

Plant, soil and faunal responses to a contrived pH gradient

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Plant, soil and faunal responses to a contrived pH gradient

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Abstract

Purpose To build a more holistic understanding of soil pH change we assessed the synchronised effects of a contrived soil pH change on soil chemistry, vegetation growth and nutrition, and soil faunal abundance and diversity.

Methods We established a fifteen year old field experiment with a contrived pH gradient (pH 4.3 to 6.3) and measured the effect on soil chemistry, plant biomass and elemental composition and the impact of these changes on soil fauna (earthworms, nematodes, rotifers and tardigrades) and biological indices (based on ecological group structures of earthworms and nematodes). A single $20 \times 20 \times 20$ cm soil block was excavated from each sample site to directly attribute biotic parameters in the block to the abiotic (soil) conditions.

Results Acidification affected the extractable concentrations of Al, Ca, Mn and P and the C:N ratio of the soil

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and caused a reduction in plant Ca (r_s for pH vs Ca = 0.804 p < 0.01), an increase in plant Mn ($r_s = -0.450 p = 0.019$), along with significant decrease in root:shoot ratio ($r_s = 0.638$, p < 0.01). There was a significant positive correlation between pH and earthworm index ($r_s = 0.606$, p < 0.01), and a negative correlation between pH and nematode index ($r_s = -0.515$, p < 0.01). *Conclusion* Soil pH influenced the mobility of Ca, Al, Mn and P, which in turn has impacted on plant tissue chemistry and plant biomass ratios. Linked changes in soil chemistry and vegetation had a corresponding effect on the abundance and diversity of nematodes and earthworms in the soil blocks.

 $\label{eq:constraint} \begin{array}{l} \textbf{Keywords} \hspace{0.1cm} \text{Soil acidification} \cdot \text{Sulphur} \cdot \text{Sulfur} \cdot \text{Soil} \\ \text{biology} \cdot \text{Soil biodiversity} \cdot \text{Soil biological indices} \cdot \\ \text{Earthworms} \cdot \text{Nematodes} \cdot \text{Grassland} \cdot \text{Pasture} \cdot \\ \text{Heathland} \end{array}$

Introduction

Soil acidification is a major global problem (Sullivan and Gadd 2019), which contributes to the widespread degradation of soils. A number of natural and anthropogenic processes lead to decreased soil pH, including the mineralisation of organic matter, natural precipitation, nutrient uptake by plants, excretion of organic acids by plants roots and fungi, deposition of acid gases (e.g. acid rain) and the use of acidifying fertilisers, particularly those containing sulphur (S), ammonium and urea (Gadd 1999; Goulding 2016; Rowell and Wild 1984). Ironically, as deposition of sulphuric gases has decreased due to emission controls, the use of S fertilisers has increased to prevent S deficiency in crops (DEFRA 2019). Ecosystems differ in response to acidifying substances with grasslands being among the most sensitive to change (Tian and Niu 2015) and therefore most likely to be responsive to changes in edaphic conditions.

Biological parameters, such as soil organism abundance and diversity, have long been recognised as potentially useful indicators of soil quality (Doran and Zeiss 2000; Ritz et al. 2009). This is largely due to the sensitivity of organisms such as nematodes and earthworms to soil disturbance (Falco et al. 2015). Soil abiotic conditions, including pH, have been reported to impact soil biodiversity and organism distribution (Bardgett and Van Der Putten 2014; Griffiths et al. 2011; Tibbett et al. 2020). Manipulation of soil pH in both arable and grassland systems through liming, for example, has been shown to increase the abundance of bacteria, nematodes and earthworms, and reduce fungal abundance (Holland et al. 2018). Subsequent work, however, suggests that liming had a limited effect on soil microbial and nematode community structure (Neilson et al. 2020) and soil pH is negatively correlated with nematode abundance (van den Hoogen et al. 2019). Soil pH can affect the abundance and diversity of soil organisms through a range of mechanisms: (i) A direct effect on their survival and growth. Some bacterial taxa, for example, have growth tolerances within a narrow pH range. This small niche can lead to competitive exclusion if pH lies beyond the optimum range i.e. broad pH niche bacteria outcompete those with narrow ranges (Rousk et al. 2010); (ii) The modification of their metabolism (Hartman and Richardson 2013), including the activity of microbial enzymes (Neina 2019; Turbé et al. 2010); (iii) The alteration of the bioavailability of both nutrients and contaminants (Turbé et al. 2010). Soil pH affects a number of processes that will influence the solubility, mobility and bioavailability of elements through mineralisation of organic bound elements, chemical adsorption and precipitation reactions (Holland et al. 2018; Neina 2019). An increase in soil pH through liming, for example, can enhance the availability of key nutrients such as N and P, and reduce the availability of elements that can be toxic to microbial growth such as Al, Cu, Fe and Mn (Holland et al. 2018). Finally there are (iv) indirect effects on primary producers and lower trophic organisms. The mechanisms above can affect abundances and diversity of plant communities and microorganisms. Therefore, higher trophic organisms that feed on these will be secondarily affected by the change in resource availability and quality (including elemental composition). Soil pH, and its resultant biogeochemical effects, is commonly considered in relation to microbial communities, with relatively little consideration given to soil fauna by comparison.

Soil organisms live in a heterogeneous ecosystem where small spatial scales are associated with biological trends that are spatially unpredictable (Nielsen et al. 2010), and where random point auger sampling may often neglect key species and hence field-scale experimental responses. Therefore, in order to sample a representative portion of the faunal and plant habitat we excavated soil blocks from a long-term field experiment with a contrived pH gradient. Single $20 \times 20 \times 20$ cm blocks were retrieved from each of the sampling sites, fifteen years after application of acidifying agents, and all analysis is conducted on the sample soil block. We assessed how soil acidification influenced soil chemistry in the block, and how this affected plants and particularly soil faunal responses. The abundance data obtained, therefore, can be directly attributed to the abiotic conditions in which those organisms were found. By using a contrived pH gradient using amendments, on a single soil type, rather than comparing multiple sites across a naturally occurring pH gradient, we limit the confounding effects of edaphic and environmental changes across landscapes (i.e. soil texture, rainfall etc.) driving changes in the soil biology.

