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Plastic-degrading potential across the global microbiome correlates with recent pollution trends

Jan Zrimec¹, Mariia Kokina^{1,2}, Sara Jonasson¹, Francisco Zorrilla^{1,3}, Aleksej Zelezniak^{1,4*}

- 5 1 - Department of Biology and Biological Engineering, Chalmers University of Technology, Kemivägen 10, SE-412 96, Gothenburg, Sweden
- 2 - Department of Chemistry and Biochemistry, Faculty of Agriculture, University of Belgrade, Nemanjina 6, 11080, Belgrade, Serbia
- 3 - Current address: MRC Toxicology Unit, Tennis Ct Rd, Cambridge CB2 1QR, United
10 Kingdom
- 4 - Science for Life Laboratory, Tomtebodavägen 23a, SE-171 65, Stockholm, Sweden
- * corresponding author (email: aleksej.zelezniak@chalmers.se)

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Abstract

Poor recycling has accumulated millions of tons of plastic waste in terrestrial and marine environments. While biodegradation is a plausible route towards sustainable management of plastic waste, the global diversity of plastic-degrading enzymes remains poorly understood.

5 Taking advantage of global environmental DNA sampling projects, here we construct HMM models from experimentally-verified enzyme sequences and mine ocean and soil metagenomes to assess the global potential of microorganisms to degrade plastics. By controlling for false-positives using gut microbiome data, we compile a catalogue of over 30,000 non-redundant enzyme homologues with the potential to degrade 10 different plastic types. While differences

10 between the ocean and soil microbiomes likely reflect the base compositions of these environments, we find that ocean enzyme abundance might increase with depth as a response to plastic pollution and not merely taxonomic composition. By obtaining further pollution measurements, we reveal that the abundance of the uncovered enzymes in both ocean and soil habitats significantly correlates with marine and country-specific plastic pollution trends. Our

15 study thus uncovers the Earth microbiome's potential to degrade plastics, providing evidence of a measurable effect of plastic pollution on the global microbial ecology as well as a useful resource for further applied research.

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1. Introduction

Despite demands for plastic production increasing annually, the problem of plastic waste management remains largely unresolved and presents a global ecological problem ^{1,2}. The majority of plastic waste ends up in landfills or dispersed in the environment, resulting in over 5 150 million metric tons currently circulating in marine environments with an estimated 4.8–12.7 million tons of plastic entering the ocean every year ³. Even monomer additives such as Phthalate compounds, frequently used as plasticizers, are a major source of concern due to their overuse in a variety of different products and adverse health effects ^{4,5}. While some thermoplastics (PE, PP, PET, PVC, PA) can be recycled, contaminated and composite plastics 10 as well as thermosets (PU, vinyl esters) cannot be remolded or heated after the initial forming ^{6,7}. However, although man-made synthetic plastics were designed to remain persistent in the environments, the synthetic polymers, just as natural polymers, can serve as a microbial carbon source ^{8–10}. Microorganisms thus mediate a number of plastic biodegradation reactions and even the toughest plastics including PET ¹⁰ and PU ¹¹, can be transformed and metabolized by 15 microbial species across different environments ^{12–17}. Yet, despite their involvement in the global biogeochemical cycle, the true microbial potential for plastic degradation across different global habitats is not yet fully understood ⁹.

The isolation, identification and characterization of microorganisms with plastic-degrading 20 potential are frequently conducted from aquatic environments ^{18–21}, waste disposal landfills ^{22–25} or places that are in direct contact with the plastic, such as plastic refineries ^{26–28}. However, growing microorganisms outside of their natural environments using conventional approaches is extremely challenging ²⁹ and limits the amount of isolated species that can be cultured and studied to as little as 1% or lower ³⁰. Studying single microbial isolates also limits our 25 understanding of the microbial ecology of plastic degradation, where microbial consortia were found to act synergistically, producing more enzymes and degrading plastics more efficiently than individual species ^{31,32}. Likewise, localized analyses from single locations hinder our understanding of the global environmental impact of plastic materials ³³. On the other hand, with advances in environmental DNA sequencing and computational algorithms, metagenomic 30 approaches enable studying the taxonomic diversity and identifying the functional genetic potential of microbial communities in their natural habitats ^{33–35}. For example, global ocean sampling revealed over 40 million mostly novel non-redundant genes from 35,000 species ³⁵, whereas over 99% of the ~160 million genes identified in global topsoil cannot be found in any

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previous microbial gene catalogue³⁴. This indicates that global microbiomes carry an enormous unexplored functional potential with unculturable organisms as a source of many novel enzymes³⁰. Identification of such enzymes involved in the biological breakdown of plastics is an important first step towards a sustainable solution for plastic-waste utilisation^{36,37}. However, despite the availability of experimentally determined protein sequence data on plastic-degrading enzymes^{10,38-43}, no large-scale global analysis of the microbial plastic-degrading potential has yet been performed.

