Sample richness and genetic diversity as drivers of chimera formation in nSSU metagenetic analyses

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ABSTRACT

Eukaryotic diversity in environmental samples is often assessed via PCR-based amplification of nSSU genes. However, estimates of diversity derived from pyrosequencing environmental data sets are often inflated, mainly because of the formation of chimeric sequences during PCR amplification. Chimeras are hybrid products composed of distinct parental sequences that can lead to the misinterpretation of diversity estimates. We have analyzed the effect of sample richness, evenness and phylogenetic diversity on the formation of chimeras using a nSSU data set derived from 454 Roche pyrosequencing of replicated, large control pools of closely and distantly related nematode mock communities, of known intragenomic identity and richness. To further investigate how chimeric molecules are formed, the nSSU gene secondary structure was analyzed in several individuals. For the first time in eukaryotes, chimera formation proved to be higher in both richer and more genetically diverse samples, thus providing a novel perspective of chimera formation in pyrosequenced environmental data sets. Findings contribute to a better understanding of the nature and mechanisms involved in chimera formation during PCR amplification of environmentally derived DNA. Moreover, given the similarities between biodiversity analyses using amplicon sequencing and those used to assess genomic variation, our findings have potential broad application for identifying genetic variation in homologous loci or multigene families in general.

INTRODUCTION

Second-generation pyrosequencing of environmental DNA has provided unique insights into prokaryotic (1,2) and eukaryotic (3,4) molecular diversity and ecology. Massive parallel pyrosequencing has the potential to produce a large volume of data relatively cheaply and with an unprecedented read depth, generating millions of DNA sequences within a matter of hours (5). Despite advantages of high throughput sequencing, a major challenge is to determine the extent to which sequences produced from pyrosequencing-amplified regions of marker genes correspond to biological diversity. Recently, studies have recognized that biodiversity levels have become inflated due to artifacts associated with sample processing including both the PCR amplification and the pyrosequencing itself (6–8). PCR amplification with universal primers applied to genes conserved across phyla, such as the ribosomal nuclear small subunit (nSSU), is commonly used to identify microbial eukaryotes in natural environments. The extreme conservation of primer binding sites (9) and the availability of extensive database resources (10) has resulted in the nSSU being the most widely used marker for studying the molecular taxonomy of a diverse range of eukaryotes. Target taxa range from all protist kingdoms (11) to metazoan microorganisms (4), that are dominated by the Nematoda (12). In such analyses, one of the most commonly reported sources of sequence artifacts associated with highly homologous nSSU genes from environmental DNA samples is the formation of chimeric sequences during PCR amplification (8,13–16).

Chimeric sequences, or chimeras, are generated when incomplete extension occurs during PCR amplification and the resulting amplicon re-anneals to a foreign DNA strand and is copied to completion in the following PCR cycles. Chimeras are composed of two or more
phylogenetically distinct parental sequences and have been shown to occur in PCR-amplified nSSU data sets with frequencies of 30–70% (6,17,18) thus leading to false diversity estimates and false novel taxa. The critical factors that seem to affect PCR-generated recombination are the number of PCR cycles, PCR extension time, template concentration, Taq DNA polymerases and amplicon size (18–21). Chimera formation can be minimized experimentally by PCR optimization, nonetheless, no method has yet proved to be entirely effective. The importance of detecting chimeras is such that a plethora of bioinformatic software has also been developed, such as Chimera_Check (22), Bellerophon (13), CCode (23), Pintail (24), Mallard (17), Chimera Slayer (6) and Perseus (8). With the exception of Perseus, such approaches will only detect evident induced chimeras (25) and their accuracy for chimera detection has not been rigorously tested (6) or is still at an early stage, especially given recent advances in environmental DNA sequencing approaches. Although metagenetic (4,9) analyses are clearly based on complex and phylogenetically diverse assemblages, the roles of sample richness and phylogenetic diversity in driving chimera formation are largely unknown.

