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³¹ **Abstract**

32 Species identification using DNA barcodes has revolutionized biodiversity sciences and 33 society at large. However, conventional barcoding methods may lack power and universal 34 applicability across the Tree of Life. Alternative methods based on whole genome 35 sequencing are hard to scale due to large data requirements. Here, we develop a novel 36 DNA-based identification method, varKoding, using exceptionally low-coverage genome 37 skim data to create two-dimensional images representing the genomic signature of a 38 species. Using these representations, we train neural networks for taxonomic 39 identification. Applying a taxonomically verified novel genomic dataset of Malpighiales 40 plant accessions, we optimize training hyperparameters and find the highest performance 41 by combining a transformer architecture with a new modified chaos game representation. 42 Remarkably, >91% precision is achieved despite minimal input data, exceeding alternative 43 methods tested. We illustrate the broad utility of varKoding across several focal clades of 44 eukaryotes and prokaryotes. We also train a model capable of identifying all species in 45 NCBI SRA using less than 10 Mbp sequencing data with 96% precision and 95% recall and 46 robust to sequencing platforms. Enhanced computational efficiency and scalability, 47 minimal data inputs robust to sequencing details, and modularity for further development 48 make varKoding an ideal approach for biodiversity science. 49 50 **Keywords:** biodiversity science, computer vision, DNA barcoding, DNA signature,

51 Malpighiaceae, natural history collections, neural networks, species identification,

52 taxonomy

⁵⁴ **Introduction**

55 For two decades, conventional DNA barcoding, which relies on standardized short 56 sequences (400–800 bp) for species identification¹⁻⁵, has enabled novel and massively 57 scalable science spanning evolution^{4,6-9}; ecology¹⁰⁻¹⁴ and paleontology¹⁵⁻¹⁹. Practical 58 applications of barcoding have also made major contributions to environmental health, 59 including the ability to authenticate medicinal plants²⁰, detect agricultural pests²¹, and 60 monitor poaching and the trade of endangered species^{22–27}. Despite these remarkable 61 achievements, conventional DNA barcoding suffers from at least four limitations. First, 62 barcodes are customized specifically for a taxon (e.g., plants, animals, and fungi), and 63 therefore are not universal. For example, commonly used plant barcodes from chloroplast 64 genes such as *mat*K and *rbcL* cannot be applied as barcodes for all plants^{28,29}, or for animals 65 and fungi. Second, conventional barcode loci may fail to distinguish closely related taxa, a 66 pervasive shortcoming in plants^{2,30}. Third, reliance on a single locus may lead to spurious 67 results in the case of complex evolutionary scenarios such as hybridization in deep or 68 shallow time^{31–34}. And fourth, the necessary comparison of homologous genes may fail 69 when PCR primers are not universal³⁵, the source DNA is fragmented²⁷, or paralogy and the 70 presence of pseudogenes confounds accurate orthology assessments $36,37$.

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72 Newer alternatives to conventional barcoding have begun to address these challenges by 73 leveraging high-throughput sequencing and machine-learning powered by deep neural 74 networks. High-throughput sequencing facilitates more comprehensive assessments of 75 total genomic space^{38,39}. For example, presence and absence patterns among short DNA 76 sequences (k-mers) from low-coverage reads (i.e., genome skims) can estimate overall 77 sequence distances, bypassing genome alignments entirely as implemented in *Skmer*⁴⁰. 78 Machine learning enables more complex sequence comparisons than conventional methods 79 that rely on homology and simple metrics⁴¹. Machine-learning models can cluster DNA 80 sequences without supervision^{42,43} or classify sequences based on reference datasets⁴⁴⁻⁴⁹.

81 In particular, neural networks are exceptionally powerful for sophisticated computer-

82 vision tasks, such as image classification⁵⁰. Thus, the combination of low-coverage genome

83 skimming data and neural networks holds enormous promise for accurate and scalable

84 DNA barcoding, but its potential has yet to be fully realized³⁹.

85

86 Genomes differ substantially in many features beyond the simple nucleotide divergence 87 commonly used in conventional barcoding, but these genomic features have been 88 overlooked in species identification⁵¹⁻⁵⁵. We propose that (1) relevant genomic features 89 can be captured by nucleotide composition with short k-mer counts and very small 90 sequence coverage; and (2) these counts can be used to distinguish species and higher taxa 91 efficiently and accurately using machine learning. Prior work on k-mer-based 92 representations of genome composition (i. e. DNA signatures) has shown high accuracy can 93 be achieved with high-coverage data or a large number of replicates per taxon, particularly 94 for identification at higher taxonomic ranks^{42-47,56-63}. However, given the millions of 95 existing species and the sparse genetic data available, a practical scalable method would 96 require: (1) consistently high accuracy despite limited evolutionary divergence; (2) fast 97 computations; and (3) high accuracy with small training datasets (both in number of 98 samples and DNA data per sample). Here we developed a novel DNA signature method, 99 which we call **varKoding**, that integrates very low-coverage genome skim data with 100 optimized training of machine-learning models using two-dimensional images 101 representing genome composition (**Figure 1A**). To develop and optimize varKoding for 102 accurate species identification, we generated a *de novo* genome skim dataset including 103 hundreds of samples derived primarily from historical herbarium specimens for the 104 diverse plant genus *Stigmaphyllon* (Malpighiaceae), which has received extensive 105 phylogenetic and taxonomic treatment⁶⁴⁻⁶⁸. Next, we explored the utility of varKoding and 106 compared it to alternatives at different phylogenetic depths from families to species within 107 the flowering plant order Malpighiales (Malpighiaceae, Chrysobalanaceae, and 108 Elatinaceae). Finally, we demonstrate the scalability of varKoding and its potential 109 application in forensics and related fields by testing it on (1) species-level datasets from

- 110 fungi, plants, animals, and bacteria; (2) massive datasets retrieved from the NCBI sequence
- 111 read archive (SRA); and (3) a previously published environmental DNA (eDNA) dataset.

113 Figure 1. Overview of varKoding. (A) Image generation workflow, depicting varKodes. Images are natively 114 grayscale, but here they are mapped to a rainbow color scale for increased contrast. **(B)** Phylogeny and 115 example varKodes of *Stigmaphyllon* species. (C) Phylogeny and example varKodes of Malpighiaceae genera 116 including their closest outgroup (*Elatine*, Elatinaceae). Time trees in 1B and 1C (D) Examples of varKodes 117 from across plant families of Malpighiales, and (E) across kingdoms. Chronograms depicted for each 118 representative set with timelines in millions of years (Myr) at the bottom of **B** and **C**. These were derived 119 from an ongoing family-wide phylogenomic investigation of the family Malpighiaceae (C. C. Davis personal 120 communication) using methods and fossil constraints described in Cai et al.⁶⁵; dates inferred are consistent 121 with earlier findings⁶⁵ and were not applied in this study for quantitative analyses.

