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## An introduction to machine learning tools for the analysis of microplastics in complex matrices

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As microplastic (MP) particles continue to spread globally, their pervasive presence is increasingly problematic. Analyzing MPs in matrices as varied as soil, river water, and biosolid fertilizers is critical, as these matrices directly impact the food sources of plants, animals, and humans. Current analytical methods for quantifying and identifying MPs are limited due to labor-intensive extraction processes and the time and effort required for counting and analysis. Recently, Machine Learning (ML) has been introduced to the analysis of MPs in complex matrices, significantly reducing the need for extensive extraction and increasing analysis speeds. This work aims to illuminate various ML techniques for new researchers entering this field. It highlights numerous examples in the application of these models, with a particular focus on spectroscopic techniques such as infrared and Raman spectroscopy; tools which are used to quantify and identify MPs in complex matrices. By demonstrating the effectiveness of these computer-based tools alongside the hands-on techniques currently used in the field, we are confident that these ML methodologies will soon become integral to all aspects of microplastic analysis in the environmental sciences.

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### Environmental significance

The work presented herein presents the use of Machine Learning (ML) tools in the analysis of microplastics (MPs) in terrestrial and aquatic matrices. Microplastics have been known to cause harm to both fauna and flora and, as more and more plastics are produced each day everyday, our ability to count and identify these plastics with speed and precision becomes essential. ML represents an emerging tool for analyzing MPs, but oftentimes those that study computer science and environmental science do not overlap. This review paper intends to introduce ML tools to environmental scientists, allowing them to add rapid counting and identification methods to their tool kit.

## 1 Introduction

For better or worse, plastics have become ubiquitous in our society. From bottles to tires to airbags and pacemakers, the benefits of plastic cannot be overstated. But with their ubiquity comes a number of unintended consequences. Microplastics (MPs) have been found everywhere from the Antarctic ice sheets<sup>1</sup> to bottled water<sup>2</sup> to the inside of our very lungs.<sup>3</sup> From a human perspective, the prevalence of MPs in agricultural soil and riverine systems has raised a number of concerns about how MPs may be affecting agricultural practices and toxicity towards microbes,<sup>4–6</sup> animals,<sup>7–10</sup> plants,<sup>11–14</sup> and humans.<sup>15,16</sup> These MPs have also been found to be co-carriers of known carcinogens, such as PFAS<sup>17,18</sup> or heavy metals.<sup>19,20</sup> Specific emphasis in recent years has been placed on examining the role of biosolids as vectors for MP propagation in terrestrial environments such as agricultural soil,<sup>21,22</sup> where biosolids collected from wastewater treatment plants are often amended to the soil

as a fertilizer. This direct application of MPs has raised a number of concerns about how this affects agriculture and the food we eat.<sup>23</sup> Given that human health may be at risk, the pervasive nature of MPs must be given the attention it deserves.

The various matrices in which MPs are found present unique issues when it comes to identifying and quantifying the MPs found within, as they are incredibly complex. Plastic-free glassware and laboratory equipment are essential for analysis, regardless of the matrix, as they present easy sources of contamination. Less complex matrices, such as ocean and river samples, will undergo numerous filtering and sieving steps, often augmented with manual collection of particulates to cumulate plastic and debris.<sup>24</sup> Sand, soil, and biosolid matrices display even greater complexity as they vary drastically between geographic locations. They contain different mixtures of inorganic (silt, sand, clay) and organic (plant and animal matter) components affecting the density, relative humidity, pH, and myriad other properties of the soil. More recently, the use of synthetic fertilizer and the application of biosolids<sup>25,26</sup> to agricultural fields has increased the amount of plastic that percolates into the soil. Adhikari *et al.*<sup>27</sup> found that soils that had been amended with biosolids over a 23 year period contained a mean

