





Antibiotic resistance patterns in soils across the Scottish landscape

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The environment disseminates antimicrobial-resistance genes; however, it remains challenging to distinguish whether human activities exacerbate antimicrobial resistance or what is natural. Here, we quantified ~300 resistance-related genes in 200+ Scottish soil samples. Location or land use does not explain gene differences, but nutrient levels reduce gene richness. Elevated levels of metals increased gene richness, and selenium increased transposase levels. Rainfall and persistent organic pollutants also increased transposase relative abundance, possibly promoting conditions conducive to the horizontal transfer of antimicrobial-resistance genes. Selenium and polychlorinated biphenyls were primary factors in gene abundance, while polychlorinated biphenyls, polycyclic aromatic hydrocarbons, and pH influenced gene diversity. Polychlorinated biphenyls are derived from anthropogenic activities, highlighting human activities' potential impact on gene prevalence. This is the first national-scale, high spatial resolution dataset of antimicrobial-resistance genes in Scottish soils and provides a novel resource on which to build future studies.

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Antimicrobial resistance (AMR) threatens the effectiveness of drug therapies to treat bacterial infections. It is predicted to become the leading cause of mortality in the future¹, including economic consequences². It was presumed that AMR was primarily acquired in the clinical environment for years. However, it is now accepted that the acquisition of AMR genes or mutations might also occur elsewhere, including soil, water, wastewater, food and gut^{3–5}.

Analyses of AMR genes in pristine soils^{6,7} demonstrated that the natural environment has an innate level of AMR; however, this environmental reservoir of AMR can be altered through anthropogenic activities that exert environmental pressures on microbial communities. These pressures include biocides, heavy metals and organic pollutants³. However, the relative importance of different human activities—such as clinical and veterinary medicine, agriculture and environmental pollution—in increasing AMR is unknown due to the difficulty in linking direct AMR consequences to specific causative actions. This is because most experimental studies and sites have confounding factors, such as antibiotic residues, toxic organic pollution and potentially toxic elements (PTE; e.g., metals)^{3,8,9} making associated results less conclusive. Furthermore, many datasets cover a limited range of environmental conditions, have limited background information, and quantify a narrow range of AMR gene biomarkers; and examination of specific “human impacted” locations can create a myopic view of AMR and parochial bias. Moreover, there are no agreed international reporting standards or meta-data standards for AMR, meaning that it is usually impossible to produce composite datasets to test hypotheses. As a result, the debate lingers, and we must strive to delineate relative contributions and influences of different factors that correlate with the emergence and propagation of AMR.

Extending the scale and depth of monitoring, including places with no apparent human impact, is critical to distinguish between natural and anthropogenically driven AMR. This will help us better understand background conditions and define factors that can alter the environmental resistome. Here, we quantified the abundance and distribution of AMR genes belonging to eleven antibiotic or mobile genetic element classes across Scotland, including clinically relevant genes associated with vancomycin and ESBL resistance, using a high throughput multi-array qPCR chip. Specifically, over 200 archived soil samples from the National Soils Inventory of Scotland (NSIS2 at James Hutton Institute) were characterised, ranging from low-impacted, relatively pristine soils (such as those in the Highlands) to high-impacted agricultural soils. Therefore, this study represents the first Scottish national-scale, high spatial resolution (20 km grid) assessment of AMR genes in soils. The results should form a foundation towards understanding the impact of human activities and future scenarios, such as climate change, on AMR gene prevalence and whether future interventions effectively reduce environmental AMR.

Results and discussion

AMR genes in Scottish soils. AMR genes were measured in soil samples collected between 2007 and 2009 as part of the re-sampling campaign for the second National Soil Inventory for Scotland (NSIS2). AMR genes were clustered into classes according to antibiotic or genetic element type to facilitate the analysis ($5 < n < 54$ genes in each class). AMR gene distributions and relative abundances found in the Scottish soils are summarised in Table 1, sorted according to antibiotic class. AMR genes conferring resistance to beta-lactam, multidrug resistance and vancomycin were ubiquitous in soils (i.e. 100% of the locations tested contained those resistances). Results for vancomycin

are concerning, given that this is a last-resort antibiotic reserved for treating multidrug-resistant infections¹⁰. Conversely, the distribution of sulphonamide resistance was very variable, with only 46% of the locations containing detectable levels of those genes.

The mean relative abundance and richness of resistance genes belonging to each antibiotic class were determined, respectively. We defined AMR gene richness as the number of AMR genes detected per antibiotic class. Genes conferring multidrug resistance were the most abundant at each location, while sulphonamide resistance was the least abundant. The highest gene richness at each location conferred multidrug resistance (20 genes, 39%) and beta-lactam resistance (13 genes; 24%), while the lowest richness of genes conferred aminoglycoside resistance (6 genes, 13%) and sulphonamide resistance (1 gene, 14%).

The three most frequently detected resistance genes from each antibiotic class or MGE group were also identified (Table 1). In many cases, these were the most abundant resistance genes. Of note is the dominance of clinically important AMR genes such as *bla*_{TEM}, *bla*_{CTX-M}, and *ampC* and the potentially clinically relevant *int11*.

While it is impossible to know which of these resistance genes are expressed, this dataset shows the reservoir of resistance genes in soil that could be horizontally transferred or expressed by resistant microbes under certain conditions. The genes represent three broad mechanisms of resistance: efflux pumps ($n = 88$ genes), antibiotic deactivation ($n = 115$ genes) and cellular protection ($n = 71$ genes), while a fourth group includes mobile genetic elements and resistance genes with unknown function ($n = 24$ genes)¹¹. The average number of genes per sample conferring these different resistance mechanisms was found across four broad categories of land use (arable, semi-natural, managed grassland and woodland/forestry) (Fig. 1a). This showed that more genes conferring efflux pumps were detected, followed by antibiotic deactivation, then cellular protection, with others being the least detected. There were no significant differences in the number of genes according to land use ($p > 0.05$).

The mean relative abundance of genes conferring the different resistance mechanisms was also found across the four land use categories (Fig. 1b). This showed that efflux pumps were the most abundant, followed by antibiotic deactivation, then others, with cellular protection being the least abundant. There were significant differences according to land use, with arable land having significantly lower abundances of genes for efflux pumps and other genes than other land uses ($p < 0.05$). This is surprising given that genes with specific resistance mechanisms are enriched in farm environments.¹¹

AMR genes are ubiquitous but uneven. Mapping the total relative abundances and richness of AMR genes shows that AMR genes are ubiquitous in Scottish soils (see Fig. 2), which is unsurprising given that AMR is a natural phenomenon¹². Furthermore, AMR genes have been found in all environmental matrices, including in relatively pristine environments⁶. Almost all gene classes displayed no significant relationships with geographical features regarding gene richness or abundance (i.e., northing, easting, latitude or longitude; Spearman correlations, $p > 0.01$). Furthermore, there were no statistical differences when split among Scottish councils (i.e., 32 local-authority regions; Supplementary Tables 1–2; *K-W* test, $p > 0.01$), consistent with Spearman’s rank results.

