Accelerated low water corrosion: the microbial sulfur cycle in microcosm

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Accelerated low water corrosion is a form of marine steel corrosion caused by bacterial activity. It has a global spread and is potentially responsible for billions of pounds of damage. We have determined in detail both the chemistry of corrosion products and the associated microbiology at a UK site. The corrosion products form a layered structure with iron sulfides at the steel surface and iron oxides and sulfates in contact with water. The iron sulfides are formed by reaction of steel with hydrogen sulfide formed by sulfate-reducing bacteria and are oxidised through a series of sulfur oxidation states by sulfate-oxidising bacteria, forming acid at all stages and encompassing the whole of the bacterial sulfur cycle. The bacteria involved are endemic in anoxic bed sediment, and the process is a response to the presence of steel as an electron donor, and the generation of anoxic microenvironments within corrosion products.

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INTRODUCTION

Accelerated low water corrosion (ALWC) (Fig. 1) is an aggressive form of microbially influenced corrosion that may occur on steel in estuarine and marine structures.1-3 Corrosion rates on marine steel as high as 1 mm/y, and typically around 0.3 mm/y have been reported, compared to electrochemical corrosion rates in marine environments of around 0.05 mm/y.3 This makes ALWC a major threat to marine and coastal infrastructure internationally. Previous work has indicated the role of both sulfate-reducing and sulfur-oxidising bacteria in the formation of ALWC4-12 and highlighted the role of marine nutrient contamination in promoting microbial activity and hence enhanced corrosion rates,13 alongside the electrochemical environment within the intertidal zone.14 A recent study by Malard15 identified at least fifteen genera of the Order Desulfobacterales associated with ALWC tubercles. Sulfate-reducing bacteria utilise sulfate, thiosulfate and sulfur as electron acceptors and produce hydrogen sulfide (H2S) as the product of both SO42− and S0 reduction,11,15 which may then react with steel to produce iron sulfides11 and ultimately pyrite (FeS2). Both H2S and pyrite may then be oxidised, producing sulfuric acid, and Fe oxhydroxides and sulfates.16 However, the full mechanism of ALWC has yet to be elucidated and aspects of its development over time remain unclear.17 The introduction of a conductive electron donor (steel) into the marine environment also has the potential to provide a substrate for metabolic mechanisms that have only recently been identified, notably extracellular electron transfer.16,19 The phenomenon itself is global and has been reported from all climates, but it is unclear if any apparent spread of the issue is related to transmission of causative bacteria via vectors such as international shipping, or caused by bacteria in local environmental reservoirs such as marine sediment.1

In this study, we have combined spectroscopic (Fourier transform infrared (FTIR) and X-ray photoelectron spectroscopy (XPS)) and crystallographic (X-ray diffraction (XRD)) techniques with metagenomic analysis using the GeoChip microarray system20,21 and related the results to associated bacterial communities in order to determine the precise mechanism of ALWC for a UK site (Shoreham Harbour, Supplementary Fig. 1). This is important not just for its engineering and economic implications, but because it also represents a phenomenon whereby natural microbial communities respond in their development to the presence of an electron donor, anthropogenically introduced into the environment. We have also identified the residence of the individual components of the microbial community within the surrounding environment in order to identify infection vectors for structural steel and environmental risk factors for ALWC. Traditionally, microbiological culture has been employed to determine the composition of microbial communities associated with MIC, but it is time consuming and has a degree of artificiality due to its isolation of the subject microorganism from the original community; a bacterium which is readily cultivatable in the laboratory may be dormant or otherwise have a negligible influence within its community.22 In addition, analysis of community interactions is further limited by our current inability to cultivate the majority of bacterial species present in nature; more than 99% remain uncultivated.23,24 The use of a microarray system here gives us the potential to investigate a much wider range of the microbial community and their interaction with steel and the local environment than culture based studies allow.

RESULTS

Corrosion mineralogy

The results of SEM analysis of ALWC and more typical marine corrosion products from Shoreham Harbour, UK, are shown in Fig. 1b–e. The measurements confirmed the presence of iron sulfides on the very inner surface of blisters, and their intergrowth with iron oxhydroxides-sulfates (identified from XRD and FTIR data) within the blister layer structure (Fig. 1b). The outer surface of the blisters was shown to be composed of iron oxides overlain by a biofilm (Fig. 1c), possibly colonised by a mat of rod-like bacilli, and with widely distributed spheroidal structures composed of Fe oxides. These possibly result from crystallisation from gel, or are a
product of microbial activity\(^2\) (Fig. 1d). The association of these with biofilms was also established as diatomaceous algae were observed cemented into the oxide and sulfate.