In the present study, we explore the global potential of microorganisms to degrade plastics. We compile a dataset of all known plastic-degrading enzymes with sequence-based experimental evidence and construct a library of HMM models, which we use to mine global metagenomic datasets covering a diverse collection of oceans, seas and soil habitats^{34,35,44,45}. By controlling for false-positives using gut microbiome data⁴⁶, we compile a catalogue of over 30,000 non-redundant enzyme homologues with the potential to degrade 10 different plastic types. Comparison of the ocean and soil fractions shows that the uncovered enzymatic potential likely reflects the major differences related to the composition of these two environments. Further analysis of metagenome-assembled genomes in the ocean reveals a significant enrichment of plastic-degrading enzymes within members of the Alpha- and Gamma-proteobacteria classes, and supports the notion that enzyme abundance increases with depth as a response to plastic pollution and not merely taxonomic composition⁴⁷⁻⁴⁹. By relating the identified enzymes to the respective habitats and measured environmental variables within the soil and ocean environments, we further show that the abundance of the uncovered enzymes significantly correlates with both marine and country-specific plastic pollution measurements⁵⁰⁻⁵⁵, suggesting that the earth's microbiome might already be adapting to current global plastic pollution trends.

2. Results

Global microbiome harbours thousands of potential plastic-degrading enzymes

To probe the potential for plastic degradation across the global microbiome, we mined published studies ^{10,38–42,56–61} and databases ⁴³ and compiled a dataset of known enzymes with experimentally observed evidence of plastic modifying or degrading activity, representing a total of 95 sequenced plastic enzymes spanning 17 different plastic types from 56 distinct microbial species (Figure 1a, Methods M1, Dataset S1). The types of plastics (13 types) and plastic additives (4 types of phthalate-based compounds, see Figure 1a: additives marked with a star) spanned the main types of globally produced plastics that constitute the major fraction of global plastic waste ¹, except for PP and PVC, for which no representatives could be found (Figure S1). To enable efficient searching across global metagenomic datasets we built Hidden markov models (HMMs) ⁶² by including the known homologous sequences from the Uniprot Trembl database ⁶³ (Figure 1a,b, Figure S2). Briefly, we clustered the known enzymes to obtain representative sequences (95% seq. id., Figure 1a) and used these to query the Uniprot Trembl database and obtain an expanded dataset of a total of 16,834 homologous enzyme sequences (E -value < $1e-10$, Methods M2, Figure S2). Each group of enzyme sequences at a given Blast sequence identity cutoff ranging from 60% ⁶⁴ to 90% was then clustered (95% seq. id.) to obtain groups of representative sequences that were used to construct a total of 1204 HMM models (Figure 1a, Figure S3, Methods M2).

The HMMs were then used to search for homologous sequences from the metagenomes spanning 236 sampling locations (Methods M3, Figure 1e) that included global ocean ³⁵, global topsoil ³⁴ and additional Australian ⁴⁵ and Chinese topsoil projects ⁴⁴ (Methods M1, Table S1). With over 73% of orthologous groups shared between gut and ocean microbiomes ³⁵, a high number of false positive identifications would be expected, as certain enzymes might have related evolutionary ancestry but no plastic degradation activity. Thus, as a control, we filtered the environmental hits by comparing them to those in the gut microbiome ⁴⁶, where little to none plastic enzyme coding potential should exist. Briefly, for each HMM model *precision* and *recall* were computed by comparing the corresponding hits in the global microbiomes to those in the gut microbiome and, to minimize the risk of false positives, models with hits in the global

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microbiomes with scores above a precision threshold of 99.99% and AUC of 75% were retained (Figure 1c, Methods M3). The final filtered results with the global microbiomes contained 121 unique HMM models, of which 99 HMM models matched (E -value $< 1e-16$) to ocean samples and 105 to soil samples, representing 10% of the initial HMM models used prior to filtering (Table S1). Consequently, an average of 1 in 4 organisms in the analysed global microbiome was found to carry a potential plastic-degrading enzyme (Table S1, Dataset S2).

