OLIGONUCLEOTIDES WITH CATIONIC PHOSPHORAMIDATE INTERNUCLEOSIDE LINKAGES AND METHODS OF USE

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Continuation-in-part of application No. 08/619,301, filed on Mar. 21, 1996, now Pat. No. 5,734,040.

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This invention relates to the use of oligonucleotides with cationic phosphoramidate internucleoside linkages for the identification of nucleic acid under DNA—DNA duplex denaturing conditions.

16 Claims, 3 Drawing Sheets
OTHER PUBLICATIONS


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Fig. 1A

\[
\begin{align*}
O &= P - NCH_2CH_2OCH_3 \\
O &= P - NCH_2CH_2N
\end{align*}
\]

**METHOXYETHYLAMINE PHOSPHORAMIDATE**

\((*)\)

**N,N'-DIETHYLETHYLENEDIAMINE PHOSPHORAMIDATE**

\((+)\)

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**Fig. 1B**

| SEQ. ID. NO.: 1 | 5' A G T T T T G T G T C C C C C T C T C A G G T G T C A C A G |
|                | \(0\% \text{ MODIFIED}\) |
| P-1            | A+A+A+A*T+A-T+A-G+G-G+A-G+A+G |
| P-2            | A+A+A+A*T+A+T+A-G+G-G+G+A+G |
| P-3            | A+A+A+A*T+A+T+A+G+G+G+A+G+A+G |
| P-4            | A+A+A+A*T+A+T+A+T+A+G+G+G+A+G+A+G |

| SEQ. ID. NO.: 3 | 5' G C C C C C T G G C C C C C T C C C C T T T G T T C C A T T T |
| SEQ. ID. NO.: 9 | 3' C G G G G A C C G G G G G A A A C A A G G T A A A |
Cross-correlation
Sep021758

bin = 4
offset = 8.0E+6
amplitude = 5.4E+5
width = 9.34
M = 190.73
SNR = 18.24

Fig. 2A
OLIGONUCLEOTIDES WITH CATIONIC PHOSPHORAMIDATE INTERNUCLEOSIDE LINKAGES AND METHODS OF USE

CONTINUING APPLICATION DATA

This application is a continuation-in-part of parent application Ser. No. 08/619,301, filing date Mar. 21, 1996 which issued as U.S. Pat. No. 5,734,040 on Mar. 31, 1998.

GOVERNMENT FUNDING

This work was performed, in part, by funding from the National Institutes of Health Grant No. HLA 42266. The government may have certain rights to this invention.

FIELD OF THE INVENTION

This invention relates to oligonucleotides used to control gene expression or other cellular events. In particular, this invention relates to oligonucleotides with cationic phosphoramidate internucleoside linkages.

BACKGROUND OF THE INVENTION

Synthetic oligonucleotides have been shown to regulate gene expression in cells. Thus, these molecules bind to target regions of nucleic acids and inhibit gene expression. They are useful to control infectious disease and have been used and tested in a variety of diseases or conditions associated with altered or aberrant gene expression such as cancers, hereditary disorders, and the like.

There are two major oligonucleotide (ODN) based strategies designed to inhibit gene expression. One method uses antisense ODNs complementary to a specific mRNA to form DNA-RNA hybrids. These hybrids are stabilized by Watson-Crick base pairing and are substrates for cellular ribonuclease H (RNase H), an enzyme that degrades the RNA portion of the duplex rendering the mRNA untranslatable (Walter et al., Proc. Natl. Acad. Sci. USA, 85:5011–5015, 1988).

Because RNase H does not degrade the ODN, the ODN is able to hybridize to another copy of the target mRNA.

A second ODN-based strategy for altering gene expression targets DNA by the formation of a triple helical structure. The use of ODNs to form triple helix structures was initially reported by Moser et al. (Science, 238:645–650, 1987, and see LeDoan et al., Nucl. Acids Res., 15:7749–7760, 1987). Under suitable conditions, an ODN will bind in the major groove of a DNA duplex. The presence of a third strand may either sterically block transcription, prevent the sequence specific interactions of regulatory proteins with DNA, and/or alter the conformation of the bound duplex.


The second triplex motif involves a purine rich triplex forming oligonucleotide (TFO). Thymidine or adenosine residues of the third strand bind to the adenine of an A:T duplex and guanosine in the third strand interacts with the guanosine of a G:C duplex (Beal et al., Science, 251:1360–1363, 1991; Durland et al., Biochem., 30:9246–9255, 1955, and Cooney et al., Science, 241:456–459, 1988). The orientation of the third strand has been shown to be antiparallel to the purine rich strand of the duplex (Beal, et al., supra). The major drawback to using this approach in vivo is the tendency of G-rich ODNs to self associate into quartets at physiologic potassium concentrations. However, recent studies indicate that the use of a GT rich ODN to affect transcription of a transfected CAT plasmid in vivo indicates that GT rich combinations may minimize quartet formation. Unmodified negatively charged oligonucleotides do not generally stably participate in triplex formation in cells. Triplex forming oligonucleotide strategies must account for the physiologic concentrations of Mg\(^{2+}\) and the level of potassium tolerated for stable triplex formation.

To improve the stability and cellular uptake of oligonucleotides, oligonucleotides have been prepared having modifications to the phosphate backbone. For example, phosphorothiate and methylphosphonate derivatives of oligonucleotides have been synthesized and have sequence specificity and hybridization strengths similar to that of unmodified oligonucleotides. Aminoethyl phosphonate derivatives of oligonucleotides have also been synthesized and demonstrate enhanced stability in aqueous solution as compared with aminoethylphosphonate linkages (Fathi et al., Nucl. Acids Res., 22:5416–5424, 1994). The uncharged character of the methylphosphonate derivatives permits an enhanced uptake of the oligonucleotides by the cell and increased resistance to nucleases as compared with unmodified oligonucleotides. Methoxyethyl-phosphoramidate linkages that have been incorporated into the internucleoside backbone at 3' and 5' linkages to inhibit exonuclease degradation. The synthesis of ODNs with some modified and some unmodified linkages allowed both increased nuclelease resistance while maintaining RNase H mediated target RNA degradation. (Dagle et al., Antisense Res. and Devel., 1:11–20, 1991).

Intercalating agents have also been conjugated to oligonucleotides to increase the stability of the conjugate with the complementary strand (see, for example, Wilson, et al., Biochem., 32:10614–10621, 1993, and Orson et al., Nucl. Acids Res., 22:479–484, 1994). In addition, molecules are often attached to the oligonucleotides to modify the net charge. Examples of these agents include polycationic, cationic peptides, polyamines and polycationic polymers. Intercalators and polycations have shown an increased resistance to nucleuse degradation.

Nucleosomonomers can also be modified to improve triplex formation and PCT International Publication No. WO 94/21444 discloses oligomers with 7-deaza-7-substituted purines.

The formation of a DNA triplex using a purine rich ODN is inhibited by monovalent cations, particularly potassium ions (Olivas, et al., Biochem., 34:278–284, 1995). Intracellular K\(^+\) concentrations inhibit triplex formation using unmodified oligonucleotides. Potassium ions are the predominant intracellular cations. One problem with forming triplexes at physiologic K\(^+\) levels is the self association of guanosine-rich ODNs into aggregates which are stabilized by guanine quartets (Olivas, et al., Biochem., 34:278–284, 1995; Olivas, et al., Nucl. Acids Res., 23:1936–1941, 1995).
In addition to decreasing the rate of triplex association, K⁺ increases the rate of triplex dissociation in vitro (Olivas, Biochem., supra). The inhibitory effect of K⁺ can be partially diminished by chemical modification of ODNs. For example, incorporation of the modified base 6-thioguanine in place of native guanine into TFOs decreases the association of ODNs into quartets and increases triplex formation in the presence of monovalent cations (Olivas, Nucl. Acids Res., supra, and Rao, et al., Biochem., 34:765–772, 1995).

Another hurdle associated with the use of ODNs in vivo is that the oligonucleotides are rapidly degraded by intracellular nucleases (Rebagliata, et al., Cell, 48:599–605, 1987; Dagle et al., Nucl. Acids Res., 18:4751–4757, 1990; Dagle, et al., Antisense Res. and Dev., 1:11–20, 1991). The chemical modification of ODNs provides resistance to nuclease degradation (Dagle, et al., supra), potentially increasing the overall activity of these compounds in vivo (Dagle, et al., supra, and Weeks, et al., Development, 111:1173–1178, 1991). The type and degree of chemical modification of ODNs, however, is limited when strategies require the action of cellular RNase H (Dagle, et al., Antisense, supra).

Additionally, some modification of ODNs can result in nonspecific toxicity mediated through non-nucleic acid interactions, such as has recently been reported for phosphorothioate ODNs and the basic fibroblast growth factor receptor (Guvakova, et al., J. Biol. Chem., 270:2620–2627, 1995). In contrast, the formation of triplex structures does not require an enzymatic activity and thus allows greater flexibility with regard to ODN design. The enhanced nucleolytic stability of an ODN with many or all internucleoside linkages modified would be useful for in vivo applications if these compounds are able to form stable triplex structures. The present invention discloses a class of oligonucleotides with enhanced stability for duplex and triplex formation.