Wang and Wang (18,26) tested how sequence similarity between cloned 16S rRNA genes or mixed bacteria genomic DNA can influence PCR-based chimera formation. Nonetheless, these investigations were performed on a very small scale, did not consider sample richness and pre-dated the current second-generation sequencing perspective of amplicon pool diversity. The overarching aim here is to (i) analyze the effect of richness, evenness and genetic diversity on chimera formation and link this to diversity estimates and (ii) understand how chimeras are formed with respect to variable genetic diversity and secondary structure of the parent nSSU molecule. To this end, a nSSU metagenetic data set was generated by 454 Roche pyrosequencing of control pools of closely and distantly related nematode mock communities of known identity and richness.

MATERIAL AND METHODS

Sample preparation
To test if chimera formation during PCR reactions was associated with taxon richness or with phylogenetic distance, 74 Sanger-sequenced single nematode species were blast aligned to a contemporary Nematoda phylogenetic framework (27). Subsequently, the sequences were aligned using ClustalX and pairwise distances (p-distance) were calculated using MEGA 4.1 (28). Based on the phylogenetic affinities of the nematode sequences, subsets of closely related [mean pairwise divergence (MPD) of 25%, referred to as ‘phylogenetically close’] and distantly related (MPD of 40%, referred to as ‘phylogenetically distant’) nSSU controls were generated by pooling the DNA extracts of 12, 24 or 48 individuals.

DNA extraction and preparation
DNA extraction from DESS-preserved (29) single worms was performed using a DNeasy Blood & Tissue Kit (Qiagen Inc), following the manufacturer’s instructions. After extraction, all DNA was eluted in 40 μl of AE buffer and samples were stored at −20°C until use. The DNA extracts from all single individuals were quantified using a Nanodrop spectrophotometer and diluted to 0.5 ng/μl, and five replicates of the 12, 24 and 48 individuals were selected for the closely and distantly related treatments.

PCR amplification and sequencing analysis
The primers SSUFO4 forward (5’-GCTTGTCTCAAAAG ATTAAGCC-3’) and SSUR22 reverse (5’-GCCTGCTGC CTTCCTTGGA-3’) were used to amplify ~450 bp of the nSSU rDNA (18S rDNA) region (30). Fusion primers were then developed according to Fonseca et al. (4). PCR amplification reactions and the thermocycle for the targeted nSSU region were optimized. Optimized reactions were performed using 0.25 ng/μl of genomic DNA template in 3 x 40 μl reactions using Pfu DNA polymerase (Promega) for each of the closely and distantly related nematode pools (12, 24 and 48 individuals) and all individual DNA extracts. PCR thermocycle conditions consisted of a 2-min denaturation step at 95°C followed by 35 cycles (thus facilitating the generation of chimeras) (6,18,26) of 1 min at 95°C, 45 s at 55°C, 3 min at 72°C and a final extension of 10 min at 72°C. Negative controls (ultrapure water only) were included for all amplification reactions. Electrophoresis of triplicate PCR products was undertaken on a 2% gel with Top Vision™ LM GQ Agarose (Fermentas), and the expected 450-bp fragment was purified using the QIAquick Gel Extraction Kit (Qiagen), following the manufacturer’s instruction. All purified PCR products were quantified with an Agilent Bioanalyser 2100 and diluted to the same concentration (10 ng/μl). PCR amplifications from single nematodes and pooled nematodes were sequenced in a single direction (A-Amplicon) on a quarter and three-quarters of a plate, respectively, using a 454 Roche GSFLX (454 Life Sciences, Roche Applied Science) sequencing platform at Liverpool University’s Centre for Genomic Research, UK.

Denoised reads and detection of chimeric PCR molecules
Pyrosequencing reads derived from 454 Roche data contain a substantial number of errors (referred to as noise), which includes sequencing errors mainly derived from the inclusion or deletion of single bases in homopolymer runs of 3 bp or longer, PCR single base substitutions and PCR chimeras (14,31). AmpliconNoise was used to remove noise from the pyrosequencing data; this comprises filtering, flowgram and sequence clustering steps. It has been shown to reduce noise by ~50% in environmental data sets (8). Subsequently, chimeras were identified using Perseus (8); this algorithm generates a Chimera Index (CI) for each read that is ≥0 with higher values corresponding to reads that are most likely to be chimeric. Perseus by pairwise alignments to all sequenced of greater than or equal abundance identifies the most likely parent sequences of the candidate read and the most likely break point. Logistic regression is then used.
to classify chimeras so that the pyrosequencing data output lists chimeric and non-chimeric sequences. The lower the probability of the sequence evolving naturally, the higher the CI (8).

