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microntology: a lightweight, data-driven controlled vocabulary to describe earth's microbial habitats

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Abstract

Motivation: Data-enabled studies of microbial ecology and evolution depend on high-quality descriptions of microbial habitats, based on curated and consolidated vocabularies.

Results: We introduce *microntology* v1.0, a pragmatic controlled vocabulary of 148 terms to describe microbial habitats and lifestyles, and provide manually curated *microntology* annotations for >300k metagenomic samples from public repositories.

Availability: *microntology* controlled vocabulary terms and term hierarchies (doi: [10.5281/zenodo.19730167](https://doi.org/10.5281/zenodo.19730167)), and curated annotations for 305 626 metagenomic samples (doi: [10.5281/zenodo.18164252](https://doi.org/10.5281/zenodo.18164252)) are available via Zenodo and spire.embl.de/downloads. Underlying code is available via github.com/grp-schmidt/microntology and Zenodo (doi: [10.5281/zenodo.20323497](https://doi.org/10.5281/zenodo.20323497)). User feedback, suggestions and bug reports are welcome at github.com/grp-schmidt/microntology/issues.

1 Introduction

Data on microbial life continues to accrue in public repositories: at the time of writing, nearly 6 million amplicon sequencing and 1.2 million metagenomic experiments on microbial communities are freely accessible via the International Nucleotide Sequence Database Collaboration (INSDC, [Karsch-Mizrachi et al. 2025](#)), with rapid growth also in other 'omic data types ([Yurekten et al. 2024](#), [Perez-Riverol et al. 2025](#)). This wealth of information enables studies of microbial ecology and evolution at unprecedented *depth* (i.e. with deep readouts on individual samples or habitats) and *breadth* (across many samples, often across different habitats), facilitated by dedicated resources that integrate microbial data across Earth's biomes, such as catalogues of genes ([Coelho et al. 2022](#)), gene families ([Rodríguez Del Río et al. 2024](#)), metabolic pathways ([Kautsar et al. 2021](#)), metagenome-assembled genomes (MAGs, [Gurbich et al. 2023](#), [Nayfach et al. 2021](#), [Schmidt et al. 2024](#), [Sun et al. 2026](#)) or taxonomic profiles ([Woodcroft et al. 2025](#), [Matias Rodrigues et al. 2026](#)).

A key challenge in navigating repositories and identifying datasets relevant to specific research questions is the varying availability of consistent contextual data, which is why INSDC databases require submissions to conform with MixS standards by providing Minimum Information about Genome/Metagenome Sequences (MIGS/MIMS, [Field et al. 2008](#)), MAGs (MIMAGS, [Bowers et al. 2017](#)) or MARKer gene Sequences (MIMARKS, [Yilmaz et al.](#)

[2011](#)). These include descriptions of the broad-scale and local environmental context [formerly defined as annotation fields *environment* (*biome*) and *environment* (*feature*)], as well as the sampled environmental medium [*environment* (*material*)], with a recommendation to use terms from corresponding classes in the Environment Ontology (EnvO, [Buttigieg et al. 2013, 2016](#)). However, annotations are often incomplete, uninformative (using generic terms) or indeed erroneous ([Kuhn et al. 2026](#)) and require extensive curation and harmonization, provided via dedicated databases such as curatedMetagenomicData ([Pasolli et al. 2017](#)) or Metalog ([Kuhn et al. 2026](#)). Moreover, EnvO and other available controlled vocabularies to describe microbial habitats, such as Omnicrobe ([Dérozier et al. 2023](#)) or the GOLD Ecosystem Classification ([Mukherjee et al. 2025](#)) used for IMG/M resources ([Chen et al. 2023](#)), are designed towards maximum resolution, containing thousands of classes that can be difficult to navigate and summarise into broader categories of related terms. Here, we introduce *microntology*, a lightweight controlled vocabulary to describe microbial habitats and lifestyles that addresses these challenges and complements existing ontologies.

2 Results and discussion

microntology follows five core design principles: it is (i) data-driven, encompassing terms to describe frequent sample types

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in existing data; (ii) pragmatic and query-oriented, defining terms at intermediate resolutions to facilitate user access and browsing; (iii) shallow, with few hierarchical levels of classes; (iv) cross-linked between related terms from different hierarchical paths (e.g. a *lake* [MICRONT: 02010100] would also be considered a *lentic water body* [MICRONT: 02060100]); and (v) designed for parsimonious multi-tagging where individual samples are described by multiple independent terms, rather than a single highly detailed ‘best match’ term (see examples below).