Candidate oligonucleotides should produce triplex formation at physiologic salt concentrations. The close association of two nucleic acid strands creates a highly negatively charged environment. Many oligonucleotides disclosed in the literature to date require the presence of magnesium ions above physiologic concentrations to produce stable triplex formation. The obligate presence of magnesium ions for triplex association most likely reduces interstrand charge repulsions. However, magnesium concentrations within the cell cannot be altered without altering cell physiology. Therefore, oligonucleotides dependent on minimum concentrations of magnesium ions may not function well in the cellular milieu.

There is a need for oligonucleotides with improved binding stability and for methods that take advantage of the increased binding stability of these oligonucleotides.

**SUMMARY OF THE INVENTION**

The present invention relates to the selection, production, and use of oligonucleotides with cationic phosphoramide modified internucleoside linkages.

In a preferred aspect of this invention, the invention relates to a method for hybridizing an oligonucleotide to denatured nucleic acid to form a duplex comprising the steps of hybridizing a duplex-forming oligonucleotide comprising at least one ethylenediamine-class phosphoramide internucleoside linkage to single-stranded nucleic acid; and detecting hybridization of the oligonucleotides to the single-stranded nucleic acid. In one embodiment the oligonucleotide comprises at least about 30% to about 100% ethylenediamine-class phosphoramide internucleoside linkages. The nucleic acid can be DNA or RNA and in one embodiment the denatured nucleic acid is denatured using heat, denaturing concentrations of salt or at least one chaotrope. Preferably the chaotrope is selected from the group consisting of formamide, urea or a guanidinium salt. In one embodiment, the oligonucleotide is about 12 to about 50 nucleotides in length and preferably about 15 to about 50 nucleotides in length. Preferably the oligonucleotide comprises N,N-diethyl-ethylenediamine phosphoramide internucleoside linkages. In one embodiment the oligonucleotide is labeled with a tag and the tag is selected from the group consisting of a radioactively labeled tag or a fluorescent tag. In a preferred method the oligonucleotide further comprises a nucleic acid sequence capable of hybridizing to a portion of a gene, wherein the detecting step indicates the presence of the gene in denatured DNA. In one embodiment, the denatured DNA is genomic DNA. In another embodiment, the method is used to detect single-nucleotide polymorphisms.

The invention also relates to a method for detecting a nucleic acid fragment comprising the steps of hybridizing an oligonucleotide comprising at least 30% ethylenediamine-class phosphoramide internucleoside linkage under DNA—DNA duplex denaturing conditions to nucleic acid to form a hybridization complex and detecting the hybridization complex. In one embodiment the oligonucleotide is at least about 12 nucleotides in length. The DNA—DNA duplex denaturing conditions preferably comprise at least one chaotrope and preferably the chaotrope is selected from the group consisting of formamide, urea or a guanidinium salt. In a preferred embodiment, the oligonucleotide is labeled such as with a radioisotope, a fluorescent tag or another tag capable of detecting the oligonucleotide including an enzyme, an antibody, and the like. In one embodiment the nucleic acid fragment comprises genomic DNA.

**BRIEF DESCRIPTION OF THE FIGURES**

FIG. 1A details the structures of phosphoramide modified internucleoside linkages compared in this invention in one set of studies according to this invention. The neutral 2-methoxyethylamine derivative on the left and the N,N-diethyl-ethylenediamine derivative (shown in an uncharged state) on the right.

FIG. 1B lists exemplary oligonucleotides (ODN) used to form triplex structures according to this invention. ODNs U-1, N-1, N-2, P-1, P-2, P-3 and P-4 bind in the major groove of Duplex I, as shown. All of the triplex forming oligonucleotides (TFOs) have the same sequence but are modified as follows: –2phosphodiester linkages, –N,N-diethyl-ethylenediamine phosphoramide linkages and +2,N,N-diethyl-ethylenediamine phosphoramide linkages. The percent of modified bonds are listed to the right of each ODN. Duplex II represents an unrelated, nontarget polyurine-polyuridine duplex.

FIG. 2A is a graph produced using a single-molecule detection apparatus detecting hybridization of the oligonucleotide in the presence of the λ target in a single detection zone by crosscorrelation analysis. FIG. 2B is a control also using the single-molecule detection apparatus showing that no correlated signal is seen in a negative control sample identical to that of FIG. 2A but without the λ target DNA. “SNR” refers to (amplitude)/(standard deviation(values including the base line)). “bin” refers to data collected in 1 ms (milliseconds) intervals and taking a 4 ms collection time to determine the photons detected over 1 ms. “offset” refers to the baseline average on either side of the peak; “amplitude” refers to the peak value minus (offset). Numbers such
as 5.4E+6 refers to scientific notation and is equivalent to 5.4x10^6. "width" refers to the full width at half maximum (FWHM) of the fitted curve. “M” refers to (amplitude)/(sqrt(root(offset))).

DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to the selection, production, and use of oligonucleotides with cationic phosphoramidate modified internucleoside linkages to form stable triplex or duplex DNA structures. The oligonucleotides of this invention, that is, oligonucleotides with cationic phosphoramidate modified internucleoside linkages, can be prepared to bind to a variety of nucleic acids including RNA or DNA and single stranded or double stranded nucleic acid and therefore have a wide range of applications for inhibiting transcription or for detecting duplex formation.

The term "in vitro" is used herein to cell-free experiments (i.e., either as a cell lysate or in solution). The term "in vivo" refers to intracellular events, experiments or work performed on intact cells. That is, those of ordinary skill in this art will recognize that whether or not a particular cell is existing in culture, ex vivo, or in situ (i.e., within a particular tissue or as part of an organism), the ability of the oligonucleotides of this invention to bind to target DNA within a cell is independent of the location of the cell.

For example, transcription level-altering oligonucleotide-directed strategies can be and have been directed to a variety of cancers that have resulted from inappropriate gene activation, including leukemias (such as the use of oligonucleotides to purge bone marrow cells of leukemic progenitors prior to bone marrow transplantation) or cancers associated with oncogene activation. These types of strategies can benefit from the oligonucleotides of this invention. The oligonucleotides of this invention can also be used to control or limit viral infection (including, but not limited to, HIV, members of the Herpesvirus family including HSV, CMV, or VZV; as well as other viruses where transcription is directed from duplex DNA) or other intracellular pathogens through the suppression, alteration, or reduction of gene expression (for a general discussion see Askari, et al., New Engl. J. Med., 334(5):316-318, 1996). The oligonucleotides of this invention can also be used in triplex-forming strategies for a variety of metabolic diseases including inborn errors of metabolism, hypertension, and cardiovascular disease (such as oligonucleotides engineered to inhibit cellular proliferation to prevent restenosis due to cellular hyperplasia).

As a first step in practicing one embodiment of this invention, target DNA is selected. Preferably, the target DNA for triplex formation is selected from a gene, preferably a regulatory region of a gene whose expression is targeted for depletion. The particular target regions can come from any of a variety of regulatory regions within the DNA including, but not limited to, the promoter, the transcriptional start site or 5' of the transcriptional start site, enhancers, promoters or from transcribed portions of the gene adjacent, proximate or relatively close to the transcription initiation site. Promoter regions are preferred and methods for identifying the promoter region of a particular gene are known in the art.

Currently, regions appropriate for triplex formation require the identification of consecutive purines on one strand in a region that overlaps or encompasses the binding site for a protein-DNA complex. The mechanism of action in this case depends upon competition for the same hydrogen binding sites found in the major groove. This region is preferably upstream from the methionine start site. In general, the longer the stretch of purines, the more stable the complex and the better the binding of the oligonucleotide. Our in vitro evidence suggests that a region with at least fifteen purines with two pyrimidine interruptions is adequate for triplex formation.

Where the target site is within a promoter, formation of a triplex in that promoter region adjacent to a protein binding region can also disrupt binding, therefore, sites around the DNA-protein interaction domains are also suitable target sites (for example, see Maher, et al., (1991), “Inhibition of DNA-protein Interactions,” in Prospects for Nucleic Acid Therapy of Cancer and AIDS, pp. 227-242, Wiley Liss, Inc.). The effect of binding of the oligonucleotide to a target region within a promoter could interfere with the proper association of the promoter and transcriptional activators. When a specific target sequence is available for binding to protein it is likely that it will also allow binding to an oligonucleotide.

A second way that triple helix formation could inhibit gene expression is to act as a physical block to the translocation of the polymerase complex. The target area in this case is positioned after the transcriptional start site. There are in vitro examples of triplexes serving as physical blocks to transcription.

A third way that triplexes might disrupt transcription is by introducing structural changes that affect the ability of the polymerase complex to initiate transcription, thus, targeting areas upstream but close to the transcriptional initiation site may allow testing of this type of transcription initiation.

To summarize, the more that is known about a promoter, the better the chance of altering gene expression. In many cases, the site of binding of an activator protein is known and the sequence conforms to the definition of purine rich (a duplex region of at least 14 purines with preferably fewer than two pyrimidine bases). The term GA rich refers to at least 10 of the 17 purines being either G or A. This sequence may be part of an enhancer or adjacent to a known protein binding domain. When little is known about the regulation of the gene, any purine stretch is a potential target that can be tested in vitro using the exemplary methods disclosed in the examples that can be adapted to any of a variety of target duplex DNA sequences and potential triplex forming oligonucleotides.

Methods for sequencing a particular gene or the upstream regulatory regions of a gene are known in the art; therefore, these methods will not be detailed here. Once the 5' start site and/or regulatory regions of the gene are identified, this region is sequenced and reviewed. A target region of preferably between 15-100 nucleotides of double stranded DNA is selected and more preferably a region of between 15-50 nucleotides of double stranded DNA is selected. One or more oligonucleotides are designed to bind to one or more regions of the target double stranded DNA.

