

**LAURIS KAPLINSKI**

The application of oligonucleotide  
hybridization model for PCR and  
microarray optimization





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## LIST OF ORIGINAL PUBLICATIONS

- I. Kaplinski L, Andreson R, Puurand T, Remm M (2005). MultiPLX: automatic grouping and evaluation of PCR primers. *Bioinformatics* 21(8): 1701–2.
- II. Kaplinski L, Scheler O, Parkel S, Palta P, Toome K, Kurg A, Remm M. (2010) Detection of tmRNA molecules on microarrays at low temperatures using helper oligonucleotides. *BMC Biotechnol.* 10:34
- III. Scheler O, Kaplinski L, Glynn B, Palta P, Parkel S, Toome K, Maher M, Barry T, Remm M, Kurg A (2011) Detection of NASBA amplified bacterial tmRNA molecules on SLICSel designed microarray probes. *BMC Biotechnology* 11:17
- IV. Kaplinski L, Möls M, Möls T, Remm M. Predicting failure rate of Multiplex PCR (submitted)

Author's contributions:

- Ref. I: created the grouping algorithm, wrote the software implementation and was responsible for drafting the manuscript.
- Ref. II: implemented the probe selection algorithm, wrote the software, designed the detection probes, performed data analysis and was responsible for drafting the manuscript.
- Ref. III: conducted this study, designed the chaperon probes, performed data analysis and was responsible for drafting the manuscript.
- Ref. IV: conducted this study, performed data analysis and was responsible for drafting the manuscript

## LIST OF ABBREVIATIONS

NN	nearest neighbor
NP	nondeterministic polynomial time
$T_m$	melting temperature
tmRNA	transfer-messenger RNA
PCR	polymerase chain reaction
ROC	receiver operating characteristic
$\Delta G$	free energy change

## INTRODUCTION

Nucleic acid based technologies are becoming increasingly important in each year at fields as diverse as medical diagnosis, nanoscale engineering and information storage. Although the basis of nucleic acid structure was determined over half a century ago, the study of more complex structures and interactions is ongoing. The practically infinite variability of the primary structure of nucleic acids makes the case-by-case study of their properties practical only for few important biological molecules, such as tRNAs and rRNAs. For the overall majority of cases, scientists rely on statistical models that predict with varying success the important properties of nucleic acid structure and interactions. The greater the importance of nucleic acid-based technologies, the more precise the models need to be.

The first part of this thesis gives a brief overview of the thermodynamic and kinetic properties of nucleic acids hybridization. The second part concentrates on the nearest-neighbor thermodynamic model that is currently the state-of-the-art method for calculating hybridization parameters. The third part gives an overview of the application of thermodynamic model on the design of PCR primers, with special attention paid to multiplex PCR and microarray hybridization probes.

The research part of this dissertation covers the following topics: 1) development of efficient method for grouping primer pairs in multiplex PCR, 2) determination of the most important primer-specific factors that cause the failure of multiplex PCR, 3) development of an automated procedure for designing an exhaustive set of probes for the detection of bacterial RNA, and 4) development of helper oligonucleotides for breaking the secondary structure of tmRNA.



# REVIEW OF LITERATURE

## The digital code

Nucleic acids form the most basic “digital code” of life. The unique property of a forming stable perfectly aligned double-helical structure, discovered by James Watson and Francis Crick, is the basis of both the storage and transmission of genetic information in living organisms.

In addition to information storage and transmission, nucleic acids, or more specifically RNA, have many metabolic and control functions in cells. Although the complexity of nucleic acid sequences is many orders of magnitude lower than the complexity of amino acid sequences of comparative length (Levitt 2009), they can nevertheless fold into complex 3-dimensional structures that have specific interactions within cell metabolic system. Like for proteins, the exact 3-D structure of a nucleic acid fragment is determined by both its primary structure and environment, where it was synthesized and operates.

## Chemical basis of the digital code

In many biological applications, it is sufficient to think of nucleic acid sequences as a purely digital code built on a 4-letter alphabet. Nevertheless, it is important to understand, that its “digital” nature is determined by the chemical properties of the nucleic acids. These properties determine the specificity of chemical reactions – replication, transcription and translation – that result in copying or interpreting the genetic information.

In the context of information processing, the most important of those chemical properties are:

- a) The polar structure of the nucleic acids. The chains are composed of negatively charged hydrophilic pentose phosphate “backbone”, and neutral or positively charged hydrophobic bases. Although the negative charges of the phosphate groups repel each other thereby destabilizing the double helix, the containment of hydrophobic bases and hydrogen bonds in the internal region stabilizes the structure. The double-helix is also stabilized by mono- and divalent cations that partially neutralize the repulsive negative charges of the phosphates (Levitt 2009; Cisse, Kim, and Ha 2012)
- b) The hydrogen bonds between aligned bases from 2 antiparallel strands. The preferred pairings are such that the aligned bases “fit” with each other, forming 2 (A/T) or 3 (G/C) hydrogen bonds. These bonds are stronger than in solution due to the hydrophobic nature of the inner part of dihelical structure (Levitt 2009).
- c)  $\pi$ -stacking, van der Waal’s forces and hydrophobic interactions between adjacent aromatic rings. Nucleic acid chains, even in the absence of double-helical structure, often take such a form that the planes of aromatic bases are stacked parallel to each other and partially overlap. In this configuration, the

interactions between p-electrons between adjacent bases are strongest and stabilize the structure of the whole nucleic acid sequence (Borer et al. 1974; Levitt 2009).

## Hybridization of nucleic acids

The 3 above-mentioned chemical features contribute to the spontaneous formation of double-helical structure between 2 single-stranded nucleic acid chains or 2 regions of the same chain. This process, known as hybridization, is the basis of the formation of most of the ordered structures of nucleic acids, such as the near-perfect double helix of genomic DNA or clover-leaf structure of tRNA.

The strength and speed of spontaneous hybridization of specific nucleic acid sequence depends mostly on 3 factors, namely enthalpy, entropy and steric effects.

**The enthalpy (the “pure” energetic effect) of hybridization ( $\Delta H$ ).** This is the potential energy change during the formation of double helical structure, mostly caused by the formation of various bonds – hydrophobic, aromatic and hydrogen bonds and electromagnetic interactions between charged parts of molecule. In practically all environments, the double-helical structure is energetically advantageous for semi-complementary strands of nucleic acids and the double helix with its perfectly aligned Watson Crick pairs has the highest change of enthalpy (Levitt 2009).

**The entropy effect (i.e. change in the degree of freedom of configurations) of hybridization ( $\Delta S$ ).** The double helical structure always has a much lower entropy than random distribution of nucleic acid fragments or chains in solution. The more ordered the alignment in double helix, the lower is the entropy and the smaller is the probability of nucleic acid chain randomly taking this configuration (Levitt 2009). The statistical effect of entropy to the actual hybridization is strongly temperature-dependent.

**Steric effects.** Nucleic acid chains being macromolecules, the formation of certain configurations may be improbable or impossible simply due to their size. One very common effect is steric blocking of the hybridization with the inner part of single-stranded molecule due to its 3-D structure.

For the actual calculation of the properties of chemical reactions, including nucleic acid hybridization at constant temperature, it is usually sufficient to use the composite free energy value ( $\Delta G$ , Gibbs’ free energy).

$$\Delta G = \Delta H - T\Delta S$$

Gibbs’ free energy corresponds to the actual amount of energy that can be extracted or is needed to perform certain reaction or transition. Its value determines the equilibrium constant of a reaction – if  $\Delta G$  of a reaction is negative, the equilibrium is shifted towards products; if it is positive, towards reagents. As  $\Delta G$  contains both enthalpy and entropy components, it is also temperature-dependent.

The hybridization of nucleic acids is a complex process that depends on the type of molecules (DNA or RNA) and their primary sequence. Longer nucleotide chains hybridize in zipper-like fashion, starting from short “seeds”. The hybridization of shorter oligonucleotides is usually modeled on a one-step process, although even in this case it seems to start from a contiguous perfectly aligned seed of 7 base pairs (Cisse, Kim, and Ha 2012). The dissociation seems to start from the ends of hybridized oligonucleotides (Wienken and Baaske 2011).

For the complementary molecules with the identical primary sequence, the hybridization strength normally increases in the following order: DNA/DNA < DNA/RNA < RNA/RNA; but there are exceptions, as certain Watson-Crick or mismatched pairs and motifs have very different pairing strengths in different hybridization types (Sugimoto, Nakano, and Katoh 1995; T. Xia et al. 1998).

Steric effects are the cause of long nucleotide sequences not forming perfectly paired hybridized duplexes. Although the perfect duplex between such strands would be globally most energetically advantageous, its spontaneous formation would require extremely improbable random formation of exactly paired duplex between 2 long sequences and unwinding of the helical structure of single-stranded molecules. Instead, certain regions of single stranded molecules form locally hybridized double-helical structures that represent a local energy minimum and block the formation of global alignment (Levitt 2009). As the composition of locally hybridized regions is dependent on the folding history of nucleotide chain, the secondary structure of long DNA and RNA chains is unpredictable (Chavali et al. 2005).

Steric effects become the stronger the longer nucleic acid chains are. In living cells topoisomerases are needed to clear out steric blockage of nucleotide chains during replication.

Hybridization is stochastic process; there are no objective factors that force a nucleic acid fragment to search out and hybridize only with its “correct” partner. Any 2 fragments can associate and form hybridized pair, and any 2 hybridized fragments can dissociate. What differs between the “correct” and “incorrect” hybridizations are their energetic effects. Usually the  $\Delta G$  of hybridization is lowest (i.e. most energetically favorable) for the hybridization between perfect Watson-Crick complementary sequences and highest (i.e. energetically unfavorable) between certain sterically incompatible sequences. As the spontaneous processes are statistically favored towards the state with lower  $\Delta G$ , perfect or near-perfect hybridized states are normally favorable and eventually prevail when the hybridization process runs long enough.

Hybridization is also a reversible process; at any temperature above absolute zero, both the hybridization and dissociation processes can take place between arbitrary fragments of nucleic acids. The speed of both processes, as expressed by the reaction rates of annealing ( $K_{\text{ann}}$ ) and melting ( $K_{\text{melt}}$ ), are determined by the change in free energy ( $\Delta G$ ) during hybridization and the concentrations of free and hybridized nucleic acid fragments. The balance of hybridized and free

states as expressed by the equilibrium constant  $K_d$  is determined by the ratio of corresponding reaction speeds – and thus depends on the  $\Delta G$  of hybridization.

Hybridization and melting go through different intermediary stages, as indicated by the fact that the effects of internal mismatches in oligonucleotide sequences are different for annealing and melting. While the dissociation speed is highest when mismatches occur in the middle of sequence, the association speed has a step-like dependence on the number of consecutive perfectly aligned nucleotides. This is in accordance with the melting models, where dissociation proceeds over the intermediate form where only the middle of the sequence is in the hybridized state (Wienken and Baaske 2011). Hybridization, on the other hand starts with a seed of certain minimal length in any region of nucleotide chain. Seven contiguous base-pairs increases the speed of the annealing of DNA or RNA by several orders of magnitude compared to alignments with shorter perfectly aligned spans (Cisse, Kim, and Ha 2012).

