Annotation-free Learning of Plankton for Classification and Anomaly Detection

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11 Abstract

12 The acquisition of increasingly large plankton digital image datasets requires automatic methods 13 of recognition and classification. As data size and collection speed increases, manual annotation 14 and database representation are often bottlenecks for utilization of machine learning algorithms 15 for taxonomic classification of plankton species in field studies. In this paper we present a novel 16 set of algorithms to perform accurate detection and classification of plankton species with minimal 17 supervision. Our algorithms approach the performance of existing supervised machine learning 18 algorithms when tested on a plankton dataset generated from a custom-built lensless digital device. 19 Similar results are obtained on a larger image dataset obtained from the Woods Hole 20 Oceanographic Institution. Our algorithms are designed to provide a new way to monitor the 21 environment with a class of rapid online intelligent detectors.

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24 Author Summary

25 Plankton are at the bottom of the aquatic food chain and marine phytoplankton are estimated to be 26 responsible for over 50% of all global primary production [1] and play a fundamental role in 27 climate regulation. Thus, changes in plankton ecology may have a profound impact on global 28 climate, as well as deep social and economic consequences. It seems therefore paramount to collect and analyze real time plankton data to understand the relationship between the health of plankton 29 30 and the health of the environment they live in. In this paper, we present a novel set of algorithms 31 to perform accurate detection and classification of plankton species with minimal supervision. The 32 proposed pipeline is designed to provide a new way to monitor the environment with a class of 33 rapid online intelligent detectors.

34 Introduction

35 Plankton are a class of aquatic microorganisms, composed of both drifters and swimmers, which 36 can vary significantly in size, morphology and behavior. The exact number of plankton species is 37 not known, but an estimation of oceanic plankton puts the number between 3444 and 4375 [2]. 38 Traditionally, plankton are surveyed using either satellite remote sensing, where leftover biomass 39 is inferred indirectly through measurement of total chlorophyll concentration, or with large net 40 tows via oceanic vessels [3], with subsequent microscopic analysis of the preserved samples. 41 Satellite imaging methods are extremely accurate in terms of global geographic association and 42 very useful for broad species characterization but may present practical challenges in terms of 43 accuracy of the performed counts, species preservation and fine-grained characterization. The 44 analysis of preserved samples, instead, allows for fine grained classification and accurate counting 45 with narrow spatial sampling. More recently, real time observation of plankton species has been

46 made possible by novel instruments for high-throughput *in situ* autonomous and semi-autonomous 47 microscopy [4]. Such high-resolution imaging instruments make it possible to observe and study 48 spatio-temporal changes in plankton morphology and behavior, which can be correlated with 49 environmental perturbations. Sudden or unexpected changes in number, shape, aggregation 50 patterns, population composition or collective behavior may be used to infer anomalous conditions 51 related to potentially catastrophic events, either natural, like harmful algal blooms, or man-made, 52 like industrial run offs or oil spills. Intelligent systems trained on curated data could help establish 53 the characteristics of a healthy ecosystem and detect perturbations that may represent potential 54 threats. More importantly, given the diversity of plankton morphology and behavior across species 55 and the growing but still limited availability of high-quality labeled data sources, there is a need 56 for algorithms which require minimal supervision to classify and monitor plankton species with a 57 performance approaching that of supervised algorithms. Moreover, it is also desirable for such 58 algorithms to aid the discovery of new plankton classes, which cannot generally happen with 59 supervised classification techniques.

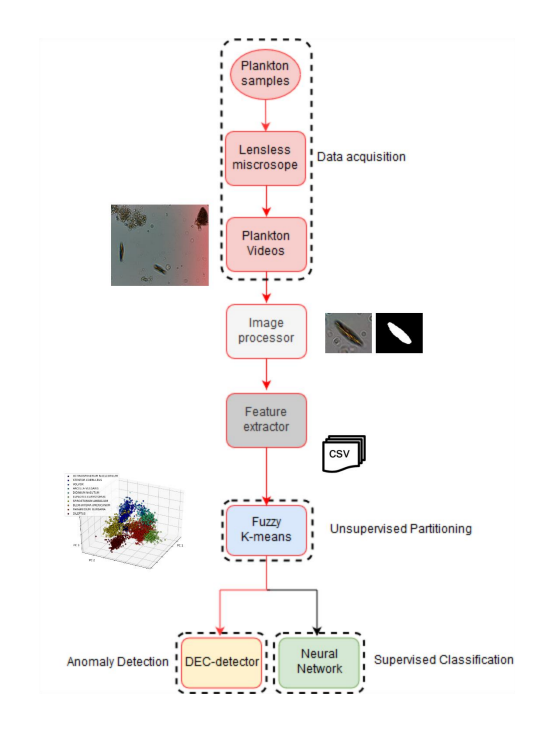
60 In this paper we propose a set of novel algorithms to reliably characterize and classify plankton 61 data. Our method is based on an unsupervised approach to overcome the limits of supervised 62 machine learning techniques, and designed to dynamically classify plankton from instruments that 63 continuously acquire plankton images. First, we evaluate the performances of our algorithms on 64 a mixture of ten freshwater plankton species imaged with a lensless microscope designed for in 65 situ data collection [5]. Next, we evaluate the performance of our algorithms on an image dataset 66 extracted from the Woods Hole Oceanographic Institution (WHOI) plankton database [6]. 67 Machine learning methods are becoming a popular way to characterize and classify plankton [7]– 68 [14]. A recent paper [15] explores the use of Convolutional Neural Networks to classify species of

zooplankton, by introducing an architecture named ZooplanktoNet. The authors claim that their customized architecture can reach higher accuracy compared to standard deep learning configurations, like VGG, AlexNet, CaffeNet, and GoogleNet. In [16] and [17], the authors use an SVM based algorithm to classify species with high accuracy from the WHOI dataset. In a recent Kaggle competition contest (http://www.kaggle.com/c/datasciencebowl), the authors developed a deep learning architecture named DeepSea [18] to perform accurate classification of plankton collected with an underwater camera. In [19] the authors combine features obtained with multiple kernel learning to achieve higher accuracy than classic machine learning algorithms. However, all these advancements use supervised learning algorithms that rely on large labeled training sets which are very difficult and time consuming to create. Although recent computational advances may reduce the annotation burden for large biological datasets [20], a high-performance unsupervised learning algorithm can provide an alternative for real time unbiased in situ analysis.

