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Nicotinamide adenine dinucleotide (NAD) is a pivotal redox cofactor of primary metabolism. Its redox reactivity is based on the nicotinamide mononucleotide (NMN) moiety. We investigated whether NMN+ can engage in pairing interactions, when incorporated into an oligonucleotide. Here we describe the incorporation of NMN+ at the 3′-terminus of an oligodeoxynucleotide via a phosphoramidate coupling in solution. The stability of duplexes and triplexes with the NMN+-containing strand was measured in UV-melting curves. While the melting points of duplexes with different bases facing the nicotinamide were similar, triplex stabilities varied greatly between different base combinations, suggesting specific pairing. The most stable triplexes were found when a guanine and an adenine were facing the NMN+ residue. Their triplex melting points were higher than those of the corresponding triplexes with a thymidine residue at the same position. These results show that NMN+ residues can be recognized selectively in DNA helices and are thus compatible with the molecular recognition in nucleic acids.

Introduction

Nicotinamide adenine dinucleotide (NAD) is a pivotal cofactor in the cell. It consists of nicotinamide mononucleotide (NMN) and adenosine monophosphate, linked by a pyrophosphate bond. The reduced form of NAD+, NADH, is generated during glycolysis and the citric acid cycle, and is consumed in the respiratory chain. Further, NADH and its oxidized form, NAD+, are coenzymes for a host of oxidoreductase enzymes. Also, NAD+ is a substrate for the ADP-ribose transfer to enzymes. Moreover, NAD+ serves as a precursor of cyclic ADP-ribose, a second messenger. Bacterial DNA ligases consume the NAD+ when joining two DNA strands. Nicotinamide residues are also found as a cap for regulatory RNAs. Additionally, it is becoming increasingly clear that NAD+ and its phosphorylated form, nicotinamide adenine dinucleotide phosphate (NADP), have roles in signal transduction and in cellular defence systems. On the extracellular level NAD+ is assumed to act as a neurotransmitter.

It is not surprising that NAD+ is conserved in all kingdoms of life. Apparently, it originated early in evolution. Structurally, NAD+ consists of two ribonucleotides. This points to a possible origin in the so-called ‘RNA world’.

If so, it is likely that nature first used NAD+ in biomolecular complexes that formed, directed by base pairing. But while the adenosine monophosphate portion of NAD+ is well known to engage in hydrogen bonding interactions underlying duplex and triplex formation, it is not clear whether the NMN+ residue is able to engage in base pairing with other nucleobases in a duplex or triplex context (Fig. 1). If it does, this would be most useful for the design of novel functional DNA nanostructures with tailor-made active sites.

We therefore sought a methodology for incorporating nicotinamide mononucleotide residues in oligonucleotides. Liu and Orgel had previously attempted to incorporate NAD+ residues at the terminus of primers terminating in a ribonucleoside. Enzymatic extension of oligoribonucleotides and
untemplated oligomerization was achieved, using di- or triphosphates of pyridine nucleosides. Attempts to induce in vitro transcription or enzyme-free primer extension were unsuccessful, though. The authors noted that NMN should, in principle, be able to undergo base pairing, but because both enzymatic and enzyme-free template-directed incorporation of NMN failed, they concluded that NMN does not act like an ‘informational unit of a polymer’.

More recently, Mutschler and Holliger described a ribozyme that catalyzes addition of nicotinamide adenine dinucleotide 2′,3′-cyclic phosphate to the 5′-terminus of RNA strands through a nucleotidyl transferase activity, but the resulting constructs contained the NMN residue in the pyrophosphate-linked form, not as part of a continuous phosphodiester chain. Here we report a methodology for incorporating NMN residues at the 3′-terminus of DNA strands via a phosphoramidate linkage, together with binding data that show what base combinations best accommodate the NMN residue in a double helix or triplex. The challenge to link the labile NMN substructure to a synthetic oligonucleotide sequence was met by converting the NMN residue into an active ester and subsequent coupling to the aminoterminal oligonucleotide in aqueous solution.