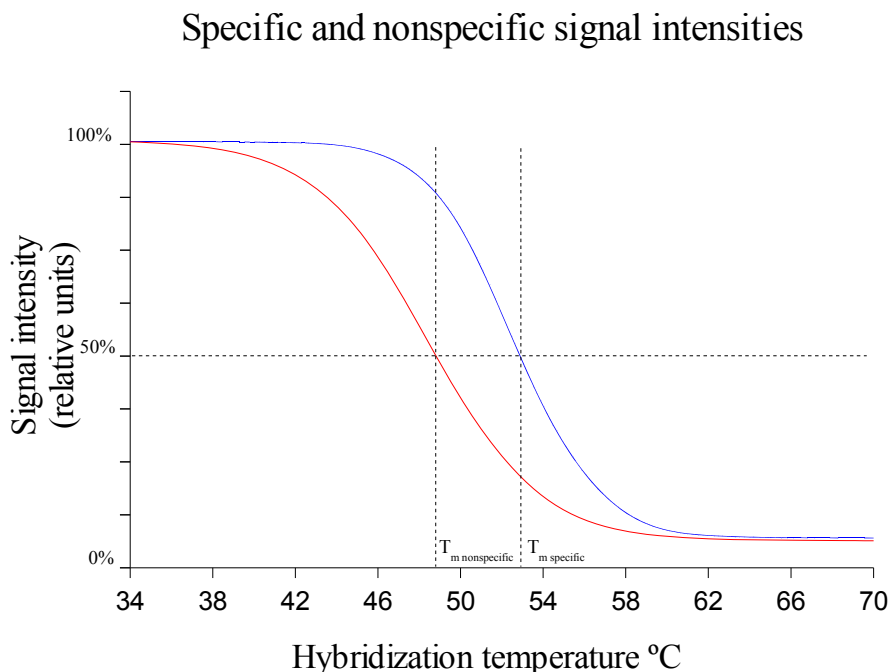
The positions of mismatches relative to each other in oligonucleotide sequence also affect the hybridization equilibrium. Mismatches closer than 5 bp have a smaller effect on the stability of hybridization than mismatches with wider spacing. Also mismatches close to the ends of oligonucleotide chain have less effect (Hadiwikarta et al. 2012).

The term ‘melting temperature’ ( $T_m$ ) is often used to characterize certain oligonucleotides, such as PCR primers and detection probes. It corresponds to the temperature at which the rates of both hybridization and dissociation are equal, and thus exactly half of the oligonucleotides are in the hybridized and half in the free state. The  $\Delta G$  of hybridization reaction is roughly 0 at this temperature, because the equilibrium depends on the concentrations of the hybridizing oligonucleotides in addition to the free energy of helix formation.

In many practical applications of oligonucleotide hybridization like PCR, the exact value of equilibrium constant  $K_d$  can be ignored as long as it is sufficiently high or low. For example, if the temperature is sufficiently below  $T_m$ , the  $\Delta G$  of hybridization is significantly negative and the equilibrium is strongly shifted towards the hybridized state; and vice versa, i.e. if the temperature is sufficiently above  $T_m$ , most nucleic acids are in free state.

While such approach is often sufficient for experimental work, there are other cases where it may not be possible to guarantee a sufficient temperature difference from  $T_m$ . In such cases more precise calculation of  $\Delta G$  value is beneficial, as it allow the experimenter to take into account the actual equilibrium ratio and find optimal reaction parameters. One such example is the discrimination, either by PCR or hybridization array, between closely related organisms (e.g., bacterial strains). These often differ only by few nucleotides. Thus primers or probes designed to be complementary to the DNA sequence of one organism hybridize only slightly more strongly to their intended target than to control group (Figure 1). To achieve the maximal possible discriminatory power, annealing should take place at the precise temperature at which the difference between actual detectable hybridization to target and nontarget is maximum. Another example is the discrimination between many target mole-

cules in a single fluorescence channel by the differences in the melting temperatures of their corresponding hybridization probes (Fu, Miles, and Alphey 2012), i.e. where the precise prediction of melting curves of hybridization probes is essential.



**Figure 1:** Theoretical melting curve of a perfect hybridization and nonspecific hybridization with one mismatch of the same probe. Blue – melting curve of perfect hybridization. Red – melting curve of nonspecific hybridization. X axis – annealing temperature. Y axis – relative signal intensity (proportional to the amount of hybridized probes). The melting temperatures of both specific and non-specific hybridization are marked with vertical lines.

Certain non-perfect alignments, e.g. terminal G/T (and especially G/U in case of RNA) mismatches, are only slightly disadvantageous ( $\Delta G$  in the range of 0.2 kcal/mol) compared to perfect matches in aqueous solution. A similar effect is also seen for the antiparallel combination of G/A and A/G mismatches, which may be very difficult to discriminate by only hybridization because the equilibrium constants are very similar for specific and nonspecific hybridization. But if subsequent PCR reaction is used and thus DNA strands are bound to the active center of DNA polymerase, the  $\Delta G$  difference between perfect and mismatched pairing of a terminal nucleotide is amplified by an order of magnitude (Petruska et al. 1988).

The concentration of mono- and divalent cations has a strong effect on the hybridization of nucleic acids. Higher salt concentrations make all hybridized states, both specific and non-specific, more stable, possibly by shielding negative charges of phosphate backbones.

## **High-level structure of nucleic acids**

The best-known structure of nucleic acids is the linear double-helix, of which there are 2 alternative configurations – the A and B forms that differ by the angle of the bases and the distance of phosphate backbones (Levitt 2009). Double helices are prevalent in genomic DNA, being formed during the synthesis of nucleotide strands and staying in the hybridized state for most of the time. On the other hand, long single-stranded nucleic acid chains normally do not form double helix even when their complementary chain is present because such a configuration has extremely low entropy (and thus also an extremely low probability of spontaneous hybridization).

Nevertheless, nucleic acid chains can spontaneously form complex structures due to partial self-complementarity of the different regions of the same molecule. The self-complementary parts of the molecule tend to form double-helical stems that are connected by single-stranded loops. Both local and distant parts of the molecule can hybridize, creating complex 3-D configurations that tend to stabilize the conformation of the molecules (Levitt 2009). The 3-D configuration itself is highly dependent on both the primary structure of the nucleic acid chain and its folding history. There is often no single prevalent configuration but many variants. The actual configurations of nucleic acids are also sensitive to changes in primary sequence as a single mutation can make it to fold in a completely different configuration. This phenomenon has been used for the detection of biological variants based on their structural properties instead of sequence differences (Dong et al. 2001).

It should be noted that all the properties of nucleic acid fragments are ultimately determined by their primary structure and local environment. For short oligonucleotides, the primary structure is all that needs to be known to predict precisely their hybridization behavior and secondary structure. For longer nucleotide chains, the secondary structure becomes dependent on the folding history of the molecule, and thus it is not strictly predictable from the primary structure. Nevertheless, the stability of various global and local conformations is determined by primary sequence (Breslauer et al. 1986; Dong et al. 2001).

The detection of the secondary structure of nucleic acids is complex process. For short sequences, the most energetically favorable configurations can be determined computationally. The most popular program for secondary structure prediction is UNAFold (Markham and Zuker 2008). Computational methods usually only predict the overall energetic potential of different structures and cannot estimate the relative abundance of actual configurations because the

latter depend on folding history. For native molecules, the most precise method is often by using sequence alignment with a phylogenetically close molecule of known structure. For new molecules without known reference structures various physical and chemical methods are used with varying success, for example X-ray crystallography and enzymatic cleavage. Often many methods, both experimental and computational, have to be combined to determine the true structure of nucleic acids (Dong et al. 2001).

## **Methods for calculating the hybridization parameters of nucleic acids**

The oldest and frequently used method for estimating the melting temperature of 2 nucleotide strands with complementary sequences is by their GC content. It is practical for long nucleotide chains where it is not possible to use precise thermodynamic calculations. In long molecules, the statistical effect averages out the effects of local context. For short oligonucleotides, this method is imprecise because the local context influences the strength of G/C and A/T pairings considerably (Borer et al. 1974). The widely used formula is:

$$T_m = 81.5 + 16.6 \log \left\{ \frac{[Na^+]}{1.0 + 0.7[Na^+]} \right\} + 0.41(\%G + C) - \frac{500}{D} - P$$

where D is duplex length and P is the percentage of mismatching positions (Wetmur 1991).

A similar method was traditionally used in probe design – hybridization probes had fixed length, a GC content within specified range, and a certain minimum number of mismatches with non-target sequences. By fulfilling these conditions, it is expected that a given oligonucleotide hybridizes strongly with a target sequence and does not hybridize with non-target sequences at annealing temperature, i.e. the hybridization is specific.

The G/C and A/T match and mismatch counting only takes into account the effect of hydrogen bonds between Watson-Crick pairs and ignores stacking effects between neighboring bases. Thus it is not precise, and in the worst cases the estimated  $T_m$  may be significantly inaccurate. This may result, for example, in poorly or nonspecifically annealing PCR primers.

Although not designed for the evaluation of hybridization strength BLASTN (Altschul et al. 1990) is a popular tool for estimating the strength and specificity of nucleic acid hybridization, especially the number of nonspecific hybridization sites in full genomes. The main benefits of BLASTN compared to more precise methods is the speed of the alignment search and the possibility of evaluating longer sequences where the nearest-neighbor thermodynamic algorithm does not work. BLASTN is designed for calculating the evolutionary distance between 2 sequences and uses 4x4 substitution matrix of transition

probabilities for scoring alignments. By modifying the substitution matrix, it is possible to make BLASTN estimate the true non-specific hybridization probability of nucleotides with higher precision than with the default evolutionary matrix (Eklund et al. 2010).

## **Nearest neighbor thermodynamic calculation**

In theory, knowing  $\Delta G$  of hybridization at given temperature makes it possible to calculate precisely the equilibrium constant of hybridization, and thus find the optimal nucleotide sequences and parameters for experiment. In practice, the calculation of true  $\Delta G$  is extremely complicated or impossible because of the entropy contribution of the steric effects caused by the secondary or tertiary structure of nucleic acids. Thus a simplified model of  $\Delta G$  calculation is used, most commonly the nearest neighbor algorithm (NN algorithm). It gives very precise results for short oligonucleotide fragments that hybridize in almost a single-step reaction and do not form strong secondary structures. NN algorithm is not appropriate for longer sequences ( $\geq 40\text{bp}$ ).