91 **Results**

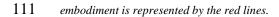
92 Plankton Classifier

93 We developed an unsupervised customized pipeline for plankton classification and anomaly 94 detection, that we named *plankton classifier*. The pipeline, shown in Fig 1, is tested on a collection 95 of videos containing ten fresh water species of plankton captured with a lensless microscope [5]. 96 Each video is ten seconds long and contains one or more species. As the method is unsupervised, 97 no labels are provided to the classifier during training. The plankton classifier consists of four 98 modules: an **image processor**, a feature extractor, an unsupervised partitioning module and a 99 classification module. The image processor examines each frame of video and generates cropped 100 images of each plankter. The feature extractor examines each plankter image and generates a 101 collection of features. The **unsupervised partitioning module** clusters samples by features into 102 classes. The classification module comprises of a neural network-based anomaly detector to both 103 perform classification based on the inferred labels and provide information to extend the database 104 in an unsupervised manner. A sample is considered an anomaly with respect to a class if the 105 extracted features are significantly different from the class average, as described below. The 106 classification module also includes a standard neural network classifier, for performance 107 comparison. See section materials and methods for a description of the modules in more details, 108 along with the methods considered and tested that led to our final design.





110 Fig 1. Schematic overview of the pipeline used to detect and classify plankton species with minimal supervision. Our preferred

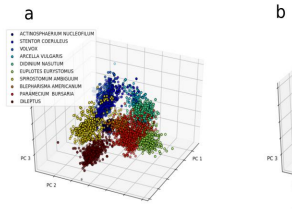


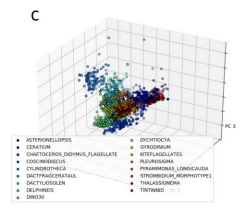
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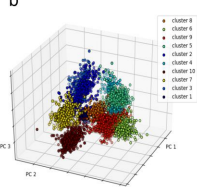
114 Unsupervised partitioning performance

115 First, the plankton classifier examines each frame of an acquired video and generates cropped 116 images of each plankter. A set of 131 features is then extracted, as described in Materials and 117 Methods. The unsupervised partitioning module uses such features to place each plankton sample 118 into one of Z classes. To automatically obtain the number of classes from the dataset, we have 119 designed a custom algorithm based on partition entropy (see Materials and Methods). We 120 evaluated the robustness of the implemented method on random subsets of the lensless dataset 121 with different sizes, ranging from three to ten species. The box plot indicating the distribution for 122 the estimated number of clusters Z among ten iterations can be observed in Fig 2e. The inferred 123 number of classes, Z, is correctly identified in every case. A comparison of the performance of this 124 algorithm against other existing methods is reported in the Supporting Information. Once we have 125 obtained the number of clusters, we compared three clustering algorithms (see Supporting 126 Information): k-Means, Fuzzy k-Means and Gaussian Mixture Model (GMM). Clustering 127 accuracy is evaluated using purity (see materials and methods). The Fuzzy k-Means algorithm 128 reaches a purity value of 0.934 (see Figs 2a, 2b), outperforming the standard k-Means (purity value 129 = 0.887) and GMM [21] (purity value = 0.886). A posterior analysis of the results of the GMM 130 reveals that this algorithm is not able to distinguish between *Blepharisma americanum* and 131 *Paramecium bursaria*, due to their nearly identical appearance in the acquired videos. The Fuzzy 132 k-Means algorithm is able to match the fuzziness exhibited by the plankton classes in parameter 133 space which explains the lower accuracy of the crisp algorithms (k-Means and GMM). Therefore, 134 we use the Fuzzy k-Means for our unsupervised classifier. A potentially important effect on the 135 performance of any clustering algorithm is the class imbalance. The lensless microscope dataset is 136 composed of 500 training samples for each of the ten considered species. To evaluate the impact

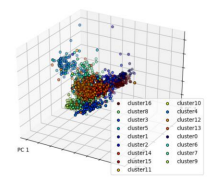
137 of class imbalance, we performed the following experiment: We have built a dataset where the 138 number of images of a species is a fraction (between 10% and 80%) of the number of images of 139 the other species. We then evaluate the purity of this dataset and repeat the procedure for all the 140 other species. Fig 2f reports the average performance over the ten datasets obtained as described 141 above, as measured by the purity. The algorithm is always able to infer the correct number of 142 species, without any overlap, with a minimum average purity value of 0.74 ± 0.09 (corresponding 143 to 80% of class imbalance) and a maximum average purity value equal to 0.90 ± 0.08 144 (corresponding to 10% of class imbalance), with a maximum purity value of 0.972. This result 145 shows that our pipeline can accurately cluster the data even in the case of strong class imbalance.



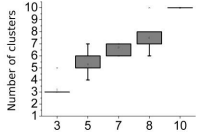


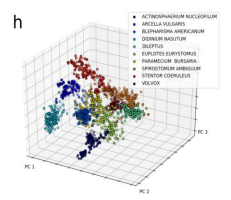


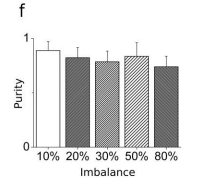


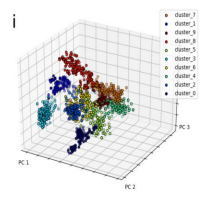












147 Fig 2. Unsupervised clustering results. a, b We performed a PCA analysis on the lensless digital microscope dataset to provide 148 a graphical representation of the data distribution into the features space. We plot the first three principal components that account 149 for $\sim 67\%$ of the total variance. We assigned different colors to the different plankton species. **a** Species are assigned using ground 150 truth labels. **b** Species are assigned to the most overlapping cluster resulting from the unsupervised partitioning procedure. **c**, **d** 151 Same analysis and procedure applied on the WHOI dataset. c Species are assigned using ground truth labels. d Species are assigned 152 to the most overlapping cluster, resulting from the unsupervised partitioning procedure. e Distribution of number of clusters 153 computed using our PE algorithm for a random subset of species in the lensless microscope dataset. Results are reported for different 154 initial number of species. f Effect of class imbalance. For each of the ten species included into the lensless microscope dataset, we 155 simulated class imbalance by increasing the number of images available to the clustering algorithm for the considered species. h, i 156 PCA analysis on the lensless digital microscope dataset provides a graphical representation of the data distribution into the deep 157 features space. The unsupervised partitioning using deep features is highly accurate. The first three principal components are plotted 158 and different colors to the different plankton species are assigned. h Species are assigned using ground truth labels. i Species are 159 assigned to the most overlapping cluster resulting from the unsupervised partitioning.

