Universal strategies for the DNA-encoding of libraries of small molecules using the chemical ligation of oligonucleotide tags

Alexander Litovchick, Matthew A Clark, and Anthony D Keefe*
X-Chem Pharmaceuticals; Waltham, MA USA

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The affinity-mediated selection of large libraries of DNA-encoded small molecules is increasingly being used to initiate drug discovery programs. We present universal methods for the encoding of such libraries using the chemical ligation of oligonucleotides. These methods may be used to record the chemical history of individual library members during combinatorial synthesis processes. We demonstrate three different chemical ligation methods as examples of information recording processes (writing) for such libraries and two different cDNA-generation methods as examples of information retrieval processes (reading) from such libraries. The example writing methods include uncatalyzed and Cu(I)-catalyzed alkyne-azide cycloadditions and a novel photochemical thymidine-psoralen cycloaddition. The first reading method “relay primer-dependent bypass” utilizes a relay primer that hybridizes across a chemical ligation junction embedded in a fixed-sequence and is extended at its 3′-terminus prior to ligation to adjacent oligonucleotides. The second reading method “repeat-dependent bypass” utilizes chemical ligation junctions that are flanked by repeated sequences. The upstream repeat is copied prior to a rearrangement event during which the 3′-terminus of the cDNA hybridizes to the downstream repeat and polymerization continues. In principle these reading methods may be used with any ligation chemistry and offer universal strategies for the encoding (writing) and interpretation (reading) of DNA-encoded chemical libraries.

Introduction

The use of DNA-encoded chemical libraries permits the simultaneous interrogation of large numbers of small-molecules using affinity-mediated selection methodologies.1-11 Such libraries have recently been reported to have yielded potent small-molecule inhibitors for a range of therapeutic target proteins10,12-18 including for challenging targets such as the inhibition of protein-protein interactions.19-22 During the chemical synthesis of DNA-encoded libraries of small-molecules the chemical history of individual library members is either recorded using the sequences of oligonucleotide tags that are ligated to the library member within each of multiple chemistry-specific compartments or directed using a pre-existing oligonucleotide template. During DNA-recorded chemical library generation successive cycles of split, encoding by tagging, chemistry and pool generate large combinatorial mixtures of different small-molecules, the identities of each of which may be inferred by determining the sequence of the attached concatenated oligonucleotide tag set. A range of oligonucleotide tagging strategies has been reported to be suitable for the generation of DNA-recorded libraries including solid-phase coupled peptide and oligonucleotide synthesis,1,23 the enzymatic ligation of single-stranded oligonucleotides,7 the enzymatic extension of hybridized single-stranded oligonucleotides,24 the enzymatic ligation of double-stranded oligonucleotides10 and the chemical ligation of single-stranded oligonucleotides.25 Related strategies have been reported for the generation of DNA-directed libraries using template-directed chemical synthesis on a pre-existing template3,5,11 or hybridization-dependent routing of pre-existing templates to chemistry-specific compartments.5-8

Many of these oligonucleotide tagging schemes suffer from significant limitations. Hybridization-mediated schemes suffer from small split sizes because large differences need to exist between the sequences of all members of the encoding oligonucleotide set in order to reduce mis-hybridization rates. And some of these strategies require the covalent conjugation of each chemical building-block to a different oligonucleotide tag which will become laborious for large sets of chemical building blocks. Also, the use of enzymes to effect ligation or extension requires

*Correspondence to: Anthony Keefe; Email: akeefe@x-chemrx.com
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the alternate establishment of chemistry-compatible and enzyme-
compatible conditions often within large numbers of low-volume
compartments and can suffer from the inhibition of enzyme
activity by residual chemistry components.