The NN method takes into account both the hydrogen bonds between bases of opposite strands and stacking interactions between adjacent bases of the same strand, generalizing these into a single set of thermodynamic values ( $\Delta H$  and  $\Delta S$ ) that characterize hybridized dinucleotide (Borer et al. 1974). While calculating the hybridization energy of oligonucleotide sequence, it is necessary to sum the entropy and enthalpy or free energy values of each dinucleotide pair in the hybridized sequence (i.e. move along the sequence one nucleotide at a time with the 2 nucleotide-wide window). The contribution of each single base pair into global enthalpy and entropy (and  $\Delta G$ ) is distributed between 2 values – one of the left-side dinucleotide and the other the right-side dinucleotide. The thermodynamic parameters of the full sequence is the sum of the contribution of all dinucleotide pairs (Breslauer et al. 1986).

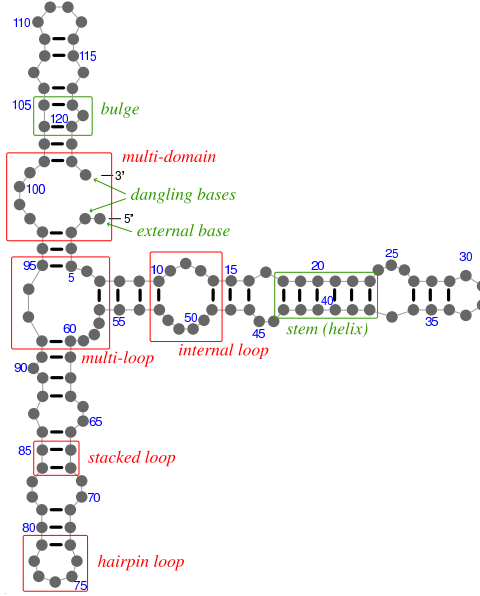
The NN algorithm can be used not only for calculating the hybridization energy of perfect reverse complementary strands, but also strands with mismatches. For the latter, energy and entropy values of mismatched dinucleotide pairs have to be known.

In addition to the contribution of paired dinucleotides, the NN method has been extended to take into account other features (SantaLucia and Hicks 2004):

- The contribution of single terminating nucleotide pair
- Terminal mismatches
- Dangling ends (i.e., where one nucleotide chain is longer than other)
- Gaps (i.e., the 2 strands are not perfectly aligned but one forms loop)
- Internal loops (i.e., a certain inner part of alignment is not hybridized state)
- Coaxial stacking (i.e., the hybridization of 2 distinct middle regions of nucleic acid chain to a single complementary template)

Examples of structural features in RNA secondary structure are shown on Figure 2.





Example of a pseudoknot-free secondary structure containing all elementary structures (Androneanu M.R. 2000)

**Figure 2.** Structural features of RNA secondary structure.

For example, the  $\Delta G$  of internal loop can be calculated using the following formula (SantaLucia and Hicks 2004):

$$\Delta G_{\text{total}} = \Delta G_{\text{loopN}} + \Delta G_{\text{asymmetry}} + \Delta G_{\text{left mismatch}} + \Delta G_{\text{right mismatch}}$$

where  $\Delta G_{\text{loopN}}$  is constant,  $\Delta G_{\text{asymmetry}}$  depends on the difference between the lengths of both sides of the internal loop, and  $\Delta G_{\text{left mismatch}}$  and  $\Delta G_{\text{right mismatch}}$  are the sequence-dependent contributions of mismatches at the ends of the loop.

The NN method can be extended to next-nearest-neighbor (NNN) method, which uses trinucleotides as elementary blocks. Although in some situations it gives even better predictions than the NN model, the gains are small compared to the difficulties in determining the correct parameters for mismatched trinucleotides (Owczarzy et al. 1999; SantaLucia and Hicks 2004).

Nearest neighbor thermodynamics can be successfully used for DNA/DNA, DNA/RNA and RNA/RNA interactions when the relevant dinucleotide hybridization parameters are known. For the ones involving RNA, this method is particularly important because the energetic contributions of nucleotide and dinucleotide pairs in the case of RNA interactions vary much more than in DNA/DNA interactions. For example, G/U mismatch in double-stranded RNA is almost as energetically favorable as the A/U Watson-Crick pair. Thus the probes designed by counting matches and mismatches may work very poorly unless the specific properties of RNA are taken into account.

By predicting the  $\Delta H$  and  $\Delta S$  of hybridization reaction, the NN thermodynamic model is also the most precise method for estimating the  $T_m$  of oligonucleotides. The most widely used formula also takes into account the concentration of cations and the nature of oligonucleotides:

$$T_m \text{ primer} = \frac{\Delta H}{\Delta S + R \ln(C_T/F)} + 12.5 \log[Na^+] - 273.15$$

where  $\Delta H$  and  $\Delta S$  are the enthalpy and entropy for helix formation respectively,  $R$  is the molar gas constant (1.987 cal/K° mol), and  $C_T$  is the total molar concentration of the annealing oligonucleotides.  $F$  describes the “symmetry” of oligonucleotide concentration. It is 1 for self-complementary oligonucleotides and 4 for non self-complementary oligonucleotides in equimolar concentration. The formula uses the concentration of sodium; if other cations are used in the buffer their concentrations can be converted to equivalent  $Na^+$  concentration with empirical formulas (Stellwagen, Muse, and Stellwagen 2011; SantaLucia, Allawi, and Seneviratne 1996).

In most cases NN thermodynamics predicts very well the energy and  $T_m$  of oligonucleotide hybridization (Breslauer et al. 1986; Rychlik, Spencer, and Rhoads 1990; Chavali et al. 2005; Hughesman, Turner, and Haynes 2011). But it is important to keep in mind that the commonly used NN thermodynamic model treats  $\Delta H$  and  $\Delta S$  values as temperature-independent. This is incorrect, overestimating oligonucleotide  $T_m$  at high temperatures. At low temperatures the NN method also loses precision because it is unable to take into account the potential secondary and tertiary structure of single-stranded oligonucleotides (Hughesman, Turner, and Haynes 2011).

A more precise estimation of  $T_m$  can be done by using the experimentally determined heat capacity change:

$$T_m = \frac{\Delta H^\circ(T_{ref}) + \Delta C_p(T_m - T_{ref})}{\Delta S^\circ(T_{ref}) + \Delta C_p \ln(T_m/T_{ref}) + R \ln(C_T/4)}$$

where  $\Delta H^\circ(T_{ref})$  and  $\Delta S^\circ(T_{ref})$  are the thermodynamic values calculated with the temperature independent NN parameters, determined at the reference temperature (usually at 37°C) and  $\Delta C_p$  is the heat capacity change ( $\sim 42$  cal mol<sup>-1</sup>K<sup>-1</sup> for a single nucleotide pair). This formula gives a significantly better estimation of oligonucleotide  $T_m$  at temperatures up to 70°C, but still loses precision at low temperatures (Hughesman, Turner, and Haynes 2011).

The NN method is also the most widely used and precise method for predicting the possible secondary structures of oligonucleotide strands. It can give precise prediction about the free energy (and thus stability) of different structures (Borer et al. 1974), but since it relies on local energy minimization, it cannot always predict the relative abundance of different structures, especially

if competition between different hybridizations is involved (Chavali et al. 2005).

NN thermodynamic calculations give accurate results only if the hybridization behaves as a one-step reaction. This mostly holds true only for nucleotide sequences of <40 base-pairs. Longer sequences hybridize in a multi-step process in which initially small double-stranded “seed” is formed. This “seed” elongates from the initial hybridization point in 2 directions, given that the flanking regions of both strands are complementary (Levitt 2009). As the reaction kinetics of initial hybridization and elongation are quite different, the NN method cannot be used to calculate hybridization properties of long nucleotide chains.

Considerable effort has gone into determining the correct NN hybridization parameters. The most complete unified hybridization model and set of NN parameters was published (SantaLucia 1998). Although the most common cases have been solved with great accuracy, the work remains on-going, e.g. for the  $\Delta H$  and  $\Delta S$  temperature dependence (Hughesman, Turner, and Haynes 2011), unnatural bases, secondary structure variants and microarray-aligned probes (Hooyberghs, Van Hummelen, and Carlon 2009). The web-based database project NNDB (<http://rna.urmc.rochester.edu/NNDB>) collects recently published NN parameters together with tutorials and references about their application (Turner and Mathews 2010).

Many programs have been devised for calculating hybridization parameters ( $\Delta H$ ,  $\Delta S$ ,  $\Delta G$  and  $T_m$ ) of nucleic acids, usually in the context of some other task (PCR primer design, hybridization probe design). **MELTING** is an extensible specialized tool for calculating hybridization properties; it supports all major types of hybridization (DNA/DNA, DNA/RNA and RNA/RNA), allows the specification of the concentrations various cation types, denaturing agents and unnatural bases (inosine and azobenzene). It uses NN thermodynamic model for short oligonucleotides and several published empirical formulas for longer sequences where NN is unreliable (Dumousseau et al. 2012). **Visual-OMP** uses very precise physical model of oligonucleotide hybridization and multi-stage equilibrium algorithms for the design and evaluation of PCR primers and probes (SantaLucia 2007).

## **Practical applications of nucleic acid hybridization prediction**

### **PCR primer design**

Polymerase Chain Reaction (PCR) primers need to have certain desirable properties:

- They have to hybridize strongly at target sites at the annealing step
- They should not hybridize non-specifically at the annealing step
- They should completely dissociate at the heating step

- They should not form strong dimers with themselves or opposite primers
- They should not form strong hairpin loops

All these requirements are statistical, being related to average primer behavior. Some primers almost always hybridize non-specifically, some form weak dimers and so on. The important point is that the strength and frequency of incorrect hybridizations have to be low to ensure that only the correct reaction is carried out to a significant extent. Thus the efficiency of PCR relies fundamentally on the thermodynamic properties of oligonucleotide hybridization.

It is crucial to avoid mispriming in the first cycles of PCR as these will be amplified at a similar concentration to the actual products. On the other hand, artefacts such as partial products and chimaeras, formed at the later stages of PCR, will remain in low concentration compared to the product (Kanagawa 2003).

In common applications, the most important parameters of the PCR primer pair are melting temperature and secondary structure formation.

Primer  $T_m$  has to be in an optimal range depending on the PCR conditions (annealing temperature) and the  $T_m$  of template DNA (Mitsubishi 1996); too low an annealing temperature (relative to the melting temperature of primers and product) can lead to the formation of false products, and too high an annealing temperature decreases the PCR yield (Rychlik, Spencer, and Rhoads 1990). Currently the best algorithm for  $T_m$  calculation is the NN thermodynamic model (Chavali et al. 2005).

PCR primers are normally 20–24 nucleotides long, i.e. well below the length threshold of the applicability of the NN model. Thus the NN algorithm can be used to predict precisely their hybridization strengths and eliminate candidate primers with wrong  $T_m$ . The most common all-purpose primer design software Primer3 uses this approach. NN thermodynamics also allow the precise calculation of the hybridization strength of the 3' sub-region of primer, which is the most important in PCR specificity (Mitsubishi 1996; Miura et al. 2005).