Results

The chemical synthesis of oligodeoxynucleotides with an NAD+ residue is complicated due to the lability of the pyridine nucleotides. In the cationic, oxidized form, lability to a base is a significant problem, while the reduced form reacts quickly under acidic conditions. This was confirmed in exploratory reactions involving NADH, NAD+, and their pyrophosphate cleavage products. The labilities precluded a synthetic approach involving conventional chain assembly by solid-phase DNA synthesis, which employs both acidic and strongly basic conditions. Further, it was not immediately clear how a phosphoramidite, H-phosphonate, or a phosphodiester building block could be generated, starting from commercially available starting materials. This prompted us to study the conversion of fully deprotected NMN under conditions established for enzyme-free primer extension.

Liu and Orgel had reported the synthesis of a methylimidazolide of NMN using a redox condensation in DMSO, but no characterization of the product was given. This was also true for the oligomerization products of their enzymatically catalyzed conversions that gave electrophoretic bands contaminated with broad smears that the authors attributed to the partial loss of the nicotinamide from the nucleotide under the basic conditions employed. When we prepared the methylimidazolide of NMN using conditions optimized for ribonucleotides, we observed the expected signal by ESI mass spectrometric analysis of the precipitated product (Fig. S1, ESif). Attempts to bring about a coupling reaction with a primer were unsuccessful, though, both with an RNA primer, and with a more reactive DNA primer with a 3′-terminal, 3′-amino-2′,3′-dideoxynucleoside. The latter gives a phosphoramidate linkage that is isoelectronic and largely isosteric to natural phosphodiester linkages. The aminoterminal primer had been prepared by solid-phase synthesis, starting from controlled pore glass loaded with the aminonucleoside.

Scheme 1 shows the successful synthetic approach. It starts with the activation of NMN+ to the oxyazabenzotriazolide (OAt ester) I, using hydroxyazabenzotriazolide (HOAt) and N-ethyl-N′-dimethylaminopropyl carbodiimide (EDC) as the condensation agent in aqueous solution at pH 5.5. The formation of the active ester was confirmed by 31P-NMR and UV-Vis spectroscopy (Fig. S2 and S3, ESIf). An aliquot of the solution of I was then added to a solution of aminoterminal strand 2 in primer extension buffer containing MgCl2 and NaCl at pH 8.9. The concentration of the buffer was increased to 0.5 M to ensure a proper pH shift from that of the activation mixture and thus high-yielding reactions. Typically, 0.1 M OAt ester I was used, and the reaction was allowed to proceed for 48 h at 20 °C. As expected, the isolation and purification of NMN-containing oligonucleotide 3 proved far from trivial. While gel permeation chromatography or affinity chromatography with an immobilized complementary strand to 3 and subsequent dialysis gave unsatisfactory results, anion exchange HPLC, followed by desalting on a C4-phase gave a solution showing pure 3 (Fig. 2). This solution was evaporated to dryness without decomposition by using a gentle stream of nitrogen.
With NMN\(^+\)-containing oligonucleotide 3 in hand, we then proceeded toward studying the molecular recognition of the nicotinamide mononucleotide in duplex and triplex contexts. For this, the four-strand triplex shown in Fig. 3 was designed. Besides 3 or unmodified T-terminated control strand 4, it features a long strand (5a–t) engaging in Watson–Crick base pairing, a triplex-forming oligonucleotide (TFO, 6a–t) that bridges the two duplexes, and the purine-rich central strands (3/4 and 7).

Three subsequent melting transitions were expected for the triplexes (Fig. 4a). The first releases the TFO from the duplexes, the second involves the dissociation of the short purine-rich strand, and the third transition leads to the dissociation of the remaining long duplex. Fig. 4b shows that all three transitions are indeed observable experimentally. This allowed us to screen for pairing interactions with the nicotinamide residue, both in a duplex and in a triplex environment, by offering each of the four different nucleobases (A/C/G/T) in either of the two pairing strands (5 and 6) and measuring the thermal stability of the resulting complexes. The third melting transition (\(T_{m3}\), Fig. 4a) should be largely unaffected by the sixteen possible base combinations of 5a–t and 6a–t, thus serving as an internal control. This was indeed the case, with a \(T_{m3}\) of 66 ± 1 °C in all cases.