Perseus finds break point positions in the two parent sequences. To do this, a four-way alignment between each chimeric sequence, its two parents and the Caenorhabditis elegans reference sequence was formed (GenBank/EMBL accession number EU196001). The most likely break point was identified by minimizing the number of differences between the sections of the parents contributing to the chimera and the chimera itself. The position of each break point on the reference sequence was then recorded and from this, the frequency breaks occurring at each position could be calculated. MFold RNA-folding software was used to predict the potential role on chimera formation of the secondary structure of the 18S rDNA amplicon region (32).

Generation of operational taxonomic units

Denoised mock nematode community data from which chimeras had been removed was used to identify Operational Taxonomic Units (OTUs). OTUs were calculated using a complete linkage-clustering algorithm, measuring the distance between the most distant members in each cluster, at a 99% identity cut-off. The number of OTUs generated was then used to determine the effect of taxa richness on chimera formation within a sample. Although numbers of reads within treatments varied this did not have a significant (ANOVA, P > 0.05) effect on chimera frequencies, number of OTUs and/or Shannon Index. However, all the analyses were also performed on a normalized data set by subsampling equal read numbers from each treatment and observations were found to be congruent with the non-normalized data (data not shown).

Statistical analysis

Species richness, or in this case OTU richness, takes no account of the evenness of the distribution. An index with better properties is the Shannon index (33). This increases with more taxa but also as the distribution of abundances across taxa becomes more even. The Shannon Index of biodiversity was established for each sample using Vegan R (34). To analyze the relationship between overall chimera percentage and the explanatory variables (e.g. phylogenetic relatedness, richness, diversity, number of reads), a linear model was fitted to the data, giving a multiplicative coefficient for each explanatory variable. An ANOVA was then performed to statistically determine which of the variables had an effect on the chimera percentage. Variables without a significant effect on chimera percentage were removed and the model was refitted to give accurate ANOVA results. A probability (P-value) <0.05 was considered significant.

RESULTS AND DISCUSSION

Here, it was possible to assess the effect of OTU richness, evenness and phylogenetic relatedness on chimera formation in a nSSU second-generation sequencing environmental data set. Moreover, a rigorous experimental design and bioinformatic analysis facilitated the identification of chimera breakpoint frequencies within the parent nSSU molecule. The `mock community’ contained equivalent concentration of 18S rDNA genes of individual nematodes of known identity and richness (GenBank Accession Numbers JN968213–JN968286). Nematodes are the most abundant phylum of meiofaunal environmental samples (4) representing a major part of biodiversity and perform numerous essential roles in ecosystems processes (35,36). The nematodes that were chosen included both phylogenetically distant and closely related species to emulate a likely environmental assemblage. Amplicons were sequenced on a Roche 454 GSFLX platform and generated a total of 339515 pyrosequence reads (Genbank/EMBL/DDBJ short read archive accession number SRA043810.1). AmpliconNoise (14) generated 236406 reads after removing errors arising from PCR and pyrosequencing errors and truncating sequences to a uniform 200bp. Chimeras were detected in denoised `mock community’ data using the Perseus algorithm (8) and ~42% of the sequences were classified as chimeric and were removed before taxon richness was assessed by clustering sequences into OTUs at a 99% identity threshold for each data set. A summary of the mean number of reads for each data set, denoised sequences, chimera percentages and OTUs is given in Table 1.

Denoised sequences contained between 15 and 60% of chimeras in some pools, confirming that 35 cycle PCRs do indeed generate numerous chimeras as already confirmed by previous studies (18,19,25) even within a small mock environmental data set. The results stress the importance of a chimera removal step to allow an accurate estimation of OTU numbers and robust estimates of biodiversity levels in environmental samples (8,15,37).