The NN model cannot be used to calculate the melting temperature of product since it is not applicable to long DNA sequences (Rychlik, Spencer, and Rhoads 1990).

Primer secondary structures, such as hairpins and primer dimers lower the success rate of PCR. NN thermodynamics is also used to calculate the strengths of possible primer dimers and hairpins, and eliminate those that form too strong structures. Among these structures, the most unwanted ones are those that involve 3'-ends of primers (Mitsubishi 1996). Although published NN thermodynamic models allow also the estimation of the free energy of different secondary structures of primers (SantaLucia and Hicks 2004), they are not always directly correlated with PCR success. One has to take into account that primer-product hybridization competes with the secondary structures of both primers and products, and thus the real effect on PCR success depends on the precise equilibrium between the different states (Chavali et al. 2005).

Depending on the PCR application, there often are additional requirements for primers like constraints to their length, need for specific tag at one end and so on.

Good primers should also avoid extreme GC content and certain nucleotide patterns, such as single nucleotide and CA repeats that are easily enforceable in primer design software (Yamada, Soma, and Morishita 2006; Mallona, Weiss, and Egea-Cortines 2011). The synthesis of a new DNA strand starts from the 3'-end of the primer and thus it is thought that the nucleotide composition at the 3'-end strongly influences the success rate of PCR. Both too high and too low GC content at the 3'-end is detrimental to PCR specificity (Mitsubishi 1996). Certain nucleotide triplets (WSS and SWS) are over-represented and some (WCG) underrepresented in the 3'-ends of successful primers (Onodera and Melcher 2004).

Predicting the strength and number of nonspecific hybridization sites is more complicated, because not only has the actual sequence of interest to be taken into account (PCR product with flanking regions), but the sequence of the whole DNA that may be present in PCR tube. This may, for example, be the whole human genome and/or several bacterial genomes.

A perfect match between the whole primer and template DNA is not needed for the PCR reaction to start. Hybridization is a probabilistic process and even weak non-specific hybridizations can start polymerase reaction as long as the 3'-end of the primer is in a hybridized state (Mitsubishi 1996; Miura et al. 2005; Yamada, Soma, and Morishita 2006; Kalendar, Lee, and Schulman 2011). As the primer sequence is incorporated into such false product, subsequent PCR cycles with this product work at an efficiency similar to the proper product. Thus it is often advisable to test not only the whole primer for uniqueness in source genome, but also shorter subsequences, especially the 3'-end (Mitsubishi 1996; Andreson, Möls, and Remm 2008).

On the other hand, if the hybridization between primer and template is strong but the 3' end of alignment contains mismatches, the probability of a successful start of the replication is many orders of magnitude lower (Petruska et al. 1988). But it is known that certain DNA polymerases have exonucleolytic (proof-reading) activity and can remove mismatched nucleotide from the 3'-end of the primer and then start the elongation (Skerra 1992).

To design the most specific primers, the balance between primer melting temperature and annealing temperature has to be kept in the optimal range to maximize specific and minimize the non-specific hybridization ratio. The true hybridization ratio of primers for a given temperature can be predicted from the  $\Delta G$  for both specific and nonspecific hybridization. Unfortunately, calculating the hybridization  $\Delta G$  for all nonspecific positions in the genome is a computationally costly process (Yamada, Soma, and Morishita 2006). One way to speed up the test for non-specific hybridization is to find the shortest 3' subsequence that gives significant probability of mispriming, and test for the uniqueness of such a subsequence. The precise length of such subsequences

(SDSS – specificity determining subsequence) varies and can be calculated using NN thermodynamics (Miura et al. 2005).

The simplest method to avoid primers that may give too many non-specific hybridizations with the genome is to test the source sequence for motifs that are repeated in the whole genome. Often this is done in the form of masking repeats, i.e. marking in the source sequence the regions that are locally similar to known repeating motifs and should be excluded from primer design. It is expected that the remaining parts of the sequence are locally sufficiently unique, so that even if primers designed to such regions have few nonspecific binding sites in genome, the probability of any 2 of these sites being close enough together to start a false reaction is very low.

A common and most widely used method is masking known repeating motifs in the human genome (or other known genomes). The most widely used program for this is RepeatMasker, which uses precompiled libraries of repeated sequences.

A more precise method is to find the abundance of different oligonucleotide motifs in genome and mask all those words that have too many identical or similar motifs in the genome. Many algorithms and programs have been developed over the years, but only a few of them are applicable to routine PCR primer design (Andreson, Möls, and Remm 2008; Morgulis et al. 2006).

The most popular PCR primer design tool is **Primer3**. It can design both PCR primers and (optionally) hybridization probes, taking into account many experimentally determined factors in addition to the NN model (Untergasser et al. 2012).

**Pythia** uses a small set of thermodynamic parameters to estimate the PCR efficiency and precomputed index of non-specific hybridization sites with specified thermodynamic stability (Mann et al. 2009).

**pcrEfficiency** is a Web-based tool that uses statistical model to predict the efficiency of PCR reactions (Mallona, Weiss, and Egea-Cortines 2011).

**RexPrimer** uses Primer3 as the primer design backend and uses a set of databases of genomic variance (SNPs, indels, pseudogenes) to avoid designing primers to non-unique regions (Piriyaongsa et al. 2009).

Although the in-silico primer design algorithms have evolved considerably in last 20 years they cannot guarantee the success of PCR using designed primers. Thus if failure is not an option, for example, if the primer pair is used for diagnostic purposes, experimental verification of primers cannot be avoided (Carter et al. 2010).

Primer evaluation is a time-consuming process if either a large number of primers is required or an extensive test for nonspecific hybridization is needed. Thus for certain common tasks, such as human gene expression analysis, prebuilt primer databases are created (Wang et al. 2012).

## **Multiplex PCR**

Multiplexing several PCR reactions into single tube is a very popular method to save time, chemicals, source DNA, and provide internal control of reaction success. Multiplex PCR has been successfully used, for example, in bridging gaps in shotgun sequencing (Sorokin et al. 1996; Tettelin et al. 1999), mutation and indel detection (Edwards and Gibbs 1994; Shen et al. 2010), pathogen detection (Elnifro et al. 2000; Strommenger et al. 2003; Pinar et al. 2004; Gardner et al. 2009; Shi et al. 2012), quantitative analysis of GM food (Rudi, Rud, and Holck 2003; Jinxia et al. 2011) and genotyping (Edwards and Gibbs 1994).

The number of concurrent PCR reactions in a single tube (multiplexing level) is usually in the low 10's (Edwards and Gibbs 1994; Nicodème and Steyaert 1997; Tettelin et al. 1999; Syvänen 2005) but much larger levels can be used in specific protocols (Fredriksson et al. 2007).

The average quality of a PCR is lower in a multiplex environment and is negatively correlated with the multiplex level (Edwards and Gibbs 1994; Henegariu et al. 1997; Rudi, Rud, and Holck 2003; Shum and Paul 2009). The precise causes are not well understood (Kanagawa 2003), but some probable factors can nevertheless be outlined.

### **Nonspecific hybridization of primers to source DNA that creates false products**

Although this can happen with single-plex PCR as well, its effect on multiplex PCR may be much stronger simply because there are more different primer sequences in the reaction (Sorokin et al. 1996; Rudi, Rud, and Holck 2003; Shum and Paul 2009). Each primer-primer combination can potentially initiate the synthesis of a false product if the nonspecific hybridization sites of those primers fall close enough together and have correct directionality. The number of possible combinations of primer pairs in a multiplex environment is proportional to  $2^n$ , where  $n$  is the number of PCR reactions in single tube (Syvänen 2005).

Although primer design programs can successfully exclude those that give significant probability of forming a false product between the primers of the same PCR, they cannot predict the false product between primers from different PCR.

### **Formation of primer-primer dimers**

Most primer design programs test for dimers between the primers in the same PCR and exclude pairs that hybridize too strongly with each other. But in a multiplex environment dimers between primers from different reactions can form (Edwards and Gibbs 1994; Nicodème and Steyaert 1997; Syvänen 2005). Although the hybridization strength of primer dimers is normally low, the concentration of primers is initially very high compared to the concentration of target DNA. Thus even weak hybridizations can start a polymerase reaction

(Brownie 1997). Among all possible hybridizations between PCR primers the ones involving the 3'-end are considered the most dangerous. It is proposed that the  $\Delta G$  of hybridization which includes 3'-end of both primers should be kept below 6 kcal/mol (Rachlin et al. 2005b).

### **Nonspecific interactions between primers and products**

As the concentration of all products increases with time, so does the probability of nonspecific interactions between any primer and product. Normally primers are tested so as not to form too strong alignments with their own product, but interactions with the product of other PCR are possible in a multiplex environment.

### **Differences between PCR efficiency**

Different PCR primer pairs have different efficiency (i.e. amplification speed), which introduces bias to multiplex PCR. PCR drift is variation in the product yield that is unbiased and most probably caused by the random fluctuations in the first few PCR cycles that subsequently become amplified (Kanagawa 2003). PCR selection is the biased difference between speeds of PCR reactions caused by the properties of primer or product sequences. Faster reactions may out-compete slower reactions in multiplex environment by depleting reagents. Also, if quantitative measurements are needed, the bias will cause either over- or underestimation of the specific product.

Primers with similar melting temperatures have more similar rates of amplification. Thus to achieve a more uniform amplification, it is useful to limit the maximum difference between the primer melting temperatures and product lengths in groups (Edwards and Gibbs 1994; Nicodème and Steyaert 1997; Elnifro et al. 2000; Syvänen 2005). Also calibrating primer concentrations of individual PCR reactions can be used to make reaction rates uniform (Jinxia et al. 2011). The relative speed of PCR reactions is dependent on the concentration of primers and products. In the final stage of PCR there is a strong tendency towards a 1:1 ratio of products (Kanagawa 2003).

There is greater probability of the occurrence of heteroduplexes and chimeras in multiplex PCR. A heteroduplex forms if 2 different products partially hybridize with each other. A chimera is a linear conglomerate of partial products of the same or different PCR. As these artefacts happen with higher probability in the later stages of PCR when the concentration of products is high and primers are depleted, it is advisable to terminate multiplex PCR earlier than single-plex (Kanagawa 2003).

Although not related to PCR failure, it is sometimes necessary to impose additional constraints to product length in a multiplex group. If gel electrophoresis is used to detect the result of PCR, it is necessary to limit the minimum difference between any product lengths in a multiplex group (Pinar et al. 2004).

For these reasons, it is often challenging to achieve high multiplex levels. Difficulty of distributing PCR primers into multiplexing groups is critically dependent on the average compatibility between any 2 primer pairs and the



multiplexing level required. For each compatibility and group size combination, there exists computational “phase transition”, i.e. the maximum practical group size from which, upwards, the size of necessary calculations for finding usable groups starts to increase exponentially (Rachlin et al. 2005b).

