Four PCR primers necessary for the detection of periplasmic nitrate reductase genes in all groups of Proteobacteria and in environmental DNA

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Abstract

Generic primers are available for detecting bacterial genes required for almost every reaction of the biological nitrogen cycle, the one notable exception being napA (gene for the molybdoprotein of the periplasmic nitrate reductase) encoding periplasmic nitrate reductases. Using an iterative approach, we report the first successful design of three forward oligonucleotide primers and one reverse primer that, in three separate PCRs, can amplify napA DNA from all five groups of Proteobacteria. All 140 napA sequences currently listed in the NCBI (National Center for Biotechnology Information) database are predicted to be amplified by one or more of these primer pairs. We demonstrate that two pairs of these primers also amplify PCR products of the predicted sizes from DNA isolated from human faeces, confirming their ability to direct the amplification of napA fragments from mixed populations. Analysis of the resulting amplicons by high-throughput sequencing will enable a good estimate to be made of both the range and relative abundance of nitrate-reducing bacteria in any community, subject only to any unavoidable bias inherent in a PCR approach to molecular characterization of a highly diverse target.

The missing link in our ability to detect nitrogen cycle genes in community DNA

The availability of oxygen, phosphate, sulfate and reduced carbon compounds are major determinants of the balance between beneficial and detrimental chemical changes in soils, sediments and water. However, the productivity of many ecosystems, especially soils, is limited by biologically available nitrogen. Micro-organisms, especially bacteria, are critical in determining the balance between nitrification, denitrification, nitrate assimilation, anammox, nitrogen fixation and the reduction of nitrate to ammonia. The range of species involved in these processes is extremely diverse. Many groups of bacteria synthesize multiple enzymes for each reaction. Nevertheless, considerable progress has been achieved in developing oligonucleotide PCR primers to detect, amplify and quantify the capacity of any given community to catalyse each process. A notable exception to this generalization is the current inability to detect the complete range of genes encoding periplasmic nitrate reductases from Gram-negative bacteria.

Three types of bacterial nitrate reductase have generally been recognized, though detailed analysis of their subcellular location and associated electron transfer components justifies more detailed subdivisions. Assimilatory nitrate reductases located in the cytoplasm catalyse the first step in nitrate reduction to ammonia, which is then used for amino acid biosynthesis. During anaerobic growth, respiratory nitrate reductases typified by the NarGHI (membrane-associated, energy-conserving nitrate reductase) complexes from Escherichia coli and Paracoccus denitrificans are associated with the cytoplasmic membrane where they conserve energy released during nitrate reduction as proton motive force [1,2]. Although the active site of most NarG proteins is located in the cytoplasm, exceptions to this have recently been recognized, creating a further subdivision of nitrate reductase, typified by Nar from various Archaea (reviewed in [3]). The third group, the periplasmic nitrate reductases, are biochemically and genetically distinct from the other groups. They fulfil diverse roles: the maintenance of redox balance during growth on reduced carbon sources [4–10]; maintenance of redox balance during photosynthetic growth [11]; scavenging low concentrations of nitrate [12]; or nitrate respiration in the first stage of denitrification or nitrate reduction to ammonia [5,13,14].

Due to their diverse roles, there is much greater sequence variation in napA genes encoding the periplasmic nitrate reductases than those of the assimilatory and membrane-associated nitrate reductases. Consequently, oligonucleotide primers designed for detecting napA from some groups of bacteria are unsuitable for detecting napA in other groups. Based on very limited sequence data available at the time, primers were designed that successfully amplified napA sequences from DNA isolated from soils, but the range of bacteria detected was limited [15]. Conversely, primers that amplified napA from the δ-proteobacterium Desulfovibrio desulfuricans strain 27774 would not recognize napA from E. coli or P. denitrificans [16]. The Phillipot group subsequently redesigned primers suitable for detecting a...
A standardized PCR protocol to detect periplasmic nitrate gene products

The PCR protocol was developed using as templates chromosomal DNA isolated from E. coli K-12, D. desulfuricans 27774, Campylobacter jejuni, Haemophilus influenzae, Habella chejuensis, Wolinella succinogenes, Campylobacter lari, Campylobacter hominis, Campylobacter curvus, Campylobacter concisus, Bortetella bronchiseptica, Bradyrhizobium japonicum and Paracoccus pantotrophus. DNA sequences of species that were known to contain the napA gene were downloaded from the NCBI (National Center for Biotechnology Information) database. They were aligned with the ClustalX tool and displayed by the BoxShade tool in order to detect conserved regions. Degenerate primers were designed based on these alignments and produced by Alta Biosciences.

After many preliminary experiments that included trials with hot start, touchdown and gradient PCR, a standardized touchdown protocol was developed that included a 5 min denaturation step at 95°C followed by 32 touchdown cycles in which the annealing temperature was decreased by 0.5°C per cycle from 55°C down to 39°C. Samples were held at the annealing temperature for 1 min. In each subsequent cycle, the sample was denatured for 30 s at 95°C and extended for 150 s at 72°C. The amount of product was amplified in 30 steps in which the denaturation and extension steps were exactly as in the touchdown cycles, but primers were annealed for 1 min at 39°C. After a final 10 min extension step at 72°C the sample was stored at 4°C. Products were checked by electrophoresis in a 100 ml 1% agarose gel.

