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5-Bromo-2'-deoxycytidine—a potential DNA photosensitizer†

Magdalena Zdrowowicz, Paweł Wityk, Barbara Michalska and Janusz Rak*

A double-stranded oligonucleotide, 80 base pairs in length, was multiply labeled with 5-bromo-2'-deoxycytidine (BrdC) using polymerase chain reaction (PCR). The modified oligonucleotide was irradiated with 300 nm photons and its damage was assayed by employing DHPLC, LC-MS and denaturing polyacrylamide gel electrophoresis (PAGE). Two types of damage were demonstrated, namely, single strand breaks (SSBs) and intrastrand cross-links (ICLs); the ICLs were in the form of d(G^C) and d(C^C) dimers. The former species are probably formed due to photoinduced electron transfer between the photoexcited BrdC and the ground state 2'-deoxyguanosine (dG), whereas the latter is a result of a cycloaddition reaction. Since SSBs and ICLs are potentially lethal to the cell, BrdC could be considered as a nucleoside with possible clinical applications.

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1. Introduction

In the late 1950s, it was already demonstrated that 5-bromo-2'deoxyuridine (BrdU) sensitizes E. coli cells to ultraviolet light,^{1,2} which is related to the incorporation of BrdU into cellular DNA. The strand breaks and cross-links belong to the most common damage induced by UV photons in BrdU-substituted DNA.³ As indicated by several photochemical studies, single strand breaks in such a system are formed primarily via the UV-induced long-range electron transfer (ET) from a distant ground-state guanine to the photo-excited BrdU.⁴ This process leads to the formation of a short-lived BrdU anion that releases the bromide anion⁵⁻⁷ and leaves the reactive uridine-5-yl radical in the DNA. The latter species is stabilized by abstracting a hydrogen atom from the adjacent sugar moiety that ultimately gives a strand break.8 On the other hand, cross-links, both intra- (intra-CLs) and interstrand (inter-CLs), appear, due to the attack of the uridine-5-yl radical on an adjacent base, giving a covalent base-base dimer as the end-product.9

Much less is known about the photochemical properties of the remaining bromonucleosides (BrNucs). Discussing sparse data concerning the photochemistry of non-BrdU labeled DNA, one should mention the studies of Wang *et al.*,^{10,11} which demonstrate the formation of UV-induced intra-CLs in duplex oligonucleotides, point labeled with 5-bromo-2'-deoxycytidine (BrdC). Furthermore, the Chatgilialoglu group^{12,13} has shown the efficient photodebromination of 8-bromo-2'-deoxyadenosine (BrdA) and 8-bromo-2'-deoxyguanosine (BrdG) incorporated into dsDNA, induced by long-range electron transfer from a flavin electron donor to the brominated base. Although the rate of debromination was significantly smaller for bromopurine nucleosides than for BrdU, the overall process proceeds in a similar fashion in all the systems compared, which suggests that the brominated purines may play the role of DNA sensitizers. The properties of BrNucs mentioned above suggest that they could be exploited in a photodynamic therapy (PDT) that does not involve singlet oxygen-a species inherently related to PDT treatment.¹⁴ Since tumors are frequently hypoxic,¹⁵ the oxygen-independent mode of action is a clear advantage of BrNucs, as compared to most PDT drugs.

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In the current work, we study the UVB damage to a doublestranded oligonucleotide, multiply substituted with 5-bromo-2'-deoxycytidine, in order to assess its photosensitizing properties. By comparison to the BrdU photosensitizer whose main mode of action is related to long range photoinduced ET,¹⁶ we assumed that BrdC should possess similar properties. Employing the PCR method, we synthesized an 80 bp DNA fragment in which all but the dC nucleosides present in the primers were replaced by BrdC. Using denaturing HPLC (DHPLC), gel electrophoresis and high-resolution mass spectrometry against the UV irradiated samples, we demonstrated the photochemically induced formation of strand breaks and intrastrand cross-links in the BrdC labeled oligonucleotide. Interestingly, all SSBs were formed in "hot spot" sequences, similar to those characteristic of photochemically induced SSBs in DNA labeled with BrdU (i.e. in the fragments

Faculty of Chemistry, University of Gdańsk, Wita Stwosza 63, 80-308 Gdańsk,

Poland. E-mail: janusz.rak@ug.edu.pl; Fax: +4858523 5771; Tel: +4858 523 5118 † Electronic supplementary information (ESI) available: (i) PCR protocol (ii) Fig. S1 – native PAGE, (iii) Fig. S2 – comparison of UV spectra for BrdC and native DNA, (iv) Fig. S3 – degree of damage vs. irradiation time and (v) Fig. S4 – chromatogram employing the total ion current observed in the irradiated photolyte. See DOI: 10.1039/c6ob01446a

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where BrdC was separated from the G base by several AT base pairs). Since SSBs may lead to double strand breaks (DSBs; even a single unrepaired DSB is sufficient to kill a eukaryotic cell¹⁷) during DNA repair or biosynthesis,¹⁸ and intra-CL causes the most cytotoxic¹⁹ damage; the studied nucleoside (possibly in the form of an appropriate pro-drug) could broaden a very modest pool of PDT drugs used nowadays.²⁰

Experimental section 2.

