IDT SciTools: a suite for analysis and design of nucleic acid oligomers

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ABSTRACT

DNA and RNA oligomers are used in a myriad of diverse biological and biochemical experiments. These oligonucleotides are designed to have unique biophysical, chemical and hybridization properties. We have created an integrated set of bioinformatics tools that predict the properties of native and chemically modified nucleic acids and assist in their design. Researchers can select PCR primers, probes and antisense oligonucleotides, find the most suitable sequences for RNA interference, calculate stable secondary structures, and evaluate the potential for two sequences to interact. The latest, most accurate thermodynamic algorithms and models are implemented. This free software is available at http://www.idtdna.com/SciTools/SciTools.aspx.

INTRODUCTION

Synthetic oligonucleotides are widely employed in various molecular biology applications, e.g. polymerase chain reaction (PCR), molecular beacons, microarrays, mutagenesis, RNAi, antisense and de novo gene construction (1–7). Published bioinformatics algorithms can predict biophysical properties of oligonucleotides from their sequence and estimate performance of oligonucleotides in specific assays both singly and together with other sequences (8,9). Here, we describe an online suite of computational software tools that enable molecular biologists to design, evaluate and make informed decisions about the properties of nucleic acid sequences. The IDT SciTools receives over 7000 unique visitors and 1.5 million hits every month. The web servers consist of several independent applications summarized in Table 1. Instructions and help to each software tool can be found at the top of web input forms. The code is regularly updated when more accurate models and algorithms are published. New applications will be added in the future.

ONLINE TOOLS

OligoAnalyzer 3.1

The OligoAnalyzer is the central calculator where various kinds of information about an oligonucleotide sequence can be predicted. The interface is presented on Figure 1. A user can input a nucleotide sequence and conditions, i.e. the concentrations of DNA, Na+, K+, Mg2+ and deoxynucleoside triphosphates. Melting temperature is predicted under these conditions for the duplex where the oligonucleotide hybridizes to the complementary sequence. This complementary strand can be either RNA or DNA; this is selected using the Target Type option. The oligonucleotide sequence can be modified with over 150 different labels and chemical groups (e.g. biotin, phosphorothioate, fluorescent dyes) using symbols listed in the tabbed sections below the sequence box. Seven different analyses can be performed when the specific button is selected on the right side of the interface. Selection of the ANALYZE button results in the physical properties of the oligonucleotide, such as a complementary sequence, oligonucleotide length, content of G and C bases, melting temperature, extinction coefficient at 260 nm and molecular weight (Figure 1). Published nearest-neighbor parameters are employed to calculate the extinction coefficient (10–12). Using values obtained from the published literature or coefficients estimated at Integrated DNA
Table 1. Summary of IDT SciTools

| Software tool       | Features                                                                 |
|---------------------|---------------------------------------------------------------------------|
| OligoAnalyzer       | Comprehensive oligonucleotide analysis (molecular weight, extinction coefficient, melting temperature, folding and hybridization of strands, effects of mismatches). |
| PrimerQuest         | Select optimal probes and primers for PCR assays.                         |
| LNA design          | Tool to design LNA modified probes and primers having specific duplex stability. |
| ddRNAi design       | Design sequences for DNA-directed RNA interference.                       |
| RNAi design         | Tool to design siRNA duplex oligomers.                                    |
| TriFECTa RNAi Kits  | Predesigned dicer-substrate RNAi duplexes, gene knockdown kits.           |
| Antisense design    | Antisense oligonucleotide selection tool.                                 |
| mFold               | Prediction of oligonucleotide secondary structure.                        |
| DilutionCalc        | Calculate volumes to dilute oligonucleotide to the desired concentration. |
| ResuspensionCalc    | Calculate volumes to dissolve a dry lyophilized nucleic acid to the desired concentration. |

Figure 1. Interface of OligoAnalyzer 3.1. Instructions and help files are at the top of the calculator. The sequence and conditions of experiments are entered in the upper region. Many chemical modifications can be added to a sequence using symbols from tabbed sections in the middle segment. The results are displayed in the lower left segment.
Technologies, the effects of modifications are included in the oligonucleotide extinction coefficient.

