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<td><strong>Author(s)</strong></td>
<td>Fernandes, Neil; Steinberg, Peter; Rusch, Doug; Kjelleberg, Staffan; Thomas, Torsten</td>
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Community Structure and Functional Gene Profile of Bacteria on Healthy and Diseased Thalli of the Red Seaweed Delisea pulchra

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Abstract

Disease is increasingly viewed as a major factor in the ecology of marine communities and its impact appears to be increasing with environmental change, such as global warming. The temperate macroalga Delisea pulchra bleaches in Southeast Australia during warm summer periods, a phenomenon which previous studies have indicated is caused by a temperature induced bacterial disease. In order to better understand the ecology of this disease, the bacterial communities associated with three types of samples was investigated using 16S rRNA gene and environmental shotgun sequencing: 1) unbleached (healthy) D. pulchra 2) bleached parts of D. pulchra and 3) apparently healthy tissue adjacent to bleached regions. Phylogenetic differences between healthy and bleached communities mainly reflected relative changes in the taxa Colwelliaceae, Rhodobacteraceae, Thalassomonas and Parvularcula. Comparative metagenomics showed clear difference in the communities of healthy and diseased D. pulchra as reflected by changes in functions associated with transcriptional regulation, cation/multidrug efflux and non-ribosomal peptide synthesis. Importantly, the phylogenetic and functional composition of apparently healthy tissue adjacent to bleached sections of the thallus indicated that changes in the microbial communities already occur in the absence of visible tissue damage. This shift in unbleached sections might be due to the decrease in furanones, algal metabolites which are antagonists of bacterial quorum sensing. This study reveals the complex shift in the community composition associated with bleaching of Delisea pulchra and together with previous studies is consistent with a model in which elevated temperatures reduce levels of chemical defenses in stressed thallii, leading to colonization or proliferation by opportunistic pathogens or scavengers.


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Introduction

Disease in natural communities is increasingly seen as a major ecological factor. Moreover, a number of studies have argued that the frequency and impact of disease on natural communities is on the rise, likely due to the increasing impact of environmental stressors, such as global warming or other anthropogenic effects [1,2]. The impact of disease is arguably felt most strongly when the affected hosts are biogenic habitat formers, or so-called “ecosystem engineers”, because decline in these organisms results in a fundamental change in the physical structure of the habitat, and the loss of not just the hosts, but of the substantial biodiversity associated with habitat forming species.

In marine systems, to date the most prominent example of disease impacting habitat-forming organisms are tropical reef-building corals [3,4]. However, on temperate and boreal rocky shorelines, macroalgae (i.e. kelps and other seaweeds) dominate, and there they form the basis for extensive and highly diverse communities [5]. There is now evidence that these macroalgal forests are in decline globally, and one suggested mechanism is that of an increased impact of disease [6].

The red macroalga Delisea pulchra [7] is common and in places dominant in the shallow sub-tidal zone in southern Australia, New Zealand, Japan and Antarctica [8]. We have recently described a bleaching phenomenon in this alga, in which portions of the thallus lose their pigmentation and decay [9,10]. The consequences of this bleaching disease are severe; because of the pattern of bleaching on the thallus relative to the production of reproductive tissue, bleached D. pulchra are essentially neutered, with the amount of reproductive tissue an order of magnitude less than that of healthy individuals [11]. Bleaching is most common in summer, but rather than being a direct effect of light or temperature or other environmental stressors, it appears to be due to bacterial infection of (in particular) temperature stressed plants [10,11]. Two bacteria from the Rosebacter clade, namely Ruegeria sp. R11 and
Phaeobacter sp. LSS9, have been identified from the surface of D. pulchra, which can cause bleaching in the laboratory [11,12], and strain R11 has also been shown experimentally to cause bleaching in the field [11].