The sequence of the oligonucleotide will depend on the sequence of the targeted double stranded DNA. While sequences of purines are preferred for triplex formation using unmodified nucleoside bases, oligonucleotides having, for example, N2-protonated deoxycytidine modifications can be used to make pyrimidine based triplex forming oligonucleotides that escape pH dependent binding to DNA duplexes (Karpwezki, et al., supra). It is believed that base modifications to accommodate triplex formation by pyrimidine triplex forming oligonucleotides can be combined with the cationic phosphoramidates of this invention and that
such modifications can be made and tested in view of this disclosure without undue experimentation. All triplex forming oligonucleotides are expected to benefit from the stability imparted by the cationic phosphoramidate internucleoside linkages of this invention.

The purine triplex motif adenosine bases in the target duplex can interact with either A or T residues. The TFOs (triplex-forming oligonucleotides) in this invention are designed to be GA rich instead of GT rich, although either motif will benefit from the use of the cationic backbone modification. The T residues in a GT rich ODN may encourage intracellular guanine quartet formation by providing points of stable hairpins. This idea is supported by examination of human telomeres, structures known to form G quartets (Williamson, et al., Cell, 59:871–880, 1989). Human telomeres contain numerous repeats of the sequence TTAGGG, exhibiting an enhancement of T residues and a paucity of A residues.

Among the trials that serve to predict efficacy for triplex forming oligonucleotides are a series of in vitro trials using the targeted duplex and purine motif triplex forming oligonucleotides. As one way of identifying preferred oligonucleotides and as disclosed in Example 5, apparent binding constants for each triplex forming oligonucleotide may be determined and the triplex forming oligonucleotide with the best binding is used for in vitro and in vivo analysis of transcriptional inhibition.

The length of the oligonucleotide can vary. Preferably, the oligonucleotides are from about 15 bases in length to about 50 bases in length with a most preferable length of about 15 to about 30 and still more preferably of about 17 to about 24 bases.

Once the sequence of the oligonucleotide has been designed or selected, the oligonucleotide can be synthesized. There are a variety of methods known in the art for synthesizing oligonucleotides. Most notably, oligonucleotides can be synthesized manually or using automated DNA synthesizers employing H-phosphonate monomers and chemistry. These methods are known in the art and for that reason will not be detailed here. The oligonucleotides of this invention incorporate modified internucleoside linkages. Cationic phosphoramidates are used to replace at least one phosphodiester linkage. In a preferred embodiment, the cationic phosphoramidate is N,N-diethyl-ethylendiamine; however, other classes of ethylenediamines are also contemplated in this invention, including, but not limited to, ethylenediamine and N-ethyl-ethylendiamine. Essentially, any compound that can be added by oxidative amidation could be tested using the guidelines provided in this disclosure. Other cationic phosphoramidates suitable as substitutes for phosphodiester internucleoside linkages include, but are not limited to, diaminobutane and polylysine.

This invention incorporates cationic phosphoramidate linkages during oligonucleotide synthesis. Oligonucleotides can be prepared with a single cationic phosphoramidate internucleoside linkage, or up to 100% of the internucleoside linkages can be prepared using cationic phosphoramidates. In a preferred embodiment, the range of substituted linkages is between 30–100% and in another particularly preferred embodiment, the range is between 60–100% cationic phosphoramidate substituted linkages.

Example 1 details a preferred method for preparing oligonucleotides with cationic phosphoramidate linkages, where the cationic phosphoramidate is N,N-diethyl-ethylendiamine. Since the class of cationic phosphoramidates of this invention are used to create internucleoside linkages by oxidative amidation, the chemistry for synthesizing oligonucleotides using other cationic phosphoramidates will not differ significantly and those of ordinary skill in the art of oligonucleotide modifications will be readily able to prepare a variety of other oligonucleotides with other cationic phosphoramidate linkages without undue experimentation.

A number of oligonucleotides were prepared that varied in the amount of cationic phosphoramidate internucleoside linkages, as illustrated in FIG. 1. Oligonucleotides with these modified cationic linkages were compared to oligonucleotides containing neutral linkages. FIG. 1A illustrates the structure of a preferred cationic phosphoramidate modified used in this study. The methoxyethylamine derivative was used to generate oligonucleotides with neutral internucleoside linkages. Solubility of the derivative was enhanced through hydrogen bonding between water and the ether oxygen. The N,N-diethyl-ethylendiamine derivative is shown uncharged but is protonated (and, hence, positively charged) at physiologic pH.

In one example of this invention, the sequence of the target region was derived from the enhancer of the GS17 gene of Xenopus laevis (Ovesen et al., Development, 115:649–655, 1992). Duplex 1 (SEQ ID NO:1) is 30 bp in length and contains a 17 bp purine rich region. The presence of pyrimidines on the purine rich strand makes this region an imperfect triplex forming consensus sequence but likely represents a more commonly available target than the 25–35 bp target region with the myc gene containing an unmodified polyuridine-polyurimidine sequence that is often used for in vitro studies.

The oligonucleotides used in one example in this study are identical in nucleic acid sequence and correspond to SEQ ID NO:2. Each oligonucleotide illustrated in FIG. 1B contains either no modifications (oligo U-1), neutral internucleoside linkages (N-1 and N-2), or cationic phosphoramidate internucleoside linkages (P-1 through P-4). In designing the TFOs, thymidine was chosen to interact with the 2:3:1 G inversion based on a report demonstrating significant T:C:G binding (Durland et al., Nucleic Acids Research, 22:3233–3240, 1994). N-1 and N-2 represent identical oligonucleotides containing 7 and 11 neutral methoxyethyl phosphoramidate linkages, respectively. These neutral bonds are represented by ∗ in FIG. 1. ODNs P-1 through P-4 contain increasing numbers of positively charged N,N-diethyl-ethylendiamine phosphoramidate linkages, from 9 to 16. The positive linkages are represented as + in FIG. 1. The extent of modification of each ODN is indicated to the right of the sequence.

The positive internucleoside linkage was made by amidation of the corresponding hydrogen phosphonate diester with N,N-diethyl-ethylendiamine as described in Example 1. N,N-diethyl-ethylendiamine is preferred to N-ethyl-ethylendiamine or ethylenediamine for triplex formation under physiologic conditions because it possesses both primary and tertiary amine moieties. The primary and tertiary amine moieties allow for a more specific reaction and minimize oxidative amidation at each end of the diamine. Oxidative amidation has the potential for crosslinking oligonucleotides. Moreover, the pKa of a tertiary amine is higher than that of the corresponding primary or secondary amine. The resulting increase in pKa of the ODN produces a greater net positive charge at physiologic pH.

Once target duplex has been identified and oligonucleotides are synthesized, synthetic target duplex can be pre-
pared to test the ability of the synthesized oligonucleotides to bind to the target duplex under physiologic conditions. The oligonucleotides provided in FIG. 1 were tested in vitro and the ability of the oligonucleotides containing neutral internucleoside linkages to form triplex structures was compared to the triplex forming ability of oligonucleotides with cationic phosphoramidate substituted linkages. Example 3 details methods for testing triplex formation in vitro using both negatively charged, neutral, and positively charged oligonucleotides.

Experiments were performed to test for the ability of a particular modified oligonucleotide to bind to duplex DNA. Example 4 details the results of binding studies to assess triplex formation between an unmodified negatively charged oligonucleotide and oligonucleotides N-1 and N-2, where negative charges were replaced with uncharged phosphoramidates. These experiments were performed in 10 mM Mg$^{2+}$ without potassium with ODN concentrations ranging from 20 nM to 2 µM. Triplex formation was found to be more efficient as negative phosphodiester linkages were converted to neutral phosphoramidate derivatives. Without limiting the scope of this invention, the most likely explanation for this observation is that the oligonucleotides containing the uncharged phosphoramidates carry a significant net negative charge and are repulsed, although less than unmodified oligonucleotide U-1, by the negative charge density of Duplex I.

The requirement for relatively high concentrations of Mg$^{2+}$ in order to form triplex DNA using the neutrally modified ODNs supports the hypothesis that charge remains a major factor affecting triplex strand association. Increases in triplex formation can result from either increased affinity for the DNA duplex or decreased tendency toward formation of guanine quartets. The latter could increase the effective concentration of the TFO. Any ODN modification, whether directed at the bases, the sugar moiety, or the phosphate backbone, has the potential to affect either or both of these equilibria.

In contrast to the oligonucleotides containing neutral modifications, nearly complete conversion of duplex DNA to triplex DNA was observed with the cationic phosphoramidates tested (see Example 5). The formation of triplex DNA in the presence of K$^+$ is important in determining whether the cationic phosphoramidates of this invention are useful in vivo. Example 6 details the effect of potassium chloride on triplex formation using the cationic phosphoramidates of this invention. The cationic phosphoramidate substituted oligonucleotides were significantly better at forming triplex structures than their unmodified or neutral counterpart. Moreover, increasing the amount of cationic internucleoside linkages increased the amount of triplex formation.

Triplex formation in the presence of LiCl, NaCl, and KCl was evaluated to determine whether the observed triplex inhibition was potassium-specific or resulted from increases in the ionic strength of the solution. Example 7 discloses methods for assessing the effect of various monovalent cations at concentrations of 40, 80, and 120 mM on triplex formation. Triplex formation was at least affected by increasing concentrations of LiCl. NaCl caused an intermediate level of inhibition and KCl had the greatest effect on triplex formation.