XRD results are shown in Fig. 2. ALWC blisters are characterised by the presence of sulfate green rust (\(\text{Fe}^{3+}\text{Fe}^{2+}\cdot\text{OH})_{12}\text{SO}_4\cdot2\text{H}_2\text{O}\) \(^2\)), magnetite, maghemite, lepidocrocite (\(\gamma\text{FeOOH}\)) and/or goethite (\(\alpha\text{FeOOH}\)). In addition the inner layers show the presence of pyrite (\(\text{FeS}_2\)). Aragonite derived from invertebrate shells is variably present in all blisters analysed. Iron fluorescence caused by the use of Cu K\(\alpha\) radiation and the presence of amorphous material means that XRD cannot fully identify all the chemical components of the corrosion products.

A summary of the results of FTIR analyses are shown in Fig. 3. Despite broad similarities between patterns from ALWC and typical marine corrosion, there are some differences relating to distinctions in mineralogy and sulfur content, most notably around 850 cm\(^{-1}\) and 1050 cm\(^{-1}\) (shown in representative samples in Fig. 3a). Both sets of analyses show absorption from bond stretching modes characteristic of Fe oxide and oxyhydroxide minerals and Fe hydroxide gels. They also both show strong absorbance by sulfate (\(\text{SO}_4^{2-}\)) and bisulfate (\(\text{HSO}_4^-\)) functional groups, and indications of the presence of amaranthite (\(\text{Fe}^{3+}\text{Fe}^{2+}\cdot\text{O} \cdot \text{SO}_4\cdot7\text{H}_2\text{O}\)) and Rozenite (\(\text{Fe}^{2+}\text{SO}_4\cdot4\text{H}_2\text{O}\)), both related to sulfate green rust. The ALWC spectra include, in addition, absorption bands characteristic of pyrite (\(\text{FeS}_2\)), thiosulfate (\(\text{S}_2\text{O}_3^-\)) and in some instances sulfite (\(\text{SO}_3^{2-}\)). Figure 3b shows data from 56 samples in terms of band ratios between low absorbance regions and absorbance bands for sulfate and thiosulfate. The thiosulfate absorbance bands clearly differentiate ALWC, although there is a continuum in spectral characteristics reflecting the progressive development of sulfur species in corrosion. Supplementary Table 1 shows the main absorption peak positions identified in each sample, and potential causative molecular groupings.

The utility of XPS in analysis of steel surfaces undergoing corrosion was highlighted by Seyeus et al.\(^2\) Two samples were selected for X-ray photoelectron analysis, one of ALWC and one of more typical corrosion. The samples were imaged using backscattered electrons after XPS analysis. The outer surface of the ALWC sample had a clearly defined biofilm with a significant chloride content (Fig. 1c). This covered a microcrystalline to amorphous layer of Fe oxyhydroxides, which in turn covered a layer of crystalline Fe sulfates and oxyhydroxides. The inner surface of the ALWC sample was also dominated by Fe oxyhydroxides, but also a well crystalline, Fe–S–O phase identified as sulfate green rust. The normal corrosion sample was uniformly dominated by Fe oxyhydroxides, with poor crystal structures. The inner surface showed a microporous structure that may be important in maintaining fluid contact with the underlying steel and allowing continued corrosion (Fig. 1d).

The results of XPS depth profiling of representative samples of ALWC and normal marine corrosion are shown in Figs 4–6. Analytical data is presented in Supplementary Table 2 and binding energy peak positions are given in Supplementary Table 3. In each case the sample was split into two and mounted for analysis of the outermost and innermost surfaces. In the ALWC sample the

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Fig. 1 Images of steel corrosion. a Macroscale ALWC blisters developed on steel pile wall in intertidal zone. Scale bar 50 cm. b High magnification view of inner surface ALWC blister showing amorphous FeS overlying well crystallised Fe sulfates (green rust). Scale bar 10 \(\mu\)m. c Outer surface of AWLC showing biofilm overlying star like acicular goethite formations (c.f.\(^2\)). The depth profile is given by the edge of an Ar\(^+\) focussed ion beam ablation crater. Scale bar 10 \(\mu\)m. d Outer surface of ALWC blister showing rod-like bacilli and the development of Fe oxide or carbonate nodules. Scale bar 10 \(\mu\)m. e Outer surface of more typical marine corrosion showing microporous toroidal magnetite formation. Scale bar 25 \(\mu\)m.
profiles show significant concentrations of Mg and Cl throughout, derived from drying of initially seawater-wet samples, and trace levels of N related to the presence of biofilm material. Iron is more or less constant in concentration throughout the sputtering of the outer surface (Fig. 4a). Alongside iron the dominant component is oxygen, and there is a notable lack of sulfur in the outer layers (typically close to the detection limit of 1 atom %). Profiles from the very inner surface of the blister outward (Fig. 4a) show sulfur concentrations are significant and reach a maximum (11.5 atom %) close to the innermost layer. The initial low in S content on the inner surface may reflect post-sampling surface oxidation. Oxygen and iron are still the dominant components. Deconvolution of the Fe 2p binding energy spectrum (Fig. 5a–d) shows characteristic peak shifts in relation to the Fe compounds and bonding environment present. The outermost layer shows the corrosion material is dominated by Fe\(_3^+\) sulfates (Fe\(_2\)(SO\(_4\))\(_3\)·XH\(_2\)O), goethite/lepidocrocite (FeO(OH)), magnetite (Fe\(_3\)O\(_4\)), Fe\(_2\)O\(_3\) (probably as maghemite by comparison with XRD data), iron carbide (Fe\(_3\)C) and FeO (Fig. 5a). The same range of compounds was identified after sputtering, but with metallic iron particles, pyrite (FeS\(_2\)) and FeS present (Fig. 5b). FeS may be present as amorphous iron sulfide, or as mackinawite,\(^1\) but mackinawite has not been identified by XRD in this study. Despite the presence of sulfate compounds the sulfur concentration in the outer layers is low (<5 atom % compared to up to 15 atom % close to the inner surface). The same process for the innermost layers shows the presence of magnetite, maghemite/hematite, and iron sulfide compounds (Fig. 5c, d). Examining the S 2p binding energy fine structure (Fig. 5e, f), shows that after sputtering the sulfur speciation can be seen to be dominated almost entirely by iron monosulfide FeS, with a smaller component of pyrite (FeS\(_2\)) and cluster type compounds. All major phases and non-crystalline components identified by XPS are confirmed by the XRD and FTIR data.