Further evidence for the role of bacteria in causing bleaching comes relates to the alga's antibacterial chemical defenses. D. pulchra produces halo-brominated furanones at its surface [13,14]. These compounds are strong antibacterials [14–20] and one important mechanism for their antibacterial activity is the inhibition of N-acyl homoserine lactone (AHL) based quorum sensing QS [13,16]. Furanones in D. pulchra are typically at their lowest in summer [21], corresponding to the peak in the incidence of bleaching. A decrease in the furanone concentration of the thallus is also correlated with tissue bleaching [9]. Most compellingly, direct experimental manipulation of furanones results in rapid bleaching of D. pulchra in the laboratory [10,11] and in the field [11].

A number of significant questions remain about the nature of this bacteria-alga interaction, but a critical one is: What is the nature of changes in the microbial community associated with bleaching? To address these questions we performed an in-depth microbial community analysis using 16S rRNA gene sequencing and metagenomics, expanding upon previous community studies based on relatively low-resolution techniques (tRFLP), which showed consistent differences in the community composition of healthy and bleached D. pulchra across sampling years, location and depth [9]. We compared bacterial communities from unbleached algae, from bleached tissue, and from apparently healthy (pigmented) tissue adjacent to bleached tissue. This later category is of particular interest, because such tissue, although visibly unaffected, contains furanone levels comparable to bleached tissue [9]. Thus by understanding the microbial communities on chemically poorly defended, but apparently unaffected, contains furanone levels comparable to bleached tissue, and from apparently unbleached algae, from bleached tissue, and from apparently healthy (pigmented) tissue adjacent to bleached tissue. This later category is of particular interest, because such tissue, although visibly unaffected, contains furanone levels comparable to bleached tissue [9]. Thus by understanding the microbial communities on chemically poorly defended, but apparently otherwise undamaged tissue, we may gain insight into the progression of the infection process.

Materials and Methods

 Samples and Community DNA Isolation

TriPLICATE samples of healthy and bleached individuals of D. pulchra were collected at depths of 2–4 m from Bare Island (33°39′38″S, 151°14′00″E), off the coast of Sydney, Australia, during the late Australian summer (23rd March 2008). While this sampling in this study is confined to one time and place, we note that the goal here was a more detailed examination of shifts of microbial community composition on D. pulchra associated with bleaching, which have already been demonstrated to occur across multiple years and locations [9]. The sampling was covered by permit no P00/0034-6.0 of the New South Wales Department of Industry and Investment. The algae were transported to the laboratory in polyethylene bags containing seawater kept cool by ice packs. Within two hours of collection, tissue samples were individually excised from three healthy (H) and three bleached (B) algae, as well as three tissue samples adjacent (A) to bleached tissue of the same thallus (Figure S1). Community DNA from these fresh tissue samples was extracted by a modified enzymatic-detergent lysis protocol [22] (see supplementary information for details).

Phylogenetic Analysis

A modification of the nested PCR-denaturing gradient gel electrophoresis (DGGE) approach [23] was used to obtain a community fingerprint of all replicate DNA extracted from samples types H, B and A. Further details of the community fingerprint experiments are given in the supplementary information. Clone libraries for the 16S rRNA gene were constructed as previously described [24]. Briefly, 16S rRNA genes were amplified separately from DNA extracted from duplicates of each sample type using PCR primers 27F and 1492R [25]. PCR amplicons were cloned into the Zero blunt TOPO PCR cloning kit (Invitrogen, Carlsbad, Ca, USA) following the manufacturer’s instructions and libraries were sequenced. Forward and reverse reads were assembled and 16S rRNA genes greater 1200 bp were aligned with the SILVA web aligner tool using default parameters [26]. Putative chimeric sequences were identified and removed using the program Mallard [27]. A total of 4670 aligned sequences were clustered by the program MOTHUR [28] into operational taxonomic units (OTUs) using the furthest neighbor method and identity cut-offs of 0.03 representing approximately species-level clustering. In order to identify discriminating species from 16S rRNA gene analysis, OTU-abundance tables were subjected to multivariate statistical analysis using the Similarity Percentage Analysis program (SIMPER) [29] implemented in the PRIMER software package (Plymouth Routines in Multivariate Ecological Research) [30]. Bray-Curtis similarity matrices were used to cluster sample types and SIMPER analysis was used to determine the OTUs that contribute to greater than 1% of the total difference between samples from bleached, adjacent and healthy tissues. Because Bray-Curtis similarity measures the contribution of each species, the average dissimilarity between different samples can be expressed in terms of the average contribution from each species. The Ribosomal Database Project (RDP) classifier [31] was then used with default parameters to classify all sequences.

