with the theoretical model of the same system [30]. Single strands can be considered much

less structurally regular than the double helices because of their local and especially long-range deviations from the regular helical arrangement. Reduced stacking interactions then decrease the potential overlap of their π -electron systems and overall conductivity. Also the first solvation shell, known to play a crucial role in the structural integrity of the double-helical DNA [31,32], is likely to be much less ordered in ss than in dsDNA. Double-stranded DNA has a more regular structure including the first solvation shell, and its long-range periodicity decreases the dispersion of the polarization energy and makes the distribution of hopping states narrower. Considering these effects, the charge-carrier mobility, and thus conductivity, increases in more regular dsDNA systems.

3.3. Base pairing — Watson-Crick vs. "mismatched" pairs

Non-Watson-Crick or "mismatched" base pairs are the type of DNA damage where two non-complementary bases are paired, e.g. adenine with cytosine or thymine with guanine. The role of the non-W-C base pairs and their different stacking in DNA conductivity was estimated by comparing the conductivity of two dsDNA, the canonical 32-nucleotide-long double-stranded DNA molecules (Table 1, ds 5 and 8) with analogical systems containing two non-W-C pairs in the middle (Table 1, sequences 6 and 7). The unequivocal conclusion is that the molecules containing non-W-C pairs are less conductive than the DNA double strands containing only W-C pairs (Fig. 7, Table 2). The mismatched molecules are expected to be under mechanical stress, and the disrupted regular periodicity of the stacked W-C pairs and their possibly worse stacking interaction can lead to a decrease in the electron transport efficiency [33].

3.4. Abasic nucleotides — structure change and interrupted stacking

The role of base stacking in DNA conductivity was also estimated by comparing the conductivity of two double-stranded sequences with a mixed G/A/C/T sequence, one containing all bases in W–C pairs (number 3 in Table 1) and the other with the same sequences except for three bases in the middle, at positions 15–17, which were substituted with an abasic nucleotide spacer (number 4 in Table 1). The measurements consistently showed that samples with the abasic spacer had a lower conductivity than the standard DNA sample. The I(V) curves obtained for double-helical DNA samples 3 and 4 are compared in Fig. 5. In this case, the conductivity of the sample with the gap in the middle of the sequence was much lower than the conductivity of a standard DNA chain. The same result – a lower conductivity of an abasic dsDNA sample when compared with an intact dsDNA – was obtained in the case of the ds AT chain with two bases missing in the middle (sample 9).

There are no examples of atomic-resolution structures of DNA molecules with two or even three base pairs replaced by an abasic nucleotide. However, structures missing just one base in one of the two strands in a double-helical construct may be significantly deformed, as for example in the crystal structure of human APE1 bound to abasic DNA [34]. Therefore, it is not possible simply to dissect the effect of the overall ds structure and quality of the stacking; both effects, less regular double-helical structure and sub-optimal stacking interactions, are likely to cause the lower conductivity of the abasic samples. The DNA backbone constitutes a quasi-one-dimensional periodic system, which is essentially independent of the base pair sequence and could allow for extended Bloch states. Nevertheless, conductivity through the backbone seems to be low because of the insulating sugar groups separating the phosphate groups from each other.

3.5. Phosphate charge

To study the influence of the DNA negative charge on its conductivity, we synthesized DNA chains with phosphates modified by isopropoxy groups, thus eliminating phosphate negative charges (*p*-isopropoxy-modified sequences 10, 11, 12 in Table 1). The measurements undertaken for the pairs of double-stranded samples 1–10, 2–11, and 3–12 (Table 1) demonstrated that in all three cases the *p*-isopropoxy-modified DNA sequences had a lower conductivity than their counterparts of natural DNA (Table 2, Fig. 6).

