# Increasing flooding frequency alters soil microbial communities and functions under laboratory conditions.

Richard J. Randle-Boggis1\*, Peter D. Ashton1and Thorunn Helgason1

1Department of Biology, University of York, York, YO10 5DD, UK

\* Corresponding author’s e-mail address: r.randle-boggis@lancaster.ac.uk

Running title: Effects of flooding on soil microbial ecosystems.

Keywords: Climate change, DNA sequencing, Flooding, Metagenomics, Microbial ecology.

## Originality-Significance Statement

The authors confirm that all of the reported work is original. To our knowledge, this is the first report showing the impacts of increased flooding frequency on microbial community composition and functional gene abundances. This work contributes significantly to the understanding of how microbial communities and functions will respond to a future climate with more frequent precipitation events.

## Summary

The impacts of increased flooding frequency on soil microbial communities and potential functions, in line with predicted environmental changes, were investigated in a laboratory-controlled environment. More frequent flooding events altered microbial community composition and significantly increased the resolved species alpha-diversity (Shannon index). The Bacteria:Archaea ratio was greater at the end of the experiment than at the start, more-so after only one flood. Significant changes in taxa and functional gene abundances were identified and quantified. These include genes related to the reduction and oxidation of substances associated with anoxia, for example those involved in nitrogen and sulphur cycling. No significant changes were observed in the methanogenesis pathway, another function associated with anoxia and which contributes to the emission of greenhouse gases.

## Introduction

### Climate change and flooding

It is predicted that climatic changes will increase the frequency of extreme precipitation events in places such as Northern Europe, North America and Asia, particularly in winter, and that this will result in an increase in flooding frequency (Min et al., 2011; Collins et al., 2013; Kirtman et al., 2013). This will alter soil microbial ecosystems and biogeochemical cycles (e.g. N, C, Fe and S), at least transiently. Complex microbial communities, such as those found in soil, can be highly responsive to environmental changes (Schmidt *et al.*, 2000; Waldrop and Firestone, 2006; Rinnan *et al.*, 2007).

### Flooding and microbial ecosystems

Anoxic conditions resulting from flooding will affect soil properties and ecosystems (Ponnamperuma, 1984; Stams and Plugge, 2010). Zhou *et al.* (2002) reported that soils saturated in water have reduced bacterial diversities. Microorganisms dominate most biogeochemical cycles, and alterations to community structure and function may result in changes to these cycles. As the frequencies of extreme weather conditions are predicted to increase, it is necessary to understand how these changes will affect ecosystems and their functions.

Some studies research the effects of flooding on microbial ecosystems using targeted approaches. Studying four sites with varying flooding patterns along a river, Bodelier *et al.* (2012) discovered that the abundance of methanotrophs increased with the increase in flooding using denaturing gel gradient electrophoresis (DGGE) and phospholipid fatty acid analysis (PLFA). Kemnitz *et al.* (2004) identified an increase in methanogen diversity in samples from the same river using terminal-restriction fragment length polymorphism (T-RFLP). Unger *et al.* (2009) found that flooding decreased the bacteria:fungi ratio using PLFA. These studies provide a useful insight into the effects of flooding on microbial diversity and community composition; however it is clear that a deeper understanding of the impacts of environmental stressors on the whole community is required. Furthermore, a gene-oriented analysis is required to understand the functional responses to flooding in a pasture field.

Alternating flooding and draining will perturb microbial communities, as the anoxia will kill or supress some populations and allow others to develop (Denef et al., 2001). Cycling between the two states will inhibit the community from stabilising with a predominantly aerobic or anaerobic population, and those that thrive will be able to tolerate both conditions. Flood duration will impact the community as redox potentials take time to decrease during anoxia (Wang *et al.*, 1993; Mohanty *et al.*, 2013), with denitrification occurring, then iron and sulphur reduction, then finally methanogenesis (Reddy and Patrick, 1975; Patrick and Jugsujinda, 1992). Drainage oxidises these compounds again, increasing the redox potential and inhibiting downstream reduction processes. Baldwin and Mitchell (2000) found that nitrification and denitrification decreased after periods of desiccation but increased again after rewetting, and Morillas *et al.* (2015) found that increased dry/wetting frequency decreased nitrification.

Anaerobic soils may contain methanogens, Archaea that produce CH4 under strictly anaerobic conditions, and flooding could increase their populations (Conrad, 2007). Methanotrophs, found both aerobically and anaerobically, metabolise CH4. Methane has a 100-year global warming potential that is 32 times greater than CO2 (Myhre *et al.*, 2013), thus studying the factors that increase CH4 flux is essential for understanding climate change risks. Studies of rice paddies (Yagi *et al.*, 1996; Sigren *et al.*, 1997; Ratering and Conrad, 1998) found that short-term drainage of floods resulted in a sharp decrease in CH4 emissions. This is expected because methanogens are intolerant to even low levels of oxygen (Conrad, 2007). However, once flooding re-occurred, CH4 emissions were still supressed. This may have been caused by the oxidation of reduced sulphate and ferric iron during drainage (Patrick and Jugsujinda, 1992) providing a fresh source of substrates for sulphate/iron reducing bacteria. These would outcompete methanogens for H2 and acetate (Conrad, 2007). How microbial communities will respond to frequent flooding and drainage on pasture soil is yet to be investigated.

