Canonical nucleosides can be utilized by T4 DNA ligase as universal template bases at ligation junctions

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ABSTRACT
T4 DNA ligase catalyzes the template-dependent ligation of DNA. Using T4 DNA ligase under specific experimental conditions, we demonstrate that each of the four canonical nucleosides, centrally located on a template molecule such that they flank the site of ligation, can direct the ligation of nucleic acids regardless of the identity of the terminal nucleosides being covalently joined. This universal templating capability extends to those positions adjacent to the ligation junction. This is the first report, irrespective of the ligation method used or the identity of the template nucleosides (including analogs), which shows that nucleosides can act essentially as universal templates at ligation junctions in vitro. The canonical nucleosides do, however, differ in their ability to template sequence-independent ligations, with thymidine and guanosine being equally effective, yet more effective than adenosine and cytidine. Results indicate that hybridization strength surrounding the ligation junction is an important factor. The implications of this previously undiscovered property of T4 DNA ligase with canonical nucleosides are discussed.

INTRODUCTION
DNA ligases catalyze the ligation of an acceptor oligonucleotide, which contains a required 3'-OH group, and a donor oligonucleotide, which contains a required 5'-phosphate (Fig. 1), both of which typically are base paired to a nucleic acid template (1–4). Ligase is an essential enzyme in vivo, playing a critical role in recombination, replication and excision repair. High fidelity in these reactions is important for maintaining genetic integrity (5). For example, ligation of non-base-paired segments in replication would lead to mutagenesis. Ligase is also an essential enzyme in a wide variety of in vitro biotechniques, including standard cloning protocols, the ligase chain reaction (LCR) (6) and the ligation detection reaction (LDR) (7–9). High fidelity in these reactions is important for experimental accuracy.

Ligases with low fidelity could also be useful for certain experimental strategies, including ligating together degenerate or unknown oligonucleotides for purposes such as site-directed mutagenesis or sequence identification (10,11). To this end, various ligases have been studied under a wide variety of conditions in order to analyze the fidelity of these ligation reactions. The results show that some ligases have the ability, at least in vitro, to ligate oligonucleotides that form certain mismatched base pairs with their templates, both at the ligation junction and at other positions in the resultant helices (9,10,12–22). Under no circumstances, however, has a ligase been shown to catalyze ligation reactions in a sequence-independent manner at ligation junctions, which would be required for developing new types of sequence-independent experimental strategies. Moreover, chemical methods of template-dependent oligonucleotide ligation have been developed (23–25) and have not been shown to be any more effective at catalyzing low fidelity ligation reactions (24–28).

Another route that is being actively pursued is the development of nucleoside analogs that could act as universal bases (29). By definition, universal bases should be able to bind each of the four canonical nucleosides equally, although not necessarily tightly. In this way, ligases (or chemical methods) would be expected to exploit a universal base template region at ligation junctions to catalyze sequence-independent ligations. Perhaps surprisingly, under no circumstances have known universal base analogs been shown to be effective in these low fidelity reactions, especially at ligation junctions (20,30). Moreover, nucleoside analogs are not likely to be practical for subsequent work-up experiments that rely on standard molecular biology protocols.

It is well known that the properties of enzymes depend on the experimental conditions under which reactions are conducted. With this in mind, we evaluated whether a DNA ligase, under a definable set of reaction conditions, could catalyze sequence-independent ligation reactions. For this study, we chose to evaluate T4 DNA ligase, as it can ligate oligonucleotides that contain at least some mismatches (9–13,15,17), in addition to it being one of the most useful enzymes in nucleic acids research. In addition, we analyzed whether canonical nucleosides could act essentially as universal bases in the template at the ligation junctions.
junction (i.e. promoting sequence-independent ligation with T4 DNA ligase).