Table 1 lists the \(T_{m2}\) and \(T_{m1}\) values determined in our screening process. We measured at least two melting curves for each base combination and analyzed the transitions with two different methods (first derivative of numerical values, or graphical determination of \(T_{m}\) as the point at which 50% of the hyperchromicity is reached). The duplex melting points for the thermal release of 3 were very similar for each of the four possible bases in 5a–t, ranging from 39.4 to 42.5 °C. This suggested that the NMN\(^+\) residue does not engage in highly selective pairing in a duplex context.

However, triplex stabilities varied greatly with the nucleobases in the pairing strands (5/6). Here, the lowest melting...
measured a series of melting curves with control strand 4.

triplet interactions between the two purines, we then

20 °C. The triplexes of adenine, guanine and NMN+ is due to non-specific or non-

duplex transition is shifted for a stabilized 0.4 °C were obtained. Fig. 5 shows how much the triplex-to-

Expansion of melting curves showing the triplex-to-duplex Fig. 5

<table>
<thead>
<tr>
<th>Partner strands</th>
<th>Triplet</th>
<th>$T_m$ (d)°C</th>
<th>$T_m$ (t)°C</th>
<th>Hyperchromicity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5a, 6a</td>
<td>A:N:C</td>
<td>40.6 ± 0.5</td>
<td>14.3 ± 0.8</td>
<td>6.9 ± 0.8</td>
</tr>
<tr>
<td>5a, 6c</td>
<td>A:N:C</td>
<td>40.7 ± 0.3</td>
<td>10.4 ± 0.5</td>
<td>6.8 ± 0.3</td>
</tr>
<tr>
<td>5a, 6g</td>
<td>A:N:C</td>
<td>41.7 ± 0.3</td>
<td>19.9 ± 0.1</td>
<td>7.6 ± 0.2</td>
</tr>
<tr>
<td>5a, 6t</td>
<td>A:N:C</td>
<td>40.7 ± 0.3</td>
<td>12.7 ± 0.4</td>
<td>6.6 ± 0.9</td>
</tr>
<tr>
<td>5c, 6a</td>
<td>C:N:T</td>
<td>40.9 ± 0.9</td>
<td>14.0 ± 0.5</td>
<td>5.2 ± 0.7</td>
</tr>
<tr>
<td>5c, 6c</td>
<td>C:N:T</td>
<td>41.9 ± 0.4</td>
<td>14.2</td>
<td>6.5</td>
</tr>
<tr>
<td>5c, 6g</td>
<td>C:N:T</td>
<td>42.5 ± 0.4</td>
<td>&lt;10°</td>
<td>&lt;5°</td>
</tr>
<tr>
<td>5c, 6t</td>
<td>C:N:T</td>
<td>41.9 ± 0.3</td>
<td>12.8</td>
<td>6.0 ± 0.9</td>
</tr>
<tr>
<td>5g, 6a</td>
<td>G:N:T</td>
<td>40.3 ± 0.3</td>
<td>18.5 ± 0.2</td>
<td>10.3 ± 0.1</td>
</tr>
<tr>
<td>5g, 6c</td>
<td>G:N:T</td>
<td>39.4 ± 0.4</td>
<td>11.5 ± 0.6</td>
<td>6.4 ± 1.0</td>
</tr>
<tr>
<td>5g, 6g</td>
<td>G:N:T</td>
<td>40.1 ± 0.2</td>
<td>14.5 ± 0.2</td>
<td>8.0 ± 0.0</td>
</tr>
<tr>
<td>5g, 6t</td>
<td>G:N:T</td>
<td>40.4 ± 0.3</td>
<td>12.7</td>
<td>5.7 ± 0.2</td>
</tr>
<tr>
<td>5t, 6a</td>
<td>T:N:C</td>
<td>42.1 ± 0.1</td>
<td>15.6 ± 0.6</td>
<td>5.0 ± 0.6</td>
</tr>
<tr>
<td>5t, 6c</td>
<td>T:N:C</td>
<td>40.6 ± 0.0</td>
<td>(10.2 ± 0.7)</td>
<td>&lt;5°</td>
</tr>
<tr>
<td>5t, 6g</td>
<td>T:N:C</td>
<td>41.7 ± 0.3</td>
<td>11.5</td>
<td>5.0</td>
</tr>
<tr>
<td>5t, 6t</td>
<td>T:N:C</td>
<td>41.4 ± 0.5</td>
<td>(10.1)</td>
<td>5.0</td>
</tr>
</tbody>
</table>