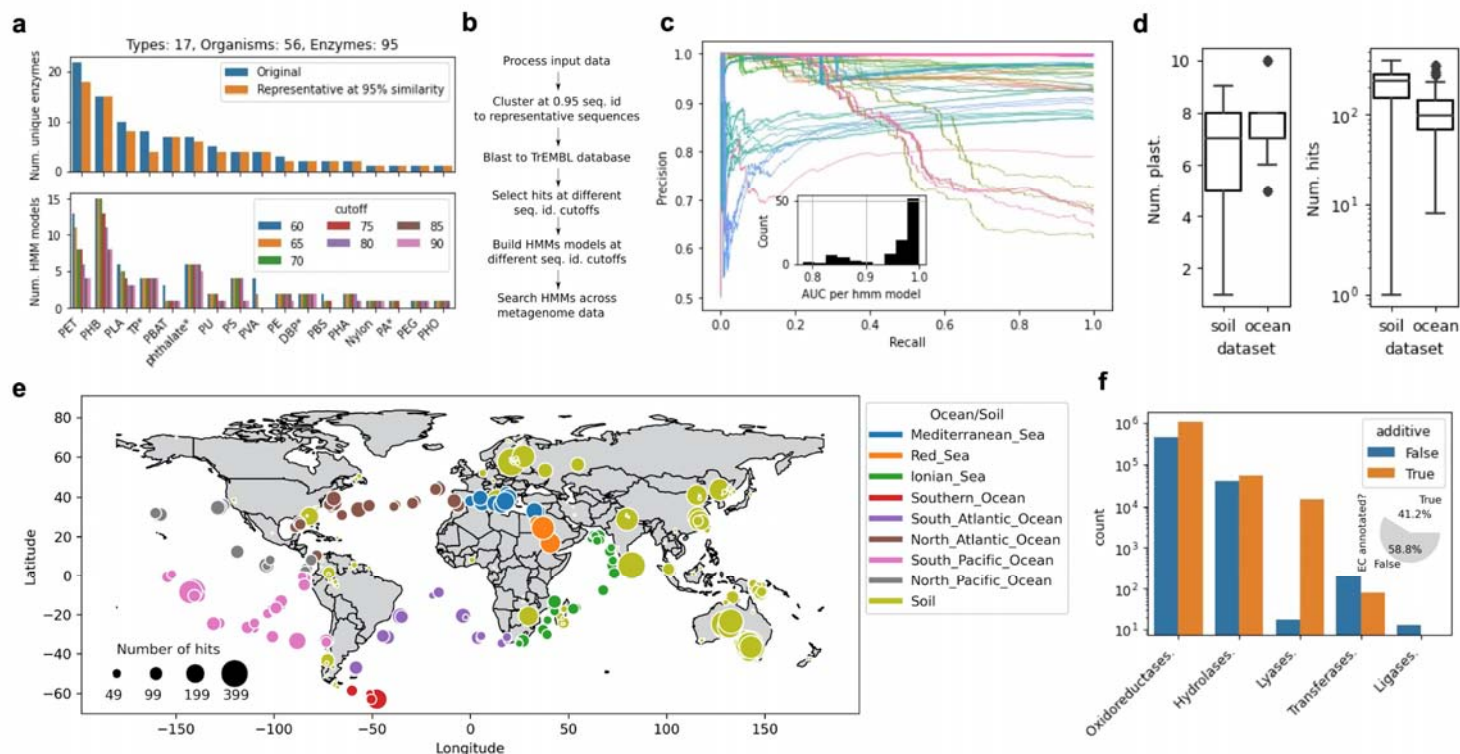
The recovery of unique plastic types was ~60%, 10 in ocean and 9 in soil datasets, respectively, identifying altogether 11,906 hits in the ocean and 18,119 in the soil datasets (Figure 1e). Of these, 38 HMM models matched 43% of hits corresponding to the 6 plastic polymers (Figure S4a: PBAT, PEG, PET, PHB, PLA, PU) and 83 HMM models identified 57% of hits corresponding to the 4 additives (Figure S4a: DBP, PA, TP, phthalate). Specifically, of the plastic polymer enzyme hits, PU was found only in the ocean and not in the soil microbiome, whereas over 2-fold higher amounts of PEG, PBAT and PHB and a 2-fold lower amount of PET were found in the ocean fraction compared to the soil (Figure S4a). The amount of hits corresponding to additives was significantly (Fisher's exact test one-tailed p -value = $5.4e-6$) larger in the soil fraction than the ocean fraction, representing 69% of the total amount of soil hits compared to 39% in the ocean fraction and resulting in an almost 4-fold increase in the average amount of additives across the soil sampling sites (Figure S4b). On the other hand, the overall number of plastic polymers across the samples was relatively similar in both the soil and ocean fractions, with a 15% larger amount observed in the soil samples (Figure S4b). The resulting amount of all hits including polymers and additives was thus, on average, over 2-fold larger across the soil samples than in the ocean samples, whereas the amount of distinct plastic types was equal (Figure 1d). These results were however much more variable across the soil fraction, where, for instance, the variability of the number of hits across soil sampling sites was over 4-fold larger compared to the ocean fraction (Figure 1d).