However, the data mapping showed localised hot spots of some AMR genes (and some not), influencing relative abundances and richness patterns (Figs. 3, 4, Supplementary Figs. 1–16). For

Table 1 Summary of the distribution and relative abundance of AMR genes according to antibiotic class in Scottish soils.

Antibiotic class	The proportion of locations with resistance detected (%)	Mean gene frequency at each location	Mean log relative abundance at each location (std. dev. in parenthesis)	Top genes	
				Presence	Abundance
Aminoglycoside	89	6 (13%)	-6.37 (1.01)	<i>aacC</i> , <i>spcN</i> , <i>aadA9</i>	<i>aacC</i> , <i>aacC4</i> , <i>spcN</i>
Beta-lactam	100	13 (24%)	-5.16 (0.49)	<i>bla</i> _{CTX-M-04} , <i>ampC</i> , <i>cphA</i>	<i>cphA</i> , <i>fox5</i> , <i>bla</i> _{TEM}
FCA	94	2 (22%)	-6.14 (0.96)	<i>cmxA</i> , <i>floR</i> , <i>cmlA1</i>	<i>cmxA</i> , <i>floR</i> , <i>cmlA1</i>
MLSB	100	9 (20%)	-5.81 (0.49)	<i>mphA</i> , <i>pikR2</i> , <i>erm36</i>	<i>mphA</i> , <i>pikR2</i> , <i>erm36</i>
Multidrug	100	20 (39%)	-4.85 (0.36)	<i>tolC</i> , <i>oprD</i>	<i>mexF</i> , <i>oprD</i> , <i>oprJ</i>
Sulfonamide	46	1 (14%)	-7.35 (0.50)	<i>sul2</i> , <i>sulA-folP</i> , <i>sul1</i>	<i>sulA-folP</i> , <i>sul2</i> , <i>sul1</i>
Tetracycline	98	7 (18%)	-6.30 (0.54)	<i>tetG</i> , <i>tetR</i>	<i>tetB</i> , <i>tetR</i> , <i>tetG</i>
Vancomycin	100	7 (22%)	-5.76 (0.51)	<i>vanC</i> , <i>vanHB</i> , <i>vanTC</i>	<i>vanC</i> , <i>vanXD</i> , <i>vanHB</i>
Other	94	2 (18%)	-6.11 (0.67)	<i>pncA</i> , <i>bacA</i>	<i>pncA</i> , <i>bacA</i>
<i>Mobile genetic elements</i>					
Integrons	94	1 (20%)	-5.21 (0.56)	<i>intI_1clinic</i> , <i>intI3</i> , <i>intI_1LC</i>	<i>intI_1clinic</i> , <i>intI3</i> , <i>intI_1LC</i>
Transposons	97	3 (38%)	-6.85 (0.62)	<i>tnpA</i>	<i>tnpA</i>

example, the relative abundance and richness of vancomycin genes predominate in NE Scotland (Fig. 3a, b), though the correlations were weak ($r_s = 0.20$, $p < 0.01$ for both). Conversely, other antibiotic resistance classes, such as beta-lactam genes, show no significant spatial variation in relative abundance and richness as this was not significant ($p > 0.01$) (Fig. 3c, d).

Despite spatial heterogeneity in some AMR gene classes, there were no significant differences among land-use patterns (e.g., arable, grassland, etc.; Supplementary Tables 3, 4; $K-W$, $p > 0.01$). Since the NSIS soil archive encompasses a range of soils from highly impacted to relatively pristine soils, it was surprising that land use did not significantly influence endemic AMR genes, especially since several studies have shown that impacted land (e.g. from agriculture¹³) can increase relative abundances of AMR genes.

Here, land-use data were limited at the time of sampling, i.e., no historical information was associated with the sites, even though past anthropogenic activities have been shown to leave legacy pollutants that continue influencing AMR genes in the environment¹⁴. Furthermore, some relatively pristine environments could be impacted by agriculture, e.g. peatlands can be used for grazing. These factors may confound detecting AMR gene patterns relative to land use.

Least squares regression analysis was undertaken to explore the potential influence of land use fully. Data were log-transformed, and categorical variables were dummy-coded as appropriate. The analysis showed no significant associations between AMR gene abundance or richness with any of the land use types recorded as part of the original soil survey (17 different categories, dummy coded) or when these were aggregated into four major classes (arable, managed grassland, semi-natural, woodland/forestry). This suggests that other factors not measured in the original survey might be more explanatory or that the categorisation used while suitable for the original purpose of the survey is less appropriate when it comes to explaining spatial variation in AMR gene richness/abundance (i.e., traditional land use categories might not adequately reflect impacts that promote AMR). This

may be reflected in the fact that specific indicators of anthropogenic impact (carbon, nitrogen, pH, persistent organic pollutants, etc.) did show significant correlations with AMR gene richness/abundance (see below).

Rainfall increases transposon relative abundance. Climate change has been predicted to affect AMR gene prevalence, given that higher temperatures and moisture can accelerate bacterial growth rates and increase horizontal gene transfer, possibly promoting increased environmental transmission¹⁵. Therefore, we looked for potential correlations between temperature and rainfall. The temperature was not significantly correlated with either AMR gene relative abundance or richness, possibly because temperature ranges within Scotland are too limited to see a correlation. However, rainfall significantly correlated with transposon relative abundance (Table 2) but not transposon richness. While this is consistent with a previous study that showed that rainfall can be associated with AMR gene transmission and proliferation from air to soil¹⁶, it may also reflect the additional influence of non-point pollution in run-off from high rainfall. We cannot discern the difference from the archived soil samples.

Factors affecting bacterial richness also affect AMR genes. Here, the biggest contributor to AMR gene richness was bacterial abundance. Significant positive correlations existed between bacterial abundance and the resistance gene richness for aminoglycoside, beta-lactam, am-/chlor-/fluoro-quinolones, multidrug resistance, sulphonamide, tetracycline and transposase genes. We also measured bacterial richness using the Chao Index. There were significant positive correlations between bacterial richness and the richness of resistance genes for beta-lactam and sulfonamide ($0.20 < r_s < 0.31$, $p < 0.01$). In other words, the greater the bacterial abundance and richness, the greater the AMR gene richness.