$^a$ Conditions: 1 μM oligonucleotides, each, 10 mM phosphate buffer, pH = 7.0, 1 M NaCl, 10 mM MgCl2; melting points are averages of at least two measurements. All duplex-to-single strand transitions of the remaining long duplex (5a-t:7) were within 0.3 °C of 65.1 °C. $^b$ ±1 standard deviation when ≥ three data points were available. $^c$ Melting transition not sigmodal.

Table 1 UV-melting results for triplexes of strand 3 with Watson–Crick partner strands 5a, 5c, 5g or 5t and Hoogsteen partner strands 6a, 6c, 6g or 6t, in the presence of downstream-binding strand 7.

Fig. 5 Expansion of melting curves showing the triplex-to-duplex melting curves for a weakly stabilized triplex (squares, strands 5t, 3, 6c and 7) and a strongly stabilized triplex (triangles, strands 5a, 3, 6g and 7).

point was below 10 °C, and the highest $T_m$1 was just below 20 °C. The triplexes of 3 with 5a/6g and 5g/6a were significantly more stable than other triplexes. This was confirmed by the melting points obtained by the graphical data analysis mode (Table S1, ESI†), where $T_m$1's of 19.5 ± 0.2 °C and 19.6 ± 0.4 °C were obtained. Fig. 5 shows how much the triplex-to-
duplex transition is shifted for a stabilized versus a non-stabilized triplex.

To exclude that the high stability of the triplexes with adenine, guanine and NMN+ is due to non-specific or non-triplet interactions between the two purines, we then measured a series of melting curves with control strand 4 replacing 3 in the triplex. We chose seven sequences, focusing on the base combination with a purine in at least one of the pairing positions. The thymine residue of 4 is similar in size and shape to the nicotinamide residue of 3 but offers very different hydrogen bonding functionalities. Table 2 lists the results from these experiments.

As expected, the thymidine residue of 4 pairs sequence-
specifically with the A of 5a, following the base-pairing rules of Watson and Crick. Mismatched bases in 5 (C, G or T) lead to a depression in $T_m$2 of 7.1–8.9 °C (Table 2). However, neither of the four possible bases of 6a-t gave melting points as high as those found for triplexes with 3 and the combinations of A and G as pairing partners. This was confirmed by the results of the graphical data analysis method, as detailed in Table S2 of the ESI.† Here, the melting points of the most stable triplexes with 4 gave $T_m$1's of 15.9 ± 0.3 and 16.8 ± 0.1 °C.

Discussion

Thus far, it was assumed that nicotinamide mononucleotide residues do not act as base pairing partners when incorporated into nucleic acid chains.24 Our results now show that a combination of adenine and guanine as pairing partners for an NMN+ residue gives specific and surprisingly stable triplexes (Table 1). It is therefore interesting to ask what the interactions may be that lead to these stabilized triplexes. Two different hydrogen bonding patterns present themselves as likely inter-
action modes with the NMN+ at the center position of a triplet. One of these patterns features two hydrogen bonds with guanine on the Watson–Crick side of the pairing (Fig. 6a). It involves the so-called cis-conformation of the NMN+ that places the carbonyl group of the amide moiety towards the C2 carbon of the pyridinium ring.25 Computational work and crystal structures of nicotinamides show that this is the pre-
ferred conformation of free NAD+.26,27 The second pairing pattern is shown in Fig. 6b. It involves the trans-conformation of the amide of the NMN which has been described as the main conformation of NAD+ in complexes with proteins.28 Both con-
formers are close enough in energy to make it likely that they are populated to a significant extent.\textsuperscript{27,28} In the second pairing arrangement, adenine is found on the Watson–Crick side of the triplet, whereas guanine is the base facing the nicotinamide in the triplex-forming oligonucleotide.