In the normal marine corrosion sample there are significant differences in the XPS profiles. Carbon is abundant in the outermost layer (Fig. 4b), partly present as aragonite/calcite from the biofouling of the steel. After sputtering the outer part of the corrosion layer is shown to be dominantly Fe and O, but N and S occur at trace levels throughout. The maximum sulfur...
concentration is ~2 atom % and mainly around ~1 atom % making it significantly less abundant than in the ALWC blister (Fig. 4b). The inner surface is also dominantly Fe–O, but with up to ~8 atom % S (Fig. 4b). The Fe 2p spectra on the inner side of the corrosion layer after sputtering shows the dominant sulfur compound is Fe$^{3+}$ sulfate (Fe$_2$(SO$_4$)$_3$·XH$_2$O), goethite/lepidocrocite, magnetite and trace pyrite (Fig. 6a, b). The innermost layer of the corrosion (i.e. with no sputtering) consists of maghemite, green rust and goethite/lepidocrocite. Examination of the S 2p binding energy fine structure (Fig. 6c, d) shows the S in the innermost layer to be dominantly Fe$_2$(SO$_4$)$_3$·XH$_2$O, whilst after sputtering a more complex mineralogy is revealed. This includes Fe$_3$(SO$_4$)$_3$·XH$_2$O, sulfite and pyrite. Further analysis to determine the relative proportion of sulfur compounds was carried out. The very innermost layer (i.e. with no sputtering) is almost entirely Fe$_2$(SO$_4$)$_3$·XH$_2$O, whilst in the interior of the corrosion layer, the 1% total S is composed of ~20% FeS, 40% Fe$_2$(SO$_4$)$_3$ and 40% SO$_3$.$^{-2}$. Iron mineralogy is always dominated by Fe oxide species.

The iron species binding energy peaks show a number of complexities in the interpretations above. The observed binding energy depends not only upon initial state effects (the charge on the atom), but also final state effects, especially electron relaxation. In this case, S$_{2-}$ in FeS$_2$ is highly polarisable and therefore has large relaxation effects. Due to this final state effect, FeS$_2$ has lower binding energy than FeS. In Fe$_2$O$_3$, both Fe$^{2+}$ and Fe$^{3+}$ are present. Fe 2p$_{3/2}$ core level BE peaks of Fe(II) and Fe(III) appear at ~709.8 eV and ~710.8 eV respectively. On the other hand, in Fe$_2$O$_3$, Fe is only in the +3 oxidation state. So the Fe 2p$_{3/2}$ core level peak should appear around ~710.8 eV. For that reason, Fe$_3$O$_4$ is assigned to 709.8 eV and Fe$_2$O$_3$ is assigned to 710.8 eV. References for binding energy peak positions are given in the Supplementary Table.

The results indicate the inner surfaces of ALWC blisters are dominated by iron monosulfide and pyrite (up to 15 atom% S). These are oxidised on their outer surfaces to a range of green rust compounds. The outer materials also include goethite, magnetite and maghemite/hematite, and minor amounts of relict steel phases. Fourier transform infrared spectra of corresponding samples show the presence of the same Fe phases, but also sulfite (SO$_3^{2-}$) and thiosulfate (S$_2$O$_3^{2-}$). X-ray photoelectron spectroscopic analyses of more typical marine corrosion products show much lower S concentrations (1–5 atom % S), with S dominantly present as sulfite in mixed Fe oxidation state sulfate, green rust, amarantite and rozenite. Pyrite is still present in the normal marine corrosion, but in much lower concentrations associated with the inner surface of corrosion.