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161 Algorithm performance on features extracted using deep feature extraction

162 Feature selection is an important part of any unsupervised learning pipeline. Indeed, hand 163 engineering features introduces a degree of arbitrariness, which can be removed using a method 164 of automated feature selection. Deep feature extraction, which consists in training a neural network 165 architecture on either in- or out-of-domain data and use the last layer before prediction to extract 166 features [9][22], is one such method. We trained the model described in section *Convolutional* 167 *Neural Network (CNN) for deep features extraction* using the ten classes included in our lensless 168 microscope dataset. The model reached 99% of training accuracy, 99% of validation accuracy and 169 98% of testing accuracy on the dataset obtained using our lensless microscope. Finally, the 128 170 neurons from the fully connected layers preceding the output are extracted and used as features for 171 our pipeline. The PCA computed for the lensless microscope testing set among these features can 172 be visualized in Fig 2h. Fig 2i shows the results of the unsupervised partitioning procedure. The

underlying structure of the data set is very accurately captured, with a purity value of 0.98. Despite the fact that the accuracy obtained using deep feature extraction is slightly higher than the one obtained using the hand engineered features (purity of 0.980 vs 0.934), we decide to use the interpretable features described in Table 1. In fact, we think it is important that interpretability is maintained for the purpose of establishing a causal link between environmental perturbations and

morphological modifications. However, for the purpose of organism classification, the customized

179 deep feature extraction algorithm we implemented is a very viable alternative to the one proposed.

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182 Classification

183 **Supervised Classifier.** At this stage of the pipeline, all samples have been assigned labels which 184 have no correspondence to the actual plankton classes. We use the same trained clustering 185 algorithm to classify the test samples, assigning each sample to the closest centroid. Using the 186 trained Fuzzy k-means algorithm we reach a testing accuracy of 89%. Alternatively, one can use 187 the labels obtained by our unsupervised partitioning algorithm to train a supervised classifier. We 188 evaluated two algorithms: An Artificial Neural Network (ANN) and a Random Forest (RF) 189 classifier. Our ANN architecture consists of a collection of classifiers, each trained to detect one 190 plankton class. The RF approach consists in a set of decision trees to separate the training step 191 samples into the correct classes.

For comparison, a simple ANN classifier is trained using the labels provided by the unsupervised partitioning algorithm. The ANN is a massive parallel combination of single processing units which can learn the structure of the data and store the knowledge in its connections [23]. See

195 Materials and Methods for further information and for a detailed description of the implemented 196 architecture. The network is very shallow, providing an efficient feature selection process. The 197 ANN classifier reaches a validation accuracy of 99% and a testing accuracy of 94.5%. Figs 3c and 198 3d report the ROC curves and the confusion matrix obtained by testing the trained ANN classifier 199 on our ten species plankton dataset. The ROC curves are close to a perfect classifier and the 200 confusion matrix is almost diagonal with minor overlap between two pairs of species: *Blepharisma* 201 americanuum-Paramecium bursaria and Spirostomum ambiguum-Stentor coerouleus. This 202 misclassification is primarily due to the similarity in the shape, size and texture of the two pairs of 203 species, influencing both the unsupervised training clustering and the subsequent testing of the 204 supervised classifier.

An alternative classifier method employs a Random Forest (RF) approach, a popular ensemble
learning method used for classification and regression tasks.

We train an RF algorithm using the labels provided by the unsupervised classifier and reach an accuracy of 94%. For comparison, we train the same RF algorithm using the actual labels (ground truth) of the training set and reach an accuracy around 98%, proving that our unsupervised classification approach performs comparably well with respect to the correspondent supervised approaches for the trained classifier. Since the ANN performs marginally better than the RF classifier, we propose the former for a pipeline. In the next section, we will present an alternative classification method

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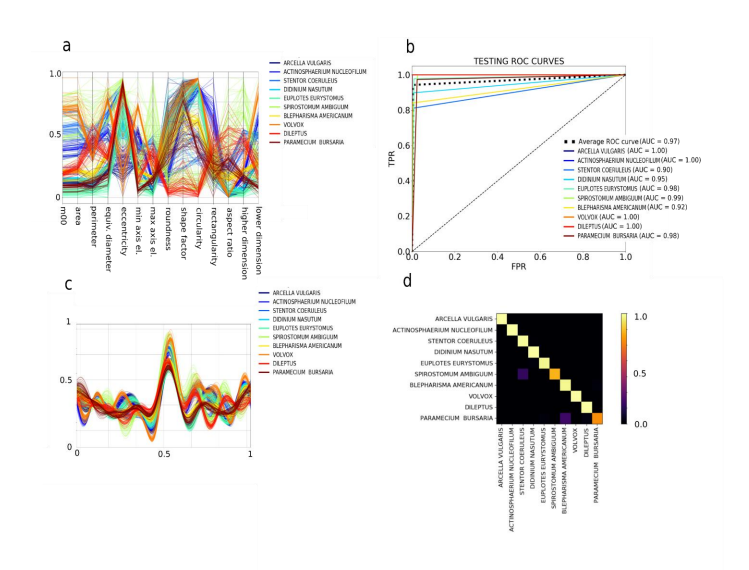
215 Anomaly Detector

When deployed in the field, microscopes will encounter species that have never been seen before,so it is essential that such samples are detected and correctly identified as anomalies. For a given

218 class, a sample is considered an anomaly if the sample features are significantly different from the 219 feature average for the class. Algorithms for anomaly detection based on the separation of the 220 features space have been successfully used to identify the intrusion in computer networks for 221 security purposes [24]. Two anomaly detectors are implemented and compared; a state of the art 222 one-class SVM¹⁵ and a customized neural network we call a Delta-Enhanced Class (DEC) detector 223 that combines classification with anomaly detection. The one-class SVM algorithm uses a kernel 224 to project the data onto a multidimensional space and can be interpreted as a two class SVM 225 assigning the origin to one class and the rest of the data to another class. It then solves an 226 optimization problem determining a hyperplane with maximum geometric margin, i.e., a surface where the separation between the two sets of points is maximal, that will be used as decision rule 227 228 during the testing step.

A customized one-class SVM is implemented by normalizing the testing samples using the training data belonging to a single class. In this way, there will be a significant difference in the absolute value obtained for the anomaly (out-of-class) samples compared to the in-class samples, improving the accuracy of the SVM. The one-class SVM so designed reaches an average testing accuracy of (93.5 \pm 6.0) %, with high accuracy in both anomaly detection and classification.

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Fig 3. Feature space representation and classification performances. a, b Multidimensional visualization of the geometric subset of the ten species in the lensless microscope dataset, obtained using the following methods (see Supporting Information): a Andrew's curve. b Parallel coordinates. c ROC curves obtained for the neural network classifier trained on the labels provided by the clustering algorithm for the lensless microscope dataset. d Corresponding confusion matrix.

