PrimerStation: a highly specific multiplex genomic
PCR primer design server for the human genome

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ABSTRACT
PrimerStation (http://ps.cb.k.u-tokyo.ac.jp) is a web service that calculates primer sets guaranteeing
high specificity against the entire human genome. To achieve high accuracy, we used the hybridization
ratio of primers in liquid solution. Calculating the status of sequence hybridization in terms of the stringent
hybridization ratio is computationally costly, and no web service checks the entire human genome and
returns a highly specific primer set calculated using a precise physicochemical model. To shorten the
response time, we precomputed candidates for specific primers using a massively parallel computer with
100 CPUs (SunFire 15 K) about 3 months in advance. This enables PrimerStation to search and output
qualified primers interactively. PrimerStation can select highly specific primers suitable for multiplex
PCR by seeking a wider temperature range that minimizes the possibility of cross-reaction. It also allows
users to add heuristic rules to the primer design, e.g. the exclusion of single nucleotide polymorphisms
(SNPs) in primers, the avoidance of poly(A) and CA-repeats in the PCR products, and the elimination
of defective primers using the secondary structure prediction. We performed several tests to verify the
PCR amplification of randomly selected primers for ChrX, and we confirmed that the primers amplify specific
PCR products perfectly.

INTRODUCTION
There are pressing needs to detect genomic polymorphisms
and alterations with high accuracy. For example, recent studies
have revealed that large-scale copy number polymorphisms
in the human genome contribute to human genetic variation
(1), and genome alterations could be a latent cause of cancer
(2,3). Despite the long history of research, the selection of
highly specific genomic PCR primers that do not hybridize
anywhere else in the genome, except for the target, remains
a challenging task. Although some software programs con-
sider the specificity against target sequences, some of
these programs (4–8) simply check the specificity in terms
of sequence similarity using text-matching algorithms for
the prompt selection of primers but fails to take into account
real primer hybridization in a liquid solution.

Since the precise quantification of real primer hybridiza-
tion is difficult to achieve, many software programs (9–18)
evaluate the specificity of a primer in terms of a rough indi-
cator, the melting temperature at which 50% of the copies of
the primer hybridize the target sequence in a liquid solution.
With this alternative standard, a better primer has a larger
margin that separates the melting temperature for the target
sequence and those for off-targets. Therefore, we should
select the target-specific primer that maximizes the margin.
Currently this physicochemical model is accepted as the
best for designing primers.

Stringent requirements for genomic PCR primers
Care has to be taken to select an annealing temperature
because a lower annealing temperature may yield unrelated
PCR products of off-target genes while a higher value is
likely to fail to amplify the target. Annealing temperature
must be in between the two extremes in order to guarantee
effective amplification of the target genomic sequence
while minimizing the risk of cross-reaction with off-targets.
To amplify the target genomic sequence effectively, the
hybridization ratio (19) of the primer to the target should be
close to 1, e.g. >0.99, while to avoid generating false-positive
PCR products, the hybridization ratio to any off-target should
be minimized, e.g. <0.05. Let us call the temperature for
the primer ‘executable’ if this requirement is satisfied. We
require that the annealing temperature should be executable. This demand may appear to be too stringent. Nevertheless, we hold that it is a prerequisite for designing genomic PCR primers with executable temperatures because the human genome is always "expressed" and is much longer than the total of its coding regions. Hence, cross-reactions with off-targets must be eliminated effectively. Unfortunately, the design scheme based on melting temperature may not meet this prerequisite because it only ensures the weaker condition that the hybridization ratio of the primer to the target is > 0.5, while that to any off-target is < 0.5.

Considering the stringency of executable temperatures, one might be concerned with the feasibility of selecting primers with executable temperatures. However, we obtained a positive answer to this question. Figure 1 illustrates how the hybridization ratios (y-axis) of a particular primer to the target itself and four representative off-targets drop as the temperature (x-axis) increases. Note that when the temperature is below 60°C, the hybridization ratios approach 1.0, indicating that the primer is likely to hybridize to off-targets as well as to the target. The vertical band in the middle that ranges from 71 to 73°C displays executable temperatures because the hybridization ratio of the primer to the target is over 0.99 and the hybridization ratios to off-targets are no more than 0.05. In this respect, a better primer has a wider range of executable temperatures. Therefore, we developed a program for selecting the best primer with the widest range of executable temperatures, which allowed us to design qualified primers for human genes.