Limited applicability of previously described primers

In preliminary experiments, various primers described in the literature or developed in-house were tested for their ability to direct the synthesis of a limited range of napA target genes. The design of primers V16 and V17 was based on the known sequences of napA genes from a limited number of bacteria that included E. coli and P. denitrificans [15] (Figure 1). The ability of these primers to recognize these templates, but not napA from D. desulfuricans strain 27774, was confirmed. Similar results were obtained with related primers described by Bru et al. [17]. Conversely, primers DdV16 and DdV67, which were used in the determination of the sequence of D. desulfuricans nap operon [16], were unable to recognize E. coli or P. denitrificans templates.

Design of primers that recognize both D. desulfuricans and E. coli napA templates

We then attempted to design universal primers that would amplify napA DNA from all groups of proteobacteria by comparing the sequences of highly divergent napA genes from 3α-, 3β-, 6γ-, 25- and 46-proteobacteria (Figure 1). Although very few regions of sequence identity were found, two primers resulting from this analysis, 1173b and R2, were predicted to amplify napA fragments from both E. coli and D. desulfuricans. This prediction was confirmed. However, none of the primer combinations available recognized napA genes in C. jejuni or C. lari.

Design of primers that recognize napA templates from all groups of proteobacteria

Further analysis revealed that this limited range of sequences could be split into two subgroups within which were more regions of sequence similarity. Two pairs of degenerate primers based on these regions of similarity were designed. Primers SF1173d and SR2294 successfully amplified the expected napA fragments of both E. coli and D. desulfuricans DNA, but failed to recognize C. jejuni DNA (Figures 1 and 2). Primers LF716 and LR1837 amplified napA fragments from both E. coli and C. jejuni DNA, but not from D. desulfuricans DNA. When primer SF1173d was combined with LR1837, the expected 0.5 kb fragments were generated, but no product was obtained with D. desulfuricans DNA as template (Figure 2). The primer pair LF716 and SR2294 generated the expected 1.5 and 1.8 kb products with E. coli and C. jejuni DNA templates (Figure 2). However, not every expected product was generated, and some templates directed the synthesis of more than one product.

The sequence analysis was then extended to all 140 napA sequences from 70 bacterial species in the NCBI database in February 2010. Each of the four primers was checked against the database sequences to determine whether there was a mismatch in the last six bases at the 3′-end of each primer sequence that might prevent effective amplification of the required fragment (Table 1). This analysis revealed only three sequences, those from Geobacter lovleyi, C. lari and C. hominis, that failed to meet the stringent criteria set with the forward primers SF1173d and LF716. One of the reverse primers, SR2294, was predicted to recognize every napA sequence in the database, and was therefore potentially truly generic. Furthermore, this primer allowed problems predicted when using primer LR1837 to be avoided (Table 1).

In a final attempt to design a universal forward primer, alignments were first generated for all five subdivisions of proteobacteria, with a further alignment of sequences that posed the greatest problems. This resulted in the design of...
Figure 1 | Oligonucleotide primers used to detect napA genes in all groups of proteobacteria

The top five segments of the Figure show the sequences of five of the primers used to develop the generic protocol for the detection of napA in various groups of proteobacteria above the typical sequences they were designed to recognize. The lower diagrams show where within the napA genes of *Escherichia coli*, *D. desulfuricans* and *C. jejuni* the various primers mentioned in the text are predicted to bind.

Primer OF640: GSNKHAAYATGSC
  C. hominis 620 T7ACCTGGGTGGTCAATGCTGAAATG 648
  G. lovleyi 608 TCTCTGGGGGCGCAATGCGAGGCC 636

Primer LF716: GCGNARATGCACC
  *D. desulfuricans* 626 CAGCGAGGCCCAAGGCTCTTTTCGC 654
  *C. jejuni* 638 TGGCTAATGCGCGCAATCCTTGGTC 666

Primer SF1173d: GTATGGGNNMRAACC
  *E. coli* 1105 TACATGGACATGGCTACACGACATAC 1133
  *D. desulfuricans* 1069 TTCTGGGTATCAGCAGACCGAAGAGT 1097

Primer LR1837: TCCWCTRADGCWCC
  *F. marina* 1773 ATGGGAGCTTATGGTACGTGAAAGGA 1801
  *C. jejuni* 1725 ATGGGAGCTTATGGTACGTGAAAGGA 1753

Primer SR2294: GWRGGCGARTGNC
  *C. jejuni* 2448 GTTTTTTAGAGGACATGGCATAAGACTA 2476
  *D. desulfuricans* 1947 GGTATCGACATCGGATACGGCAACCA 1975

Table 1 | Overview of napA sequences that might not be amplified using each of the four primers