122 Results and Discussion

123 **DNA signature images can be classified with generalized neural networks**

124 We first generated a novel kind of image representation of a DNA signature, which we 125 termed a **varKode**. varKodes map k-mers onto pixels of a 2-D image based on their 126 similarity and represent ranked k-mer frequencies as pixel brightness. Variation in 127 varKodes can be small but remain visually perceptible among species (**Figure 1B**) and 128 genera (**Figure 1C**). Variation is more striking among higher levels of phylogenetic 129 divergence, such as between families in the order Malpighiales (**Figure 1D**) or different 130 kingdoms of eukaryotes and prokaryotes (Figure 1E). We expected, therefore, that neural 131 network architectures developed for image classification, (e.g., deep residual networks, 132 resnets⁶⁹ or vision transformers, ViT^{70,71}) would be able to differentiate varKodes.

133

134 We first optimized hyperparameters and training conditions to maximize accuracy for 135 species-level identification of *Stigmaphyllon*. We identified that varKodes depicting k-mer 136 length = 7 struck a good balance between accuracy and the amount of input sequence data 137 (**Figure 2A**). Furthermore, models trained with augmented data from several subsampled 138 sequences drawn from each individual exhibited substantially better performance (**Figure**) 139 **2A**). A linear model demonstrated that neural network architectures and training methods 140 designed for image classification of photographs^{69,72-75} are extremely useful for varKode-141 based identification. Specifically, we observed increased accuracy with more parameter-142 rich neural network architectures (*ResNeXt101*⁷⁶, among those tested), augmentation with 143 lighting transformations, *CutMix*⁷⁵ and *MixUp*⁷⁴. Label smoothing⁷⁷ and pretraining models 144 on generalized photographs decreased accuracy (**Figure 3**). Contrary to the widely held 145 idea that deep neural networks require very large training datasets^{60,78}, the 146 aforementioned approaches enabled training with very modest data amounts: four 147 biological replicates per taxon was sufficient for 100% median accuracy (**Figure 2B**). 148 Errors in species-level identification were concentrated among sequences derived from 149 herbarium samples that demonstrated evidence of DNA damage, as is sometimes reported **150** for ancient DNA⁷⁹ (**Figure 2B**). However, including low-quality training samples slightly

- 151 decreased mean validation accuracy—from 73% to 71%—for low-quality validation
- 152 samples, but had no effect on high-quality validation samples (89-90% mean accuracy,
- **Figure 4A**).

165

- 166 **Figure 3**. Marginal effects of neural network model and training options. Dots represent individual replicates,
- 167 and bars depict averages. All parameters were identified to be significant in a linear model: more complex
- 168 model architectures, lighting transformations, and augmentation methods *MixUp* and *CutMix* improved
- 169 accuracy. However, pretraining with large image datasets and label smoothing decreased accuracy.

170

171 **Figure 4.** Effect of the inclusion of low-quality training samples, inferred from variation in base pair content 172 (A, B) or insert size (C, D) . Increasing the fraction of samples in the training set that were low-quality did not 173 strongly affect the average validation accuracy, but it increased dispersion. Low-quality samples are the four 174 samples with highest variation in base-pair content or shortest insert size in raw reads for each species. 175 Panels **B** and **D** show the correlation of each quality metric with DNA extraction yield.

177 We hypothesized that lower-quality samples shared similar sequences resulting from 178 common patterns of DNA damage and greater levels of microbial or human contaminants, 179 resulting in spurious similarities in varKodes (**Figure 5**). Contaminants also are thought to 180 increase errors in other genome skim methods⁸⁰. To mitigate this problem, we applied 181 multi-label classification⁸¹ to our neural network models. Although single-label 182 classification models always return a single prediction (that is, an inferred label), multi-183 label models can return zero or more predictions, avoiding spurious results when there is 184 uncertainty. For a set of samples with known labels used for validation, a prediction is a 185 true positive if the predicted label matches the actual label, and a false positive if not.

- 186 Failure to predict an actual label is deemed a false negative. For each validation sample, we
- 187 summarized predictions as (1) correct (true positives only); (2) incorrect (false positives
- 188 only); (3) ambiguous (multiple predictions, including true and false positives); or (4)
- 189 inconclusive (i. e. no prediction above the confidence threshold of 0.7). For each test, we
- 190 summarized results across all validation samples using two metrics: precision (the sum of
- 191 all true positives divided by the sum of all true and false positives) and recall (the sum of all
- 192 true positives divided by the sum of all true positives and negatives).

varKodes for species of *Stigmaphyllon*

193

194 Figure 5. Low-quality DNA may lead to spurious patterns of similarity in varKodes. Samples with lower 195 quality show varKode patterns divergent from their species more often than high-quality ones. These 196 divergent patterns may be similar between low-quality samples across species. These samples also show 197 reduced validation accuracy in a single-label model. For each sample, we show the varKodes produced from 198 all DNA data available. Within each species, samples are organized from lowest (left) to highest (right) DNA 199 quality. Bounding boxes around each sample indicate the average validation accuracy across 30 random 200 replicates with 7 training samples per species.

202 After optimizing these training conditions, we directly compared varKodes to an existing 203 method of DNA signature representation: the frequency chaos game representation 204 *(fCGR*)^{56,59}. In *fCGR*s, k-mers are mapped to pixels based on their oriented sequence and 205 pixel brightness represents the rescaled k-mer frequency. To isolate the effects of pixel 206 mapping and brightness, we created a new representation combining *fCGR* mapping with 207 *varKode* ranked frequency transformation (*rfCGR*). By directly comparing these 3 kinds of 208 representation combined with four neural network architectures, including (1) two 209 previously employed with *fCGR*s^{42,44,60}, (2) the optimal architecture in our initial tests 210 (ResNeXt101⁷⁶), and (3) a Vision Transformer (ViT^{70,71}), we found that ViT combined with 211 *rfCGR* representation maximizes performance (**Figure 6**). A multilayer perceptron, as 212 employed in previous work^{42,60}, could not identify any species correctly here (**Figure 6**). 213 Similarly, a previously employed shallow 1D convolutional neural network⁴⁴ 214 underperformed more complex architectures (Figure 6). *fCGR* showed much higher error 215 rates than either *rfCGR* or *varKodes*, which yielded similar results but with slightly higher 216 accuracy for *rfCGR* (**Figure** 6). These results indicate that deep complex neural networks, 217 while not explicitly developed for DNA signature, are necessary to extract features from 218 very low-coverage data and distinguish closely related species. Moreover, the method of k-219 mer frequency data transformation seems more consequential than the mapping of k-mers 220 to pixels for the performance of different image representations. Due to its higher 221 performance, we adopt the combination of *ViT* and *rfCGRs* for subsequent tests.

Stigmaphyllon species cross-validation accuracy

222

223 **Figure 6.** Effect of image representation and neural network architecture on cross-validation accuracy of 224 species identification in *Stigmaphyllon*. One example for each image representation is shown, drawn from the 225 same DNA data (SRA accession XXXX) and mapped to a rainbow color scale for increased contrast. See text for 226 details on architectures.