In v1.0, *microntology* encompasses a controlled vocabulary of 148 terms that fall into five broad habitat categories (terrestrial, aquatic, aerial, host-associated and anthropogenic environments) and additional categories that describe broad physicochemical properties (sample salinity, temperature, pH and oxygen level) which are often associated with microbial community structure in different habitats, morphology (biofilms), human host properties to further differentiate within this dominant sample group (age group, birth term, place of residence, non-industrialized lifestyles), animal host captivity status, and miscellaneous informative features such as (chemical) contamination or ancient (paleontological) sample provenance. Terms are moreover linked to matching classes in EnVO (Buttigieg *et al.* 2013, 2016), UBERON (Mungall *et al.* 2012) and other existing ontologies, as well as to NCBI Taxonomy IDs (Schoch *et al.* 2020) where applicable. An overview of *microntology* terms (and their frequency among currently available metagenomes, see below) is shown in Fig. 1; a full table including resolved hierarchies, cross-linked terms and definitions is available as Supplementary Table S1 and in Fullam *et al.* (2026a).

Major habitat categories are further resolved along parallel yet cross-linked paths. *Terrestrial* habitats (MICRONT: 01000000) can be described based on sample type (e.g. *soil*, *rock*, *subsurface*) and environmental context (*forest*, *grassland*, *tundra*, etc.). Following *microntology*’s underlying philosophy of using multiple descriptive tags, a forest soil sample would be annotated using the terms *soil* (MICRONT: 01010000) and *forest* (MICRONT: 01020400), rather than a specific term ‘forest soil’ as a subclass of one or the other. *Aquatic* habitats (MICRONT: 02000000) are further categorized based on general salinity ecosystem (*fresh water*, *brackish*, *marine*, *non-marine saline*, *brine*), water body type (*lentic* or *lotic*), broad water layer (*littoral*, *pelagic*, *benthic*), and dedicated categories describing *springs* (including *hydrothermal vents*), *ice*, and *groundwater* samples. *Aquatic sediments* are considered as a distinct subcategory, cross-linked to *benthic* features (*lake and stream beds*, *seafloor*). Descriptions of habitats at terrestrial-aquatic interfaces are resolved via both cross-linking of terms and multiple tagging: *groundwater* (MICRONT: 02070100) is considered an *aquatic* habitat, but automatically cross-tagged as *terrestrial* and *subsurface* (MICRONT: 01030000); *wetlands* (MICRONT: 01020700) are categorized as *terrestrial*, but cross-tagged as *lentic* (MICRONT: 02060100) and *paludal* (MICRONT: 02090000) water bodies; *soils* from intertidal zones or ocean coasts would in practice be co-tagged as *littoral* (MICRONT: 02080100) and *marine* (MICRONT: 02020000).

Host-associated samples (MICRONT: 03000000) are broadly split into *plant host* (MICRONT: 03010000) and *animal host* (MICRONT: 03020000) groups. The latter is further resolved based on broad host taxonomy (*mammalian*, *avian*, *sponge hosts*, etc), body site (*digestive tract*, *urogenital tract*, *skin*,

airways, etc) and sample type (*bodily fluids* or *mucosa*). *Feces* (MICRONT: 03030141), *saliva* (MICRONT: 03030111) and *ancient dental calculus* (MICRONT: 03030112) are sample types that are treated as special cases, as they are unambiguously *intestinal* (MICRONT: 03030140) and *oral* samples (MICRONT: 03030110), respectively. Following *microntology*’s multi-tag approach, a dog fecal sample would be annotated as *canine* (MICRONT: 03020131) and *feces* (MICRONT: 03030141), whereas caecal contents of chicken would be considered as *avian* (MICRONT: 03020210) *intestine* (MICRONT: 03030140). For *human* (MICRONT: 03020111) samples, by far the most data-rich category in practice, additional information can be added by tagging age group (from *infant* to *elderly*), birth term (*full* or *preterm birth*), place of residence (*urban* or *rural*) and host lifestyle (e.g. *non-industrialized*).

The *anthropogenic* (MICRONT: 04000000) category includes engineered or directly human-impacted habitats, such as the *built environment* (MICRONT: 04020000), *agriculture-associated* environments (MICRONT: 04040000), *wastewater* (MICRONT: 04030000) and *food* (MICRONT: 04060000). *microntology* includes a special *artificial sample* category (MICRONT: 04010000) to flag mock communities or negative controls, but also to describe experimentally derived samples from raw environmental starting materials, e.g. using controlled environments, cell sorting or enrichment cultures. For example, microcosm experiments from lake sediments would be annotated as *lake bed* (MICRONT: 02010101), but qualified with a *controlled environment* (MICRONT: 04010300) tag.