The effect of extensive cationic phosphoramidate modification on triplex formation was examined with oligonucleotides containing 88% modified linkages and 100% modified linkages (see Example 8). The ability of these oligonucleotides to associate with target duplex DNA was compared to unmodified oligonucleotides with the same base sequence at 130 mM K$^+$ and 1 mM Mg$^{2+}$ concentrations that approach physiologic salt concentrations. Oligonucleotides with extensive cationic phosphoramidate linkage modification demonstrated an improved capacity to form triplex structures.

The oligonucleotides were also tested for their capacity to form triplex DNA structures under stringent salt conditions (see Example 9). Results demonstrated that triplex formation with oligonucleotides containing increasing levels of cationic phosphoramidate linkages were less sensitive to K$^+$ concentration. Unexpectedly, almost 90% of Duplex I was in the triplex form at 250 mM KCl. Lane 13 shows Duplex I in the absence of any TFOs. Increasingly positively-charged internucleoside linkages from 11 of 16 (69%) in P-2 to 14 of 16 (88%) in P-3 is associated with a significant decrease in potassium-mediated inhibition of triplex formation. In addition, and also unexpectedly, although magnesium is known to stabilize traditional triplex DNA, the results of Example 10 indicated that oligonucleotides with extensive cationic phosphoramidate modifications were essentially unaffected by Mg$^{2+}$ at physiologic K$^+$ concentrations.

This invention demonstrates that positively-charged ODNs with cationic phosphoramidate modified internucleoside linkages are significantly more effective in forming triplex DNA than the corresponding unmodified or neutrally-modified compounds. When an ODN is designed to interfere with intracellular nucleic acid metabolism, specificity becomes a crucial issue. Nonspecific electrostatic interactions between positively-charged ODNs and nontarget duplex DNA are a potential concern with these compounds. Example 11 discloses methods for testing the specificity of the oligonucleotide for its target using unmodified and cationic phosphoramidate modified oligonucleotides.

A second double-stranded DNA strand nonspecific for the oligonucleotide (SEQ ID NO:3) was used to demonstrate the specificity of the oligonucleotides of this invention for their target duplex DNA. Duplex II, illustrated in FIG. 1, is a random, purine-pyrimidine 30 bp sequence with 5 inversions. This duplex is used to measure nonspecific binding of a positively-charged ODN to a DNA duplex and, as disclosed in the examples, is used to assess triplex forming oligonucleotide specificity. The sequence is found in the GS17 promoter, just downstream of the oligonucleotide target site. This sequence is random with respect to the selected oligonucleotides designed to bind to the GS17 promoter and can therefore be used to assess specificity for any oligonucleotide that is targeted for triplex formation at a particular duplex DNA sequence; however, those skilled in the art will also appreciate that other duplex sequences can be used to assess the binding specificity of other oligonucleotides and that it is possible to produce nonspecific duplex DNA fragments from regions near or adjacent to a given targeted duplex DNA sequence to assess nonspecific binding, as well.

To further investigate these nonspecific interactions under salt concentrations more closely resembling in vivo conditions, the experiment was repeated in 130 mM KCl. Oligonucleotides with extensive cationic phosphoramidate modifications had reduced nonspecific binding to Duplex II under physiologic concentrations of K$^+$.

The inability of purine rich ODNs to form triplex DNA in the presence of physiologic concentrations of K$^+$ is a major hurdle in the development of oligonucleotide compounds as regulators of gene expression. This invention describes a
class of positively charged ODNs that can efficiently form triplexes in the presence of physiologic and even supra-physiologic levels of potassium.

Replacing negative phosphodiester bonds with positively-charged phosphoramide linkages resulted in compounds that more efficiently formed triplexes with double stranded target DNA. A progressive increase in triplex forming ability was observed with increasing positive charge. As discussed above, this was demonstrated by: (1) a decrease in $K_d$ with increasing ODN modification, (2) a decreased sensitivity to the inhibitory effects of monovalent cations on triplex formation, and (3) a diminished requirement of Mg$^{2+}$ in order to form triplex DNA. Without intending to limit the scope of this invention, because of the Mg$^{2+}$ independence of triplex association observed using oligonucleotides with at least 80% cationic phosphoramide modified linkages, it is likely that electrostatic attraction between the ODN and the target duplex is a predominant force stabilizing the triplex. In addition to electrostatic attraction, it is possible that positively charged ODNs will have a reduced capacity to form guanine quartets and that both mechanisms could act to promote triplex formation.

A second benefit of the oligonucleotides of this invention as compared to other positively-charged oligonucleotides is the reduced levels of nonspecific association between cationic phosphoramide modified oligonucleotides with non-target duplex DNA. Under physiologic levels of potassium chloride, the nonspecific binding of oligonucleotides with greater than 80% cationic phosphoramide modification was nearly undetectable. Thus, under conditions that more closely approach physiologic, the extent of nonspecific triplex formation becomes minimal.

A third benefit of the oligonucleotides of this invention is that the cationic phosphoramide modified internucleoside linkages, like other internucleoside modifications, are resistant to RNase activity. This resistance enhances stability of the oligonucleotide in the cell.

The cationic phosphoramide modified oligonucleotides of this invention can additionally incorporate other oligonucleotide modifications known in the art. For example, the oligonucleotide may include base modifications including but not limited to, 6-thioguanine base substitutions for purine:purine motif triplex forming oligonucleotides (Olivas, W. M. and Maher, L. J. III, Nucleic Acids Res., 23:1936–1941, 1995, and Rao, T. S., et al., Biochemistry, 34:765–772, 1995). Alternatively, N3 protonated deoxycytidine analogs can be incorporated into oligonucleotides for pyrimidine:purine motif triplex forming oligonucleotides (see Frechler, et al., Nucleic Acids Res., 14:5399–5407, 1986). Other base modifications that could also be incorporated into the oligonucleotides of this invention include, but are not limited to, methyl cytidine and alkynyl base modifications.

In addition, the oligonucleotides can be labeled with tags such as enzymatic tags, radiolabelled tags, fluorescent tags, including, but not limited to, fluorescein, Rhodamine Green, Rhodamine Red, Texas Red, Biotin, DNP, and the like (See, for example, FluorReportol Oligonucleotide Amine Labeling Kits provided by Molecular Probes, Eugene, Oregon). Suitable chromophores are those with amine groups that can be added by oxidative amidation. In addition, chromophores with isothiocyanate groups can also be used. These can be added to the oligonucleotide during synthesis using oxidative amidation of H-phosphonate diester with diamine or can be reacted with available amines. Similarly, the oligonucleotides of this invention can be coupled to hapten or any of a variety of fluoroprobes that do not interfere with the binding characteristics of the modified oligonucleotides for their target DNA. These oligonucleotides can be used to monitor binding of the labeled oligonucleotide to duplex DNA.

Once the oligonucleotides have been tested for their binding affinity to target duplex DNA, the cationic phosphoramide modified probes of this invention can be tested to determine in vivo efficacy. There are a variety of methods known in the art for introducing oligonucleotides into cells both in culture and in an organism. These methods include, but are not limited to, microinjection, liposome technology including antibody directed liposomes or antibody directed oligonucleotides as well as non-targeted liposomes.

As a preferred testing regime, the oligonucleotides are microinjected into cells obtained from animal or man. The cell type chosen is determined by the presence or absence of the selected target DNA in the cell type. As one example, the target DNA is a sequence of duplex DNA from the enhancer region of the GSI7 gene of Xenopus laevis, corresponding to Duplex I of FIG. 1, and the oligonucleotides tested have the nucleic acid sequence and base modifications of oligonucleotides P-3 and P-4 of FIG. 1. Although embryos and oocytes of Xenopus laevis are used in these studies and the target duplex DNA is the GSI7 gene, oligonucleotide delivery to a variety of cells is known in the art and those skilled in the art will recognize the applications of these examples to a wide variety of target duplex DNA sequences resident in a variety of eukaryotic cells.

There are two factors that are assessed in determining the ability of a particular oligonucleotide to modulate an inhibitory effect on transcription; toxicity of the oligonucleotide to the cell and effect of the oligonucleotide on transcriptional inhibition. Example 12 details methods for assessing the cellular toxicity of the oligonucleotide and Example 13 discusses methods for assessing the effect of the cationic phosphoramide modified oligonucleotide on transcriptional inhibition. Reduction in transcription rates is analyzed by measuring either mRNA levels produced by the targeted genes or by assaying for levels of the protein product encoded by the mRNA. Methods for measuring both mRNA and protein production from a cell are well known in the art.

Once an inhibitory effect has been observed in vitro, it is contemplated that the oligonucleotides of this invention can be used to impart a transcriptional impact on eukaryotic cells ex vivo and in vivo. Thus, in one embodiment of a method for inhibiting expression of a gene in an animal, a cell sample is removed from an animal and processed to produce a cell sample that can be treated with the oligonucleotides of this invention using, for example, the methods disclosed in PCT patent application WO 9415524 to Anderson, et al. In a second embodiment, the oligonucleotides of this invention are encapsulated in lipids, using the methods and techniques disclosed in PCT published patent applications WO 9517373 to Ciccarene, et al., WO 9014074 to Abai, et al., U.S. Pat. No. 5,194,654. The lipid/oligonucleotide formulation is delivered to the animal or human by parenteral or oral routes of administration including, but not limited to, intravenous, subcutaneous, intramuscular, mucosal introduction such as through the use of aerosols or drips, and alimentary introduction. Those skilled in the art will recognize that the choice of delivery will be determined by the source of cells containing duplex DNA targeted for gene expression inhibition.

