# **OpenMeta: A Comprehensive Multi-Task Benchmark** for Metagenomics Understanding

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# Abstract

Metagenomics is essential for exploring the vast diversity and intricate interactions 1 of microbes that impact health, agriculture, and environmental sciences. Despite 2 3 the surge of machine learning-based metagenomic models addressing these questions, evaluating their respective benefits is challenging due to the use of distinct, 4 5 experimental datasets, partly contrived, and varying model performance across different tasks. To this end, we introduce OpenMeta, the first comprehensive 6 benchmark tailored for metagenomic function prediction, which integrates diverse 7 datasets ranging from 1,000 to 213,000 sequences and incorporates hierarchical 8 data. We highlight the inadequacies of current genomic models and the superior 9 performance of metagenomic pre-trained models for handling complex metage-10 nomic data. Furthermore, we identify a critical research gap: the lack of unified 11 12 models that process both sequence and hierarchical data. Addressing this could significantly advance metagenomic analyses. OpenMeta sets a new standard for 13 metagenomic analysis, offering insights that could enhance the understanding and 14 application of microbial ecology in biotechnology and environmental science. 15

#### 16 **1** Introduction

Metagenomics is a discipline that studies the genetic composition and functional dynamics of all 17 microorganisms in environmental samples [41, 44]. By directly extracting the entire DNA from these 18 microorganisms, metagenomics captures a broad spectrum of life forms, including viruses, viroids, 19 and free DNA present in diverse habitats such as soil, seawater, and human microbiomes [53, 40]. 20 Unlike traditional genomics, which focuses on sequencing DNA from single species in isolation [57, 21 74], metagenomics eliminates the need for isolating each organism, allowing research of uncultivable 22 microorganisms [83, 48, 23]. Consequently, metagenomics unveils the vast diversity of microbial 23 24 communities, enabling the interpretation of gene interactions and essential biological processes within ecosystems [3, 37, 39]. 25

Deep learning techniques have significantly advanced metagenomics research, enabling more pre-26 cise function prediction and complex relationship elucidation in metagenomics [1, 60, 45, 77, 22, 27 38](Sec. 2). However, the field lacks standardized benchmarks, making it difficult to evaluate the 28 efficacy of various metagenomic models that often rely on distinct, artificially constructed datasets. 29 Existing genomic benchmarks primarily focus on single-species genomic analysis. The GUE bench-30 mark [84], built upon DNABERT2, encompasses multiple datasets ranging from humans to viruses 31 and includes 7 binary classification tasks like promoter detection and transcription factor prediction. 32 GenomicBenchmarks [49] address gene regulation and chromatin accessibility tasks. Nucleotide 33

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Figure 1: Framework of OpenMeta, including dataset preprocessing, data encoding, general and specialized models pre-trained or not, and target tasks.

Transformer (NT) [9], pre-trained on multi-species genomic data, concentrates on transcription factor binding and enhancer-promoter interactions within the human genome [32] (Sec. 3.1).

However, these methods often fail to address the unique challenges in metagenomics. *Firstly*, 36 metagenomic data is distinct as it comprises genomic collections from multiple organisms sequenced 37 simultaneously from numerous individuals, unlike genomic studies that focus on single organisms. 38 Secondly, metagenomic tasks are unique, involving the prediction of complex interactions among 39 various microorganisms, including bacteria and viruses, influenced by dynamic environments -40 a complexity that surpasses single-species genomic analysis. Lastly, existing genomic models 41 often perform poorly on metagenomic tasks, as models trained on genomic data cannot be directly 42 transferred to metagenomic data. Hence, there is an urgent need to develop a metagenomic reference 43 to deepen our understanding of microbial ecology and provide a powerful tool for biotechnology and 44 45 environmental science research (Sec. 3.2).

To this end, we develop OpenMeta, the first comprehensive benchmark specifically designed for 46 metagenomics based on the FGBERT model [13]. OpenMeta integrates a wide range of tasks across 47 48 genetic, functional, bacterial, and environmental levels and can handle datasets with sequences ranging from 1,000 to 213,000, as well as richer hierarchical data of phylogenetic tree structures, 49 reflecting the diversity and complexity of metagenomic data. We compare metagenomic pre-trained 50 models with genomic pre-trained models (Sec. 5.2), and although the latter provided new insights, 51 their performance on metagenome data tended to decrease, highlighting the need for the development 52 of metagenome pre-trained models. Our main contributions are as follows: 53

- i. We establish OpenMeta, the first comprehensive benchmark for metagenomic research encompassing 23 representative models. OpenMeta sets a new standard in the evaluation of metagenomic models using various collected standardized datasets and metrics across three dimensions: pre-trained vs. not pre-trained models, general vs. specialized models, and sequence
- <sup>58</sup> data-based vs. hierarchical data-based models.
- ii. We conduct extensive experiments on various tasks ranging from metagenomic sequences to
   hierarchical data, covering small-scale, large-scale, and fine-grained scopes.
- 61 iii. Our findings lead us to reconsider the potential of metagenomic pre-trained models, advocat 62 ing for architectures like FGBERT that better accommodate the diversity and complexity of
   63 metagenomic data.
- iv. We identify a significant research gap: the lack of a unified model capable of simultaneously
   processing sequence and hierarchical data. Addressing this could significantly advance compre-
- hensive metagenomic analyses and represent a promising direction for future research.

# 67 2 Related Work

Gene Representation Learning. For sequence metagenomic data, while the K-mer method [17] 68 efficiently captures characteristics of short sequences, it struggles with longer sequences due to 69 its inherent limitations. Alternatively, one-hot encoding, despite its high-dimensional and sparse 70 nature, restricts its utility for large-scale applications. In contrast, deep learning-based embeddings, 71 such as those from Transformer models, enhance sequence representation by capturing contextual 72 and global features, offering biologically meaningful insights. For hierarchical metagenomic 73 data, constructing phylogenetic trees provides an effective framework for delineating hierarchical 74 and evolutionary relationships among microbial taxa [2, 79]. This process begins with analyzing 75

microbial genomes through multiple sequence alignment, organizing them into a phylogenetic
structure. Microbial taxa abundances are then mapped to corresponding nodes on the tree. By
aggregating abundance values from child nodes to their respective parent nodes and transforming this
phylogenetic tree into a matrix format [18, 58], the structure is adapted for input into CNN models
for disease phenotype prediction.

Metagenomic Methods. Traditional alignment-based methods like MetaPhlAn5 [66] aim to match 81 similarities between query sequences and known reference genomes and are common for taxonomic 82 profiling. Advancements in deep learning have led to new methods like DeepVirFinder [61], which 83 use CNNs for viral classifications with one-hot encoding. K-mer tokenization [17], employed in 84 approaches like MDL4Microbiome [33], is a standard for DNA sequence characterization. Addi-85 tionally, Virtifier [45] maps a nucleotide sequence using a codon dictionary combined with LSTM 86 to predict viral genes. DeepMicrobes [35] employs a self-attention mechanism, while DeepTE [81] 87 uses K-mer inputs with CNNs for element classification, and Genomic-nlp [46] applies word2vec 88 89 for gene function analysis. MetaTransformer [77] uses K-mer embedding for species classification with Transformer. For pre-training models, LookingGlass [24] uses a three-layer LSTM model for 90 functional prediction in short DNA reads. ViBE [22] employs a K-mer token-based BERT model 91 pre-trained with Masked Language Modeling for virus identification. 92