Overall, the mean OTU numbers were approximately double the number of unique nematode species in each pool. This could be associated not only with sequencing artifacts but also because organisms frequently contain multiple copies of heterogeneous nSSU genes (38). To assess the impact on the data set of multi-copy nSSUs, all single nematode PCR products were amplified

| Species phylogeny | Species number | OTUs at 99% | Denoised sequences | Chimera (%) | Read number |
|-------------------|----------------|-------------|--------------------|-------------|-------------|
| Close             | 48             | 87.6        | 138.40             | 35.60       | 13882.20    |
| Close             | 24             | 40.4        | 63.20              | 34.55       | 3809.00     |
| Close             | 12             | 35.8        | 42.80              | 14.57       | 6159.80     |
| Distant           | 48             | 63.2        | 161.00             | 58.98       | 5657.80     |
| Distant           | 24             | 53.6        | 119.00             | 53.57       | 10134.20    |
| Distant           | 12             | 34.4        | 58.20              | 39.93       | 7638.20     |
with unique MID-tag sequences. Of the 74 MID-tagged single nematode amplifications, 61 were single copy 18S rDNA and 10 were double copy but all taxa were represented by a similar total number of sequences in PCR reactions (data not shown).

A striking observation was the difference in chimera formation between close and distantly related nematode assemblages. In the latter, the mean percentage of sequences classified as chimeric was 55% for the 24 and 48 species pools and was significantly higher (ANOVA, \( P < 0.001 \)) than the equivalent pools of closely related nematode assemblages that had 35% chimeras. In line with the previous observation, the mean percentage of chimeras was significantly lower (ANOVA, \( P < 0.01 \)) in the 12 species pools irrespective of the similarity or distance of the individuals in the nematode assemblages (Table 1). These results suggest that chimera formation in the 5′ region of nSSU amplicon pools is significantly higher in more phylogenetically diverse and richer data sets.

Although the studies were on a smaller scale, Qiu et al. (19) and Wang and Wang (26) analyzed chimera formation with bacterial rRNA clones and also found that PCR artifacts and chimera frequency increased as species diversity increased. To further confirm this, OTU diversity within the two nematode assemblages (close and distant pools) was expressed using the Shannon Index (33). OTU diversity (Shannon Index) and OTU richness (OTUs numbers) showed a significant effect (ANOVA, \( P < 0.01, \ P < 0.001 \)) on chimera frequency further supporting the hypothesis that more diverse and richer samples generate a higher frequency of chimeric molecules. Additionally, diversity (Shannon index) had a significant effect both on the closely \( (P < 0.05, \ P = 0.0324) \) and distantly \( (P < 0.01, \ P = 0.0023) \) related nematode assemblages, and had a positive relationship with chimera frequency (Figure 1).

Analysis of chimera breakpoint occurrence in nSSU amplicon sequences revealed that regions with higher nucleotide sequence similarity had significantly higher breakpoint frequencies \( (P < 0.001, \ P = 0.00039) \) (Figure 2a and b). Indeed, in studies with bacteria using the 16S rRNA gene a large number of competing templates with fairly high sequence similarity generated more chimeras (6,16,26). Presumably, one explanation for this phenomenon may be the priming of strand synthesis by prematurely terminated templates in the next PCR round.

Different copies of the nSSU genes from the same organism may differ by up to 6.5% (18,38), and in the present study alignment of close and distantly related nematodes indicated an overall inter-specific sequence divergence of 10%, evidently enough to generate chimeras. To better reflect an environmental data set, in the present study alignment of close and distantly related nematodes was performed and had a positive relationship with chimera frequency (Figure 1).

Figure 1. Chimera percentage and Shannon Index of closely and distantly related pools of nematodes.
chimera frequencies. To further investigate the breakpoint region, the secondary structure of the amplified nSSU fragment was modeled in 12 single nematode sequences at 55°C and 65°C folding temperatures (Supplementary Table S1). Analysis of the nSSU secondary structure showed that the regions where the breakpoints occurred coincided with hairpin loop structures at both temperatures, although at 65°C regions of secondary structure were less abundant (Supplementary Table S1 and Figure 3).