We hypothesised that: (i) Soil chemistry would be substantially altered by sulphurous acidifying treatments and that changes would be seen across the pH gradient; (ii) Plant biomass and elemental composition would be significantly different between treatments and across the pH gradient and (iii) Soil fauna abundance (earthworms, nematodes, rotifers and tardigrades) and biological indices (based on earthworm and nematode ecological group structure), would be negatively affected in acidified soils. We tested these hypotheses using a contrived pH gradient of experimental field plots using sulphurous amendments (elemental sulphur and ferrous sulphate) as acidifying treatments on circumneutral grassland soil. The subsequent pH gradient was used to examine the effect of pH on abundance and biological indices based on the nematode and earthworm functional group structure. These two invertebrate groups have been previously identified as potential indicators of soil health (Stork and Eggleton 1992).

Materials and methods

Site description and approach

Thirty 50×50 m experimental plots were established in 1999 as part of a heathland restoration experiment, on the Isle of Purbeck, UK (50.658754 °N, -2.066839 °W). The plots were established on agricultural pasture, created during the 1950s and 1960s through application of rock phosphate, manure and chalk marl for agricultural improvement of the podzolic soil of the native heathland. This application increased the pH and soil nutrients to support the growth of grassland for grazing. Across this improved pasture, plots were amended with either: (i) elemental sulphur, as Brimstone 90 (90% S); (ii) ferrous sulphate as Wet Copperas 50TM (19% Fe and 13% S); or (iii) left as an unamended control (each replicated 10 times). Treatments were applied in 2000 at a rate of 2000 kg ha⁻¹, with an additional 1600 kg ha⁻¹ applied in 2001. See Diaz et al. (2008) and Tibbett et al. (2019) for further details on the experimental design and landscape features.

A single $20 \times 20 \times 20$ cm cube of soil or 'soil block' was excavated from each one of the thirty sampling plots, and all analysis (abiotic and biotic) conducted on each soil block, allowing for ten replicate blocks for each of the three treatments. This way the faunal abundance data obtained from each block can be directly attributed to the abiotic conditions (the habitat) in which those organisms were found (Fig. 1). The position of the block within each plot was random, but at least 2 m from the plot boundary to account for possible boundary effects.

Soil, plant and earthworm sampling

Soil block samples were collected in November 2016. For each plot, the soil block was removed using a flat shovel and placed in trays in the field for hand sorting. Earthworms were carefully removed, counted and placed in a subsample of the same soil to be transported back to the lab for classification. Specimens were rinsed, blotted dry, individually weighed, recorded as juvenile or adult, and classified as epigeic, endogeic or anecic. Adult earthworms were also identified to species level following Sherlock (2012).

Soil from the block was homogenized prior to collecting subsamples for analysis. Soil samples were sieved to 3.35 mm and $\sim 100 \text{ g}$ was stored at $4 \text{ }^\circ\text{C}$ for

nematode extractions. The remainder of the soil was airdried for 5 d, a 10 g subsample of dry soil was then ground to \sim 0.2 mm using a disc mill for C and N determination (detailed below) and the remainder retained for pH and extractable elements analysis (detailed below).

All plant biomass, including roots, were collected in the field from the 20×20 cm surface area of the soil block. Whilst every effort was made to sample all roots, it was not possible to obtain all fine roots. Shoots and roots were separated, washed with water to remove adhered soil, dried to a constant mass at 60 °C, and dry weights recorded. Shoot samples (all aboveground vegetation) were then ground to ~0.2 mm using a disc mill and analysed to give a complete plant community chemical composition.

Nematode, rotifer and tardigrade extractions

Nematodes, rotifers and tardigrades were extracted from ~ 100 g fresh soil with a modified Baermann funnel technique by substituting extraction trays for funnels, with samples collected after 24 and 72 h. Nematodes, rotifers and tardigrades in the extracts were counted while alive on a Leitz Wilovert inverted microscope at 4x magnification. The two sampling times were counted separately and then combined. Identification to feeding group level was completed on 100 nematodes per sample as specified by Yeates (2003) and Yeates et al. (1993). Abundance was expressed as the number of individuals per 100 g soil dry weight equivalent.

Soil analysis

Soil pH for each soil block was measured as a 2.5:1 water-soil slurry after shaking for 15 min at 120 rpm (Rowell 1994).

Extraction of Al, Ca, Cu, Fe, K, Mg, Mn, P and S was conducted on a subsample of each soil block using a Mehlich III extraction (Mehlich 1984). 2.0 ± 0.05 g soil (air dried; < 3.35 mm sieved) was weighed into a 50 mL centrifuge tube with 20.0 mL of Mehlich III extracting solution. This consisted of 0.2 M CH₃COOH; 0.25 M NH₄NO₃; 0.015 M NH₄F; 0.013 M HNO₃; and 0.001 M EDTA adjusted to a pH of 2.50 ± 0.05 . Centrifuge tubes were placed on an end-over-end shaker (40 rpm) for 5 min. After shaking, the suspension was filtered immediately through Whatman No. 42 filter paper. Extractions were analysed for the abovementioned elements



Fig. 1 Schematic of soil block approach and data obtained

using a Perkin Elmer 7300 Dual View inductively coupled plasma - optical emission spectrometer (ICP-OES) (Perkin Elmer Inc., Massachusetts, USA). In house (University of Reading - Sonning Series) reference samples were run alongside with a mean $100.9\% \pm 6.4$ recovery for the elements analysed.

Total C and N content of dried, ground, soil samples from each block were determined using a Thermo Scientific Flash 2000 C/N Analyser (Thermo Fisher Scientific, Massachusetts, USA). Six replicates of an in-house QC material that is traceable to GBW07412 (certified for N by State Bureau of Technical Supervision, The People's Republic of China) and AR-4016 (certified for C by Alpha Resources Inc. with ISO 17025 accreditation) were run alongside samples with recoveries of $100\% \pm 4$ and $99\% \pm 12$ for N and C, respectively.

