

# Locked Nucleic Acid (LNA)

# Increased thermal stability and hybridization specificity Improved signal-to-noise ratio in qPCR assays Enhanced single nucleotide discrimination

## **Background Information**

Locked Nucleic Acid (LNA) is a synthetic nucleic acid analogue containing a bridged, bicyclic sugar moiety. The extra methylene group attached between the 2'-O- and the 4'-positions "locks" the ribofuranosyl-ring in its 3'-endo conformation (see *Figure 2*). This conformation leads to the characteristic structure of A-form RNA. As a consequence of the constraint bicyclic sugar skeleton, LNA exclusively forms A-type duplexes. Furthermore, LNA fully complies with Watson-Crick base pairing rules. LNA:DNA hybrid duplexes are formed spontaneously from complementary DNAand LNA-sequences, and it was found that LNA:DNA hybrids show strongly improved annealing temperatures

compared to their DNA:DNA counterparts [1]. Since the synthesis of LNA is compatible with standard oligonucleotide synthesis, site-selective incorporation of single or multiple LNA nucleotides into DNA sequences can be achieved straightaway. These LNAcontaining oligonucleotides anneal with their DNA complements to form chimeric LNA:DNA hybrids. Any such duplex adopt A-form conformation, and again Tm are substantially increased compared to analogous DNA:DNA double-strands. As a rough guess, the incorporation of LNAnucleotides into short DNA primers (< 30 nt) increases the Tm by 3-8 °C for each substituted nucleotide [2].

All in all, the major advantage of LNA lies in the design options for primers and probes as the Tm can be finetuned according to the needs of a desired oligonucleotide. Due to the enhanced binding affinity, shorter probes can be realized and as a result, binding specificity to the target DNA is increased. Therefore, LNA is recommended for use in any hybridization assay that requires high specificity and/or reproducibility, e.g. PCR primers, dual-labeled probes, in-situ hybridization probes, and molecular beacons. Furthermore, but for the same reasons, LNA modified oligonucleotides are equally interesting as candidates in antisense drug development [3].

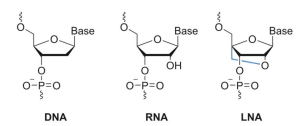


Figure 1: Structural drawings of DNA, RNA and LNA nucleotides

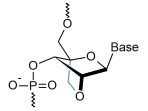


Figure 2: LNA 3'-endo conformation

# **Increased Thermal Stability of qPCR Probes**

The superior hybridization characteristics of LNA allow a fine-tuning of the Tm in the design of qPCR probes [4]. This significantly broadens the scope of assay conditions and permits

more successful qPCR. When probes consist of LNA nucleotides, affinity and specificity of the hybridization to the target sequence are improved. This in turn **reduces background** 

**fluorescence** from spurious binding and leads to a **better signal-to-noise** ratio.



#### **Multiplex qPCR Systems**

As the Tm of qPCR probes can be adjusted using LNA, **normalization of the Tm** across several short sequences with varying GC-content becomes accessible. For example, AT-rich nativestate DNA qPCR probes often need to be over 30 bases long (sometimes

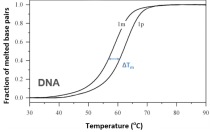
over 40 nt) to satisfy amplicon design guidelines but may still perform poorly. With LNA qPCR probes, the selective positioning of LNA nucleotides **facilitates the optimal design** of highly-specific, shorter probes. A narrow Tm range is **in particular ben** 

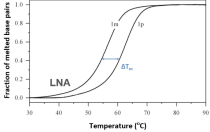
eficial for microarray and multiplex PCR applications, where simultaneous binding of probes to many different targets must occur under the same conditions.

#### **Enhanced Single Nucleotide Discrimination**

The ability of qPCR probes to discriminate between alleles via single nucleotide polymorphism (SNP) is greatly enhanced by the incorporation of LNA nucleotides since there is strong preference of LNA to form Watson-Crick base pairs over

mismatches [5]. This is reflected by thermal denaturation studies that show a remarkable difference in Tm between a perfect match and a mismatch when LNA is incorporated at the particular position of the oligonucleotide (see Figure 3). Therefore, in SNP assays, LNA qPCR probes have an **enhanced destabilizing effect** on target hybridization and permit **better mismatch discrimination** compared to native-state DNA probes.





**Figure 3**: Influence of LNA on the melting temperature (Tm) and the resulting larger difference between specific (1p) and non-specific (1m) signals

#### **Antisense Technology**

Due to the enhanced binding affinity to complementary nucleic acids, there is great potential for LNA to be used in antisense technology. The concomitant high nuclease resistance of LNAs is an important benefit for *in vivo* 

and *in vitro* applications. Numerous studies confirm the superior properties of LNA as antisense agents. LNA oligonucleotides are transfected by the conventional techniques. For knockdown of microRNA or other

small RNAs, LNA antisense oligonucleotides are designed to hybridize to their target sequence by base pairing. A further increase in nuclease resistance is obtained by the introduction of a phosphorothioate backbone.

#### Literature

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