The chemical ligation of oligonucleotides is an attractive alter-
native encoding strategy. Chemical reactions are generally more
versatile than enzymatic reactions and in many instances may be
performed under a range of solution conditions which could obvi-
ate the need for buffer exchange within thousands of low-volume
compartments - as has been reported for some enzymatic tagging
strategies.10 One of the more significant limitations of chemical
ligation-mediated encoding schemes is the apparent need to gen-
erate a chemical linkage between two oligonucleotides that may
be traversed by a template-dependent polymerase in order that tag
identities and associations may be determined by sequencing. In
this paper we demonstrate two different cDNA-generation (read-
ing) strategies for chemically ligated oligonucleotides—neither of
which require that a polymerase be able to translocate through
the chemical ligation junction. We also demonstrate three differ-
ent example chemical ligation strategies that may be read by these
processes. In principle both of these cDNA-generation strategies
permit the use of any chemical bond-forming reaction to support
an oligonucleotide tagging strategy for DNA-encoded chemical
library generation.

Strategies for generating chemically ligated oligonucleotides
that polymerases may translocate through have also been reported
and also offer alternative approaches for the synthesis of DNA-
encoded chemical libraries. Examples include the use of cyano-
ogen bromide or water-soluble carbodiimides to generate native
phosphodiester linkages from 5′-monophospho and 3′-hydroxyl
oligonucleotides,26,27 the use of 5′-Iodo and 3′-Phosphothio
oligonucleotides to generate phosphorothodiester linkages,28
the use of Cu(I)-catalyzed triazole synthesis with 5′-propargyl
and 3′-azido oligonucleotides29 and the use of Cu(I)-catalyzed
triazole synthesis with 5′-azido and 3′-propargyl oligonucle-
otides.25,30 The utility of such approaches for the generation of
DNA-encoded libraries of small-molecules is dependent upon the
extent to which tag polymerization can be avoided by for example
by making chemical ligations template-dependent, or the devel-
opment of a suitable protection strategy, or the use of pairs of
orthogonal chemistries. Challenges to these approaches are likely
to be experienced with regard to yield, removal of unreacted tags
and interference with the reaction scheme used to generate the
encoded chemical entity, but we believe that these can be over-
come and we are also exploring such approaches.25 In this paper
we report on strategies for the generation of soluble unimolecular
oligonucleotide-encoded information (writing) and the deter-
mination of this information (reading) using chemical linkages
between oligonucleotides that are not traversable by polymerases.
We present three chemical ligation methods and two cDNA-
generation methods by which this may be achieved and support
each with experimental demonstration of both the writing and
the reading processes.

Figure 1. Mechanism of cDNA generation from chemically ligated tags using relay primer-mediated bypass on a photochemically ligated non-travers-
able template. A terminal primer is annealed to the 3′-terminus of the chemically ligated oligonucleotide and a second 5′-monophosphorylated primer
(relay primer) is annealed across the chemical ligation junction. Extension of both primers and ligation generates full-length cDNA including potentially
encoding sequences from both upstream and downstream of the photochemical ligation junction.

Table 1. Yields of psoralen-mediated oligonucleotide-templated photo-
chemical oligonucleotide ligations under different solution conditions

| Buffer          | Psoralen-thymidine ligation conversion |
|-----------------|---------------------------------------|
| pH 5.5 Phosphate| 65%                                   |
| pH 7.0 Phosphate| 88%                                   |
| pH 9.4 Borate   | 80%                                   |
Results

Generation of cDNA from a chemically ligated conjugate using relay primer-dependent bypass

Relay-primer dependent bypass utilizes a “relay primer” and coupled template-dependent polymerization and ligation as a means to generate contiguous cDNA sequences that can in turn be amplified, cloned and sequenced. In this approach both termini of the concatenated tag set contain sufficient fixed sequence for the purposes of primer-binding and each chemical ligation junction is flanked on either side by sufficient fixed sequence for the purposes of relay primer-binding. A relay primer is our term for an oligonucleotide that hybridizes to a pair of chemically ligated tags in a manner that bridges their ligation junction. In the presence of a non-strand displacing template-dependent polymerase and annealed 5′-terminal and 5′-phosphorylated relay primers each primer is extended up to the 5′-terminus of the next. The addition of a ligase then seals the nick between these and generates a contiguous cDNA that may then be amplified, cloned and sequenced as part of the reading process. This process is shown in Figure 1. With multiple relay primers the sequences of multiple concatenated tags may be copied into a single cDNA. We refer to this method as relay primer-dependent bypass and note its similarity to the manner in which Okazaki fragments are generated on the lagging template strand and then ligated together during DNA replication.31