We note that a single hydrogen bond may not provide a decisive directing force and that shape complementarity and stacking interactions with neighboring bases in the same strand may be important factors for pairing. They cannot be predicted as easily as hydrogen bonding, though. Elucidating them will require a detailed three-dimensional structure, to be predicted as easily as hydrogen bonding, though. Elucidating the Watson–Crick-pairing strand of the triplex, whereas the base of the triplex-forming strand engages in no more than a single hydrogen bond. Alternative pairing arrangements may exist for conformers of \( \text{NMN}^{\text{+}} \) that are resulting from a 180° rotation about the glycosidic bond.

Conclusions

Independent of what the finer structural details of the pairing interactions are at the site of the \( \text{NMN}^{\text{+}} \) in the triplex, it is now clear that a preferred sequence contexts exist for accommodating \( \text{NMN}^{\text{+}} \) in a helical DNA structure. This is important as \( \text{NMN}^{\text{+}} \) is the reactive part of \( \text{NAD}^{\text{+}} \), a pivotal redox cofactor in the metabolism of cells. Together with the synthetic methodology to covalently incorporate nicotinamide mononucleotide residues into oligonucleotides, our results may thus help to design new functional DNA structures, including structures with catalytic activity that rely on \( \text{NMN}^{\text{+}} \) residues as prosthetic groups.

Experimental

General

Nicotinamide mononucleotide was obtained from Sigma-Aldrich (Deisenhofen, Germany). Both HOAt and EDC were purchased from Carbolution (Saarbrücken, Germany). All chemicals used for automated solid-phase DNA synthesis were from Proligo (Hamburg, Germany), including the phosphoramidite building blocks of deoxynucleosides, DCI activator (0.25 M 4,5-dicyanoimidazol in acetonitrile), amidite diluent (acetonitrile, \( \leq 90 \) ppm water), tetrazole activator solution (0.45 M in abs. acetonitrile), cap B (THF/pyridine/1-methylimidazole 80 : 10 : 10 v/v/v), oxidizer solution (0.1 M iodine in THF/pyridine/water 77 : 21 : 2, v/v/v), and deblock solution (trichloroacetic acid in CH\(_3\)Cl; 3 : 97 v/v). Controlled-pore glass loaded with 5′-protected 3′-amino-2′,3′-dideoxyadenosine, linked with a hexafluoroglutaric acid linker (pore size 1000 Å, 25 µmol g\(^{-1}\) loading) was prepared as reported.\textsuperscript{23} Yields of oligonucleotides were determined by UV-absorption at 260 nm. MALDI-TOF mass spectra were acquired on a Bruker Reflex IV spectrometer in linear negative mode using 0.3 M 2,4,6-trihydroxyacetophenone monohydrate as a matrix and 0.1 M diammonium citrate in water as a co-matrix solution.

Oxyazabenzotriazolide of \( \text{NMN}^{\text{+}} \) (1). To obtain a 0.2 M stock solution of the activated nicotinamide mononucleotide 1, \( \text{NMN}^{\text{+}} \) (2.7 mg, 8.0 µmol, 1 eq.), EDC hydrochloride (7.7 mg, 40.0 µmol, 5 eq.) and HOAt (3.3 mg, 24.0 µmol, 3 eq.) were dissolved in H\(_2\)O (40 µL). Then, NaOH solution (1 M, 2 µL) was added to adjust the pH to 5.5. The reaction mixture was vortexed for 1 h. \textsuperscript{31}P-NMR and UV-Vis showed conversion to the desired oxyazabenzotriazolide. \textsuperscript{31}P-NMR (202 MHz, D\(_2\)O); \( \delta = -1.1 \) ppm; UV, \( \lambda_{\text{max}} = 260 \) nm. The resulting solution was either used immediately for extension of 2 or frozen in liquid nitrogen for later use.