Microbiology of ALWC

Microbial community functional structures of eighteen samples: three duplicates of two microbially influenced corrosion samples (BT and DBMIC); three duplicates of two seawater samples (BT and DBWAT); and three duplicates of two bed sediment samples (BT and DBSED) were analysed with GeoChip 5.0 (Glomics Inc.) and the results are summarised in Supplementary Table 4. Details of the sample sites are given in the methods section—the Brighton Terminal (BT) is in a non-tidal lagoon and the Dredger Berth (DB) is in a tidal area connecting to the main estuary. The number of detected genes sequences were between 30,294 and 51,748 depending on the samples. The alpha-diversity richness and evenness) of the microbial communities was assessed using the Shannon–Wiener index (H), Simpson index (1/$D$) and Simpson evenness ($E$).$^{28}$ The Shannon–Weaver index values were from 8.47 to 9.91 across the samples, showing a high diversity in all samples and no significant differences among them. Consistently, the 1/$D$ (values from 713 to 5468) and $E$ (values from 0.02 to 0.11) indices did not show any significant differences between the samples. At the phylogenetic level, 32 bacterial phyla and up to 1948 bacterial species (depending on the sample) were detected in the samples.
with the most predominant phyla being Proteobacteria 57%, Actinobacteria 21.3%, Firmicutes 9.3%, Bacteroidetes 3.5%, Cyanobacteria 2.5%, Deinococcus-Thermus 1.4%, and Chloroflexi 1.1%. All the others were represented by less than 1%. These data show a high diversity of microbial communities in the different samples.

When looking at the microbial biogeochemical cycles gene categories, genes related to the carbon cycle were the most present in all samples, with variations in the intensity depending on the sample. Genes related to nitrogen, phosphorus and sulfur cycles were found in lower intensity than carbon cycle and showed no significant differences between all samples. We specifically focused the analyses on the sulfur cycle functional genes as these are inferred to be involved in the key biochemical processes in ALWC. The genes and related mechanisms studied were apr, and cysH (sulfate reduction); dsr, sir, and cysI (sulfite reduction); fcc, and sqr (sulfide oxidation). All these genes were derived from 461 different bacterial species belonging to 29 phyla, with the most abundant being the Proteobacteria (31.1%) followed by the Firmicutes (28.3%); Actinobacteria (21.3%); and the Chloroflexi (21.3%). Genes involved in sulfate reduction, sulfite reduction, sulfide oxidation, and thiosulfate oxidation show a broadly similar pattern between samples for each of the two environments (Fig. 7). Within BT, seawater displays the greatest level of gene expression between each metabolic category, whereas sediment and ALWC show broadly similar levels of activity. The DB samples indicate the highest activity is within

![Fig. 5 XPS narrow scan spectroscopy of typical ALWC products. References for chemical shifts are given in supplementary file.](image-url)
the marine sediment, with seawater displaying the lowest degree of gene expression. When focusing the analysis on genes involved in sulfide and thiosulfate oxidation, and sulfate and sulfite reduction, we observed that gene expression was broadly similar between categories, with no dominant gene or metabolic pathway being apparent.

The aprA, aprB and cysH genes were active in 51, 30 and 9 species respectively; composed on average of Proteobacteriacea (53.4%); Firmicutes (8.2%); Ascomycota (4.8%); and Crenarchaeota (3.4%). Thirty-four percent of the detected genes were from uncultured bacteria. The species distribution was Proteobacteriacea (31.7%); Firmicutes (6.9%); Actinobacteria (3.9%); Euryarchaeota (2%); Crenarchaeota (1.3%). Twenty-six of the detected genes were found in uncultured bacteria. The fccA gene was found in 38 species; the sir gene, only 14 species. These were found to belonging mainly to Proteobacteriacea (62.7%); Chlorobia (7.9%); Actinobacteria (3.2%); Aquilicace (3.2%) and Deinococcus-Thermus (2.4%). Twenty-nine percent of the detected genes were found in uncultured bacteria. Of the sox genes, the soxA and sox B genes were observed in 14 and 5 species only. The soxC and soxY genes were found in 37 and 24 species. The dominant phylum was Proteobacteriacea (86.4%); with Chlorobia comprising 7%; Deinococcus-Thermus (2%) and Aquilicace (1%). Most of the genes could be attributed to a species (only 2.4% of the genes were found in uncultured bacteria).