Plant tissue analysis

Total Al, Ca, Cu, Fe, K, Mg, Mn, P and S in the shoots of the plant community were determined using nitric acid digestion. 0.5 g of ground, milled, dried, plant material was weighed into a MARSXpress (CEM Microwave Technology Ltd., Buckingham, UK), digestion tube with 2 mL ultra-pure water and 8 mL trace element grade concentrated nitric acid. Tubes were left to stand open for 15 min to predigest before being capped and placed into a MARS 6 microwave digestion system (CEM Microwave Technology Ltd., Buckingham, UK). Tubes were ramped to 200 °C for 20 min, held at 200 °C for 10 min, and cooled for 15 min. Digested material was then filtered through Whatman No. 540 filter paper and diluted prior to analysis with Perkin Elmer 7300 Dual View ICP-OES (Perkin Elmer Inc., Massachusetts, USA). Four replicates of IPE 898 Cabbage (*Brassica oleracea*) were run alongside samples (certified by Wageningen Evaluating Programs for Analytical Laboratories, WEPAL), with an average recovery of 98% \pm 11 for the 9 elements measured.

Soil biological indices

Two biological indices were calculated using the functional groups identified for earthworms and nematodes in each block. An index of trophic diversity (ITD) was calculated for nematodes, as described by (Bianchelli et al. 2013). Calculated as:

1-ITD, where ITD = $g_1^2 + g_2^2 + g_n^2$ (1)

g is the relative contribution of each trophic group (bacterial feeder, fungal feeder, plant parasite, omnivore and predator) to the total number of individuals, and n is the number of groups. A higher 1-ITD is indicative of greater diversity.

A Soil Biological Quality Index based on earthworms, or QBS-e (Fusaro et al. 2018), was calculated for each soil block using the abundance of adult and juveniles in each ecological group (epigeic, endogeic and anecic) as follows:

$$\begin{split} \mathsf{QBS-e} &= \left(\mathsf{EMI}_{\mathsf{epi},\mathsf{ad}}*\mathsf{N}\right) + \left(\mathsf{EMI}_{\mathsf{epi},\mathsf{juv}}*\mathsf{N}\right) \\ &+ \left(\mathsf{EMI}_{\mathsf{end},\mathsf{ad}}*\mathsf{N}\right) + \left(\mathsf{EMI}_{\mathsf{end},\mathsf{juv}}*\mathsf{N}\right) \\ &+ \left(\mathsf{EMI}_{\mathsf{ane},\mathsf{ad}}*\mathsf{N}\right) + \left(\mathsf{EMI}_{\mathsf{ane},\mathsf{juv}}*\mathsf{N}\right) \quad (2a) \end{split}$$

Where N is the number of individuals per m^2 of each ecological group. The EMI is the EcoMorphological score attributed to each ecological category, as assigned and described by Fusaro et al. (2018). Resulting in the following:

$$QBS-e = (3*N_{epi,ad}) + (2.5*N_{epi,juv}) + (3.2*N_{end,ad}) + (2.5*N_{end,juv}) + (14.4*N_{ane,ad}) + (10*N_{ane,juv})$$
(2b)

These EMI scores have been designated to each group according to their ecological role, physiological functions and body mass. For example, larger earthworms contribute more in terms of physical function (such as burrowing), resulting in adult worms having larger scores than juvenile ones. Anecic earthworms have the highest score because of their deep-burrowing and large body size, therefore creating 'optimal' soil conditions. Full details of the designation of these scores can be found in Fusaro et al. (2018).

Statistics

We have an experiment that allows an analysis by categorical treatment (i.e. sulphurous amendment) but the experiment also provides a unique opportunity to analyse as a pH gradient. Results presented in the following sections have been explored both as a mean per treatment (tested for equal variance using a Levene's test), and as linear data across the pH gradient (tested for normality using an Anderson-Darling test). All statistical analyses were conducted using Minitab (Version 18.0, Minitab LLC, Pennsylvania, USA). Significance of treatment effects of sulphurous amendments were determined using Kruskal-Wallis one-way ANOVA on ranks, with Bonferroni post-hoc testing. Significance of correlative relationships was investigated using Spearman's rank order correlation.

Multivariate analysis of soil and plant tissue chemistry was also conducted using PRIMER Version 6.0 (PRIMER-e Quest Research Ltd. Auckland, New Zealand). Using the softwares 'normalise pre-treatment' function, all data were normalised before generating a resemblance matrix of Euclidean distances between samples (Clarke and Gorley 2006). Similarities between treatments were assessed through multidimensional scaling (MDS) and analysis of similarities (ANOSIM). All tests for significance were performed at the 95% level.

Results

Soil pH

Application of sulphurous amendments resulted in a pH gradient across the 30 plots, ranging from 4.28 to 6.27 (Fig. 2). Application of elemental sulphur resulted in significantly lower soil pH than control plots, but application of ferrous sulphate did not alter pH significantly from control plots (Fig. 2).

Soil and plant chemistry

Table 1 outlines soil and plant nutrient concentrations as a function of both sulphurous amendment and pH gradient. Soil extractable concentrations of Ca were significantly reduced in the elemental sulphur treatment compared to the control. As with pH, application of ferrous sulphate did not alter extractable concentrations of any of the soil elements measured compared to the control (Table 1). Although a large amount of sulphur was applied to the elemental sulphur and ferrous sulphate treatments, there was no significant difference in S concentrations in the soil 15 years after the final application. Application of ferrous sulphate resulted in higher levels of Fe than the soils of the elemental sulphur treatment, but did not significantly raise the extractable Fe concentration compared to the control. Application Fig. 2 Soil pH in amended soils. Including individual values for each of the 30 plots (•) and mean pH of each treatment ($\land \blacksquare \bullet$) with error bars for standard error (n = 10). Means labelled with the same letter (a, b) indicate treatments that have no significant difference according to Kruskal-Wallis and Bonferroni post-hoc testing (p >0.05)



of ferrous sulphate resulted in significantly lower extractable Cu, compared to elemental sulphur, but not to an extent to be significantly lower than the control. There was a significant increase in soil extractable Ca, Mn and P as pH increases along the gradient, and a significant reduction in soil extractable Al and C:N ratio (Table 1).

Only the Al and Ca concentration of the plant community shoots were significantly different between treatments (Table 1). Extractable Al and Ca was reduced in the elemental sulphur treatment, when compared to the control. The only significant correlations with soil pH observed in the plant community shoots were found for Ca, Mn and S. A significant increase in the plant community shoot Ca and S was seen with increasing pH, and a decrease in shoot Mn (Table 1).