The identified enzyme hits were annotated using orthologous function mapping^{65,66} (Methods M3), which assigned EC enzyme classifications for 41% of the hits (Figure 1f inset) with the majority of the annotated enzyme classes corresponding to oxidoreductases, hydrolases and lyases (Figure 1f). An over 2-fold larger fraction of monomer additives were annotated compared to the general polymer plastics, meaning that, whereas ~1/2 of the additives were annotated, this was the case with only 29% of the general polymers (Figure S5a). Despite similarities in distributions of the general classes across the ocean and soil fractions (Figure 1f),

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37% less hits were annotated with the soil fraction (Figure S5b). Further analysis showed that indeed differences in function were present, with the ocean fraction possessing an 11% larger diversity of enzyme functions than soil (Figure S6a: 40 vs 36 distinct enzyme types with at least 3 occurrences) and 27% of the enzyme functions differing among the two microbiome fractions.

5 The difference between additives and polymer plastics was however discernible already at the level of general enzyme classes (Figure S5c). Similarly, in both ocean and soil fractions, an almost 3-fold larger amount of functional diversity was present with the additives than with the



polymers, and only a single function (2%) was shared among the additive and polymer

Figure 1. Global microbiome harbours thousands of potential plastic-degrading enzymes.

10 enzyme dataset and representative sequences obtained by clustering (95% seq. id. cutoff), covering the major types of pollutant plastics (PVA, polyvinyl alcohol; PLA, polylactic acid; PU, polyurethane; PHB, polyhydroxybutyrate; PBS, polybutylene succinate; PET, polyethylene terephthalate; Nylon; PBAT, polybutylene adipate terephthalate; PE, polyethylene; PEG, polyethylene glycol) and additives/plasticisers (phthalate; PA, phthalic acid; DBP, di-n-butyl phthalate; TP, terephthalic acid). The lower plot shows the final constructed HMM models across the different sequence Identity cutoffs. (b) Overview of the procedure to construct the HMM models. (c) Precision-recall curves with the 99 HMM models that returned results in the ocean fraction. Inset: area under the curve (AUC) with these HMM models. (d) Number of plastic-degrading enzyme hits and plastic types across the ocean and soil microbiome fractions. (e) Plastic-degrading enzyme hits across the global microbiome. (f) Enzyme classes (EC) predicted with orthologous function mapping⁶⁵ at the topmost EC level. Inset shows the amount of EC annotated results.

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Earth microbiome's plastic-degrading potential might already be adapting to global pollution trends

The analysed ocean microbiome spanned 67 locations sampled at 3 depth layers and across 8 oceans (Figure 1e, Methods M1). A significant (Rank Sum test p -value $< 2.9e-2$) increase of plastic-degrading enzyme hits was identified in samples obtained from the Mediterranean Sea and South Pacific Ocean compared to the other locations (Figure 2a, Table S3), which might reflect the relatively high plastic pollution in these areas^{52,67}. A higher amount of pollution in sampling areas in the lower longitudinal region, however, might be indicated by the significant negative correlation (Spearman r was 0.393 and 0.357, p -value $< 1.6e-5$) of both the plastic types and enzyme hits, respectively, with longitude (Figure 2b, Figure S7). Whereas the majority of plastic polymer and monomer additive types were found across all oceans, PU was only present in the Ionian Sea and South Pacific Ocean, whereas PLA only in the Ionian Sea, likely reflecting their overall 6-fold lower content than the other plastic types (Figure S8a).

As expected according to published results showing an increasing amount of taxonomic and functional richness with depth³⁵, we observed measurable depth stratification of the enzyme hits in the ocean samples (Figure 2c). Both the amount of plastic types and enzyme hits were positively correlated with depth (Spearman r was 0.552 and 0.384, p -value $< 4.3e-6$, respectively) as well as negatively correlated with temperature (Spearman r was 0.451 and 0.336, p -value $< 6.7e-5$, respectively, Figure 2b,c, Figure S7). This was also supported by Principal coordinate analysis (PCoA) on enzyme hits across samples (Methods M3), where the first principal coordinate carrying 25% of the data variance correlated significantly (Spearman r was 0.453 and -0.420, p -value $< 4e-7$) with both depth and temperature, respectively (Figure S9,10). We therefore next reconstructed metagenome-assembled genomes (MAGs) in the ocean samples and predicted their taxonomies (Methods M1). The results corroborated a significant correlation (Spearman r of 0.392 and 0.548, p -value $< 2.5e-6$) between the number of plastic types and enzyme hits, respectively, with the number of unique organisms at the family level (Figure S11; similar results with other taxonomic levels). We found that, although the majority (62%) of organisms (MAGs) were associated with a single plastic type (Figure 2d inset), 2.5% of them carried enzymes corresponding to 4 or more different plastic types (Figure 2d inset, Figure S12). Analysis of the plastic distribution across species showed that the number of enzyme hits was significantly (Fisher's exact test one-tailed p -value $< 1.4e-05$) enriched within Alpha- and Gamma-proteobacteria, which can be expected since this is the most