Chemicals and reagents

The enzymes for enzymatic digestion [DNase I, snake venom phosphodiesterase (SVP), calf spleen phosphodiesterase (CSP), the bacterial alkaline phosphatase (BAP) and nuclease P1], the mobile phases components [LC-MS grade acetonitrile, methanol, hexafluoroisopropanol (HFIP), formic acid, acetic acid and triethylamine] and standard deoxynucleotide set were purchased from Sigma Aldrich; the DNA purification kit from Eurx Molecular Biology Products (Gdansk, Poland), and Marathon DNA polymerase from A&A Biotechnology (Gdynia, Poland). The 5-bromo-2'-deoxycytidine-5'-triphosphate (BrdCTP) was bought from TriLink Biotechnologies (San Diego, CA, United States). The oligonucleotides (PCR primers and template) were obtained from GeneSys (Wroclaw, Poland). Ultrapure water was obtained using the Millipore system (Merck Millipore).

PCR Reaction with BrdCTP

The double-stranded oligonucleotide labeled with BrdC was obtained in the PCR reaction. The resulting PCR products were 80 bp in length (see Fig. 1). The amplified DNA fragments were purified with the use of a PCR/DNA clean up purification kit and then with HPLC.

UV irradiation

Photolysis was carried out in guartz capillaries $(3 \text{ mm} \times 3 \text{ mm})$ filled with a DNA solution (200 ng μ l⁻¹, 4.54 × 10⁻⁶ M) containing Tris-HCl (pH ~ 8.0) in a total volume of 50 µl with the use of a 500 W high-pressure mercury lamp. The 300 nm wavelength of incident light (half-width 2.5 nm) was selected using a grating monochromator (M250 Optel, Poland). The incident light intensity for the above irradiation setup was c. 41 W m⁻². This value was determined with a Maestro11 laser power/ energy meter, equipped with an 11PD photodetector (Standa, Lithuania).

Denaturing PAGE

15% denaturing (7 M urea) polyacrylamide gels were prepared in 1× TBE buffer. Samples were incubated at 95 °C for 10 min before loading onto the gel. The gel was electrophoresed at 200 V for 2 hours. Then the gel was visualized after staining with Sybr Gold using a Fusion FX imaging system (Vilber Lourmat, Germany).

Enzymatic digestion

0.2 U of nuclease P1 and 0.005 U of calf spleen phosphodiesterase were added to a solution containing 10 µg of photoirradiated DNA, 300 mM sodium acetate (pH 5.0) and 10 mM zinc acetate. The digestion was carried out at 37 °C for 3 h. After the incubation, 40 U of DNase I, 10 mM CaCl₂, 100 mM MgCl₂, 100 mM tris-HCl (pH 8.9), 400 µM DTPA and water were added to the mixture. The digestion was continued at 37 °C for 1 h. Then, 1 U of bacterial alkaline phosphatase was added and the sample was incubated at 37 °C for an additional hour. In the final step of the protocol, 0.00004 U of snake venom phosphodiesterase was added and the incubation was prolonged for one more hour. Finally, the solution was chloroform-extracted to remove the enzymes. The aqueous layer was lyophilized and then resuspended in water for further analysis.



Forward primer ₃, **TGCTGTCCAAAGGGCTGACCT T TCGCCC GTCA CTCG CGTTGCGTTAATT ACACTCAATCGA**GTGAGTAATCCGTGGGGTC **Reverse primer** В. A40 A39 C17 C62 **B73**

B6 ⁵′ A C G A C A G G T T T C C C G A C T G G A A A G B G G G B A G T G A G B G B A A T T A A T G T G A G T T A G B T B A T T A G G B A B B B A Y T G B T G T B <mark>B A A A</mark> G G G B T G A B B T T T B G B B B G T <mark>B A B</mark> T B G B G T T G B G T A A T T A <mark>B A B</mark> T B G A G T G A G T G A G T A A T CGTGGGGT D31 D48 G51 G28 E55 E24 F71

Fig. 1 (A) DNA sequence of non-labeled DNA fragment, 80 bp in length. (B) DNA sequence of DNA fragment labeled with 5'-bromo-2'-deoxycytidine. The incorporated 5'-bromo-2'-deoxycytidines are marked in blue. The DNA sequences responsible for long-range electron transfer are shown in the yellow boxes; the numbers of the nucleotides in the particular fragments formed as a result of secondary single strand break are given in brackets.