While flooding induces anoxia in the bulk soil, the oxic state present during and after drainage may restore the community to its previous state. Ponnamperuma (1984) reported that most of the changes to the physical, chemical and biological processes of soil in response the flooding are reversed with draining and drying, however the rate at which this occurs depends on many factors, such as the proliferation rates of species, redox potentials, the quantities of metabolic substrates present, and the flood subsidence rate. Obligate aerobic and facultative anaerobic bacteria grow best in aerobic conditions, but some can survive periods of hypoxia or anoxia, e.g. *Methylosinus trichosporium* (Roslev and King, 1994) and *Mycobacterium smegmatis* (Berney *et al.*, 2014). Frequent flooding interspersed with drainage periods will therefore only inhibit the growth of many bacteria species, rather than kill them. Furthermore, as a moist environment is preferable for many aerobic bacterial species (Roberson et al., 1993; Potts, 1994; Fredrickson et al., 2008), occasional flooding will provide a suitable environment for these species during drained periods.

### Aims and hypotheses

We investigate the impacts of increased laboratory-controlled flooding frequency on microbial communities and their functions. We hypothesise that increased flooding frequency will significantly change the composition and decrease alpha-diversity of microbial communities and their potential functions. Significant increases in abundances of genes involved in methane production and sulphate reduction are predicted following greater flooding frequencies, with decreases in methane oxidation.

## Results

### Sequencing

8,408,535 paired-end sequences were generated with a mean sample sequence count of 934,300 ± 664,308. PEAR merged 78.98 ± 4.45 % of reads. All samples maintained a mean phred score greater than 30 (Supporting Information Figure 1). The mean sequence length after merging and trimming was 231 ± 131 bases (Supporting Information Figure 2). The rarefaction curves suggest that sequence coverage was sufficient in all samples to represent the microbial community at the genus level; an enhanced sampling effort would yield only a few additional genera (Supporting Information Figure 3). Three x Floods replicate 1 would benefit the most from enhanced sampling

### Diversity and Bacteria:Archaea ratio

There was a significant difference between the order α-diversities of the samples (Start: 4.478 ± 0.010, 1 x Flood: 4.465 ± 0.005, 3 x Floods: 4.492 ± 0.007; ANOVA, F = 8.486, df = 2, p = 0.018). Post-hoc testing revealed that the 3 x Floods treatment was significantly more diverse from than the 1 x Flood treatment (Tukey’s HSD, p = 0.015).

The Bacteria:Archaea ratio significantly increased in response to flooding (square-root transformed (n:1): Start: 12.01 ± 0.15, 1 x Flood: 13.26 ± 0.31, 3 x Floods: 12.74 ± 0.11; ANOVA, F = 26.85, df = 2, p = 0.001; Tukey’s HSD, Start & 1 x Flood: p = 0.001, Start & 3 x Floods, p = 0.012).

### Sample dissimilarities

Flood frequency had a significant effect on the microbial community taxonomic composition (ANOSIM, R: 0.679, *p* = 0.023) and function (ANOSIM, R: 0.251, *p* = 0.003). Both treatments were taxonomically dissimilar from each other and the Start samples (Figures 1 & 2). The 3 x Floods samples were functionally dissimilar from the Start and from 1 x Flood samples, both of which were not dissimilar from each other (Figure 3, Tables 1 and 2).

### Taxonomic and functional abundances

The most abundant phyla across the samples were Proteobacteria, Actinobacteria, Acidobacteria and Verrucomicrobia (Figure 4), which are often dominant phyla (Janssen, 2006).

The relative abundances of 29 orders (out of 223) were significantly different among the treatments, after correcting *p*-values for multiple comparison corrections (ANOVA and Benjamini Hochberg) (Table 3). Most significant differences occur between the 1 x Flood treatment and the 3 x Floods (Tukey’s HSD). There were no significant differences between Subsystems level 3 functions, however at level 2, 14 out of 166 functions were significantly different (Table 4).

Of the 206 orders detected at the start, 66 population relative abundances increased, 122 population relative abundances decreased and 18 populations were undetected after receiving one flood (Supporting Information Table 1). 107 increased, 78 decreased and 21 were undetected after receiving three floods (Supporting Information Table 2). 17 orders were undetected in the starting soil but were detected at the end of the experiment. Supporting Information Tables 3 and 4 show the fold changes between orders (Figure 5). Of the 1,080 level 3 functions detected at the start, 537 relative abundances increased, 471 relative abundances decreased and 46 relative abundances were undetected after receiving one flood (Supporting Information Table 5). 512 level 3 functions increased, 483 decreased and 46 were undetected after receiving three floods (Supporting Information Table 6). 39 level 3 functions were undetected in the starting soil but were detected at the end of the experiment.

### Relative abundance of selected functional groups

The relative abundances of genes involved in methanogenesis and CH4 oxidation were not significantly different (ANOVA, methanogenesis: F = 1.681, df = 2, p = 0.263; CH4 oxidation: F = 2.535, df = 2, p = 0.159). There was a significant difference in the relative abundances of genes involved in Sulphate reduction (ANOVA, F = 11.07, df = 2, p = 0.001, Tukey’s HSD: Start & 3 x Floods: p = 0.014, 1 x Flood & 3 x Flood: p = 0.017) (Figure 6).

## Discussion

### Diversity and Bacteria:Archaea ratio

The order α-diversities were significantly greater in the samples that received three floods than those that received one. While it is hypothesised that anaerobic environments would tend towards a lower α-diversity over time, the short-term repeated shift between aerobic and anaerobic conditions would inhibit a community from stabilising, allowing populations of facultative anaerobic organisms to develop and aerobic populations to recover after flooding. Pett-Ridge and Firestone (2005) and Frindte *et al.* (2015) discuss how redox fluctuations structure microbial communities in tropical soils, and Forth *et al.* (2015) report how anoxic microbial communities shift towards those found in oxic conditions upon oxygenation. Our results may partly be due to community shifts over time, and stochastic shifts are not accounted for. The differences between the treatments, however, indicate the effects of increased flooding frequency.