The oligonucleotide molecular weight also includes the weights of any chemical modifications. These weights have been experimentally validated (±2 g/mol) for thousands of synthesized sequences by electrospray-ionization liquid chromatography mass spectrometry (13).

Melting temperatures are calculated from the nearest-neighbor model (14–16) and the duplex is assumed to melt in two-state fashion,

\[ T_m = \frac{\Delta H^o}{\Delta S^o + R\ln C_{oligo}} \]

Oligonucleotide concentration, [S1], is assumed to be significantly larger (at least 6x) than the concentration of the complementary target, [S2], as this is seen in many molecular biology assays. In that case, \( C_{oligo} \) is equal to [S1] and the concentration of the target can be neglected (17). If [S1] is not significantly larger than [S2], but \( [S1] \geq [S2] \), the following concentration should be entered into the calculator,

\[ C_{oligo} = [S1] - \frac{[S2]}{2} \]

If \( [S2] < [S1] \), Equation (2) is valid when the designation of strands is switched. Transition enthalpy, \( \Delta H^o \), and entropy, \( \Delta S^o \), are calculated from the latest nearest-neighbor parameters for DNAs (15,16) and RNAs (18,19). The effects of counterions are modeled using the improved corrections for monovalent ions (20) and magnesium ions (53),

\[ \frac{1}{T_m(Mg^{2+})} = \frac{1}{T_m(1M Na^+)} + \left\{ 3.92 - 0.911 \ln[Mg^{2+}] ight\} + f_{GC} \times (6.26 + 1.42 \ln[Mg^{2+}]) + \frac{1}{2(N_{bp} - 1)} \times \left\{ -48.2 + 52.5\ln[Mg^{2+}] \right\} + 8.31(\ln[Mg^{2+}]^2) \times 10^{-5} \]

This unique biophysical model employed for various counterions is not implemented elsewhere (21–23). Thermodynamic parameters are not available for many modifications that were demonstrated to change duplex stability (e.g. \( 2'-O \)-methyl RNA, 2-aminopurine, Cy3 dye) (13), and their effects on melting temperature are therefore neglected in the current version. When these parameters are published, they will be implemented in the predictive algorithm. If a sequence contains degenerate bases, the minimum and the maximum melting temperatures for the mixture of sequences are also estimated (Figure 1). The thermodynamic algorithm was validated using an independent set of over 100 different sequences ranging in length from 8 to 60 base pairs that were not used to derive the algorithm (20).

Selection of the TM MISMATCH button will allow the user to examine the effects of single base mismatches on duplex stability and oligonucleotide hybridization. Several published sets of nearest-neighbor parameters from SantaLucia’s lab are employed to make these predictions (16,28–32). Dangling unpaired bases usually stabilize the duplex, so the predictive algorithm also takes these effects into account (33). If a red target base is clicked, a dropdown box will appear and allow the user to select the desired base mismatch. The target concentration can be set to zero when the target concentration is negligible in comparison with the oligonucleotide concentration. Results will show melting temperatures of perfectly matched and mismatched duplexes as well as the fractions of oligonucleotide bound to the targets.

The LNA CONVERSION button will be described later. The tool allows the user to position LNA modifications within a sequence, so that the desired melting temperature of the duplex sequence is achieved.

**PrimerQuestSM**

Primer and probe selection for the PCR-based assays are important activities in molecular biology. Several software packages were therefore designed for this procedure (34–39). PrimerQuestSM is based on the Primer3 code (37). However, the selection method was improved and a graphical user interface was created. The algorithm finds sequences having desired oligonucleotide length, GC content, melting temperature, content of consecutive GC base pairs and sequence stability at the 3’ end. The intramolecular secondary structures, long repeats of the same bases and cross-hybridization between primers and the probes are minimized in the primer selection model. Furthermore, the oligonucleotide melting temperature is calculated using the same thermodynamic model employed in the OligoAnalyzer. Once the nucleotide sequence is entered in the sequence box, the name and design criteria for the sequence of interest can be set using the appropriate fields under the basic, standard and advanced tabs. The basic interface exposes the minimal information that needs to be entered and hides detailed criteria. These basic settings are suitable for typical PCR experiments. The standard and advanced tabs show increasing amounts of settings that an advanced user can configure to customize their predictive model. The **CALCULATE** button submits the data for the prediction of primers with the desired properties.