We now report that template-directed ligation reactions can be catalyzed in an essentially sequence-independent manner at ligation junctions. For this we exploit reaction conditions under which T4 DNA ligase becomes an extremely low fidelity enzyme. Moreover, we show that each of the four canonical nucleosides can act essentially as universal template bases at and adjacent to the ligation junction. The canonical nucleosides do, however, differ in their ability to template sequence-independent ligations, with thymidine and guanosine being equally effective, yet more effective than adenosine and cytidine. These results suggest that stable base pairing at and adjacent to the ligation junction is not fundamentally required for T4 DNA ligase activity. Hybridization strength surrounding the ligation junction, however, does appear to be important for effective ligation. The implications of this previously undiscovered property of T4 DNA ligase with canonical nucleosides are discussed.

### MATERIALS AND METHODS

#### Oligonucleotide synthesis and preparation

Oligonucleotides were synthesized either on an Applied Biosystems 380B DNA Synthesizer or purchased from Integrated DNA Technologies (Coralville, IA). Oligonucleotides were purified either by reverse phase HPLC (Integrated DNA Technologies) or thin layer chromatography, as described (31–33). Designated oligonucleotides were 5'-end radiolabeled by incubating 0.4 μM DNA, 0.43 μM [γ-32P]rATP (Amersham Pharmacia, Piscataway, NJ), 70 mM Tris-HCl (pH 7.5), 50 mM KCl, 10 mM MgCl2, 5 mM dithiothreitol (DTT), 0.1 mM EDTA, 0.1 μM rATP, 10% glycerol and 10 U T4 polynucleotide kinase (New England Biolabs, Beverly, MA) in a reaction volume of 10 μl for 45 min at 37°C. The products were isolated and purified via a 17% native polyacrylamide gel, using 0.25× TBE as the running buffer (1× TBE is 100 mM Tris, 90 mM boric acid and 1 mM EDTA, pH 8.4). The products were then isolated from the gel

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**Figure 1.** Ligating two decamers using DNA ligase and templates containing tandem canonical bases at the site of ligation. (Left) Studies on only the n positions, showing both catalytic steps. (Right) Studies on the n-1 positions, which also consist of experiments where the n positions are not base paired. The phosphorylated (represented by p) decamers are radiolabeled (represented by an asterisk) and are in italics. The names of the decamers to be ligated are represented by lower case lettering according to what nucleoside (from 5' to 3') is paired with the universal template positions (in the gray box). The template names are in upper case and represent the central universal template positions (underlined). Note that the adenylation step consists of a 5'→5' phosphate bond between the phosphorylated decamer and an AMP cofactor. Unless otherwise noted, all oligonucleotides are written in the 5'→3' direction. Be aware that, although we have no evidence to the contrary, hybridization of each oligonucleotide to the template might not be precisely as drawn here.
matrix via overnight pulverization in 2 ml sterile water with a stir bar, followed by centrifugation to partition the gel matrix from the solution. The radiolabeled product solutions were then evaporated to a final concentration of 8 nM.

Ligation reactions
All ligation reactions consist of 3.125 pmol template, 2.5 pmol non-radiolabeled oligonucleotide (the acceptor molecule), 70 fmol radiolabeled oligonucleotide (the donor molecule) and 3 U T4 DNA ligase (Promega, Madison, WI). In all ligation reactions the template and donor molecules were mixed in an appropriate buffer solution and denatured at 90°C for 30 s. After a 5 min annealing period at room temperature, the ligase and the acceptor molecules were added to the reaction mixture (10 μl total volume). This was then incubated for 18 h at an appropriate temperature (listed in the figures and tables). The reactions were terminated by adding 7 μl stop buffer (10 mM urea, 0.1× TBE, 3 mM EDTA). Initial reactions were conducted under Promega’s standard recommended conditions, which consist of 30 mM Tris–HCl (pH 7.8), 10 mM DTT, 10 mM MgCl2 and 1 mM rATP at 30°C. These conditions were then optimized to allow for universal templating abilities by altering the MgCl2 concentration (varied from 1 to 100 mM), the ATP concentration (varied from 0.25 μM to 10 mM), temperature (varied from 4 to 44°C) and by including additives such as dimethyl sulfoxide (DMSO), polyethylene-glycol 8000 (PEG) and single-stranded binding protein (from 5 to 50%). The final optimized buffer conditions were 30 mM Tris–HCl (pH 7.5), 10 mM DTT, 3 mM MgCl2, 10 μM ATP and 20% DMSO at either 30 or 22°C for 18 h. In certain cases, 1 U calf intestinal alkaline phosphatase (Promega) was added to individual reactions. Reaction products and intermediates were isolated using 11% denaturing polyacrylamide gels and the reactions were quantified using a Storm 860 Molecular Dynamics PhosphorImager. Observed rate constants were quantified by fitting the time-dependent data to a simple exponential function.