ANOSIM of soil chemical parameters show the elemental sulphur treatment was significantly different to the ferrous sulphate and control treatments (Table 2). Ferrous sulphate soil chemistry did not differ from the control. Elemental composition of in the plant community shoot biomass did not show any significant treatment effect (Table 2).

Plant biomass

Plants growing on plots treated with elemental sulphur had significantly higher shoot biomass than those growing in the ferrous sulphate or the control treated plots (Fig. 3a). There was a significant decrease in root:shoot ratio with acidification (Fig. 3b).

Fauna abundance

Total, adult and juvenile earthworm abundance was significantly lower in the elemental sulphur treatment compared to the control (Table 3). This effect was not shown between ferrous sulphate and control treatments. Endogeic earthworms were the only functional group that showed a significant treatment effect in abundance between the elemental sulphur and control (Table 3).

There was a significant positive correlation between soil pH and total, adult, and juvenile earthworm abundance. This change in total earthworm abundance over the pH gradient appeared to be driven by changes in the endogeic and anecic earthworm population, rather than the epigeic earthworms. Only the total anecic, and total adult and juvenile abundance of endogeic earthworms were significantly correlated to soil pH.

Total earthworm abundance also had significant positive correlations with soil extractable P and Ca, and plant tissue Ca. There was a significant negative correlation between plant tissue Mn and total earthworm abundance. Epigeic earthworm abundance had the fewest significant correlations with soil or plant chemistry of all three earthworm functional groups. Only soil extractable K and Fe showed a significant correlation with epigeic earthworm abundance (Table 3).
 Table 1
 Soil and plant community shoot tissue nutrient concentration as a function of sulphurous amendment and pH gradient.

 Means with the same suffix letter indicate treatments that are not significantly different according to Kruskal-Wallis and Bonferroni

post-hoc testing (p > 0.05). Spearman's rank order correlation of each variable against soil pH are also displayed with p value, all significant correlation (p < 0.05) are displayed in **bold**

	Soil*			Plant **		
Variable	Mean±S.E.		Spearman's rank order correlation (Vs pH)	Mean±S.E.		Spearman's rank order correlation (Vs pH)
Al (mg/kg)	Control Elemental Sulphur Ferrous sulphate	173.9±64.1a 109.3±19.4a 163.2±92.2a	-0.497 <i>p</i> < 0.01	Control Elemental Sulphur Ferrous sulphate	1652.0±521.5a 720.0±277.7b 693.6±201.4ab	0.247 <i>p</i> =0.215
Ca (mg/kg)	Control Elemental Sulphur Ferrous sulphate	962.1±126.5a 570.5±136.7b 922.5±146.5ab	0.639 <i>p</i> < 0.01	Control Elemental Sulphur Ferrous sulphate	5342.8±498.9a 3375.5±446.0b 4955.3±652.0a	0.804 <i>p</i> < 0.01
Cu (mg/kg)	Control Elemental Sulphur Ferrous sulphate	5.1±1.1ab 6.2±0.8a 3.1±0.4b	-0.199 <i>p</i> =0.291	Control Elemental Sulphur Ferrous sulphate	4.4±0.5a 4.7±0.4a 1.1±0.1a	0.129 <i>p</i> =0.523
Fe (mg/kg)	Control Elemental Sulphur Ferrous sulphate	264.0±44.2ab 211.7±24.7b 334.8±47.5a	0.102 p =0.593	Control Elemental Sulphur Ferrous sulphate	1007.7±241.3a 659.8±152.8a 830.0±137.3a	0.123 <i>p</i> =0.540
K (mg/kg)	Control Elemental Sulphur Ferrous sulphate	25.6±3.4a 45.5±10.8a 35.8±4.4a	-0.282 p = 0.132	Control Elemental Sulphur Ferrous sulphate	7208.8±1214.5a 7551.9±590.6a 7152.8±1468.9a	-0.151 p = 0.454
Mg (mg/kg)	Control Elemental Sulphur Ferrous sulphate	54.4±9.4a 48.3±7.0a 70.4±8.0a	0.175 <i>p</i> =0.355	Control Elemental Sulphur Ferrous sulphate	1299.0±123.6a 1160.2±81.3a 1371.4±143.0a	0.232 <i>p</i> =0.244
Mn (mg/kg)	Control Elemental Sulphur Ferrous sulphate	3.1±1.2a 2.9±0.8a 4.4±1.0a	0.413 <i>p</i> = 0.023	Control Elemental Sulphur Ferrous sulphate	257.4±38.8a 351.2±69.0a 229.2±36.2a	-0.450 <i>p</i> =0.019
P (mg/kg)	Control Elemental Sulphur Ferrous sulphate	15.4±1.5a 17.3±3.2a 17.7±2.2a	0.364 <i>p</i> = 0.048	Control Elemental Sulphur Ferrous sulphate	1690.9±259.7a 1770.8±133.5a 2065.0±436.5a	0.144 <i>p</i> =0.474
S (mg/kg)	Control Elemental Sulphur Ferrous sulphate	8.1±1.0a 9.8±1.8a 9.6±1.2a	0.193 <i>p</i> =0.306	Control Elemental Sulphur Ferrous sulphate	1532.9±108.9a 1380.4±71.0a 1522.2±197.5a	0.384 <i>p</i> = 0.048
Total N (%)	Control Elemental Sulphur Ferrous sulphate	$0.2\pm0.0a$ $0.2\pm0.0a$ $0.2\pm0.0a$	0.129 <i>p</i> =0.498			
Total C (%)	Control Elemental Sulphur Ferrous sulphate	3.5±0.3a 3.9±0.4a 4.2±0.4a	-0.002 p = 0.992			
C:N Ratio	Control Elemental Sulphur Ferrous sulphate	19.0±1.2a 20.9±1.7a 18.2±0.9a	-0.412 <i>p</i> =0.024			

*Soil nutrients are Mehlich III extractable concentrations, unless otherwise stated

**Plant tissue nutrients are total concentrations

Table 2 ANOSIM pairwise tests of soil and plant communitytissue chemistry. R and p value generated by ANOSIM conductedon resemblance matrix of data presented in Table 1

	Soil		Plant	
ANOSIM Pairwise Test	R	p value	R	p value
Control vs Elemental Sulphur	0.211	0.023	-0.05	0.750
Control vs Ferrous Sulphate	0.035	0.262	-0.03	0.653
Elemental Sulphur vs Ferrous Sulphate	0.188	0.019	-0.09	0.956

