PrimerProspector: de novo design and taxonomic analysis of barcoded PCR primers

William A. Walters¹,6, J. Gregory Caporaso²,6, Christian L. Lauber³, Donna Berg-Lyons³, Noah Fierer³,4, and Rob Knight²,5,*

¹ Department of Molecular, Cellular, and Developmental Biology, University of Colorado at Boulder, Boulder, CO, USA.
² Department of Chemistry and Biochemistry, University of Colorado at Boulder, Boulder, CO, USA.
³ Cooperative Institute for Research in Environmental Sciences, University of Colorado at Boulder, Boulder, CO, USA.
⁴ Department of Ecology and Evolutionary Biology, University of Colorado at Boulder, Boulder, CO, USA.
⁵ Howard Hughes Medical Institute, Boulder, CO, USA.
⁶ These authors contributed equally to this work.

ABSTRACT

Motivation: PCR amplification of DNA is a key preliminary step in many applications of high-throughput sequencing technologies, yet design of novel barcoded primers and taxonomic analysis of novel or existing primers remains a challenging task.

Results: PrimerProspector is an open source software package that allows researchers to develop new primers from collections of sequences, and to evaluate existing primers in the context of taxonomic data.

Availability: PrimerProspector is open source software available at http://pprospector.sourceforge.net

Contact: rob.knight@colorado.edu

1 INTRODUCTION

Using next-generation sequencing methods to characterize hundreds of samples simultaneously in a single sequencing run has revolutionized microbial ecology (Tringe and Hugenholtz, 2008). However, primer design for such studies remains challenging. The primers must amplify an appropriate region of DNA that is the right length for sequencing and also taxonomically informative (Liu, et al., 2008; Wang, et al., 2007); a linker that is not complementary to the target in any one of many diverse species must be inserted before the barcode to avoid differential amplification (Hamady, et al., 2008); and the set of barcodes must be checked to avoid formation of secondary structure within or between primers (i.e., primer-dimers) or between the barcodes and the primers. Additionally, the techniques need to be generic rather than tied to one taxonomic outline or database, so that many different target genes can be studied.

Here we present PrimerProspector, an open source software package for primer design and analysis built using the PyCogent toolkit (Knight, et al., 2007), that resolves these issues. We recently applied PrimerProspector to identify the 16S rRNA 515f/806r primer pair as nearly universal to Archaea and Bacteria, and to optimize this primer pair for increased sensitivity across these domains. This optimized primer pair, applied successfully in several recent studies (Bates, et al., 2010; Bergmann, et al., 2010; Caporaso, et al., 2010), has provided novel insight into archaeal and bacterial community membership in soils by allowing for more accurate determination of the abundances of taxa missed by many commonly-used canonical primer pairs, e.g. the Verrucomicrobia.

No existing tools specifically address the issues associated with designing barcoded PCR primers for community analysis. Primer design is a large field and we cannot survey it comprehensively in this article, but among a selection of related tools, PrimerValidator (http://bioinfo.unice.fr/454) allows taxonomic assessment but does not generate de novo primers, or allow a customizable 3' weighted scoring system to predict successful amplification of tested primers. BarCrawl (Frank, 2009) allows design of barcodes for specified PCR primers but not design of the primers themselves, so is a useful complement to PrimerProspector. RDP’s Probe Match (Cole, et al., 2005) will report sequences matching a probe, as does Greengenes’ probe function (DeSantis, et al., 2006), but these tools are tied to the respective 16S rRNA databases and do not have support for barcodes. Primrose and OligoCheck (Ashelford, et al., 2002) are useful for small numbers of target sequences, but do not scale well to thousands or tens of thousands of sequences, as is necessary when designing universal or near-universal primers, and do not incorporate differential weighting of 5’ and 3’ bases in primer scoring. Primer BLAST uses Primer3 software to build primers of a specified length against one target sequence, then BLASTs the results against other databases to ensure that putative primers do not target BLAST hits. This function-
ality is also a useful complement to that provided in PrimerProspector.

While applications of PrimerProspector to date have focused on SSU rRNA primer design, PrimerProspector can be used for any nucleic acid sequences and allows users to design de novo primers based upon arbitrary multiple sequence alignments. User-specified design parameters include primer length, degeneracy, and targeted regions for generation of primers. Existing or de novo primers can be analyzed for predicted taxonomic coverage, as shown in Figure 1. Finally, common pitfalls in primer design can be identified, such as likely barcode-primer secondary structure, regions susceptible to primer dimerization, and disparate GC content between primer pairs. Convenient reports show amplicons or simulated reads that cover regions of sequences that are not phylogenetically informative or are of unsuitable lengths for sequencing.