RESULTS
Experimental strategy
To test whether canonical nucleosides, in the template at the ligation junction, can direct the ligation of oligonucleotides irrespective of the identity of the terminal nucleosides being ligated, we developed a model system that allows the quantification of oligonucleotide ligation as a function of the sequence of the oligonucleotides being ligated. A schematic representation of this experimental strategy is shown in Figure 1. The oligonucleotides to be ligated are decamers, each of which begin and end with the same nucleoside. The acceptor oligonucleotides to be ligated are decamers, each of which begin and end with the same nucleoside. The acceptor molecules contain a 3′-OH group and the donor molecules contain a radiolabeled 5′-phosphate. When the decamers bind to a template, T4 DNA ligase catalyzes the formation of a covalent bond between the 5′-phosphate and 3′-OH group (example in Fig. 1). By phosphorylating the 5′-end of only one of the decamers (the donor molecule) and adding an excess of unphosphorylated decamer (the acceptor molecule), we can control the sequence of the oligonucleotides at the ligation junction to be ligated. The templates contain a centrally located region consisting of a single type of nucleoside in tandem, which will flank the ligation junction when bound to the donor and acceptor molecules. In this way, we can analyze the ability of the ligase to ligate particular oligonucleotides as a function of template sequence. In this report we focus on the sequences immediately flanking the ligation junction (the n positions in Fig. 1, left) and their adjacent positions (the n – 1 positions in Fig. 1, right).

Investigation of the n position: standard conditions
We analyzed the 16 dinucleotide n position combinations using four different templates with either A, C, T or G as the templating nucleoside at the site of ligation. For this we utilized T4 DNA ligase (Promega) under the standard manufacturer’s conditions. We found that the ratio of 1/0.8/0.02 template, non-radiolabeled acceptor and radiolabeled donor, respectively, yielded the best results (data not shown). A representative gel using the T2 template is shown in Figure 2. All experimental indications are that the assays used to study the ligation reactions are working correctly, as the numbers and sizes of the products are as expected. One unique ligation product stems from the ligation of (c + ρα). With each of the four templates, this product runs noticeably lower than the other ligation products (Fig. 2). A synthetic size control of the (c + ρα) product was run on the gel and was found to also migrate faster than other products. Evidently, this specific sequence is an anomaly as far as its migration behavior. Note that the (a + ρα) and (c + ρα) size controls run marginally faster than the actual ligation products, which is due to the presence of 5′-phosphates in the controls that are absent in the actual ligation products. A further indication that the assays are working correctly is that formation of both products and intermediates are template dependent. In addition, dephosphorylation of the product band results in a minor (if any) reduction in its intensity, which is expected if the product is

Figure 2. Representative gel of ligation reactions run under standard conditions using the T2 template. The standard reaction conditions are 30 mM Tris–HCl (pH 7.5), 10 mM DTT, 10 mM MgCl2, 1 mM ATP, 2.5 pmol acceptor molecule, 70 fmol donor molecule, 3.125 pmol T2 template and 3 U T4 DNA ligase at 30°C for 18 h. P represents product, I represents adenylylated intermediate and S represents starting material. The reaction components are shown above each lane. The first lane contains a generic 25 base size control, the next four lanes are the four unreacted donor molecules and the next two lanes are the synthetic (a + ρα) and (c + ρα) size controls. The next four panels show the 16 possible reactions. The next three lanes show the ligation of (a + ρα) without template, with template and treated with calf alkaline intestinal phosphatase after the reaction has gone to completion, and without ligase.
only internally radiolabeled. In other words, we are not ligating together two radiolabeled donor molecules.