Chromatography

The HPLC and DHPLC separation was performed on a Dionex UltiMate 3000 System with diode array detector, which was set at 260 nm for monitoring the effluents. An XBridge OST, reversed-phase C18 column (2.5 μ m in particle size, 4.6 × 50 mm) was used. For the purification of PCR products, a 20 min gradient of 0–16% acetonitrile in 50 mM triethylamine acetate (pH 7) was employed, and the flow rate was 1 mL min⁻¹. For the DHPLC separations, the conditions were as follows: a 20 min gradient from 8.4 to 9.2% acetonitrile in 100 mM triethylamine acetate (pH 7), a flow rate of 0.5 ml min⁻¹ and a column temperature of 70 °C.

Mass spectrometry

LC-MS identification of single strand breaks. LC-MS analysis of DNA fragments was carried out using a ultra high performance liquid chromatography (UHPLC) system, Nexera X2, coupled to a mass spectrometer, TripleTOF 5600+ (AB SCIEX), equipped with a duo-electrospray interface operated in negative ionization mode. Chromatographic separations of oligonucleotides were performed using an Acquity UPLC BEH C18 1.0×50 mm column (Waters) at a flow rate of 0.1 mL min⁻¹. The mobile phase A consisted of 400 mM HFIP and 15 mM TEA in deionized water, and mobile phase B consisted of the same concentration of HFIP and TEA in the water-methanol mixture (50:50, v/v). We separated 20 µl of irradiated sample using the following gradient conditions [time (min), % mobile phase B]: (0, 0) (20, 20) (50, 40) (65, 40) (70, 60). In the case of the nonirradiated sample, the linear gradient 0-60% of phase B in 40 min was used. The effluent was diverted to waste for 1 min after injection during each analysis. The column temperature was set at 80 °C. MS operation parameters were as follows: the spray voltage -4.5 kV, the nebulizer gas (N₂) pressure 30 psi, the flow rate 11 L min⁻¹ and the source temperature 300 °C.

LC-MS identification of intrastrand cross-links. LC-MS/MS analysis of irradiated and digested DNA fragments labeled with BrdC was performed using an Agilent 1200 Technologies HPLC System. A Wakopak® Handy ODS column (4.6 mm × 150 mm; 5 µm in particle size and 100 Å in pore size) with a mobile phase consisting of deionized water, acetonitrile and 0.1% formic acid (pH 2.55; 87.7 : 2 : 10.3, v/v/v) was used. 20 µL of digested sample was injected in each run. The effluent was coupled to a Bruker Daltonics HCT Ultra Ion Trap MS with an electrospray (ESI) ion source, which was operated in negative-ion mode. The ion source parameters were set as follows: the source temperature of 360 °C, spray voltage of -4.0 kV, the drying gas (N₂) pressure of 50 psi and the drying gas flow rate at 11 l min⁻¹.

3. Results and discussion

Photochemical studies on labeled DNA are frequently carried out with short oligonucleotides comprising a single molecule of the sensitizer.³ Such an approach enables the sequence dependency of the studied process to be comprehended. However, due to a significant number of possible sequences, even in short DNA, one has to study a multitude of systems in order to get a complete picture. Even more important is that in cellular environments, where DNA labeling proceeds in a statistical fashion (the chance for a modified nucleotide to be incorporated into a newly synthesized strand depends on the relative amount of its triphosphate in the overall pool of triphosphates), all possible sequences occur in the biosynthesized polymer. Therefore, to study a situation possibly close to that occurring in the cell, we decided to synthesize an oligonucleotide similar to that produced during DNA biosynthesis. Hence, we employed the PCR reaction in order to obtain a labeled DNA fragment in which all the 2'-deoxycytidines, besides those present in the primers, are substituted with BrdC. A similar approach to the labeled DNA synthesis was used in the past by us,^{16,21} as well as by other groups.²²

In comparison to our recent studies on the BrdU-labeled dsDNA fragment,¹⁶ and also in the current studies, we amplified the same part of the pUC19 plasmid by PCR (for the PCR protocol, see ESI†). This time, however, all dCTPs were substituted with BrdCTPs, rather than TTPs with BrdUTPs, as we did in ref. 16. In Fig. 1A, the amplified sequence, primers and the sites where BrdC have been incorporated are displayed. One can note that altogether, 34 2'-deoxycytidines were substituted with BrdC.