2 METHODS

De novo design of primers is performed by finding short conserved sequences in a given multiple sequence alignment to act as a 3’ binding site for new primers. Once these sites have been identified, full-length forward or reverse de novo primers are generated by incorporating the N upstream or downstream bases, where N is 15 by default. De novo full-length primers can then be sorted according to sensitivity, specificity, or degeneracy, and compared to known primers to find matches or significant overlap. Specificity for particular target groups, such as Archaea, can be obtained by supplying an optional alignment of sequences from which to exclude matches. Primer analyses, including the prediction of taxonomic coverage, relies upon scoring primers against target sequences. To predict its taxonomic coverage, a primer is locally aligned to full length target sequences with known taxonomies, and scored based on gap, 3’ mismatch, and non-3’ mismatch counts. An example of
the graphical output is provided in Figure S3. The final five bases are considered to be the 3' region by default, and are considered to be the most important for PCR amplification. The scoring scheme is parameterizable. The RDP Classifier (Wang, et al., 2007) is used to classify the resulting sequence fragments, and the accuracy is displayed both in terms of which taxa are amplified and in terms of classification level of the resulting fragments. PrimerProspector supports retraining of the RDP Classifier for taxa coverage analysis based on different reference taxonomies.

Descriptions of the scripts included in PrimerProspector, the various outputs generated by PrimerProspector, and an example based on the F515/R806 primer pair, are included in the online documentation at http://pprospector.sourceforge.net/.

3 CONCLUSIONS

PCR amplification continues to be a key step in many high-throughput sequencing applications such as barcoded marker-gene-based microbial community analyses. PrimerProspector represents a significant advance over prior work in this area by providing a single tool to facilitate primer design and analysis, including support for barcodes (and associated linkers). PrimerProspector is a fast and extensible framework for primer design and analysis, and has already been successfully applied to help researchers identify the most relevant and useful primers for their application, starting with multiple sequence alignments for any nucleic acid sequence.

ACKNOWLEDGEMENTS

Funding: This work was funded in part by the Bill and Melinda Gates foundation, the CCFA, HHMI, and the NIH SCR Training Grant (T32GM008759).

REFERENCES

Ashelford, K.E., Weightman, A.J. and Fry, J.C. (2002) PRIMROSE: a computer program for generating and estimating the phylogenetic range of 16S rRNA oligonucleotide probes and primers in conjunction with the RDP-II database, Nucleic Acids Res, 30, 3481-3489.

Bates, S.T., et al. (2010) Examining the global distribution of dominant archaeal populations in soil, ISME J.

Bergmann, G., Bates, S., Eilers, K., Lauber, C., Caporaso, J., Walters, W., Knight, R., and Fierer, N. (2010) The under-recognized dominance of Verrucomicrobia in soil bacterial communities. Under review.

Caporaso, J.G., et al. (2010) Global patterns of 16S rRNA diversity at a depth of millions of sequences per sample, Proc Natl Acad Sci U S A.

Cole, J.R., et al. (2005) The Ribosomal Database Project (RDP-II): sequences and tools for high-throughput rRNA analysis, Nucleic Acids Res, 33, D294-296.

DeSantis, T.Z., et al. (2006) Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB, Appl Environ Microbiol, 72, 5069-5072.

Edgar, R.C. (2010) Search and clustering orders of magnitude faster than BLAST, Bioinformatics, 26, 2460-2461.

Frank, D.N. (2009) BARCRAWL and BARTAB: software tools for the design and implementation of barcoded primers for highly multiplexed DNA sequencing, BMC Bioinformatics, 10, 362.

Hamady, M., et al. (2008) Error-correcting barcoded primers for pyrosequencing hundreds of samples in multiplex, Nat Methods, 5, 235-237.

Knight, R., et al. (2007) PyCognent: a toolkit for making sense from sequence, Genome Biol, 8, R171.

Liu, Z., et al. (2008) Accurate taxonomy assignments from 16S rRNA sequences produced by highly parallel pyrosequencers, Nucleic Acids Res, 36, e120.

Tringe, S.G. and Hugenholtz, P. (2008) A renaissance for the pioneering 16S rRNA gene, Curr Opin Microbiol, 11, 442-446.

Wang, Q., et al. (2007) Naive Bayesian classifier for rapid assignment of rRNA sequences into the new bacterial taxonomy, Appl Environ Microbiol, 73, 5261-5267.