Table 1 lists the results of duplicate trials using each of the four templates. The results show that in cases of complete complementarity, the ligation extent is 95% or more, as would be expected. In cases where one mismatch occurs between the universal template region and the ligation junction, most ligations react greater than 80% (the few exceptions still reacting over 45%). In cases where no complementary base pairing occurs at the ligation junction, the results vary widely, with some dinucleotide combinations being ineffective [for example, 11% for (a-pG)/A2] while some are quite effective [for example, 77% for (c-pA)/A2]. Nevertheless, each of the canonical nucleosides display at least some ability to template the ligation of all 16 dinucleotide combinations. As simply judged by the average product generated as a function of template, the effectiveness of each template reacting with each of the 16 possible sequence combinations follows the order G2 (63%) = T2 (62%) = A2 (58%) > C2 (47%).

Investigation of the α position: optimized conditions

The previous results show that under standard conditions T4 DNA ligase is a sequence-dependent enzyme. Where product formation was low, however, adenylated intermediate formation was usually substantial. Therefore, to increase product formation, only the second step of the reaction had to be enhanced (the ligation step in Fig. 1). We found that altering three components of the reaction enhanced product formation.

Figure 3A and B shows that optimizing the MgCl2 concentration and then the ATP concentration in the reaction enhanced product formation of (c+pc)/T2 from 12% (Table 1) to >90%. Under these conditions, however, (c+pc)/C2 ligates poorly (data not shown). Figure 3C shows that the inclusion of DMSO to this reaction enhances the ligation of (c+pc)/C2 from <10% to >90%. These reactions were optimal at 30°C, although only marginally better than at 22°C (data not shown). Figure 3D shows a typical time-dependence plot of the reaction, which indicates that under the optimized conditions products do not break down over time. The addition of PEG, which is a molecular crowding agent, does not further stimulate the reaction, nor does the inclusion of single-stranded binding protein (data not shown).

A representative gel under final optimized conditions with the T2 template is shown in Figure 4. Table 2 shows the results of duplicate trials of all 64 combinations of decamers and template under the optimized conditions (to be compared with the standard conditions in Table 1). These optimized
conditions dramatically increase the ligation of mismatched nucleosides at the site of ligation without reducing the effectiveness of ligation with matched base pairs. The template-dependent average for each of the 16 dinucleotide combinations is T2 (97%) > G2 (96%) > A2 (93%) > C2 (92%). Under these optimized conditions, each of the four canonical nucleosides can direct the effective ligation of all 16 possible sequence combinations at the n position. Note that C-G base pairs at the n±1 positions are not required for effective ligation, as replacing them with A-T pairs does not reduce ligation extent (data not shown).

Investigation of the n±1 position: optimized conditions

We analyzed whether these optimal experimental conditions can allow ligation of decamers that are mismatched at the positions one nucleotide away from the ligation junction (the n±1 donor and acceptor positions in Fig. 1) when the ligation junction itself (the n position) is base paired. In these cases, the templates have a central tandem segment of four identical nucleosides. The results (Table 3) show that these optimized experimental conditions do allow the effective ligation of mismatches at positions adjacent to the ligation junction. Note that, while nearly all combinations worked well, 4 out of 64 yielded <50% product. Dropping the reaction temperature from 30 to 22°C in these instances significantly enhanced these reactions [(gt-pc)/A4 went from 6.2 to 83.1%, (gt-pc)/A4 went from 24.6 to 87.2%, (cg-pcc)/C4 went from 34.7 to 90.9% and (tg-pgc)/C4 went from 42.2 to 97.4%]. Dropping the reaction temperature presumably aids in the hybridization of the decamers to the template and is not expected to significantly hinder the other reactions. The effectiveness of the template nucleosides follows a trend similar to that seen for the n position; T4 (92%) > G4 (92%) > A4 (80%) > C4 (79%).