227

228 In summary, we developed and tested a robust and scalable method of DNA barcoding 229 capable of training with small amounts of data, and implemented it in the **varKoder** 230 software, which can process sequence data, train an image-classification neural network 231 using varKodes or rfCGRs, query new data with a trained neural network, and convert 232 between the alternative k-mer mappings. These tasks are accomplished with widely used

233 tools for sequence processing^{82–86} and for neural network training^{87–90}.

235 varKodes are highly accurate for identification of species, genera, and families 236 To test varKoder under a real-world scenario with heterogeneous data (e.g., large numbers 237 of taxa, multiple replicates per taxon, varying sequence depth and sample quality), our *de* 238 *novo* genomic data set included 287 accessions: 100 samples of *Stigmaphyllon* from our 239 initial development outlined above, plus additional genera in the families Malpighiaceae 240 (31 genera; 151 samples), Chrysobalanaceae (8 genera; 30 samples), and Elatinaceae (1 241 genus; 6 samples) in the order Malpighiales. We found high cross-validation accuracies for 242 species identity of *Stigmaphyllon* (87.0–96.7% correct, 94.6%–98.9% precision, 88.0%– 243 96.7% recall depending on data input amount; **Figure 7A**). Most errors were inconclusive 244 predictions $(2.2-10\%)$, instead of ambiguous $(0-3\%)$ or incorrect $(1-4\%)$ predictions. 245 *varKoder* is robust to the amount of input sequence data necessary for model training, 246 performing well even at the lower range of input data (**Figure 7A**). Assuming an average 247 genome size of about 2 Gbp for the average species of Malpighiaceae⁹¹, the 500Kbp-248 200Mbp of data used here represented exceptionally low coverages of about $\sim 0.0002 \times -$ 249 $0.107\times$. Moreover, when compared to cross-validation accuracies of alternative barcoding 250 methods, *varKoder* accuracy is higher than *Skmer*, which showed 46% correct predictions 251 (57.5% precision, 46% recall) with minimal data amounts and peaked at 79.1% for the 252 larger data amounts (80% precision, 79.1% recall, **Figure 7A**). On the other hand, 253 conventional barcodes including individual plastid genes and nuclear ribosomal ITS 254 regions performed well for both BLAST-based (25-97% correct, 66.6-97.3% precision, 25-255 97% recall depending on the gene) and phylogenetic-based (94–95% correct, >99% 256 precision, 97.2–98.4% recall for concatenated matrices) approaches when at least 50 Mbp 257 of data was provided (**Figure 7A, Figure 8**). However, these results were much worse 258 when <50 Mbp of data were available (down to zero correct for BLAST), with unsuccessful 259 locus assembly leading to inconclusive predictions as the primary reason for the failure 260 (**Figure 7A, Figure 8**). Finally, an unsupervised clustering method based on neural 261 networks applied to *fCGRs* (*iDeLUCS*⁹²) reached 24–60% clustering accuracy depending on 262 input data amount when prompted to cluster *Stigmaphyllon* sequences into 10 groups 263 (Table 1). In summary, *varKoder* reaches much higher accuracy for species determination

264 than existing methods for unprecedentedly small amounts of data and demonstrates

265 similar accuracies when greater amounts of sequence data are available.

267

268 **Figure 7.** Performance of *varKoder* and alternative barcoding methodologies across different data sets. (A) 269 Leave-one-out cross-validation to identify species of Malpighiales using different approaches and amounts of 270 data to assemble query samples. (**B**) Same as (A), but for genera. (C) Performance for species-level

- 271 identification across different publicly-available datasets: *Bembidion* beetles, *Corallorhiza* orchids,
- 272 *Mycobacterium tuberculosis* bacteria, and *Xanthoparmelia* fungi. All query samples used as much data as were
- 273 available. (D) Performance for Eukaryote family-level identification for different amounts of input data.
- 274

Conventional barcode accuracy across different taxonomic levels

Base pairs in query images

276 **Figure 8.** Accuracy of conventional barcode loci for species, genera and families within the Malpighiales. 277

- 285 Table 1. Accuracy in deuces classification by data amount and plastid genes included.
- 286
- 287 Genus-level identification yielded similar high accuracies with *varKoder* (86.1–93.3%)
- 288 correct, 97.2%-97.7% precision, 86.4%-94.7% recall depending on input amount, **Figure**
- 289 **7B**), but with a higher rate of inconclusive predictions (4.5–11.5%). A linear model
- 290 demonstrated that this higher uncertainty can be attributed to two factors: (1) samples

291 exhibiting higher levels of DNA damage in genera other than *Stigmaphyllon*; and (2) genera 292 trained with fewer replicates (e.g., down to 3 samples for some genera; **Figures** 9-10). 293 Despite this trend, the vast majority of genera with fewer replicates and lower DNA quality 294 can still be correctly predicted, resulting in the $>97\%$ prediction and $>86\%$ recall across 295 the whole dataset. Additionally, samples within genera share fewer genetic similarities 296 than samples within species, which likely poses a more challenging classification problem. 297 However, the incorrect rate was very small in all cases $(0.7-2.1\%)$, with most errors being 298 inconclusive or ambiguous predictions. In contrast, *Skmer* exhibited better performance 299 when larger amounts of data were used (99.2% correct, 99.2% precision, 99.2% recall for 300 200 Mbp), but performed poorly for lower amounts of data like those commonly generated 301 from genome skim experiments (58.2% correct, 58.2% precision, 58.2% recall for 500 302 Kbp) (**Figure 7B**). Genus-level identifications using conventional barcodes in a 303 concatenated phylogeny were up to 98.1% correct (99.2% precision, 97.2%% recall) when 304 a large amount of data (200 Mbp) was available (**Figure 7B**). But like its application at 305 species-level identification, most predictions were inconclusive when less than 20 Mbp 306 reads were used (**Figure 7B**). Although genome skimming can be used to sequence 307 conventional barcodes, they are more often obtained with amplicon sequencing, which has 308 failure rates ranging from $15-75\%$ even with highly optimized protocols⁹³, leading to an 309 even higher number of inconclusive predictions. At the family level, *Skmer* and *varKoder* 310 had near-perfect accuracy across all data amounts (>97% correct), while conventional 311 barcodes performed well when there were sufficiently large amounts of data (**Figures 8,** 312 **11**).

Factors affecting varKode prediction accuracy

314 **Figure 9.** Predictors of confidence in correct genus. A) Confidence increases with more training samples per 315 genus. B) Amount of data per validation image has little effect. C) Validation samples with low quality have 316 lower confidence.

317

Number of samples available for different data amounts

Post−cleaning base pairs available

- 319 **Figure 10.** Number of samples available for different data amounts in the Malpighiales and Eukaryote
- 320 families datasets. Arbitrary colors are assigned to individual taxa.