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241 We now describe an alternative ANN-based approach that simultaneously performs classification

and anomaly detection. As demonstrated above, a single layer ANN is able to satisfactorily classify

243 plankton data from our in-house dataset. However, to effectively approach the anomaly detection

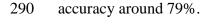
244 step, we designed a deep neural network called Delta-Enhanced Class (DEC) detector (see 245 materials and methods for further details). One DEC detector must be trained for each of the 246 training species. Therefore, we train ten DEC detectors, one for each of the species of plankton 247 identified in the unsupervised learning step. This procedure affords excellent accuracy on both 248 classification and anomaly detection, on both real and simulated plankton data (see Fig 4), with an 249 average testing accuracy on real data of 98.8 \pm 2.4 %, an average anomaly detection testing 250 accuracy of 99.2 \pm 0.7 % and an average overall testing accuracy of 99.1 \pm 0.9 % (see Fig 4b for 251 details). The confusion matrices in Fig 4a demonstrate the discrimination power of our algorithm. 252 The DEC detector outperforms the alternative one-class SVM classifier in both supervised 253 (average accuracy equal to 95%) and unsupervised (average accuracy equal to 93.5%) 254 configurations. It is worth reporting that the unsupervised one-class SVM reached a minimum 255 overall accuracy of 79%, compared to 97.2% for the DEC detector (minimum values correspond 256 to *Paramecium bursaria* detector). To test the overall performance of our method, we produce a 257 dataset of surrogate plankton organisms. For each different species, we test the corresponding DEC 258 detector architecture using a surrogate species created with a feature-by-feature weighted average 259 of all the species in our dataset. Starting with a uniform weight distribution, we increase the weight 260 for the species corresponding to the trained DEC detector architecture up to 0.9 (steps of 0.1), 261 obtaining 9 different surrogate species (see Fig 4d for an average parallel coordinates plot, showing 262 the resulting distributions for the species *Spirostomum ambiguum*). The aim of this robustness test 263 is to simulate the acquisition of an unknown species, whose features are increasingly closer to the 264 features of the class correspondent to the detector, up to a maximum of 90% similarity. As Fig 4e 265 shows, our classifier can recognize the synthetic species as an anomaly with an average accuracy 266 higher than 98% if the similarity between the synthetic and the real species is up to 30%, and it

can maintain an average accuracy of over 82.6% if the species similarity is up to 50%. Accuracy
of anomaly detection severely decreases if the species similarity is over 50%, reaching the
minimum value of 37.5%.

270 Plankton classifier performance on the WHOI dataset

271 The WHOI provides a public dataset comprising millions of still monochromatic images of 272 microscopic marine plankton, captured with optical Imaging FlowCytobot an (https://mclanelabs.com/imaging-flowcytobot/). To use this dataset as a benchmark to test our 273 274 unsupervised classifier, we extract a set of 128 features from a collection of 40 species of plankton 275 (100 images per species, randomly selected), using both the segmented binary image and the 276 portion of the gray-scale image containing the plankton cell body. A full description of the species 277 selection process is reported in the Supporting Information. The features set is identical to the one used for the lensless microscope dataset, except for the absence of three-color features, as the 278 279 lensless microscope is a color-based sensor, while the Imaging FlowCytobot is monochromatic. 280 Figs 2c, 2d show the results of our pipeline applied on the normalized features set. The algorithm 281 reaches an overall purity value of 0.715 for the 40 WHOI species that we selected. The ability of 282 our pipeline to distinguish between inter-species plankton morphology can be further observed 283 comparing Fig 2c, which represents the PCA space corresponding to a subset of 18 of the 40 284 species for the ground truth dataset, and Fig 2d, which represents the corresponding PCA space 285 resulting from the unsupervised partitioning algorithm. A complete PCA representation for the 40 286 species can be found in Supporting Information. We trained a random forest algorithm using the 287 labels provided by the unsupervised partitioning with a train-test ratio of 80:20, obtaining a 288 classification accuracy around 63%. For comparison, we have trained a supervised random forest

algorithm using the ground truth labels on the extracted features, obtaining a classification



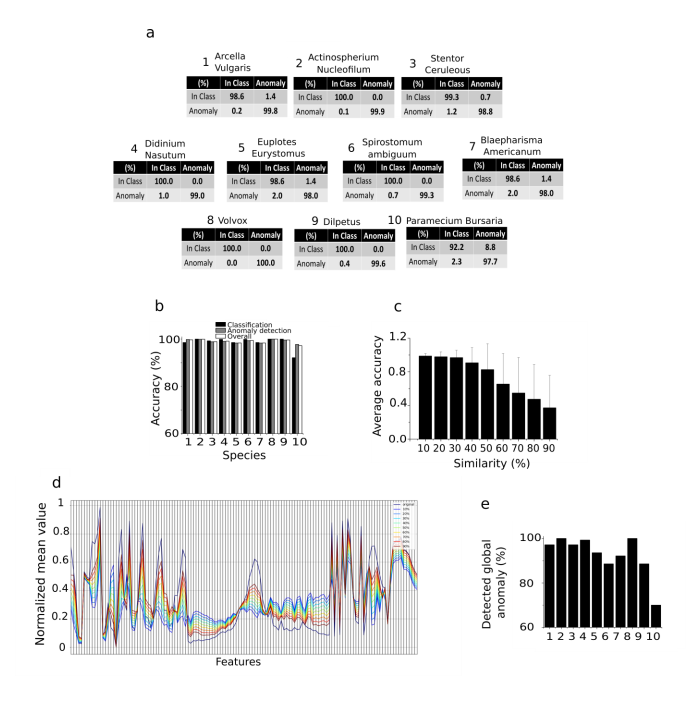


Fig 4. Delta-Enhanced Class detector performances and results. a Confusion matrix corresponding to each of the ten neural
 networks trained on the lensless microscope dataset. b Overall testing accuracy performances for each of the ten testing classes.
 The number used on x axis to label each species correspond to the species number in panel a. c-d DEC detector anomaly detection

295 performances tested on in silico generated data. d Testing accuracy performances for varying percentage values of in silico species 296 similarity with the trained species. e Example of average features space parallel coordinates plot for the in-silico species obtained 297 using the species Spirostomum Ambiguum. By increasing the similarity, the features of the surrogate species approach the features 298 of the real species, resulting in an increased average anomaly misclassification rate, decreasing the overall accuracy levels. e 299 Detection of unknown species. The panel shows the percentage of samples detected by all the DEC detectors as anomaly, when 300 removing one training species from the set, for each of the ten training species. These numbers reflect the level of accuracy of the 301 proposed algorithm in detecting unseen species. The number used on x axis to label each species correspond to the species number 302 in panel **a**.