We curated *microntology* annotations for 305 626 publicly available shotgun metagenomic samples from 1688 studies [Fig. 1; per-sample data available under (Fullam *et al.* 2026b)]. We initially included all metagenomes (*source: METAGENOMIC* and *strategy: WGS*) from projects containing ≥ 20 samples with $\geq 1\text{M}$ reads and $\geq 1\text{Gbp}$ sequencing depth, generated on Illumina machines or compatible platforms (DNBseq, BGISEQ), that were available on the European Nucleotide Archive [ENA, O’Cathail *et al.* 2025] on 31st October 2024. From this initial list, datasets were excluded upon manual inspection if they were (i) misannotated amplicon or genomic sequencing data; (ii) artificial samples such as synthetic metagenomes or mock communities; (iii) obtained from *in vivo* models in a laboratory context, e.g. mouse experiments; (iv) enriched for viral particles or specific prokaryotic community members. Additional datasets were included on a per-study basis, in particular from IMG/M (Chen *et al.* 2023) and SPIRE (Schmidt *et al.* 2024). For 1501 studies (representing 95% of samples), ENA project IDs were matched to publications following a previously described approach (Schmidt *et al.* 2024, Kuhn *et al.* 2026). *microntology* terms were annotated in multiple steps (see Supplementary Discussion for details). First, for a subset of 129 904 samples, available manually curated information was parsed from Metalog (Kuhn *et al.* 2026) and matched to *microntology* tags. Next, we mapped tags to entries from ENA contextual data fields (such as environmental context, scientific name and host tax ID). Finally, for a subset of 163 studies and an additional 22 862 samples, *microntology* tags were manually added based on information in associated publications or additional (not automatically parsed) ENA contextual data fields.

In our survey (Fig. 1; see Supplementary Fig. S1 for an alternative view), publicly available metagenomic data remains