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abundant and diverse phylum in the dataset (Figure 2d, Table S4). Nevertheless, the results suggested that the observed plastic-degrading enzyme abundance (Figure 2d) might not be a reflection of merely taxonomic and functional richness, but also of recently uncovered large amounts of plastic pollution below the ocean surface^{47–49}.

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The analysed soil microbiome spanned 169 sampling locations across 38 countries and 11 distinct environmental habitats (Figure 1e, Methods M1). To ensure the accuracy of cross-habitat and cross-country comparisons, due to the different technical specifications of sample acquisition and processing across the metagenomes^{34,44,45}, here we focused on the uniformly processed global topsoil dataset³⁴, which also represented the largest fraction of the data (163 sampling locations) covering all given countries and habitats. A significant (Rank Sum test p -value < 4.8e-3) increase of plastic-degrading enzyme hits was identified in samples from the Moist tropical forests and Tropical montane forests habitats compared to the other habitats (Figure 2e). This was corroborated by a significant correlation (Spearman r was 0.248 and 10 0.332, p -value < 5e-5) of both the amount of plastic types and enzyme hits, respectively, with longitude as well as the amount of enzyme hits with both the measured annual moisture content (Spearman r = 0.292, p -value = 6.8e-6) and precipitation levels (Spearman r = 0.330, p -value = 4.6e-8, Figure 2f, Figure S13,14). Interestingly, the soil habitats contained the most distinct differences of plastic content compared to the ocean microbiome, with all plastic types present 15 only in the Moist tropical forests and Temperate deciduous forests (Figure S8b). Besides these two areas, PET for example was additionally found only in the Mediterranean habitat (Figure S8b).

Since the results suggested that the plastic-degrading enzyme hits might reflect actual global 25 pollution trends (Figure 2a,e), and considering that global pollution with plastics and microplastics has been an ongoing and steadily increasing problem for over 5 decades^{68,69}, we next determined if the global potential for plastic degradation reflected the current plastic pollution trends. We obtained data from 4 ocean expeditions^{50–54}, pooling the data to cover 61% of the ocean sampling locations at the surface depth layer, and matched the closest data points 30 to those of the ocean sampling locations at a maximum radius of 400 km (see sensitivity analysis in Table S2, Methods M1). Similarly, by obtaining a dataset of mismanaged and inadequately managed plastic waste across different countries^{55,70} we achieved a 72% coverage of the soil samples across 35 countries. Using these common pollution datasets, we indeed observed significant correlation (Spearman r of 0.492 and 0.407, p -value < 1.1e-3)

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between the numbers of identified enzymes and pollution trends within both the ocean and soil microbiomes, respectively (Figure 2g,h). Strikingly, this observed correlation between the abundance of plastic-degrading enzymes with global pollution suggests that the global microbiome might already be adapting to the effects of global plastic pollution.