Metagenomic Analysis

Metagenomic clone libraries were produced and sequenced from algal community DNA using previously described procedures [32,33]. Briefly, large-scale sequencing was performed on ABI3730XL sequencers for duplicate samples of sample types H, B and A, which corresponded to the ones used for preparation of the 16S rRNA gene libraries. Shotgun sequences were assembled using the Celera Assembler software version 5.1 with twelve percentage errors permitted in the unitigger (utgErrorRate) and 14% error allowed in the overlapper (ovlErrorRate), consensus (consErrorRate), and scaffold (cgwErrorRate) modules. Seed length (merSizeOvl) was set to 14. Overlap trimming, extended clear ranges, and surrogates were turned on. Fragment correction and bubble popping were turned off. In order to filter out eukaryotic DNA contamination, arising from co-extraction of algal DNA, the assembled data was filtered by searching them against The National Center for Biotechnology Information (NCBI) nucleotide database (NT) using BLASTN [34]. Further information on the filtering and the annotation of the metagenomic sequences can be found in the supplementary information. Annotated matches to the COG databases [35] were organized into COG categories. Matrices of raw counts of COGs were standardized to account for the unequal sequence coverage between samples. Bray-Curtis similarity matrices were used to cluster sample types and SIMPER was employed to determine the contribution of each COG to differences between the sample types. Shotgun and 16S rRNA gene sequencing data are available through the Community Cyberinfrastructure for Advanced Microbial Ecology Research and Analysis (accession: CAM_PROJ_BotanyBay).
Results

Taxonomic Composition and Differences of Bacterial Communities

To confirm that the samples collected in this study show similar patterns of bacterial community shift as seen previously [9], we performed finger-printing analysis (DGGE). Clear differences in the healthy (H) versus bleached (B) samples could be visualized on the DGGE gels (Figure S2) and were reflected in the sample dendograms (Figure 1a). Also, the community associated with healthy tissue adjacent to diseased parts of the thalli showed similarities to both the healthy and bleached samples, consistent with previous observations [9].

Filtered and quality-checked 16S rRNA gene libraries (see Material and Methods) consisted of 694 and 1091 sequences for the two healthy tissues of D. pulchra, 767 and 812 sequences from bleached tissues and 416 and 890 sequences from tissues extracted from the region adjacent to bleached tissue (Table 1). Chao1-estimates at 0.03 sequence differences showed lower community richness for healthy samples (216 to 151), than for adjacent samples (507 to 937) or bleached thalli (846 and 937) (Table 1). This shows that healthy D. pulchra supports less diverse communities than bleached samples (p = 0.0062 for H versus B; unpaired student t-test), which is also reflected in the rarefaction curve analysis (see Figure S3).

Bray-Curtis similarity analysis and clustering of 16S rRNA gene libraries reflected the patterns seen in the DGGE fingerprinting (Figure 1b). At OTU cut-offs of 0.03 dissimilarity, healthy samples showed less than 20% similarity with the other two samples types, while bleached and adjacent samples shared at least 35% of the sequences found. Adjacent and bleached samples shared as many as 143 sequences, while each of these samples shared only around 30 sequences each with the community of healthy samples (see Figure S4).