Significant differences (p < 0.05) are displayed in **bold**

Five species of adult earthworm were identified across the soil blocks. These were: Lumbricus rubellus (epigeic); Allolobophora chlorotica (endogeic); Aporrectodea caliginosa (endogeic); Aporrectodea longa (anecic); and Lumbricus terrestris (anecic). The relationship between the abundance of each species and the abiotic soil properties can be found in the supplementary information (Table S1 and S2). There was no treatment effect on species richness or Simpson and Shannon diversity indices for adult earthworms (Table S2) or for any of the species individually, with the exception of acid intolerant Allolobophora chlorotica, which was significantly reduced in the elemental sulphur treatment compared to the control and had a positive correlation with the soil pH gradient (Table S1). No adult species richness or Simpson and Shannon index diversity was significantly correlated to pH (Table S2). Juvenile earthworms were not identified to species level. There was also no significant treatment effect found in multivariate MDS and ANOSIM analysis of adult earthworm species (data not shown).

In contrast to earthworm populations, there were no significant correlations between soil pH and total nematode abundance or the abundance of any nematode functional group (Table 4). However, there was a significant treatment effect, with a significantly higher total abundance of nematodes in the ferrous sulphate treatment compared to the elemental sulphur treatment. This effect was a result of significantly higher numbers of both bacterial and fungal feeders in the ferrous sulphate treatment, compared to the elemental sulphur treatment (Table 4). There were also significantly fewer nematodes in total in the elemental sulphur treatment than the control. However, there was no significant reduction in any particular functional group when elemental sulphur was applied to the control. Significant negative correlations were found between soil Cu concentration and the total bacterial feeding and fungal feeding nematode abundance (Table 4). Shoot biomass (and therefore total biomass) was also significantly negatively correlated with abundance of fungal feeders and omnivores.

Whether abundance is examined as a treatment effect or as a correlation between pH and abundance, the results suggested that pH, as imposed by sulphurous amendments, did not significantly influence rotifer or tardigrade abundance (Table 5).

Biological indices

Table 6 shows the calculated nematode (1-ITD) and earthworm (QBS-e) indices, both of which are based on functional groups rather than species. A significant treatment effect was only observed for QBS-e, with a significantly lower earthworm QBS-e in the elemental sulphur treatment. There was no treatment effect seen for the nematode 1-ITD index.

Both indices significantly correlated with soil pH (Table 6). However, while earthworm QBS-e increased with increasing pH, conversely, the nematode 1-ITD decreased. There was a significant negative correlation between nematode 1-ITD and both total nematode abundance, and bacterial feeder abundance ($r_s = -0.436$ and $r_s = -0.616$ respectively, p < 0.05, data not shown).

Results synthesis

Effect of sulphurous amendment compared to control soil blocks

There was no single variable that differed significantly between the ferrous sulphate and control treatments according to Kruskal-Wallis and Bonferroni post-hoc testing (p > 0.05). Characteristics that were altered as a result of elemental sulphur application are summarised in Fig. 4.

Fig. 3 Effect of pH on total plant biomass **a** Root and shoot biomass according to sulphurous treatment. Error bars for standard error (n = 10). Letters in boxes denote treatments that are not statistically different according to Kruskal-Wallis and Bonferroni post-hoc testing (p > 0.05); **b** Root:Shoot ratio across pH gradient. Spearman's rank order correlation = 0.638, p < 0.01 (n = 30)



Difference between soil treated with different sulphurous amendments

Although there were no differences between ferrous sulphate and the control, there were some significant differences between ferrous sulphate and elemental sulphur treated soils, summarised in Fig. 5.

Effect of pH gradient

Table 7 shows soil chemistry, plant biomass, plant tissue elemental composition and soil fauna parameters that significantly correlated (p < 0.05) with pH.

Discussion

Soil chemistry altered by acidifying treatment

Soil chemistry was significantly altered as a result of acidifying treatment, but only in the case of elemental sulphur. Soil pH reduction in experimental sulphur treated plots has been demonstrated in the past on these plots (Tibbett et al. 2019; Tibbett and Diaz 2005) and elsewhere (Lawson et al. 2004; Owen et al. 1999; Owen and Marrs 2000; van der Bij et al. 2018). This reduction in soil pH led to a subsequent significant reduction in extractable Ca in the soil of the elemental sulphur treatment compared to the control, reiterated by the

s				Earthworm	Abundance							
				Epigeic			Endogeic			Anecic		
	Total Earthworms	Total Adults	Total Juveniles	Adult	Juvenile	Total	Adult	Juvenile	Total	Adult	Juvenile	Total
Mean $(n=10)\pm S.E.$												
Control	8.2±2.4a	$2.1\pm0.7a$	$6.1\!\pm\!1.9a$	$0.1\pm0.1a$	$1.7\pm0.8a$	$1.8{\pm}0.8a$	1.7±0.8a	4.1±1.7a	5.8±2.3a	$0.3\pm0.2a$	$0.3 {\pm} 0.3a$	$0.6\!\pm\!0.4a$
Elemental Sulphur	2.0±0.5b	$0.5 \pm 0.3b$	$1.5 \pm 0.5b$	$0.0{\pm}0.0a$	0.6±0.3a	$0.6\pm0.3a$	$0.2\pm0.2b$	$0.7 {\pm} 0.3b$	$0.9\pm0.4b$	$0.3\pm0.2a$	$0.2\pm0.1a$	$0.5\!\pm\!0.2a$
Ferrous Sulphate	6.3±1.3a	$1.3\pm0.3ab$	5.0±1.2a	$0.1\pm0.1a$	1.4±0.5a	$1.5\pm0.5a$	$0.7\pm0.3ab$	2.8±0.9ab	3.5±1.1a	$0.5\pm0.3a$	0.8±0.4a	$1.3\!\pm\!0.5a$
Spearman's rank coi	relation											
Soil pH	0.616	0.470	0.528	Ι	I	I	0.447	0.477	0.528	I	I	0.375
Soil Al	1	Ι	-0.380	I	I	I	I	-0.394	I	I	-0.504	I
Soil Ca	0.490	0.450	0.378	Ι	I	I	0.548	0.526	0.607	I	I	I
Soil Cu	I	I	I	I	I	Ι	I	I	I	I	I	-0.366
Soil Fe	I	I	I	0.401	I	Ι	I	I	I	I	Ι	Ι
Soil K	I	Ι	I	I	-0.533	-0.532	I	Ι	I	Ι	Ι	Ι
Soil Mg	I	Ι	Ι	Ι	Ι	I	Ι	0.415	0.381	Ι	Ι	0.382
Soil Mn	I	Ι	Ι	Ι	Ι	I	Ι	0.370	Ι	0.419	0.549	0.634
Soil P	0.423	0.464	I	Ι	I	I	0.454	I	0.402	I	I	Ι
Soil C	I	I	I	I	I	Ι	0.335	Ι	Ι	Ι	Ι	Ι
Shoot biomass	I	I	I	Ι	Ι	Ι	Ι	Ι	I	-0.365	Ι	Ι
Plant Tissue Al	I	I	I	Ι	Ι	I	Ι	Ι	Ι	Ι	-0.383	Ι
Plant Tissue Ca	0.529	0.460	0.457	Ι	Ι	Ι	0.588	0.479	0.564	Ι	Ι	Ι
Plant Tissue Cu	I	Ι	Ι	Ι	Ι	I	Ι	0.435	Ι	Ι	I	Ι
Plant Tissue Mg	I	Ι	Ι	Ι	Ι	Ι	0.387	Ι	Ι	Ι	I	Ι
Plant Tissue Mn	-0.448	I	-0.492	Ι	I	I	I	-0.418	-0.420	I	I	Ι
Plant Tissue S	1	Ι	Ι	Ι	Ι	I	0.465	0.354	0.441	I	I	I