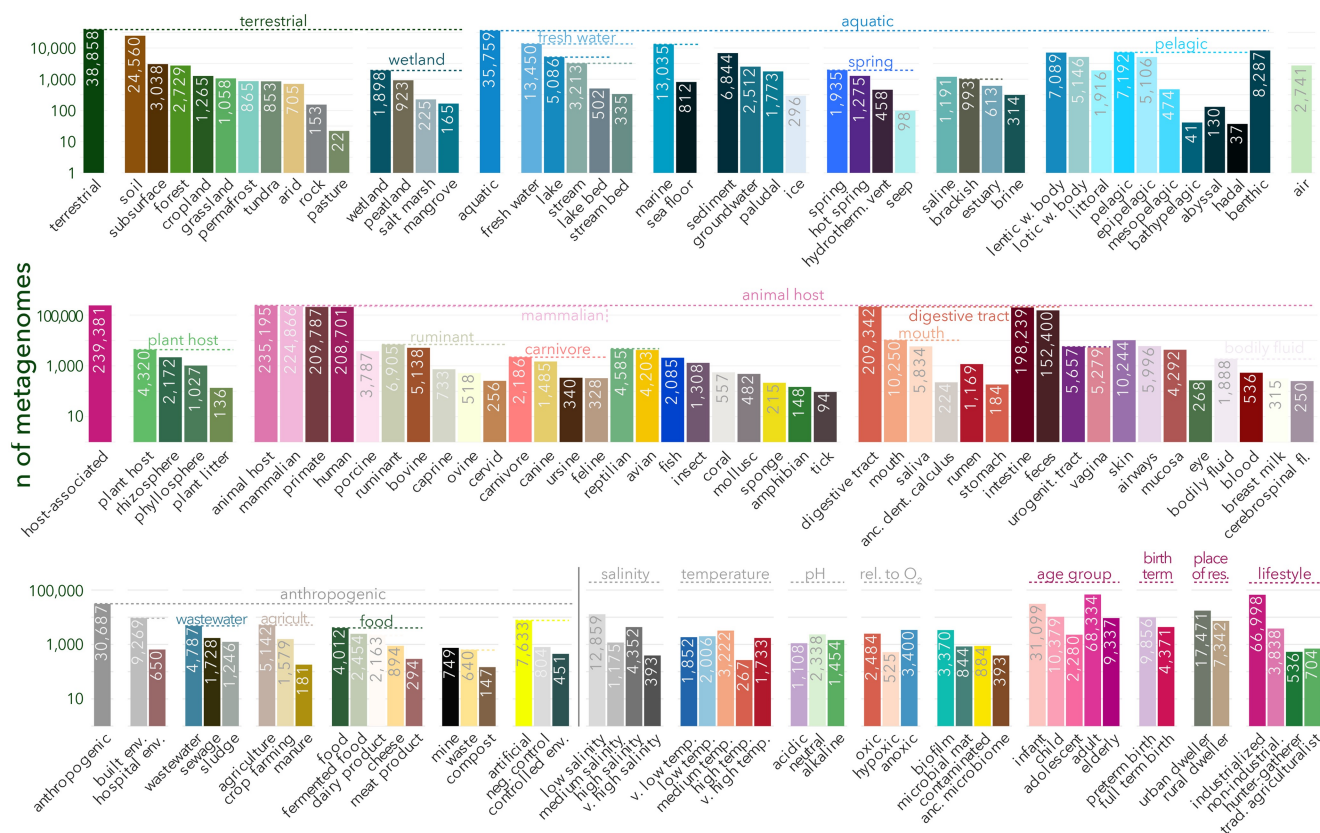


Figure 1 Overview of curated *microntology* annotations for 305 626 publicly available metagenomes. For each *microntology* term (categorical x axis), the number of annotated metagenomic samples (y axis) is shown. Categories are organized by sample group. Note that y axis scales differ between rows for improved visibility. As annotations follow hierarchical categories and individual samples are annotated with multiple tags (see main text), total counts do not sum to the total number of annotated metagenomes.

strongly biased towards animal-associated habitats (235k samples, or 77% of total), and in particular to human (209k, 68.5%) and intestinal (198k, 65%) samples. However, the representation of other body sites (such as mouth, skin, airways and urogenital tract), non-human hosts (in particular bovine, porcine and avian), but also of soils (25k samples), fresh water (13.5k), ocean (13k), the built environment (9k) and food-associated habitats (4k) has increased significantly compared to recent surveys (e.g. Schmidt et al. 2024). We note that for additional descriptive categories, such as salinity, temperature, human age group or place of residence, annotation recall is necessarily limited, as the corresponding information is often not available in public repositories, nor can it be extracted from publications. In such instances, we annotated conservatively, i.e. assigning only tags unambiguously supported by publicly available data, without extrapolation.

microntology annotations have previously been used to facilitate data exploration e.g. in SPIRE v1 (Schmidt et al. 2024) and the Global Microbial SmORFs Catalogue (Duan et al. 2024), to benchmark global-scale microbial habitat unsupervised taxonomic clusterings and modeling (Kim et al. 2026), and to categorize the main reservoirs of discoverable microbial diversity in existing data (Prasoodanan Pk et al. 2026). The resource is continuously developed and iteratively refined, following the basic principles laid out above, i.e. driven by data availability and user requirements in practice, relying on multiple parsimonious tags to describe

microbial habitats at an intermediate and pragmatic resolution. *microntology* is intended as a lightweight complement to established highly detailed ontologies such as EnvO (Buttigieg et al. 2016) or the GOLD Ecosystem Classification (Mukherjee et al. 2023) while retaining interoperable links, yet to provide broader coverage than project-specific vocabularies such as the Earth Microbiome Project (Thompson et al. 2017) or Microflora Danica ontologies (Singleton et al. 2026). By design, *microntology* describes microbial habitats at a resolution determined by the availability and scope of underlying data, in particular metagenomic samples. Therefore, *microntology* annotations can be browsed as a rough classification of datasets towards explorative analyses, e.g. to track microbial responses to rapidly changing conditions across Earth's habitats. Other resources such as curatedMetagenomicData (Pasoli et al. 2017) or Metalog (Kuhn et al. 2026) provide detailed descriptions of individual samples and their ecological niches with detailed contextual data. In summary, *microntology* annotations may facilitate studies of microbial ecology and evolution at large scales, integrated across Earth's microbial habitats and lifestyles.

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Author contributions

Anthony Fullam (Data Curation [equal], Methodology [equal], Validation [equal]), Vishnu Prasoodanan PK (Data Curation [equal], Formal Analysis [equal], Visualization [equal], Writing Review Editing [equal]), Michael Kuhn (Data Curation [equal], Resources [equal], Writing Review Editing [equal]), Peer Bork (Resources [equal], Supervision [equal], Writing Review Editing [equal]), and Thomas SB Schmidt (Conceptualization [lead], Data Curation [lead], Formal Analysis [equal], Methodology [equal], Project Administration [equal], Software [equal], Supervision [lead], Writing Original Draft [lead], Writing Review Editing [equal])

Supplementary material

Supplementary material is available at *Bioinformatics* online.

Conflict of Interests

None declared.

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