Therefore, the environmental factors often correlated with bacterial communities (Supplementary Table 5) were also

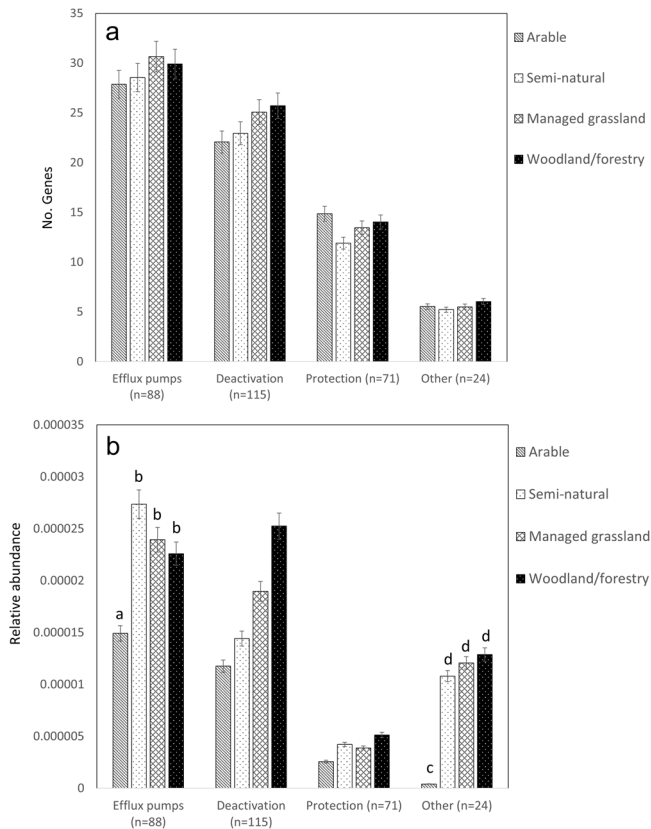


Fig. 1 Groups of resistance mechanisms across four broad land use types.

Resistance mechanisms (efflux pumps, antibiotic ‘deactivation,’ cellular ‘protection’ and ‘other’ denoting mobile genetic elements and genes of unknown function) are displayed according to (a) the number of genes detected or (b) relative abundance of genes across four broad land use types (arable, semi-natural, managed grassland and woodland/forestry). Values of n refer to the number of genes belonging to that group of resistance mechanisms assayed in the soil samples. The error bars represent standard errors of the mean, and the letters above the bars show significant differences, where $p < 0.05$.

examined with AMR genes (Tables 2 and 3). There were no consistent correlations between carbon, nitrogen and pH with AMR gene abundance (Table 2); however, these factors often correlate with AMR gene richness (Table 3). Carbon was negatively correlated with bacterial abundance and richness (Supplementary Table 5; $r_s = -0.34$ and $r_s = -0.78$, respectively, both $p < 0.01$) and, therefore, negatively correlated with the richness of the AMR genes for beta-lactam, multidrug resistance, sulfonamide, tetracycline and vancomycin.

Similar correlations were observed with LOI (loss on ignition; $r_s = -0.36$ and $r_s = -0.75$, $p < 0.01$), which is unsurprising since LOI also indicates the amount of organic carbon in soils. Nitrogen was also negatively correlated with bacterial abundance and richness (Supplementary Table 5; $r_s = -0.19$ and $r_s = -0.58$, respectively, $p < 0.01$) and sulfonamide gene richness. On the other hand, soil pH was positively correlated with bacterial abundance and richness (Supplementary Table 5; $r_s = 0.33$ and $r_s = 0.84$, respectively, $p < 0.01$) and positively correlated with the richness of the AMR genes associated with beta-lactams and sulphonamides.

While we cannot explicitly assign how much carbon and nitrogen came from human activity versus natural sources, our results suggest that human activities that alter concentrations of crucial nutrients in the environment, such as those in fertilisers,

can influence AMR gene richness. Nutrient availability impacts bacterial abundance and richness, so it is reasonable to expect that these factors also affect AMR genes. Nitrogen application to soil can decrease microbial biomass¹⁷, reducing the pool of potential hosts for AMR genes. Increased assimilable carbon can diminish soil biodiversity¹⁸, whereas in water, it can decrease microbial richness as eutrophication tends to select for fewer dominant, typically opportunistic species¹⁹. This selection process potentially explains why AMR richness declines with increased nutrient loadings in our study.

Our results are consistent with other studies that found soil carbon significantly influenced microbial community composition and AMR gene prevalence. For example, Li et al.²⁰ showed that higher total organic carbon led to lower AMR gene abundance in Tibetan soils, which was attributed to higher autochthonous microbial abundances that out-compete and prevent invasions of antibiotic-resistant bacteria. In a different context, higher carbon concentrations in a biofilm from a drinking water sand filter also decreased bacterial diversity and AMR gene richness, attributed to a few species’ dominance at higher nutrient concentrations¹⁹. An alternative explanation is the reversion of resistant bacteria to susceptible forms (i.e. plasmid curing) without stress conditions, especially when the carriage of resistance genes comes at a fitness cost²¹.

Our results also showed that higher soil pH is associated with increased bacterial abundance and richness (Supplementary Table 5) and AMR gene richness (Table 3). Many studies have shown soil pH to be the overarching factor influencing microbial community composition^{22–25}, so it is reasonable that it might also affect AMR gene prevalence. For example, one study on rice paddy soils showed that changes in microbial community structure were strongly influenced by pH, which consequently influenced local AMR gene spread²⁶. The positive correlation between soil pH and AMR gene richness found in this study is contrary to other studies on soil AMR^{27,28}; however, soils in those studies encompassed a broader pH range, whereas Scottish soils are mostly acidic (the average was pH 4.7 and the range was pH 3.2–7.9 within our sample set).

Metals influence AMR gene richness. Metals can influence AMR prevalence and selection; however, their bioavailability remains the critical factor in the relationship²⁹. Despite this, we looked for correlations between total metal concentrations and AMR genes in the soils studied here. Surprisingly, Spearman’s rank correlation showed that metal concentrations were largely negatively correlated with AMR gene relative abundances. Specifically, negative correlations were observed between copper, strontium, mercury, nickel, zinc, boron, magnesium, manganese and chromium and the relative abundance of resistance genes conferring multidrug resistance, vancomycin and transposons. Only selenium was positively correlated with transposase relative abundance (Table 2). The predominantly negative correlations with metals may result from antimicrobial activity and fewer hosts for AMR genes³⁰. Moreover, metal bioavailability and the abundance and types of anions consequentially alter the influence of heavy metals on microbial communities³¹, including relative AMR gene abundances²⁹, which might also explain some of the negative correlations.