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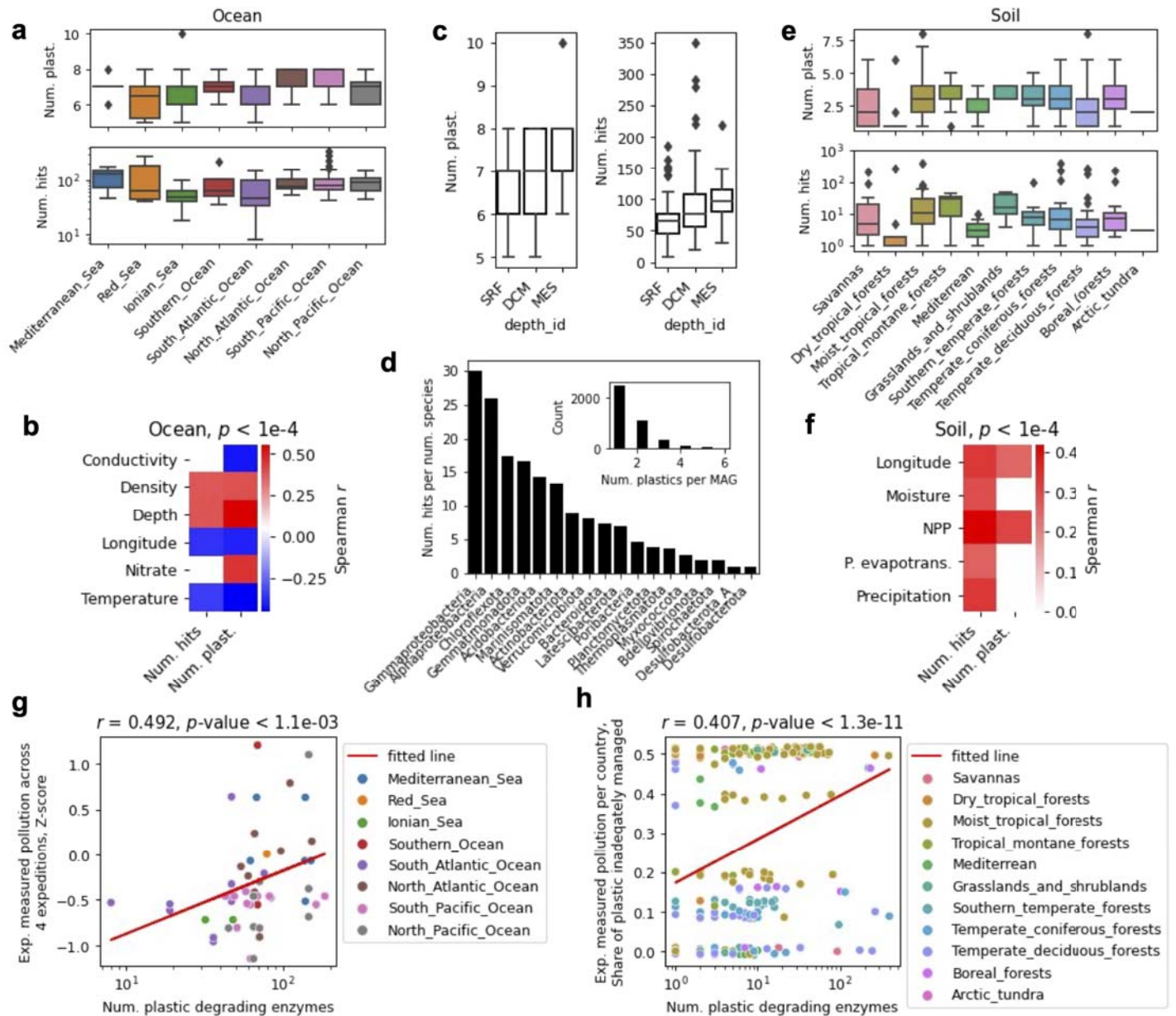


Figure 2. Earth microbiome's plastic-degrading potential might already be adapting to global pollution trends. (a) Number of plastic-degrading enzyme hits and different plastic types found across 8 oceans. (b) Correlation between the number of enzyme hits and different plastic types with ocean environmental variables: longitude [°], depth [m], conductivity [mS/cm], temperature [°C], water density [kg/m] and nitrate content [$\mu\text{mol/l}$] ³⁵. All p -values $< 1e-4$. (c) Number of enzyme hits and different plastic types across the ocean sampling depth layers ³⁵. (d)

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Number of enzyme hits relative to the number of species obtained with the metagenome-assembled genome (MAG) analysis at the phylum level (class-level for Proteobacteria) (Methods M1, see text). Inset: number of plastic types per MAG. (e) Number of plastic-degrading enzyme hits and different plastic types found across 11 soil habitats. (f) Correlation between the number of enzyme hits and different plastic types with soil environmental variables: longitude [°], avg. monthly moisture content [%], net primary productivity (NPP) [$\text{gcm}^{-2}\text{yr}^{-1}$], avg. yearly potential evapotranspiration and precipitation [L/m^2]³⁴. All p -values $< 1\text{e-}4$. (g) Correlation of ocean plastic-degrading enzyme hits with experimentally measured plastic pollution across 4 ocean expeditions⁵⁰⁻⁵⁴ (Methods M1). (h) Correlation of soil plastic-degrading enzyme hits with the share of inadequately managed plastic per country⁵⁵.

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3. Discussion

Here, we catalogued potential plastic-degrading enzymes, including the majority of massively produced and globally polluting polymers (Figure 1a, Figure S1) as well as the major additives involved in plastic production, identified from metagenomes sampled from soils and oceans across the globe^{34,35,44,45} (Figure 1e). We used an initial set of 95 experimentally verified published sequences (Dataset S1) and expanded it with Uniprot sequences to build enzyme sequence models (hidden Markov models⁶²) for mining metagenomic data (Figure 1a,b). We identified a total of 30,000 enzyme hits in the ocean and soil microbiomes (Figure 1e: 11,906 and 18,119, respectively) corresponding to 10 major plastics types, including 6 polymers and 4 additives (Figure 1d, Figure S4). To minimize the number of false positive hits, we used the gut microbiome⁴⁶ as a negative control (Figure 1c), that is, we assumed that the gut microbiome is not evolved to degrade plastics and thus enzyme hits that are similar to the ones found in the human gut would indicate false positives. Nearly 60% of identified plastic-degrading enzymes did not map to any known enzyme classes (Figure 1f), suggesting that novel plastic-degrading functional content was uncovered, which is not surprising considering the vast amounts of novel functions being uncovered in recent large-scale metagenomic studies^{33-35,49}.