Assessment of the PCR Product by LC-MS Analysis

The outcome of the PCR reaction was confirmed by employing PAGE (see Fig. S1 in ESI[†]). The significant yield of the PCR product (see a distinctive band corresponding to the mass marker of 80 bases in lane 3 of Fig. S1[†]) demonstrates that Marathon DNA polymerase accepts the brominated analogue of dC triphosphate. The formation of the expected band in gel electrophoresis is a generally accepted confirmation of an efficient PCR reaction. One should note however, that such confirmation constitutes only an indirect proof for the accuracy of the actual PCR process. Due to the small resolution of the gel, all PCR products having the same length, but differing in sequence, bases identity, etc., form bands that move with more or less the same velocity in an electric field. In order to unequivocally demonstrate that the required PCR product was actually synthesized, we employed a powerful analytical tool, which is recently becoming a golden standard of DNA analysis, i.e. liquid chromatography coupled to high-resolution electrospray mass spectrometry (LC-ESI-MS).²³ In Fig. 2, several variants of the LC-MS spectra for the PCR product are presented. The synthesized oligonucleotide was separated using a mobile phase dedicated to MS analysis of oligonucleotides, i.e., containing TEA and HFIP (see Methods). As shown in Fig. 2A, where the total ion current is plotted against the time of separation, the particular DNA strands are quite well separated. On the other hand, the MS spectra obtained in the ESI-MS experiments are shown in Fig. 2C and D-the particular oligonucleotides form multiply ionized ions and the deconvoluted masses remain in very good accordance with the theoretical ones (cf. Fig. 2C and D with Table 1). Finally, the extracted ion chromatograms for m/z equal to 891.8 and 871.8, respectively, are

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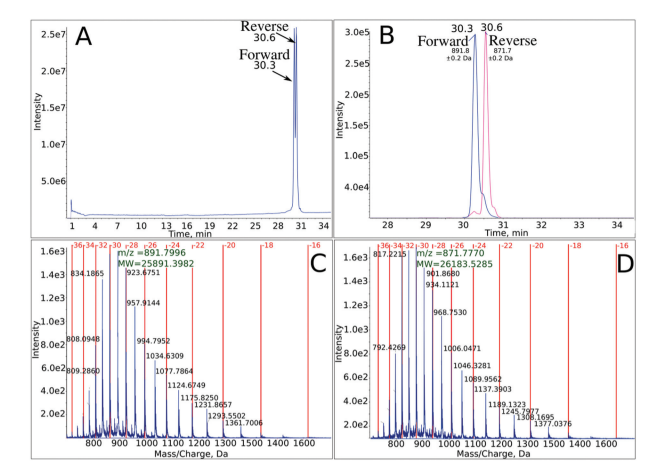


Fig. 2 LC-MS analysis of the PCR product - completely labeled with BrdC 80-mer oligonucleotide, with the sequence depicted in Fig. 1. (A) Total ion current chromatogram showing two separated strands. (B) Extracted ion chromatograms for m/z equal to 891.8 and 871.8 (for these most intensive signals, the charge state of the analyzed oligonucleotides is equal to -32) for the forward and reverse strands, respectively. (C) and (D) Mass spectra showing multiply ionized ions and the deconvoluted masses corresponding to the forward and reverse strands, respectively.

Table 1 Measured and calculated masses of strands of PCR product and fragments formed as a result of secondary single strand breaks (see Fig. 1)

routinely used for DNA identification, delivers a difficult to overestimate approval for the accuracy of PCR synthesis.

Identities	Calculated mass [Da]	Measured mass [Da]
Forward strand	25 890.6	25 891.4
Reverse strand	26 183.1	26 183.5
A40	13 109.0	13 109.6
B6	2053.9	2054.1
C17	5668.8	5668.5
D48	15 580.1	15 580.2
E28	9041.4	9041.3
F8	2614.3	2613.3
G24	7587.8	7687.4

displayed in Fig. 2B. All these results constitute a direct confirmation that the PCR process produced the expected doublestranded DNA fragment containing forward and reverse strands (see Fig. 1).

Thus, the above-described LC-MS identification of the PCR product complements gel electrophoresis and although not

UV irradiation

DHPLC analysis. In order to demonstrate the photoreactivity of the synthesized DNA, its irradiation with 300 nm photons was carried out in a buffered 4.54 µM solution. Similar to BrdU, the UV spectrum of BrdC is red-shifted compared to the absorption maximum of native DNA that occurs at 260 nm (see Fig. S2[†]). As a result, the absorption of labeled DNA, due to its substitution with BrdC is still substantial at around 300 nm, where the absorbance of the native oligonucleotide is relatively weak. Therefore, a significant fraction of 300 nm photons is absorbed by BrdC, rather than by the remaining native nucleosides. Indeed, using the absorbance of native and substituted PCR product measured at 300 nm, one can calculate that over 55% of the energy absorbed excites the BrdC molecules in the labeled PCR product.