321

325 varKodes are universal and scalable across the Tree of Life

- 326 To further test the universality of varKodes, we expanded to sequencing data from diverse
- 327 clades of plants, fungi, animals, and bacteria (**Figure 7C**). These tests included species-level
- 328 identification in insects (*Bembidion* beetles^{54,94}) and lichen-forming fungi
- 329 *(Xanthoparmelia*⁹⁵), species and infra–specific taxon identification in coralroot orchids
- 330 *(Corallorhiza*⁹⁶), and clinical isolate identification of strains of human pathogenic bacteria
- 331 *(Mycobacterium tuberculosis⁹⁷)*. In all cases, we tested the performance of *varKoder* on taxa
- 332 included in the training set and on taxa not included in the training set. We identified

333 perfect species identification (100% correct, 100% precision, 100% recall) for beetles and 334 coralroot orchids included in the training set. For bacteria, 5.6% of the validation set 335 returned ambiguous predictions; the remaining samples were correctly identified (94.7%) 336 precision, 100% recall). In lichen-forming fungi, which include DNA from both the fungal 337 and algal partners, and thus are more challenging, 10% of the test samples returned 338 incorrect predictions and another 10% were inclusive; the remainder were correct (89% 339 precision, 80% recall). For all cases, species or varieties not included in the training set 340 generally resulted in inconclusive results, with a minority yielding incorrect predictions 341 **(Figure 7C)**.

342

343 Finally, we tested the scalability of varKodes in three large-scale datasets: (1) all 861 344 eukaryotic families with Illumina data on NCBI SRA, (2) all taxa with multiple accessions on 345 NCBI SRA, including different sequencing platforms and library strategies (254,819) 346 accessions and $14,151$ taxa across all taxonomic ranks), and (3) a previously published 347 dataset of 2916 soil eDNA samples from all seven continents⁹⁸. Owing to NCBI download 348 speed bottlenecks, we restricted varKode construction to a very limited maximum of 10 349 Mbp of DNA data in the former 2 cases. The family-level eukaryote data achieved a rate of 350 correct predictions of 65.2–81.3% across all kingdoms when families were included in the 351 training set (**Figure 7D**), with most errors being inconclusive predictions (17.5–33.1%). 352 Precision varied from 95.3% to 97.3% and recall from 67.9% to 78.3%. Similarly to the 353 species- and variety-level exercise, families not included in the training set often yielded 354 inconclusive predictions **(Figure 7D)**, suggesting a potential for varKoding to be used as a 355 discovery tool when reasonably well-sampled training data sets are available. The 356 expanded data with all taxa from NCBI SRA revealed that varKoding is robust to sequencing 357 platform and library preparation method (Figure 12). Predictions at the family level or 358 pooled for all the taxonomic hierarchy are accurate regardless of sequencing details (>94% 359 precision, >86% recall). The much higher accuracy when compared to the dataset based on 360 Eukaryotic families alone may be an effect of a completely random validation set instead of 361 stratified by family, resulting in higher representation of commonly sampled families. At 362 the genus and species level, results are more dependent on the sequencing method (**Figure**)

363 12). For genera, precision/recall using 10Mbp of data varies from 90.8%/90.8% with 364 whole genome shotgun libraries in PacBio to 97.9%/97.6% with genotype-by-sequencing 365 in Illumina. Finally, the eDNA data shows promise in using varKoding to identify the 366 geographical origin of an environmental sample: in the validation set, at 10Mbp of DNA 367 data, 94.0% of the samples had continent correctly identified, with 2.6% being incorrect, 368 1.9% being ambiguous, and 1.5% being inconclusive (84.7% prediction, 84.5% recall) 369 **(Figure 13)**.

Metric - precision - recall

370

371 **Figure 12.** varKoder performance in predicting taxonomy for all data on SRA. Sample sizes refer to the

- 372 number of validation accessions available for each combination of platform, sequencing strategy and
- 373 taxonomic rank.
- 374
- 375

377 Figure 13. Varkoder performance in identifying the geographical origin of a soil metabarcoding sample. A) 378 Performance across the whole dataset. B) Performance for each continent.

379

380 A single model classifying all of life is not possible with conventional barcodes. *Skmer*, the 381 state-of-the-art genome skimming alternative, cannot be scaled to a dataset of this size: our 382 attempt to apply it to Eukaryote families could not be finished after more than 40 days 383 using 32 high-performance computing cores. In general, conventional barcodes, when 384 derived from genome skimming data, require memory- and processor-intensive sequence 385 assembly, and *Skmer* relies on pairwise all-by-all sample comparisons; its computing time 386 and required storage both increase quadratically with the number of samples. Neural 387 network models, on the other hand, have a fixed size, independent of the number of 388 samples used in training, and training time scales linearly with the number of input 389 samples. Our most complex model, trained on all taxa available from the NCBI SRA, has 390 about 1.3GB of disk size. varKode images also are tiny replacements (8.2 KB on average for 391 k-mer length of 7) for much larger genomic data sets (on average, 144 MB per sample 392 here). Downloading up to 20Mb of sequence data for over 250,000 accessions from the 393 NCBI SRA was the bottleneck, taking over 70 days. By parallelizing processing over 40 394 cores, processing this data into varKodes was about 10 times faster, resulting in

395 approximately 18GB of data including all of these accessions. Training a model on more 396 than 1.3 million images took about 45 hours using only 2 GPUs. Although training on large 397 datasets requires powerful GPUs and large memory, training on small datasets and 398 querying is possible on personal computers in a few seconds to minutes. To reduce the 399 computational resources required for training new datasets, we provide a pre-trained 400 model from both varKodes and rfCGRs from all taxa on SRA using the huggingface hub 401 (https://huggingface.co/brunoasm/vit_large_patch32_224.NCBI_SRA). See Asprino et al.⁹⁹ 402 for details on the data used for this model. Whenever the data become available, a model 403 potentially trained on millions of species easily can be ported to devices without 404 continuous internet access. Moreover, the minimal data amounts needed for identification 405 could be generated in seconds in a portable Nanopore device. Finally, the library 406 preparation method based on shotgun sequencing is very simple and can be automated 407 with portable consumer devices, such as the Nanopore Voltrax. Together, these properties 408 allow for more widely distributed applications of varKoding, such as field-laboratory 409 environments¹⁰⁰ or proposed distributed genetic databases¹⁰¹.

410

411 **Conclusions**

412 varKoding is universal, accurate, efficient, and holds tremendous promise for documenting 413 and discovering Earth's biodiversity. It achieves accurate identification with minimal data 414 compared to existing next-generation sequencing methods, while maintaining universal 415 applicability across the Tree of Life. Its modular framework can evolve alongside advances 416 in sequencing technologies, bioinformatics, and machine learning, as exemplified here by 417 the update in image representation (*varKodes* to *rfCGRs*) and neural network architecture 418 (resnext to ViT) after initial testing. For these reasons, we expect it will contribute for the 419 wider adoption of DNA signatures on biodiversity assessments and ecological research, 420 overcoming current challenges³⁹. Reference data for varKoding will be increasingly 421 available from ambitious efforts in genome sequencing¹⁰²⁻¹⁰⁶. However, we note that 422 reference data for varKoding is much easier and cost-effective to obtain from low-coverage 423 genome skims than high-quality contiguous genomes: the robustness to minimal levels of 424 coverage a central advantage of our method. For example, our cost for a $3\times$ skim of

425 herbarium samples is about \$34 per sample, versus a high-quality genome which may cost 426 tens-of-thousands of dollars each. Thus, varKoding shows tremendous promise for further 427 automating species identification from natural history collections¹⁰⁷⁻¹⁰⁹.