The Bacteria:Archaea ratio increased in response to flooding, with the greatest ratio observed in the 1 x Flood treatment. The relative abundance of archaea decreased. Archaeal RNA polymerase initiation factors and archaeal thermosomes, involved in transcription and protein structuring respectively, also decreased. Most archaea are either strict anaerobes or can only tolerate low levels of oxygen (Berg *et al.*, 2010), thus the drainage period in between the floods would kill several of the strict anaerobes. Some bacteria, on the other hand, can survive periods of hypoxia or anoxia (Roslev and King, 1994; Berney *et al.*, 2014) and some would thrive in the moist environment provided by the initial flood (Roberson *et al.*, 1993; Potts, 1994; Fredrickson *et al.*, 2008).

### Sample dissimilarities

The community compositions were all dissimilar, revealing that flood frequency has a strong impact on community structure; the 3 x Flooded communities were the most distinct. This was expected, as soil microbial communities can be highly responsive to environmental changes (Schmidt *et al.*, 2000; Waldrop and Firestone, 2006; Rinnan *et al.*, 2007). While the 3 x Flooded communities were functionally different from the other two, the start and 1 x Flood communities were not significantly different. The discrepancy between taxonomic and functional results is most likely due to functional complementarity among taxonomically different microbial communities, a reminder that taxonomic fluctuations do not necessarily imply functional shifts. The 3 x Flood samples are less aerobically stable than the other samples and experience greater perturbation, thus communities will undergo a much greater shift that would likely include significant functional differences.

### Taxonomic and functional shifts

Orders that decreased in response to both treatments include several eukaryotes such as fungi (Capnodiales, Mucorales and Polyporales) and algae (Cyanidiales). As this was a controlled laboratory experiment using homogenised soil, the loss of free organic matter due to consumption may cause the populations of many fungi and algae to decrease. Algae orders Chroococcales and Oscillatoriales both decreased after receiving one flood, but not three. The repeated flooding would limit the effects of desiccation between floods, allowing organisms that prefer moist environments to survive.

Most bacteria involved in the nitrogen cycle that declined in abundance decreased in response to one flood and not to three floods. These include Enterobacteriales, which are largely facultative anaerobes and nitrate reducers (Imhoff, 2005), and Nitrospirales and Nostocales, both aerobic nitrifying bacteria. Furthermore, the relative abundances of RNA polymerase sigma-54 factor RpoN, a function involved in nitrogen assimilation and fixation (Powell *et al.*, 1995; Gardner *et al.*, 2003), and its response regulator both increased in response to three floods but not one, as did nitrosative stress genes. Rhizobiales populations, which include four nitrogen-fixing families, declined in response to both treatments, whereas populations of nitrifying Nitrosomonadales increased in response to both treatments. Initial wetting releases nitrogen that becomes available for nitrification, but after long periods of desiccation several bacteria die off (De Groot and Van Wijck, 1993). As the initial influx of nitrites is assimilated there will be less available for nitrifying bacteria to oxidise. The abundance of genomes containing heterocyst (nitrogen-fixing cells) formation genes in cyanobacteria increased in response to one flood; heterocysts are formed during nitrogen stress, supporting most of our taxonomic findings (with the exception of Nitrosomonadales). Rewetting allows oxidised material to be reduced again, continuing the cycle, and the additional periods of anoxia will permit denitrification (Baldwin and Mitchell, 2000). Verhoeven *et al.* (2014) discovered a decrease in nitrification and denitrification in mangroves after increased flooding frequency, opposing these findings. Nutrient cycling is influenced by a variety of factors, for example: nutrient availability, redox potential, microbial community composition and temperature. Discrepancies between results are therefore expected due to differing conditions, such as those in saline mangroves versus those in terrestrial pasture soils.

We predicted that methanogen and methanotroph populations would increase in response to a greater flooding frequency, but the patterns were complex. The methanotrophic family Methylococcaceae within the Rhizobiales, which are important oxidisers of CH4 in flooded soils (Conrad, 1996) declined in response to both treatments, as did all the other families in this order. Methanotrophic Methylococcales populations, however, did increase as predicted after three floods. Methanotrophic Rhizobiales can survive anaerobic conditions, so it is expected that their populations would also have increased in response to flooding and associated CH4 emissions. One explanation for the relatively weak response of methanotrophs to flooding may be that a corresponding increase in methanogen populations was not observed (and indeed the proportion of archaea also declined, against our original prediction). Therefore, an increase in CH4 production may not have occurred, resulting in little response from methanotroph populations such as Methylococcaceae. In fact, genes involved in the serine-glyoxylate cycle, a part of methylotrophic metabolism (Ensign, 2006), decreased in response to three floods. The lack of developing methanogen populations could be explained by the observed increase in sulphate-reducing bacteria after three floods, as these initially out-compete methanogens for substrates to metabolise (Conrad, 2007).

The greater taxonomic resolution achievable by NGS, compared to other methods such as DGGE and T-RFLP, allows for more detailed understandings of ecosystems to be made. However, the complexity of interactions and responses means that environmental data such as nutrient content and gas fluxes is necessary to make reliable conclusions. To further understand the methanogen/methanotroph results discussed above, sulphur compound content, hydrogen content and CH4 fluxes need to be measured. This would verify the potential functional responses observed in the DNA.

Populations of several strict anaerobic organisms decreased after one flood followed by oxygenation, and many increased in response to three floods. Syntrophobacterales, Chlorobiales, Clostridiales, and Desulfovibrionales are all obligate anaerobes that decreased after the one flood treatment and increased after three floods. Chlorobiales oxidise sulphur compounds, H2 or Fe(II) (Bryant and Frigaard, 2006), and Desulfovibrionales reduce sulphates, thus are important in mineral cycling. Alkanesulfonate assimilation, involved in sulphur assimilation during limited sulphur availability (Ellis, 2011), decreased after three floods; this supports our taxonomic findings. Genes involved in organic sulphur assimilation decreased overall in response to both treatments.