Fig. 3 compares the DHPLC chromatograms of non-irradiated (A) and irradiated samples (B, C). The elevated temperature of HPLC separation brings about the analyzed double-

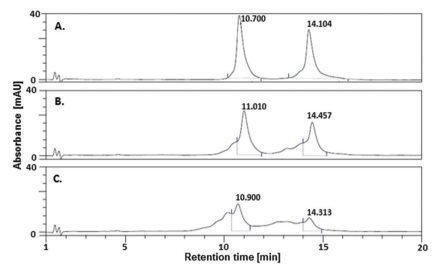


Fig. 3 DHPLC chromatogram of (A) a non-irradiated DNA fragment labeled with 5-bromo-2'-deoxycytidine and irradiated for (B) 3 min (C) 10 min.

stranded oligonucleotide moving through the column in the form of its single stranded components, giving two well-separated signals at 10.7 and 14.1 min (see Fig. 3A). Even 3 min irradiation leads to a visible degradation of the UV-exposed oligonucleotides (cf. Fig. 3A with B). The substrate signals, at 10.7 and 14.1 min, gradually disappear and a number of products are formed, resulting in several broad peaks with retention times shorter than those of substrates. Employing the incident beam intensity of 41 W m⁻², the actual absorbance of the sample at 300 nm, its degree of decay and the irradiated geometry, one can estimate the quantum yield of the overall process to be c. 2.5×10^{-3} . This value was estimated for 1 min irradiation, i.e., in the time range when the substrate decays linearly with the total incident energy. Indeed, we found out that for our experimental set-up, substrate decay against irradiation, time runs in a linear regime within the first seven minutes (see Fig. S3[†]). Hence, changes in the absorbance due to a change in the system's constitution with the time of irradiation should not significantly perturb the quantum yield calculated for the first minute of exposure.

It is worthy of emphasis that the quantum yield calculated for the same 80-mer sequence labeled with BrdU was as much as 8.4×10^{-3} (measured for 1 min irradiation, *i.e.*, the exposure time for which the degree of degradation *versus* irradiation time displays almost perfect linearity),¹⁶ which is *ca*. 3-fold larger than that for DNA labeled with BrdC. This result clearly indicates that BrdC is a significantly weaker photosensitizer than BrdU.

Denaturing electrophoresis and LC-MS analysis of single strand breaks. Although BrdC photosensitizes DNA to a lesser extent than BrdU does, the pattern of fragments generated in the 80-mer double-stranded oligonucleotide labeled with the former (see Fig. 1) is more complicated than that observed in ref. 16. In both labeled fragments, 300 nm photons produce the electronically excited $\pi\pi^*$ oxidizing state of the bromopyrimidine that can oxidize a distant guanine. Indeed, $\Delta G_{\rm ET}$ =

 $AIP_G - AEA_{BrC} - E_{h\nu}$, where ΔG_{ET} , AIP_G , AEA_{BrC} and $E_{h\nu}$ stand for the free energy of charge transfer between the excited BrC and ground state guanine, adiabatic ionization potential of aqueous guanine, adiabatic electron affinity of BrC, and excitation energy of BrC, respectively. Hence, for AIPG, AEABrC and $E_{h\nu}$ equal to 5.8,²⁴ 2.3⁶ and 4.1 ($h\nu$ = 300 nm) eV, respectively, one obtains a substantially negative $\Delta G_{\rm FT}$ of -0.6 eV. Therefore, one may expect that similar to the BrdU case, also in the BrdC-labeled DNA, long-range electron transfer from a distant guanine to the electronically excited bromopyrimidine occurs. By the same analogy, one can assume that substitution of Tris-HCl by a buffer lacking hydrogens like phosphate or cacodylate buffer would have no effect on the studied process as we demonstrated some time ago.¹⁶ Looking at Fig. 1, one can identify seven "hot spots", i.e., sequences in which BrdC is separated from the guanine located in the same or complementary strand by 1-2 AT base pairs. In all these configurations, the electronic excitation of BrdC should trigger ET, which in the sequence of secondary chemical reactions leads to the release of the neighboring adenine and the formation of thermally unstable 2'-deoxyribonolactone. Ultimately, the latter damage causes cleavage, giving two well-defined fragments. For the studied 80 bp sequence, 14 fragments with the phosphate group at the 5' or 3'-ends, respectively, should be observed (see Fig. 1). To confirm the assumed mechanism of UV-induced damage to the PCR synthesized 80 bp BrdClabeled DNA fragment, we again employed the LC-MS identification. Since, under ESI-MS conditions, oligonucleotides are identified as multiply charged ions, we decided to search for MS signals corresponding to the shorter of two fragments that are to be produced, due to the cleavage of 2'-deoxyribonolactone. The longer the identified fragment, the higher is the ionization degree necessary to make this fragment detectable in the mass detector range. Therefore, shorter oligonucleotides lead to stronger MS signals (see Fig. 4-the shorter the oligonucleotide, the less noisy is the spectrum).