Sample discrimination analysis (SIMPER) showed that at the 0.03 difference level only 19 sequences accounted for 60% of the average dissimilarity between bleached and healthy samples (see Figure S5). The taxonomic assignment of these “discriminatory” OTUs and their relative abundance in the 16S rRNA gene clone libraries of all three sample types are given in Figure 2. An OTU belonging to the family Colwelliaceae had an abundance of 21.46% in bleached samples and was reduced to an abundance of about 2% in adjacent tissue, and was not detected in healthy thalli. Similar trends were observed for OTUs belonging to the taxa Thalassomonas, Rhodobacteraceae, Cellulophaga, Aquimarina, Marinomonas and Hyponemadaceae. In contrast, healthy samples had a very low abundance in OTUs belonging to the Rhodobacteraceae, Parvularcula, Haliscomenobacter, Flexibacteraceae and Saprospiraceae compared to the other two sample types. We also found no evidence for the presence or enrichment of Ruegeria sp. R11 and Phaeobacter sp. LSS9 on bleached samples.

Together this data show clear differences in the taxonomic composition of communities of healthy D. pulchra versus thalli that are bleached or parts that appear healthy and are located immediately adjacent to bleached regions. These changes seem to be largely driven by substantial shifts in the abundance of a few OTUs. To better understand these changes, we further analysed these bacterial communities using a metagenome sequencing approach.

Functional Gene Profiles of Healthy, Bleached and Adjacent Communities

Shotgun sequencing, functional annotation and statistical analysis of six microbial metagenomes from two samples each of diseased, healthy and adjacent tissues of D. pulchra identified a number of “discriminatory” functions and those were used to cluster samples in dendograms (Table 2, Figure 3) and to ordinate by multi-dimensional scaling (MDS) (Figure S6). Overall, functional profiles based in COG annotation from healthy samples clustered together and were distinct from the bleached and

Table 1. The number of sequences analyzed, OTUs at a cut-off of 0.03, and Chao 1 OTU richness estimates for each 16S rRNA library prepared from bleached tissue (B) healthy tissue (H) and adjacent tissue (A).

<table>
<thead>
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<th>OTUs at 0.3</th>
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<tr>
<td>H2</td>
<td>694</td>
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<td>H3</td>
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<td>364</td>
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<tr>
<td>B3</td>
<td>767</td>
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adjacent samples. This clustering based on functional profile reflected those seen for the taxonomic assignment.

A ranked plot of the relative contribution of each COG function showed that only nine COG functions make up for 60% of the average dissimilarity between bleached and healthy samples (Figure S7). The relative abundances of those discriminatory functions in the three different samples are shown in Figure 4. In general, the individual discriminatory functions became more abundant in sample types A and B versus healthy sample. Substantial enrichment in sample types A and B was observed for the functions acyl-CoA synthetases (AMP-forming)/AMP-acid ligases II (COG318), response regulators consisting of a CheY-like receiver domain and a winged-helix DNA-binding domain (COG745), signal transduction histidine kinase (COG0642), NAD-dependent aldehyde dehydrogenases (COG1012), cation/multidrug efflux pump (COG841), ABC-type multidrug/protein/lipid transport system (COG 1132) and non-ribosomal peptide synthetase modules and related proteins (COG1020).

Discussion

Bacterial Community of Bleached and Adjacent Tissue

Bleaching of *D. pulchra* is associated with clear changes in the community composition, not only directly at the affected part of the thallus, but also in the surrounding areas. Community richness of bleached and adjacent tissue was in general similar, but greatly increased (>3-fold; see Table 1) compared to healthy samples. This increased diversity could be the result of changes in the furanone concentrations, which are high in healthy *D. pulchra* [19], but greatly reduced in bleached algae and their apparently healthy tissue [9]. The loss of selective and inhibitory effect of the alga’s chemical defense could thus allow for a greater variety of bacteria to colonize and these might potentially include opportunistic pathogens and scavengers (see below). Increases in bacterial diversity have also been observed when the Caribbean coral *Montastraea faveolata* was afflicted by White Plague Disease [36] and when the common reef-building coral, *Porites cylindrica* was exposed to stressful environmental events [37].