The reduction of Fe(III) during the floods would likely have caused the increase in the Fe(II) oxidising bacteria Gallionellales observed after both treatments, due to the spike in substrate availability (Conrad, 2007). Both treatments resulted in an increase in abundance of genes involved in iron acquisition, transport and metabolism, with Ton and Tol transport systems (iron transport, (Noinaj *et al.*, 2010)) increasing after three floods only. The increase in reduced metals and other substrates due to repeated flooding would explain the increase observed in Cobalt-zinc-cadmium resistance genes and substrate uptake regulation (e.g. Ton and Tol transport systems). These increases were not observed in the one-flood samples, probably due to the resulting oxidation after drainage. Hydrogenase genes, largely involved in anaerobic metabolism (Vignais and Billoud, 2007), also increased after three floods. To further understand these interactions, the behaviour of the microbes needs to be linked to detailed soil chemistry analysis.

Not all anaerobes decreased after the one-flood treatment followed by desiccation; Rhodocyclales, which contains aerobic species but also anaerobic denitrifying oligotrophs (Brenner *et al.*, 2007), increased after both treatments. Fibrobacteres, which include many anaerobic rumen bacteria (Ransom-Jones *et al.*, 2012), increased after three floods but did not change significantly after one flood. While the soil was homogenised for the experiment, localised microbial populations from faeces may have been present.

Other orders that increased after both treatments include Euglyphida, Gemmatimonadetes, and Myxococcales. Euglyphida are amoebae common in soils, marshes and organic-rich environments that feed on bacteria (Lamentowicz *et al.*, 2011). A meta-analysis suggested that Gemmatimonadetes are adapted to arid conditions (DeBruyn *et al.*, 2011), suggesting this result is unexpected. However, Gemmatimonadetes typically make up 2.2 % of soil bacteria (Janssen, 2006), and the only characterised species was isolated from wastewater (Zhang, 2003), thus presence in moist soils is surprising. The increase in Myxococcales hints at one of the current caveats of metagenomics. Myxococcales has an exceptionally long genome (ca. 13 mb) (Schneiker *et al.*, 2007), so for a given number of individuals, sequence read abundance of large genome organisms is likely to be disproportionate and give a skewed impression of community structure. This could be accounted for using the genome sizes of all organisms present, but as yet, this information is not available for complex communities. This issue is exacerbated in eukaryotes, where not only are genomes typically much longer, but the frequency of genes and the functional complexity are not correlated with genome length – the so called C-value paradox (Thomas, 1971).

Planctomycetes, Rhodobacterales and Rhodospirillales decreased after both treatments. These are typically aquatic bacteria, and Rhodospirillales can use sulphide or hydrogen as an electron donor (sulphide is produced by sulphate reducing bacteria typically under anaerobic conditions (Barton, 1995), although they can function aerobically (Kjeldsen *et al.*, 2004; Muyzer and Stams, 2008). We might expect that Planctomycetes, Rhodobacterales and Rhodospirillales populations would increase in response to flooding due to the anoxic conditions and availability of reduced substrates. That this is not the case again suggests that a better understanding of the biology of these microbes is required and the chemical properties of the soil need to be studied throughout these experiments.

Many of the greater fold changes in relative abundances were attributed to mammals and insects, for example: Carnivora, Lagomorpha, Coleoptera, Hemiptera and Phthiraptera. In the case of the mammals, this is likely to be from residual DNA in the soil, such as from rabbit skin. Increases in total abundance of residual DNA in a laboratory experiment are not surprising as, despite homogenisation, spatial variation of relatively larger eukaryotic material is unavoidable; therefore it may not be accurately represented in the results. The soil was sieved and homogenised prior to the experiment, and no insects were observed. While some invertebrates are microscopic, caution should be taken before conclusions are made about these orders as the DNA observed could, like the mammals, be due to residual DNA rather than actual reflections of populations responding to the treatments. Without this critique of results, false conclusions about species’ responses to treatments could be made.

Genes involved in cell growth (RNA polymerase sigma-70 factor), cell signalling (bacterial cAMP signalling) and membrane transport (ABC transporters) decreased after both treatments. The reduction in carbon input due to the removal of plants would restrict growth. Cell signalling is most beneficial when bacterial cell densities are at their highest (Darch *et al.*, 2012), so a reduction in that is expected too as the carbon reduction and water stresses perturb populations. The reduction in membrane transport genes can due to the the sieving, homogenisation and removal of plants reducing the amount of extracellular compounds being available for cell uptake, thus favouring species adapted to relatively lower nutrient environments (than *in situ* pasture soils).

Genes involved in flagellum motility and bacterial chemotaxis increased in response to three floods, but not one flood, suggesting a possible link between flooding frequency and bacterial mobility. Flooding changes the chemical composition of soil, prompting chemotaxis (Bren and Eisenbach, 2000). Transcriptomics would be advantages here to determine which genes are being expressed, rather than just observing which are present. As technology develops, studying mRNA will allow more detailed analysis of functional responses. For example, motility changes may be caused by factors unrelated to mobility that coincidentally change the abundance of organisms, and thus DNA, that utilise flagellae. This complication can be inferred from our data, as the abundance of genes involved in transcription regulation and gene expression appear to decrease in response to either one flood or both treatments. Observing the abundance of mRNA would allow us to determine if gene expression is actually decreasing. Future studies should include analyses of soil chemical properties and microbial biomass to develop a holistic understanding of the microbial ecosystems.

Several genes involved in broader functions, i.e. metabolism, fatty acid metabolism, anaerobic carbon monoxide metabolism, pathogenesis and protection, have varied results, thus broad conclusions cannot be made for these functions. Instead, our results indicate more specific responses to varying flooding frequencies that could be used as a basis for future, more targeted studies.