Those skilled in the art will recognize that there are a variety of modifications which can be made to oligonucle-
otides to facilitate cellular uptake. Similarly, a number of studies have been performed that address methods for delivery of a variety of oligonucleotides to cells in a mammal.

In another aspect of this invention, this invention relates to the use of the oligonucleotides with cationic phosphoramide internucleoside linkages to hybridize to DNA or RNA. The nucleic acid can be genomic or extrachromosomal, including, but not limited to plasmids, episomes, and linear nucleic acid. For example, the oligonucleotides of this invention can be used as probes to bind to denatured double stranded DNA or to denatured RNA. The oligonucleotides of this invention bind to single stranded RNA and DNA or denatured double stranded DNA under the same conditions that are well known in the art for oligonucleotides with naturally occurring internucleoside linkages. One advantage of the oligonucleotides of this invention is the ability of the oligonucleotides of this invention to bind to DNA or RNA under conditions where oligonucleotides with naturally occurring internucleoside linkages would not bind (e.g., denaturing concentrations of salt, such as a reduced concentration of salt, increased temperature, in the presence of chaotrope agents, or a combination of denaturing concentrations of salt and one or more chaotropic agents).

The term “denatured nucleic acid” or “DNA—DNA denaturing conditions” is used herein to refer to conditions that promote the conversion or dissociation of nucleic acid, such as DNA or RNA from a double-stranded state (double stranded in whole or in part) into a single-stranded state. This conversion can be accomplished through a variety of methods including, but not limited to, heat, denaturing concentrations of salt, such as lower salt concentrations, detergents, or the addition of chaotropic agents such as for example, formamide, urea or a guanidinium salt, and the like. Preferred denaturing concentrations of salt, detergents or chaotropic agents will vary and those skilled in the art of nucleic acid association and dissociation kinetics will be able to optimize the DNA—DNA denaturing conditions as needed. In one example, the concentration of urea used varied from about 0.5M to about 7M.

Advantageously, the oligonucleotides of this invention can bind to single stranded DNA or RNA, including denatured double-stranded DNA, under conditions that would not support the formation of a hybridization complex if oligonucleotides with only phosphodiester internucleoside linkages were used. The oligonucleotides of this invention that are capable of hybridizing to DNA or RNA under DNA—DNA duplex denaturing conditions comprise at least one cationic phosphoramide internucleoside linkage and preferably at least about 30% cationic phosphoramide internucleoside linkages. A preferred oligonucleotide includes at least about 50% to 100% cationic phosphoramide internucleoside linkages. A preferred type of cationic phosphoramide internucleoside linkage for duplex formation includes oligonucleotides with ethylenediamine-class phosphoramide internucleoside linkages, and in particular oligonucleotides with N,N-diethyl ethylenediamine phosphoramide internucleoside linkages.

The cationic phosphoramide-modified oligonucleotides used as probes for binding to DNA and RNA under DNA—DNA denaturing conditions are preferably at least about 10 nucleotides and preferably at least about 15 nucleotides in length. In one embodiment, the oligonucleotides are less than about 50 nucleotides in length and preferably less than 100 nucleotides in length.

In one example, using thermal denaturation studies where salt concentration was varied to analyze the stability of duplex formation using the oligonucleotides of this invention, in 150 mM NaCl, unmodified oligonucleotides and cationic oligonucleotides both had Tm’s of about 50°C, however, as salt concentration was lowered to 0 mM NaCl, unmodified oligonucleotides exhibited a decreasing Tm while the oligonucleotides of this invention with 100% modified internucleoside linkages were unaffected. Thus, the formation of duplex using oligonucleotides with the internucleoside linkages of this invention forms complexes with target nucleic acid independent of salt concentration (see Example 15).

In a preferred method for detecting nucleic acid using the modified oligonucleotides of this invention, the oligonucleotides are contacted with single-stranded or substantially single-stranded nucleic acid (i.e., where the double-stranded character of the nucleic acid is sufficiently disrupted to permit duplex formation with the oligonucleotide), such as DNA or RNA under conditions suitable for the oligonucleotides of this invention to form duplex structures with the nucleic acid. In a preferred embodiment, the hybridizing step is performed under DNA—DNA duplex denaturing conditions. Following hybridization, the hybridization complex is detected using any of a variety of methods known in the art. For example, the oligonucleotides of this invention can be modified before, after or during oligonucleotide synthesis to include one or more tagged nucleotides such as nucleotides incorporating a radioactive tag, a fluorescent tag or an enzyme-labeled nucleotide. The detection step can then include detecting the tagged oligonucleotide. Methods for incorporating tagged nucleotides or for tagging oligonucleotides are known in the art and in one example are taught, for example, by Sambrook et al. (Molecular Cloning, A Laboratory Manual, 1989 Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.).

In a preferred embodiment, the nucleic acid detected under DNA—DNA duplex denaturing conditions is part of a sample of genomic nucleic acid, such as DNA from a host. The genomic nucleic acid can be obtained from bacteria, virus, animals, or plants.

In a preferred embodiment of this invention, the oligonucleotides including cationic phosphoramide internucleoside linkages, such as ethylenediamine-class internucleoside linkages, are hybridized to nucleic acid under DNA—DNA duplex denaturing conditions to detect nucleotide polymorphisms within a gene, including single-nucleotide polymorphisms. There is interest in being able to distinguish between highly similar target nucleic acid sequences. Oligonucleotide probes differ in their ability to efficiently recognize a similar target. The length of the oligonucleotide as well as the nucleic acid content of the oligonucleotide, the modifications to the oligonucleotide and the hybridization conditions all affect the ability of an oligonucleotide to differentiate between very similar target sequences. For example, oligohistidine peptide nucleic acids (oligohistidine-PNA) have been used to detect and/or differentiate between highly similar target sequences (Orum et al. BioTechniques 19:472—480, 1995). These oligonucleotides could bind to a DNA target in the presence of 2M urea but, unlike the oligonucleotides of this invention, the oligohistidine-PNAs were unable to bind and/or to differentiate target DNA in the presence of 2M urea and denaturing concentrations of salt.

The oligonucleotides of this invention can be used to quantitate gene copy number, quantitate the level of gene expression and to detect polymorphisms in target nucleic acid.

In experiments to detect the ability of the oligonucleotides of this invention to bind to DNA under DNA—DNA dena-
turing conditions, labeled cationic phosphoramidate-containing oligonucleotides capable of binding to λ phage DNA were combined with salmon sperm DNA with or without λ phage DNA under DNA—DNA denaturing conditions. The sample without λ phage DNA had increased scatter as evidenced by an increased signal to noise ratio as compared to the sample containing λ phage DNA. This provided evidence that the oligonucleotide was binding specifically to the λ phage DNA (see Examples 15 and 16).

All references and publications cited herein are expressly incorporated by reference into this disclosure. Particular embodiments of this invention will be discussed in detail and reference has been made to possible variations within the scope of this invention. There are a variety of alternative techniques and procedures available to those of skill in the art which would similarly permit one to successfully perform the intended invention.

EXAMPLE 1
Oligonucleotide Synthesis

Modified ODNs were synthesized on an ABI model 391 PCR-mate DNA synthesizer using hydrogen phosphonate chemistry Froehler et al., Nucleic Acids Res., 14:5399–5407, 1986. All reagents used for automated DNA synthesis were obtained from Glen Research (Sterling, Va.). To generate unmodified phosphodiester bonds, hydrogen phosphate diesters were oxidized for 4 minutes with freshly prepared 5% iodine in THF-pyridine:water (15:2:2) and then 3 minutes with the same solution diluted 1:1 with 8% TEA in THF-water (43:3). Oxidative amidation of hydrogen phosphate diesters was performed manually using a 10% solution of either 2-methoxyethamine or N,N-diethyl-ethylenediamine (Aldrich) in anhydrous CCl₄, as previously described (Dagle et al., Nucleic Acids Res., 18:4751–4757, 1990). ODNs containing both phosphodiester and phosphoramidate linkages were synthesized in blocks. The desired number of 3’ residues was first coupled and then either oxidized or oxidatively amidated. The next block of residues was then individually condensed and subsequently oxidized or oxidatively amidated. Purification of ODNs was performed as described previously (Dagle, Nucl. Acids Res., supra). Following Sephadex G-25 column chromatography (Pharmacia) to remove small molecular weight impurities, ODNs were dissolved in sterile water and quantified by UV spectroscopy. It was assumed that the modified linkages did not significantly affect the extinction coefficients of the individual ODNs. ODN heterogeneity was assessed using denaturing reversed-polarity gel electrophoresis. The presence of a distinct band was indicative of a homogenous oligonucleotide population.