Conversely, elevated total metal concentrations were associated with increased AMR gene richness. Specifically, the richness of resistance genes for beta-lactam, sulphonamide, tetracycline, and vancomycin increased with increasing concentrations of manganese, barium and aluminium (Table 3). Therefore, higher metal concentrations for the soils in this study increased AMR gene

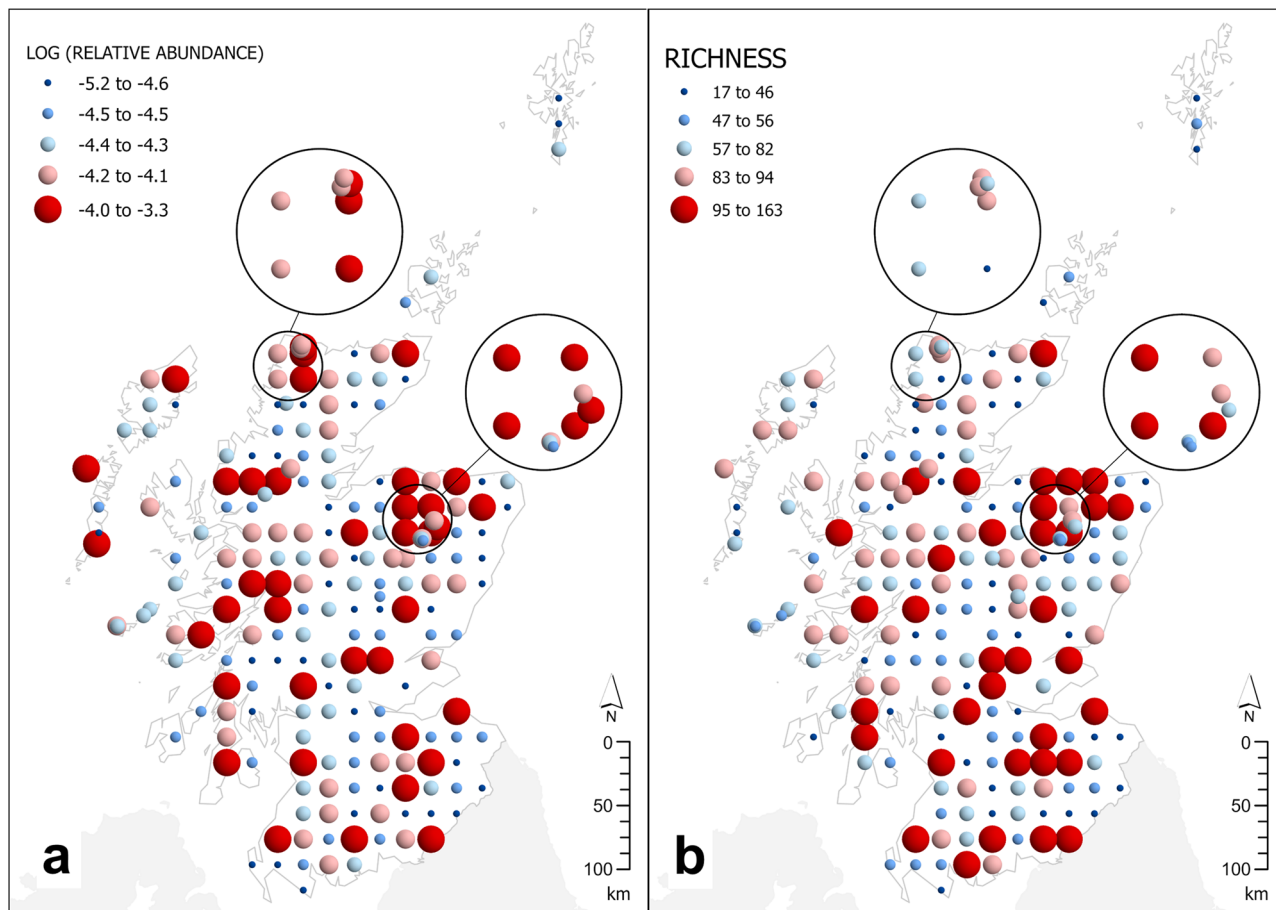


Fig. 2 Total antimicrobial resistance genes. Total AMR genes are denoted in terms of (a) relative abundance and (b) richness across Scotland using quintiles.

richness but did not necessarily increase overall AMR gene abundance (apart from the case with selenium).

Several studies have shown that metals enhance AMR genes and mobile genetic elements through co-resistance (inheritance of different resistance determinants on the same genetic element) and cross-resistance (the same resistance determinant confers resistance to metal and antibiotic)^{8,32}. For example, resistance to both barium and aluminium was prevalent in antibiotic-resistant bacterial isolates from a polluted river in Turkey³³. Furthermore, Robins et al.²⁹ discovered that mobile genetic elements in NE England were associated with AMR genes in rural settings, whereas in urban samples, the AMR genes directly correlated with metal pollutants.

In agricultural soils, the metal composition tends to be more closely governed by the parent material³⁴, i.e. from natural sources. Analysis of historical samples from the National Soils Archive of Scotland (1940–1970, representing the early antibiotic era) showed a baseline level of AMR gene relative abundance that correlated with a combination of baseline levels of copper, chromium, nickel, lead and iron³⁵. Since this new study focussed on rural locations, probable anthropogenic sources of metals in these soils include fossil fuel combustion, organic material application, fertilisers and atmospheric deposition³⁴. Although it is impossible to determine how many specific metals are from natural or anthropogenic sources here, the results suggest that local metals influence AMR gene richness; therefore, reducing anthropogenic metals to Scottish landscapes will help decrease AMR gene richness in the soils.

Persistent organic pollutants drive transposase abundance.

There has been increasing interest in the potential for persistent organic pollutants (POPs) to drive AMR^{9,36,37}. The concentrations of several POPs, including polychlorinated biphenyls (PCBs), polycyclic aromatic hydrocarbons (PAHs) and polybrominated diphenyl ethers (PBDEs) were previously measured in the NSIS soils³⁸ so these were used to determine any possible relationships with AMR genes. There are no known natural sources of PCBs or PBDEs, and they are an explicit measure of environmental anthropogenic pollution.

Spearman's rank showed that there were significant positive correlations between the relative abundance (but not richness) of transposase genes and the PCBs, 2,3',4,4',5-pentachlorobiphenyl, 2,2',3,4,4',5'-hexachlorobiphenyl, 2,2',4,4',5,5'-hexachlorobiphenyl and 2,2',3,4,4',5,5'-heptachlorobiphenyl, and the PAH fluorene (Tables 2 and 3). On the other hand, significant negative correlations were observed between the richness of resistance genes for beta-lactam, sulfonamide, tetracycline and vancomycin with the PCBs 2,2',4,5,5'-pentachlorobiphenyl, 2,3',4,4',5-pentachlorobiphenyl, 2,2',3,4,4',5'-hexachlorobiphenyl, 2,2',4,4',5,5'-hexachlorobiphenyl and 2,2',3,4,4',5,5'-heptachlorobiphenyl and the PAH chrysene (Tables 2 and 3). Only one correlation with PBDEs was observed with AMR genes, where the relative abundance of tetracycline was negatively correlated with 2,2',4,4', and 6-pentabromo-diphenyl ether.