### Conclusion

In this study we identify some of the impacts that increasing flooding frequency has on microbial communities and their functions. Communities appear to change significantly when they are subjected to additional floods, and functional changes reflect this. Many differences identified relate to the reduction and oxidation of substances associated with anoxia. Changes were not observed in methanogen populations, therefore as long as water drains between floods, an increase in flooding frequency is not expected to increase CH4 emissions (for floods lasting a couple of weeks, at least).

Conducting a laboratory experiment allows variables to be controlled and specific mechanisms tested. To more accurately represent environmental applications, further experiments in the field need to be conducted to investigate the impacts of flooding on *in-situ* communities. Some key advantages of this would be 1) the lack of additional anthropogenic soil disturbance, 2) the inclusion of plants that act as a carbon source (amongst many other things), and 3) the inclusion of diurnal variations in environmental factors such as temperature and light irradiance.

## Experimental Procedures

### Experimental design

Topsoil was collected from within a pasture field in Wiltshire located next to a river confluence (Lat. 51.044770, Long. -2.111945). The soil was collected away from the river and hedges. The soil association is Wickham 2: fine loamy over clayey soil (Supporting Information 1) (National Soil Resources Institute (NSRI), 2013). The mean air temperature for the area is 10.0oC and the mean rainfall is 770.4 mm (Met Office, 2017)

The soil was passed through a 6 mm sieve and stored at room temperature for one week. It was then homogenised and placed in six 8 (h) x 10 (d) cm plastic pots (700 g per pot).

### Treatment

All replicates were subjected to an initial flood for two weeks. The pots were placed in open 1.8 l containers and filled with deionised water to a soil-surface depth of 20 mm. The experiment was conducted in complete darkness at 18oC. After two weeks, all replicates were drained and their GWC brought to field capacity. For the remainder of the experiment, the 1 x flood treatments were not flooded again. The 3 x flood treatments were left drained for two weeks, then subjected to two further two-week flooding treatments, with a two-week period in between and at the end where they were left to drain freely (Table 5).

### DNA sampling

Three randomly selected soil samples (2 g) were extracted from the homogenised soil prior to filling the treatment containers, representing the starting soil community. At the end of the experiment, soil samples were randomly extracted from each container from a depth of 5 cm using a 2 cm corer. DNA was extracted within two hours of sample collection using a PowerSoil® DNA Isolation kit (250 mg) (Mo Bio Laboratories Inc., Carlsbad, CA, USA) following the manufacturer’s protocol and stored at -80 °C.

### Sequencing

DNA quantities were determined using a Qubit® Fluorometer (Life Technologies Corporation, Carlsbad, CA, USA) and the technical extraction replicates were pooled together in equal quantities to form biological replicates. Samples were purified using an Agencourt AMPure XP bead clean-up kit following the manufacture’s protocol (Beckman Coulter (UK) Ltd., High Wycombe, UK). Samples with concentrations greater than 10 ng/μl were diluted 1:10 using RNase-free water to ensure that the quantities were appropriate for use with the Nextera XT DNA sample preparation kit (Illumina UK, Little Chesterfield, UK); concentrations too high result in fragment lengths that are too long for sequencing. The samples were further diluted with RNase-free water to make 5 μl of solution with approximately 10 ng of DNA. DNA libraries were produced using the Nextera XT DNA sample preparation kit following the manufacture’s protocol. The samples were pooled, resulting in a DNA concentration of 17.5 ng/µl. The libraries were sequenced using a MiSeq Personal Sequencer (Illumina UK, Little Chesterfield, UK) following the manufacturer’s protocol. The v3 reagent kit was used, generating paired-end reads of 600 bp.

### Analyses

The paired-end reads were merged with PEAR (Zhang *et al.*, 2014). Unmerged forward reads were trimmed with Sickle (<https://github.com/najoshi/sickle>) using a mean phred score threshold of 25. Unmerged reverse reads were discarded to remove abundance bias when included with merged reads. The merged and trimmed forward read files were concatenated and uploaded to MG-RAST (Table 6) (Meyer *et al.*, 2008).

Sequences were annotated with a representative hit annotation technique, which selects a single, unambiguous annotation for each feature. The RefSeq database was used for taxonomic identification and Subsystems for functional assignment. The maximum E-value was 1e-15, providing a strict search parameter. The minimum sequence identity was 60 %, and the minimum alignment length was 20 bases. These parameters were selected to maximise annotation sensitivity without significantly sacrificing precision (Randle-Boggis *et al.*, 2016). Taxa and functions with a total abundance below five sequences across all samples were removed, as confident conclusions cannot be drawn for such low representations as there is a greater chance that they are influenced by annotation errors (Randle-Boggis *et al.*, 2016). Relative abundance values were generated and arc-sin square root transformed to normalise the proportional data. Rarefaction curves (1,000 resamples) display the taxa richness per sequence count, visualising the effectiveness of sequence coverage. The lowest taxonomic level studied was the order level. Below this we consider the proportion of potential incorrect annotations unacceptable (Randle-Boggis *et al.*, 2016).

The Bacteria:Archaea abundance ratio was calculated and the α-diversity of each sample calculated using the Shannon index, an abundance-weighted average of the logarithm of the relative abundances of taxa. Treatment dissimilarities were tested with Analysis Of Similarity (ANOSIM, 100,000 permutations), Principal Coordinates Analysis (PCoA) and hierarchical clustering, all using the Bray-Curtis dissimilarity method. Taxa and function PCoA weightings were ranked and plotted (Supporting Information Figure 4); those before or after the curve plateaus, at > 0.02 or < -0.02 respectively, were considered for further analysis. Changes in relative abundance of orders and gene functions were analysed using ANOVA. Multiple comparison corrections were made using Benjamini-Hochberg. Significant differences in the abundances of methanogenesis, CH4 oxidation and sulphur reduction genes were selectively tested for using ANOVA.