428

429 We expect that varKoding will be invaluable to the biodiversity science community in 430 numerous ways, with many avenues remaining to be explored. One of them is the 431 identification of samples with poor-quality and degraded DNA, such as unidentified 432 fragmentary fossil and subfossil remains in natural history collections^{107,110}. For example, 433 Malpighiales samples with signs of DNA damage could be correctly identified using 434 *varKoder* to species or genus in many cases and to family in almost every case. Future 435 research could explore the lower limits of sample quality and sequence coverage to achieve 436 accurate identification at different divergence levels. Finally, we expect that new neural 437 network architectures and forms of DNA representation will continue to be explored. One 438 limitation of varKoding, as applied here, is the challenging identification of samples within 439 mixed components such as lichens or environmental DNA. However, with long-read 440 sequencing, *varKodes* and *rfCGRs* from single reads could potentially include sufficient data 441 for that end. Moreover, mixed samples could be useful for other ends: varKodes could be 442 used to classify a set of sequences based on any kind of metadata, beyond taxonomy as 443 demonstrated by our test on the geographical origin of a soil sample. 444

445 Author contributions

446 BdM conceived varKodes and wrote the program *varKoder*. BdM and CCD designed the 447 research. CCD, XD, YY, LCM, and CA collected the new sequence data. BdM, CCD, LC, YY, PJF 448 analyzed and interpreted the data. LCM prepared the figures. BdM and CCD wrote the 449 manuscript with key contributions from LC, YY and PJF. All authors approved the 450 manuscript.

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⁸²² **Online Methods**

⁸²³ **Sequence data**

825 *data*—The newly generated plant data used here and the methods to obtain these data are

826 described in detail in a data descriptor article⁹⁹. Briefly, they included members of the large

827 and diverse order Malpighiales³⁴: Malpighiaceae (251 accessions representing 31 genera),

828 Elatinaceae (6 accession for 1 genus), and Chrysobalanaceae (30 accessions for 8 genera).

829 Malpighiaceae includes *Stigmaphyllon* with the most comprehensive species sampling: 10

830 species and 10 accessions sampled per species. All 100 *Stigmaphyllon* samples were

831 sequenced specifically to build, validate, and test our identification models at shallower

832 phylogenetic depths, since their taxonomy has been extensively revised by coauthor C.

833 Anderson^{67,68}. Each of these samples was labeled with species, genus, and family names.

834 The focus for the remainder of the Malpighiaceae, Chrysobalanaceae, and Elatinaceae

835 sampling was to identify a given sample to genus. In this case, among the non-

836 *Stigmaphyllon* samples we included 3–9 species per genus. Each accession in this case was

837 labeled with its corresponding genus and family identification. Unlike *Stigmaphyllon*, where

838 we included multiple accessions per species, there were no additional replicates per

839 species for our genus-level sampling. For this dataset, we used leave-one-out cross

840 validation in all assessments, and therefore there are no train and validation sets. For

841 additional information see Asprino et al.⁹⁹.

842 Public genomic data compilation—To further understand the versatility of varKodes more 843 broadly across the Tree of Life, we tested species identification using genome skim data 844 sets from four genera of plants, animals, fungi, and a bacterial species. This involved a plant 845 data set from coralroot orchids (genus *Corallorhiza*)⁹⁶, a beetle data set in the genus 846 *Bembidion*^{54,94}, a lichen-forming fungus in the genus *Xanthoparmelia*⁹⁵, and a bacterial data 847 set of clinical isolates from *Mycobacterium tuberculosis*, the species of pathogenic bacteria 848 \cdot that causes tuberculosis⁹⁷. In all these cases, we labeled samples with the lowest-level 849 taxonomic identification available (species, subspecies or isolates). For taxa with two or 850 more samples available, 20% (with a minimum of 1) were randomly selected for the 851 validation set, which also included all taxa represented by a single sample (therefore, 852 absent from the training set). The remaining accessions were used in the training set. See 853 Asprino et al.⁹⁹ for further information.

854

855 We also compiled two broad datasets from the NCBI SRA. The first consists of all 861 856 eukaryotic families with sufficient sequence read data using the Illumina platform and 857 whole genome shotgun sequencing. We labeled samples with family name only and 858 included taxa with at least two associated accessions in the training set. Our validation set 859 consisted of 20% randomly selected accessions from each family (with a minimum of one), 860 plus all accessions in families with a single accession available (therefore not part of the 861 training set). The second broad-scale dataset includes all taxa on NCBI SRA that could be 862 represented by at least 3 independent accessions. In this case, we included different 863 sequencing platforms (Illumina, PacBio, Nanopore, BGIseq) and library preparation 864 methods (whole genome shotgun, RADseq, GBS). For taxa with too many sequences (such 865 as humans, crops, etc.), we randomly chose up to 20 accessions for each combination of

⁸⁷⁹ **varKode design and testing**

880 *Sequence data preprocessing—Prior* to the construction of images, raw reads were lightly 881 cleaned using the following steps: identical reads were de-duplicated using *clumpify.sh* as 882 implemented in *BBtools*^{83,111}, adapters were removed, low-quality tails trimmed, and 883 overlapping read pairs merged using *fastp*⁸⁵ with options "--detect_adapter_for_pe", "--884 dedup", "--dup_calc_accuracy 1", "--disable_quality_filtering", "--disable_length_filtering", "--885 trim_poly_g", "--merge", "--include_unmerged", . Next, we randomly selected subsets of 886 cleaned reads with predefined data amounts, ranging from 500 kbp to 200 Mbp, with 887 *BBtools*. These data subsets were used to generate a variety of input varKodes for a single 888 sample and all such images were used for training (see main text and Figure 2A). Finally, 889 we applied dsk^{84} to count k-mers of a given length based on clean raw reads (i. e. k-mers 890 are counted for each read and their frequencies are pooled across reads). dsk exhibits good 891 performance with low memory requirements, which is ideal for potential applications 892 using varKodes on low-memory devices. We note that analyses for species-level public 893 datasets have low computational requirements and were performed on an Apple MacBook 894 with ARM processor architecture.