EXAMPLE 2
Oligonucleotide Labeling

Target duplexes are formed from a 1:1 mixture of complementary unmodified 30-mer ODNs which are heated to 80°C for 5 minutes and allowed to slowly cool to room temperature. These duplexes were 5’ end-labeled with T4 poly nucleotide kinase (Promega, Madison, Wis.) and γ-³²P-ATP (6000 Ci/mmol, Amersham). 2 pmole of DNA duplex (4 pmol of 5’ ends) was incubated at 37°C for 45 minutes under the following conditions: 70 mM Tris pH 7.6, 10 mM MgCl₂, 5 mM DTT, 8 pmole ATP and 3 U T4 polynucleotide kinase in a total of 10 μl. The reaction was stopped by the addition of 90 μl of 7.5 M ammonium acetate and subsequent phenol:chloroform extraction. The aqueous phase was added to 20 μg of carrier glycogen. To this was added 300 μl ethanol and the solution was kept at −70°C for at least 30 minutes. Following centrifugation, the pellet was resuspended in water to a concentration of 10 nM.

EXAMPLE 3
Triplex Formation Assay

Triplex formation was initiated by the addition (in order) of 5 μl H₂O, 2 μl 5x buffer (100 mM Tris-HCl pH 7.5, 0 to 50 mM MgCl₂), 1 μl labeled DNA duplex (2–10 fmole/μl), 1 μl yeast tRNA (1 mg/ml), and 1 μl triplex forming ODN. The buffer solutions and ODN concentrations were altered to examine triplex formation under various conditions. The mixtures were incubated at ambient temperature overnight. After the addition of 1 μl of gel loading buffer (0 to 10 mM MgCl₂ to match the assay conditions, 20 mM Tris-HCl pH 7.5, 50% glycerol and 0.05% each bromophenol blue and xylene cyanol), the samples were loaded onto a 15% polyacrylamide (acrylamide:bisacrylamide= 100:1) gel and analyzed by nondenaturing gel electrophoresis. The electrophoresis buffer was TMB (10 mM Tris base, 90 mM boric acid, and 10 mM MgCl₂). Electrophoresis was performed at 4°C for 4–6 hours. The gel was dried under vacuum and exposed at −70°C to Kodak X-OMAT AR film using an intensifying screen. The amount of radioactivity present in the duplex and triplex forms was determined by electronic autoradiography (InstantImager, Packard, Downers Grove, Ill.). The fraction of target duplex bound by a TFO, θ, was calculated using the equation:

\[ \theta = \frac{S_{duplex} - S_{triplex}}{S_{duplex}} \]

where \( S_{duplex} \) and \( S_{triplex} \) represent the electronic autoradiographic signal for the duplex and triplex bands, respectively. The Kd for an ODN in triplex formation was determined from the concentration of the compound which causes ½ of the target duplex to shift to the triplex form.

EXAMPLE 4
Effect of Charge Neutralization on Triplex Formation

Negative charges were removed from the oligonucleotide of FIG. 1 and replaced with neutral charges using the methoxyethamine derivative illustrated in FIG. 1. ODNs-1 and N-2 were compared with their unmodified counterpart, U-1, for the ability to form triplexes with Duplex I. These experiments were performed in 10 mM Mg²⁺ and no K⁺, with ODN concentrations of 0 nM, 20 nM, 200 nM, and 2 μM for each of the three oligonucleotide samples. A gel shift assay was performed. Triplex formation was seen with increasing concentrations of U-1, N-1, and N-2. Triplex formation was more efficient as negative phosphodiester linkages were converted to neutral phosphoramidate linkages. 2 μM of unmodified oligonucleotide shifted a small amount of Duplex I. With decreasing negative charge, an increasing fraction of Duplex I was shifted to the more slowly migrating triplex form.

EXAMPLE 5
Triplex Formation with ODNs Containing Positively-Charged Internucleoside Linkages

ODNs P-1 and P-2 were also compared to the unmodified oligonucleotide, U-1, in a gel shift assay. The assay was
performed in 1 mM MgCl$_2$ and no K$^+$ and demonstrated a significant increase in triplex formation in those oligonucleotides with positively-charged internucleoside linkages. Interestingly, U-1 is unable to shift more than 15% of Duplex I to a more slowly migrating form at 2 μM, the highest ODN concentrations examined. No distinct triplex band was visible. In contrast, nearly complete conversion of duplex to a distinct triplex was observed with oligonucleotides P-3 and P-4 over the 2 nM to 2 μM concentration range tested (samples were tested at 2 nM, 20 nM, 200 nM, and 2 μM). The Kd for U-1 could not be measured under these conditions. The Kd for P-1 was about 8×10$^{-7}$ M, while that of P-2 was about 8×10$^{-8}$ M.

EXAMPLE 6

Triplex Stability in the Presence of Potassium

At 1 mM MgCl$_2$ and 0 mM KCl, ODNs containing several positively-charged internucleoside linkages were superior to their unmodified counterpart in triplex formation. The formation of triplex DNA in the presence of K$^+$, however, is crucial if these oligonucleotides are useful in inhibiting gene expression in vivo. Positively-charged ODNs P-1 and P-2 are compared to U-1 at 2 concentrations each of both MgCl$_2$ and KCl. The ODN concentration used was 200 nM. Oligonucleotide U-1 was examined in 1 mM and 10 mM MgCl$_2$, while oligonucleotides P-1 and P-2 were examined at the more stringent conditions of 0.1 mM MgCl$_2$ and 1 mM MgCl$_2$. A trace amount of shifted duplex was seen by gel shift assay with ODN U-1 at 10 mM MgCl$_2$, but not at 1 mM MgCl$_2$. In contrast, ODNs P-1 and P-2 formed triplex DNA at both 0.1 mM MgCl$_2$ and 1 mM MgCl$_2$. In both cases, there was only a slight increase in triplex at 1 mM MgCl$_2$ compared to 0.1 mM MgCl$_2$. The response of P-1 compared to P-2 with respect to K$^+$ inhibition was striking, especially at 1 mM MgCl$_2$. Triplex DNA formation seen with P-2 at 1 mM MgCl$_2$ and 100 mM KCl was approximately 80% of that seen without KCl. In contrast, triplex DNA formation with P-2 at 0.1 mM MgCl$_2$ and 100 mM KCl was reduced to less than 60% of that seen without KCl. This non-competitive inhibition of triplex formation is similar to that reported by other laboratories with unmodified ODNs (Milligan, et al., Nucleic Acids Res., 21:327–333, 1993; Cheng, A. J., et al., Nucleic Acids Res., 21:5630–5635, 1993; Olivas, et al., Biochem., supra).

EXAMPLE 7

Triplex Formation in the Presence of Monovalent Cations

Triplex formation in the presence of LiCl, NaCl, and KCl was evaluated to determine whether the observed triplex inhibition was potassium-specific or resulted from increases in the ionic strength of the solution. ODN P-2 was studied at a concentration of 2 μM with a Mg$^{2+}$ concentration of 0.1 mM using 40, 80, and 120 mM concentrations of LiCl, NaCl, or KCl. Triplex formation was least affected by increasing concentrations of LiCl. NaCl caused an intermediate level of inhibition and KCl had the greatest effect on triplex formation. This pattern of monovalent cation inhibition was identical to that seen with unmodified ODNs suggesting a similar mode of association of the oligonucleotide to the duplex.

EXAMPLE 8

Triplex Formation Using Oligonucleotides With Extensive Cationic Phosphoromidate Modifications

The effect of extensive ODN modification on triplex formation was examined with oligonucleotides containing 88% modified linkages or 100% cationic phosphoromidate modified linkages (P-3 and P-4, respectively). The ability of ODNs P-3 and P-4 to associate with Duplex I was compared to that of compounds U-1 and P-2. The assay was performed using 130 mM K$^+$ and 1 mM Mg$^{2+}$, concentrations that approximate physiologic salt concentrations. The ODN concentrations used were 20 nM, 200 nM, and 2 μM. Samples were processed with the various oligonucleotides at the various concentrations and a sample containing no oligonucleotide was used as a background control. Shading in the control lane, located in the region where the triplex band migrated, was subtracted from the triplex bands during data analysis. Triplex formation with U-1 was essentially undetectable under the concentrations tested. Both P-3 and P-4 showed a greater affinity and, therefore, improved stability, for Duplex I than did P-2. The dissociation constants for triplex formation were 8×10$^{-7}$ M for P-2, 1×10$^{-7}$ M for P-4, and 7×10$^{-8}$ M for P-3. The migration of the triplex formed with P-3 was slightly slower than that with P-2, a result of the increased cationic nature of P-3.

EXAMPLE 9

Triplex Formation Under Stringent Salt Conditions

The effect of increasing KCl concentrations on triplex formation was also assessed. Oligonucleotides P-2 and P-3 were incubated with Duplex I under increasing concentrations of KCl (0, 50, 100, 150, 200, and 250 mM). The concentration of MgCl$_2$ was held at 1 mM and the ODN concentration was 2 μM. Increasing concentrations of KCl gradually reduced the amount of triplex DNA observed for oligonucleotide P-2; however, significant triplex formation was observed at physiologic levels of potassium. The inhibitory effect of KCl appeared to plateau at 200 mM, as no further triplex inhibition was observed at 250 mM. Triplex formation with ODN P-3 was significantly less sensitive to K$^+$ concentration. In fact, at 250 mM KCl, almost 90% of Duplex I remained in the triplex form. Increasingly positively-charged internucleoside linkages from 11 of 16 (69%) in P-2 to 14 of 16 (88%) in P-3 is associated with a significant decrease in potassium-mediated inhibition of triplex formation.