Studies have shown that catabolic genes responsible for the degradation of organic compounds are closely associated with mobile genetic elements, such as transposons, which promote

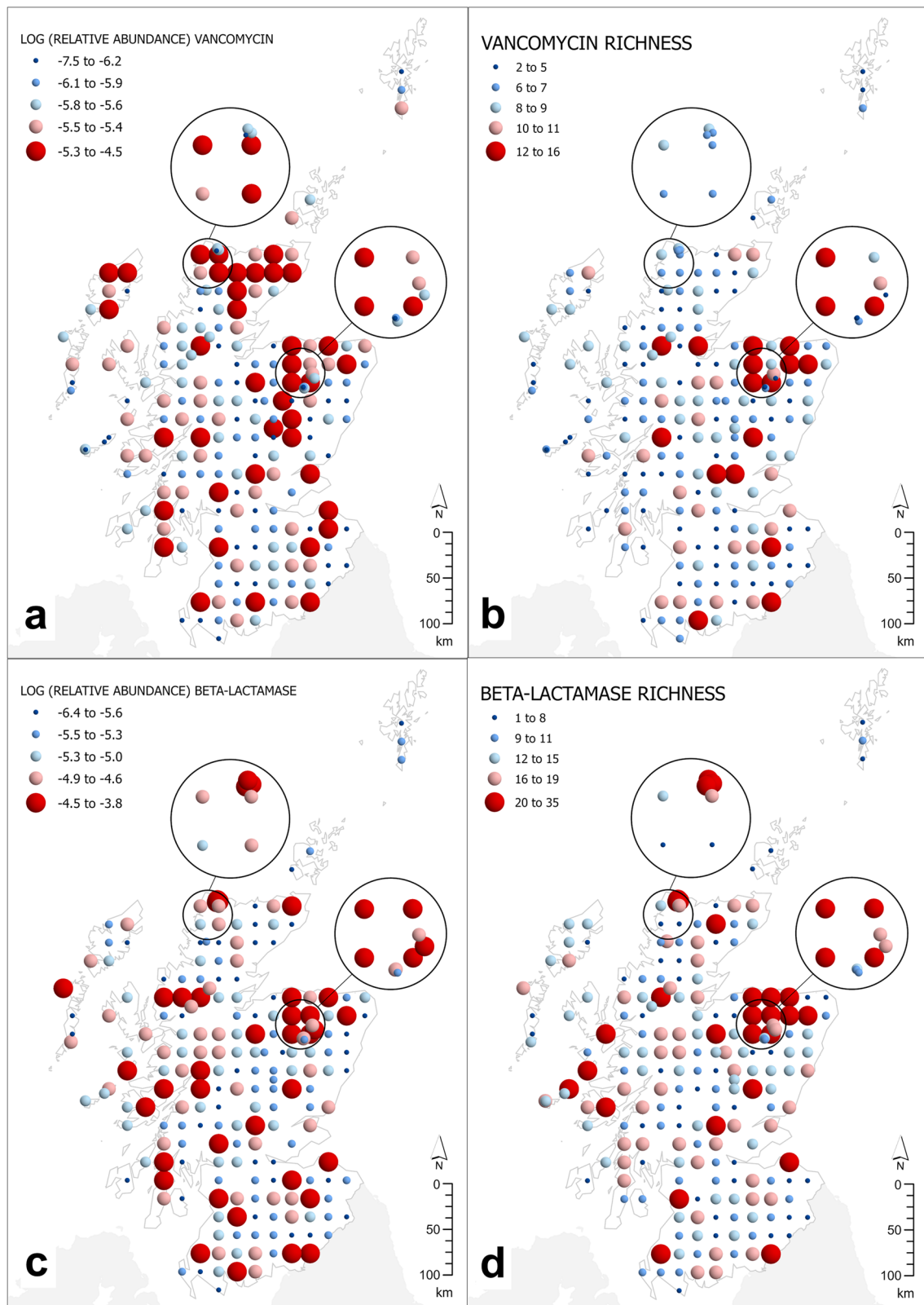


Fig. 3 Relative AMR gene abundances and richness for selected antibiotic classes. Levels of AMR genes are denoted for (a) vancomycin relative abundance, (b) vancomycin richness, (c) beta-lactam relative abundance, and (d) beta-lactam richness across Scotland using quintiles.