Genomic Benchmark. To our knowledge, there is no benchmark in the field of metagenomics. 93 Due to the scarcity of specialized benchmarks in metagenomics and the inherent similarities in data 94 structure and content between genomic and metagenomic datasets, comparing the two allows us 95 to leverage advancements in genomics to address metagenomics' unique needs. Existing genomic 96 benchmarks, such as GUE, GenomicBenchmarks, and Nucleotide Transformer [9], each have their 97 distinct focus but primarily address single-species genome analysis. GUE, as part of DNABERT2 [9], 98 addresses challenges in genome tokenization and pre-training, covering multi-species datasets from 99 humans to viruses and involving tasks like promoter detection and transcription factor prediction. 100 GenomicBenchmarks, through HyenaDNA [49], focuses on improving long-sequence genome 101 modeling and handling ultra-long sequences, including gene regulation and chromatin accessibility 102 analysis. NT is pre-trained on multi-species genomic data, emphasizing transcription factor binding 103 and enhancer-promoter interaction in the human genome. 104

# **105 3 Background**

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#### 106 3.1 Difference Between Genomics and Metagenomics

The Main distinction between genomics 107 and metagenomics lies in the Number of 108 organisms evaluated in a sample in Tab. 1. 109 Genomics focuses on the genome of a sin-110 gle organism, whereas metagenomics ex-111 amines the collective genomes of differ-112 ent organisms within a sample [31, 63, 52]. 113 Sample type: Genomics targets the com-114 plete genetic information of a single or-115

ganism, typically from individual cells or

Table	1:	Differences	in	Metagenomics	and	Genomics.
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Comparison Factor	Metagenomics	Genomics
Main difference: Organisms number	Many	One
	Genomes from	Individual
Sample type	many individuals	organism's
	within an environment	genetic makeup
	Large, mixed data	Relatively small,
Data complexity	from multiple organisms	well-structured data
	mixed together	from single organism
Sequencing depth	3-100M Reads	3-6M Reads
Sample types	Stool, Skin, Soil, Water	Cell Culture
Cost	Higher	Lower

species, while metagenomics analyzes mixed DNA from multiple organisms in an environmental sam-117 ple, allowing scientists to study unculturable microorganisms [48, 23]. Data complexity: Genomic 118 data involves well-structured genetic information from a single organism, making it relatively simple. 119 Metagenomic data includes genomes from various organisms, leading to large, mixed, and complex 120 data. Sequencing depth: Metagenomic samples require significantly greater sequencing depth 121 than single-genome sequencing. Common sample types: Metagenomic samples are derived from 122 environments containing multiple genomes, such as soil, water, and human microbiomes, and are 123 inherently more complex than genomic samples, which typically come from single-cell cultures [15]. 124

Cost of sequencing: The extensive sequencing depth required for metagenomics generally results in
 higher costs than genomic sequencing.

#### 127 3.2 Necessity of Developing Metagenomic Benchmarks

Tab. 2 provides a detailed comparison of downstream tasks for genomic and metagenomic pre-128 trained models. DNABERT2 [84] primarily engages in binary classification tasks such as promoter 129 detection in human, mouse, and yeast species. Notably, Transcription Factor Prediction task recurs 130 identically for both human and mouse, indicating consistent yet singular difficulty, which may reduce 131 the overall challenge. HyenaDNA [49] is limited to regulatory elements classification tasks, reflecting 132 a narrow scope. Additionally, Demo and Dummy Datasets are typically used for initial testing, 133 lacking authenticity and practical value, reflecting the simplicity and limitations of its datasets. NT [9] 134 covers 18 downstream tasks centered on splice site prediction and chromatin accessibility analysis. 135

While these genomic benchmarks perform 136 well in single-species analyses, they of-137 ten fail to capture the inherent complexi-138 ties of multi-species interactions present 139 in metagenomics. Specifically, the limita-140 tions include (1) reliance on single-species 141 data, overlooking the complex interactions 142 in metagenomics; (2) lack of data diver-143 sity, insufficient environmental diversity re-144 quired for metagenomics; (3) limited func-145 tional prediction, focusing on sequence-146 based predictions without integrating cru-147 cial functional annotations; and (4) inade-148

Table 2: Comparison	of pre-trained	models f	or genome
and metagenome.			

	Model	Category	Task	#Class
			Transcription Factor Pred.	2
	DNADEDTO	Human	Promoter Detection	2
enome	DNABERIZ		Splice Site Detection	3
		Mouse	Transcription Factor Pred.	2
		Virus	Covid Variant Class.	9
		Human	Regulatory	2
9	HyenaDNA	Demo Dataset	Elements	2
		Dummy Dataset	Classification	2
		Yeast	Epigenetic marks Pred.	10
	NT	-	Splice site Pred.	2
		-	Chromatin Profiles Pred.	919
			Gene Structure Pred.	1379
eta.	ECDEDT	Mixed	ARG Pred. on Gene Family	269
Й	FUDERI	Multi-Species	Virulence Factor Pred.	15
			Pathogenic Genes Pred.	110

quate model adaptability, as models trained on single-species genomic data struggle to adapt to multi-species metagenomic data. In contrast, FGBERT [13], a metagenomic pre-trained model, aims to address interactions within different microbial communities. Its downstream tasks span multiple species, with a large number of classification categories, reflecting the diversity and complexity of metagenomic data. Therefore, incorporating FGBERT's multi-species metagenomic datasets into OpenMeta can enhance its ability to decipher complex microbial functions. For detailed analysis, please refer to Appendix B.

#### 156 4 OpenMeta

#### 157 4.1 Supported Methods

OpenMeta supports 23 methods for comprehensive analysis and performance evaluation across various tasks and data types, detailed in Tab. 3 with their respective publication years. To present them systematically, we categorize them along three dimensions: General vs. Specialized Models, Pre-trained vs. Not Pre-Trained Models, and Sequence data-based vs. Hierarchical data-based Models, acknowledging that some methods may span multiple categories.

**General vs. Specialized Models:** General models include widely used machine learning and deep learning models, such as SVM [69], Random Forest (RF) [64], CNN, LSTM [25], and Transformer [76], providing robust foundations for various tasks. Moreover, models like DNABERT2 [84] will be discussed in detail as pre-trained models in the next part. Conversely, specialized models are designed for specific metagenomic tasks such as functional gene prediction (PLM-ARG [78], DeepARG [5], RGI [4], ViBE [22], CLEAN [82]), and prototype prediction (PopPhy-CNN [59]).

Pre-trained vs. Not Pre-trained Models: OpenMeta includes DNABERT2 [84], HyenaDNA [49],
and NT [9] genomic pre-trained models, alongside FGBERT [13] metagenomic pre-trained model.
Although ViBE [22] and PLM-ARG [78] are not pre-trained from scratch, they use BERT model [11]
and protein language model [36], respectively, to enhance their functional prediction capabilities.

Table 3: Categorizations of all supported metagenomic methods in our OpenMeta. RF denotes Random Forest, and VT represents Vanilla Transformer.

Model	Pre- Trained	Not Pre- Trained	General	Specialized	Sequence- based	Hierarchical- based	Multi- Classification	Binary- Classification	Year	Description
RF		1	1		1		1			Machine Learning
SVM		1	1		1		1			Machine Learning
AdaBoost		1	1		1		1			Machine Learning
LSTM		1	~		1		1			Deep Learning
BiLSTM		1	1		1		1			Deep Learning
VT [76]		1	~		1		1			Deep Learning
FGBERT [13]	1		~		1		1		2024	Metagenomic pre-trained model for functional prediction.
HyenaDNA [49]	1		1		1		1		2023	Genomic pre-trained model trained on multi-species genomes.
NT [9]	1		1		1		1		2023	Genomic pre-trained model trained over human reference genome.
DNABERT2 [84]	~		~		1		1		2023	Genomic pre-trained model trained on diverse human genomes.
CNN-MGP [1]		1		1	1			1	2019	Gene prediction using CNN network.
PlasGUN [16]	,	~		1	1		,	~	2020	Gene prediction tool using multiple CNN network.
PLM-ARG [78]	~	,		1	1		1		2023	ARG identification framework using a pretrained protein language model.
DeepAKG [5]		1		1	×,		1		2018	ARG prediction software by alignment and metagenomic sequences.
KGI [4] Daan Vin Eindan [61]		1		1	1		~	,	2023	Viral accurate mediation with reference and alignment from CNNs
Van (22)	/	~		1	×,		,	~	2020	Fully sequences prediction with reference and anglinent-free CNNs.
VIDE [22] ViroMinor [71]	•	1		·,	· /		•	1	2022	Viral ganamas identification in human samples
DeepVE [90]		· /		·,	· /			· /	2019	Viral factor identification with hybrid framework using stacking strategy
HyperVP [27]		1		1	1			1	2021	Viral factors and mixing of APG simultaneous prediction
CLEAN [82]		1		1	1		1	v	2023	Enzyme function prediction using contrastive learning
DeenMicrobes [35]		1		1	1		v	1	2023	Taxonomic classification for metagenomics with self-attention model
PopPhy-CNN [59]		1		1	•	1	1	•	2020	Host Phenotypes prediction by systematic tree embedded CNN network.

