



Product Specifications

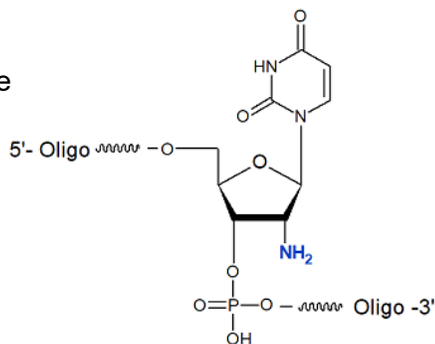
Custom Oligo Synthesis, antisense oligos, RNA oligos, chimeric oligos, Fluorescent dyes, Affinity Ligands, Spacers & Linkers, Duplex Stabilizers, Minor bases, labeled oligos, Molecular Beacons, siRNA, phosphonates Locked Nucleic Acids (LNA); 2'-5' linked Oligos

Oligo Modifications

For research use only. Not for use in diagnostic procedures for clinical purposes.

2'-Amino-U

Category	Nuclease Resistance
Modification Code	2-AmU
Reference Catalog Number	27-6507
5 Prime	Y
3 Prime	Y
Internal	Y
Molecular Weight(mw)	304.2



2'-Amino U

[27-6507-XX]

Oligonucleotide based diagnostics and therapy usually requires chemically modified bases for improved binding affinity/duplex stability, high selectivity and increased nuclease resistance.

The modification at 2' ribose position has shown to have stronger base-pairing with target, which lead to increase duplex stability, enhanced specificity and nuclease resistance (1). These modifications includes 2'F, 2'O methyl, 2'O MOE, and 2' amino bases.

Libraries of RNA molecules containing 2'amino-(2'NH₂)- or 2'fluoro-(2'F)-2'-deoxyuridines could yield ligands with similar nuclease resistance but not necessarily with similar affinities. This is because the intramolecular helices containing 2'NH₂ have lower melting temperatures (T_m) compared with helices containing 2'F, giving them thermodynamically less stable structures and possibly weaker affinities. These were tested by isolating high-affinity ligands to human keratinocyte growth factor from libraries containing modified RNA molecules with either 2'-NH₂ or 2'F pyrimidines. It was demonstrated that 2'F RNA ligands have affinities (K_d approximately 0.3-3 pM) and bioactivities (K_i approximately 34 pM) superior to 2'NH₂ ligands (K_d approximately 400 pM and K_i approximately 10 nM). In addition, 2'F ligands have extreme thermostabilities (T_m approximately 78C in low salt (1).

Oligonucleotide modified with 2'-amino C and U bases has shown to increase half-life 10 times more than other modified base in serum stability. However binding capacity is not as strong as other modified base.

2'-Amino pyrimidine Applications

- Antisense Oligos
- Aptamers
- siRNA

ASO's and siRNA Modifications.

Click this link to view ASO's and siRNA Modifications.

ASO's and siRNA Delivery. The development of effective delivery systems for antisense oligonucleotides is essential for their clinical therapeutic application. The most common delivery system involves a relatively hydrophobic molecule that can cross the lipid membrane. Cholesterol TEG, alpha-Tocopherol TEG (a natural isomer of vitamin E), stearyl and GalNac modifications have been shown to effective for delivery of ASO's and siRNA in addition to cell penetrating peptides.

[com/newsite/products/mod_detail.asp?modid=431](http://www.genelink.com/newsite/products/mod_detail.asp?modid=431)"> Click this link to view these modifications.

REFERENCES:

1. Pagratis, N.C., et al., Potent 2'-amino- , and 2'-fluoro-2' deoxyribonucleotide RNA inhibitors of keratinocyte growth factor. (1997), **15**: 68-73.
2. Lou, C. et al., Oligonucleotides Containing Aminated 2'-Amino-LNA Nucleotides: Synthesis and Strong Binding to Complementary DNA and RNA. *Bioconjugate Chem.* (2017). **28**: 1214-1220