This positive result motivated us to develop a system called PrimerStation for designing multiplex genomic PCR primers because there is no such comprehensive web service for the human genome. In addition, these primers are applicable to a variety of other studies, such as designing sequence tagged site (STS) markers. Although there exist some software programs (20,21) that calculate the hybridization ratio, PrimerStation focused on the possible annealing temperature considering the hybridization ratio.

### Multiplex genomic PCR primers

To design primers for multiplex PCR, we need to adjust the amplification conditions for each primer so that the PCR product sizes of individual target sequences are separated sufficiently. Most existing multiplex primer design software programs use melting temperature for this adjustment. However, the large discrepancy between the melting temperatures of designed primers makes it difficult to perform PCR amplification under the same conditions. To overcome this problem, we need to look for a set of primers for individual target sequences in the human genome that share common executable temperatures. Once such a set of primers is found, any common executable temperature suffices for PCR amplification using all the primers in the set under the same conditions. Further improvement can be achieved by selecting the optimal set of primers that maximize the range of common executable temperatures. PrimerStation attempts to output the optimal set of primers for the given target sequences.

### Experimental result

Before we describe PrimerStation in detail, we present an experimental result that uses primers designed using PrimerStation. Figure 2 shows the result of 7% acrylamide gel electrophoresis for randomly selected multiplex PCR primers on human ChrX. We confirmed that the primers magnified specific PCR products perfectly. Lanes b–q are the results of the amplification. Lanes l–q are the result of multiplex genomic PCR; each band in the lane was amplified by the primers corresponding to the lane \{d,e,h\}, \{g,j,k\}, \{b,c,i\}, \{d,e,g,h,k\}, \{b,c,f,i,j\}, \{b,c,d,e,f,g,h,i,j,k\}.

### Pre-calculation of qualified primer candidates

PrimerStation calculates multiplex genomic PCR primers for given query target sequences in the human genome. Since the calculation demands extremely costly computations, it cannot be executed online. To guarantee a prompt response on the web server, we preprocessed the human genome to enumerate

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**Figure 1.** The relationship between the hybridization ratio and temperature. The horizontal axis denotes the temperature and the vertical axis the hybridization ratio. The red line depicts the hybridization ratio of the primer to the target sequence. The black lines show the hybridization ratio of the primer to the off-target sequences. The primer and target sequences are TCGCCAGGAAGTAACTGGGAGCAG and the off-target sequences are (a) CAGCTCCCACTTACTCCAGCTG, (b) CTGCCGCACTTCTCCCTCCGTGGAG, (c) CTCGCTCCAGTTATTCTCTCTTGGTGG, and (d) TGGGCAAGTTACTGGAGGCAG. The blue region indicates the executable temperatures at which the hybridization ratio exceeds 0.99 to the target sequence and the hybridization ratios to the off-targets are no more than 0.05.
highly specific 25 base substrings in gene coding regions, such that their hybridization ratios to any off-target sequences are less than 0.05. The pre-calculation took 80 days using a massively parallel computer with 100 CPUs (SunFire 15 K) because off-target candidates were searched for as carefully as possible in the pre-calculation step. In particular, we examined off-target sequence candidates with at most four mismatches before the thermodynamic calculation using an efficient text-matching algorithm (22).

Seven percent of 25 base substrings of the gene coding regions were qualified. Although the figure 7% may appear to be small, in reality, these qualified 25 base substrings suffice to cover most of base pairs in the gene coding regions because 74% of base pairs in the gene coding regions are covered by at least one 60–600 base-long PCR product of qualified primers. The remaining 26% of the gene coding regions are difficult to amplify due to repetitive elements.

**PrimerStation web service**

Figure 3 shows a snapshot of PrimerStation. To specify specific positions in the human genome, PrimerStation accepts a list of RefSeq accession numbers of genes to design primers for multiplex genomic PCR. The users can also input configure options for the PCR product involving the product size range, the minimum product size differences among the set of designed primer pairs, the exclusion of primers with known single nucleotide polymorphisms (SNPs), the avoidance of (A)$_n$ and (CA)$_n$ repeats and the elimination of defective primers using the secondary structure prediction by Mfold (23). It also allows us to set PCR condition options, such as the cation concentration and primer concentration. We are able to fit these parameters to the real experiment by changing the values.