303 The plankton classifier can reveal unseen species

304 We have demonstrated that our DEC neural networks are able to classify a sample as either a 305 training class (i.e., the plankton species used to train the detector) or as an anomaly. If a sample is 306 discarded by all the implemented detectors, it could either represent an intra-species anomaly (i.e., 307 species included into the training set) or a sample belonging to an unseen species (i.e., species not 308 included in the training set). The former represents the basis for using the proposed pipeline for 309 real-time environmental monitoring, and its implications are discussed in the next section. We now 310 test the potential of our pipeline to detect new species. We remove one class from our unsupervised 311 partitioning ensemble set, consider it as never before seen and compute the number of testing 312 samples detected as anomaly by all the remaining DEC detectors. This number indicates the 313 algorithm accuracy in detecting new species. We repeat the procedure for each class. The average 314 detection accuracy is 98.3 ± 10.1 % (see Fig 4e), demonstrating the ability of the pipeline to detect 315 the presence of a new species. If two or more unseen species are detected, they will be stored as 316 anomalies. As this group of anomalies grows, a human expert may determine offline the actual 317 labels for these new species, thus allowing a DEC detector to be trained for each new species. 318 Alternatively, the samples corresponding to unseen species may be clustered and classified by the

unsupervised partitioning step of our pipeline, reducing the number of new species that must beexamined by a human.

321 **Discussion**

The plankton classifier described in this paper provides the foundation for a robust, accurate and scalable mean to autonomously survey plankton in the field. We have identified interpretable and non-interpretable image features that work with our algorithms to perform an efficient clustering and classification on plankton data using minimal supervision and with a performance accuracy comparable to supervised learning algorithms [16]. Instead of labeling thousands of samples, an expert need only identifying one member of cluster to label all the samples of the cluster.

We introduced a neural network that performs classification by learning the shape of the feature space and uses this information to identify anomalies. The network uses a novel unbiased methodology of feature-to-feature comparison of a test sample to a random set of training samples. While most of the existing classification methods require various degrees of user input, our method is automated, without sacrificing performance accuracy or efficiency.

All features the plankton classifier relies upon are extracted from static images. However, our custom lensless microscope captures 2D and 3D dynamic of plankton. While this dynamic information is not considered in the analysis presented here, motion data can increase the dimensionality of the feature space, by adding spatio-temporal "behavioral" components, and may improve the performance of classifiers and anomaly detectors. This is particularly valuable in cases where species have considerable overlap in morphology feature space, as seen with *Blepharisma americanuum* and *Paramecium bursaria*, and *Spirostomum ambiguum* and *Stentor coerouleus*,

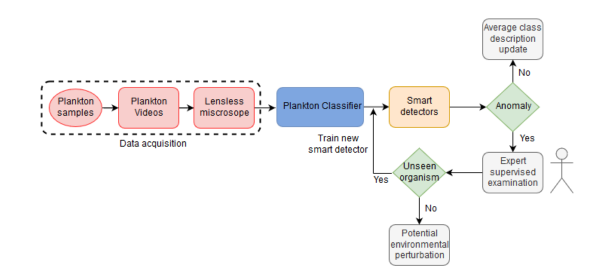
340 shown in the confusion matrices in Fig 3d. Currently, existing large plankton datasets, like the 341 WHOI used in our validation experiments, are based on static images, but as the cost of video-342 based *in situ* microscopes drops and their deployment increases, we believe datasets that include 343 spatio-temporal data will become available and the use of such features will gain importance.

344 Deploying smart microscopes capable of real-time continuous monitoring will give biologist an 345 unprecedented view of plankton in situ. The adoption of an unsupervised unbiased pipeline is a 346 significant step ahead in the development of a real-time "smart" detector for environmental 347 monitoring. Several high-resolution acquisition systems for real-time plankton imaging already 348 exist [25] and could adopt the pipeline proposed into this paper. Fig 5 shows a high-level 349 representation of a continuous environmental monitoring system in the form of a flow chart, 350 showing an example of how the detector could be coupled to the computational pipeline we 351 designed. Once the descriptors have been extracted from the acquired videos, it is possible to use 352 them to build a set of DEC detectors. It is important to stress that the size of the data likely to be 353 acquired, or already present in the databases, makes neural networks the obvious choice to carry 354 out the analysis due to their unsurpassed scalability. Our newly designed and customized DEC 355 detector neural architecture for plankton classification and anomaly detection is a functional and 356 efficient example of such algorithm. Moreover, neural algorithms can infer non-linear 357 relationships between features (input) and correlate them with the class description (output) 358 without making any assumptions on the underlying learning model. Hence, the classification 359 depends only on the extracted features. Every time the network identifies a species belonging to a 360 specific class, the average set of morphological features is then updated, thereby further qualifying 361 the class morphology phase space. If an anomaly is detected, it may be sent to an expert for a 362 supervised examination. The expert will determine whether that sample could be a species not

363 represented in the training set, or if it belongs to an existing training class, but its morphological 364 features deviate significantly from the average features space of the corresponding class. In the 365 former case, a new smart detector will be trained offline, so that the training set is dynamically 366 expanded, and the system will provide a continuous monitoring of the aquatic environment using 367 the human expert-in-the-loop paradigm. In the latter case, the identified anomalies may represent 368 local environmental perturbations, either natural or man-made. Further work is needed to assess 369 the validity of such hypothesis. An additional re-training step may be necessary to update the 370 algorithms. Our pipeline is based on local analysis using a low powered device, capable of image 371 capture and processing, classification and anomaly detection. Coupling such platform with a local 372 (laptop, server) or cloud-based system where the training step may occur could provide the 373 flexibility and resources needed to close the loop and generate the training data the low power 374 platform can use for classification. Examples of systems that use this paradigm are already present 375 in the literature [26], and we hope the availability of computational paradigms like the one we 376 propose may increase the research in the field. A high-resolution plankton acquisition system 377 placed in the water and powered with our unsupervised pipeline may enable the development of 378 real time continuous smart environmental monitoring systems that are fundamentally needed to 379 stakeholders and decision-making bodies to monitor plankton microorganisms and, consequently, 380 the entire aquatic ecosystem [27].

Finally, it is interesting to consider if such unsupervised approach can be utilized for different data types, thus widening the potential applicability and interest of the technique. While an extensive analysis of the performance of our pipeline on diverse set of data is beyond the scope of this work, it is worth commenting that the algorithms we use are general and pose no evident drawback to their application to other cell types. Particularly, the features our classifier uses to cluster the

images do not include anything specific to plankton species (e.g. detection and estimation of number of flagella or other organelles.) Moreover, the proposed Deep Feature extraction method is even less dependent on the kind of data under study and may increase the applicability to other cell types. Thus, we expect the method to be potentially useful to other biological imaging fields.



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391 Fig 5. Proposed real-time smart environmental monitoring pipeline.

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401 Material and methods

The proposed unsupervised pipeline (i.e., the plankton classifier) shown in Fig 1, consists of four modules: an **image processor, a feature extractor, an unsupervised partitioning module** and a **classification module**. In the following paragraphs we provide a description of the modules in more details, along with the methods considered and tested that led to our final design.