173 Sequence Data-based vs. Hierarchical Data-based Models: This work further integrates models trained on hierarchical data, such as PopPhy-CNN [59], which leverages the phylogenetic tree structures of microbial communities to enhance the understanding of microbial interactions, con-

trasting with sequence-based models essential for raw genetic data analysis without considering the

177 hierarchical relational structure among microbial taxa.

#### 178 4.2 Supported Tasks

Table 4: Detailed information of metagenomic sequence datasets in OpenMeta.

Туре	Dataset	Category	#Seq	ŀ	#Cat	e.	Seqs/ (Min	Cate -Max	Range	Avg	. Len.	Description
	E-K12 [65]	Gene-pair Cls.	4,315	5	1,379	)	1-106	5		510	.96	Tasks involving
Small Saala	CARD-A [28]		1,966	5	269		1-229	)		108	8.1	smaller datasets
Classification	CARD-D [28]	Cana usiaa Cla	1,966	5	37		1-513	;		108	8.1	focusing on high
Classification	CARD-R [28]	Gene-wise Cis.	1,966	5	7		1-126	53		108	8.1	accuracy in
	PATRIC [19]		5,000	)	110		1-108	31		307	.82	narrow contexts.
Laura Carla	ENZYME [6]		5,761	1	7		288-2	2055		426	.76	Tasks requiring handling
Large-Scale	VFDB [7]	Gene-wise Cls.	8,945	5	15		5-168	3		415	.47	of large data volumes,
Classification	NCycDB [75]		219,0	219,089 69			1-205	548		347.03		broad pattern extraction.
	NCRD-N [42]		104,3	363	1912		1-183	70		407	.44	Focused on detailed
Fine-Grained	NCRD-F [42]		104,3	363	420		2-353	64		407	.44	differentiation within
Classification	NCRD-C [42]	Gene-wise Cis.	104,3	363	29		1-141	59		407	.44	closely related
	NCRD-R [42]		104,3	363	10		166-3	8073		409	.79	categories.
Tab	le 5: Detaile	d informatio	n of m	neta	Igeno	mi	c hie	rarc	hical	datas	sets in	openMeta.
T	Defend	C. A.	#G			Tax	onom	ic Lev	vels		D	
Туре	Dataset	Category	#Seq.	Κ	Р	С	0	F	G	S	Desci	ription
Hierarchical	Cirrhosis [56]	Metagenome	542	3	15	27	40	76	186	531	Detail divers	led profiling of microbial sity linked to cirrhosis.
Classification	T2D [30]	-wise CIS.	606	3	17	29	48	94	216	587	Detail Diabe	led analysis for Type 2

We conduct extensive experiments across various multi-classification tasks, including gene structure
analysis, functional gene prediction, pathogenicity assessment, nitrogen cycle prediction, and disease
phenotype prediction, utilizing diverse datasets that range from metagenomic sequences to hierarchical data We provide detailed descriptions of the following 14 datasets in Tab. 4 and 5, covering
small-scale, large-scale, and fine-grained scopes. Seqs/Cate Range provides the range of sequences
in each category, from minimum to maximum. Details can be found in Appendix C.

(1) Small-Scale Classification: Gene Operon Prediction Task. This task aims to identify transcrip-185 tion factor binding sites with the strongest correlation with operon regulation in the gene regulatory 186 network [8, 14, 51] This gene-pair classification utilizes E-K12 dataset [65], consisting of 4.315 oper-187 ons, each detailed with operon names, descriptions, and gene components. Antibiotic Resistance 188 Genes (ARGs) Prediction Task. Accurate identification of ARGs is essential for understand-189 ing the relationship between the microbiome and disease, as pathogenic microorganisms threaten 190 public health by exacerbating ARGs to invade the host [50]. This gene-wise classification uses 191 CARD dataset [28], categorizing genes by 269 AMR Gene Families (CARD-A), 37 Drug Classes 192

Table 7: Enzyme function Pre-<br/>diction on ENZYME.Table 8: Virus factor Prediction<br/>on VFDB.Table 9: N Cycling Prediction<br/>on NcycDB

			UEDD	. 0	п неусвв.	
Method	ENZYME	Method	VFDB	_	Method	NCycDB
RF (3-mer)	33.6	RF (3-mer)	22.4			-
SVM (3-mer)	31.3	SVM (3-mer)	28		RF (3-mer)	67
AdaBoost (3-mer)	31.4	AdaBoost (3-mer)	27.3		SVM (3-mer)	66.9
I STM (w2v)	42.8	$I STM (w^2 y)$	36.7		AdaBoost (3-mer)	68.8
LSTM (one-hot)	34.1	LSTM (w2v)	32.9		LSTM (w2v)	71.9
BI STM (w2y)	29.7	Dil STM (w2w)	46.1		LSTM (one-hot)	65
BiLSTM (w2v)	21.6	BiLSTM (w2v) BiLSTM (one hot)	21.2		BiLSTM (w2v)	66.9
DILSTIM (one-not)	31.0	DILSTM (one-not)	27.7		BiLSTM (one-hot)	82
BILSTM-Att. (W2V)	30.9	BILSTM-Att. (W2V)	31.1		BiLSTM-Att. (w2v)	69
BiLSIM-Att. (one-hot)	43.6	BiLSIM-Att. (one-not)	36.7		BiLSTM-Att. (one-hot)	67.3
VT	68.2	VT	58		VT	84 5
HyenaDNA	79.6	HyenaDNA	61.1		HvenaDNA	92.4
NT	74.1	NT	58.3		NT	75.1
DNABert2	85.4	DNABert2	58.2		DNAPort?	996
FGBERT	99.1	FGBERT	75.7		ECDEDT	00.5
CLEAN	92.3	ViBE	50.9		FUDERI	99.5

<sup>193</sup> (CARD-D), and 7 Resistance Mechanisms (CARD-R). **Pathogens Prediction Task.** This task <sup>194</sup> assesses the pathogenic potential of pathogens to cope with the public health risks [29]. We use

195 <u>PATRIC core dataset</u> [19], which has 5000 pathogenic bacterial sequences across 110 classes.