After inputting all of the parameters, the design of multiplex PCR primers is requested by pressing the submit button located below the gene ID textbox. PrimerStation attempts to output an optimal primer set for multiplex genomic PCR for the given gene set, or it reports the failure to find a set. The result includes primer sequences for individual genes, their product size, the second maximum hybridization ratio against off-targets, the melting temperatures of the forward and reverse primers, the minimum executable temperature and information about amplifying the target chromosome. The primer sequences can be downloaded as a fasta or csv formatted file.

**Figure 2.** Multiplex genomic PCR. Lanes b–q are the electrophoresis results of multiplex genomic PCR on the human ChrX. Ten primers were used, and the multiplicity of primers for each result is (b–k) 1, (l–n) 3, (o and p) 5 and (q) 10. Primers were mixed before PCR amplification. The bands in lanes a and r are ladder markers. Observe that the designed primers amplified a single target sequence from the human genome, and that primers b–k amplified highly specific bands.
MATERIALS AND METHODS

Hybridization ratio

Here, we describe the idea of the hybridization ratio in detail. The primer–template dissociation reaction is given by the following equation:

$$\frac{1}{2} \text{Primer} + \text{Template} \leftrightarrow \frac{1}{2} \text{Primer} + \text{Template}.$$ 

The left side of the equation means that the primer and template sequences are dissociated. The right side of the equation indicates that the primer and template sequences are hybridized to each other. In a real solution of primer and template sequences, equilibrium of the two states is established.

To calculate the hybridization ratio, it is essential to know the dissociation constant, a value expressing the extent to which a primer–template complex dissociates in solution:

$$K = e^{-\frac{\Delta G}{RT}} = \frac{f}{(C_p - fC_t)(1 - f)},$$

where $\Delta G$ is the free energy change, which can be calculated using the nearest neighbor method (24); $R$ is the molar gas constant; $T$ is the temperature; $C_{\text{pK}}$, $C_p$ and $C_t$ are the molar concentrations of the duplex of primer and template, the single strand of primer and the single strand of template, respectively; and $f$ denotes the hybridization ratio, which is the ratio of hybridized primer to the total amount of primer.

If $C_p$ is much larger than $C_t$, the hybridization ratio $f$ is approximated by

$$f \approx \frac{C_pK}{1 + C_pK} = \frac{C_p e^{-\frac{\Delta G}{RT}}}{1 + C_p e^{-\frac{\Delta G}{RT}}}.$$ 

The hybridization ratio $f$ is determined by the primer concentration $C_p$, the free energy change $\Delta G$ and the temperature $T$. The temperature at which the hybridization ratio becomes the specified value can also be calculated from $C_p$, $\Delta G$ and the hybridization ratio $f$. The temperature at which the hybridization ratio is 50% ($f = 0.5$) is the melting temperature.

The conditions for the multiplex PCR in Figure 2

The reaction consisted of 200 mM Tris–HCl (pH 8.4), 500 mM KCl, 100 mM MgCl$_2$, 2.5 mM dNTPs, 6.0 µl ddH$_2$O, 10 µM concentration of each primer, 36 ng of human genomic DNA and 2 U of Bio _Tag_ HS DNA polymerase (Takara Bio, Tokyo, Japan). The final reaction volume was 10 µl. The cycling conditions were 1 min at 94°C, 40 cycles of 30 s melting at 94°C, and 2 min of polymerization at 72°C, followed by a final 7 min extension at 72°C.

DISCUSSION

To evaluate the completeness of the design using PrimerStation, we attempted to process 28 516 human RefSeq genes
(November 2004, RefSeq gene set). We found that our primer sets were able to amplify 74% of base pairs in the gene coding regions. From the candidates of the primers, PrimerStation selects the best primer set from the candidates. The maximum hybridization ratio against off-targets was set to 0.05, although relaxing the threshold did not improve the coverage of genes for which primers were designed successfully because these genes have highly homologous sequences in their family genes.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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