406 Image Processing

407 Each video consists of ten seconds of color video (1920x1080) captured at 30 frames per second. 408 Background subtraction is applied to each frame to detect the swimming plankton in the image. A 409 contour detector is applied to the processed image to create a bounding box around each plankter. 410 Because of instrument design, organisms can swim in and out of the field of view (FOV) during 411 acquisition. Our algorithm automatically selects organisms which are fully contained inside the 412 FOV by checking whether the bounding box touches the borders of the FOV. In this way, the 413 images we obtain will be only of fully visible organisms. The resulting cropped image is then 414 saved. From this collection of images, a training set of 640 images (500 training and 140 testing) 415 is selected for each class. An image processor module for static images has also been implemented 416 for benchmarking the plankton classifier on existing plankton datasets (e.g., the WHOI dataset; 417 See Supporting Information for further details.).

418 **Feature Extraction**

For each plankter image, 131 features are extracted from four categories: geometric (14), invariant
moments (32), texture (67) and Fourier descriptors (10). Geometric features include area,
eccentricity, rectangularity and other morphological descriptors, that have been used to distinguish

422 plankton by shape and size [16]. The invariant Hu [28](7) and Zernike moments [29] (25) are 423 widely used in shape representation, recognition and reconstruction. Texture based features encode 424 the structural diversity of plankton. Fourier Descriptors (FD) are widely used in shape analysis as 425 they encode both local fine-grained features (high frequency FD) and global shapes (low frequency 426 FD). A full list of the features we have selected is reported in Table 1. These features span a 131-427 dimensional space, capturing the biological diversity of the acquired plankton images. Figs 3a and 428 3b demonstrate as an example, the discriminating power of the geometrical features for the ten 429 evaluated species.

Class	Number	Description
Geometric feature	14	Area (pixels), area (0-th order moment), perimeter,
		eccentricity, rectangularity, roundness, shape factor,
		width and height (minimum fitting rectangle),
		circularity, major and minor axis (fitting ellipse),
		equivalent diameter, convexity.
Hu moments	7	Hu moments computed from the normalized central
		image moments.
Zernike moments	25	Zernike moments up to order 5.
Image intensity features	8	Blue/green channels ratio, red/green channels ratio,
		red/blue channels ratio, gray levels histogram
		statistical features (skewness, kurtosis, mean value,
		standard deviation and entropy)
Haralick features	13	The first 13 features as proposed by Haralick in ¹³ ,
		computed from the Gray Scale Co-occurrence Matrix
		(GSCM).
Local binary patterns	54	Local binary patterns summarize structures of the
		image comparing each pixel to its neighborhood
Fourier descriptors	10	Fourier descriptors are contour-based features
		invariant with respect to rotation, scaling and
		translation.

Table 1: List of morphological features extracted from the processed images. See Supporting Information for a detailed
 explanation.

433 Convolutional Neural Network (CNN) for deep features extraction

434 We implemented a deep CNN using eight convolutional layers and two fully connected layers, as 435 described in Fig 6. We customized our architecture to be invariant with respect to rotation, similar 436 to what has been done in [18]. Each input sample is rotated four times at multiples of 90 degrees, 437 and all the tensors resulting from the features extraction module are concatenated and used to train 438 the fully connected layers. The neural network has been trained for 60 epochs, using stochastic gradient descent with learning rate equal to 10^{-5} , using data augmentation by means of translation, 439 440 zooming, and rotation. It is worth noticing that the implemented rotational invariance module actually performs a data augmentation operation, and it is indeed useful when partial training data 441 442 are available.

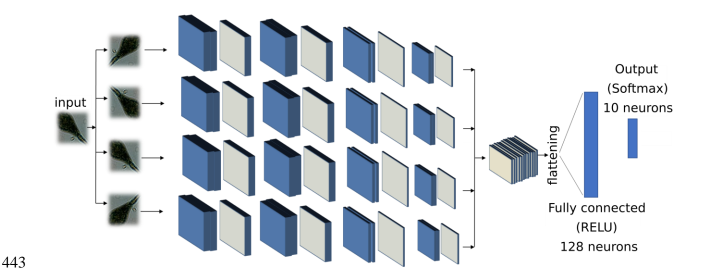


Fig 6. Deep features extraction. Deep CNN implemented for the purpose of deep features extraction. The blue layers represent convolutional layers, the grey ones represent a max pooling 2D operation. The fully connected layer with 128 neurons output has been used as feature set to the subsequent modules in our pipeline.

447 Unsupervised Partitioning

448 **Partition Entropy (PE)**

449 The Partition Entropy (PE) coefficient is defined as:

450
$$PE = -\frac{1}{N} \sum_{i=1}^{N} \sum_{j=1}^{K} u_{ij} * \log(u_{ij})$$
 (1)
451

452

460

The coefficient is computed for every j in [0, K] and takes values in range [0, log(K)]. The estimated number of clusters is assigned to the index j^* corresponding to the maximum PE value, PE(j^*). The lower the PE(j^*), the higher the uncertainty of the clustering. We repeat this procedure ten times and obtain a distribution of j^* . Finally, the estimation of the number of clusters *Z* is the mode of this distribution.

458 Clustering accuracy

459 Clustering accuracy is evaluated using purity:

$$purity = \frac{1}{N} \sum_{k} \max_{j} |w_k \cap c_j|$$
(1)

461 where the class k is associated to the cluster j with the highest number of occurrences. A purity 462 value of one corresponds to clusters that perfectly overlap the ground truth. Purity decreases when 463 samples belonging to the same class are split between different clusters, or when two or more 464 clusters overlap with the same species. We have implemented a purity algorithm capable of 465 checking for these occurrences and automatically adapt to the correct number of non-overlapping 466 clusters (see Supporting Information).

467 Classification algorithms

468 Random Forest

469 Random Forests (RF) is a popular ensemble learning method [30] used for classification and 470 regression tasks, introduced in 2001 by Breiman. Random forests model providing estimators of 471 either the Bayes classifier or the regression function. Basically, RF work building several binary 472 decision trees using a bootstrap subset of samples coming from the learning sample and choosing 473 randomly at each node a subset of features or explanatory variables [31]. Random forests are often 474 used for classification of large set of observations. Each observation is given as input at each of 475 the decision tree, which will output a predicted class. The model outputs the class that is the mode 476 of the class output by individual trees [32].