Finding the optimal groups for PCR multiplexing is NP complete problem, but simple heuristic algorithms can achieve a near optimal solution (Nicodème and Steyaert 1997).

In many cases, the experimental conditions, such as the concentration of primers, enzyme and  $Mg^{++}$ , annealing and extension times, and temperatures, can be adjusted to increase the efficiency of multiplex PCR (Henegariu et al. 1997; Edwards and Gibbs 1994; Strommenger et al. 2003; Carter et al. 2010). Increasing PCR cycle lengths guarantees that all parallel reactions have time to be completed and reduces the differences between product yields (Shum and Paul 2009). These approaches have to be adjusted for each experimental situation. Thus they are time-consuming and cannot replace the stringent primer design and grouping (Elnifro et al. 2000). Using chemically modified bases in primers can help to suppress false products and make amplification speeds more uniform (Shum and Paul 2009).

Creating optimal or near-optimal multiplex groups is a multi-objective problem (Rachlin et al. 2005a; Lee, Shin, and Zhang 2007). Simultaneously there is a need to consider the number of groups, multiplexing levels and the quality or inclusiveness of PCR. Although the achievable grouping level depends on the list of primers, it is practical to separate primer design and grouping to make the algorithm simpler. Given the initial set of chemically similar primers, the multi-objective evolutionary algorithm (MOEA) performs good grouping with a reasonable performance (Lee, Shin, and Zhang 2007). For a fixed coverage, the maximum achievable multiplexing level is proportional to  $\log(N)$  where  $N$  is the number of PCR reactions (Rachlin et al. 2005b).

Several tools and algorithms have been developed for multiplex primer design.

**MultiPLX** calculates the primer-dimer strengths using NN thermodynamic model and uses greedy algorithm to distribute a set of primer pairs into optimal groups based on their compatibility. In addition to hybridization strengths it can use user-specified compatibility values that makes it very flexible (Kaplinski et al. 2007).

**MuPlex** integrates primer design and multiplex group creation for the design of SNP genotyping primers using various criteria for multi-objective optimization (Rachlin et al. 2005a). Among the factors to be considered are  $T_m$  difference, the strengths of primer dimers, the number of non-specific hybridization sites, group sizes and SNP coverage.

**PrimerStation** uses an algorithm that predicts the actual primer hybridization ratio in liquid solution at annealing temperature, using the thermodynamic model. Good primers for multiplexing are required to have a hybridization ratio of  $>0.99$  for specific binding and  $<0.05$  for non-specific binding. As the calculation of precise  $\Delta G$  is computationally costly, PrimerStation includes

a precomputed database of potential primer candidates with good specificity (Yamada, Soma, and Morishita 2006).

**The MPP algorithm** allows the design of multiplex primers to large number of diverse targets where multiple sequence alignment (MSA) is not practical or possible (Gardner et al. 2009).

One method for reducing potential primer-dimer formation is to minimize the number of distinct primers needed for multiplexed PCR. In this case, certain primers are chosen such that they participate in many individual PCR reactions, but each pair of primers can be amplified only in a single reaction. An implementation of such primer selection algorithm is **G-POT** (Konwar et al. 2005).

The average probability of primer compatibility is also crucial to multiplex PCR design. Thus for high multiplexing levels, specific experimental techniques that increase primer compatibility with each other are needed, for example, using universal primers (Rachlin et al. 2005b).

Another option to increase the success rate of multiplex PCR is to customize PCR protocol in such a way that only the first cycles use longer specific primers and the main amplification involves universal primer sequences. This method gives more uniform amplification speeds, allowing using it for quantitative PCR (Rudi, Rud, and Holck 2003). As the universal primers are specifically designed to be not complementary with any region of source DNA and to not form dimers the formation of nonspecific products is also suppressed. But even in this case, the initial specific primers have to be designed taking into account all possible cross-reactions (Brownie 1997; Shi et al. 2012).

One method for increasing the detection rate of multiplex PCR products is to selectively pick only correct products from the amplified mixture. Correct products have terminal sequences composed of primers from single pair, whereas most false products have terminal sequences from different pairs. Specific ligation probes can be designed that hybridize to both ends of the product sequences, generating circular products that can be further amplified (Fredriksson et al. 2007).

Although considerable interest for multiplex PCR exists, and many algorithms are created for designing optimized multiplex groups, there is no fool-proof method. Thus in most cases, there is no alternative to experimental validation of multiplex groups (Henegariu et al. 1997; Elnifro et al. 2000).

## Hybridization probe design

Hybridization based technologies have become popular methods for rapid and accurate analysis of biological samples. They have been successfully used for the detection of pathogenic viruses and bacteria (Wilson, Strout, and DeSantis 2002), analyzing species composition of environmental and medical samples (Guschin et al. 1997), quantitative analysis of GM food (Rudi, Rud, and Holck 2003) and for many other types of analyzes.

Ideally hybridization probes need the following properties (Kaderali and Schliep 2002; Harrison et al. 2013; X.-Q. Xia et al. 2010)

- They should hybridize strongly with the target sequence at the annealing temperature
- They should not hybridize with other sequences that may be present in the solution (non-target sequences) at annealing temperature
- They should not form hairpin loops

Compared to PCR primers, there are a few differences in the behavior of hybridization probes.

First they are usually attached to some kind of solid surface or gel matrix. This means that their movements are constrained and the hybridization energy formula applicable to solution is not precise, although there is a clear correlation between  $\Delta G$  values obtained for probes in solution and on surfaces (Hooyberghs, Van Hummelen, and Carlon 2009; Hadiwikarta et al. 2012). How big is the difference between the thermodynamic properties of the oligonucleotides in solution and the identical nucleotides attached to surface depends on the technology used and is hard to derive analytically (J. Fuchs et al. 2010). Thus it is advisable to experimentally determine the correction factor for  $\Delta G$  and  $T_m$  values calculated using the solution formula or to determine separate sets of NN parameters for immobilized oligonucleotides (Hooyberghs, Van Hummelen, and Carlon 2009; X.-Q. Xia et al. 2010).

During hybridization, the local concentration of hybridized probes increases much more rapidly than in solution and the electrostatic repulsion between the negatively charged phosphate chains affects the hybridization kinetics (J. Fuchs et al. 2010). The uneven distribution of probes influences the hybridization behavior (Vainrub and Pettitt 2011). The surface has limited accessibility and mass transport of target molecules to probes may become the limiting step in hybridization kinetics (J. Fuchs et al. 2010).

Fluorescent labeling, often used in microarray experiments, affects the hybridization properties of oligonucleotides. Immobilization of probes to surface also reduces the difference in the intensities of fluorescent signals of specific and nonspecific hybridization compared to solution hybridization (Hooyberghs, Van Hummelen, and Carlon 2009).

If hybridization probes are used for discriminating between closely related target molecules, such as the sequences from 2 bacterial strains, the precise calculation of hybridization properties becomes extremely important. Such sequences often differ only by few nucleotides, and consequently their hybridization energies and  $T_m$  values are very close (Dong et al. 2001; Kaderali and Schliep 2002). It is not sufficient to simply design highest-affinity probes to target sequence because these probes may have considerable affinity with the control sequence, thereby giving false positive signals (Bernhard Maximilian Fuchs et al. 1998). To achieve maximum discriminatory power, the  $\Delta G$  of target hybridization has to be as negative, and  $\Delta G$  of control hybridization, as positive as possible at the precise annealing temperature. In such a case the equilibrium

of probe-target interaction is mostly in the hybridized state and the equilibrium of probe-control interaction is mostly in the free state (Figure 1).

Unlike PCR, microarray hybridization occurs at a constant temperature. Thus hybridization probes have to compete with the secondary structure of target molecules that may exist at that temperature. If this secondary structure is too strong, hybridization to probes may be suppressed because their complementary regions in target molecule are either directly or indirectly blocked (Ratushna, Weller, and Gibas 2005).

If the hybridization energy ( $\Delta G$ ) of the secondary structure of target molecule is higher ( $\Delta G$  is more negative) than  $\Delta G$  between probe and target, the secondary structure out-competes hybridization probes, and a given target will not hybridize at all. If  $\Delta G$  of the secondary structure is higher (i.e. hybridization is weaker) than between the target and the probe, the probes will eventually hybridize, but the process may be very slow. The cause of this is that at every moment most target molecules have secondary structures, and the regions complementary to probes are blocked. For hybridization to occur the target molecule has first to loosen and then come to the proximity of a hybridization probe. Both of these processes have a low probability and thus the combination of 2 has even lower (Dong et al. 2001).

The blockage of hybridization by the secondary structure of a target molecule is more prevalent in RNA targets because their secondary structure is stronger than that of DNA (T. Xia et al. 1998). Certain native RNA molecules (e.g., rRNA and tmRNA), have strong evolutionary conserved structures that make their hybridization challenging (Ratushna, Weller, and Gibas 2005; Yilmaz, Okten, and Noguera 2006). This is especially true if the hybridization has to take place at lower temperatures, at which RNA secondary structure is very stable (Peplies, Glockner, and Amann 2003).

There are several approaches to break RNA secondary structure and make their hybridization possible. The simplest and most used method is either to increase hybridization temperature or use chemicals that lower the  $T_m$  (i.e. lower the free energy of nucleic acid hybridization). Both of these affect not only the secondary structure formation, but also probe-target hybridization energy and  $T_m$ , but as this hybridization is normally stronger than the secondary structure of target, the overall hybridization speed increases (Ratushna, Weller, and Gibas 2005; Small et al. 2001).

Another method is breaking the target molecule into smaller parts. Although the effective concentration of the target is lowered this way, since breakage sometimes occurs in the complementary region, the remaining fragments have much weaker structures and the hybridization is more effective (Pozhitkov et al. 2006; Ratushna, Weller, and Gibas 2005).

If the precise secondary structure of the target molecule can be determined, it is possible to design hybridization probes with this taken into account (Ratushna, Weller, and Gibas 2005). As the secondary structure is often critically dependent on few mutations in primary sequence such probes may have great discriminatory power even under non-stringent conditions. For

example, probes that extend both sides of stem-loop structures can be designed. These probes can only hybridize on the correct structure, and are discriminatory even at temperatures as low as 4°C (Dong et al. 2001).

Another method is using small specific helper nucleotide molecules (chaperones) that hybridize to certain regions of the target molecule. They block the complementary or semi-complementary regions inside the target and thus the secondary structure cannot form. Although chaperones themselves block certain regions in the target, other regions become available for hybridization (Peplies, Glockner, and Amann 2003; Small et al. 2001; B. M. Fuchs et al. 2000; Kaplinski et al. 2010).

Compared to hybridization probes, chaperones are much more capable of competing with native secondary structure for 2 reasons. First, they may be applied at a temperature higher than the hybridization (annealing) temperature when the secondary structure is much weaker. If the temperature is subsequently lowered, a strong secondary structure in the target molecule cannot form. Second, unlike bound probes, chaperones are added to the solution at much higher concentration than that of the effective hybridization probes; thus they can interact with target molecules much more freely compared to hybridization probes that are attached to a solid matrix and are in low concentration.