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concentration of 500 plastic particles per kg of dry soil, up to 3 times higher than in non-biosolid amended soils. Corradini *et al.*<sup>28</sup> found that after 10 years of biosolid application in Chilean field soils, the fields they studied had anywhere from 1100 to 3500 plastic particles per kg soil, up to 20x higher than expected. As agricultural practices are actively adding MPs to soil, it becomes necessary to isolate the MPs from the major inorganic and organic soil components in order to identify and quantify the MPs. Techniques ranging from magnetic extraction,<sup>29,30</sup> centrifugation,<sup>31</sup> and solvent extraction separation<sup>32,33</sup> have been employed to carry out the separation, but density separation techniques<sup>34,35</sup> have become the most prevalent due to their ease of use, cost, and effectiveness. In this scenario, the soil is placed into a salt solution of high density, such as NaI (1.8 g cm<sup>-3</sup>). Denser inorganic material will sink, while lighter organic material and polymers will rise, allowing the two layers to be separated. Finally, the organic material is broken down using acids,<sup>36</sup> enzymes,<sup>37,38</sup> or peroxide solutions<sup>39-41</sup> that leave the MPs unscathed, providing us with just the plastics to be characterized. It should be noted that these same techniques can be applied to concentrate the MPs found in marine and ocean samples which can also include soil and organic matter, but on the whole, terrestrial matrices have a much larger component of their collected bulk comprising non-MP elements.

One of the first steps in the process of analyzing MPs is the classification of the particles. What are these MPs made of? What shape are they in? What colour are they? Identification of polymers is often performed using spectroscopic techniques such as IR<sup>42-46</sup> or Raman<sup>47-49</sup> spectroscopy, often combined with an imaging method, such as microscopy. For example, Chouchene *et al.* used microscopy to select MPs based on size, colour, and morphology for future identification by FTIR.<sup>50</sup> Sobhani *et al.* extracted MPs from Australian garden soil, using Raman to identify the plastics.<sup>51</sup> Liu *et al.* extracted soils from Shanghai and not only identified the plastics using  $\mu$ FTIR, but also noted the variation in size and concentration of MPs when looking at top soil *versus* soil collected deeper in the earth (>6 cm).<sup>52</sup> As most polymers have a unique spectroscopic fingerprint, these techniques are highly regarded for their easy identification of MPs, often through comparison with libraries. These processes unfortunately require either time-consuming mapping scans of the specimen, or countless individual scans throughout the sample. The library searches can also have difficulties identifying individual polymers when the samples are a mixture of polymers, or when samples contain contaminants from the soil.

It is also necessary to quantify the number of MPs found in a given sample. As the MPs may eventually reach our bodies through the food we eat, understanding how saturated a water or soil sample is with MPs can give us clues about the risk presented. Unfortunately, the best current technique is to manually count all the microplastics in the sample through a microscope. For example, Fakour *et al.*<sup>53</sup> used a stereomicroscope to count the number of particles found in agricultural fields in Taiwan, as well as examining their colour, shape and morphology. Zhang *et al.*<sup>54</sup> paired a camera with a simple light microscope to perform their counting. Jia *et al.*<sup>55</sup> attempted to

automate the process by using a Laser Directed Infrared (LDI) Spectrometer, but admits that there are limitations on identifying particles below 100  $\mu$ m with great accuracy. As one parses the literature, it becomes clear that new tools are needed to speed up the counting rate compared to a human, as well as improve the collection of data for MPs as they move from the micro to the nanoscale.

As the characterization of MPs has become a time consuming and laborious affair, scientists have once again turned to computing power to find a solution. Giant leaps in computing power and the maturation of the field of Machine Learning (ML) has opened up unforeseen avenues of exploration in a multitude of scientific fields, and MP analysis is no exception. This work hopes to untangle some of the mysteries of ML for those without computer science backgrounds, and show how these *in silico* approaches are being applied in soil samples.

## 2 Machine learning

ML isn't merely a single approach, but refers to a collection of various computational models that can learn from a given data set in order to extrapolate and eventually make predictions about new information that is provided. There are a number of different ML models that can be employed, and often it is best to apply multiple algorithms to the problem and compare the results. In the case of environmental samples, goals tend to focus on classification and quantification, and many different ML algorithms can achieve these goals. In this section, some of the basics of ML will be discussed, as well as several commonly used models and how they work. This is by no means an exhaustive list of techniques, but serves to show the many ways ML can attempt to solve the same problem. As the terminology in this field can often be confusing for new computer scientists, a glossary of terms can be found in Table 1.

### 2.1 Classification *vs.* regression

ML is often employed to solve one of two problems: classification or regression. Regression problems tend to focus on making a prediction based on historical data. For example, if one has a data set that shows the pH of the soil and the total amount of microplastics in that soil, a best fit line can be created to show the relationship between these two features. That best fit line can then be employed to predict the amount of microplastics in a soil sample if the pH is known.