In order to demonstrate relay primer-dependent bypass we used a psoralen-mediated photochemical ligation strategy to generate a cyclobutane adduct between an adjacent 3′-thymidine and a 5′-psoralen in a template-dependent reaction using a splint oligonucleotide to co-locate the photo-reactive oligonucleotide termini as shown in Figure 2A and B. The stereochemistry of the thymidine-psoralen adduct may be either cis or trans.32 The linkage thus formed lacks a contiguous sugar-phosphate backbone and does not resemble a wild-type internucleotide linkage. This ligation is efficient as indicated in Figure 2B with conversions exceeding 90% and the conjugate of (1) and (4) was synthesized with a yield exceeding 80% and an observed deconvoluted MW of 15,819 Da (calculated MW 15,813 Da). Photochemistry of this kind is orthogonal to most other chemical reactions and may be performed under a range of solution conditions - we have determined that it occurs with high yield at pH values as low as 5.5 and as high as 9.4 as shown in Table 1.

Oligonucleotides modified with psoralen are easily obtained commercially and photochemistry is readily applied to multi-well plates containing many low-volume samples. Accordingly, a photochemical ligation strategy such as this appears to be well-suited for the generation of DNA-recorded chemical libraries so long as tag identities and associations can be determined by sequencing. The first step in the process of determining the sequence of photoligated concatenated psoralen oligonucleotide tags is to generate a contiguous cDNA from which the sequences and associations of the photoligated tags can be deduced. To facilitate the subsequent separation of the cDNA from the photoligated template for analysis purposes the 5′-terminus of the template was biotinylated, and to facilitate detection of the cDNA the 5′-terminus of the terminal primer was fluoresceinylated. The photoligated conjugate was hybridized to the 5′-terminal primer and the 5′-monophosphorylated relay primer. This hybridized complex was then incubated with T4 DNA Polymerase to extend both the relay primer and the terminal primer. T4 DNA Polymerase was chosen because it does not exhibit strand-displacement activity27 and so was expected to extend the 3′-terminus of the terminal primer up to the...
5′-terminus of the relay primer without displacing it. Because the 5′-terminus of the relay primer is 5′-monophosphorylated the junction between the extended relay primer and the extended terminal primer is a substrate for ligation with T4 DNA Ligase which was then added to the incubating mixture along with terminal primer is a substrate for ligation with T4 DNA Ligase once the incubation was complete the hybridized complex was incubated with immobilized streptavidin to capture the chemically ligated conjugate and the cDNA was eluted with 0.1 M NaOH and neutralized. LCMS analysis is shown in Figure 3A and B with detection of the fluorescein at 495 nm indicating that approximately 50% of the terminal primer had now been extended to generate a product with a mass close to that of full-length cDNA (observed deconvoluted MW 15,172 Da, calculated MW 15,459 Da) with the observed mass suggesting the likely loss of one dT. The remaining approximately 50% corresponds to the extension of the terminal primer up to the psoralen ligation junction (observed deconvoluted MW 11,157 Da, calculated MW 11,154 Da). A control experiment in which the T4 DNA Ligase was omitted gave no full-length cDNA as expected. Sequencing indicated the cDNA had the expected sequence except for the absence of a 3′-terminal dT along with a shorter cDNA that was terminated at the psoralen ligation junction. Of 34 full-length cDNA, all were missing a single dT from the 3′-terminal region with 33 of these containing no more than one additional deletion or substitution.