When analysing the intensity of each gene, eight species represented 56% of the total intensity of all dsrAB genes that could be attributed to a species: Desulfovibulobus propionicus (9%), Holdemania filiformis (8%), Clostridium leptum (6%), Moorella thermoaceticota (6%), Desulfovibrio alaskensis (6%), Desulfovibrio vulgaris (5%), Thioalkaliibibrio sulfodophilus (5%) and Acetobacterium woodii (3%). For sqr gene, six species represented 79% of the total intensity that could be attributed to a species: Chlorobaculum tepidum (17%), Bradyrhizobium diaeoffiticiis (17%), Thiorhodospira sibirica (15%), Teredinibacter turnerai (14%), Mycobacterium para- scrofulaceae (9%) and Rhodospirillum centenum (2%). For soxY gene, eight species represented 61% of the total intensity that could be attributed to a species: Nitrobacter hamburgensis (19%), Methylobacterium populi (19%), Methylococcus nodulens (7%), Ruegeria pomeroyi (6%), Sideroxylans littoratus (5%), Chloro- obium phaeovibrioides (5%), Myceliosinus trichomosporium (3%), Halothiobacillus neapolitanus (3%). Interestingly, soxABCV genes were present in different species: Thermus aquaticus (20%), Methylobacterium extorquens (14%), Hyphomonas neptunim (5%), except Ruegeria pomeroyi (4%) that carried both soxY and soxABCV genes. None of the species was specific to any samples type (seawater, sediment or ALWC). The full results of metagenomic analyses using Geochip and 16s RNA analysis are shown in Supplementary Table 4 and summarised in Fig. 7. The range of species and gene activity identified covers most of the active sulfur cycle, including species involved in intermediate sulfur species disproportionation, and those implicated in the use of extracellular electron transfer.
Discussion—mineralogy, microbiology and the mechanism of ALWC

The data presented here allow identification of the mechanism of ALWC within the local harbour environment. In this setting normal marine corrosion typically has an outer layer, including calcium carbonate, from a macrobiofilm, overlying layers of goethite and magnetite associated with minor amounts of a range of iron-sulfur compounds. These are not limited to sulfate green rusts, but also include small amounts of FeS and sulfate compounds (SO$_4^{2-}$). In ALWC samples outer layers of goethite, magnetite, maghemite and Fe (III) sulfate with progressively higher S contents moving inwards, form over a layer of FeS (as mackinawite or amorphous iron sulfide) and pyrite (FeS$_2$), with minor green rust components. More strongly developed minor components in ALWC include sulfate (SO$_4^{2-}$) and thiosulfate (S$_2$O$_3^{2-}$) compounds. Iron (II) oxide (FeO) also occurs as a minor phase, alongside relics of un-oxidised steel including iron metal (Fe$^0$) and iron carbide (Fe$_3$C) and highly metastable phases including Fe(III)-S compounds. The inferred major mineralogy is in keeping with previous studies of ALWC mineralogy.

The formation of the FeS layer in ALWC in particular is widely recognised as the key factor in driving accelerated corrosion rates.$^{19,25}$ These studies have proposed 2 main mechanisms for microbially mediated corrosion:

Electrical microbially influenced corrosion (EMIC):

$$4\text{Fe}^0 + \text{SO}_4^{2-} + 3\text{HCO}_3^- + 5\text{H}^+ = \text{FeS} + 3\text{FeCO}_3 + 4\text{H}_2\text{O}$$

Chemical microbially influenced corrosion (CMIC):

$$3[\text{CH}_3\text{O}^-] + 2\text{Fe}^0 + 2\text{SO}_4^{2-} + \text{H}^+ = 3\text{HCO}_3^- + 2\text{FeS} + 2\text{H}_2\text{O}$$

$$\text{H}_2\text{S} + \text{Fe}^0 = \text{H}_2 + \text{FeS}$$

In the former (EMIC, Eq. 1) SRB attack iron via withdrawal of electrons directly by metabolic coupling and the formation of an electrically conductive FeS layer. Although this has the potential to greatly increase corrosion rates it has been suggested to be less common, and most proposed mechanisms include the formation of Fe carbonates which have not been observed here. It is also proposed to occur in the absence of organic matter, which observation of blisters and data from XPS suggest is not the case. The initial corrosion of the steel surface by SRBs in the ALWC samples is by the generation of H$_2$S by sulfate reduction, followed by reaction to form FeS,$^{11}$ and by the overall process of organic matter oxidation and sulfate reduction (CMIC, Eqs. 2 and 3). The enhancement of corrosion by direct attack of steel by H$_2$S to generate mackinawite was noted by Little et al.$^{11}$ The production of pyrite is either direct, or as recrystallisation and oxidation of FeS. An observation from this study is the widespread presence of intermediate sulfur oxidation products within ALWC. These could be the result of oxidation of FeS on exposure to the atmosphere. However, all samples were stored in anoxic jars post-collection, and both iron and sulfide-oxidising bacteria have been detected associated with ALWC in this and other studies.$^4$ The formation of reactive intermediate oxidation state sulfur compounds via microbial sulfur oxidation therefore has a key role to play in the development of ALWC. Simple thermochemical oxidation of H$_2$S may play a major role in enhancing corrosion through the generation of low pH.$^{40}$ Direct oxidation of pyrite and FeS may have a similar effect, as may sulfide oxidation in the presence of Fe$^{3+}$ from the outer layers of the blisters.$^{41}$ These reactions are illustrated in Fig. 8. However, pyrite (and by analogy FeS) oxidation has been demonstrated to be a multistage process,$^{41,42}$ with potential microbial involvement throughout.$^{39,43}$ The reaction sequence involved has been summarised by [42] and is shown in relation to S compounds identified in ALWC in Fig. 8. Acidity (H$^+$) is generated throughout this process. It also involves disproportionation, whereby microbial metabolism uses intermediate oxidation state sulfur compounds as both electron acceptors and donators and hence produces both oxidised and reduced sulfur species as products.$^{19}$ Within this process the formation of sulfated iron compounds of varying stoichiometry is further driven by the oxidation, in the presence of seawater, of Fe$^{2+}$ released to