(2) Large-Scale Classification: Enzymes Prediction Task. Enzymes are important catalysts in living 196 cells that produce essential molecules needed by living organisms through chemical reactions [73]. 197 ENZYME dataset [6] contains 5,761 enzyme sequences, which are grouped into 7 classes according 198 to their corresponding EC numbers. Virulence Factors (VFs) Prediction Task. Viruses are common 199 in both humans and different habitats, and they are always changing. Therefore, accurately identifying 200 VFs is extremely important for understanding the relationship between the microbiome and disease. 201 VFDB dataset [7] for virulence factors prediction contains 8,945 VF sequences across 15 categories, 202 detailing structural features, functions, and mechanisms of major bacterial pathogens. Nitrogen (N) 203 Cycling Process Prediction Task. The N cycle is a collection of important biogeochemical pathways 204 in the Earth's ecosystems, and quantitatively studying the functional genes related to the N cycle [20]. 205 NCycDB dataset [75] contains 68 genes (sub)families and covers 8 N cycle processes with 219,089 206 representative sequences, each involving a specific gene family. 207 (3) Fine-Grained Classification: ARG Prediction. Targets a more precise and detailed prediction of 208

antibiotic resistance properties, which helps in exploring ARG characteristics comprehensively and
 detecting potential resistant mechanisms [34]. <u>NCRD dataset</u> [42] is dedicated to the fine-grained
 categorization of microbial resistance genes, differentiating in detail between 420 <u>Gene Families</u>,
 1,912 specific <u>Gene Names</u>, 30 major <u>Resistance</u>, and 10 different <u>Mechanisms</u>.

(4) Hierarchical-Data Classification: Disease Prediction Task. Predicting host phenotypes and identifying relevant markers are pivotal for unraveling the complexities of host-microbiome interactions [72, 55], and the impact of such interactions on disease [26, 43, 47] can be explored using the phylogenetic structure and relative abundance of microbial taxa [21]. <u>Cirrhosis dataset</u> [56] comprises
232 data cases on microbiome liver disease. <u>Type 2 Diabetes (T2D) dataset</u> [30] comprises 440 data

218 cases on glucose metabolism disorders.

#### 219 4.3 Evaluation Metrics

For multi-classification tasks, we use the Macro F1-220 score (M.F1) as the primary metric to accommodate 221 the inherent class imbalance present within datasets. 222 For the Fine-grained Benchmark, Accuracy, Preci-223 sion, Recall, and False Negative Rate (FNR) are in-224 corporated. FNR is particularly critical for ARG 225 prediction scenarios where the consequences of over-226 looking true positives are severe, necessitating nu-227 anced assessments of the model's ability to identify 228 them reliably. 229

Table 6: Comparison of ARG prediction meth-
ods on CARD. (- means inability to predict
specific category).

Mathad	ARG Prediction						
Method	CARD-A	CARD-D	CARD-R				
RF (3-mer)	22.4	36.1	47.8				
SVM (3-mer)	27.6	33.6	43.3				
AdaBoost (3-mer)	36.9	36.4	36.2				
LSTM (w2v)	47.1	37.5	47.5				
LSTM (one-hot)	46.2	39.1	41.5				
BiLSTM (w2v)	43.3	35.5	36.3				
BiLSTM (one-hot)	47.4	38.9	58.9				
BiLSTM-Att. (w2v)	31.9	43.5	35.1				
BiLSTM-Att. (one-hot)	46.7	31.2	41.6				
VT	57.1	49.8	55.7				
HyenaDNA	50.9	53.6	66.2				
NT	58.5	56.2	68				
DNABERT2	65.2	51.5	61.2				
FGBERT	78.6	57.4	69.4				
DeepARG	-	52.2	65.3				
PLM-ARG	-	-	68.1				
RGI	-	-	-				

# 230 5 Results and Insights

#### 231 5.1 Results

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Small-Scale Benchmarks. Appendix Tab. A5 and A6 show M.F1 for gene operon and pathogen 232 prediction on two small-scale datasets, E-K12 and PATRIC (sequence length less than 5000). General 233 models (RF to VT) test K-mer (K=3), one-hot, and w2v data encoding methods, and BiLSTM 234 (one-hot) and LSTM (w2v) performed best. FGBERT achieves superior results, far exceeding 235 other methods, highlighting the necessity of metagenomic pre-training. Models like CNN-MGP [1], 236 PlasGUN [16], and DeepMicrobes [35], designed for binary classification, are unsuitable for our 237 multi-classification benchmark. Tab. 6 evaluates various models on three ARG prediction datasets: 238 CARD-A, CARD-D, and CARD-R. In General models, the word2vec data encoding method performs 239 better. FGBERT outperforms HyenaDNA, NT, and DNABERT2. This suggests that genomic models 240 may not be sufficient to cope with the complexity of metagenomic data, which involves multiple 241 microbial interactions and environments. In Specialized models, DeepARG [5] performs well on 242 CARD-D and CARD-R but unable to predict CARD-A. Since RGI [4] itself is based on CARD 243 dataset, it is not included. 244

Large-Scale Benchmarks. Tab. 7, 8and 9 245 summarize M.F1 across three large-scale 246 datasets (sequence length more than 5000): EN-247 ZYME, VFDB, and NCycDB, the analysis re-248 flects a similar trend. FGBERT demonstrates 249 exceptional efficacy across all datasets, signifi-250 251 cantly outperforming specialized models such as ViBE [22] and CLEAN [82]. Conversely, 252 models designed for binary classification, such 253 as ViraMiner [71], deepVF [80], HyperVR [27] 254 and DeepVirFinder [61] do not align well with 255 our benchmark's requirements. However, Hy-256 perVR's innovative approach to predicting VF 257 and ENZYME concurrently has inspired poten-258

tial developments in OpenMeta for simultaneous



Figure 2: Comparison of different data encoding methods across tasks.

multi-task predictions. Fig. 2 compares different data encoding methods. Details can be found in
 Appendix E. From K-mer to language model representation, model performance gradually improves,

indicating that capturing contextual information in the sequence is important.

Fine-Grained Benchmarks. The fine-grained 263 NCRD dataset provides a more rigorous test for ARG 264 prediction tasks. As depicted in Fig. 3, FGBERT 265 performs well in all resistance categories, identify-266 ing 30 antibiotic classes. In comparison, DeepARG 267 identified 17, and RGI identified 22, with the unde-268 tected classes shown in light gray. Moreover, the 4 269 methods have high accuracy in beta-lactam, amino-270 glycoside, and multidrug categories. Furthermore, 271 Tab. 11 shows that FGBERT and RGI cover all NCRD 272 classes, while DeepARG is limited to specific classes, 273 likely due to limitations in its training data. PLM-274 ARG has a high false negative rate, as shown in 275 Tab. 10, indicating its limitations in application. 276



Figure 3: Comparison of different methods on each ARG category.

277 Hierarchical-Data Benchmark. In the host phe-

<sup>278</sup> notype prediction task, we use microbial taxa's relative abundance values as input. Since abundance

Table 10: Comparison of M.F1 for ARG Prediction on different categories of NCRD Datasets. (– indicates inability to predict specific categories. FNR is the false negative rate, lower is better).

Mathad	N	CRD-G	ene Fa	mily (42	20)	N	CRD-G	ene Na	me (190	)0)		NCRD	Resista	ance (30	))	N	ICRD-N	Mechan	isms (1	1)
Method	Acc.	Pre.	Re.	F1	FNR	Acc.	Pre.	Re.	F1	FNR	Acc.	Pre.	Re.	F1	FNR	Acc.	Pre.	Re.	F1	FNR
DeepARG	-	-	-	-	-	0.63	1.00	0.43	0.60	0.57	0.97	0.99	0.51	0.65	0.42	-	-	-	-	-
RGÎ	0.56	0.97	0.36	0.50	0.61	0.50	1.00	0.26	0.42	0.74	0.56	1.00	0.36	0.52	0.64	0.61	0.62	0.59	0.57	0.51
PLM-ARG	-	-	-	-	-	-	-	-	-	-	0.96	0.87	0.74	0.88	12.38	-	-	-	-	-
FGBERT	0.94	0.96	0.94	0.93	0.03	0.93	0.93	0.92	0.93	0.05	0.99	0.99	0.99	0.99	0.31	0.99	0.99	0.98	0.99	0.01

Table 14: Comparative Analysis of Pre-training Strategies in Genomic and Metagenomic Models.