3′-Aminoterminal oligonucleotide 2. The 3′-aminoterminal strand 2 was synthesized on an 8909 Expedite synthesizer according to a published protocol\textsuperscript{22} on the 1 µmol scale, using 40 mg of controlled pore glass with a nucleoside loading of 0.025 µmol mg\(^{-1}\). At the end of the chain assembly, the 3′-aminoterminal strand was deprotected and cleaved from the solid support by treatment with aqueous ammonia (25%) at 60 °C for 20 h. Residual ammonia was removed with a gentle stream of nitrogen for 20 min, and the resulting aqueous solution was reduced to dryness by lyophilization. The crude product was

![Fig. 6 Possible hydrogen bonding interactions between the nicotinamide residue (\( \text{N}^{\text{+}} \)) and adenine (A) and guanine (G) as pairing partners.](image-url)
dissolved in 0.1 M TEAA buffer, pH 7.0, and any residual solid was removed by filtration. The DMT-bearing oligonucleotide was then purified by reversed-phase HPLC, followed by cartridge purification with detritylation in the adsorbed state.

5′-DMT-GCCAGAGAA-NH2-3′. HPLC (Rainin Dynamax system, with Macherey–Nagel EC 250 × 4.6 mm Nucleosil 120-5 C4 column) and a gradient of acetonitrile in 0.1 M triethylammonium acetate buffer (TEAA), pH 7.0, starting from 0% CH3CN for 5 min, then from 0% to 30% CH3CN in 45 min, and then 30–80% in 5 min, tR = 23 min. MALDI-TOF MS, calc. for C109H227N45O54P9 [M – H]– 3057.3 (average mass), found 3056.4 (maximum of unresolved isotope pattern). Product-containing fractions were lyophilized to dryness.

5′-GCCAGAGAA-NH2-3′ (2). The deprotection meaning purification was performed on an RP C-18 cartridge from Waters (Milford, USA). A solution of the product in 0.1 M TEAA was loaded onto the cartridge, followed by washing with brine, and eluting with water/acetonitrile (9:1, v/v). Oligonucleotide 2 eluted at 10% acetonitrile; overall yield 10% (based on the loading of the support), MALDI-TOF MS, calc. for C68H109N42O47P8 [M – H]– 2753.5 (average mass), found 2752.7 (maximum of unresolved isotope pattern).

5′-GCCAGAGAA-NH2-3′ (3). The extension of 3′-aminoterminal 2 was carried out in a final volume of 10 µL. A solution of strand 2 (2 mM in 10 µL, 20 nmol) was lyophilized to dryness and then dissolved in HEPES buffer (50 µL, 1 M, 800 mM NaCl, 160 mM MgCl2, pH 8.9), and treated with an aliquot of the aqueous stock solution of OAt-NMN+ (50 µL) to avoid evaporation. Analitical samples (0.5 µL) were diluted with water (20-fold) and desalted over Dowex cation exchange resin (ammonium form). Full conversion was observed after 48 h, as monitored by MALDI-TOF mass spectrometry. The elongated DNA strand 3 was then purified by a two-step process, starting with ion exchange HPLC, and followed by RP-HPLC. First, the reaction mixture was subjected to ion-exchange HPLC (Waters 1525 series pumps and 2489 UV/Vis detector; Macherey–Nagel Nucleogel SAX 1000-8 column) and eluted with a gradient of 0% to 20% CH3CN in 50 min, 20% to 80% in 10 min, tR = 28 min. Yield: 12%, MALDI-TOF MS, calc. for C99H122N44O547P9 [M – H]– 3070.4 (maximum of unresolved isotope pattern).

UV-melting curves

The UV-melting curves were measured on a Lambda 25 spectrophotometer (Perkin Elmer) with Peltier-based temperature control, using Suprasil precision microcuvettes (Hellma, Müllheim, Germany) and sample volumes of 80 µL, with a protective layer of liquid paraffin (50 µL) to avoid evaporation. Control experiments without the protective layer showed identical melting points. Absorption was measured at 260 nm in the temperature range of 3–85 °C with a heating or cooling rate of 1 °C min−1. Melting points are extrema of the first derivative of heating or cooling curves, determined numerically, using UV-WinLab software. In some instances, even though the overall appearance of a curve was similar to other melting curves of the same sample, the transition was not sigmoidal enough to allow for unambiguous determination of the lowest melting point. If so, the number of available data points was sometimes insufficient to calculate an accurate standard deviation and an average value is being reported instead. Melting points, determined graphically as the temperature at which 50% of the hyperchromicity of the triplex-to-duplex transition was reached, are given in Tables S1 and S2 in the ESL†.

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Notes and references