Table 4 Relationship between nematode abundance in soil blocks (total in $20 \times 20 \times 20$ cm) and soil and plant parameters. Treatments with the same prefix letter in the same column

indicates no significant difference according to Kruskal-Wallis and Bonferroni post-hoc testing (p > 0.05)

	Nematode Abundance (100 g DWE)					
	Total	Plant Parasite	Bacterial Feeder	Fungal Feeder	Omnivore	Predator
Mean $(n=10)\pm S.E.$						
Control	1434.5±363.3a	572.7±136.6a	715.9±274.0ab	89.2±24.7ab	47.3±16.6a	9.3±5.3a
Elemental Sulphur	705.9±169.2b	300.4±75.1a	297.9±82.4b	61.8±24.7b	36.0±12.8a	9.7±3.4a
Ferrous Sulphate	1546.1±234.5a	494.1±148.6a	857.5±188.7a	127.6±24.2a	39.4±10.2a	27.6±20.2a
Spearman's rank correld	ition					
Soil pH	_	_	_	-	_	-
Soil C	0.391	_	_	0.570	_	-
Soil N	0.389	_	_	0.555	_	-
Soil Cu	-0.513	_	-0.557	-0.382	_	-
Soil Fe	_	_	_	_	_	-0.335
Soil Mg	0.584	_	0.700	_	0.585	-
Soil Mn	_	_	0.503	_	_	-
Plant Tissue Ca	0.399	_	_	_	_	-
Plant Tissue K	_	_	_	_	0.397	_
Plant Tissue Mg	0.650	_	0.700	_	0.635	_
Plant Tissue P	_	_	0.490	_	_	_
Shoot Biomass	_	_	_	-0.516	-0.364	-
Total Plant Biomass	_	_	_	-0.422	-0.374	_
Root:Shoot	0.441	_	_	0.404	_	-

Spearman's rank order correlations were calculated for all plant biomass, plant and soil chemistry variables against nematode abundance but only variables that have a significant correlation (p < 0.05) are shown

significant correlation between pH and extractable Ca. Reduced availability of base cations, such as calcium, with acidification is a result of cation exchange with H⁺ occurring as an acid-buffering mechanism, leading to release of bound cations and eventual leaching from the system (Xu et al. 2020; Zhang et al. 2016).

Elemental sulphur treated soil had significantly lower concentrations of Fe than the ferrous sulphate treated soil, which is to be expected given the Fe content of the ferrous sulphate. The concentration of extractable Cu was significantly lower in the ferrous sulphate treated soils than the elemental sulphur treated soils, despite the lower pH in the elemental sulphur treated soils. A possible explanation for this observation is that, ferrous sulphate application may contain increased Fe oxide or oxyhydroxide concentrations, resulting in increased Cu sorption (Nachtigall et al. 2007).

Univariate analysis of the soil's chemistry may not resolve overall differences in the chemical environment, a key part of the habitat, in which soil organisms and plant reside. While other individual soil chemical parameters did not vary significantly between elemental sulphur and the control, multivariate analysis of soil chemical parameters revealed elemental sulphur treated soil was significantly different to the control and ferrous sulphate treated soil.

Across the pH gradient, regardless of treatment, soil extractable Ca, Al and P were significantly and positively correlated with pH. The positive correlation between extractable Ca and pH, could be a result of H⁺ during acidification competing with Ca for binding sites (discussed above). This could be further perpetuated by a higher Al³⁺ ions, which are more soluble in acidic conditions, competing with base cations for binding sites (Lucas et al. 2011), in turn leading to loss of calcium through leaching.

Acidification is often associated with an increase in extractable Mn (Kidd and Proctor 2001) and a decrease

Table 5 Relationship between rotifer and tardigrade abundance in soil blocks (total in $20 \times 20 \times 20$ cm) and soil and plant parameters

	Total Abundance (100 g DWE)	
	Rotifers	Tardigrades
Mean $(n=10)\pm S.E.$		
Control	12.0±5.4a	1.2±0.8a
Elemental Sulphur	10.8±3.9a	1.2±0.6a
Ferrous Sulphate	8.9±1.8a	1.3±0.6a
Spearman's rank correla	tion	
Soil pH	_	_
Soil Fe	-0.462	_
Soil Mg	_	0.371
Plant Tissue Al	-0.480	_
Plant Tissue Fe	-0.443	_
Plant Tissue K	0.498	_
Root Biomass	_	-0.336

Treatments with the same prefix letter in the same column indicates no significant difference according to Kruskal-Wallis and Bonferroni post-hoc testing (p > 0.05 Spearman's rank order correlations were calculated for all plant biomass, plant and soil chemistry variables against rotifer and tardigrade abundance but only variables that have a significant correlation (p < 0.05) are shown

in available P (Ch'ng et al. 2014). In our research the reverse was found, extractable Mn increased significantly with pH, and extractable P decreased. The increase in extractable P in the low pH samples has been observed in previous studies on this site, and is likely a result of acidification causing a release of P from historical rock phosphate amendments during agricultural improvement (Tibbett et al. 2019; Tibbett and Diaz 2005). The mobilisation of P and Mn in the rhizosphere takes place though a similar mechanism. In a naturally low pH system, low extractable P in the soil promotes plants to release exudates to mobilise P, which results in the coliberation of Mn (Alejandro et al. 2020). This has not happened in our research because our low pH soils, as discussed, have an excess of extractable P. Therefore, the release of exudates would not occur to liberate P and, inadvertently mobilise Mn.