EXAMPLE 10

Magnesium Requirements for Triplex Formation

Mg$^{2+}$ is known to stabilize traditional triplex DNA. The effect of decreasing Mg$^{2+}$ and increasing K$^+$ on triplex formation was assessed using ODNs P-2 and P-3 by gel shift assay. In the absence of K$^+$, triplex formation with both P-2 and P-3 was essentially unaffected by Mg$^{2+}$ concentrations ranging from 0 to 1 mM. Triplex formation was greater than 80%, even in the absence of Mg$^{2+}$. Although increased, Mg$^{2+}$ concentration had no effect on the amount of triplex formed with P-3, in the absence of K$^+$ a small fraction of the triplex DNA changed to a more rapidly migrating form with increased Mg$^{2+}$. The appearance of this band with intermediate electrophoretic mobility suggests the possibility of different conformations of triplex DNA. In 130 mM KCl, triplex formation with ODN P-2 became sensitive to Mg$^{2+}$ concentration. There was a significant reduction in triplex DNA formed with oligonucleotide P-2 as Mg$^{2+}$ decreased from 1 mM to 0 mM. Triplex formation with P-3 remained essentially independent of Mg$^{2+}$ concentration in the presence of physiologic concentrations of KCl.

EXAMPLE 11

Oligonucleotide Specificity for Target DNA

Oligonucleotide P-4 was used to examine the specificity of the oligonucleotide for its target versus a random oligo-
nucleotide such as Duplex II. The nonspecific interaction of oligonucleotides U-1 and P-4 with the nontarget Duplex II in the presence of either 1 mM or 10 mM MgCl₂ and in the absence of KCl were assessed by gel shift analysis. Binding of U-1 to Duplex II was not detected and nonspecific interactions between P-4 and Duplex II were not detected at ODN concentrations of 20 nM to 200 nM of oligonucleotide. The complex formed at about equal levels at both 10 mM MgCl₂ and 1 mM MgCl₂, indicating a lack of MgCl₂ dependence in this range of ion concentration. To further investigate these nonspecific interactions under salt concentrations more closely resembling in vivo conditions, the experiment was repeated in 130 mM KCl. Under conditions approximating physiological, oligonucleotide U-1 again showed no evidence of binding to Duplex II at either 1 mM or 10 mM MgCl₂. At an ODN concentration of 2 μM, the extent of nonspecific binding of P-4 to Duplex II was greatly attenuated by increasing the K⁺ concentration of the triplex buffer. In both cases, the degree of nonspecific binding was too low for accurate quantitation. The decrease in nonspecific binding seen in 130 mM KCl was likely related to a general increase in ionic strength, however, it is also possible that there was a specific effect of K⁺.

EXAMPLE 12

Toxicity Testing of Oligonucleotide for Target Cell

As an example of a method to determine whether or not the cationic phosphoramidate modified oligonucleotides are toxic to target cells containing the target duplex DNA, Xenopus embryos were microinjected with varying concentrations of oligonucleotide. Xenopus laevis were purchased from Xenopus I (Ann Arbor, Mich.). Eggs were obtained from mature frogs fertilized in vitro and maintained in 0.1x MBSH (Colman, et al., 1984) In Hames, D. and Higgins, S. (ed.), Transcription and translation—A Practical Approach, IRL Press, Oxford, pp. 271-302). Oligonucleotides were injected by microinjection into the cytoplasm of the embryos as described by Colman (supra). To study the degradation of the oligonucleotides, at various times following injection of labeled oligonucleotides (see Example 2), the injected embryos were frozen in dry ice and thawed in 200 μl chloroform and 400 μl 0.2% SDS, the phenol layer was extracted with water twice more. Aqueous fractions were pooled, extracted once with chloroform, and dried in a SpeedVac Concentrator. The residue was resuspended in water and the amount of radioactivity was determined by scintillation counter or analyzed by electrophoresis using a 20% polyacrylamide-7M urea gel which was subsequently exposed at ~70°C to Kodak X-OMAT AR film with an intensifying screen.

To determine the cellular toxicity of the oligonucleotides, various concentrations of oligonucleotide were microinjected into matched cell samples. Defects in activation of gene expression, reflecting toxicity related to general inhibition of transcription, is assessed by looking at the activation of specific genes that are expressed at the 4000 cell stage including GS17 or EF-1α gene expression. In addition, defects in gastrulation, neurulation, or other early developmental events serves as a sensitive measure of nonspecific toxicity; therefore, this model is useful to test toxicity for oligonucleotides in general. (See Vize, et al., Methods in Cell Biol., 36:367-387, 1991). Embryos obtained as outlined in Weeks, et al. (1991), supra, have been injected with up to 10 ng of modified oligonucleotides as described herein without nonspecific toxic effects. Further, experiments to determine the toxic dose of these modified oligonucleotides in vivo are under way. Uninjected and control injected (injection buffer alone) embryos are compared to oligonucleotides injected using groups of 50 embryos in experiments performed in triplicate. Defects are noted by comparison to Xenopus laevis normal tables (Nieuwkoop, et al., 1967), Normal table of Xenopus laevis (Daudin), North Holland Publishing Co., Amsterdam).

EXAMPLE 13

Effect of Oligonucleotides on Gene Expression

The most direct way of examining the effect of a TFO on gene expression is to measure RNA levels derived from the gene that is targeted. This measurement might be made by sampling tissue and extracting RNA followed by northern blot analysis, RNase protection assays, or Reverse transcriptase PCR. All techniques are well known in the art and discussed in detail in Molecular Cloning: A Laboratory Manual (Sambrook, et al., 1989, Cold Spring Harbor Laboratory, Cold Spring, N.Y.).

Alternatively, levels of RNA can be examined by in situ analysis of tissues of interest. These techniques are also known in the art. A less direct, but often useful, way to look at transcription is to assay for the protein products of the gene. This would include Western analyses, polyclonal antibodies, gel electrophoresis, and a variety of antibody based strategies including radioimmunoassays and enzyme linked immunosorbent assays. All well known in the art. Sensitive measures of protein levels both from extracted tissue (including blood) or immunohistochemistry are known in the art. The effectiveness of introduction of TFOs into the bloodstream (as described for clinical trials on antisense oligo introduction for AIDS therapy) or at the site of tumor growth can be assayed by regression of tumor size, or by sampling the tumor for decreases in growth rate, induced apoptotic events, or changes in myc RNA levels.

EXAMPLE 14

Using TFOs to Alter the Expression of the Myc Oncogene

One of the most common oncogenes expressed in transformed cells is the c-myc oncogene. In vitro inhibition of the c-myc oncogene has been reported (Kim, H-G and Miller, D. (1995), “Inhibition of In vitro transcription by a triplex forming oligonucleotide targeted to human c-myc P2 promoter,” Biochemistry. 34, 8165-8171). The oligonucleotides AGGGAGGAGGGAAGA[AGG] (SEQ ID NO:4) and GG[AGAGGAGGGAAGGAGGAGGGA] (SEQ ID NO:5) are prepared having 87% of the linkages substituted with N,N-diethyl-ethylendiamine as disclosed in Example 1. Triplex formation is demonstrated by electrophoretic mobility shift analysis of triplex formation and DNase I footprinting as disclosed by Kim, et al. (supra). In vitro transcription alteration is performed using a HeLa nuclear extract in vitro transcription system (Promega) as disclosed by Kim, et al.

EXAMPLE 15

Duplex Studies

The sequences of the oligonucleotides used in these studies are found in FIG. 1B. To form duplex DNA, 1 nmol of target oligonucleotides was mixed with 1 nmol of each modified oligonucleotide to be tested. Three different conditions were tested, 150 mM NaCl/10 mM NaPO₄, 10%
formamide or 500 mM urea. The reactions comprising the different solutions, duplex DNA and target were heated to 85°C for 5 minutes and cooled slowly to room temperature. Absorbance of the oligonucleotide solutions were measured every 0.5°C from 15°C to 75°C at 200 nM using a Giford Model Response II thermal spectrophotometer. Melting temperature was obtained from the maximum value of the first derivative plots of absorbance vs. temperature.

Duplexes formed with an unmodified complementary strand of the oligonucleotides listed in FIG. 1 have the following thermal denaturation profiles in 150 nM NaCl/10 mM NaPO4, 10% formamide or 500 mM urea.

<table>
<thead>
<tr>
<th>Oligo</th>
<th>150 nM NaCl</th>
<th>10% formamide</th>
<th>500 nM urea</th>
</tr>
</thead>
<tbody>
<tr>
<td>U-1</td>
<td>52</td>
<td>29</td>
<td>31</td>
</tr>
<tr>
<td>N-1</td>
<td>40.5</td>
<td>22.5</td>
<td>24</td>
</tr>
<tr>
<td>N-2</td>
<td>42</td>
<td>24</td>
<td>29</td>
</tr>
<tr>
<td>P-1</td>
<td>49.5</td>
<td>44</td>
<td>48.5</td>
</tr>
</tbody>
</table>

In 150 mM NaCl, unmodified oligonucleotides and cationic oligonucleotides both had Tm’s of about 50°C, however, as salt concentration was lowered to 0 mM NaCl, unmodified oligonucleotides exhibited a decreasing Tm while the oligonucleotides of this invention with 100% modified internucleoside linkages were unaffected. Thus, the formation of duplex using oligonucleotides with the internucleoside linkages of this invention formed duplexes with target nucleic acid independent of salt concentration.