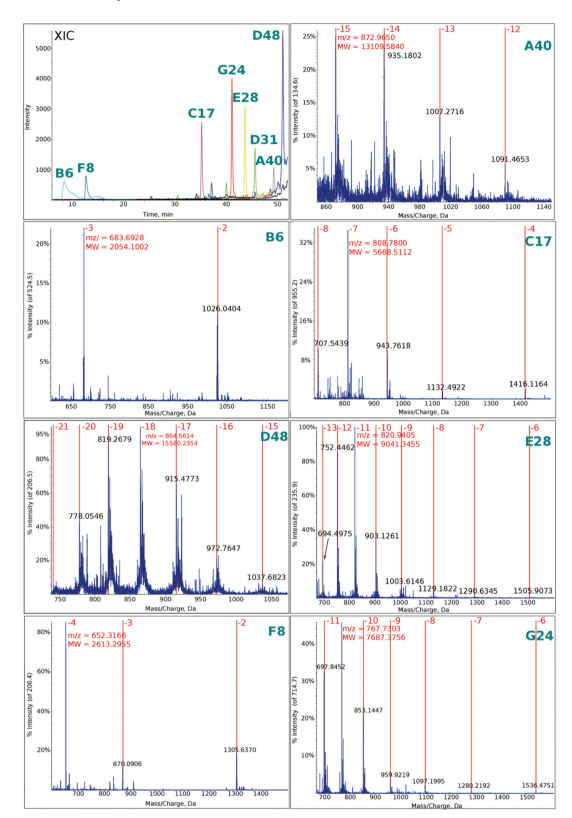


Fig. 4 Extracted ion chromatogram (XIC) and mass spectra of multiply charged ions (for fragment symbols, see Fig. 1) observed in the photolyte obtained by irradiation for 15 min.

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In Fig. 4 the ESI-MS spectra for the fragments denoted as in Fig. 1 are presented. These spectra were recorded for the photolyte irradiated for 15 min. Note that we were able to observe all the fragments that should be formed from the "hot spot" sequences shown in Fig. 1 (note that the deconvoluted masses agree very well with the calculated ones, see Table 1).

Additionally, in Fig. S4[†] a chromatogram employing the total ion current is demonstrated. Comparing the irradiated and non-irradiated samples, one observes a tail in the range of retention times where the particular fragments occur in the ion-extracted chromatogram (see XIC in Fig. 4).

Although the above-described LC-MS analysis unequivocally reveals the formation of the assumed DNA fragments, confirming that the photoinduced long range electron transfer is also operative in the double-stranded oligonucleotide labeled with BrdC, we decided to further corroborate these results by using denaturing PAGE. First, all the assumed fragments were synthesized chemically. Then, denaturing electrophoresis was carried out for a sample irradiated for 60 min. In Fig. 5, the

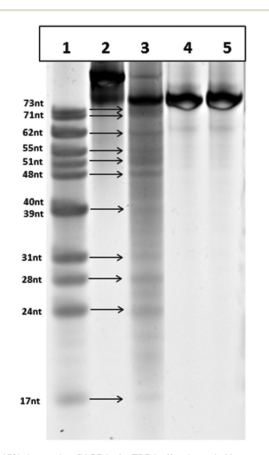


Fig. 5 15% denaturing PAGE in 1x TBE buffer. Lane 1: Mass standard an equimolar mixture of oligonucleotides that can occur if photoinduced single strand breaks concern the "hot spots" presented in Fig. 1 (for fragment labels see Fig. 1). Lane 2: Non-irradiated BrdC-labeled PCR fragment. Lane 3: BrdC-labeled PCR fragment irradiated for 60 minutes. Lane 4: Non-irradiated and non-labeled (not containing BrdC) PCR fragment. Lane 5: Non-labeled (not containing BrdC) PCR fragment irradiated for 60 minutes.

picture of the denaturing gel compares irradiated (lanes 3 and 5) and non-irradiated (lanes 2 and 4) samples to the mass standard, based on synthetic oligonucleotides (lane 1). For the irradiated, labeled sample, one can notice a series of bands that correspond pretty well to those originating from the synthetic oligonucleotides (*cf.* lane 3 to lane 1). These findings again confirm that the assumed photoinduced long-range ET does occur in the irradiated sample labeled with BrdC oligonucleotide, and leads to SSBs.

Yet another argument for the occurrence of the photoinduced ET process comes from the following reasoning. Strand breaks may occur in our system as a result of two mechanisms: (i) homolytic photodissociation of the C-Br bond in the excited BrdC or (ii) electron transfer to the excited halogenated nucleobase from a distant guanine. In both mechanisms, a reactive cytidine radical is formed that stabilizes, forming a strand break. It is easy to differentiate between these two mechanisms, since they lead to different products. In mechanism (i) the strand break is formed via the β-elimination of the phosphate group, which leads to two fragments - one with the 5'-phosphatae and another one without phosphate.²⁵ The sum length of these two fragments is equal to the length of the non-damaged PCR product, in our case, 80 bases. On the other hand, for mechanism (ii), ET goes via the formation of 2'-deoxyribonolactone, followed by the elimination of adenine. Ultimately, the thermal degradation of 2'-deoxyribonolactone leads to two fragments, one with 5'-phosphate and another one with 3'-phospahte.²⁶ The sum length of these two fragments is one base smaller than that of the original PCR product, and both fragments resulting from a strand break, contain ending phosphates. Since the sum length of pairs of fragments, A39/A40, B73/B6, C17/C62 etc., is one smaller than that of the PCR product, and both fragments end with a phosphate, one can conclude that the strand breaks observed in our system result from the long-range ET between a distant guanine and the photoexcited BrdC.