Let us consider a set of observations $x_1, x_2, ..., x_N$, with $x \in \mathbb{R}^m$. The decision tree is designed as 477 478 follows: we extract N times from the set of training observations (with replacement), for a each of the total number of decision tree. We specify the number of features m^* to consider for the tree 479 480 growing, with $m^* \ll m$. For each of the nodes in the tree, the algorithm randomly selects m^* 481 features and calculates the best split for that node. The trees are only grown and not pruned (as in 482 a normal tree classifier [33]. The split's aim is to reduce the classification error at each branch. In 483 detail, the algorithm considers an entropy-based measure trying to reduce the amount of entropy 484 at each branch, selecting, with such a procedure, the best split. A possible choice is the Gini index:

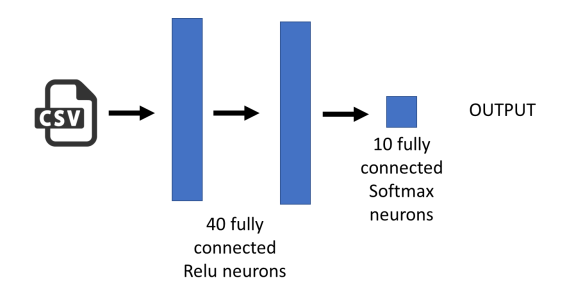
485

486
$$G_m = \sum_{i=1}^{K} p_{im}(1 - p_{im})$$
 (27)

Where G_m is the Gini Index for branch at level m in the decision tree, and p_{im} is the proportion of observations assigned to class *i*. Minimizing G_m , means to decrease the heterogeneity at each branch, i.e., a best split will correspond to a lower number of class in the children nodes. The algorithms continue in growing trees until convergence on the entropy-based on the generalization error [32].

493 Neural Networks

494 An artificial neural network (or multi-layer perceptron) is a massive parallel combination of single 495 processing unit which can acquire knowledge from environment through a learning process and 496 store the knowledge in its connections [23]. Classification is one of the most active research and 497 application areas of neural networks. In this work we used an artificial neural network to build a 498 classifier able to predict the species for each observation extracted using the shadow microscope. 499 Fig. 2 shows the developed architecture. The network is very shallow, with two hidden layers of 500 40 neurons and an output layer with as much neurons as the number of species to classify. As 501 reported in the main text of this manuscript, we used a training dataset with 10 species, thus the output layer is made up of k neurons, where k is the number of clusters obtained using the 502 503 unsupervised clustering. As Fig 7 shows, the developed NN uses RELU activation function and 504 dropout to reduce the overfitting. The network was trained using 200 epochs, Root mean square 505 as an optimizer, a learning rate $\lambda = 0,005$ and categorical cross-entropy as loss function. The 506 training requires 50 seconds on a MAC book PRO, core i7 – 2.9 GHz, solid state disk and 16 GB 507 of RAM. The neural network has been implemented using KERAS, a powerful high-level neural 508 network API running on top of TensorFlow.





511 Fig 7. ANN architectures implemented for classification based on the extracted features.

512 Anomaly Detection

513 One Class SVM

We adopted the one class SVM described by Scholpoff in [34]. Let us consider a set of N observations: $\{x_i, y_i | x_i \in \mathbb{R}^m, y_i = \pm 1\}$. Where x_i is a m-dimensional real vector and $y_i = \pm 1$ simply imply that the set contains normal observations belonging to a certain class. The one-class SVM is a classification algorithm returning a function which takes ± 1 in a "small" region capturing most of the data points, and ± 1 elsewhere. Let ϕ be a feature map that map our observations set x_i , into an inner product space such as the inner product for the image of ϕ can be evaluated using some simple kernel:

521

522
$$k(x,y) = \phi(x)\phi(y)$$
(28)

524 The strategy of the one class SVM is to map the data into the kernel space and separate the data 525 from the origin with maximum margin, defining a hyperplane as: 526 $H(x) = w \phi(x) - \rho$ 527 (29)528 Meaning that we want to maximize the ratio $\frac{\rho}{\|w\|}$, corresponding to the hyperplane's distance 529 530 from the origin. In order to solve this maximization problem, we have to solve a quadratic 531 problem: 532 $\min_{w,\xi,\rho}\frac{1}{2} \|w\|^2 + \frac{1}{\nu l} \sum_i \xi_i - \rho$ 533 (30)534 subject to $w \phi(x) \ge \rho - \xi_i, \ \xi_i \ge 0$. 535 536 Where $\phi(x)$ is the feature mapping function that maps observations x into a feature space, ξ_i is a 537 538 slack variable for outlier that allows observations to fall on the other side of the hyperplane , $\nu \in [0,1)$ is a regularization parameter determining the bounding for the fractions of outliers 539 540 and support vectors. If w and ρ solve this problem, then the decision function: 541 542 f(x) = sgn(H(x))543 (31)544

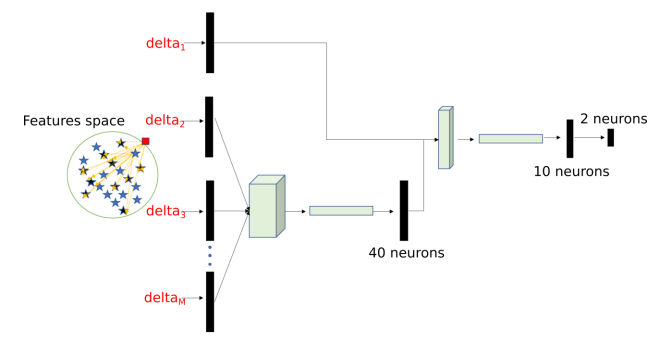
545	will be positive for most of the training observation, while w will be still small. The param	eter
546	influences the trade-off between the reported properties. To solve the quadratic form, we c	an use
547	Lagrangian multipliers, obtaining:	
548		
549	$L(w,\xi,\rho,\alpha,\beta) = \frac{1}{2} \ w\ ^2 + \frac{1}{\nu l} \sum_i \xi_i - \rho - \sum_{i=1}^l \alpha_i (w * \phi(x_i) - \rho + \xi_i - \sum_{i=1}^l \beta_i \xi_i)$	(32)
550	And set the derivatives with respect to w, ξ and ρ and expanding using the kernel expression	on
551	yields:	
552		
553		
554 555	$f(x) = sgn\left(\sum_{i} \alpha_{i} k(x_{i}, x) - \rho\right) (a)$	
	$\alpha_i = \frac{1}{\nu l} - \beta_i \leftrightarrow 0 \le \alpha_i \le \frac{1}{\nu l} \qquad (b)$	(33)
558 559	$\sum_{i=1}^{l} \alpha_i = 1 \tag{c}$	
560		
561	We used a Radial Basis Function kernel (RBF):	
562		
563	$k(x_{i},x) = e^{-\frac{\ x_{i}-x\ ^{2}}{2\sigma^{2}}}$	(34)
564		
565	And then the original quadratic problem is solved substituting Eq. 16 into Eq. 15, yielding 31	:

566
567
568
$$\min_{\alpha} \sum_{i=1}^{l} \alpha_i \alpha_j k(x_i, x_j)$$
 (35)
569
570 under the constraint of Eq. (16b) and (16c).
571
572
573 We finally use the support vectors x_i to recover the parameter ρ needed to compute the
574 hyperplane:
575
576
577
578 $\rho = w \phi(x_i) = \sum_j \alpha_j k(x_j, x_i)$ (36)
579

580 **DEC detectors**

We designed a deep neural network that we named Delta-Enhanced Class (DEC) detector for the purpose of anomaly detection. The DEC detector's architecture is represented in Fig 8, and shows a 2-neurons output, indicating that the sample is a member of the class or is an anomaly (i.e. not a member of the class). For each observation, we train such neural network with the actual features vector and extract randomly select a set of points from the training class in our dataset. For each of these selected points, we define a custom network layer (delta layer) that computes the difference in absolute value (as a vector, feature by feature) between the actual observation and

- 588 the extracted random set. The vector of differences and the actual observations are used as inputs
- 589 to the neural network (Fig 8), which assigns the proper weights to either one during training. The
- set of points to select is a hyperparameter which needs to be tuned. Through testing we determine
- that 25 points is the optimal tradeoff accuracy and computational cost.