Generation of cDNA from a chemically ligated conjugate using repeat-dependent bypass

Repeat-dependent bypass utilizes a “repeat” architecture in which the chemical ligation junction is flanked upon either side by the same repeated sequence. In the presence of a hybridized upstream primer and a suitable polymerase the primer will be extended up to the chemical ligation junction and then stall having copied the first of the repeated sequences. If the rearrangement of this stalled complex results in hybridization of the 3′-terminus of the newly synthesized cDNA to the second of the repeated sequences then template-dependent polymerization may continue. This process is shown in Figure 4. With multiple different repeats flanking multiple chemical ligation junctions multiple concatenated tags may be copied into a single cDNA. We have shown by experiment that this process readily occurs, does not lead to the shuffling of tag combinations in free solution, and we propose a mechanism by which it may occur. We refer to this method as repeat-dependent bypass and note its similarity to the manner in which repeated sequences within genomic DNA can be contracted or expanded in processes that can lead to genomic instability and have been observed to occur in vitro to different extents with different polymerases.29

In order to demonstrate repeat-dependent bypass we successively used a Cu(I)-free triazole formation reaction followed by a Cu(I)-catalyzed triazole formation reaction thereby enabling the controlled sequential conjugation of a central oligonucleotide functionalized with both a 5′-alkynyl and a 3′-azide first by reacting it at its 3′-terminus and then at its 5′-terminus as shown in Figure 5A. The chemical structure of each linkage is different and neither was expected to be traversable by a polymerase in the absence of a bypass mechanism because of the relatively large distance between the ligated bases and the fact that similar length triazole linkages have been shown not to function as templates for PCR amplification.29 This was tested experimentally as described below. The Cu(I)-free ligation conversion was over 90% (Fig. 5B) and the Cu(I)-catalyzed ligation conversion was over 75% (Fig. 5C). After gel-purification and extraction we isolated 6 and 8 nmoles (40 and 53%) of each of two doubly conjugated oligonucleotides—one constructed by conjugating (7) to (8) and (9) ("short-short") and the other by conjugating (7) to (10) and (11) (“long-long”). The final products were analyzed by LCMS with observed deconvoluted MW values of 38,572 and 49,915 Da (calculated MW 38,560 and 49,917 Da) as shown in Figure 5D and E. We screened a panel of DNA polymerases for their ability to read through the Cu(I)-catalyzed linkage formed by conjugating (7) and (9) including SuperScript III (Invitrogen),

![Figure 3. cDNA generation from chemically ligated tags using relay primer-mediated bypass on a photochemically ligated non-traversable template. (A) LC analysis showing approximately equal amounts of two primer-extension products after purification with detection at 495 nm. (B) Deconvoluted MS analysis of mixed primer-extension products showing two principal components with masses of 15,172 Da and 11,157 Da which correspond to full-length cDNA minus one dT, and primer extension up to the photochemical ligation junction respectively.](image-url)
Taq (NEB), Thermoscript (Invitrogen), KOD (Novagen), Vent (NEB), Vent (exo-) (NEB), Deep Vent (NEB) and Deep Vent (exo-) (NEB). We observed that PCR products were only generated with Vent (exo-) and Deep Vent (exo-). Similar constructs without a repeated sequence flanking the chemical ligation junction gave no PCR products under the same conditions. PCR amplification was conducted for each conjugate separately and for an equimolar mixture of the two conjugates using Deep Vent (exo-) DNA Polymerase. Analysis of the PCR products by agarose gel electrophoresis showed that the amplification of each of the isolated conjugates generates double-stranded PCR products with electrophoretic mobilities that correspond in each case to the loss of one of each of the repeated sequences in each of the pairs of ligation junctions—as shown in lanes 3 and 4 of Figure 6. The electrophoretic mobility values observed are consistent with the operation of the repeat-dependent bypass mechanism shown in Figure 5. The short-short conjugate is 120 nucleotides in length with an expected PCR amplification product length of 96 base-pairs and the long-long conjugate is 156 nucleotides in length with an expected PCR amplification product length of 132 base-pairs. The mixed conjugate sample was also analyzed by cloning and sequencing. Of 35 sequence reads, 25 were observed to be of the long-long combination and 10 of the short-short combination with no chimeric sequences observed with all sequence reads showing only one of each of the repeated sequences at each junction.