Data processing was performed in python (Python Software Foundation, v. 2.7.10) and statistical analyses in R (R Core Team, 2012, v. 3.0.3).

## Acknowledgements

We sincerely thank Sandy Macdonald for his technical support and assistance with data analysis techniques, and Phil Ineson for providing advice with experimental design and the use of laboratory equipment.

## References

Baldwin, D.S., and Mitchell, A.M. (2000). The effects of drying and re-flooding on the sediment and soil nutrient dynamics of lowland river–floodplain systems: a synthesis. Regul. Rivers Res. Manag. *16*, 457–467.

Barton, L. (1995). Biotechnology Handbooks: Sulfate-Reducing Bacteria. (New York, USA: Plenum Press).

Berg, I.A., Kockelkorn, D., Ramos-Vera, W.H., Say, R.F., Zarzycki, J., Hügler, M., Alber, B.E., and Fuchs, G. (2010). Autotrophic carbon fixation in archaea. Nat. Rev. Microbiol. *8*, 447–460.

Berney, M., Greening, C., Conrad, R., Jacobs, W.R., and Cook, G.M. (2014). An obligately aerobic soil bacterium activates fermentative hydrogen production to survive reductive stress during hypoxia. Proc. Natl. Acad. Sci. U. S. A. *111*, 11479–11484.

Bodelier, P.L., Bar-Gilissen, M.-J., Meima-Franke, M., and Hordijk, K. (2012). Structural and functional response of methane-consuming microbial communities to different flooding regimes in riparian soils. Ecol. Evol. *2*, 106–127.

Bren, A., and Eisenbach, M. (2000). How Signals Are Heard during Bacterial Chemotaxis: Protein-Protein Interactions in Sensory Signal Propagation. J. Bacteriol. *182*, 6865–6873.

Brenner, D.J., Krieg, N.R., and Staley, J.R. (2007). Bergey’s Manual® of Systematic Bacteriology: Volume 2: The Proteobacteria, Part B: The Gammaproteobacteria (Springer Science & Business Media).

Bryant, D.A., and Frigaard, N.-U. (2006). Prokaryotic photosynthesis and phototrophy illuminated. Trends Microbiol. *14*, 488–496.

Collins, M., Knutti, R., Arblaster, J., Dufresne, J.-L., Fichefet, T., Friedlingstein, P., Gao, X., Gutowski Jr., W.J., Johns, T., Krinner, G., et al. (2013). Long-term Climate Change: Projections, Commitments and Irreversibility. In Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, T. Stocker, D. Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex, and P.M. Midgley, eds. (Cambridge University Press),.

Conrad, R. (1996). Soil microorganisms as controllers of atmospheric trace gases (H2, CO, CH4, OCS, N2O, and NO). Microbiol. Rev. *60*, 609–640.

Conrad, R. (2007). Microbial Ecology of Methanogens and Methanotrophs. In Advances in Agronomy, Donald L. Sparks, ed. (Academic Press), pp. 1–63.

Darch, S.E., West, S.A., Winzer, K., and Diggle, S.P. (2012). Density-dependent fitness benefits in quorum-sensing bacterial populations. Proc. Natl. Acad. Sci. U. S. A. *109*, 8259–8263.

DeBruyn, J.M., Nixon, L.T., Fawaz, M.N., Johnson, A.M., and Radosevich, M. (2011). Global Biogeography and Quantitative Seasonal Dynamics of Gemmatimonadetes in Soil. Appl. Environ. Microbiol. *77*, 6295–6300.

De Groot, C.-J., and Van Wijck, C. (1993). The impact of desiccation of a freshwater marsh (Garcines Nord, Camargue, France) on sediment-water-vegetation interactions. Hydrobiologia *252*, 83–94.

Denef, K., Six, J., Bossuyt, H., Frey, S.D., Elliott, E.T., Merckx, R., and Paustian, K. (2001). Influence of dry–wet cycles on the interrelationship between aggregate, particulate organic matter, and microbial community dynamics. Soil Biol. Biochem. *33*, 1599–1611.

Ellis, H.R. (2011). Mechanism for sulfur acquisition by the alkanesulfonate monooxygenase system. Bioorganic Chem. *39*, 178–184.

Ensign, S.A. (2006). Revisiting the glyoxylate cycle: alternate pathways for microbial acetate assimilation. Mol. Microbiol. *61*, 274–276.

Forth, M., Liljebladh, B., Stigebrandt, A., Hall, P.O.J., and Treusch, A.H. (2015). Effects of ecological engineered oxygenation on the bacterial community structure in an anoxic fjord in western Sweden. ISME J. *9*, 656–669.

Fredrickson, J.K., Li, S.W., Gaidamakova, E.K., Matrosova, V.Y., Zhai, M., Sulloway, H.M., Scholten, J.C., Brown, M.G., Balkwill, D.L., and Daly, M.J. (2008). Protein oxidation: key to bacterial desiccation resistance? ISME J. *2*, 393–403.

Frindte, K., Allgaier, M., Grossart, H.-P., and Eckert, W. (2015). Microbial Response to Experimentally Controlled Redox Transitions at the Sediment Water Interface. PLOS ONE *10*, e0143428.

Gardner, A.M., Gessner, C.R., and Gardner, P.R. (2003). Regulation of the Nitric Oxide Reduction Operon (norRVW) in Escherichia coli Role of NorR and ς54 in the nitric oxide stress response. J. Biol. Chem. *278*, 10081–10086.

Imhoff, J.F. (2005). “Enterobacteriales.” In Bergey’s Manual® of Systematic Bacteriology, D.J. Brenner, N.R. Krieg, J.T. Staley, G.M.G. Sc.D, D.R. Boone, P.D. Vos, M. Goodfellow, F.A. Rainey, and K.-H. Schleifer, eds. (Springer US), pp. 587–850.