Model	Pre-training Dataset	Token	Network Architecture	Application Tasks	Benchmark
DNABERT2	Human and multi-species genome	BPE	Advanced BERT with MLM+ ALiBi	Promoter detection, Transcription factor, binding site prediction	28 datasets on GUE benchmark
HyenaDNA	Human reference genome	6-mer	Simple stack of Hyena operators with NTP	Gene regulation prediction, chromatin accessibility analysis	8 datasets on GenomicBenchmarks +18 prediction tasks on NT
NT	Human reference genome	6-mer	Encoder-only Transformer + RoPE	Transcription factor binding, enhancer-promoter interaction prediction	18 prediction tasks
FGBERT	Multi-species metagenome	Protein-based genomic representation	Advanced BERT with contrastive learning	Metagenomic sequences and functions analysis	14 datasets on Metagenomic benchmark

data alone cannot reveal the hierarchical structure among species and introduces data redundancy, we 279 adopt a phylogenetic tree-based modeling approach to process abundance data [12], effectively reduc-280 ing redundancy and retaining species information. After constructing a phylogenetic tree through 281 multiple sequence alignment, abundance values are filtered and assigned to the tree's nodes, and the 282 values of child nodes are summed to their parent nodes. Finally, the phylogenetic tree is converted 283 into a matrix format for analysis. Tab. 12 shows that the specialized model PopPhy outperforms the 284 general models on Cirrhosis dataset. LSTM and Transformer models are not tested because they 285 are mainly applicable to sequence data and have difficulty capturing the hierarchical structure and 286 phylogenetic relationships between species. At present, no model can process both metagenomic 287 sequence and hierarchical phylogenetic tree data, indicating a key direction for future research. 288

#### 289 5.2 Observations and Insights

(A) Metagenomic Pre-trained Models vs. Genomic 290 Pre-trained Models: Tab. 14 compares genomic 291 and metagenomic pre-trained models, including pre-292 training datasets, token embedding methods, network 293 architectures, application tasks, and benchmarks. In 294 terms of Pre-Training Datasets, DNABERT2 [84] 295 utilizes human and multi-species genomes for its 296 foundational model pre-training, covering a vast 297 dataset of 27.5 billion nucleotide bases from the Hu-298 man reference genome [32] and 135 species genomes 299 across seven categories. HyenaDNA [49], on the 300 other hand, is pre-trained solely on a single hu-301 man reference genome. NT [9] pre-trains on three 302 datasets: the human reference genome [32], 3,202 303 diverse human genomes, and 850 genomes from sev-304 eral species. In contrast, FGBERT [13] employs 305 MGnify database [62], comprising 2,973,257,435 306 metagenomic sequences from various microbial com-307 munities. For Token Embedding, DNABERT2 ap-308 plies Byte Pair Encoding (BPE) [67], while both Hye-309 naDNA and NT use 6-mer tokenization. FGBERT 310

Table 11: Comparisons of different ARG prediction methods on NCRD.

Method	Categories of NCRD Dataset								
wiethou	Gene Family	Gene Name	Resistance	Mechanisms					
DeepARG		1	1	1					
RGI	1	1	1	1					
PLM-ARG			1						
FGBERT	1	~	1	1					

Table 12: Performance of General vs. Specialized Models on Cirrhosis dataset.

Method	<b>M.F1</b>	AUC	MCC	Pre.	Re.
RF	0.79	0.93	0.61	0.88	0.87
SVM	0.77	0.89	0.57	0.85	0.84
AdaBoost	0.70	0.71	0.43	0.72	0.71
CNN	0.84	0.89	0.68	0.81	0.80
PopPhy	0.81	0.90	0.61	0.83	0.82

Table 13: Performance of General vs. Specialized Models on T2d dataset.

Method	<b>M.F1</b>	AUC	MCC	Pre.	Re.
RF	0.66	0.72	0.33	0.67	0.67
SVM	0.61	0.63	0.23	0.61	0.61
AdaBoost	0.70	0.70	0.42	0.70	0.70
CNN	0.59	0.65	0.19	0.60	0.59
PopPhy	0.58	0.64	0.18	0.59	0.58

utilizes a unique protein-based genomic representation tailored for metagenomic sequences. Regarding Network Architecture, both DNABERT2 and FGBERT adopt BERT-like structures [11];
DNABERT2 enhances its predecessor by replacing learned positional embeddings with Attention with
Linear Biases (ALiBi) [54] to eliminate input length limitations and incorporates Flash Attention [10]

to boost computational efficiency. FGBERT introduces contrastive learning to strengthen the intricate
relationships between metagenomic sequences and functions, proposing two pre-training tasks to
enhance co-representation learning of metagenomic gene sequences and functions. HyenaDNA
employs a simple stack of Hyena operators for next token prediction, while NT uses an encoder-only
Transformer architecture with Rotary Positional Embeddings (RoPE) [70] to enable reasoning over
longer sequences during training.

(B) Sequence Data vs. Hierarchical Data: 321 Why Use Hierarchical Data? Hierarchical 322 data introduce an additional dimension by pro-323 viding interrelationships and evolutionary con-324 text among microbial communities, enriching 325 326 metagenomic research. Unlike traditional abundance data, hierarchical data offer not only 327 328 quantitative information but also capture the complex hierarchical relationships between mi-329



Figure 4: Phylogenetic Tree Representation of Microbial Communities for Hierarchical Data.

crobes, which is crucial for exploring host-microbe interactions [68].

Why Use Phylogenetic Tree Structures for Hierarchical Data? Tree structures naturally represent 331 the hierarchical and phylogenetic relationships among microbial taxa. Each node represents a 332 microbial taxon, and the connections between nodes reflect their evolutionary relationships. This 333 helps to reveal the evolutionary links between different microbial taxa, integrating complex biological 334 information (such as abundance and hierarchical data) into a unified data structure. By accumulating 335 the abundance values from child nodes to their parent nodes and converting the phylogenetic tree into 336 a matrix format, each row represents a level in the tree, and columns represent different microbes 337 or attributes. As shown in Fig. 4, this matrix-based representation effectively combines abundance 338 339 and hierarchical information. This approach is particularly useful in disease prediction tasks, such as studies on Cirrhosis and T2D, demonstrating how understanding the hierarchical structure of 340 microbial communities can elucidate the complexity of host-microbe interactions. This hierarchical 341 method provides powerful tools for the precise identification and functional analysis of disease-related 342 microbial communities. Our benchmark framework underscores the importance and benefits of using 343 hierarchical data to enhance the accuracy and depth of metagenomic analysis. 344

### 345 6 Conclusion and Future Work

**Conclusion.** In this paper, we introduce OpenMeta, the first comprehensive benchmark tailored for 346 metagenomic function prediction. This benchmark standardizes the evaluation process across various 347 metagenomic tasks and facilitates the design of metagenomic models through a unified approach. 348 Our extensive analysis includes comparisons between pre-trained and not pre-trained models, general 349 versus specialized models, and sequence data-based versus hierarchical data-based models. Inspired 350 by OpenMeta, we emphasize the necessity of pre-trained metagenomic models in this field and 351 advocate for the community's engagement with metagenomic models trained on hierarchical data 352 such as phylogenetic trees. This approach can profoundly enhance our understanding of the complex 353 relationships and interactions within microbial communities. 354

Limitations. It is crucial to note that OpenMeta primarily serves as an evaluative tool that aggregates and assesses a wide array of multi-class datasets, including both sequence and hierarchical data. While this benchmark significantly contributes to the field, it does not involve the development of new models but focuses on the assessment of existing methodologies. This limitation underscores the necessity for further research and development in creating comprehensive models that can process both sequence and hierarchical inputs simultaneously.

**Future Work.** We identify a significant gap in the current landscape: the absence of a unified metagenomic model capable of simultaneously processing sequence and hierarchical data from phylogenetic trees. Addressing this gap represents a promising direction for future work and could significantly advance our holistic understanding of metagenomics.