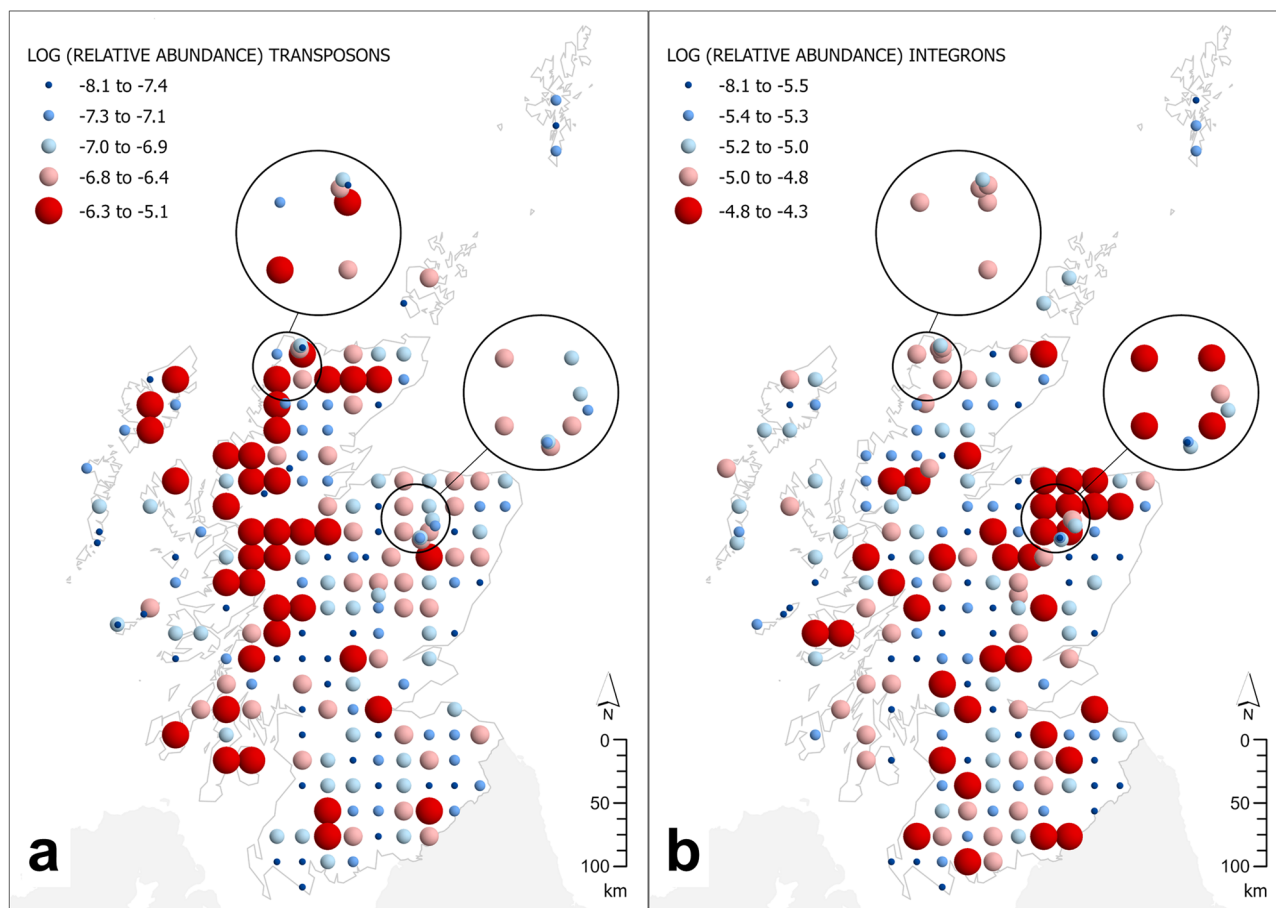


Fig. 4 Mobile genetic element gene relative abundances. Levels of MGE are denoted for (a) transposon relative abundance and (b) integron relative abundance across Scotland using quintiles.

their spread and de novo construction of catabolic pathways, allowing bacteria to adapt to such pollutants³⁹. Our results corroborate transposons' possible role in the bioremediation of organic compounds in the environment⁴⁰. The increase in transposase abundance driven by PCBs would be expected to promote AMR transmission. However, the apparent decrease in AMR gene richness with PCBs and PAHs possibly results from toxicity to host cells⁴¹. These confounding effects have unclear consequences for AMR in the environment. As noted by others⁹, there is a gap in the literature on the possible enrichment of AMR genes in PCB-degrading bacteria in soil. As such, further work is needed to help elucidate the effects of PCBs on AMR genes.

Transposon-mediated co-selection of PAH-degrading genes has been noted previously⁴², which may explain higher transposon relative abundances with increased fluorene concentrations. Field studies have shown that PAH-contaminated environments can select for AMR genes^{37,43}, possibly mediated by horizontal gene transfer³⁶, while in vitro studies have shown that PAHs can block plasmid-mediated transfer of AMR genes⁴⁴. Again, these confounding effects have unclear consequences for AMR in the environment, so further studies are required to determine the effects of PAHs on AMR genes.

A principal components analysis (PCA) determined the relative associations between environmental parameters (climate, pH, carbon/LOI, nitrogen, metals/metalloids and POPs) and AMR gene mean relative abundance and richness. Overall, the plots show no clustering according to land use (Fig. 5), which agrees with our correlation and regression analysis. There was also no clustering according to location from the correlation

analysis. For AMR relative abundance, the axes explain 71.6% of the variation and are most influenced by selenium, followed by PCBs. For AMR richness, the axes explain 79.5% of the variation and are most influenced by PCBs and PAHs, followed by soil pH. This indicates that human activities are one of the strongest drivers of AMR abundance and richness since PCBs can only be derived from human activity.

Integrons and transposons drive AMR gene richness and abundance. Since integrons and transposons can be vectors for the AMR gene to spread^{45,46}, they were used as explanatory variables to determine whether they correlated with AMR gene richness and abundance (Tables 2, 3). Interestingly, the relative spatial distribution of transposons versus integrons is quite different across Scottish soils (Fig. 4a, b). Transposon relative abundances are much higher in west Scotland than east, whereas integron relative abundances are more evenly distributed across the country.

Relative integron and transposon abundances were positively correlated with the relative abundances of most AMR gene classes (Table 2). The exception was for AMR genes conferring resistance to beta-lactams and am-/chlor-/fluoro-quinolones, where correlations with relative transposon levels were not significant ($p > 0.01$). Overall, integron relative abundance had the strongest positive correlation with AMR gene relative abundance compared to the correlation between transposon relative abundance and AMR gene relative abundance ($r_s = 0.34$ – 0.66 for integrons vs $r_s = 0.19$ – 0.34 for transposons), indicating that they may have a greater influence on possible AMR gene transfer.

Table 2 Significant correlations between AMR gene relative abundance and soil biological and chemical characteristics.

Antibiotic Class	Climate	Bacteria (16S rRNA)	pH and nutrients	Metals	PAH	PCB	PBDE	Mobile genetic elements
Aminoglycoside								Int (0.56), Tn (0.29) Int (0.57)
Beta-lactam		Abundance (0.19)	pH (0.20) LOI (-0.20) C (-0.18)					
FCA MLSB								Int (0.34) Int (0.69), Tn (0.24)
Multidrug				Cu (-0.19) Sr (0.20)				
Sulfonamide								
Tetracycline							2,2',4,4',6-pentabromo-diphenyl ether (-0.21)	Int (0.48), Tn (0.19) Int (0.66), Tn (0.2)
Vancomycin		Abundance (-0.28) Richness (-0.20)	pH (-0.23) C (0.19)	Cr (-0.29) Hg (-0.21) Ni (-0.18) Mg (-0.20)				Int (0.63), Tn (0.34)
Other								Int (0.51), Tn (0.22)
Mobile genetic elements								NA NA
Integrons								
Transposons	Rainfall (0.20)		pH (-0.21) N (0.20)	Hg (-0.26) Ni (-0.24) Se (0.19) Zn (-0.19) B (-0.20) Mg (-0.27) Mn (-0.28)	Fluorene (0.24)	2,3',4,4',5'-Pentachlorobiphenyl (0.23) 2,2',3,4,4',5'-hexachlorobiphenyl (0.20) 2,2',4,4',5,5'-hexachlorobiphenyl (0.24) 2,2',3,4,4',5,5'-heptachlorobiphenyl (0.28)		

Values of r_s are in brackets. In all cases, $p < 0.01$.

Table 3 Significant correlations between AMR gene richness and biological and chemical characteristics of the soil.

