Alternative splicing is widely found for eukaryotic genes, and it plays important roles in biological processes such as increasing protein diversity. The variant-specific PCR technique requires a unique region of a transcript variant for the primer design to distinguish between the specific variant from other isoforms. Existing programs, such as Primer3 (Rozen and Skaletsky, 2000), PerlPrime (Marshall, 2004) and Primer Premier (Singh et al., 1998), usually require users to enter or parse the DNA template sequence and to specify the unique region manually. Another program, PRIMEGENS-v2, added a new algorithm, HSPD (Srivastava et al., 2011), to design homolog-specific PCR primers without any prior information of the splice variants, but the algorithm requires much longer search times to generate homolog-specific primers. Importantly, none of these programs supports the graphic display of the splice variants of eukaryotic genes. However, the visualization of gene structure makes a more direct impression and provides more information on the spatial relationships to users than the currently used, sequence-based information.

In fact, with the rapid development of information technology (IT) tools, especially HTML5 and JavaScript, the user experience of PCR primer design can be largely improved. The configuration of complicated parameters and options can be avoided by some mouse clicks and simple dragging operations.

Therefore, we developed VizPrimer, a web server for visualized PCR primer design based on known gene structure. There are several features that distinguish VizPrimer from other traditional PCR primer design programs: (i) instead of pasting the template DNA sequence, VizPrimer accepts official gene symbols, NCBI GenBank accession numbers, official full gene names and ‘fuzzy’ words as key words to fetch the gene records from the server; (ii) instead of directly displaying the sequence on the screen, VizPrimer graphically displays the gene structure (for all of the splice variants) with the genome location, as other genome browsers do (Fig. 1), to help users find the unique region of a splice variant for PCR primer design; (iii) rather than asking users to configure or input the complicated parameters, VizPrimer provides an intuitive and easy-control platform to help users to specify the primer-picking region and other operations; (iv) designing PCR primers in VizPrimer is flexible and, depending on the primer design strategy and positions of primer icons, VizPrimer can design PCR primers for the amplification of specific exons and open reading frames (ORFs) and the identification of transcript variants and single nucleotide polymorphisms (SNPs) and (v) in combination with our previous work, MFEprimer (Qu et al., 2009), VizPrimer acts as a single package, which includes gene structure visualization, a primer design and a specificity-checking service to help users design high-quality PCR primers.

1 INTRODUCTION

Alternative splicing is widely found for eukaryotic genes, and it plays important roles in biological processes such as increasing protein diversity. The variant-specific PCR technique requires a unique region of a transcript variant for the primer design to distinguish between the specific variant from other isoforms. Existing programs, such as Primer3 (Rozen and Skaltsky, 2000), PerlPrime (Marshall, 2004) and Primer Premier (Singh et al., 1998), usually require users to enter or parse the DNA template sequence and to specify the unique region manually. Another program, PRIMEGENS-v2, added a new algorithm, HSPD (Srivastava et al., 2011), to design homolog-specific PCR primers without any prior information of the splice variants, but the algorithm requires much longer search times to generate homolog-specific primers. Importantly, none of these programs supports the graphic display of the splice variants of eukaryotic genes. However, the visualization of gene structure makes a more direct impression and provides more information on the spatial relationships to users than the currently used, sequence-based information.

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2 IMPLEMENTATION

There are five modules in VizPrimer: (i) user-input query module; (ii) gene structure display module; (iii) user-interactive module; (iv) primer design module and (v) primer specificity check module.

As mentioned above, the user-input query module in VizPrimer accepts the official gene symbols, NCBI GenBank accession numbers, official full gene names and ‘fuzzy’ words as key words to fetch gene records from the server. In addition, VizPrimer allows
VizPrimer: visualized primer design server

Fig. 1. Visualization of the mouse neuronatin (Nnat) gene structure using VizPrimer. The right and left arrowheads represent the forward and reverse PCR primers, respectively.

Fig. 1

the users to upload their own cDNA and genomic DNA sequence for visualizing the gene structure. GMAP (Wu and Watanabe, 2005) was used as a backend program for the gene structure prediction.

There are two layers of gene structure that are displayed. First, a global view of the gene structure is presented, including all of the splice variants; the resolution is dynamic and can be easily controlled. Second, the detailed gene structure can be shown at a resolution of even one base pair. The Genome Mapping data (ftp://ftp.ncbi.nih.gov/genomes/MapView) was used to display the splice variant images at the gene level. The JavaScript and HTML5 'canvas' element were implemented in the gene structure visualization. A JavaScript-based graphics demonstration has been widely used in bioinformatic and cheminformatic visualizations due to its low cost on the server-side and free plug-in feature on the browser-side.

Unlike the currently and widely used programs, the PCR primer design work in VizPrimer is similar to playing video games. Without the need to calculate the position of an exon, the user merely puts/draggs the primer icons (right and left arrowheads) on the selected exon, and the amplicon size and primer position will be simultaneously displayed when moving the forward/reverse primer icons (Fig. 1). All these operations are controlled by JavaScript on the browser-side, which is fast and does not need to communicate with the server.

There are four options available in VizPrimer to select the optimal PCR primers for special demands: Exon locked′ (for splice variant analysis), 5′-terminus free′ (for gene expression analysis), 5′-terminus locked′ (for ORF analysis) and 3′-terminus locked′ (for SNP analysis). The Exon locked′ option indicates that VizPrimer will search the optimal primers from the full region of the selected exon, which is useful for the identification of the expression of the splice variant containing this exon. The 3′-terminus free′ option means that VizPrimer will search the optimal primers globally from the full-length cDNA sequence. The 5′-terminus locked′ option means that the 5′-terminus of the primer is locked on the position indicated by the primer icons, as denoted by the right and left arrowheads; this option is suitable for ORF primer design to amplify the region from the start codon (ATG) to or before the stop codon (TAA/ TAG/TGA). Of course, the users can also specify their own cDNA region to be amplified by using this option. In addition, the 3′-terminus locked′ option will be useful for SNP PCR primer design. The kernel primer design program is a Primer3 (version 2.2.3) command-line program. The result page will return five PCR primer pairs by default and the graphic gene structure with the designed primers located on it. Moreover, a link to the MFEP primer-2.0 (http://biocompute.bmi.ac.cn/CZlab/MFEPprimer-2.0) specificity evaluation results is provided for each primer pair.

The overall computing time for both the PCR primer design and primer specificity evaluation were ~1 s on average, without considering the network communication time. VizPrimer requires browsers supporting HTML5, such as Chrome (≥5.0) and Firefox (≥3.0).

3 DISCUSSION

The design of PCR primers for eukaryotic genes in the identification of splice variants is a complex and tedious task. The currently available primer design programs do not have a function of graphically displaying the gene structure to help users locate the unique regions among the splice variants. Using HTML5 and JavaScript for adding interactivity to a web page is a new paradigm of web computing nowadays. This combination is also encouraged by scientific applications due to the fast progress in the performance of JavaScript engines and HTML5-compatible browsers. There are already several scientific applications relying on the Java and HTML5 'canvas' element and which are currently available, including Anno-J (Lister et al., 2008) and ChemDoodle (http://web.chemdoodle.com/). Therefore, by integrating these new technologies, the widely used primer design program, Primer3, and the specificity-checking program, MFEP primer, VizPrimer will be a beneficial visual PCR primer design tool. In the future, we will add the function of visualized multiple PCR primer design to VizPrimer.

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