Hairpin-loops are common motifs in nSSU gene secondary structure due to their importance in ribosome folding and function (39) and their presence requires greater energy for melting to occur during PCR and

Figure 2. (a) Nucleotide diversity (Shannon Index) and (b) breakpoint frequencies occurrence in single nematodes and parental chimeric sequences, respectively.
their maintenance will make it more difficult for DNA polymerases to read through. Modeling in silico revealed that the nSSU amplicon region in the present study retained some secondary structure at the primer annealing temperature (55°C) and may have been one of the causes of premature termination of DNA synthesis. This is corroborated by results of multiple sequence alignments of PCR-induced chimeras (40) that reveal the recombinant regions were correlated with DNA template secondary structures. Fewer secondary structures of the nSSU amplicon were found at 65°C, suggesting that (i) primers should be designed with a high annealing temperature and/or (ii) genes chosen for environmental metagenetic analyses should be selected for a low tendency to secondary structure formation which should reduce the disposition of complex samples to form chimeras. Nonetheless, other factors are known to minimize frequency of PCR recombination such as adding betaine and dimethylsulfoxide (41); dNTPs should never be limited so that amplification bias is reduced and the use of

Figure 3. Most frequent predicted secondary structures found on the 18S rDNA amplicon at (a) 55°C and (b) 65°C folding temperatures, using as an example two single nematodes. Arrows indicate where the most frequent breakpoints occur generally matching hairpin-loops. 79TN11 consensus and 25LBFB8F04 indicate the labels given for each nematode. dG: free energy necessary for sequence stability at a given temperature.
shorter amplicons will reduce prematurely terminated extension. Further to this, the choice of selectively amplifying target loci from genomic libraries or designing genome-specific amplification primers that will selectively amplify a single homologue (42) also narrow the opportunity for chimera formation.

The investigative use of higher annealing temperatures in our study was not possible since the optimal thermocycling conditions used to amplify meiofaunal representatives (9,43,44) preclude more stringent annealing temperatures. In fact, an intuitive general rule for metagenetic studies would be to avoid high annealing temperatures to ensure the co-amplification of large ranges of taxa from disparate phyla. This implies not only having very high quality DNA samples within uncontaminated laboratory environments but also compulsory stringent algorithms by using protocols to remove artifacts and/ or putative chimeras after sequencing. In addition, the use of reference databases to detect chimeric molecules in environmental data sets is complicated by their unpredictable diversity, meaning that reference data may not be representative of the true diversity. On the other hand, the existence of chimeric sequences in public DNA databases is well known (24,45) and the risk of classifying chimeras as new organisms is becoming higher than the risk of neglecting non-chimeric ones.

Experience from recent studies (4,9) where ~65% of the sequences generated from a 454 Roche environmental data set were discarded leads us to suggest that metagenetic analyses are the ideal ‘breeding ground’ for recombinant DNA molecules. DNA amplification by PCR has become the main crucial step used for next-generation sequencing technologies in the analysis of environmental samples and so PCR-derived artifacts are continuously increasing. Based on our analyses, the theory of chimera formation having a stochastic distribution (46) should probably be re-evaluated because their occurrence can be influenced by several factors, namely PCR conditions, amplicon nucleotide diversity, molecule folding structure and sequencing strategies. In fact, almost all steps of the molecular approach can introduce biases or errors, including the target DNA template concentration (6,19), DNA polymerases (20,47) and thermal cycling conditions (16,19,26,48). Moreover, the interaction between nucleic acids, DNA polymerases and thermal cycling are likely to be dynamic, suggesting that the identification of optimal molecular biological parameters that reduce chimera formation are likely to be locus/taxon specific.

Overall, we anticipate that our findings will enhance our understanding of chimera formation and associated optimization of pyrosequencing strategies when conducting PCR-based DNA amplification of homologous loci or multigene families. As technologies evolve (49), sequencing is likely to be employed to de novo analyse gene partitions from a range of genomic loci in order to address questions that have been hitherto intractable using chain termination sequencing (50). Thus, insights gained here have relevance to biodiversity identification and associated fields such as gene–environment interactions in host parasite co-evolution (51); pathogen recognition (52); genes underpinning immune response (53) and predator–prey arms races (54,55).

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