Janssen, P.H. (2006). Identifying the Dominant Soil Bacterial Taxa in Libraries of 16S rRNA and 16S rRNA Genes. Appl. Environ. Microbiol. *72*, 1719–1728.

Kemnitz, D., Chin, K.-J., Bodelier, P., and Conrad, R. (2004). Community analysis of methanogenic archaea within a riparian flooding gradient. Environ. Microbiol. *6*, 449–461.

Kirtman, B., Power, S.B., Adedoyin, A.J., Boer, G.J., Bojariu, R., Camilloni, I., Doblas-Reyes, F., Fiore, A.M., Kimoto, M., Meehl, G., et al. (2013). Near-term Climate Change: Projections and Predictability. In Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, T. Stocker, D. Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex, and P.M. Midgley, eds. (Cambridge University Press),.

Kjeldsen, K.U., Joulian, C., and Ingvorsen, K. (2004). Oxygen Tolerance of Sulfate-Reducing Bacteria in Activated Sludge. Environ. Sci. Technol. *38*, 2038–2043.

Lamentowicz, Ł., Gąbka, M., Rusińska, A., Sobczyński, T., Owsianny, P.M., and Lamentowicz, M. (2011). Testate Amoeba (Arcellinida, Euglyphida) Ecology along a Poor-Rich Gradient in Fens of Western Poland. Int. Rev. Hydrobiol. *96*, 356–380.

Met Office (2017) Larkhill Climate Data, *Met Office Regional Climate data.* URL: http://www.metoffice.gov.uk/public/weather/climate/gcneqbvek. Accessed 12/08/2017.

Meyer, F., Paarmann, D., D’Souza, M., Olson, R., Glass, E.M., Kubal, M., Paczian, T., Rodriguez, A., Stevens, R., Wilke, A., et al. (2008). The metagenomics RAST server – a public resource for the automatic phylogenetic and functional analysis of metagenomes. BMC Bioinformatics *9*, 386.

Min, S.-K., Zhang, X., Zwiers, F.W., and Hegerl, G.C. (2011). Human contribution to more-intense precipitation extremes. Nature *470*, 378–381.

Mohanty, S.R., Kollah, B., Sharma, V.K., Singh, A.B., Singh, M., and Rao, A.S. (2013). Methane oxidation and methane driven redox process during sequential reduction of a flooded soil ecosystem. Ann. Microbiol. *64*, 65–74.

Morillas, L., Durán, J., Rodríguez, A., Roales, J., Gallardo, A., Lovett, G.M., and Groffman, P.M. (2015). Nitrogen supply modulates the effect of changes in drying–rewetting frequency on soil C and N cycling and greenhouse gas exchange. Glob. Change Biol. *21*, 3854–3863.

Muyzer, G., and Stams, A.J.M. (2008). The ecology and biotechnology of sulphate-reducing bacteria. Nat. Rev. Microbiol. *6*, 441–454.

Myhre, G., Shindell, D., Bréon, F.-M., Collins, W.D., Fuglestvedt, J., Huang, J., Koch, D., Lamarque, J.-F., Lee, D., Mendoza, B., et al. (2013). Anthropogenic and Natural Radiative Forcing. In Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, T. Stocker, D. Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex, and P.M. Midgley, eds. (Cambridge University Press),.

National Soil Resources Institute (NSRI) (2013). Academic Soils Site Report for location 392413E, 127330N, 1km x 1km (National Soil Resources Institute, Cranfield University).

Noinaj, N., Guillier, M., Barnard, T.J., and Buchanan, S.K. (2010). TonB-dependent transporters: regulation, structure, and function. Annu. Rev. Microbiol. *64*, 43–60.

Patrick, W.H., and Jugsujinda, A. (1992). Sequential Reduction and Oxidation of Inorganic Nitrogen, Manganese, and Iron in Flooded Soil. Soil Sci. Soc. Am. J. *56*, 1071.

Pett-Ridge, J., and Firestone, M.K. (2005). Redox Fluctuation Structures Microbial Communities in a Wet Tropical Soil. Appl. Environ. Microbiol. *71*, 6998–7007.

Ponnamperuma, F.N. (1984). Effects of flooding on soils. In Flooding and Plant Growth, G. Meurant, ed. (Academic Press),.

Potts, M. (1994). Desiccation tolerance. Microbiol. Rev. *58*, 755–805.

Powell, B.S., Court, D.L., Inada, T., Nakamura, Y., Michotey, V., Cui, X., Reizer, A., Saier, M.H., and Reizer, J. (1995). Novel Proteins of the Phosphotransferase System Encoded within the rpoN Operon of Escherichia coli. J. Biol. Chem. *270*, 4822–4839.

Python Software Foundation Python Language Reference, version 2.7.10. Available at http://www.python.org.

Randle-Boggis, R.J., Helgason, T., Sapp, M., and Ashton, P.D. (2016). Evaluating techniques for metagenome annotation using simulated sequence data. FEMS Microbiol. Ecol. *92*.

Ransom-Jones, E., Jones, D.L., McCarthy, A.J., and McDonald, J.E. (2012). The Fibrobacteres: an Important Phylum of Cellulose-Degrading Bacteria. Microb. Ecol. *63*, 267–281.

Ratering, S., and Conrad, R. (1998). Effects of short-term drainage and aeration on the production of methane in submerged rice soil. Glob. Change Biol. *4*, 397–407.

R Core Team (2012). R: A language and environment for statistical computing. R Found. Stat. Comput. Vienna Austria.