**Discussion**

**Relay primer-dependent bypass**

In order to demonstrate cDNA generation using relay primer-dependent bypass and a template interrupted by a non-polymerase traversable chemical ligation junction we first developed a method to generate a psoralen-thymidine ligated oligonucleotide. In contrast to the more well-known use of psoralen-modified oligonucleotides to generate psoralen-thymidine cyclobutane interstrand cross-links between hybridized oligonucleotides (see for example refs. 36 and 37) this method generates psoralen-thymidine cyclobutane linkages between adjacent nucleotides on the same strand and appears to be novel. This linkage is formed by co-hybridizing a 3'-thymidine oligonucleotide adjacent to a 5'-psoralen oligonucleotide to a complementary splint oligonucleotide that had no thymidines in closer proximity to the psoralen than four nucleotides away in the 3'-direction or two nucleotides away in the 5'-direction (closer proximity was not tested), and then irradiating the complex with UV light. We were able to generate this conjugate with isolated yields of approximately 80%. Yields of this magnitude translate into overall yields of 50% for the successive ligation of two tags and of 40% for the successive ligation of three tags. For DNA-encoded chemical library applications these numbers of encoded cycles are of most interest to the medicinal chemistry community as they are more likely to lead to Lipinski-compliant compounds. For cDNA generation a 5'-biotinylated version of the acceptor oligonucleotide was used (4) and the product was then isolated using denaturing PAGE. Using this template we were able to demonstrate the formation of cDNA with a yield of 50% and with the expected sequence. These observations demonstrate our ability to photochemically ligate 5'-psoralen oligonucleotides to 3'-thymidine oligonucleotides using a complementary oligonucleotide splint (writing) and to then generate cDNA that permits the determination of the sequences and associations of ligated tags (reading).

**Repeat-dependent bypass**

In order to demonstrate the generation of cDNA using repeat-dependent bypass using a conjugate interrupted by a pair of non-polymerase-traversable chemical ligation junctions we first developed a method that allows for the controlled generation of two triazole chemical ligation junctions within a single conjugate using a copper-free triazole formation from an azide and a strained alkyne followed by a Cu(I)-mediated triazole formation similar to other reported reactions. This approach
permits the specific conjugation of three oligonucleotides without the need to use any protecting groups, also in high yield. For cDNA generation an oligonucleotide conjugate was generated by the chemical ligation of (7), (8) and (9) (“short-short”) and used as a template for PCR along with a similar second conjugate generated by the chemical ligation of (7), (10) and (11) (“long-long”). Each of these conjugates has two chemical ligation junctions, within each conjugate each junction is flanked by a different repeated sequence, but these repeated sequences are the same for each of the long-long and short-short conjugates. Both conjugates were used to demonstrate that conjugates of this general design can be used to program template-dependent polymerase-dependent amplifications such as PCR. Because the conjugates are of different lengths we were also able to use them to demonstrate whether information encoded in ligated tag set combinations would be preserved during repeat-dependent bypass amplification, or alternatively whether it would be lost via recombination-mediated processes similar to those seen during DNA shuffling.42 We were able to show that amplification using each of these templates resulted in amplification products containing only one of each repeated sequence at each junction, and that mixed amplification products were not generated when the templates were mixed prior to amplification. These data are consistent with our proposed loop-formation/ 

Figure 5A and B. (A) Controlled concatenation of three oligonucleotides using successive Cu(I)-free and Cu(I)-catalyzed chemical ligation by triazole formation. First the central oligonucleotide (7) is chemically ligated to (8) using the copper-free reaction of the azide group in (7) with the strained alkyne in (8), thereby preserving the terminal alkyne in (7) which is subsequently reacted in a Cu(I)-catalyzed reaction with the azide in the subsequently added third oligonucleotide (9). (B) LC time-course of Cu(I)-free conjugation of (7) with (8) using absorbance at 260 nm.
slippage repeat-dependent bypass model (Fig. 4) and suggest that if recombination processes are occurring they are relatively infrequent. Accordingly sequence data derived from the amplification of chemically ligated conjugates using this cDNA-generation process can be used to infer the identities and co-association statistics of ligated oligonucleotides present in the input conjugates—as is required for the successful application of this strategy to the encoding of the chemical history of individual library members within DNA-encoded libraries.