Antibiotic Class	Bacteria (16S rRNA)	pH and nutrients	Metals	PAH	PCB	Mobile genetic elements
Aminoglycoside	Abundance (0.23)					Int (0.39), Tn (0.53)
Beta-lactam	Abundance (0.31) Richness (0.21)	pH (0.21) LOI (-0.25) C (-0.23)	Mn (0.22)		2,2',3,4,4',5'-hexachlorobiphenyl (-0.20) 2,2',4,4',5,5'-hexachlorobiphenyl (-0.20) 2,2',3,4,4',5,5'-heptachlorobiphenyl (-0.20)	Int (0.37), Tn (0.49)
FCA MLSB	Abundance (0.24)					Int (0.37), Tn (0.46)
Multidrug	Abundance (0.23)	LOI (-0.21) C (-0.20)				Int (0.43), Tn (0.43)
Sulfonamide	Abundance (0.22) Richness (0.20)	pH (0.19) LOI (-0.21) C (-0.21)	Mn (0.20)		2,2',4,4',5,5'-hexachlorobiphenyl (-0.20)	Int (0.42), Tn (0.39)
Tetracycline	Abundance (0.27)	N (-0.19) LOI (-0.23) C (-0.24)	Ba (0.19)	Chrysene (-0.19)	2,2',4,5,5'-pentachlorobiphenyl (-0.26) 2,3',4,4',5'-Pentachlorobiphenyl (-0.25) 2,2',3,4,4',5'-hexachlorobiphenyl (-0.28) 2,2',4,4',5,5'-hexachlorobiphenyl (-0.29) 2,2',3,4,4',5,5'-heptachlorobiphenyl (-0.26)	Int (0.34), Tn (0.46)
Vancomycin		LOI (-0.23) C (-0.24)	Ba (0.22) Al (0.19)		2,2',4,5,5'-pentachlorobiphenyl (-0.25) 2,3',4,4',5'-Pentachlorobiphenyl (-0.23) 2,2',3,4,4',5'-hexachlorobiphenyl (-0.28) 2,2',4,4',5,5'-hexachlorobiphenyl (-0.29) 2,2',3,4,4',5,5'-heptachlorobiphenyl (-0.23)	Int (0.41), Tn (0.43)
Other						Int (0.31), Tn (0.36)
Mobile genetic elements						NA
Integrons						NA
Transposons	Abundance (0.22)					NA

No significant correlations were found with PBDEs. Values of r_s are in brackets. In all cases, $p < 0.01$.

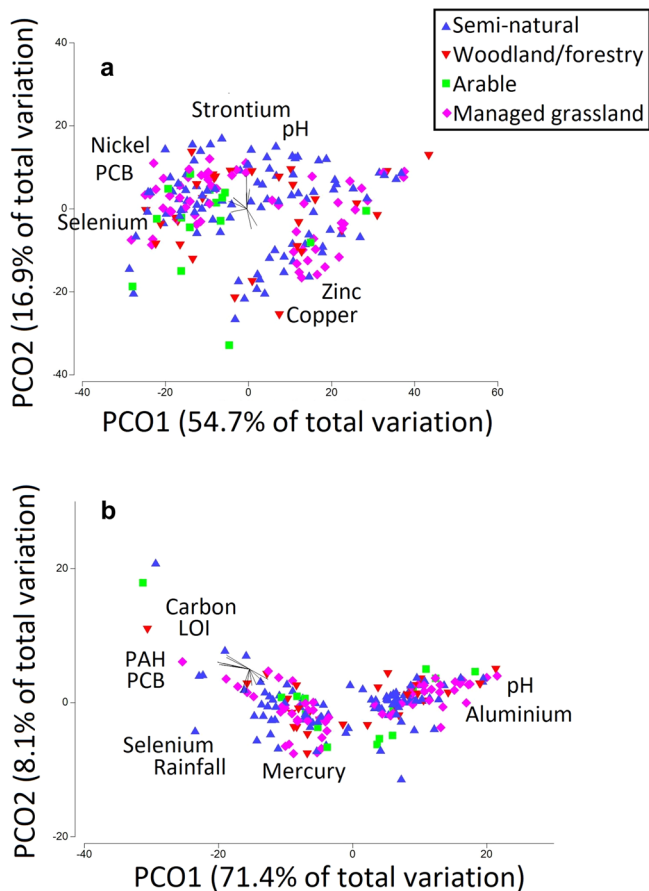


Fig. 5 Principal components analysis showing environmental-factor influences. The influence of environmental factors on (a) AMR gene abundance and (b) AMR gene richness. Colours represent different land-use conditions: semi-natural (blue triangles), woodland/forestry (red triangles), arable (green squares), and managed grasslands (pink diamonds).

For AMR gene richness (Table 3), integron and transposon richness appeared closely associated with the richness of most AMR gene classes. Only the richness of genes conferring resistance to am-/chlor-/fluoro-quinolones was not correlated with either integrons or transposons. Numerous studies have demonstrated that the abundance of the *int1* gene increases with pollution and human impact⁴⁷; therefore, such associations might be expected. Equally, transposons can increase with pollution^{29,32}. Overall, such associations between mobile genetic elements (MGEs) and AMR genes suggest an elevated potential risk of AMR gene transmission across Scottish soils via horizontal and vertical gene transfer.

Practical implications. This study shows that “intrinsic” soil resistomes in Scotland are significantly influenced by environmental factors, including climate (rainfall), soil pH, nutrients (carbon, nitrogen), metals (selenium, manganese, barium, and aluminium) and POPs (PAHs, PCBs, PBDEs), and that increased MGEs strongly associate with relative AMR gene abundances and diversity. The strongest associations with relative AMR gene abundances were with ambient selenium and PCB levels. By contrast, PCB, PAH, and soil pH conditions substantially influenced relative AMR gene richness in soil. This suggests that in situ resistance potential has been altered by pollutants from anthropogenic activities, as PCBs can only be derived from human activity. Rainfall, selenium, fluorene and PCBs positively

correlated with transposase relative abundance, which may increase the potential for AMR gene transfer. Increased AMR gene richness potentially increases the number of different mechanisms bacteria can use to evade the antimicrobial effects of antibiotics. On the other hand, increased AMR gene relative abundances increase potential human exposure via different possible soil- and water-related exposure pathways. However, it remains to be seen whether increased AMR gene richness or abundance indeed presents a greater risk to human health, either immediate or latent.

The uneven distribution of genes belonging to each antibiotic class across Scotland raises questions of when and how the AMR genes emerged in local hot spots, how the AMR genes were propagated, and why the AMR genes persisted more in some soils than others. Care needs to be taken over the interpretation of our data. Since the soil is highly heterogeneous, the results cannot necessarily be extrapolated to soils near our sampling points. Nevertheless, the extensive soil sample series shows that some geographical regions in Scotland may have a greater opportunity for environmental AMR genes to move into strains of greater health importance. This work further confirms that targeted studies using historical soil samples can help elucidate mechanisms that impact the emergence and evolution of environmental AMR. This is very important given the greater emphasis at the United Nations on broader integrated surveillance as an essential element in understanding AMR prevalence and spread⁵. This valuable national-scale dataset describes background conditions across the country, which allows the identification of locations for additional surveillance and, in turn, permits the development of mitigation strategies to control the spread of AMR.