Investigation of mismatches at both the n and n±1 positions

We analyzed the effects of having mismatches at both the n and n±1 positions, either in the donor molecule, the acceptor molecule or both. Analyzing all 256 possibilities (using a uniform universal template region) is impractical, so we analyzed a subset of 16 representative sequence combinations for each template (Table 4). Note that for this system the reaction temperature was 22°C, as it appeared to work better than 30°C.

In cases where no mismatches occur (dark gray background in Table 4) ligation went very well (89±96%), as expected. In cases where two tandem mismatches occurred with either the donor and template or the acceptor and template, ligation generally went well (55±97%), except for (gg-pcc)/C4 (31%). In cases where all four positions contain mismatches, the extents of reaction were highly variable. Using the A4, C4 and G4 templates, some donor±acceptor combinations essentially don’t work at all (<3% ligation). Using the T4 template, however, resulted in substantial ligation for all sequence combinations.

The data show percent product formation and percent intermediate formation (in parentheses). The horizontal heading denotes template used and the vertical heading denotes the new ligated product. See Figure 1 for naming conventions of acceptors, donors and templates. All reactions were run at 30°C for 18 h. Reaction conditions were as follows: 30 mM Tris–HCl (pH 7.5), 10 mM DTT, 10 mM MgCl2, 10 μM ATP, 20% DMSO, 2.5 pmol acceptor, 70 fmol donor and 3.125 pmol template.

<table>
<thead>
<tr>
<th>Template</th>
<th>A2</th>
<th>C2</th>
<th>G2</th>
<th>T2</th>
</tr>
</thead>
<tbody>
<tr>
<td>A4</td>
<td>96.2±2.7</td>
<td>97.0±0.3</td>
<td>95.5±1.2</td>
<td>97.8±0.6</td>
</tr>
<tr>
<td>C4</td>
<td>95.0±1.9</td>
<td>94.4±0.8</td>
<td>91.8±6.1</td>
<td>95.0±2.2</td>
</tr>
<tr>
<td>G4</td>
<td>93.4±5.0</td>
<td>96.0±1.4</td>
<td>95.5±1.8</td>
<td>95.8±0.1</td>
</tr>
<tr>
<td>T4</td>
<td>93.5±6.2</td>
<td>95.9±6.2</td>
<td>97.5±1.6</td>
<td>95.8±0.3</td>
</tr>
</tbody>
</table>

Table 2. Results of n position investigation under optimized conditions
are not significantly affecting our results, we synthesized and analyzed systems that mimic (ta-pat)/T4 and (tt-pat)/T4, except that the non-ligating terminal ends of each substrate are shortened and are completely complementary with the template. In this way, we found that the non-ligating terminal mismatches do not significantly affect our results, as the extent of ligation for (ta-pat)/T4 went from 95.6% in the end-matched system to 96.8% in the end-mismatched system. Similarly, (tt-pat)/T4 went from 24% in the end-matched system to 29.4% in the end-mismatched system. Therefore, the presence of terminal mismatches does not significantly affect our experimental results.