Material and Methods

Synthesis of an oligonucleotide conjugate with one psoralen-thymidine photochemical ligation junction

Oligonucleotides (1), (2) and (3) (shown in Scheme 1) were acquired from Integrated DNA Technologies. All three were mixed together in aqueous 500 mM sodium phosphate at pH 7.0 with the psoralen oligonucleotide (1) at 1 mM and the other two oligonucleotides at 1.1 mM. The mixture was then briefly heated to 95 °C followed by slow cooling to room temperature to allow hybridization to occur. Ten microliters aliquots were then cooled
to 4 °C and irradiated in 1.5 ml polypropylene microcentrifuge tubes (Fisher Scientific, 02-682-550) with UV light at 365 nm using a UVL-21 compact UV lamp (UVP). Aliquots of 1 uL were taken over a time-course and were analyzed by denaturing PAGE. The gel was visualized and photographed using the UV shadowing of a fluorescent TLC plate.

**Generation of cDNA from an oligonucleotide conjugate with one psoralen-thymidine photochemical ligation junction**

A gel-purified conjugate of (1) and (4) was then incubated in a 100 ul volume with a 5'-monophosphorylated relay primer (5) and a 5'-FAM-labeled terminal primer (6) each at 10 uM in T4 DNA Ligase Buffer (NEB) with 1 mM of each dNTP and 30 units of T4 DNA Polymerase (NEB). Following incubation at 1 h at 37 °C the mixture was supplemented with an additional 0.5 mM ATP and 10 units of T4 DNA Ligase (NEB) were added and it was then incubated for an additional 1 h at 37 °C. The reaction product was then purified by incubation with 200 µl of streptavidin-coated Dynabeads M280 (Invitrogen), washed, eluted with 35 µM of 0.1 M NaOH followed by neutralization with 1M pH 7.0 tris(hydroxymethyl) aminomethane. The product was then analyzed by LCMS on a Thermo Scientific LCQ Fleet using an ACE 3 C18–300 (50 × 2.1 mm) column and a 5 min gradient of 5–35% of buffer B using buffer A (1% hexafluoroisopropanol [HFIP], 0.1% di-isopropylethyl amine [DIEA], 10 µM EDTA in water) and buffer B (0.075% HFIP, 0.0375% DIEA, 10 µM EDTA, 65% acetonitrile/35% water). LC was monitored at 260 nm and 488 or 495 nm. MS was detected in the negative mode, and mass peak deconvolution was performed using ProMass software. In order to determine the sequence of both cDNA regions that were generated as a result of primer extensions, the 3'-terminus of the cDNA was first ligated to a new primer-binding oligonucleotide 5'-monophospho-GCTGTGCAGG TAGA GTGC-FAM-3' using single-stranded enzymatic ligation with 1 unit/ul of T4 RNA Ligase 1 (NEB,) in 1× T4 RNA Ligase buffer (NEB), supplemented with 200 µM ATP (NEB), 1 mM hexamine cobalt chloride (Sigma-Aldrich) and 25% PEG8000 (NEB) for 16 h at room temperature. This newly ligated cDNA was then purified using denaturing PAGE with fluorescence to visualize the band of interest followed by excision, extraction and PCR amplification was then conducted using the ligated cDNA at 200 pM, with the forward primer 5'-GCACCTCTACC TGCA CGACGC-3' and the reverse primer 5'-GGCGTACGG AAGCTCG-3' each at 0.5 µM and Platinum Supermix PCR mix (Invitrogen). Sixteen thermal cycles were performed with denaturation at 94 °C for 30 s, annealing at 56 °C for 30 s and extension at 72 °C for 60 s. Amplification products were visualized on ethidium bromide-stained 4% agarose E-Gels (Invitrogen) with illumination at 305 nm. Subsequently 0.2 units/µl of Klenow DNA Polymerase (NEB) were added to the PCR reaction followed by
incubation for 15 min at 37 °C. PCR products were then cloned using a Zero Blunt TOPO cloning kit (Invitrogen) and these clones were Sanger sequenced (Genewiz).