895 *varKode and rfCGR construction*— We designed novel images—**varKodes**—that portray 896 relative frequencies of k-mers from low-coverage raw Illumina reads. These are similar to a 897 frequency chaos game representation (*fCGR*) *sensu* Jeffrey⁵³, but optimized for raw reads in 898 which sequence orientation is unknown, and therefore canonical k-mers and their reverse 899 complement are indistinguishable. This averaging of canonical k-mer frequencies and their 900 reverse complements is widely used in the context of raw reads^{40,61,62,112,113}. We call these 901 images varKodes because they en*CODE* the *VARiation* in k-mer frequencies in a sample. We 902 name our method **varKoding** after varKodes, but notice that it is modular and can use 903 other kinds of DNA image representation. They are meant to represent a DNA signature by 904 mapping k-mer identity to pixel position in an image, such that k-mers with more similar 905 composition are closer together. Additionally, the brightness of these pixels represents the 906 abundance of the associated k-mer, but we use ranks instead of raw frequencies to 907 decrease the effect of overabundant and artifactual k-mers. In summary, varKodes are 908 produced by mapping k-mer counts onto a pre-computed map of k-mers to pixels, and 909 transforming frequency data to pixel brightness. varKode design employed t -SNE¹¹⁴ and 910 the python libraries *numpy*⁸⁷ and *pillow*¹¹⁵. In addition to varKodes, here we also developed 911 a new image representation that uses the same pixel mapping as *fCGRs* but represents k-912 mer abundance as ranks instead of raw frequencies. We named these ranked frequency 913 chaos game representation (*rfCGR*). Both varKodes and *fCGRs* are saved as 8-bit PNG 914 images including labels as exif metadata.

915 *Testing k-mer length and data amount—We employed fastai*⁸⁹ for, a high-level 916 implementation of neural networks based on *pytorch*⁸⁸ for training and prediction. All the 917 model architectures we applied are image classification models available from the *timm* 918 library⁹⁰, which have been widely tested using a variety of image types. To identify the 919 optimal training hyperparameters for our neural network, we conducted a series of tests 920 using the species-level data set for the genus *Stigmaphyllon*. We generated varKodes for 921 each of the *Stigmaphyllon* samples. We first tested the joint effect of k-mer length and input 922 data amount for neural network classification accuracy by selecting three samples per 923 species as a validation set; the remaining samples were used to train neural networks using 924 different amounts of input data across 10 randomly generated training sets. As input data

925 for both the validation and training sets, we randomly subsampled the original sequences 926 into fastq files containing from 500 Kb to 200 Mb (equivalent to about 1,700 to 670,000 927 $2x150$ bp Illumina reads). In this test, we only included samples that yielded at least 200 928 million base pairs after cleaning. We also tested the effect of including images for all data 929 amounts during training. For each replicate, we applied the widely used image 930 classification neural network *resnet50* architecture¹¹⁶ to classify varKodes and trained 931 models for 30 epochs. We visualized the distribution of validation accuracy for each 932 combination of input data amount and k-mer lengths to find a good balance between both. 933 Visualizations and code applied for training and evaluation is available in our GitHub 934 repository.

935 Neural network optimization—After identifying an appropriate k-mer length and input data 936 used to produce varKodes (**Figure 2**), we next tested a series of neural network training 937 conditions. We varied the neural network model complexity, choosing from seven 938 commonly used architectures: *resnet50*¹¹⁶, *resnet-D*⁶⁹ with different depths (18, 50, 101), a 939 wide *resnet50⁶⁹, efficientnet-B4*¹¹⁷, and ResNeXt101⁷⁶. We also tested the effect of the 940 following: random initial weights vs. pretrained weights from the *timm* library⁹⁰, presence 941 or absence of lighting transforms, presence or absence of label smoothing, and presence or 942 absence of augmentation strategies (i.e., \textit{CutMix}^{75} or \textit{MixUp}^{74}). Because these parameters 943 may have complex interactions, we tested all combinations of architecture, pretraining, 944 transforms, label smoothing, and augmentation, with 20 replicates for each combination of 945 conditions. In each replicate, we randomly chose 20% of the samples for each species of 946 *Stigmaphyllon* as validation and trained the model using the remainder for 30 epochs. 947 Training was performed using all varKodes available for each sample (from 500kbp to 948 200Mbp). For validation, we separately evaluated whether each varKode with a different 949 amount of data was correctly identified. For each replicate and amount of data used to 950 validate var Kodes, we recorded the average validation accuracy across the validation set. 951 We then applied a linear model to predict the effect of all training parameters and amount 952 of data in varKodes in the validation set on validation accuracy. Validation accuracy in this 953 case was arc-sin transformed for linear modeling due to its bounded range of 0–1. We 954 started from the full model containing all parameters and their interactions and reduced

955 the model step-wise based on AIC scores (i. e. Akaike Information Criteria), as implemented 956 in the R function step. Visualizations and code applied for training and evaluation is 957 available in our GitHub repository.

958

959 *Testing sample number requirements—A* legitimate concern with complex neural networks 960 is that they may require vast amounts of training data and that typical skimming data sets 961 might be insufficient for them to be useful. We tested the robustness of our models to the 962 effect of the number of samples per species included in training by using from one to seven 963 samples per species as training set and the remaining as validation, with 50 replicates per 964 number of training samples. The batch size used in training was adjusted for the cases with 965 very few samples included, so that each training epoch included about 10 batches. We 966 included varKodes from 1Mbp to 200Mbp in both training and validation sets. In this case, 967 we applied the training parameters informed by our previous analyses: a *resnext101* 968 architecture, random initial weights, *CutMix* augmentation, and label smoothing for 30 969 epochs. We visualized the effect of the number of samples by plotting the average 970 validation accuracy of each sample against the number of training samples used in each 971 case. Visualizations and code applied for training and evaluation is available in our GitHub 972 repository.

973

974 *Testing the effect of data quality*—Most of the cases with low accuracy corresponded to 975 samples with low DNA yield (**Figure 2B**). We identified that DNA extraction yield was 976 significantly correlated with two metrics of DNA quality: average insert size and variation 977 in nucleotide composition along reads⁷⁹ (Figure 4). *varKodes* produced from these samples 978 may be visually distinct from other samples of the same species (**Figure 5**). For this reason, 979 we further tested whether sample quality in training or validation impacted accuracy. 980 Using both quality metrics, we identified the five lowest quality samples for each species. 981 We next produced training sets using six randomly chosen samples per species, varying the 982 number of low-quality samples included in training from zero to four. We included

983 varKodes from 1Mbp to 200Mbp in both training and validation sets. We repeated this for 984 30 replicates for each number of low-quality samples. Like our tests with varying sample 985 numbers, we applied the following training parameters: a *resnext101* architecture, random 986 initial weights, *CutMix* augmentation, label smoothing for 30 epochs. For the validation set, 987 we separately recorded the accuracy for high- and low-quality samples. We then visualized 988 the effect of inclusion of low-quality samples in the training set by observing the 989 distribution of validation accuracies for high-quality and low-quality samples across the 990 range of number of low-quality samples included in the training set. Visualizations and 991 code applied for training and evaluation is available in our GitHub repository.