A potential reason for the observed functional differences between the soil and ocean microbiomes (Figure 1d,f, Figure S4,5,6) could arise not only from the different plastic availability and pollution trends across these environments^{50-52,55}, but also from the general mechanical and chemical differences between these two environments⁷¹. For instance, the ocean is a highly dynamic environment due to its compositional medium with a larger degree of mixing. As such, compared to soil that is in large part composed of solids, one can expect an intrinsically lower community and functional stratification per unit volume in the ocean³⁵. The increased variability of enzyme hits and plastic types across soil habitats (Figure 1d, Figure S12), for instance, was likely a reflection of such differences. Furthermore, the large fluctuations in temperature, salinity and mechanical forces in the ocean lead to it intrinsically possessing many polymer-degrading properties⁷²⁻⁷⁴, differing from those in the soil⁷¹ and possibly resulting in further preferences in the specific functional content. On the other hand, the soil generally contains a higher observed overall species richness^{34,75}, and thus it is likely that certain enzyme families are overrepresented in each environment. This, as well as the fact that additive monomers are likely easier to degrade than the general plastic polymers due to being simpler molecules, could be the reason behind the observed large differences in the additive versus

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polymer content between the ocean and soil fractions (Figure S5,6). Moreover, the uncovered additive-degrading enzymes in soil likely corresponded to overrepresented but unknown enzyme classes in soil that could not be identified using the orthogonal mapping procedure^{65,66} (Figure s5b).

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Plastics have been increasingly mass produced ever since the economic and social explosion after the 2nd world war with the first signs of global plastic pollution concern arising over half a century ago^{68,69}, giving ample evolutionary time for microbial functional adaptation to these compounds^{49,76,77}. Such adaptation was recently uncovered with PET-degrading enzymes
10 across ocean metagenomes of planktonic communities⁴⁹, where multiple fully-functional enzyme variants were found to be evolved from ancestral enzymes degrading polycyclic aromatic hydrocarbons, suggesting that the current PET exposure already provides sufficiently strong selective pressures to direct the evolution and repurposing of such enzymes. Similarly, enzymes degrading other plastic types have been shown to be widely occurring with numerous
15 homologs in diverse organisms and likely arising from well conserved general enzyme classes^{78,79}. Indeed, here we find multiple lines of evidence supporting that the global microbiome's plastic-degrading potential reflects recent measurements of environmental plastic pollution. Firstly, we find that taxonomic and functional richness is likely not the only driver of the observed depth stratification of enzyme hits (Figure 2c). The organisms found to carry the
20 largest amount of plastic-degrading enzymes (Figure 2d) do not completely reflect initial taxonomic estimates in the ocean³⁵, indicating that the plastic-degrading potential also reflects the recently uncovered trends of an increasing amount of plastic pollution below the surface (<200m)⁴⁸ with considerable microplastic pollution in the mesopelagic zone⁴⁷, which are potentially stronger drivers of the observed depth stratification⁴⁹. Secondly, certain habitats
25 containing the highest amounts of observed enzyme hits, such as the Mediterranean Sea and South Pacific Ocean (Figure 2a), are known to be highly polluted areas^{52,67}. Lastly, this prompted us to verify and uncover the significant measurable correlation of both ocean and soil enzyme hits with experimentally measured pollution across oceans and countries from multiple datasets⁵⁰⁻⁵⁵ (Figure 2g,h), suggesting that the Earth microbiome's potential for plastic
30 degradation is already evolving as a response to the rise in environmental pollution.

Considering that natural plastic-degradation processes are very slow (e.g. predicted life time of a PET bottle at ambient conditions ranges from 16 to 48 years⁸⁰), the utilisation of synthetic biology approaches to enhance current plastic-degradation processes is of crucial importance

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^{81,82}. Moreover, although there is still unexplored diversity in microbial communities, synergistic degradation of plastics by microorganisms holds great potential to revolutionise the management of global plastic waste ^{36,37}. To this end, the methods and data on novel plastic-degrading enzymes produced here can help researchers (i) gain further information about the taxonomic diversity of such enzymes as well as understanding of the mechanisms and steps involved in the biological breakdown of plastics, (ii) point toward the areas with increased availability of novel enzymes, and (iii) provide a basis for further application in industrial plastic-waste biodegradation.