Fig. 7 Results of metagenomic analysis of corrosion products and expression for biochemical pathways involved in sulfur cycling. a Gene environmental samples, Shoreham Harbour, UK, via GeoChip. b Relative abundance of genes responsible for sulfate reduction, sulfite reduction, sulfide oxidation and thiosulfate oxidation.
environmental change. This is also supported by the fact that biochemical reactions. This functional redundancy is important to ensure an ecological stability if any individual species is lost due to the action of sulfate-reducing bacteria, which is protected from the oxic water column via the development of an outer layer of iron oxides and green rusts. Half reactions are given in italics and net reactions in plain text. Every step of the sulfide oxidation is mediated by iron oxidation reaction chain generated electrons and protons that can be consumed by the sulfate reduction process. The development of the full microbial sulfur cycle occurs within corrosion microenvironments, and leads ultimately to the development of ALWC blisters with a layered structure. The emergence of blister structures generates an extremely corrosive microenvironment, conducive to the action of sulfate-reducing bacteria, which is protected from the oxic water column via the development of an outer layer of iron oxides and green rusts.

solution, which in turn contributes to formation of lepidocrocite/goethite. In addition to these processes XPS data indicate the presence of minor amounts of Fe-S cluster type compounds (Fe3S5) that may be indicative of further microbial metabolism. The XPS data also indicate low concentrations of FeS and intermediate oxidation state compounds within normal marine corrosion bioclastic. This indicates a continuum in corrosion types—the presence of anaerobic microenvironments within the splash zone allowing for a minor SRB involvement in normal corrosion generating possible initiation sites for ALWC, or a component of EMIC during periods of steel wetting above the high tide mark. Little et al. noted that abiotic aqueous synthesis of these minerals requires H2S pressures higher than those found in shallow waters. All the reactions identified from spectroscopic data above are partially or wholly microbially mediated. The high number of detected genes (up to 51,748 functional genes), the alpha-diversity results and the number of species found in the samples (32 bacterial phyla and up to 1948 bacterial species) reveal a high bacterial diversity within the corrosion samples studied here. All samples show common species, but closest correspondence is between corrosion samples and samples from 50 cm depth in bed sediment. We conclude from this that the source of corrosion causing bacteria is anoxic environments within bed sediment, with a transport vector via sediment suspension in seawater. Many functional genes from 384 species (belonging to 16 phyla) are involved in the sulfure corrosion showing a high bacterial diversity for this biochemical pathway. The high number of genes having the same function (for instance, 320 and 229 different dsrA and dsrB genes, respectively) and the high number of species carrying one type of functional gene suggest a high functional redundancy, ensured by a mix of species able to perform the same biochemical reactions. This functional redundancy is important to ensure an ecological stability if any individual species is lost due to environmental change. This is also supported by the fact that none of the species are numerically dominant and most species seem to contribute to the biochemical reaction of a gene. Genes involved in the sulfate reduction process were the most detected amongst the sulfur cycle genes, followed by sulfur and sulfide oxidation. A few genes play an important part in the sulfur cycle reactions in sediment and MIC: dsrAB, sqr and soxY. The dsrA and dsrB genes have been previously identified as involved in ALWC, mainly mediating sulfate reduction but also, for some species, sulfide oxidation. The sqr and soxY genes are known to be involved in sulfide and sulfide oxidation. From a phylogenetic point of view, dsrAB genes were mainly found in uncultured bacterial strains (74%), with the remainder derived from 61 species belonging to Proteobacteria and Firmicutes. A recent study on the phylogenetic diversity of dsrAB genes showed that at least 13 lineages gathered uncultured bacteria and 3 of them occurred mainly in marine environment, suggesting that the major taxa of SRB have not yet been identified in this context. The sqr gene was less diversified than dsrAB genes and was found in 30% of uncultured bacteria and in 21 cultivable species. The soxY gene was found almost exclusively in 63 cultivable species, 83% belonging to Proteobacteria. The genes involved in the sulfur oxidation were mainly represented by soxY gene and soxABCV genes were detected with a very low intensity. The oxidation reactions could also be attributed to other species or genes such as dsrAB genes found in Chlorobium sp. Finally, the phylogenetic data showed us that there is an exchange of bacteria between the three matrices (seawater, sediment and MIC) as they share the same species for the key genes involved in ALWC. The overall bacterial involvement in the sulfur cycling linked to ALWC is summarised in Fig. 9. The identification of this range in bacteria highlights the importance of the metagenomic approach using a microarray in characterising natural microbial populations, compared to ex situ culturing studies. The data collected here allow identification of the mechanism of ALWC in this setting. The mechanism of initial colonisation of steel surfaces within the oxic water column and intertidal zone.
remains to be directly determined, although the concentration of defects on steel pile bends has been identified as a factor. The formation of anoxic microenvironments behind macro-biofilms (algae; sea squirts, barnacles) due to the decomposition of organic matter is one possible initiation point, and bacterial adhesion is facilitated by the initial formation of extracellular polymeric substances ('slime') in the biofilm. Minor colonies of SRBs also develop in oxygen depleted pore space within normal marine corrosion, as indicated by the generation of sulfides within more typical corrosion samples. The generation of anoxic microenvironments leads to H₂S generation by sulfate-reducing bacteria against the steel surface. The subsequent oxidation of iron sulfides, via a range of reactive intermediate compounds, generates protons at every stage. This increases the corrosion potential of the process and generates an environment conducive to SRB metabolism. The generation of H₂S by sulfate-reducing bacteria and its mineralisation against the steel surface is a key factor in the enhanced corrosion rate, both as iron is oxidised as an electron donor in electrochemical reactions, and by iron sulfide mineralisation. The generation of an oxidised layer by sulfur- and sulfide-oxidising bacteria protects the SRB colonies from the atmosphere and oxy water column, allowing the process to continue. The ultimate development of high concentrations of H₂S and its oxidation to sulfuric acid results in the die off of the microorganisms in the ALWC process.