Although the ligation reactions were quantified in this report at 18 h, it is expected that the ligation rates of the various substrate–template combinations vary widely. Substrate–template combinations that are more complementary would be expected to have higher rate constants. This trend was confirmed by conducting time-dependent ligation at 22°C using the representative systems shown in Figure 5. The all matched system, (gg-gg)/C4, reacts the fastest, with ligation being complete within 1 min. The system with four mismatches, (tt-tt)/C4, reacts the slowest, with ligation being complete in ~12 h. The two systems with intermediate
observed rate constants each have mismatches at the \( n - 1 \) positions. Predictably, \((cg-pgc)/C4\), which does not ligate effectively at 30°C (Table 3), is more than 10-fold slower at 22°C than \((tg-pgt)/C4\), which does ligate effectively at 30°C (Table 3). Importantly, product breakdown is not observed during the reactions, indicating that 18 h appears to be an acceptable end point for these representative systems.

**DISCUSSION**

This is the first report showing that T4 DNA ligase can ligate oligonucleotides irrespective of the sequence of the terminal ends of the oligonucleotides being joined. In cases where mismatches only at the \( n \) position were tested, the extent of ligation was no less than 80%. In cases where mismatches only at the \( n - 1 \) position were tested, the extent of ligation was no less than 50% (after final temperature optimization). Although cases with two or more mismatches at the ligation junction can generate product in amounts above 50%, the ligation extent generally decreases as the number of mismatches increases.

**Reaction conditions**

The initial experiments indicate that T4 DNA ligase starts to lose fidelity under a relatively narrow range of \( MgCl_2 \) concentration (between 2 and 4 mM) (Fig. 3A). The reason for the narrow range is unknown, but is not due to inactivity of the enzyme at higher \( MgCl_2 \) concentrations, as intermediate formation readily occurs at up to 10 mM \( MgCl_2 \). Furthermore, since intermediate formation is dependent on duplex formation (Fig. 4), this effect is also not due to the presence or absence of duplex formation at higher \( MgCl_2 \) concentration (although it could relate to the stability of the helix, once formed). It has been previously noted that ligation of mismatches occurs most effectively when the melting temperature \( (T_m) \) of the pre-ligated duplex is slightly below the reaction temperature (5) and that only transient in-line attack is required for ligation (5,21). It could be that only the narrow range of divalent cation allows these steric conditions to occur. If true, ligation molecules that form more stable or less stable duplexes might require an adjustment to the \( MgCl_2 \) concentration.

The optimal range of ATP concentration is 10–100 \( \mu M \). Too little of this cofactor is ineffective as it will not allow for sufficient binding of ATP to the enzyme (34), leading to a lack of enzyme adenylation. At high ATP concentrations, intermediate formation is prevalent, indicating that the enzyme is active and the DNA duplex is formed; however, product formation is low. One possible explanation is that the enzyme is continuously adenylation at high ATP concentration. After the adenylation enzyme transfers the AMP to the donor molecule, the enzyme can dissociate from the complex. If the enzyme is re-adenylated before rebinding to the adenylation intermediate it can no longer catalyze the final ligation step.

The addition of 10–30% DMSO greatly enhanced the ligation of some mismatched oligonucleotides, primarily through an increase in the second step of the reaction (the ligation step in Fig. 1). It is possible, then, that nucleoside analogs developed to act as universal bases might also be more effective in this regard using buffers containing DMSO. Although the origin of the DMSO effect is unknown, it has been shown that DMSO can reduce the fidelity of nucleoside-binding restriction enzymes (35,36), which parallels our results. Nevertheless, DMSO might not be acting directly on the protein, as ligation of fully base paired nicks does not occur any faster in the presence or absence of DMSO (data not shown). Perhaps DMSO acts by interacting with the DNA in such a way as to favorably orient the mismatched components for the reaction.