Plant biomass and elemental composition

The elemental sulphur treatment resulted in a higher shoot biomass than ferrous sulphate treatment and the control. Moreover, there was a relationship between pH

Table 6 Relationship between earthworm and nematode diversity index in soil blocks (total in $20 \times 20 \times 20$ cm) soil and plant parameters. Treatments with the same suffix letter in the same column indicates no significant difference according to Kruskal-Wallis and Bonferroni post-hoc testing (p > 0.05)

	Earthworm QBS-e	Nematode 1-ITD
<i>Mean</i> $(n = 10) \pm S.E.$		
Control	689.0±194.0a	0.55±0.03a
Elemental Sulphur	$255.3 \pm 70.0b$	0.56±0.03a
Ferrous Sulphate	706.0±160.0a	0.50±0.04a
Spearman's rank corr	relation	
pH	0.606	-0.515
Soil Al	-	0.563
Soil Ca	0.434	_
Soil Cu	_	_
Soil Mg	0.368	_
Soil Mn	0.495	-0.578
Soil P	_	-0.422
Plant Tissue Ca	_	-0.488
Plant Tissue Mg	-	-0.363
Plant Tissue P	_	-0.522

Spearman's rank order correlations were calculated for all plant biomass, plant and soil chemistry variables against diversity index but only variables that have a significant correlation (p < 0.05) are shown

and plant dry matter partitioning, with roots contributing more to total biomass with increasing pH. Plants species on the soil blocks were not identified, so it is uncertain whether differences in biomass are a result of changes in growth rates of the same plants, or a shift to different plant species that have different growth patterns. However, vegetation surveys on the plots in 2014 revealed a different vegetational assemblage on the elemental sulphur plots, compared to the control and ferrous sulphate plots (Tibbett et al. 2019). Acidity is known to inhibit root development, largely as a result of increased Al solubility (Edwards 1991; Pavlů et al. 2019). Highlighted by the significant reduction in root biomass with increasing extractable Al in our data (Spearman's rank order correlations = -0.407, p < 0.05, data not shown), and extractable Al, in turn, was significantly negatively correlated with soil pH.

Within plant community shoot biomass, only Al and Ca had a significant response to sulphurous amendment treatment. Although there was no treatment effect found for available Al in the soil, the concentrations of Al in shoot tissue from plants in the elemental sulphur treated **Fig. 4** Graphical summary of the significant effects of elemental sulphur application on control soil chemistry, plant biomass, plant tissue elemental composition and soil fauna. Only significant differences between elemental sulphur treated blocks and control blocks (according to Kruskal-Wallis and Bonferroni post host testing (p < 0.05) are shown. There were no significant differences between the control blocks and ferrous sulphate treated blocks and ferrous sulphate treated blocks



soils were reduced compared to the plants from the control. Only plant tissue Ca and S concentration had a positive correlation with soil pH, whilst plant tissue Mn concentration was negatively correlated with soil pH. This negative relationship between plant tissue Mn concentration and soil pH contrasts with the relationship

between soil extractable Mn and soil pH. It has been suggested that soil Mn availability is controlled by neither soil nor plant characteristics but by the combined effects of soil properties, plant characteristics, and the interactions in the rhizosphere (Godo and Reisenauer 1980). Negative effects of elevated extractable P on

Fig. 5 Graphical summary of the significant differences between elemental sulphur and ferrous sulphate treatment on soil chemistry, plant biomass, plant tissue elemental composition and soil fauna. Only significant differences between elemental sulphur treated blocks and ferrous sulphate treated blocks (according to Kruskal-Wallis and Bonferroni post host testing (p < 0.05) are shown. There were no significant differences between the control blocks and ferrous sulphate treated blocks



plant tissue Mn accumulation have been observed in barley and wheat due to P interference directly with Mn at the uptake and/or translocation level (Neilsen et al. 1992; Pedas et al. 2011; Racz and Haluschak 1974). This suggests that the presence of extractable P in this system is impacting both the presence of extractable Mn (discussed above), and the plants ability to absorb it.

Variable	Significant positive correlation with pH	Significant negative correlation with pH
Soil Al		
Soil Ca	\checkmark	
Soil Mn	\checkmark	
Soil P	\checkmark	
Soil C:N		\checkmark
Shoot biomass		\checkmark
Root:Shoot	\checkmark	
Plant Tissue Ca	\checkmark	
Plant Tissue Mn		\checkmark
Plant Tissue S	\checkmark	
Total earthworm abundance	\checkmark	
Total adult earthworm abundance	\checkmark	
Total juvenile earthworm abundance	\checkmark	
Total endogeic earthworm abundance	\checkmark	
Total endogeic adult earthworm abundance	\checkmark	
Total endogeic juvenile earthworm abundance	\checkmark	
Total anecic earthworm abundance	\checkmark	
Earthworm diversity (QBS-e)	\checkmark	
Nematode Diversity (1-ITD)		\checkmark

Table 7 Soil chemistry, plant biomass, plant tissue elemental composition and soil fauna variables that are significantly correlated with pH

Spearman's rank order correlations were calculated for all plant biomass, plant and soil chemistry variables and diversity indices against pH, but only variables that have a significant correlation (p < 0.05) are shown