992

993 *Implementation of varKoder*—Following all the tests described above, we implemented the 994 optimal neural network training strategies in a python program named **varKoder**. 995 *varKoder* can process, train and query varKodes and is freely available on our GitHub: 996 https://github.com/brunoasm/varKoder. Because it employs standard neural network 997 frameworks (namely, *pytorch*⁸⁸, *fastai*⁸⁹, and *timm*⁹⁰), any of the image classification models 998 and training hyperparamenters available now or in the future via these libraries can be 999 easily adapted and applied to varKode classification. Moreover, we have implemented a 1000 multi-label model as the default to increase robustness to low-quality varKodes with little 1001 diagnostic information in the training set. This was done by using an asymmetric multi-1002 label loss function⁸¹ instead of the standard cross-entropy loss function used in single-label 1003 classification. Analyses used development versions of *varKoder* starting with v.0.8.0. 1004 Improvements suggested during the peer-review process are now implemented in 1005 *varKoder* v.1.1.0.

¹⁰⁰⁶ *varKoder* **evaluation and comparison to alternatives**

1007 *varKoder*—To test *varKoder* performance on a complex dataset spanning multiple 1008 taxonomic levels and varying phylogenetic depths, we used the Malpighiales dataset 1009 including genera in Elatinaceae, Chrysobalanaceae and Malpighiaceae. Species of 1010 *Stigmaphyllon* (Malpighiaceae) were labeled with species, genus, and family names; all

1011 other samples were labeled with genus and family names. We tested the performance of 1012 *varKoder* in each sample with leave-one-out cross-validation. For each sample, we retained 1013 it as validation and trained a neural network using all the other samples. In preliminary 1014 assessments, we found that a ViT⁷¹ architecture combined with a multi-label model 1015 sometimes led to instability in training for some datasets. For that reason, we used a two-1016 step approach. Models first were pre-trained for 20 epochs as single-label, using the least 1017 inclusive taxonomic assignment available for each sample and a base learning rate of 0.05. 1018 Next, we trained for an additional 10 epochs using the pre-trained weights but with a much 1019 smaller learning rate (0.005) and a multi-label output. Training samples included varKodes 1020 from 500 Kbp to 200 Mbp, and we recorded validation accuracy separately for varKodes 1021 produced from each amount of data. We used an arbitrary confidence threshold of 0.7 to 1022 make predictions in the multilabel models. For validation samples, we deemed a prediction 1023 correct if only the correct taxon was predicted for each taxonomic rank (i.e., species, genus, 1024 family). We deemed a prediction incorrect if one or more predictions passed the threshold 1025 for a taxonomic rank, but none match the actual label. When predicted labels included both 1026 the correct and incorrect taxa, we deemed it ambiguous. If the output prediction included 1027 no taxon with confidence above the threshold, we considered it as inconclusive. As metrics 1028 across all samples, we used prediction and recall, averaged across all predictions. We 1029 visualized the fraction of correct, incorrect, ambiguous, and inconclusive samples for each 1030 taxonomic rank and each amount of data used to produce varKodes. The code to reproduce 1031 training conditions and evaluation tests is available on GitHub.

1032 To test the joint effect of neural network architecture and image representation method, 1033 we applied this cross-validation approach to all combinations of three image 1034 representations and four neural network architectures. The architectures tested included: 1035 (1) *ResNeXt101⁷⁶*, the optimal convolutional neural network architecture in our initial tests, 1036 (2) ViT^{71} , a transformer-based architecture that became available after our initial testing, 1037 (3) a neural network with two convolutional layers processing vectorized k-mer counts, **1038** following Fiannaca et al⁴⁴ and (4) a multi-layer perceptron formed by a series of fully 1039 connected layers as specified in Millán Arias et al⁴². The two latter have been previously 1040 employed for *fCGR* data. The three representations tested include *varKodes* and *rfCGRs* as

1041 developed here, and *fCGRs* as estimated by iDeLUCS⁹². In the latter case, we used iDeLUCS 1042 functions to produce *fCGRs* as 2D python arrays of k-mer counts. Next, we rescaled these 1043 counts to the range of 0-255 and rounded them to the nearest integer. These arrays were 1044 then saved as 8-bit png images. All code used in *varKoder* analyses is available on GitHub.

1045 *Skmer*—To compare *varKoder* with alternative methods, we used fastq files cleaned and 1046 subsampled by *varKoder* as input files to *Skmer*. In this case, we also used leave-one-out 1047 cross-validation to evaluate performance. For each amount of input data (500Kbp to 1048 200Mbp), we cycled through all samples, constructing a *Skmer* database with the "*skmer* 1049 *reference*" command and including all samples but one and default settings. We then used 1050 the "*skmer query*" command with default settings on the sample left out and deemed the 1051 identification as correct if the sample in the reference database with closest estimated 1052 genetic distance had the correct taxon label. Because *Skmer* could always query a sample 1053 and there is no objective criterion to consider matches beyond the best match, the output 1054 predictions can only be correct or incorrect, but not inconclusive or ambiguous. We 1055 visualized the results similarly as we did with *varKoder*. The code to reproduce *Skmer* 1056 analyses is available on GitHub.

1057 *Conventional plant barcodes* — To infer phylogenies from our genome skim data (Figure 1), 1058 we applied the *PhyloHerb* bioinformatic pipeline¹¹⁸, which has been applied recently to a 1059 taxa ranging from algae to flowering plants^{119–121}. Briefly, this pipeline works as follows: for 1060 plastid loci, *PhyloHerb* maps raw short reads to a database of land plant plastid genomes. 1061 Mapped reads are then assembled into scaffolds using *SPAdes*¹²² and plastid loci are 1062 identified using nucleotide BLAST searches with a default e-value threshold of 1e-40. 1063 *PhyloHerb* then outputs orthologous plastid genes into individual FASTA files, which are fed 1064 directly into MAFFT v7.407¹²³ for alignment. Alignments are then concatenated into a 1065 super matrix using the 'conc' function within the *PhyloHerb* package. Phylogenies for both 1066 individual locus and the concatenated alignment were inferred with IQTREE v2.0.6 using 1067 the GTR+GAMMA model with 1000 ultrafast bootstrap replicates¹²⁴.

1068 To recover the traditional plant barcodes, *rbcL*, *matK*, *trnL-F*, *ndhF*, and ITS, from our 1069 Malpighiales genome skim data, we applied GetOrganelle v1.7.7.0¹²⁵ and *PhyloHerb*

1070 v1.1.1¹¹⁸ to automatically assemble and extract these DNA markers, respectively. Briefly, 1071 the complete or subsampled genome skim data were first assembled into plastid genomes 1072 or nuclear ribosomal regions using *GetOrganelle* with its default settings. Next, *PhyloHerb* 1073 was applied to extract the relevant barcode genes using its built-in BLAST database. To test 1074 whether these traditional barcodes provided accurate identification to species, genus, and 1075 family, we ran an all-by-all BLASTn analysis for each individual gene across the same data 1076 subsampling schemes as *Skmer* and *varKoder*. BLAST targets were always drawn from 1077 assemblies using all the data available for each specimen, whereas queries included 1078 assemblies from input data amounts varying from 500 Kbp to 200 Mbp. Within each BLAST 1079 analysis for each one of the Malpighiales accessions, we deemed an identification to be 1080 correct if the best non-self BLAST hit came from the same taxon, and incorrect otherwise. 1081 We deemed it inconclusive if the locus could not be assembled for that amount of data. For 1082 concatenated barcodes, we produced a phylogenetic tree for each amount of data and 1083 deemed an identification to be correct if the sample with lowest patristic distance came 1084 from the same taxon. We deemed it to be inconclusive when none of the genes in the 1085 concatenated dataset could be assembled for a sample. We visualized results similarly to 1086 *varKoder*, separately for each conventional barcoding gene and for the concatenated 1087 dataset. The code to reproduce conventional barcode analyses is available on GitHub.