Enzymatic digestion coupled to the LC-MS analysis-intrastrand cross-links. Although in the cell SSBs are relatively easily repaired via the single-strand break repair (SSBR) pathway,²⁷ they may lead to lethal double-strand breaks (DSBs) in two cases: (i) if the SSB neighbor base is damaged on the opposite strand, or (ii) if the SSB occurs in the replication fork during the S phase of the cell cycle.²⁸ It was demonstrated that even a single unrepaired DSB is sufficient to kill a eukaryotic cell.²⁹ Besides, SSBs intrastrand cross-links (ICLs) were also observed in the photoirradiated oligonucleotides labeled with BrdU.³ Such damage if not repaired, may destabilize the DNA double helix,30 block DNA replication,31,32 as well as trigger mutations at or near the cross-link.33 The previous investigations on short oligonucleotides containing the BrdC residue showed the formation of various cross-links involving cytidine and purine moieties.^{9,11} On the other hand, the presence of pyrimidine in the neighborhood of BrdU leads to an ICL consisting of pyrimidines.³⁴ A short-range electron transfer between the excited BrdC and neighboring purine is assumed to be responsible for cytidine-purine intrastrand cross-

Therefore, one can expect ICL formation also in the studied DNA fragment because besides sequences leading to SSB, several sites are present where ET is forbidden and BrdC is adjacent to a purine or pyrimidine. Since ICLs introduce a relatively small modification to the total mass of a native oligonucleotide, they are formed by the release of two hydrogen atoms (a covalent bond is then formed between the appropriate ring atoms of neighboring nucleobases); the most effective way to enable investigations of such damage is to digest the irradiated oligonucleotide and employ the LC-MS analysis against the lysate afterwards. The cross-links are not processed by the enzymes and they are "cut out" from the oligonucleotide in the form of respective dimers, which can be relatively easily separated in an HPLC run from the ordinary nucleosides and then assayed by the MS analysis. In Fig. 6, the respective chromatogram obtained for the photolyte digested by the appropriate cocktail of enzymes along with the extracted ion chromatograms registered for ions of m/z equal to 553.2 and 513.2, respectively, are displayed. The first m/zratio corresponds to the d(G^C) dimer, while the second one corresponds to the d(C^C) adduct (for the ICLs structures see Fig. 7C). The HPLC signals of the identified dimers overlap with the dC and dG peaks, as indicated by the respective XIC chromatograms (cf. Fig. 6A-C); therefore, their presence is not evident from the analysis of the chromatogram

shown in Fig. 6A. In order to confirm the identity of the measured masses, we additionally registered their MS/MS spectra.

In Fig. 7, the fragmentation spectra for both m/z, 553.2 and 513.2, are demonstrated and Table 2 shows the possible origin of particular fragments. The fragmentation of the m/z 553.2 and 513.2 ions leads to the formation of fragments with m/zequal to 357.1, 259.1, 216.1, and 495.2, 415.2, 372.1, and 219.1 (see Fig. 7b). The fragment ion of m/z 495.2 can be attributed to the loss of a H₂O molecule from m/z 513.2. The m/z 357.1 ion is due to the concomitant loss of a C5H6O2 moiety (the 2'deoxyribose component) and the phosphate group from m/z553.2, while the ions of m/z 259.1 and 216.1 are related to the d(G^C) dimer of the nucleobase and the loss of the HNCO fragment from this dimer, respectively (see Table 2). On the other hand m/z, 415.1 and 372.1 correspond to the elimination of $C_5H_6O_2$, and $C_5H_6O_2$, together with H_3PO_4 components, respectively, from m/z 513.2, while the m/z 219.1 ion is consistent with the mass of two covalently bonded molecules of cytosine $d(C^C)$ (see Table 2).

Although the above-described MS analysis does not allow the particular isomers of the analyzed dimers to be identified, we are not able to differentiate dG[8-5]C from dC[5-8]G; in Fig. 7C, we assumed that the dG[8-5]C dimer, rather than dC [5-8]G, is formed under our experimental conditions. Analyzing the sequence shown in Fig. 1, one can note that the sites where BrdC is surrounded by two G, prevail over other configurations, leading to ICL. Moreover, the literature reports

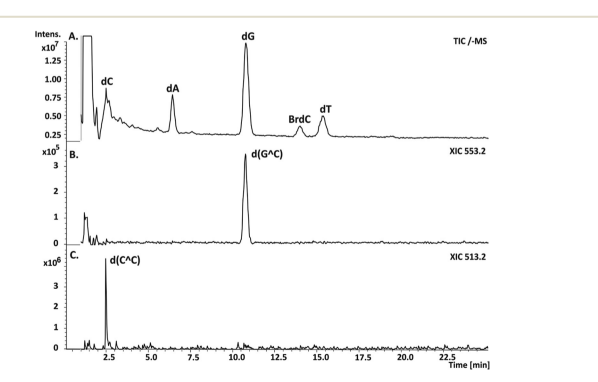


Fig. 6 LC-MS analysis of the DNA fragment (for the sequence, see Fig. 1) labeled with BrdC and irradiated for 60 min, after enzymatic digestion. (A) Chromatogram showing the total ion current (TIC) for the digested photolyte. (B) and (C) Extracted ion chromatograms registered for ions of m/z equal to 553.2 and 513.2, corresponding to d(G^C) and d(C^C) cross-links, respectively.