The SRB species implicated in ALWC have been identified in samples from 50 cm below the sediment bed surface. This indicates that the marine bed sediment environment is a reservoir for causative species, and sediment resuspension in the marine water column a vector for transport of microbes to the steel surface. Bed sediment anoxia in organic-rich, silt and clay grade sediment is a key component of the electrochemical environment of marine steel, but because of the indigenous microbial populations may also be a fundamental environmental risk factor for ALWC. The identification of ALWC linked microbial populations in the wider environment, and the diverse range of bacteria involved in each metabolic pathway indicates that the phenomenon is emergent as a result of the introduction of an electron donor into the intertidal environment, and not the result of the spread of a single causative organism, or community of organisms, over a period of time.

The range of techniques applied here have allowed the identification of the microbial mechanisms behind ALWC in at a single site. The mineralogical products have also been identified in a number of studies and are consistent with other UK and Australian sites. The presence of intermediate oxidation state sulfur compounds has not been previously reported, although they are implicated as products in sulfide corrosion. A range of SRB and SOB species have previously been reported from ALWC occurrences, and many of the species identified here are widespread in marine sediment. The overall mechanism identified here may therefore be occurring more generally in ALWC, although the precise microbial community is likely to vary from site to site. Detailed spectroscopic and metagenomic analysis of ALWC mineralogy and chemistry has provided the evidence to support an integrated model for its formation for a UK harbour site. Iron sulfides are generated against marine steel surfaces via bacterial sulfate reduction, generating H₂S, which subsequently reacts with steel. The sulfides are oxidised via a range of intermediate Fe and S oxidation state compounds representing reactive intermediates between iron monosulfide and iron oxyhydroxides and sulfate-hydroxides. These are important in the ALWC process as each oxidation step identified has the potential to further generate protons (acidity) that may enhance the steel corrosion process. Metagenomic analysis has identified microbial populations with genes for sulfate reduction, accompanied by sulfide oxidation, sulfite reduction, sulfur assimilation, thiosulfate oxidation and sulfur oxidation within ALWC blisters that are responsible for mediating this sulfide oxidation reaction train. The processes involved in ALWC encapsulate virtually the whole microbial sulfur cycle. The process has emerged as a result of the introduction of an electron donor into the marine environment rather than the spread of a causative organism, and as such is an example of the development of microbial communities in response to environmental change. Further studies are now necessary to confirm these conclusions in other anthropogenically modified marine environments. Identification of causative mechanisms is the first step in improving approaches to address a microbiologically mediated process with global economic impact. From this study both microbial community identification and the use of FTIR to identify diagnostic compounds could support early identification of the problem in the future.