There are many tools available for microarray probe design, often tailored for specific applications and technologies. **Oligoarray** is a widely used program using the NN thermodynamic model to choose probes with precise hybridization  $T_m$  (Rouillard, Zucker, and Gulari 2003). **ChipD** allows the design of thermodynamically similar probes that uniformly cover the whole length of source DNA sequence (Dufour et al. 2010). **OligoWiz** can use sequence annotation information, such as exon-intron structure in design process (Wernersson and Nielsen 2005). **Teolenn** is built as an extensible framework that allows users to prioritize between different quality scores (Jourden et al. 2010).

# PRESENT INVESTIGATION AND DISCUSSION

## Aims of the present study

The main goal of the present study has been to investigate how to apply the well-known models of oligonucleotide hybridization to high-level experimental procedures of multiplex PCR and microarray hybridization.

The specific aims were:

1. To develop an efficient algorithm and implementation for estimating primer-primer and primer-product interactions in multiplex PCR and automatic grouping of PCR primers, based on the worst-case interaction strengths.
2. To find the main primer sequence-related factors that affect the PCR success rate in the multiplex PCR environment.
3. To develop an efficient method for the design of DNA hybridization probes for closely related RNA targets.
4. To study the effect of RNA secondary structure on hybridization efficiency and to develop a method to increase the RNA/DNA hybridization at low temperatures.

## Method for automatic distribution of PCR into multiplex groups (REF. I)

Multiplexing PCR is a powerful method for lowering the cost of experiments. While primer-design software can predict the occurrence of primer-dimers and false products between the primers of single PCR, it cannot take into account the possibility of unwanted hybridizations between the primers and products of different PCR amplified together. The complexity of finding a compatible set of primer pairs to be amplified together grows exponentially, depending on the desired number of reactions in a single tube. We developed an automated tool, **MultiPLX** that calculates the strengths of different primer-dimers and primer-product interactions between all primer combinations in a given list of primer pairs using the NN thermodynamic model, and distributes these into optimized groups.

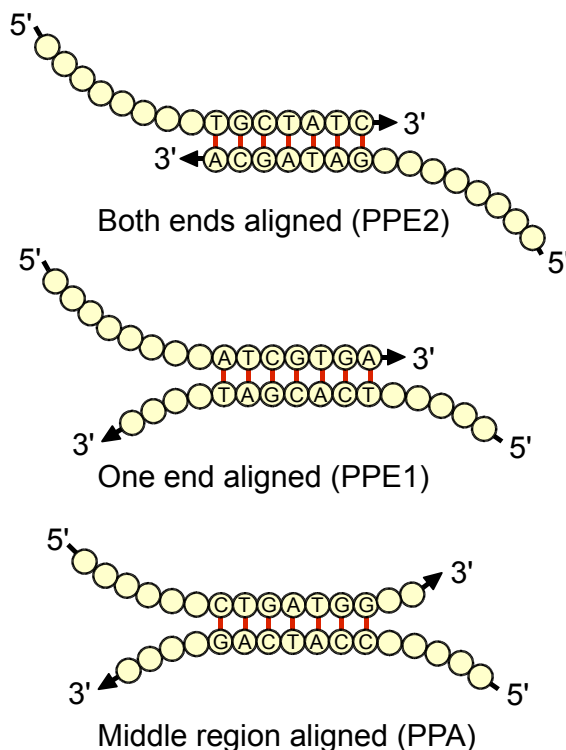
### MultiPLX Algorithm

MultiPLX calculates 8 different parameters (scores) for each combination of 2 primer pairs.

1. Maximum binding energy (minimum  $\Delta G$ ) of 2 primers including 3'-ends of both primers
2. Maximum binding energy of 3'-end of one primer with any region of another primer
3. Maximum binding energy of any region of different primers
4. Maximum binding energy of 3'-end of one primer with any region of a PCR product

5. Maximum binding energy of any region of a primer with any region of a PCR product
6. Maximum product length difference between compared PCR primer sets
7. Minimum product length difference between compared PCR primer sets
8. Maximum difference in primer melting temperatures between compared PCR primer sets.

A schema of the hybridization types corresponding to the first 3 parameters is shown in Figure 3.



**Figure 3.** Schematic diagram of different primer dimer variants whose hybridization energies have been calculated by the MultiPLX algorithm. The variants differ in whether both, one or neither 3'-ends are in the hybridized state.

MultiPLX uses the NN thermodynamic method for the calculation of both the strength of local hybridizations between primers and products, and calculating the melting temperature of primers. By default, it uses published entropy and enthalpy values (Kaderali 2001) but the user can also submit a custom thermodynamic table if a newer or better adjusted version is available.

In addition to scores calculated by MultiPLX, the grouping algorithm can also include one additional user-specified score value. This allows the incorporation of the results of other primer evaluation and compatibility tests into the

grouping; for example taking into account the probabilities of the false products caused by non-specific hybridizations of primers from different pairs.

The calculated scores are used to find optimized group layout by keeping the scores between the members of any single group below specified cut-off values. MultiPLX uses a 2-tier approach by first distributing primer pairs into groups by a greedy algorithm and then equalizing group sizes by moving compatible primer pairs from big to smaller groups. If no more primers can be moved into smaller groups, but groups remain unequal size, it can perform an additional “shuffling” step by exchanging primers randomly between groups while preserving the compatibilities in the group.

### **Determining the factors that predict the failure of multiplex PCR (REF IV)**

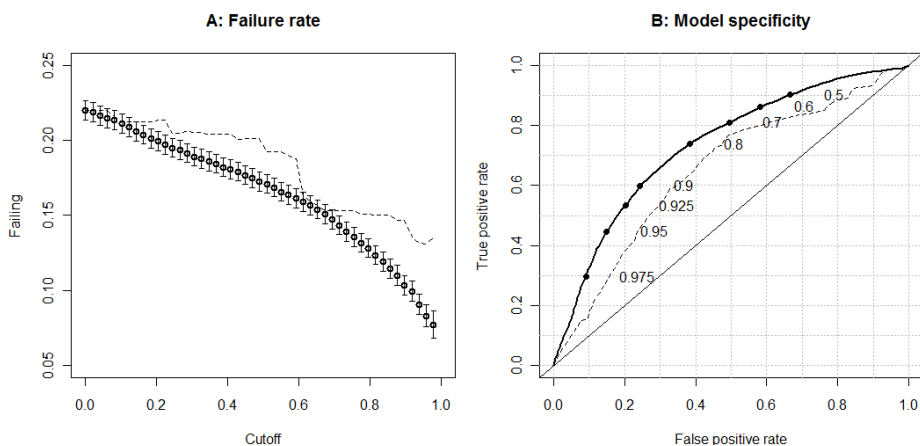
The success rate of PCR is lower in multiplex PCR, but the causes of this phenomenon are not well understood. We carried out a series of experiments to determine which parameters of primers are most strongly correlated with the failure of PCR in a multiplex environment. We were especially interested whether a) certain primer pairs disturbed (poisoned) other pairs that would normally work well, and b) whether certain combinations of otherwise good primer pairs were failing.

We created 8000 4-plex groups from primer pairs that worked flawlessly in previous single-plex control experiments. The reaction conditions were exactly the same as in the single-plex experiment, but roughly 1 in 4 reactions failed in multiplex. We created the best logistic model for predicting failure of multiplex PCR using primer pair parameters as independent variables. The factors incorporated into the best model would then correspond to the properties of primer pairs causing PCR failure in the multiplex experiment.

The analysis shows that the most important factors causing the failure of multiplex PCR are GC content of primers and the number of non-specific hybridization sites on the template DNA. Both of these factors also influence the quality of single-plex PCR in a similar way (Andreson, Möls, and Remm 2008). The more non-specific hybridization sites PCR primers had, the higher the probability of PCR failing in the multiplex environment. The detrimental effect of non-specific hybridization to PCR quality was thus similar, but much stronger, than in single-plex PCR.

We also found that the number of non-specific hybridization sites affected both the outcome of the same PCR and of other PCR in the same multiplex group (Figure 4). Primer pairs with a high number of non-specific hybridization sites caused the failure of other PCR in the same tube in addition to failing themselves. Thus these primers in particular should be avoided in multiplex experiments, and only high-quality primers should be used.





**Figure 4.** Improvement of multiplex PCR quality after applying PCR group quality model. Left: improvement of average PCR success rate after elimination of all primer pairs with predicted failure rate above a certain cut-off value. X-axis: applied cut-off value. Y axis: the ratio of failing PCR reactions. Dashed line: comparable failure rate of a model that ignores other group members. Right: ROC plot of model prediction. X-axis: the ratio of remaining failing primer pairs (of all failing pairs) after eliminating those with the prediction below the cut-off value. Y-axis: the ratio of remaining working primer pairs (of all working pairs). Dashed line: comparable plot of a model that ignores other group members. Cut-off values from 0.5–0.975 are marked on the ROC curve.

Our model indicates that single primer properties are significantly more important than interactions between primers from different pairs in predicting the failure rate of multiplex PCR. The interactions between primers had no statistically significant influence on the failure rate in our 4-plex experiments.

### SLICSeI (REF III)

Using bacterial tmRNA for species and strain discrimination poses specific challenges. First the tmRNA sequences of closely related species are very similar, making the number of possible discriminating probes limited. Second, tmRNA is known to fold into a strong secondary structure such that only certain parts of the whole molecule are exposed. And third, RNA/DNA hybridization is much stronger and less specific than DNA/DNA hybridization.

To design the probes with maximum discriminatory power, the precise hybridization energies of the probes has to be calculated with all possible target and control sequences, and ensured that all targets hybridize with  $\Delta G$  below, and all controls with  $\Delta G$  above zero. The bigger the  $\Delta G$  difference between the hybridization of a probe with 2 RNA molecules ( $\Delta\Delta G$ ), the higher will be the ratio of fluorescent signal intensities. Thus to get optimal detection probes, one

has to keep the worst case  $\Delta\Delta G$  between any target and control hybridization above a certain cut-off value.

In certain cases, it is useful to have probes that are specific to multiple target sequences – for example, this applies with a single probe that has to detect the presence of any one of multiple pathogenic strains of a bacterium, with the non-pathogenic strains being control group. In this situation, it is important to keep the hybridization levels of probe with any of these alternate targets as close as possible. This can be ensured by keeping the  $\Delta\Delta G$  of any 2 target hybridization below certain cut-off values.

We developed custom application SLICSel to do an exhaustive search among all possible hybridization probes with specified melting temperatures and choose the ones that match the abovementioned criteria. The probes picked by SLICSel are designed to discriminate between 2 sets of sequences, the target and the control sets. The  $\Delta\Delta G$  cut-offs of target and control hybridization can be specified by the user, among other relevant parameters, including annealing temperature, desired probe length and so on.