EXAMPLE 16

Probe Preparation

PNA Probe: 5′-BodipyTR-O-GCC TCC AGC CAC GTT-3′ (SEQ ID NO:6)

pDNA Probe: 5′-Rho-C6-TAT TTT AGC TGG TTT-3′ (SEQ ID NO:7)

BodipyTR is a fluorescent dye from Molecular Probes, Inc. (Eugene, Ore.). Rho (rhodamine) is a fluorescent dye from Glen Research, Sterling, Va.), C6 is a 5′-Amino-Modifier (Cat. No. 10-1906-90, Glen Research, Sterling, Va.); and O is the linker arm H-N—CH—CH—O—CH—CH—O—CH—COOH (PerSeptive Biosystems Inc., Framingham, Mass.).

The PNA (peptide nucleic acid) probe was purchased from PerSeptive Biosystems, Inc. (Framingham, Mass.). Peptide nucleic acids are uncharged analogs of DNA and RNA where the ribose-phosphate backbone is substituted by a backbone held together by amide bonds. The cationic DNA probe (pDNA) was synthesized at the 1 micromole scale with N,N-diethyl-stylenediamine phosphoramidite internucleoside linkages using H-phosphonate chemistry as described in Example 1. A C6 linker phosphoramidite was coupled to the 5′-end of the cationic probe on an automated DNA synthesizer (PE Applied Biosystems Division, Foster City, Calif.) using phosphoramidite coupling chemistry. The dye rhodamine 6G was coupled to the probe at the C6 linker as follows: The oligonucleotide (1 micromole scale) was dissolved in 200 μl of water. 50 μl of the dissolved oligo was mixed with 50 μl of 0.5 M sodium carbonate buffer, pH 9.0, plus 150 μl of water plus 4 μl of rhodamine 6G (TAMRA-NHS Ester, Cat. No. 50-5910-66, Glen Research, Sterling, Va.). The sample was vortexed and incubated at 37°C for 2 hours. Free dye was removed by precipitation of the oligonucleotide with 1 mL of ethanol followed by centrifugation. The precipitate was washed with ethanol to remove the free dye. The precipitate was dissolved in 50 μl of 50% DMSO in water and was purified by HPLC (RP-1 column, Cat. No. 79425, Hamilton Co., Reno, Nev.). Buffer A is 0.1 M triethylammonium, 5% acetonitrile, pH 8.0; Buffer B is acetonitrile; gradient is Buffer A plus 5% to 45% Buffer B in 35 min at a flow rate of 1 ml/min. Detection was at 260 nm (to detect nucleotides) and 547 nm (to detect the rhodamine). A peak was purified containing an equimolar ratio of oligonucleotide to rhodamine.

EXAMPLE 17

Detection of Phage λ DNA Target in Salmon Testes DNA Background Utilizing a DNA-Denaturing Buffer Containing Urea

Phage λ DNA (New England Biolabs, Beverly, Mass.) was mixed with salmon testes DNA (Sigma, St. Louis, Mont.) in a ratio of approximately one λ genome to one salmon genome. This DNA mixture (5×10^-11 M in genome equivalents) was incubated in 198 μl of 10 mM Tris-Cl, 0.1 mM EDTA, 7M urea, pH 7.2 at 95°C for 3 minutes. This concentration of urea prevents the formation of DNA-DNA duplexes (Orum et al., BioTechniques 19: 472 (1995)), but is permissible for the formation of pDNA-DNA hybrids as shown by the example shown.

1.0 μl of the PNA probe (2×10^-8 M) and 1.0 μl of the pDNA probe (2×10^-8 M) were added to the DNA mixture and incubated at 20°C for 20 hours. The final concentration of each probe was 1×10^-10 M. After incubation, 50 μl of sample, diluted in 5 ml of water was loaded into the sample capillary (800 μm×800 μm ID) of a single-molecule detection apparatus (Castro, A., Williams, J. G. K., “Single-Molecule Detection of Specific Nucleic Acid Sequences in Unamplified Genomic DNA” Anal. Chem. 1997, 69, 3915-3920 and Landers, E. S. Science 274:536, 1996 and the references cited therein). The sample was pushed through the capillary with a motorized syringe at 200 μm/sec. Fluorescence data were collected at 1 ms intervals. The λ target was identified by coincident detection of the two different fluorophore probes bound to the gene in a single detection zone. FIG 2A demonstrates the detection of the λ target in a single detection zone by crosscorrelation analysis. FIG. 2B is a control showing that no correlated signal is seen in a negative control sample identical to that of FIG. 2A but without the λ target DNA.

Cross-correlation analysis applies the following formula to two raw data sets $g_i$ and $h_i$.

$$ \text{Cor}(g_i, h_j) = \sum_{k=N-1}^{N} (g_k h_j) $$

where $N$ is the total number of data points in each raw data set. The raw data comprises photon counts detected in sequential 1 msec time intervals. Data is collected in separate channels for each fluorescent dye (data sets $g$ and $h$) while the sample moves past an orthogonal detection laser beam focused in the capillary. The correlation will be large at some value of $j$ if the first data set $g$ resembles the second data set $h$ at the lag time $j$. Therefore, whenever the sample contains a DNA target simultaneously hybridized to
the two probes, the coincident detection of the two fluorescent dyes produces bursts coincident in
time, which contributes to the cross-correlation peak at zero time offset. A strong cross-
correlation peak is seen in the sample of FIG. 2A, but not in the control of FIG. 2B, indicating that the pDNA probe (and also the PNA probe) hybridized to the target λ DNA under DNA-denaturing conditions.

In FIG. 2A the bin=4, offset=7.1E+6, amplitude=9.2E+4, width=10.33, M=34.47 and SNR=3.44. In FIG. 2B the bin=4, offset=8.0E+6, amplitude=5.4E+5, width=9.34, M=190.73 and SNR=18.24. In FIG. 2B the

It will be appreciated that certain variations to this invention may suggest themselves to those skilled in the art. The foregoing detailed description is to be clearly understood as given by way of illustration, the spirit and scope of this invention being limited solely by the appended claims.

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What is claimed is:

1. A method for hybridizing an oligonucleotide to denatured nucleic acid to form a duplex comprising the steps of: hybridizing a duplex-forming oligonucleotide comprising at least one cationic N,N-diethyl ethylenediamine-class phosphoramidate internucleoside linkage to single-stranded nucleic acid; and detecting hybridization of the oligonucleotide to the single-stranded nucleic acid.

2. The method of claim 1, wherein the denatured nucleic acid is DNA or RNA.

3. The method of claim 1, wherein the nucleic acid is denatured using heat, salt or a chaotrope.

4. The method of claim 3, wherein the chaotrope is selected from the group consisting of formamide, urea, or a guanidine salt.

5. The method of claim 1, wherein said oligonucleotide is about 12 to about 50 nucleotides in length.

6. The method of claim 1 wherein the oligonucleotide further comprises a tag.

7. The method of claim 1 wherein the tag is a fluorescent tag or a radiolabel.

8. The method of claim 1 wherein the denatured nucleic acid is genomic DNA.

9. The method of claim 8 wherein the oligonucleotide further comprises a nucleic acid sequence capable of hybridizing to a portion of a gene and wherein the detecting step indicates the presence of the gene in the genomic DNA.

10. The method of claim 1 wherein the method is used to detect single-nucleotide polymorphisms.

11. A method for detecting a nucleic acid fragment comprising the steps of:
   (a) hybridizing, under DNA—DNA duplex denaturing conditions, an oligonucleotide comprising about 30%
to 100% cationic N,N-diethyl ethylenediamine-class phosphoramidate internucleoside linkages to a sample of nucleic acid to form a hybridization complex; and (b) detecting said hybridization complex, wherein detecting the hybridization complex indicates the presence of the nucleic acid fragment in the sample of nucleic acid.

12. The method of claim 11, wherein the sample of nucleic acid is DNA or RNA.

13. The method of claim 11, wherein said DNA—DNA duplex denaturing conditions comprise a chaotrope.

14. The method of claim 13, wherein said chaotrope is selected from the group consisting of formamide, urea, or a guanidinium salt.

15. The method of claim 11, wherein said oligonucleotide is about 12 to about 50 nucleotides in length.

16. The method of claim 2 wherein the nucleic acid sample comprises genomic DNA.
It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Title page,
Item [56], **References Cited**, OTHER PUBLICATIONS, please delete “19(5)” and insert -- 19(15) -- therefor.
Please delete “Chaturvidi” and insert -- Chaturvedi -- therefor.
Please delete “Cusion” and insert -- Cushion -- therefor.
Please delete “Dimers, T* T and U* T” and insert -- Dimers, T* T and U* T -- therefor.
Delete “Corresponding Oligodeoxynucleotides,” and insert
-- Corresponding Oligodeoxynucleotides, -- therefor.

**Column 14.**
Line 41, please delete “ethylenediamine-class” and insert -- ethylenediamine -- therefor.

**Column 15.**
Line 24, please insert -- ( -- before “Froehl er et al.,”.
Line 25, please insert -- ) -- before “. All reagents used”.

**Column 22.**
Line 23, please delete “Mont.” and insert -- MO -- therefor.

**Column 25.**
Line 53, please delete “N,Ndiethyl ethylenediamine-class” and insert
-- N,Ndiethyl-ethylenediamine-class -- therefor.
It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Column 27,
Line 1, please delete “N,Ndiethyl ethylenediamine-class” and insert -- N,Ndiethyl-ethylenediamine-class -- therefor.

Signed and Sealed this

Third Day of December, 2002

JAMES E. ROGAN
Director of the United States Patent and Trademark Office