Communities of diseased thalli are characterized by an abundance of OTUs belonging to the genera *Cellulophaga* and *Thalassomonas* as well as the family *Colwelliaceae*, some of which are also found in adjacent tissue, but absent on healthy algae. Algal diseases are often associated with cell-wall degradation and members of the genus *Cellulophaga* have been shown to exhibit potent extra-cellular alginolytic or agarolytic activity [38]. OTUs within the *Colwelliaceae* account for most of the difference between healthy and bleached tissue and this family contains several genera with known or putative pathogenic and anti-algal representatives. For example, *Thalassomona layana* has been implicated as a pathogen for ‘white plague-like disease’ in corals in the Red Sea [39]. Bacteria within the genera *Alteromonas* and *Pseudoalteromonas* (family *Colwelliaceae*) have also been shown to exhibit algicidal activity against eukaryotic microalgae [40] and have been found associated with decaying matter of algal blooms [41].

These observations are consistent with two alternative models for the changes in microbial communities associated with bleaching in *D. pulchra*. Firstly, changes could be a result of opportunistic pathogens that either colonise or proliferate on relatively chemically undefended tissue and subsequently cause the bleaching disease. The absence of previously identified pathogens of *D. pulchra* (i.e. *Ruegeria* sp. R11 and *Phaeobacter* sp. LSS9) in the samples taken in the current study would also indicate that other bacterial taxa might have the ability to induce disease. Secondly, an alternative model would be that disease occurs independently
Loss of Microbiota from Healthy Samples

Taxonomically, healthy tissue was characterized by the abundance of bacteria belonging to the genera Parvularcula and Haliscosmenobacter, and the family Rhodobacteraceae. The most abundant Rhodobacteraceae phylotype (19% of the sequences found) could also be assigned with moderate confidence (70%) to the genus Thioclava. The genus Thioclava as well as the second most abundant group, the genus Parvularcula, harbor genes for the degradation of the abundant algal osmolyte dimethylsulfoniopropionate (DMSP) [42] and are known to oxidize inorganic sulfur metabolites [43], such as the byproducts of DMSP degradation [44]. Bacteria involved in the biogeochemical cycling of sulfur are also abundant in coral-associated communities and are thought to play a significant role in structuring the community and to be important for coral health [45,46]. Members of the family Saprospiraceae (order Bacteroidales), to which the genus Haliscosmenobacter also belongs (see Figure 2), have also been isolated from marine sponges and algae from the southern coastline of Thailand [47]. While little is known about the ecophysiology of other Saprospiraceae, members of the Bacteroidales are generally associated with the degradation of complex organic materials and Haliscosmenobacter hydrosis is abundant in activated sludge flocs and thought to be involved in the hydrolysis of polysaccharides [48].

While the ecological role of these bacterial associates that dominate the community of healthy *D. pulchra* is not clear, they belong to taxonomic groups that are commonly found on marine algae [49–52]. These groups might thus represent epiphytes that are generally not harmful to the host and are important for structuring a stable microbial community. Importantly, these community members are also in lower abundance on bleached tissue as well as on adjacent parts, where there are no visible changes. This observation would be consistent with a probiotic model, which was proposed for corals [53] and posits that the normal microbiota associated with living surfaces offers protection against microbial infection and disease. According to this concept, the community of the adjacent tissue would have already lost key members of the “normal” microbiota, a shift that may allow for subsequent proliferation of harmful bacteria in the community. This community shift might be due to the loss of algal defense (furanone) and could result in a dysbiosis of the epiphytic bacterial community prior to disease development. Thus it might be important to consider the state of the microbial community prior to disease development and define its effect on disease initiation or outcome.

Figure 3. Dendrograms comparing the level of similarity between metagenomic libraries constructed from bleached tissue (B), from tissue adjacent to bleached tissue (A) and healthy tissue (H) using a matrix containing ORFs that could be matched to COGs with E-value cut-offs of $10^{-5}$ (a), $10^{-10}$ (b) and $10^{-20}$ (b).

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bacterial communities. For example, efux pump mutants of the plant pathogens *E. chrysanthemi* and *E. amylovora* are less infectious than the wild-type strain when co-inoculated with other endophytic bacteria producing antimicrobial compounds [62,63].