1088 *iDeLUCS*—To evaluate the performance of *varKoder* with another deep learning based 1089 sequence classifier, we applied the sequences assembled from the *PhyloHerb* pipeline to **1090** *iDeLUCS*⁹². We first used concatenated sequences of five traditional plant barcodes (*rbcL*, 1091 *matK, trnL-F, ndhF, and ITS)* assembled from input reads varying from 500 Kbp to 200 1092 Mbp. *iDeLUCS* was run with k-mer length of 6, 100 training epochs, 100 data augmentations 1093 per sequence, and the SGD algorithm for neural network optimization. All input sequences 1094 were set to be clustered into 10 groups (equal to the total number of species) and the 1095 accuracy was evaluated with the *cluster_acc* function implemented in *iDeLUCS*. We also 1096 applied the entire plastid genome and the nuclear ribosomal sequence assemblies 1097 (ETS+18S+ITS1+5.8S+ITS2+28S) in *iDeLUCS* with the same parameters to evaluate the 1098 impact of input data quality.

¹⁰⁹⁹ **Application in diverse taxa**

1100 *Species-level identification in plants, animals, fungi, and bacteria—For each of the four* 1101 organismal clades, we trained a multi-label model that included five species with at least 1102 three samples per species. For *Bembidion*, we included five species with five samples per 1103 species. For *Corallorhiza*, we included five species (or varieties) with at least five samples 1104 per species, except for *C. striata* var. *vreelandii* and *C. striata* var. *striata*, for which we 1105 included six and seven samples each, respectively. For *Mycobacterium tuberculosis*, we 1106 included representatives of five monophyletic *M.* tuberculosis lineages (L1, L2, L3, 1107 L4.1.i1.2.1, and L4.3.i2) with seven clinical isolates per lineage. Samples for *Bembidion*, 1108 Corallorhiza, and *M. tuberculosis* isolates all formed monophyletic groups, whereas 1109 *Xanthoparmelia* species did not. Since the *Xanthoparmelia* species were paraphyletic, we 1110 subsampled only monophyletic groups for model training. In this case, four species 1111 included three samples per species (*X. camtschadalis, X. mexicana, X. neocumberlandia,* and 1112 *X. coloradoensis*) and one species included five samples per species (*X. chlorochroa*). One 1113 potential confounding factor for the *Xanthoparmelia* model is that *Xanthoparmelia* is a 1114 lichen-forming fungus and thus genome skim data represents a chimera of fungal and algal 1115 genomes representing both partners in this unique symbiosis. Species of the algal symbiont **1116** *Trebouxia* are flexible generalists across fungal species *Xanthoparmelia*. Since these 1117 genome skims are a mix of both algal photobiont and fungus, we hypothesize that the 1118 accuracy of our model decreased because of the more generalist nature of *Trebouxia*¹²⁶. 1119

1120 For all four test cases, we applied default *varKoder* v.0.8.0 parameters for generating *rfCGR* 1121 images, training each model, and testing the accuracy of the trained model using the 'query' 1122 function. In all cases, we included all the available data for each training or validation 1123 sample. To test if trained models accurately predicted species identity, we queried them 1124 using extra genome skim samples not used for training but from the same species included 1125 in the model. We also tested genome skim test samples of species within the same genus 1126 *not* used in model training. As in the case of Malpighiales, we set the threshold to make a 1127 prediction equal to 0.7 and used the same criteria to consider a prediction correct, 1128 incorrect, inconclusive, or ambiguous. We separately evaluated results for taxa with

1129 representatives included in the training set and taxa used only as queries, without 1130 conspecific samples in the training set. The code to reproduce these analyses is available on 1131 GitHub.

1132

1133 *All eukaryotic families data set from SRA—Each accession was labeled with its family* 1134 identification obtained from NCBI. Because of the larger size of this dataset, a leave-one-out 1135 cross-validation approach would have been intractable. Therefore, we randomly selected 1136 80% of the samples in each family as the training set and used the remainder for validation. 1137 Similarly to Malpighiales, we used a two-step training method by pre-training as a single-1138 label model and finalizing with a multi-label model. Pre-training was done with a learning 1139 rate of 0.1 and a batch size of 300 for 30 epochs. Final training was done with the same 1140 batch size but a smaller base learning rate of 0.01 in 5 epochs with frozen body weights and 1141 three epochs with unfrozen weights. The code to reproduce these analyses is available on 1142 GitHub.

1143

1144 *All taxa from SRA*—For each accession, we created *rfCGRs* from 500Kbp to 10Mbp of data. 1145 Each accession was labeled with all the taxa in its taxonomic tree, as well as library strategy 1146 (RAD, GBS or WGS) and sequencing platform (Illumina, PACBIO, Nanopore or BGISEQ). We 1147 randomly selected 10% of the samples as validation set, and eliminated from validation 1148 samples all labels absent from the training set. We used a two-step training method. First, 1149 we pre-trained using a single-label strategy, using as labels the concatenation of library 1150 strategy, sequencing platform, kingdom, family and genus. For pretraining, we used a 1151 learning rate of 0.1, a batch size of 500 and 30 epochs. We then used the weights of this 1152 pre-trained model as starting weights for a multi-label model including all labels. We 1153 trained the model for additional 50 epochs with unfrozen body weights and 10 epochs with 1154 frozen weights, learning rate of 0.05 and batch size of 600. The code to reproduce these 1155 analyses is available on GitHub.

1156

1157 Environmental metagenome global identification—The downloaded soil metagenomes from 1158 Ma et al.⁹⁸ were labeled by source continent. Similarly to the eukaryotic family data set

1159 from SRA, we randomly selected 80% of the samples as the training set and used the

- 1160 remaining 20% as the validation set. We used a two-step training method by pre-training
- 1161 as a single-label model and finalizing with a multi-label model. Pre-training was done with
- 1162 a learning rate of 0.1 and a batch size of 64 for 30 epochs. Final training was done with the
- 1163 same batch size but a smaller base learning rate of 0.01 in 5 epochs with frozen body
- 1164 weights and three epochs with unfrozen weights. The code to reproduce all these analyses
- 1165 is available on GitHub.

¹¹⁶⁶ **Methods-only references**

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1217 **Data Availability**

- 1218 New data generated for this study is described in a Data Descriptor article containing
- 1219 accession numbers (doi to be updated upon acceptance).

1220 Code Availability

- 1221 Code used in the initial development and test of *varKoder* is available on Github, including
- 1222 all code used to produce figures for this manuscript. varKoder releases and source code are
- 1223 available at https://github.com/brunoasm/varKoder.