Reddy, K.R., and Patrick, W.H. (1975). Effect of alternate aerobic and anaerobic conditions on redox potential, organic matter decomposition and nitrogen loss in a flooded soil. Soil Biol. Biochem. *7*, 87–94.

Rinnan, R., Michelsen, A., Bååth, E., and Jonasson, S. (2007). Fifteen years of climate change manipulations alter soil microbial communities in a subarctic heath ecosystem. Glob. Change Biol. *13*, 28–39.

Roberson, E.B., Chenu, C., and Firestone, M.K. (1993). Microstructural changes in bacterial exopolysaccharides during desiccation. Soil Biol. Biochem. *25*, 1299–1301.

Roslev, P., and King, G.M. (1994). Survival and recovery of methanotrophic bacteria starved under oxic and anoxic conditions. Appl. Environ. Microbiol. *60*, 2602–2608.

Schmidt, I.K., Ruess, L., Bååth, E., Michelsen, A., Ekelund, F., and Jonasson, S. (2000). Long-term manipulation of the microbes and microfauna of two subarctic heaths by addition of fungicide, bactericide, carbon and fertilizer. Soil Biol. Biochem. *32*, 707–720.

Schneiker, S., Perlova, O., Kaiser, O., Gerth, K., Alici, A., Altmeyer, M.O., Bartels, D., Bekel, T., Beyer, S., Bode, E., et al. (2007). Complete genome sequence of the myxobacterium Sorangium cellulosum. Nat. Biotechnol. *25*, 1281–1289.

Sigren, L.K., Lewis, S.T., Fisher, F.M., and Sass, R.L. (1997). Effects of field drainage on soil parameters related to methane production and emission from rice paddies. Glob. Biogeochem. Cycles *11*, 151–162.

Stams, A.J.M., and Plugge, P. (2010). The microbiology of methanogenesis. In Methane and Climate Change, (Earthscan), pp. 14–26.

Thomas, C.A. (1971). The Genetic Organization of Chromosomes. Annu. Rev. Genet. *5*, 237–256.

Unger, I.M., Kennedy, A.C., and Muzika, R.M. (2009). Flooding effects on soil microbial communities. Appl. Soil Ecol. *42*, 1–8.

Verhoeven, J.T.A., Laanbroek, H.J., Rains, M.C., and Whigham, D.F. (2014). Effects of increased summer flooding on nitrogen dynamics in impounded mangroves. J. Environ. Manage. *139*, 217–226.

Vignais, P.M., and Billoud, B. (2007). Occurrence, Classification, and Biological Function of Hydrogenases:  An Overview. Chem. Rev. *107*, 4206–4272.

Waldrop, M.P., and Firestone, M.K. (2006). Response of Microbial Community Composition and Function to Soil Climate Change. Microb. Ecol. *52*, 716–724.

Wang, Z.P., DeLaune, R.D., Patrick, W.H., and Masscheleyn, P.H. (1993). Soil Redox and pH Effects on Methane Production in a Flooded Rice Soil. Soil Sci. Soc. Am. J. *57*, 382.

Yagi, K., Tsuruta, H., Kanda, K., and Minami, K. (1996). Effect of water management on methane emission from a Japanese rice paddy field: Automated methane monitoring. Glob. Biogeochem. Cycles *10*, 255–267.

Zhang, H. (2003). Gemmatimonas aurantiaca gen. nov., sp. nov., a Gram-negative, aerobic, polyphosphate-accumulating micro-organism, the first cultured representative of the new bacterial phylum Gemmatimonadetes phyl. nov. Int. J. Syst. Evol. Microbiol. *53*, 1155–1163.

Zhang, J., Kobert, K., Flouri, T., and Stamatakis, A. (2014). PEAR: a fast and accurate Illumina Paired-End reAd mergeR. Bioinformatics *30*, 614–620.

Zhou, J., Xia, B., Treves, D.S., Wu, L.-Y., Marsh, T.L., O’Neill, R.V., Palumbo, A.V., and Tiedje, J.M. (2002). Spatial and Resource Factors Influencing High Microbial Diversity in Soil. Appl. Environ. Microbiol. *68*, 326–334.

## Table and Figure legends

Table 1. Functions with PCoA weightings > 0.02 or < -0.02 for component 1 in the functional PCoA.

Table 2. Functions with PCoA weightings > 0.02 or < -0.02 for component 2 in the functional PCoA.

Table 3. The orders with significantly different relative abundances between the samples (ANOVA). The *p*-values are adjusted for multiple comparisons (Benjamini-Hochberg) and post-hoc tests were performed (Tukey’s HSD). “*N.S.*” = not significant.

Table 4. The level 2 Subsystems functions with significantly different relative abundances between the samples (ANOVA). The *p*-values are adjusted for multiple comparisons (Benjamini-Hochberg) and post-hoc tests were performed (Tukey’s HSD). “*N.S.*” = not significant.

Table 5. The treatment regime for the laboratory experiment.

Table 6. Sample MG-RAST IDs.

Figure 1. A PCoA (A) and hierarchical clustering analysis (B) of the relative abundance of orders (Bray-Curtis distance method). Ellipses in A display 95 % confidence intervals.

Figure 2. Orders with PCoA weightings > 0.02 or < -0.02 for component 1 (A) and component 2 (B) in the taxonomic PCoA.

Figure 3. A: A PCoA of potential level 3 functions (Bray-Curtis distance method). Ellipses display 95 % confidence intervals. B: A hierarchical clustering analysis of potential level 2 functions (Bray-Curtis distance method).

Figure 4. The relative abundances of phyla in each sample.

Figure 5. The fold changes of orders, coloured by phyla, between the Start and 1 x Flood (A) and the Start and 3 x Floods (B) treatments.

Figure 6. The differences in relative abundances of genes involved in methanogenesis, methane oxidation and sulphur reduction (ANOVA). Error bars show standard deviation.