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# 585 Checklist

The checklist follows the references. Please read the checklist guidelines carefully for information on how to answer these questions. For each question, change the default **[TODO]** to **[Yes]**, **[No]**, or [N/A]. You are strongly encouraged to include a **justification to your answer**, either by referencing the appropriate section of your paper or providing a brief inline description. For example:

- Did you include the license to the code and datasets? [Yes] See Section E.
- Did you include the license to the code and datasets? [No] The code and the data are proprietary.
- Did you include the license to the code and datasets? [N/A]

Please do not modify the questions and only use the provided macros for your answers. Note that the Checklist section does not count towards the page limit. In your paper, please delete this instructions block and only keep the Checklist section heading above along with the questions/answers below.

597	1. For all authors
598 599	(a) Do the main claims made in the abstract and introduction accurately reflect the paper's contributions and scope? [Yes]
600	(b) Did you describe the limitations of your work? [Yes] See section 6
601 602	(c) Did you discuss any potential negative societal impacts of your work? [Yes] See section A
603 604	(d) Have you read the ethics review guidelines and ensured that your paper conforms to them? [Yes]
605	2. If you are including theoretical results
606	(a) Did you state the full set of assumptions of all theoretical results? [N/A]
607	(b) Did you include complete proofs of all theoretical results? [N/A]
608	3. If you ran experiments (e.g. for benchmarks)
609 610	(a) Did you include the code, data, and instructions needed to reproduce the main experi- mental results (either in the supplemental material or as a URL)? [Yes]
611 612	(b) Did you specify all the training details (e.g., data splits, hyperparameters, how they were chosen)? [Yes]
613 614	(c) Did you report error bars (e.g., with respect to the random seed after running experi- ments multiple times)? [Yes]
615 616	(d) Did you include the total amount of compute and the type of resources used (e.g., type of GPUs, internal cluster, or cloud provider)? [Yes]
617	4. If you are using existing assets (e.g., code, data, models) or curating/releasing new assets
618	(a) If your work uses existing assets, did you cite the creators? [Yes]
619 620	(b) Did you mention the license of the assets? [No] The used coda and datasets are all open-source.
621	(c) Did you include any new assets either in the supplemental material or as a URL? [Yes]
622 623	(d) Did you discuss whether and how consent was obtained from people whose data you're using/curating? [Yes]

624 625	(e) Did you discuss whether the data you are using/curating contains personally identifiable information or offensive content? [N/A] The datasets are all undergone ethical review
626	5. If you used crowdsourcing or conducted research with human subjects
627	(a) Did you include the full text of instructions given to participants and screenshots, if
628	applicable? [N/A]
629	(b) Did you describe any potential participant risks, with links to Institutional Review
630	Board (IRB) approvals, if applicable? [N/A]
631	(c) Did you include the estimated hourly wage paid to participants and the total amount
632	spent on participant compensation? [N/A]

# 633 Supplement Material

# 634 A Social Impacts

Metagenomics offers significant benefits in fields like medicine and environmental science, yet it also 635 poses dual-use concerns. For instance, technologies designed to minimize disease can theoretically 636 be repurposed for harmful uses, such as biological weapons. The advancement of metagenomic 637 benchmarks could inadvertently facilitate such misuse. Additionally, while these technologies can 638 accelerate experimental processes, the need for traditional wet lab experimentation remains crucial. 639 Thus, developing robust, clinically validated benchmarks is essential for integrating metagenomic 640 methods into medical practice responsibly. This approach will ensure technological advances support 641 health and environmental management without replacing foundational experimental techniques. 642

# 643 **B** Necessity of Developing Metagenomic Benchmarks

Table A1: Comparison of Downstream Tasks for Genomic and Metagenomic Pre-trained Models. The repetition of Transcription Factor Prediction tasks in the human and mouse categories suggests that it may not present significant challenges. Demo and Dummy datasets are usually artificially generated.

Model	Category	Dataset	Task	#Seq.	#Class
			Core Promoter Detection (3)	4,904	2
	Uumon		Transcription Factor Pred. (5)	32,378	2
	пишан		Promoter Detection (3)	4,904	2
DNABERT2			Splice Site Detection (1)	36,496	3
	Mouse		<b>Transcription Factor Pred. (5)</b>	6,478	2
	Yeast		Epigenetic Marks Pred. (10)	11,971	2
	Virus		Covid Variant Class. (1)	77,669	9
		Enhancers Cohn		27,791	2
		Enhancers Ensembl		154,842	2
	Human	Regulatory		289,061	3
HyenaDNA		Nontata Promoters	Description: Elemente Class	36,131	2
		OCR Ensembl	Regulatory Elements Class.	174,756	2
	Demo	Coding vs Intergenomic	-	100,000	2
	Demo	Human vs Worm		100,000	2
	Dummy	Mouse Enhancers		1,210	2
	Yeast		Epigenetic marks Pred. (10)	25,953	10
	-		Promoter sequence Pred. (3)	59,194	2
NT	-		Enhancer sequence Pred. (2)	14,968	3
111	-		Splice site Pred. (3)	19,775	2
	-		Chromatin Profiles Prediction (1)	-	919
	-		Enhancer Activity (1)	14,968	50
	Mul. Spe.	E. coil K12	Gene Structure Pred.	4,315	1379
	Mul. Spe.	CARD-A	ARG Pred. on AMR Gene Family	1,966	269
	Mul. Spe.	CARD-D	ARG Pred. on Drug Class	1,966	37
ECREPT	Mul. Spe.	CARD-R	ARG Pred. on Resistance Mechanism	1,966	7
IODEKI	Mul. Spe.	VFDB	Virulence Factor Pred.	8,945	15
	Mul. Spe.	ENZYME	Enzyme Function Pred.	5,761	7
	Mul. Spe.	PATRIC	Pathogenic Genes Pred.	5,000	110
	Mul. Spe.	NCycDB	N Cycling Genes Pred.	213,501	68

In our analysis of applications and benchmarks, Table A1 provides a detailed comparison of down-644 stream tasks for genomic and metagenomic pre-trained models, highlighting datasets, tasks, sequence 645 numbers, and class counts. DNABERT2 [84] primarily engages in binary classification tasks such as 646 promoter and splice site detection in human, mouse, and yeast datasets. The numbers in parentheses 647 indicate the number of independent sub-datasets for each task, and the sequence count reflects the 648 size of the first sub-dataset. Notably, the Transcription Factor Prediction task recurs identically 649 for both human and mouse species, suggesting a uniform difficulty level and potentially reducing 650 the challenge due to its repetition across similar species. HyenaDNA [49]'s downstream tasks are 651 divided into two parts: GenomicBenchmarks, which includes 8 regulatory element classification 652 datasets with sequence lengths ranging from 200 to 500, and NT's 18 prediction tasks. Beyond human 653

Table A2: The detailed information of supported datasets in OpenMeta with source link.