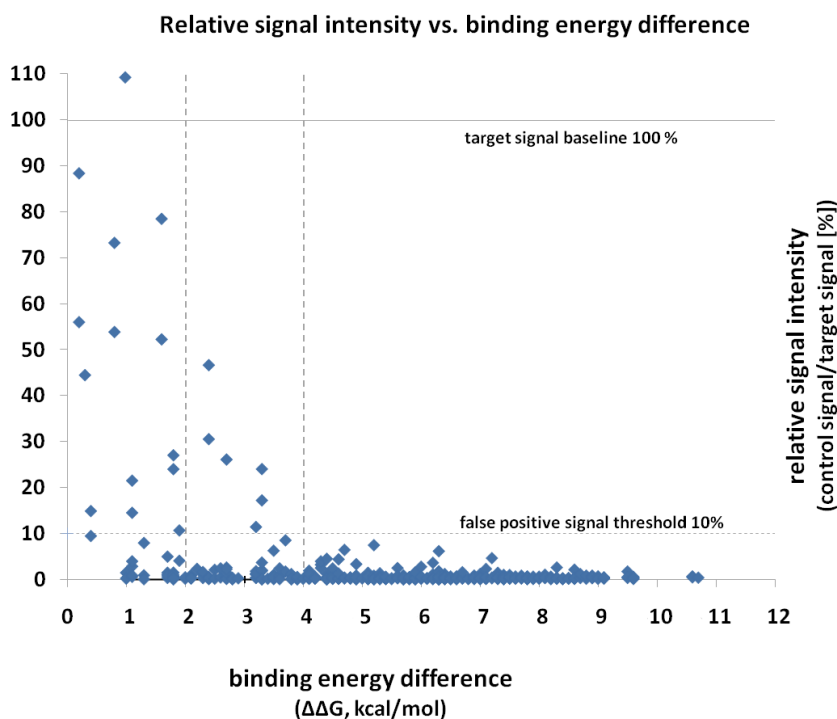
Both DNA/DNA and DNA/RNA thermodynamic tables are included with SLICSel. It also allows the user to specify a custom table when very specific reaction conditions are needed.

SLICSel also accepts degenerate nucleotide symbols in both target and control sequences. In both instances the worst case value is used – the highest  $\Delta\Delta G$  for target and lowest  $\Delta\Delta G$  for control hybridization.

SLICSel has 3 unique properties: a) it does search by brute-force and lists all probe candidates that have at least the user-provided sensitivity and specificity; b) it designs probes that selectively detect any one sequence from a list of target sequences in the environment that may contain any or all of the control sequences; and c) it allows custom thermodynamic values to be used.

To verify our algorithm, we designed 97 probes to *Streptococcus pneumoniae* tmRNA with  $\Delta\Delta G$  varying from 0 to  $-11$  kcal/mol and probe length from 9 to 26 bp. Five other bacterial tmRNA sequences were used as controls: *S.pyogenes*, *S.agalactiae*, GrC/G *Streptococcus*, *K.pneumoniae* and *M.catharralis*. All tmRNA molecules were synthesized in vitro and hybridization experiment were done by microarray.

The majority of designed probes had good specificity to *S.pneumoniae* tmRNA. Of the 463 hybridization events only 20 had false positive signals that were stronger than 10% of the true positive signal level. The relative intensity of non-specific signals was inversely proportional to  $\Delta\Delta G$ . By setting  $\Delta\Delta G$  cut-off to 2 kcal/mol, only 6 false positive hybridization events remained, and no such event was detected by setting  $\Delta\Delta G$  cut-off to 4 kcal/mol (Figure 5).



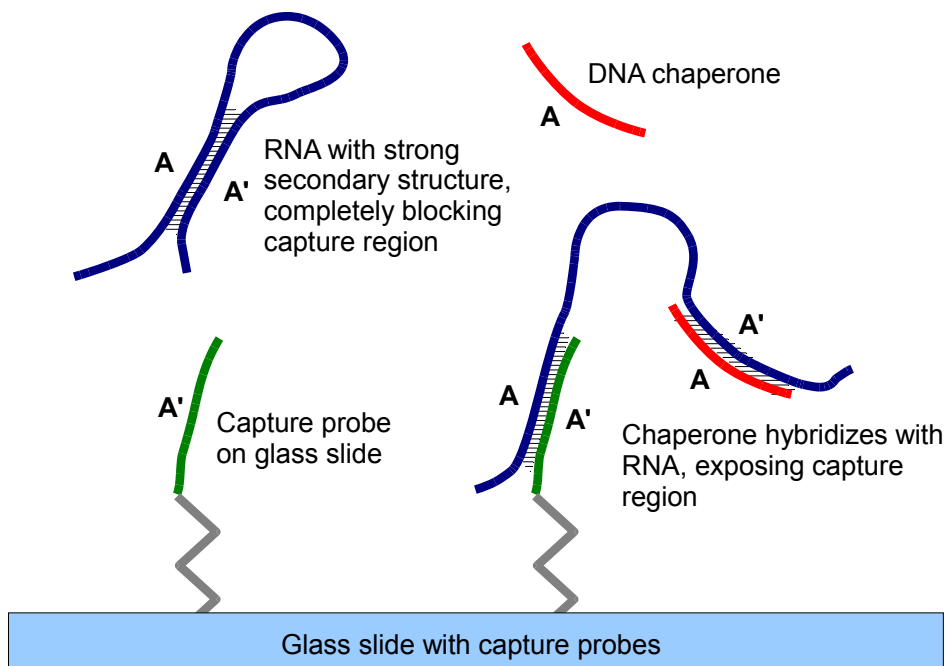
**Figure 5.** Relative intensity of false positive signals depending on the hybridization energy difference ( $\Delta\Delta G$ ) between specific and strongest nonspecific hybridization (Scheler et al. 2011).

## Chaperones (REF II)

Detection of RNA molecules with DNA hybridization probes works well at temperatures above 45°C. Hybridization signals are weak at lower temperatures, although according to NN thermodynamics, the hybridization should be stronger at lower temperatures.

We performed a series of experiments with the microbial detection microarray using different hybridization times and verified that the weak fluorescent signals were caused by a slow hybridization speed. At hybridization temperatures above 42°C, signal strengths reached plateau in < 4 h, and at 34°C they did not reach equilibrium even after 12 h.

The slow hybridization speed may be caused by strong secondary and tertiary structures of the tmRNA molecule. We assessed the possibility of improving microarray hybridization signals by designing short complementary DNA oligonucleotides, which will potentially block secondary and tertiary structure forming regions in tmRNA and thus keep the target molecule accessible to microarray probes (Figure 6).



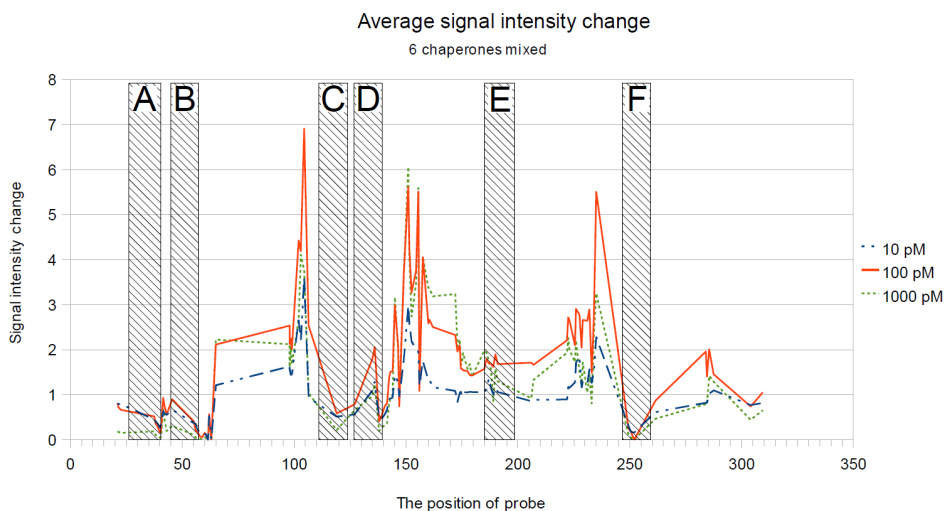
**Figure 6.** The working principle of DNA chaperones. Chaperones block the secondary structure formation and expose other parts of the molecule by hybridizing to the complementary region of RNA.

We designed 6 different helper oligonucleotides (chaperones), each complementary to one predicted functional domain of *S.pneumoniae* tmRNA. Chaperones were applied at elevated temperature, the solution was cooled and hybridization carried out.

We used previously designed detection microarray with 97 specific probes that covered most of the length of *S.pneumoniae* tmRNA. This allowed us to measure the changes in signal strength (i.e. hybridization speed) simultaneously at different regions of target molecule.

In the first experiment we applied all 6 chaperones simultaneously.

As expected, chaperone oligonucleotides suppressed these signals which had probes partially overlapping with chaperone region. All other signals increased on average, confirming our hypothesis. There were also visible peaks in the relative signal strength, suggesting regions that were almost completely blocked in the “native” state but that became well-exposed after introducing the chaperones (Figure 7).



**Figure 7.** Relative change in the signal intensities of all 97 tmRNA-specific probes after applying the mix of all 6 chaperones. X axis – position of probe midpoint on tmRNA molecule. Y axis – relative change in signal intensity of the experiments with chaperones compared to the untreated base. (Kaplinski et al. 2010)

In the next experiment chaperones were applied one-by-one, allowing us to study, which regions of tmRNA would become accessible by applying a specific chaperone.

As in the first experiment, the signals whose probes were overlapping with chaperones were suppressed. For each chaperone signals increased significantly in some other areas of tmRNA. For chaperone F, these regions were the immediate neighboring areas, suggesting the existence of a hairpin loop. For chaperones A and E, the regions with improved signal strengths were separated, suggesting the existence of more complex tertiary structure.

## CONCLUSIONS

In this investigation the oligonucleotide hybridization model was applied to multiplex PCR primer design and grouping, and to microarray hybridization.

1. We designed an efficient algorithm to partition existing PCR primer pairs into multiplex groups based on the minimum required compatibility between primers, and implemented it as a software package MultiPLX. The compatibility is defined as keeping the primer-primer or primer-product dimer hybridization energy below a specified cut-off value. MultiPLX calculates the hybridization energies using the NN thermodynamic model.
2. We analysed statistically which sequence-related properties of PCR primers are most strongly correlated with the failure of PCR in the 4-plex environment. The most important factors were primer GC content and the number of nonspecific hybridization sites of primers to genome. The more non-specific hybridization sites primer had the lower was the average success rate of the multiplex PCR. Also primers with many hybridization sites not only performed poorly themselves, but disturbed the PCR of other primer pairs in the same tube.
3. We implemented the program SLICSel that designs hybridization probes by exhaustively searching specified target sequences. By using the NN thermodynamic model, SLICSel picks all potential hybridization probes with the correct melting temperature and specified minimal difference between the specific and nonspecific hybridization strengths. We demonstrated experimentally that using a  $\Delta G$  cut-off of 4kcal/mol at the annealing temperature was sufficient to ensure probe specificity in a wide range of probe lengths and GC content.
4. We designed a set of 6 specific helper oligonucleotides (chaperones) that hybridized to a tmRNA molecule and prevented the formation of secondary structure. By using these helper oligonucleotides, we could increase the hybridization speed at 34°C up to 4-fold. We also demonstrated how the hybridization of chaperone on one part of the target molecule can increase signals in distant parts of the molecule.