In the polar aqueous environment, DNA forms a double helix with the hydrophobic base planes shielded from the aqueous solvent by the base-pair stacking in its core region with the charged phosphate fully exposed to the solvent. The phosphate negative charges are compensated by partially condensed counterions, typically mono- and bivalent alkali- and alkaline-earth metals such as Na⁺ and Mg²⁺. The metals are fully integrated into the solvation shell, which is partially ordered [35-37]. On the basis of a MALDI-TOF analysis, the presence of Na⁺ and K⁺ cations in the p-isopropoxy-modified sequences is highly unlikely, and these metal cations probably do not play any role in the conductivity of sequences 10 and 11. In natural sequences with the presence of Na⁺ or Mg²⁺ counterions, the conductivity is increased partly due to the Na⁺ or Mg^{2+} states being localized in the large π - π * energy gap. In the case of sodium, we have small activation gaps (of a few kT) between the water and sodium states, which could lead to hopping conductivity between the Na⁺ centered states. For Mg²⁺, the occupied water-state energies are not only close to the Mg^{2+} levels but also very close to the unoccupied π^* states, leading to the possibility of electron doping of DNA by water or Mg²⁺. Like in the case of the abasic dsDNA, the lower conductivity of uncharged dsDNA is caused, besides the electronic effects, also by the less regular structure of the uncharged dsDNA.

The shape of the I(V) curves can in general be interpreted as follows: The current passes through the molecules and also tunnels through the tip-molecule area. For low voltages, ohmic behavior was observed due to the Boltzmann distribution of the charge carriers and the constant position of the Fermi level. The higher the voltage is the higher is the current passing through each molecule, resulting in the nonlinear effect of the charge-carrier injection (the shift of the DNA Fermi level to the electronic tail states and their occupation). Therefore, based on our topographic data, we assume that the differences in the conductivity of different samples come from the properties of the individual molecules, not from the specific properties of the molecular monolayers.

4. Conclusions

The STM technique was used to study the conductivity of various DNA sequences in single- and double-stranded forms with one strand always covalently bonded to the gold surface. The purpose of our investigations was to provide more insight into the complex process of the charge–carrier transport in DNA molecules — the role of the sugarphosphate backbone, the role of the counterions complexed around the negatively charged phosphate oxygens, and the effect of the interactions between the bases and their pairs.

The double-helical sample with the G/C bases is a better conductor than the double-stranded sequence containing only A and T nucleotides. Easy oxidation of the guanine base makes it possible to generate the charge carriers (holes). The charges, especially holes, created on the uniform DNA chain can move by the hopping charge transport through the discrete guanine sites.

The conductivity of the double-stranded form is greater than that of the single-stranded one when complementary sequences are compared. Single strands can be considered much less structurally regular than the double helices. Regular structures could form better long-range ordering, which decreases the dispersion of the polarization energy and makes the distribution of hopping states narrower. Under these conditions, the charge–carrier mobility, and thus conductivity, in regular systems increases.

We found that the conductivity of the samples with abasic spacers was lower than that of the natural dsDNA samples because of the lack of stacking interactions and likely also because of the structural irregularities and lost periodicity of the double strand.

The DNA chains without charge-bearing phosphate groups were less conductive than natural dsDNA. The reasons are likely to be both electronic (the fact that neutral DNA lacks counterions with their effect on lowering the energy gaps between nucleotides) and structural (neutral DNA is likely to be distorted, irregular and hence more insulating than the natural dsDNA) [38].

Theoretical computations show that the orbitals occupied by the π -electrons contribute strongly to the electronic states, which arises from the electronic coupling between the neighboring bases [39,40]. Unlike the π -electrons, which can form extended states, water molecules and counterions create localized states of electrons, resulting in the hopping mechanism of conduction, with the inclusion of the electron-phonon interactions. The DNA molecule contains electronic states, which have an extended character and in which the electrons easily carry a ballistic current, and localized states of electrons, mediating the hopping mechanism of conduction. These two rather complementary pictures of conduction, π -electron overlap and hopping through energetically near states, may fit together.

Based on our measurements, we can conclude that the best conductor is the natural double-stranded DNA form with its bases forming Watson–Crick pairs and surrounded by counterions and water molecules. Since electronic states are strongly connected with molecular structure and DNA structure changes with base sequences, type and concentration of counterions, and relative humidity, the electronic states are expected to depend on all of the following parameters:

- 1. change of the regular structure;
- 2. water (as a possible ion conductor) amount;
- 3. counterion presence; in the case of DNA containing sodium, Na⁺ can mediate hopping conductivity between the Na⁺-centered states; for DNA containing magnesium, there is a possibility of electron doping of DNA by water or Mg²⁺ states.

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