592

593 Fig 8. Schematic representation of DEC detector architecture.

594

595

596

597 **Code availability**

- 598 The full source code accompanying this paper has been made available under EPL license at the
- 599 following link: <u>https://github.com/sbianco78/UnsupervisedPlanktonLearning.</u>

601 Supporting information

- 602 S1 Data. The lensless microscope dataset and the dataset extracted from the WHOI used in
- 603 this paper is available at the following link:
- 604 https://ibm.ent.box.com/s/8g2mp5knl2by7cv0ie0fx60mlb3rs6v3

S1 Text. Supplementary Information include: S1. Implemented detector to extract plankton 605 606 images from the acquired videos S2. Evaluation of purity with respect to the number of samples 607 using the lensless microscope dataset S3. Example images from the considered datasets S4. 608 Example images from the considered datasets **S5.** Estimated number of clusters adopting the 609 partition coefficient **S6.** Local Binary Pattern computation. **S7.** Multi-dimensional representation 610 for the Haralick subset of features S8. Multi-dimensional representation for the Hu-moments 611 subset of features **S9.** Multi-dimensional representation for the features extracted from the gray 612 values histogram S10. Multi-dimensional representation for the LBP subset of features S11. 613 Multi-dimensional representation for the Fourier Descriptors subset of features **S12.** Multi-614 dimensional representation for the Zernike moments subset of features **S13.** Histogram reporting 615 the normalized ranking score for the set of designed descriptors **S14.** Schematic work flow 616 describing how an observation is associated to the three possible outpus of the developed system: 617 retraining class, anomaly or belonging to a trained class

618 S1 Fig. Implemented detector to extract plankton images from the acquired videos. The
619 bounding box corresponding to the final detected contour is used to crop the plankton image.

S2 Fig. Evaluation of purity with respect to the number of samples using the lensless
 microscope dataset. The results are very accurate with number of images per sample higher or

equal to 100. Using 50 images results in an overlap between two clusters (corresponding to the
species Paramecium bursaria and Blepharisma americanuum), and in a decrease of the
performances (light gray bar). The corrected purity algorithm introduced in this supplement (see
Customized purity algorithm section), allows for a more accurate result (patterned bar).

S3 Fig. Example images from the considered datasets. a-z13 WHOI dataset (names as they are 626 627 labeled in the dataset) z14-z23 lensless microscope dataset. a Ceratium b Chrysochromulina c 628 Coscinodiscus **d** Dactyliosolen **e** Gyrodinium **f** Strombidium_morphotype1 **g** Dino30 **h** Euglena 629 i Eucampia j Flagellate sp3 k Pyramimonas longicauda l Thalassionema m Delphineis n 630 Pleurosigma o Chaetoceros didymus flagellate p Dictyocha q DactFragCerataul r 631 Emiliania_huxleyi s Corethron t Kiteflagellates u Tintinnid v Dinobryon w Ephemera x 632 Thalassiosira dirty v Skeletonema z Pseudochattonella farcimen z0 Proterythropsis sp z1 633 Heterocapsa_triquetra z2 Rhizosolenia z3 Prorocentrum z4 Pleurosigma z5 Phaeocystis z6 Laboea 634 Strobila **z7** Katodinium or Torodinium **z8** Mesodinium sp **z9** Paralia **z10** Guinardia striata **z11** 635 Asterionellopsis **z12** Amphidinium sp **z13** Pennate morphotype1 **z14** Blaepharisma Americanum 636 z15 Euplotes Eurystomus z16 Spirostomum ambiguum z17 Volvox z18 Arcella Vulgaris z19 637 Actinosphaerium Nucleofilum z20 Dileptus z21 Stentor Coeruleous z22 Paramecium Bursaria z23 638 Didinium nasutum.

639 S4 Fig. Examples of species that are incorrectly assigned to the same cluster by our algorithm
640 because of their morphological similarity in our feature space. Similarity is intended from left
641 to right a Proterythropsis_sp b Heterocapsa_triquetra c Amphidinium_sp d
642 Pseudochattonella_farcimen e Gyrodinium f Prorocentrum

643 S5 Fig. Estimated number of clusters adopting the partition coefficient. a and the XIE-BENI
644 index b as a function of sample size (species). The results are less precise if compared with the

645	partition entropy (see fig 2e in the main text). However, both the algorithms can reconstruct
646	correctly the number of clusters for subset of 3 species and 5 species. The number of clusters on
647	the y axis is the distribution of ten runs on random subsets of all species. For example, for the
648	leftmost box, 3 species have been randomly chosen from the lensless microscope database. This
649	procedure is repeated ten times and the mode is then used as the estimated number of clusters.
650	S6 Fig. Local Binary Pattern computation.
651	S7 Fig. Multi-dimensional representation for the Haralick subset of features. a Andrew's
652	curve. b Parallel coordinates
653	S8 Fig. Parallel coordinate for the Hu-moments subset of features. a Andrew's curve. b
654	Parallel coordinates
655	S9 Fig. Multi-dimensional representation for the features extracted from the gray values
656	histogram. a Andrew's curve. b Parallel coordinates
657	S10 Fig. Multi-dimensional representation for the LBP subset of features. a Andrew's curve.
658	b Parallel coordinates
659	S11 Fig. Multi-dimensional representation for the Fourier Descriptors subset of features. a
660	Andrew's curve. b Parallel coordinates
661	S12 Fig. Multi-dimensional representation for the Zernike moments subset of features. a
662	Andrew's curve. b Parallel coordinates
663	S13 Fig. Histogram reporting the normalized ranking score for the set of designed
664	descriptors.

665	S14 Fig. Schematic work flow describing how an observation is associated to the three

- 666 possible outputs of the developed system: retraining class, anomaly or belonging to a
- 667 trained class
- 668 S1 Table. Computational time on raspberry pi for the analysis of one sample. The standard
- deviation is computed among the objects contained into the 60 frames of the analyzed video.

670

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