Soil fauna abundance and biological indices

Reduction in earthworm abundance with decreased soil pH has been well documented (Curry 2004; De Wandeler et al. 2016; Huhta et al. 1986; Phillips et al. 2019; Singh et al. 2016). Total earthworm abundance in our research, was significantly affected by sulphurous amendment, with fewer earthworms found in soil treated with elemental sulphur when compared to the ferrous sulphate treated or control soil. In addition, total abundance of earthworms significantly decreased with acidification. The only other variables that were significantly correlated with total earthworm abundance (extractable P and Ca, and plant tissue Ca) were all significantly affected by soil pH (discussed above) suggesting confounding effects of extractable P, Ca and pH on earthworm abundance. However, only the endogeic and anecic earthworms were affected by sulphurous amendment or the soil pH gradient. Epigeic earthworms feed on plant litter at the soil surface above the mineral layer, whereas anecic and endogeic earthworms burrow deeper into the mineral soil, suggesting they are more susceptible to abiotic soil properties, and less tolerant to a reduction in soil pH (Blouin et al. 2013; Lavelle et al. 1995; Menta 2012; Pulleman et al. 2012; Sizmur et al. 2011). Earthworms are directly sensitive to soil pH, but tolerance varies between species (Springett and Syers 1984). For example, Allolobophora chlorotica (endogeic), the only earthworm species identified to have a significant positive correlation between abundance and pH is known to be intolerant of acidic soil conditions (McCallum et al. 2016). Different earthworm species, or functional groups, being impacted by pH in different ways had led to significant effects on the earthworm community composition. The QBS-e index showed both a significant treatment effect, and a significant positive relationship with soil pH. The elemental sulphur plots, which had an altered soil chemistry, shoot biomass and shoot chemistry, in turn, had a significantly lower QBS-e compared to the ferrous sulphate and control treatments.

As with earthworm abundance, a treatment effect was observed for total nematode abundance with a lower total abundance in the elemental sulphur treatment when compared to the ferrous sulphate treatment and control. Positive correlations between pH and total nematode abundance have often been reported (Holland et al. 2018; Neher et al. 2005), but a significant relationship was not found in our data. As discussed by Neher (2001), unlike earthworms, nematodes are ubiquitous and often persist when other organisms are lost as a result of disturbance or pollution. This may account for why soil pH was significantly correlated with total earthworm abundance, but not nematode abundance.

Biodiversity, in terms of species richness, is not as informative as a soil biological indicators based on the life strategy of key species (Moura and Franzener 2017), which is why we employed the QBS-e and nematode 1-ITD rather than indices based on the number of species present. The use of the QBS-e in the present study appears, in the context of soil pH effect on earthworm communities, to confirm this, as there was no significant relationship between soil pH and adult species diversity. Moreover, not all individual earthworm functional groups were significantly impacted by the acidifying treatments, or the pH gradient. The QBS-e index, which encompassed all groups and maturity, was significantly impacted by both sulphur treatment and the pH gradient, demonstrating the efficacy of this index in indicating change in earthworm communities in response to altered soil parameters.

Contrary to observations for earthworms, nematode abundance was not influenced by the pH gradient. This was the case for total abundance and for the abundance of individual nematode functional groups considered alone. There was, however, a significantly lower total abundance of nematodes in the elemental sulphur treatment compared to both the ferrous sulphate treatment and the control. There were also significantly higher numbers of both bacterial and fungal feeders in the ferrous sulphate treatment compared to the elemental sulphur treatment. This resulted in an unexpected decline in nematode 1-ITD with increasing pH. In all soil treatments, bacterial feeders were among the largest contributors to the total nematode population. Therefore, a decrease in bacterial feeders (if other groups are maintained or reduced to a lesser extent) will raise the relative proportion that the other trophic groups contribute, thereby raising the diversity. Hence as the abundance of bacterial feeders increases, the diversity of nematodes decreases. Bacterial feeder abundance was the only trophic group of nematodes that were significantly correlated with 1-ITD (Spearman's rank order correlations = -0.450, p < 0.05, data not shown).

A significant negative correlation was found between soil Cu concentration and both total nematode and bacterial feeder abundance. Total nematode, bacterial feeder and fungal feeder abundance was also significantly reduced in the elemental sulphur treatment, which had elevated available soil Cu compared to the ferrous sulphate treatment. This may indicate that Cu toxicity contributed to the nematode decline. A negative effect of Cu on soil nematode abundance has been reported (Korthals et al. 2000; Turbé et al. 2010). Whilst Cu concentrations in the sulphur treated soils were well below reported lethal concentration to 50% of nematodes (LC_{50}) values (31–162 mg/kg), there may be sublethal effects (Boyd and Williams 2003).

Nematodes occupy multiple levels of the soil food web (as primary, secondary and/or tertiary consumers). It has been suggested by Turbé et al. (2010) that the abundance of the different trophic groups of nematodes, or community composition, are more sensitive to change than the total abundance. Therefore, analysis of the community structure, at a trophic level, gives an indication of soil disturbance, and could be useful in establishing management targets in unique or ecologically valuable habitats (Bongers and Ferris 1999; Neher 2001; Neher et al. 2005). In addition, the use of functional nematode groups as indicators if soil quality in monitoring has practical benefits, not just for ease of identification, but because the effects of individual soil species on ecosystem processes is often unknown (Neher 2001).

Despite this, adoption of nematode community indices for large scale environmental monitoring has been hindered by a lack of tests across different systems (Neher et al. 2005), particularly as results are often contradictory (Turbé et al. 2010). The present study has demonstrated that nematode diversity at functional group level, i.e. 1-ITD, was able to show that the pH gradient had affected the nematode community in grassland, whereas measurements of abundance of both total nematodes and individual functional groups, could not. Functional group based biological indices are therefore effective tools for monitoring soil quality changes induced by soil acidification.

Conclusions

The use of soil blocks has allowed us to directly attribute changes in soil fauna across a contrived pH gradient to the abiotic conditions in which those organisms were found. Soil pH, altered through application of sulphurous amendements has influenced the mobility of Ca, Al, Mn and P, which in turn has impacted on plant tissue chemistry and plant biomass ratios. These changes in soil chemistry and vegetation have, in turn influenced the abundance and diversity of nematodes and earthworms in the soil blocks.

Biological indices based on the relative abundance of different earthworm and nematode functional groups proved to be sensitive indicators of soil pH, whilst abundance measurements were less sensitive for earthworms and not sensitive in the case of nematodes.

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