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4. Methods

M1. Datasets

We compiled the initial dataset of 95 sequenced plastic enzymes spanning 17 plastic types with experimentally observed evidence of plastic modifying or degrading activity from published studies^{10,38–42,56–61} and databases⁴³ (Dataset S1).

Metagenomic sequencing data was obtained from the Tara ocean expedition³⁵, global⁴⁴, Australian⁴⁵ and Chinese topsoil projects³⁴ and a gut microbiome study⁴⁶. From the sequencing data metagenomic assemblies were reconstructed using MEGAHIT v1.2.9⁸³ with the ‘--presets meta-sensitive’ parameter, except with Tara oceans where the published assemblies were used³⁵. Metagenome-assembled genomes (MAGs) were constructed for the ocean dataset by first cross-mapping paired end reads to assemblies with kallisto v0.46.1⁸⁴ to obtain contig coverage information across samples. This information was then input to CONCOCT v1.1.0⁸⁵ to generate a draft bin set. MetaBAT2 v.2.12.1⁸⁶ and MaxBin2 v2.2.5⁸⁷ were also used to generate additional draft bin sets. Finally, the three bin sets were de-replicated and reassembled using metaWRAP v1.2.3⁸⁸ with parameters ‘-x 10 -c 50’ to obtain the final set of MAGs. Default settings were used except where otherwise stated. Environmental data for the Tara ocean and global topsoil microbiomes was obtained as specified in the respective publications^{34,35}: (i) ocean data from the PANGAEA database (www.pangaea.de), (ii) soil data from the Atlas of the Biosphere (<https://nelson.wisc.edu/sage/data-and-models/atlas/maps.php>), except for temperature and precipitation data that was obtained from the WorldClim database (<https://www.worldclim.org/>). With the ocean data the prokaryote fraction was used³⁵. Global topsoil habitats were used as defined³⁴. Experimentally measured pollution data across the ocean from published ocean expeditions^{50–54} was pooled by normalizing the data using the Box-Cox transform⁸⁹ and computing Z-scores.

M2. Construction of HMM models

To construct the HMM models, we first obtained representative sequences from the initial input sequence data by clustering them using CD-HIT v4.8.1^{90,91} with default settings, except a word size of 5, cluster size of 5 and seq. id. cutoff of 95%. To expand the sequence space for building

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the HMM models, the Uniprot Trembl database⁶³ was queried with the representative enzyme sequences using BLAST+ v2.6⁹² with default settings except for an *E*-value cutoff of 1e-10. For each group of enzyme sequences at a given Blast sequence identity cutoff ranging from 60% to 90% in increments of 5%, representative sequences were obtained by clustering using CD-HIT
5 with the same parameters as above. Finally, HMM models were constructed using the HMMER v3.3 *hmmbuild* utility⁹³ (<http://hmmer.org/>) with default settings.

M3. Statistical and correlation analysis

For identifying homologous sequences in metagenomes *hmmsearch* from HMMER v3.3⁹³ was
10 used with default settings. To minimize the risk of false positive results, we filtered the environmental hits by comparing their bitscore to those obtained with the gut microbiome. For each HMM model *precision* and *recall* were computed by comparing the corresponding hits in the global microbiomes to those in the gut microbiome, where only models with a minimum of 20 data points and hits in the global microbiomes with an *E*-value cutoff below 1e-16 and scores
15 above a precision threshold of 99.99% and AUC of 75% were retained. Additionally, only the lowest *E*-value and bitscore hit was retained for each gene in the global metagenomes. The precision-recall analysis was performed using Scikit-learn v0.23.1⁹⁴ with default settings. Orthologous function mapping was performed using Egnog-mapper v2^{65,66} with default settings. Principal coordinate analysis (PcoA) was performed using Scikit-bio v0.5.5
20 (<http://scikit-bio.org/>) with default settings and the Bray Curtis distance. For statistical hypothesis testing, Scipy v1.1.0⁹⁵ was used with default settings. The Spearman correlation coefficient was used for correlation analysis. All tests were two-tailed except where stated otherwise.

M4. Software

25 Snakemake v5.10.0⁹⁶, Python v3.6 (www.python.org) and R v3.6 (www.r-project.org) were used for computations.

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Author contributions

JZ and AZ conceptualized the project; JZ, SJ, FZ and AZ designed the computational analysis; JZ, MK, SJ, FZ and AZ performed the computational analysis; JZ and AZ interpreted the results; JZ, MK, and AZ wrote the initial draft manuscript; JZ and AZ revised the draft and wrote the final manuscript.

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Competing Interests

The authors declare no competing interests.

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