Model	Pre-Trained	Not Pre- Trained	General	Specialized	Sequence-based	Structure-based	Year	Link	Description
SVM		51	51		51				Machine Learning
RF		51	51		51				Machine Learning
AdaBoost		51	51		51				Machine Learning
CNN		51	51		51				Deen Learning
LSTM		51	51		51				Deep Learning
Vanilla Transformer		51	51		51				Deep Learning
FGBERT	51		51		51		2024		Metagenomic pre-trained model for functional prediction.
DNABERT2	51		51		51		2023	https://github.com/MAGICS-LAB/DNABERT_2	Genomic pre-trained model trained on multi-species genomes.
HyenaDNA	51		51		51		2023	https://github.com/HazyResearch/hyena-dna	Genomic pre-trained model trained over human reference genome.
Nucleotide Transformer	51		51		51		2023	https://github.com/instadeepai/nucleotide-transformer	Genomic pre-trained model trained on diverse human genomes.
CNN-MGP		51		51	51		2019	https://github.com/rachidelfermi/cnn-mgp	Gene prediction using CNN network.
PlasGUN		51		51	51		2020	https://github.com/zhenchengfang/PlasGUN	Gene prediction tool using multiple CNN network.
PLM-ARG	51			51	51		2023	https://github.com/Junwu302/PLM-ARG	ARG identification framework using a pretrained protein language model.
DeepARG		51		51	51		2018	https://github.com/gaarangoa/deeparg	ARG prediction software by alignment and metagenomic sequences.
RGI		51		51	51		2023	https://github.com/arpcard/rgi	ARG prediction tools for annotating genes from scratch.
DeepVirFinder		51		51	51		2020	https://github.com/jessieren/DeepVirFinder	Viral sequences prediction with reference and alignment-free CNNs.
ViBE	51			51	51		2022	https://github.com/DMnBI/ViBE	Eukaryotic viruses identification with hierarchical BERT model.
ViraMiner		51		51	51		2019	https://github.com/NeuroCSUT/ViraMiner	Viral genomes identification in human samples.
DeepVF		51		51	51		2021	http://deepvf.erc.monash.edu/	Viral factor identification with hybrid framework using stacking strategy.
HyperVR		51		51	51		2023	https://github.com/jiboyalab/HyperVR	Viral factors and mixing of ARG simultaneous prediction.
CLEAN		51		51	51		2023	https://github.com/tttianhao/CLEAN	Enzyme function prediction using contrastive learning.
DeepMicrobes		51		51	51		2020	https://github.com/MicrobeLab/DeepMicrobes	Taxonomic classification for metagenomics with self-attention model.
PopPhy-CNN		51		51		51	2020	https://github.com/YDaiLab/PopPhy-CNN	Host Phenotypes prediction by systematic tree embedded CNN network.

datasets, HyenaDNA incorporates **Demo and Dummy** datasets, typically used for initial testing and 654 validation, though they may lack the data authenticity and application value of specifically collected 655 datasets. Furthermore, NT [9] covers 18 downstream tasks, primarily centered on transcription factor 656 binding, promoter prediction, and chromatin accessibility analysis, emphasizing its close relationship 657 with gene regulation mechanisms. While these genomic benchmarks perform well in single-species 658 analyses, they often fail to capture the inherent complexities of multi-species interactions present in 659 metagenomics. Specifically, the limitations of genomic benchmarks include (1) reliance on single-660 species data, which overlooks the complex interactions in metagenomics; (2) lack of data diversity, 661 as their datasets are typically structured and uniform, lacking the environmental diversity required for 662 metagenomic studies; (3) limited functional prediction, focusing on sequence-based predictions 663 without integrating crucial functional annotations; and (4) inadequate model adaptability, as models 664 trained on single-species genomic data struggle to adapt to multi-species metagenomic data. 665

In contrast, FGBERT [13], as a metagenomic pre-trained model, aims to address interactions within 666 different microbial communities and predict functions across various environments, covering diverse 667 tasks such as gene structure analysis, functional gene prediction, pathogenicity assessment, and 668 nitrogen cycle prediction. These tasks span gene, functional, bacterial, and environmental levels, 669 with input sizes ranging from 1,000 to 213,000 sequences, reflecting the diversity and complexity 670 of metagenomic data. Therefore, incorporating FGBERT's multi-species genomic datasets into our 671 OpenMeta benchmark not only substantiates its proficiency in deciphering complex microbial func-672 tions but also provides a solid framework for comparing and evaluating different models' performance 673 in practical applications. This approach enhances our understanding and utilization of metagenomic 674 pre-trained models in biotechnology and environmental science. 675

### 676 C Datasets

<sup>677</sup> We provide CSV files containing the categories and quantities of each dataset in the .zip fils.

<sup>678</sup> We provide detailed descriptions of the 12 open-source datasets as shown in Appendix Table A2.

Appendix Table A3 shows detailed statistics for all sequence datasets in OpenMeta. The 'Num. Seqs.'

column indicates the total number of sequences in the data set, and the 'Num. Cates' column shows

the number of different categories in the data set. The 'Seqs/Cate Range' column provides the range

of sequence numbers in each category, from smallest to largest. The 'Avg. Len.' column indicates the

average length of the sequences. The 'Source' column describes the source of the data. The 'Task

<sup>684</sup> Type' column indicates the type of task for which the data set was used.

Appendix Table A4 shows detailed statistics for all hierarchical datasets in OpenMeta. The 'Hierarchical Taxonomic Levels' column means the Taxonomic Distribution in Metagenomic Datasets for

Table A3: Statistical analysis of all sequence datasets.

Dataset	Num. Seqs.	Num. Cates	Seqs/Cate Range (Min-Max)	Avg. Len.	Source	Task Type
E-K12	4312	1379	1-106	510.96	Public Database	Multi-Classification
CARD-A AMR Gene Family	1966	269	1-229	1088.1	Public Database	Multi-Classification
CARD-D Drug Class	1966	37	1-513	1088.1	Public Database	Multi-Classification
CARD-R Resistance Mechanism	1966	7	1-1263	1088.1	Public Database	Multi-Classification
PATRIC Pathogenic Genes?	5000	110	1-1081	307.82	Public Database	Multi-Classification
ENZYME	5761	7	288-2055	426.76	Public Database	Multi-Classification
VFDB	8945	15	5-1683	415.47	Public Database	Multi-Classification
NCycDB Nitrogen Cycling Genes	219089	69	1-20548	347.03	Public Database	Multi-Classification
NCRD-N Gene Name	104363	1912	1-18370	407.44	Public Database	Multi-Classification
NCRD-F Gene Family	104363	420	2-35364	407.44	Public Database	Multi-Classification
NCRD-C Categories	104363	29	1-14159	407.44	Public Database	Multi-Classification
NCRD-R Resistance Mechanism	104363	10	166-38073	409.79	Public Database	Multi-Classification

Table A4: Statistical analysis of all hierarchical datasets.

Dataset	#Cog		Н	ierarchi	ical Taxo		Course	Tooka Tuna			
Dataset	#Seq.	Kindom	Phylum	Class	Order	Family	Genus	Specialized	Source	lasks Type	
Cirrhosis	542	3	15	27	40	76	186	531	Public Database	Binary-Classification	
T2D	606	3	17	29	48	94	216	587	Public Database	Binary-Classification	

Cirrhosis and T2D. 'Kingdom', 'Phylum', 'Class', 'Order', 'Family', 'Genus', and 'Specialized' are
all different taxonomic levels in the classification of organisms. Together, these taxonomic levels form
the system of taxonomy, which is commonly used to describe and classify the planet's biodiversity.
Each level represents a classification of organisms from broad to specific.

# 691 D Results

Appendix Table A5 and A6 show the M.F1 metrics for gene operon and pathogen prediction on two small-scale datasets, E-K12 and PATRIC (sequence length less than 5000).

# **694** E Implementation Details

In OpenMeta, we compare several genomic pre-trained models, including FGBERT, DNABERT2, 695 NT, and HyenaDNA. Official implementations of these models can be accessed at the follow-696 ing URL links: HyenaDNA: https://huggingface.co/LongSafari/hyenadna-medium-450k-seqlen-hf, 697 DNABERT2: https://huggingface.co/zhihan1996/DNABERT-2-117M, and Nucleotide Transformer: 698 https://github.com/instadeepai/nucleotide-transformer. We have followed the default hyperparameters 699 described in their respective publications and maintained consistent settings across all datasets, 700 evaluating models at the checkpoints where validation loss was minimized. For sequence datasets, we 701 investigate the impact of three encoding strategies on model performance: K-mer (K=3) frequency 702 features, one-hot encoding features, and mean pooling embeddings from genomic and metagenomic 703 models such as HyenaDNA, NT, DNABERT2, and FGBERT. The Macro F1-score (M.F1) is used as 704

Method	E-K12
RF (3-mer)	20.2
SVM (3-mer)	38.6
AdaBoost (3-mer)	39.9
LSTM (w2v)	40.4
LSTM (one-hot)	38.1
BiLSTM (w2v)	40
BiLSTM (one-hot)	40.1
BiLSTM-Att. (w2v)	38.2
BiLSTM-Att. (one-hot)	40.8
VT	43.3
HyenaDNA	42.4
NT	45.1
DNABert2	51.7
FGBERT	61.8

Table A5: Gene Operon prediction on E-K12.