**METHODS**

**Sampling**

Samples for analysis were collected at Shoreham, UK, and immediately stored in sealed anoxic jars in the presence of AnaeroGen™ oxygen-scavenger sachets in order to minimise atmospheric exposure. Samples of corrosion (MIC), bed sediment (SED) and water (WAT) were taken from the “DB”, a tidal location inside the port, and from the “BT”, a non-tidal location inside the port. Additional samples where obtained from dredged sediment in the main harbour channel.

Mineralogical analyses

Samples were imaged and analysed using a Zeiss EVO LS 15 SEM equipped with an Oxford Instruments XMax 80 EDX spectrometer, using an accelerating voltage of 10 kV and a beam current of 707 pA. In order to avoid destruction of microbes and associated structures under vacuum, the samples were observed uncoated at low vacuum. The samples had not been previously dried and the saturation with seawater was sufficient to render the samples conductive and avoid the effects of charge accumulation on the sample surface. In order to further reduce the effects of surface electronic charge the samples were observed in backscattered electron mode rather than using secondary electrons.

Blister samples were analysed by XRD in order to characterise the crystalline mineral phases present. Samples were analysed on a Panalytical X’Pert Pro X-ray diffractometer using 2-9 g hand ground sample mounted unbound in to a deep well powder mount. Samples were irradiated using Cu Ka radiation generated at 40 kV and 40 mA over a 10 mm analytical length, using a rotating stage (in order to maximise random orientation). Diffraction X-rays were counted using 10 s count times at 0.08° intervals with a nickel filter in place on the diffraction beam path between 5 and 70° 2θ. Iron X-ray fluorescence under Cu Ka radiation resulted in high background to X-ray peaks and low relative peak intensities. The pulse height discrimination (PHD) window of the X-ray detector was adjusted to 45–100% in order to gain the best possible signal. Phases were identified from diffraction peaks using the HighScore plus software package and the Powder Data File 2012.

Fourier transform infrared spectroscopic (FTIR) measurements were performed using a Perkin Elmer, Spectrum 65 spectrometer, fitted with an attenuated total reflectance (ATR) accessory employing a ZnSe crystal. Measurements through the ATR accessory require good contact between the sample and the ZnSe crystal. This was achieved by uniform, light, manual compaction. The samples were measured in the spectral range 4000–600 cm⁻¹ at a resolution of 4 cm⁻¹. Each spectrum was collected from 16 scans. The data were analysed using PeakFit (Jandel, Scientific Software) in order to convert data from transmission to absorbance mode, and to allow deconvolution of absorbance peaks. Analysis was focussed in the range 650–1600 cm⁻¹. XPS depth profile measurements were conducted using an ESCALAB 250xi X-ray photoelectron spectrometer (Thermo Scientific, UK) with a
MAGCIS™ Dual Beam Ion Source. XPS analysis was carried out using selected area analysis mode with a nominal width of analysis of 400 µm and monochromated Al Ka X-rays at 1486.6 eV. The MAGCIS source can generate both traditional monatomic ion beams for profiling inorganic materials and also cluster ions for organic layer profiling. In this case of depth profiling corroded iron steel samples, the gun was selected in monatomic mode with 3 keV ion energy and raster size 1.2 mm. Sputtering cycle was kept at 120 s each time with a total of 100 levels for the depth profile analysis. The charge neutraliser and X-ray source were only used during the acquisition of spectra, both being turned off during the sputtering cycle. Survey (wide) scans (step size 1 eV, pass energy 150 eV), only used during the acquisition of spectra, both being turned off during the analysis. The charge neutraliser and X-ray source were only used during the acquisition of spectra, both being turned off during the sputtering cycle. Survey (wide) scans (step size 1 eV, pass energy 150 eV), only used during the acquisition of spectra, both being turned off during the analysis. Width pro

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The normalised data were then analysed using the Microarray pipeline. To call probes positive, we used a floating signal-to-noise ratio (SNR) so that the hyperthermophile probes accounted for 5% of the positive signals. There were cases where the signal was negative if the signal was 1.3 times the background. The signal intensity was then normalised by dividing the signal intensity of each spot by the mean intensity of the microarray. The normalised data were then analysed using the Microarray Data Manager. Alpha-diversity was calculated using Shannon–Wiener index (H), Simpson index (1/D) and Simpson evenness (E). The overlapping genes between all samples were calculated by dividing the number of overlapping genes by the number of genes present in both datasets.}

**DATA AVAILABILITY**

The datasets generated during and/or analysed during the current study are available in summary form in the supplementary material and are available from the corresponding author on reasonable request. Full processed microarray and 16sRNA datasets are available at https://doi.org/10.5061/dryad.52vqt5 hosted by http://datadryad.org/. Raw metagenomic data are available at ArrayExpress (https://www.ebi.ac.uk/arrayexpress/), accession numbers E-MTAB-8364 for GeoChip Microarray data, and E-MTAB-8373 for 16sRNA data.

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