Results show several sets of probes and primers that were found to be optimal (Figure 2). The predicted sets are
ranked from best to next best. A graphical representation of the sequence is displayed with color bars for included, excluded and targeted regions. The biophysical properties of the primers and probes are also reported.

**LNA design**

Chimeric probes containing locked nucleic acid residues were demonstrated to increase duplex stability, specificity and mismatch discrimination (40,41). These properties...
Antisense design

Expression of specific genes can be suppressed with antisense oligonucleotides (43). Software can be used to select the most effective antisense oligonucleotides based on a model that discriminates between effective and ineffective antisense sequences. The nucleotide sequence of a gene or other target candidate for antisense-based knockdown can be retrieved from NCBI databases using GenBank ID or RefSeq ID. Antisense DNA oligomers are typically from 19 to 26 bases long and modified with phosphorothioates for nuclease resistance. Optionally, typically from 19 to 26 bases long and modified with phosphorothioates for nuclease resistance. Optionally, modified oligomers (40,42).

The software will attempt to decrease the length of the sequence and introduce LNA modifications, so that the desired melting temperature is achieved. The LNA residues are indicated with '+' symbol in front of the base. Melting temperatures are predicted using the nearest-neighbor two-state model (16,42), and improved salt corrections for the effects of monovalent and magnesium ions (20,53). The algorithm was tested with a published set of melting data for LNA modified oligomers (40,42).

mFold

This software tool predicts the most stable secondary structure of an oligonucleotide by minimizing folding free energy (51). Suboptimal energetic secondary structures having free energies close to minimal ΔG can be predicted as well. The mFold software was developed by and implemented in collaboration with Prof. Michael Zuker. The algorithm has been well tested and described in published sources (9,24–26). A user can input a nucleotide sequence and the conditions, including temperature and ionic concentrations. The folded structures are predicted at the specified temperature. The results show both a dot-plot diagram of possible base pairings and predicted secondary structures. These structures are ranked from the highest to lowest probability using transition free energies. Melting temperature is estimated using a two-state model. The connectivity table for each base and details of the energetics (each loop and stack ΔG contributions) can also be obtained.

Dilution and resuspension calculations

Oligonucleotide dilutions can be calculated using the dilution calculator. A user inputs initial concentration, volume and desired final concentration. The calculator returns volumes used to mix solutions. Various concentration formats and units are accepted. Similarly, the resuspension calculator determines the volume of solution needed to achieve a specific concentration when known moles or mass of dry oligonucleotide are dissolved. Both calculators can be brought directly from the Oligo Analyzer results. In that case, oligonucleotide properties predicted by OligoAnalyzer are transferred automatically to these calculators.

Biophysics.idtdna.com

Subdomain http://biophysics.idtdna.com contains advanced, unique calculators that are being developed. Software is stable and tested, but it has yet to be included into IDT SciTools. In the current version, the extinction coefficients and UV spectrum from 215 to 310 nm can be predicted for both single-stranded and double-stranded DNA oligomers. The models, parameters and their accuracy tests have been recently published (12). A user can also choose to apply Cavaluzzi–Borer correction for extinction coefficients of DNA bases at 260 nm (52). The predicted UV spectrum is plotted and extinction coefficients at each wavelength are tabulated.
The software tools help to select oligonucleotides that are most likely to exhibit the best performance in biological applications.

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Conflict of interest statement

The authors are or have been employed by Integrated DNA Technologies, Inc. (IDT). IDT is not a publicly traded company and has filed, or employed by Integrated DNA Technologies, Inc., (IDT).

CONCLUSIONS

IDT SciTools web server provides useful predictions of oligonucleotide properties under various experimental conditions. The software tools help to select oligonucleotides that are most likely to exhibit the best performance in biological applications.

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