Two-component signal transduction (TCST) systems (COG 0642) are used by many bacteria to sense environmental signals and modulate expression of genes [64], including ones involved in virulence, antibiotic resistance responses, and colonization in many pathogenic bacteria [65,66]. For example, in the plant pathogenic bacteria *Xanthomonas campestris*, *Pectobacterium carotovorum* and *Ralstonia solanacearum*, TCST systems have been shown to provide environmental signal inputs to global virulence regulation [67]. The abundance of TCST systems found in communities of bleached tissue could thus enable bacteria to modulate gene expression in response to environmental changes and regulate virulence genes responsible for bleaching. The potential importance of regulation in disease was also recently highlighted in the analysis of the genomes for the *D. pulchra* pathogens *Nautella* sp. R11 and *Phaeobacter* sp. LSS9 [12]. These two pathogens contained no unique set of putative virulence factors when compared to other closely related non-pathogenic strains, but had a unique LuxR-type regulator that was hypothesized to regulated virulence.

**Conclusion**

Marine macroalgae, including *D. pulchra*, are major habitat formers in temperate ecosystems, and diseases of these organisms have the potential to cause significant impacts on community structure, population levels and biodiversity in marine ecosystems [1]. Campbell et al [9] have proposed a model for bleaching in *D. pulchra*, in which environmental stress (particularly elevated seawater temperatures) leads to a decrease in chemical defenses (furanones), which in turn leads to a shift in bacterial community composition, which can include opportunistic pathogens or scavengers. This shift might be further exacerbated by the loss

<table>
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<th>ORFs assigned to COGs</th>
<th>% of ORFs assigned to COGs</th>
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<td>18205</td>
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</tr>
<tr>
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<td>6998</td>
<td>1542</td>
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of certain key members of the “normal” microbiota of *D. pulchra*, which might have probiotic functions. Our observations are consistent with such processes and we have identified bacterial species that might be involved in either the disease initiation or secondary scavenging. Functional gene analysis of communities of bleached sample revealed further an abundance of activities that are likely to play key roles in algal colonization and competition. However the role of these groups of bacteria and genes in the bleaching disease of *D. pulchra* needs further experimental confirmation. Further advances in understanding the bleaching disease in *D. pulchra* would require tracking temporal changes in key species, and in the expression of these key functional genes, at different stages of disease.

**Supporting Information**

**Figure S1** Bleached *Delisea pulchra* collected from Bare Island, Sydney, Australia. The bleached section in the mid thallus region is indicated by the arrowhead.

**Figure S2** Comparison of different samples by DGGE fingerprinting. Lane M: DGGE Markers, Lanes A1, A2 and A3 contain DGGE bands from tissue adjacent to bleached tissue, Lanes B1, B2 and B3, DGGE bands from bleached tissue, Lanes H1, H2 and H3, DGGE bands from healthy tissue.

**Figure S3** Rarefaction curves comparing the number of OTUs at a distance of 0.03 between 16S rDNA gene libraries constructed bleached tissue (B), from tissue adjacent to bleached tissue (A) and healthy tissue (H).

**Figure S4** Venn diagrams showing the number of OTUs shared between communities on bleached tissue (B), healthy tissue (H) and adjacent tissue (A) with OTUs at 0.03 difference.

**Figure S5** The contribution of different OTUs (at 0.03 sequence difference cut-off) to difference between 16S rDNA gene libraries from bleached and healthy samples.

**Figure S6** Multidimensional-scaling (MDS) plots comparing the level of similarity between metagenomic libraries constructed from bleached tissue (B), from tissue adjacent to bleached tissue (A) and healthy tissue (H) using a matrix containing ORFs that could be matched to COGs at E-value cut-offs smaller $10^{-5}$ (1), $10^{-10}$ (2) and $10^{-20}$ (3).

**Figure S7** The contribution of individual COGs to the difference between metagenomic libraries from bleached and healthy samples.

**Text S1** Supplementary Material and Methods.

**Acknowledgments**

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**Author Contributions**

Conceived and designed the experiments: NF PS TT. Performed the experiments: NF TT. Analyzed the data: NF DR TT. Contributed reagents/materials/analysis tools: DR. Wrote the paper: NF PS TT.

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**Author Contributions**

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