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## SUMMARY IN ESTONIAN

### **Oligonukleotiidide hübridisatsioonimudeli rakendamine PCR-i ja mikrokiipide optimeerimiseks**

Nukleiinhapped on põhiliselt tuntud kui “digitaalse” geneetilise koodi kandjad kõigis teadaolevates elusrakkudes. Nukleiinhapete monomeeride (nukleotiidide) võimalik järjestus ühes DNA ahelas on praktiliselt ilma kitsendusteta ja mõjutab kaksikhelikaalse molekuli keemilisi ja füüsikalisi omadusi ainult vähesel määral. Siiski võivad nukleiinhapped, eelkõige üheaahelaline RNA, moodustada spetsiifilisi kolmemõõtmelisi struktuure, mis võivad rakkudes omada metaboolseid või juhtfunktsioone.

Kaksikhelikaalse struktuuri spontaanne moodustumine antikomplementaarsete lõikude vahel samas DNA või RNA molekulis või erinevate molekulide vahel (hübriidiseerumine) on aluseks nukleiinhapete kasutamisele biotehnoloogias, näiteks DNA polümeraasi ahelreaktsioonis (PCR) ja hübridisatsioonikiipidel. Paljudes hübridisatsiooni rakendustes käsitletakse lühikesi nukleiinhapete fragmente “digitaalsete” koodidena mis tunnevad spetsiifiliselt ära vastavaid antikomplementaarsete fragmente. Täpsema, universaalsema ja kvaliteetsema tulemuse saamiseks on aga vaja tunda hübridisatsiooni keemilisi aluseid ning käsitleda hübridisatsiooni kui tasakaalulist ja stohhastilist protsessi. Sellisel juhul on võimalik nii spetsiifilise kui mittespetsiifilise hübridisatsiooni tugevust (sulamistemperatuuri  $T_m$ ) täpselt ennustada, kasutades hübridisatsiooni termodünaamilist mudelit. Kaasajal on kõige täpsemaks selliseks mudeliks lähima naabri mudel (Nearest Neighbour Model), mis arvestab nii antikomplementaarsete lämmastikaluste vaheliste vesiniksidemete kui kõrvutiasuvate lämmastikaluste vaheliste Wan de Waals’i jõudude mõju.

Nii PCR-is kui detektsioonikiipidel on sagedaseks probleemiks mittespetsiifiline hübridisatsioon. Ehkki see on üldjuhul nõrgem kui spetsiifiline hübridisatsioon, on potentsiaalseid mittespetsiifilisi seondumiskohti sageli palju rohkem. PCRi korral võib mittespetsiifiline hübridisatsioon põhjustada praimerite dimeeride ja valeproduktide teket. Detektsioonikiipide korral põhjustab mittespetsiifiline hübridisatsioon valepositiivseid signaale. Hübridiseerumise termodünaamiline mudel võimaldab hinnata täpselt valseondumiste sagedust teatud temperatuuril ja seega vältida PCR praimereid või detektsiooniproove, millel on suur tõenäosus mittespetsiifiliseks hübridisatsiooniks.

Käesoleva doktoritöö üheks eesmärgiks oli täiustada multipleks-PCR (mitme praimeripaari koosamplifitseerimine) metoodikat luues algoritmi ja programmi praimerite automaatseks grupeerimiseks nii, et minimeerida nendevaheliste ebasoovitavate interaktsioonide võimalusi. Töö raames loodi programm MultiPLX, mis arvutab lähima naabri termodünaamilise mudeli abil kõikvõimalike interaktsioonide hübridisatsioonienergiad ning jaotab praimeripaarid automaatselt gruppidesse nii, et nendevaheliste interaktsioonide tugevus jääks alla etteantud nivoo. Lisaks võimaldab MultiPLX arvestada grupeerimisel produktide pikkusi ja praimerite sulamistemperatuure. Grupeerimiseks kasutab

MultiPLX kiiret lähendusmeetodit paigutades praimeripaarid alguses esimesse sobivasse gruppi. Järgmise sammuna toimub gruppide suuruste ühtlustamine praimeripaaride ümbertõstmise ning vahetamise teel.

Teiseks eesmärgiks oli uurida millised PCR praimerite primaarjärjestuse omadused on seotud multiplex PCR ebaõnnestumisega. Selleks teostati 8000 4-pleks katset kasutades eelenevalt ühe praimeripaariga (single-plex) PCR katsetes verifitseeritud praimeripaare inimese genoomi juhuslike lõikude amplifitseerimiseks. Kokku ebaõnnestus PCR multipleks katsetes 22,6% reaktsioonidest.

Esimese etapina tehti kindlaks, et praimeripaaride mittetöötamine multipleks katsetes ei ole juhuslik, vaid ebaõnnestumise tõenäosus varieerub erinevate praimeripaaride vahel. Seejärel analüüsiti erinevaid praimeripaaride ja produktide primaarjärjestusega seotud parameetreid laiendatud lineaarse statistilise mudeli (GLM) abil. Kõige olulisemad PCR ebaõnnestumisega korrelatsioonis olevad parameetrid olid praimerite GC sisaldus ning praimerite mittespetsiifiliste seondumiste arv genoomile. Lisaks leiti, et praimeripaarid, millel on suur mittespetsiifiliste seondumiste arv, häirivad ka teiste samas grupid amplifitseeritavate praimeripaaride reaktsioone.

Kolmandaks eesmärgiks oli luua kiire algoritm ja programm täieliku hübridiseerimisproovide komplekti loomiseks geneetiliselt lähedaste bakteriliikide eristamiseks tmRNA primaarjärjestuse põhjal. Töö käigus loodi programm SLICSel, mis arvutab lähima naabri termodünaamika põhjal välja kõigi võimalike oligonukleotiidide potentsiaalsed seondumistugevused sihtmärk- ja kontrolljärjestustele etteantud temperatuuril. Nende hulgast on võimalik automaatselt valida alamhulk mille korral seondumistugevuste erinevus spetsiifilise ja mittespetsiifilise seondumise vahel on suurem etteantud lävest.

SLICSel programmi abil disainitud proove testiti eksperimentaalselt bakteriaalse tmRNA detekteerimiseks. Leiti, et spetsiifilise ja mittespetsiifilise seondumistugevuse  $\Delta G$  erinevus 4 kcal/mol on piisav valepositiivsete signaalide vältimiseks.

Neljandaks eesmärgiks oli leida meetoodika bakteriaalse tmRNA signaalide intensiivsuse tõstmiseks detektsioonikiibil kui hübridiseerumine toimub madalal temperatuuril. Tehti kindlaks, et madal signaalide intensiivsus oli tingitud aeglasest hübridiseerumisest, mida tõenäoliselt põhjustas tmRNA tugev sekundaarstruktuur. Töö käigus disainiti kuus spetsiifilist oligonukleotiidi (chaperoni), mis hübridiseerusid tmRNA erinevate regioonidega ning eeldatavalt pidid blokeerima sekundaarstruktuuri moodustumise. Selliselt töödeldud tmRNA hübridiseerumissignaalid olid madalatel temperatuuridel märgatavalt tugevamad. Oodatavalt blokeerisid chaperonid need signaalid, mille proovid hübridiseerusid chaperonidega kattuvatele regioonidele. Seevastu tõusid teiste signaalide intensiivsused, sealhulgas nii nende, mille proovid asusid chaperoni lähedal, kui ka nende, mis asusid tmRNA molekuli kaugetes regioonides.

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## **PUBLICATIONS**

## CURRICULUM VITAE

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2002–2004 Biodata Ltd. IT specialist  
2004– Estonian Biocentre programmer  
2007– PhD student, Department of Bioinformatics, University of Tartu,  
Institute of Molecular and Cell Biology

### Scientific work:

My main research projects have focused on applying oligonucleotide hybridization model on PCR primer design and microarray hybridization. I have implemented an algorithm for automatic calculating of inter-primer interaction and optimized grouping for multiplex PCR. For microarray hybridization I developed a probe selection algorithm and program, and studied the possibility of breaking the secondary structure of target RNA molecules. I have also analyzed the primer-specific factors related to the failure of multiplex PCR.

### List of publications:

1. Koua D, Laht S, Kaplinski L, Stöcklin R, Remm M, Favreau P, Lisacek F. (2013) Position-specific scoring matrix and hidden Markov model complement each other for the prediction of conopeptide superfamilies. *Biochim Biophys Acta*. 2013;1834(4):717–24
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4. Laht S, Koua D, Kaplinski L, Lisacek F, Stöcklin R, Remm M. (2011) Identification and classification of conopeptides using profile Hidden Markov Models. *Biochim Biophys Acta*. 2012;1824(3):488–92
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10. Mägi R, Kaplinski L, Remm M. (2006) The whole genome tagSNP selection and transferability among HapMap populations. *Pac Symp Bio-comput*. 2006:535–43
11. Andreson R, Reppo E, Kaplinski L, Remm M. (2006) GENOMEMASKER package for designing unique genomic PCR primers. *BMC Bioinformatics*. 2006;7:172
12. Kaplinski L, Andreson R, Puurand T, Remm M. (2005) MultiPLX: automatic grouping and evaluation of PCR primers. *Bioinformatics*. 2005; 21(8):1701–2

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### Teadustegevus:

Minu teadustöö on põhiliselt keskendunud oligonukleotiidide hübridisatsioonimudeli rakendamisele PCR praimerite disainis ja hübridisatsioonikiipidel. Ma olen kirjutanud algoritmi ja programmi praimeritevaheliste interaktsioonide arvutamiseks ning nende optimeeritud grupeerimiseks multipleks PCR jaoks. Detektsioonikiipide jaoks olen kirjutanud proovidisaini algoritmi ja programmi ning uurinud võimalusi RNA sekundaarstruktuuri lõhkumiseks. Viimasel ajal olen tegelnud mudelitega, mis võimaldaksid ennustada multipleks PCR ebaõnnestumise tõenäosust praimerite omaduste põhjal.

### Publikatsioonid:

1. Koua D, Laht S, Kaplinski L, Stöcklin R, Remm M, Favreau P, Lisacek F. (2013) Position-specific scoring matrix and hidden Markov model complement each other for the prediction of conopeptide superfamilies. *Biochim Biophys Acta*. 2013;1834(4):717–24
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