Table A6	: Pathegons	prediction on	PATRIC

Method	E-K12
RF (3-mer)	20.2
SVM (3-mer)	38.6
AdaBoost (3-mer)	39.9
LSTM (w2v)	40.4
LSTM (one-hot)	38.1
BiLSTM (w2v)	40
BiLSTM (one-hot)	40.1
BiLSTM-Att. (w2v)	38.2
BiLSTM-Att. (one-hot)	40.8
VT	43.3
HyenaDNA	42.4
NT	45.1
DNABert2	51.7
FGBERT	61.8

the primary evaluation metric. In fine-grained sequence datasets, particularly the NCRD dataset for 705 ARG prediction tasks, we evaluate three domain-specific models: the template-matching-based RGi, 706 the deep learning-based DeepARG, and the pre-trained language model-based approaches PLM-ARG 707 and FGBERT. Metrics used for evaluation included Accuracy, Precision, Recall, Macro F1-score, and 708 False Negative Rate. For hierarchical datasets, due to the limited number of labels per hierarchical 709 gene, we employed a variety of supervised models specifically designed for disease prediction, such 710 as RF, SVM, Adaboost, and 1D-CNN, in addition to the specialized PopPhy model. 711

#### F **Observations and Insights** 712

#### **Fine-Grained** Benchmarks. 713

ARG Regarding resistance 714 category classification on the 715 NCRD dataset, the metagenomic 716 pre-trained method FGBERT 717 outperforms the other three ARG 718 prediction methods in all per-719 formance metrics and almost all 720 resistance categories, as shown 721 in Figure A1. The performance 722 results in Table 10 show that 723 DeepARG, a combination of 724 traditional template matching 725 methods and deep learning, 726 performs well in the gene name 727

728





and resistance categories but fails to identify the gene family and mechanism categories, which is since no relevant data are 729 included in the model training process or insufficient information is available in the matching 730 dataset.RGI, as a template matching method, has a more general performance in all categories. 731 PLM-ARG based on protein language modeling provided results and high false-negative rates only 732 in the resistance category. FGBERT, as a metagenomic pre-trained model, performs well in all 733 categories, demonstrating its comprehensiveness and high performance in dealing with fine-grained 734 ARG assays, further proving the necessity and advantages of metagenomic pre-trained models. 735

#### F.1 Genomic and Metagenomic Benchmark 736

Relation between Genomic Benchmark and Metagenomic Benchmark. While these benchmarks 737 excel in single-species genome analysis, they often fail to capture the complex interactions among 738 multiple species inherent in metagenomics. In contrast, metagenomic benchmarks aim to address 739 interactions within diverse microbial communities and functional predictions in various environments, 740

covering tasks such as gene structure analysis, functional gene prediction, pathogenicity assessment, 741 and nitrogen cycle prediction. Specifically, the limitations of genomic benchmarks include (1) their 742 743 reliance on single-species data, which misses the complex interactions in metagenomics; (2) lack of data diversity, as their datasets are typically structured and uniform, lacking the environmental diver-744 sity needed for metagenomic studies; (3) limited functional prediction, focusing on sequence-based 745 predictions without integrating crucial functional annotations; and (4) insufficient model adaptabil-746 ity, as models trained on single-species genomic data struggle with multi-species metagenomic data. 747 These deficiencies underscore the urgent need to develop metagenomic benchmarks that can integrate 748 multi-species interactions and complex environmental factors. 749

#### 750 (A) Metagenomic Pre-trained Models vs. Genomic Pre-trained Models:

Table 14 compares genomic and metagenomic pre-trained models, including pre-training datasets, 751 token embedding methods, network architectures, application tasks, and benchmarks. In terms of **Pre**-752 Training Datasets, DNABERT2 [84] utilizes human and multi-species genomes for its foundational 753 model pre-training, covering a vast dataset of 27.5 billion nucleotide bases from the Human reference 754 genome [32] and 135 species genomes across seven categories. HyenaDNA [49], on the other hand, 755 is pre-trained solely on a single human reference genome. Nucleotide Transformer (NT) [9] pre-trains 756 on three datasets: the human reference genome [32], 3,202 diverse human genomes, and 850 genomes 757 from several species. In contrast, FGBERT [13] employs the MGnify database (updated February 758 2023) [62], comprising 2,973,257,435 metagenomic sequences from various microbial communities. 759

For Token Embedding, DNABERT2 applies Byte Pair Encoding (BPE) [67], while both HyenaDNA
 and NT use 6-mer tokenization. FGBERT utilizes a unique protein-based genomic representation
 tailored for metagenomic sequences.

Regarding Network Architecture, both DNABERT2 and FGBERT adopt BERT-like structures [11]; 763 DNABERT2 enhances its predecessor by replacing learned positional embeddings with Attention with 764 Linear Biases (ALiBi) [54] to eliminate input length limitations and incorporates Flash Attention [10] 765 to boost computational efficiency. FGBERT introduces contrastive learning to strengthen the intricate 766 relationships between metagenomic sequences and functions, proposing two pre-training tasks: 767 Masked Gene Modeling (MGM) and Triplet Enhanced Metagenomic Contrastive Learning (TMC) 768 to enhance co-representation learning of metagenomic gene sequences and functions. HyenaDNA 769 employs a simple stack of Hyena operators for next token prediction, while NT uses an encoder-only 770 Transformer architecture with Rotary Positional Embeddings (RoPE) [70] to enable reasoning over 771 longer sequences during training. 772

For Application and Benchmark, Table 2 provides a detailed comparison of downstream tasks for 773 genomic and metagenomic pre-trained models, highlighting the datasets, tasks, sequence numbers, 774 and class counts. DNABERT2 focuses primarily on binary classifications of promoters and splice 775 site detection tasks across human, mouse, yeast, and virus datasets, with the number in parentheses 776 indicating the number of independent sub-datasets for each task and the sequence count reflecting 777 the size of the first sub-dataset. Notably, **Transcription Factor Prediction** task recurs for both 778 human and mouse species with identical dataset numbers, class numbers, and sequence lengths, 779 suggesting a uniform level of difficulty that may not present significant challenges due to its repetitive 780 nature across similar species settings. HyenaDNA's downstream tasks are divided into two parts: 781 GenomicBenchmarks, consisting of 8 regulatory element classification datasets with sequence lengths 782 ranging from 200 to 500, and NT's 18 prediction tasks. In addition to the human datasets, HyenaDNA 783 includes Demo and Dummy datasets. The inclusion of Demo and Dummy datasets, which are 784 typically used for initial testing and validation purposes. Additionally, NT covers 18 downstream 785 tasks primarily centered around transcription factor binding, promoter prediction, and chromatin 786 accessibility analysis, underscoring its detailed engagement with gene regulation mechanisms. FG-787 BERT engages with various downstream tasks that address multi-species metagenomic sequences 788 through multi-class classification challenges. These tasks span across gene, functional, bacterial, 789 and environmental levels, accommodating input sizes that range from 1,000 to 219,000 sequences, 790 reflecting the diversity and complexity of metagenomic data. 791