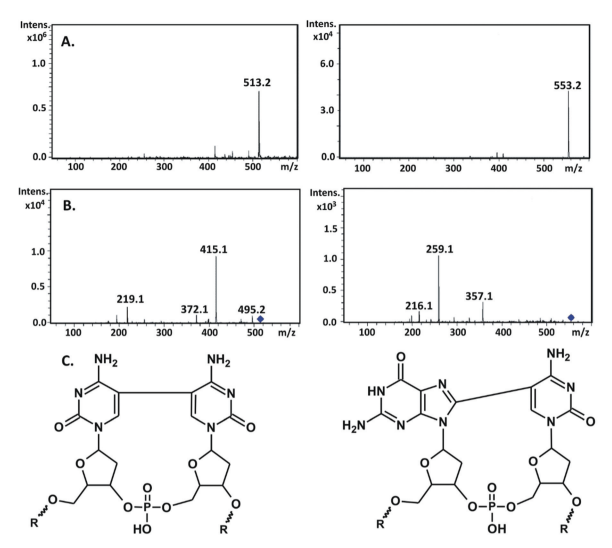


Fig. 7 (A) LC-MS analysis: negative-ion ESI-MS of the intrastrand cross-links. (B) MS/MS product-ion spectra showing the fragmentation of m/z 513.2 and m/z 553.2. (C) Proposed structures of the identified intrastrand cross-links: d(C^C) and d(G^C).

Table 2	Mass measurements of the $[M - H]^-$ ions and product ions
observed	in the MS/MS of the $[M - H]^-$ ions of d(C^C) and d(G^C)

m/z	Ion identities
d(C^C)	
513.2	$[M - H]^{-}$
495.2	-H ₂ O
415.1	-2'-Deoxyribose
372.1	-2'-Deoxyribose, -HNCO
219.1	[Nucleobases – H] ⁻
d(G^C)	
553.2	$[M - H]^{-}$
357.1	-2'-Deoxyribose, -phosphate
259.1	[Nucleobases – H] ⁻
216.1	[̈́Nucleobases – H]̈́−, -HNCO

suggest that the 5'-GBrC configuration is several times more reactive than the 5'-BrdCG.⁹ Here, it is also worth mentioning that different mechanisms are probably responsible for the formation of pyrimidine and purine dimers. Photochemical studies on DNA containing BrdC surrounded by two purines suggest electron transfer as being responsible for the ICL formation. An analogous study of sequences containing BrdU neighbored by pyrimidines at both sides indicates that the formation of the $d(U^U)$ dimer proceeds through a cyclobutapyrimidine intermediate.³⁴ This is probably the mechanism that also governs the formation of the $d(C^C)$ ICL detected in the current work. Taking into account the fact that the 5'-d(BrUU) sequence is more susceptible to photoirradiation than the 5'-d (UBrU) configuration, a similar directional preference can be expected for the formation of cytidine dimers studied in the present work.

Our attempts to identify, possibly at first glance, other ICLs like the various $d(C^A)$ dimers failed, which suggests that their quantum yield, if not zero, is so small that the amount of the produced damage falls below the limit of LC-MS detection. It is worth noting that the formation of the $d(G^C)$ dimers is

almost 20-fold more probable than the formation of the $d(A^{\Lambda}C).^9$

4. Conclusions

Using PCR, we synthesized 80 bp DNA fragments in which all cytidines but those present in primers were substituted with their brominated analogue, 5-bromo-2'-deoxycytidine. We demonstrated that the BrdC-labeled DNA is photoreactive, although its propensity for photoinduced damage is smaller than that for BrdU-labeled double-stranded oligonucleotides. Generally, two types of DNA damage were discovered in the UV-irradiated BrdC sensitized DNA: single strand breaks and intrastrand cross-links. Using the LC-MS method and denaturing PAGE, we showed the existence of seven "hot spots", sensitive to the photoinduced long range ET in the studied duplex. On the other hand, ICLs, $d(C^C)$ and $d(G^C)$, are formed if a BrdC molecule neighbors with another BrdC or dG nucleoside.

Both SSBs and ICLs pose serious damage to the DNA molecule, which, if not repaired, may lead to cell death. Therefore, due to the above-described properties, BrdC could be considered as an efficient photosensitizer of DNA damage, thus having potential clinical applications.

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