Materials and methods

Soil sampling. Soil samples were collected as part of a re-sampling campaign for the second National Soil Inventory of Scotland (NSIS2, 2007–9)⁴⁸, based on standardised methods developed for the first National Soil Inventory (NSIS1). This survey recorded the sampling sites’ location (northing, easting, longitude, latitude) and altitude⁴⁸. The collection contains representative and unbiased soil samples from 195 sites across Scotland, based on a 20 km grid sampling pattern, plus 21 rare soils off the 20 km grid pattern, giving a complete archive of 216 soil samples. Briefly, soil pits were excavated at each site, and the soil horizons were identified by visual inspection. Approximately 1.5 to 2 kg of stone-free soil was collected from the middle of each horizon, placed in bags, and sealed immediately to avoid contamination. Meteorological data (temperature and rainfall) was recorded from the Met Office⁴⁹.

Once returned to the James Hutton Institute, a sub-sample was taken for microbiological analysis; only the top horizon was considered for this study. A second sub-sample was frozen at -80°C until required, and the remaining soil was air-dried and sieved (2 mm) for chemical analyses and subsequent storage in the National Soils Archive (www.hutton.ac.uk/about/facilities/national-soils-archive). Persistent Organic Pollutants (POPs), including polychlorinated biphenyls (PCBs), polycyclic aromatic hydrocarbons (PAHs) and polybrominated diphenyl ethers (PBDEs), were determined as described in Zhang et al.²⁸ and Cui et al.³⁸. Soil pH was determined on the supernatant of a 3:1 water to the soil mix.

Total carbon and nitrogen concentrations (C%, N%) were determined using a Flash EA 1112 Series Elemental Analyser (Thermo Fisher, Germany). Loss on Ignition (LOI) was measured gravimetrically. The soil was dried at 105°C until a constant mass was achieved, then weighed into pre-weighed silica or porcelain crucibles. The sample was ignited at 450°C and 900°C for a

minimum of 2 h. For metals and metalloids (arsenic, barium, cadmium, cobalt, chromium, copper, mercury, molybdenum, nickel, lead, selenium, strontium, zinc, aluminium, boron, calcium, iron, magnesium and manganese), soils were milled to a fine powder and digested in a mixture of hydrochloric acid and nitric acid (*aqua regia*) under reflux for two hours. The digest was cooled, filtered and made up to a known volume with 0.5 M nitric acid before being analysed by inductively coupled plasma optical emission spectroscopy (ICP-OES) or Inductively Coupled plasma-mass spectrometry (ICP-MS) to determine element concentration.

DNA extraction and quantification of AMR genes. DNA was extracted from triplicate samples of 250 mg of soil using a modified CTAB method, which were pooled prior to analysis⁵⁰. Modifications to the CTAB method included carrying all steps on ice rather than at room temperature (including the lysis step), using higher centrifugation speeds for more extended periods, using Maxtract High-density Tubes (Qiagen) to separate the DNA from the phenol-chloroform, and carrying out two ethanol wash steps to remove impurities. The purity and concentrations of the extracts were verified by UV spectrometry (Nanodrop and QuBit, respectively; both ThermoFisher). The DNA were normalised to 10 ng/μL and stored long-term at -80 °C. 100 μL (1 μg) of DNA was analysed using the Applied Biosystems OpenArray platform.

All high-throughput quantitative PCR (HT-qPCR) reactions were performed in triplicate using the Applied Biosystems OpenArray system according to Looft et al.⁵¹, with threshold cycles (C_T) set at 27 cycles as the minimum detection limit. The $\Delta\Delta C_T$ method⁵² was used to calculate the relative abundances of the genes, and 16S rRNA genes were quantified to normalise the AMR gene abundances¹¹.

Primer sets for the 295 AMR and AMR-related assays were previously developed and validated¹¹. Average C_T values were calculated from triplicates; a sample without amplification was considered a “false negative” and discarded.

Individual genes were grouped according to their antibiotic class or AMR-related activity: aminoglycosides (36 genetic determinants), beta-lactam (54), am-/chor-/fluoro-quinolones (FCA, 9), macrolide-lincosamide-streptogramin B (MLSB, 46), sulfonamide (7), tetracycline (39), vancomycin (32), other antibiotic resistances (i.e. to nitroimidazole, pyrazinamide, streptothricin, bacitracin and fosfomycin, 11), multidrug mechanisms (51, here we refer to non-specific resistance mechanisms, such for efflux pumps), integrases (mobile genetic elements, 5) and transposases (mobile genetic elements, 8). The relative abundance of the AMR and AMR-related genes were expressed as the abundance of genes relative to the abundance of bacteria (i.e. the 16S rRNA gene). The total relative abundances per antibiotic class or AMR-related activity were used for statistical analysis. In addition, the total number of genes (referred to as AMR gene richness) for each antibiotic class or AMR-related activity was used for statistical analysis.

Statistical analysis. Statistical analyses were conducted in R version 4.2.1 and SPSS v26. Spearman's rank determined correlations with environmental variables⁵³. Kruskal–Wallis (K–W) test was used to compare multiple populations (a non-parametric equivalent of ANOVA), i.e. Scottish council areas and land use types including arable, bog/wetland, dunes, grassland, improved grassland, rough grassland, heath, larch plantation, moorland, scrub, broadleaf woodland and coniferous woodland. For all correlations, those with $p < 0.01$ were considered statistically significant. Least squares regression analysis was undertaken to explore the

potential influence of land use fully. Data were log-transformed, and categorical variables were dummy coded, as appropriate.

A t test was performed in Excel using a two-tailed distribution and a two-sample unequal variance to determine significant differences between the distribution of resistance mechanisms (antibiotic deactivation, cellular protection, efflux pumps and ‘other’ denoting mobile genetic elements and resistance genes of unknown function) amongst different land uses (arable, semi-natural, managed grassland and woodland/forestry).

Sequenced 16S rRNA genes were used to determine bacterial richness (Chao Index) using Primer7⁵⁴. Principal Components Analysis was done to determine the relative strength of environmental variables to explain the variation in the AMR gene abundance and richness⁵⁵. Data were square-root transformed, and the Bray-Curtis Index determined similarities. The PCA plot was drawn using Primer7, and the environmental variables with the strongest correlations ($r_s > 0.1$) were overlaid on the plot.

Data mapping. AMR gene relative abundance and richness distributions across Scotland were plotted using the ArcGIS Pro GIS software package v 3.0⁵⁶. Each dataset was summarised by quintiles (except in cases where numbers were too low, e.g. integron richness) using graduated symbols of increasing size and blue-red colour ramp to indicate increasing relative abundance and richness.

Reporting summary. Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Data has been archived at the UK Environmental Information Data Centre⁵⁷ and the James Hutton Institute. Bacterial 16S-rRNA gene sequences have been deposited to NCBI, with BioProject ID number PRJNA997939.

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Author contributions

E.P. and C.W.K. wrote the manuscript; R.H. and M.C. generated data and provided statistics support; L.R., X.Y.Z., J.Q.S., and T.P. provided laboratory support and generated data; L.A., T.F., Y.G.Z., D.W.G., and C.W.K. conceived the project and managed elements of the project. C.W.K. was the principal investigator of the project. All authors reviewed and edited the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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