**Synthesis of an oligonucleotide conjugate with two different triazole chemical ligation junctions**

Oligonucleotides (7), (8) and (9) were acquired from Integrated DNA Technologies (IA, USA). Oligonucleotides (7) and (8) were mixed together at 0.5 mM concentration in aqueous solution with 0.2 M sodium phosphate buffer at pH 7.0. The reaction progress was monitored by LCMS. To this mixture was then added oligonucleotide (9) and a Cu(I)-catalyzed triazole formation reaction was conducted by further incubation with two molar equivalents of Cu(II) acetate, four molar equivalents of sodium ascorbate and one molar equivalent of Tris-(BenzylTriazolylmethyl) Amine (TBTA) in a volume of 50 μl with oligonucleotides at a concentration of 0.5 mM. This mixture was incubated overnight at room temperature followed by LCMS and purification using denaturing PAGE.

**Generation of cDNA from an oligonucleotide conjugate with two triazole chemical ligation junctions**

PCR amplifications were conducted with conjugates at 40 pM, with the forward primer 5′-TGCGGTCTAATCCGTGTTGTGGCAGATGGGACACCTTGGAGAAGCAGG-3′ and the reverse primer 5′-AAGCATAGCA CCCGATT-3′ each at 0.5 μM using 0.1 units/μl of Deep Vent (exo-) Polymerase (NEB) in 1× Thermopol Buffer (NEB) with dNTP at 0.5 mM. Twenty-two thermal cycles were performed with denaturation at 94 °C for 30 s, annealing at 52 °C for 30 s and extension at 72 °C for 60 s. Amplification products were visualized on ethidium bromide-stained 4% agarose E-Gels (Invitrogen) with illumination at 305 nm. PCR products were cloned using a Zero Blunt

Scheme 1. Sequences and chemical modifications of oligodeoxynucleotides used in this study.
TOPO cloning kit (Invitrogen) and these clones were Sanger sequenced (Geneviz).

**Conclusion**

There is significant and increasing interest in the use of the affinity-mediated selection of DNA-encoded libraries of small molecules to discover inhibitors of proteins of therapeutic interest. In order to construct such libraries efficiently it is necessary to utilize robust methods for the ligation of oligonucleotides that are in turn able to support the determination of oligonucleotide tag sequences and associations while not interfering with the wide range of chemical reactions that will be used in the synthesis of the encoded library members. In this study we have demonstrated three different chemical ligation strategies—the uncatenated cycloaddition of a strained alkyne and an azide, the Cu(I)-catalyzed cycloaddition of a terminal alkyne and an azide and the photochemical cycloaddition of thymidine and psoralen. Each of these chemical ligations may be used as part of an encoding strategy for the chemical synthesis of encoded libraries. We have also developed two independent methods that may be used to generate cDNA that incorporates sequence information derived from oligonucleotide tags concatenated using these chemical ligations. cDNA generated by these methods may be amplified, cloned and sequenced and used to infer identity and history of individual molecules that together comprise libraries of encoded variants synthesized using split-and-mix methodologies, similar for example to (10). Both methods also appear well-suited to support the use of libraries of such compounds in affinity-mediated selection techniques to discover small molecule inhibitors of proteins of therapeutic interest as has been reported for other related technologies, also similar for example to (10). Because both of these cDNA-generation methods are agnostic to the structural details of the chemical ligation junction they are in principle applicable to any chemical ligation oligonucleotide tagging method. Therefore the demonstrated instances may be considered examples of universal strategies. We expect that these methods will find utility in support of the generation and use of DNA-encoded chemical libraries to discover inhibitors of therapeutically relevant protein targets within drug discovery programs.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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