That the ligation reactions are generally most effective at 30°C for the single and double mismatches and 22°C for the quadruple mismatches indicates that the drop in temperature permits more effective hybridization of the oligonucleotides to the template. Dropping the temperature even further, however, is ineffective, probably because the duplex becomes rigid or the enzyme loses activity. Increasing the temperature is also not helpful, as hybridization strength decreases, which also would be expected to increase the fidelity of the ligation reaction (26,37). Unfortunately, attempts to determine thermal stabilities of the various pre-ligated duplexes were unsuccessful due to the fact that the melting curves under our experimental conditions do not show a sigmoidal structural transition. We were able to obtain usable melting curves, however, using a standard thermodynamic analysis buffer (20 mM sodium cacodylate, 1 M NaCl and 0.5 mM Na2EDTA, pH 7.0) (38,39). We found that, at least for the pre-ligated representative duplex systems shown in Figure 5, those systems that ligate most effectively in general have the highest melting temperatures (data not shown). Apparently, those systems that are more stable (in terms of hybridization) are more effective in terms of reactivity, which suggests that hybridization strength is a key determinant of reactivity.

**Canonical nucleosides as universal template bases**

Each of the four canonical nucleosides act as universal template bases at both of the \( n \) positions or both of the \( n - 1 \) positions (see Fig. 1). This is also true for combined \( n \) and \( n - 1 \) position mismatches in either the donor or the acceptor (Table 4, light gray boxes), although such mismatches in the donor molecule using the C4 template are less effective than other double mismatch combinations. When mismatches occur at all four positions, however, ligation efficiencies drop off markedly.

There does not appear to be a straightforward correlation between the predicted thermodynamics of the individual mismatches (39–44) and the effectiveness of ligating oligonucleotides that contain such mismatches (at the ligation junction). For those situations where only one or two mismatches occur, all mismatches work well, irrespective of whether the individual mismatches are predicted to be stable or structured. For situations where four mismatches occur and are expected to be stable, only \((gg-pgg)/G4\) and \((tt-ptt)/G4\) work well, whereas \((aa-paa)/G4\) and \((gg-pgg)/T4\) do not. For those situations where four mismatches occur and are expected to be unstable, \((tt-ptt)/C4\) and \((cc-pcc)/T4\) work well, whereas \((cc-pcc)/C4\) does not. Therefore, either the presence of structure at these positions is not a factor in determining ligation activity or the different reaction conditions utilized between these reports preclude a direct comparison of the results.

In general, sequence-independent ligation can be accomplished using any of the canonical nucleosides in the template flanking the ligation junction. Note that these template
nucleosides do not necessarily form hydrogen bonds with the corresponding nucleosides in the donor or the acceptor molecule. In fact, it could be the lack of any specific and required interactions that permit the sequence-independent ligation activity that we observe. Indeed, previous results show that T4 DNA ligase can ligate oligonucleotides even if there is a single nucleotide gap or an abasic residue (15) at the ligation junction. Combined with the above data, it appears that the requirements for T4 DNA ligase do not include stable base pairing at the site of ligation. As suggested previously (21), transient proximity is likely sufficient.

As a template nucleoside, thymidine consistently works as well as or better than the other nucleosides. This is highlighted by the fact that the G4 template directs the ligation of tt to ggt poorly (5%) as compared with the T4 template directing the ligation of MgCl2 (3 mM) and ATP (10±100 M). This effect can be significantly enhanced by the inclusion of 10–30% DMSO.

Note that although we have shown that T4 DNA ligase can be a low fidelity enzyme in vitro, it is not expected that this is the case in vivo, as a low fidelity ligase would be highly mutagenic in terms of the genomic processes of recombination, replication and repair.

Our results show that canonical nucleosides can be utilized as universal bases in ligation templates, at least at ligation junctions. This property could potentially be exploited to develop protocols that allow for the ligation of oligonucleotides whose termini are unknown or degenerate. For example, a template molecule with tandem thymidines at the ligation junction could serve to direct the ligation of oligonucleotides of varying sequences. In such cases, if the sequence of the substrate region flanking the ligation junction is known, simple PCRs can be used to amplify ligated products for their subsequent sequencing. At its simplest, this strategy, among others, could potentially be useful for sequencing the terminal ends of oligonucleotides.

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