<table>
<thead>
<tr>
<th>Class/primer</th>
<th>LF716</th>
<th>LR1837</th>
<th>SF1173d</th>
<th>SR2294</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>α-Proteobacteria</strong> (eight sequences)</td>
<td>B. japonicum, Rhodobacterales bacterium, <em>Sinorhizobium meliloti</em>, <em>Agrobacterium tumefaciens</em>, <em>Rhodospirillum centrum</em>, Pseudovibrio sp.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>β-Proteobacteria</strong> (11 sequences)</td>
<td>Candidatus, ‘Accumulibacter phosphatis’ clade, Burkholderia xenovarans</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>δ-Proteobacteria</strong> (four sequences)</td>
<td>G. lovleyi, <em>D. desulfuricans</em></td>
<td></td>
<td>G. lovleyi</td>
<td></td>
</tr>
</tbody>
</table>

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primer OF640 that was predicted to recognize templates such as \textit{C. lari}, \textit{C. hominis} and \textit{G. lovleyi} (Figure 2B). This forward primer in combination with the universal reverse primer SR2294 directed the synthesis of the predicted 1.8 kb fragment on the two \textit{Campylobacter} templates (\textit{G. lovleyi} DNA was unavailable). Control experiments showed that the LF716 forward primer with either SR2294 or LR1837 also generated the expected fragments on these templates, but multiple products were obtained even with pure DNA templates. For this reason, use of the third forward primer, OF640, is recommended for the detection, amplification and quantification of difficult target \textit{napA} genes from \textit{\epsilon}-proteobacteria.

In summary, all known \textit{napA} genes can now be amplified using one of three forward primers, LF716, SF1173d or OF640, in combination with the single reverse primer, SR2294. Although they are all degenerate primers that also recognize sequences unrelated to \textit{napA} on some templates, the limited degeneracy restricts this problem to an acceptable degree. However, due to the generation of multiple products in some reactions, the three forward primers should not be used together in a single multiplex PCR.

**Amplification of napA sequences using community DNA as a template**

We recently described the use of community DNA from the faeces of patients suffering from the gastric diseases ulcerative colitis, irritable bowel syndrome and Crohn’s disease to compare the abundance of \textit{Faecalibacterium prausnitzii} before and after successful treatment with those of a group of healthy controls [18]. Sulfate-reducing bacteria were abundant in all of these samples, as were enteric bacteria. Virtually all of the samples yielded PCR products of the expected size range with two of the three pairs of primers (the primer pair OF640+SR2294 was not tested in these experiments). However, as expected when degenerate primers...
are used with community DNA as a template, multiple PCR products were obtained from most samples. Nevertheless, as discussed below, these PCRs can now be combined with high-throughput DNA sequencing that is increasingly widely available to determine the diversity of napA sequences in any bacterial community.

High-throughput DNA sequencing avoids a requirement for highly specific primers

Before the availability of high-throughput DNA sequencing, analysis of the variety of species capable of catalysing major environmental changes required the cloning and sequencing of individual PCR products one at a time, a process that was both time consuming and extremely expensive. Consequently, it was essential to design oligonucleotide primers that were not only sufficiently degenerate to capture all of the relevant target genes in community DNA but also sufficiently specific to avoid the generation of PCR products irrelevant to the investigation. The ability to generate up to 10^6 DNA sequences in a single experiment, for example using Roche 454 DNA sequencing technology, renders the second problem largely irrelevant because species that represent less than 0.1% of the population can be detected even in a mixture of relevant and irrelevant amplicons [19]. Primer design can now focus on ensuring the amplification of the widest range of target sequences. This has enabled us to resolve a major outstanding problem in determining the range of nitrate reducing bacteria in complex communities by designing for the first time a set of primers that have been shown to recognize all of the napA genes currently known to occur. For each sample to be analysed, three PCRs involving different forward primers but the same reverse primer are essential. Note, however, that different bar codes could be added to each of the three forward primers and the resulting PCR products could then be mixed to decrease the costs associated with sequencing products from multiple samples. In another study, we have validated this approach by determining the range of sulfate-reducing bacteria represented in the same faecal DNA samples used in the work described in the present paper (W. Jia and J.A. Cole, unpublished work). By using 15 different barcoded sequencing primers and splitting the Roche 454 sequencing plate into four segments using gaskets, up to 60 samples can be analysed in a single DNA sequencing experiment. Typically, this will generate 150 000 useful sequences per segment, or on average 10 000 sequences per barcoded product.

Even with the more relaxed objectives of the work described in the present paper, the design of universal primers to detect the wide variety of periplasmic nitrate reductase genes was a challenging task, solved by an iterative approach. The three forward primers LF716, SF1173d and OF640, in combination with the reverse primer SR2294, have either been shown or are predicted to recognize all currently known napA sequences not only from DNA from single strains of the complete range of Proteobacteria but also from DNA isolated from the faeces of patients with three different gastric diseases. Amplicons of the correct size were also obtained with primer pairs in which there was at least one mismatch in the last six bases, suggesting that the current primers might also recognize targets for which they were not designed. If necessary, further refinements can be made to these primers as additional napA sequences resulting from metagenomic studies are added to the databases.

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References


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