A classification problem is one in which an algorithm is employed to provide labels to a data set based on the training data. For example, if a series of IR spectra are measured for a set of plastics, a classification algorithm can be used to label each of the unknown polymers. When examining MPs in environmental matrices, this type of problem is most common.

### 2.2 Supervised *vs.* unsupervised learning

Another key distinction to make when discussing ML models is that of supervised *vs.* unsupervised learning. In the case of supervised learning, the training data provided to the algorithm has labels that help the model understand what the correct



Table 1 Glossary of Terms

Term	Description
Bias	The error introduced in a machine learning model by simplifying assumptions, leading to underfitting and poor accuracy on training data
Classification	A machine learning task where the goal is to categorize data into predefined classes or labels
Decision tree	A flowchart-like model used for classification and regression that splits data into branches based on feature values, leading to decision nodes or leaf nodes representing outcomes
Hyperplane	A boundary in support vector machines that separates classes by maximizing the margin between them for optimal classification
Latent variable	A hidden factor derived from the data that captures shared variation between predictors and responses, used to reduce dimensionality in partial least squares
Machine learning	A field of artificial intelligence where algorithms learn patterns from data to make predictions or decisions without being explicitly programmed for specific tasks
Regression	A machine learning task focused on predicting continuous numerical values based on input data
Supervised learning	A type of machine learning where the model is trained on labeled data, learning to predict outputs from known inputs
Unsupervised learning	A machine learning approach where the model learns patterns and structure from unlabeled data without specific guidance on outputs
Variance	The model's sensitivity to small changes in the training data, which can cause overfitting and poor generalization to new data

answer should be. For MPs, the data may be labelled as polystyrene (PS) or polypropylene (PP) so that the model knows that any spectra with these specific peak locations belong to one of those categories. In unsupervised learning, the training data is not labelled, so the model will have to make its own connections and identify patterns without help. In the case of MPs, supervised learning is far more common as much of the training data has been analyzed quite thoroughly.

### 2.3 How ML works

When setting up a ML model, a large data set related to the eventual input data is needed for training and testing of the model. For example, if the purpose of the model is to classify the different types of polymers in a soil mixture using Raman spectroscopy, then a large collection of Raman spectra of various polymers will be needed in preparation. A general rule of thumb is that 10 data points are needed for each feature you wish to examine. For example, if one wishes to examine Raman spectra of aged polyethylene (PE), ideally 10 Raman spectra of PE are required. This is by no means a hard and fast rule. Generally, more training data is better than less, but the complexity and application of the ML model will often dictate of how many data points are needed.

From here, the data needs to be processed for it to be in a useable format for ML algorithms. This often involves three steps: baseline correction, smoothing, and normalization. For Raman spectroscopy data in particular, the baseline is often distorted by background fluorescence, causing a great deal of drift along the baseline. Corrections can be performed using polynomial fitting,<sup>55</sup> least squares smoothing,<sup>56</sup> or wavelet transformations,<sup>57</sup> among others. Smoothing of the data is generally needed as Raman spectra can often have a number of

noisy signals that obscure and obfuscate the main spectral peaks. Therefore, smoothing techniques such as Savitzky–Golay filters are often used to smooth out the data.<sup>58</sup> Finally, normalization is applied to the data so that all data is comparable on a 0–1 intensity scale, allowing data from different sources to be compared on an even playing field. The combination of these three processes will generally allow multiple data sources to be used in a machine learning algorithm.

Once the data set is compiled and processed, the data is split up, with about 80% set aside for training and the remaining 20% set aside for testing. The model will then be trained on the first batch to learn trends and connections, and then it will test what it learned on the training set. From there, parameters are shifted in the algorithm in order to reduce bias and variance. Bias refers to the error between the average model prediction and the truth. High bias does not match the training data set well, while low bias will match the training set too closely, making the model unable to analyze new data that differs too greatly from the training set. Variance refers to the ability of a model to adjust to a data set. If a model is overfitting the data (Fig. 1A), that means the model is trying too hard to make the input data look like the training set. If you are underfitting (Fig. 1B), the model is doing a poor job of capturing what the data looks like, including outliers. By performing these training and testing steps, the model can be optimized to find the best fit.

## 3 Commonly used ML models

### 3.1 Partial least squares (PLS)

PLS functions best in situations where each data point has a large number of identifiable features. For example, a Raman



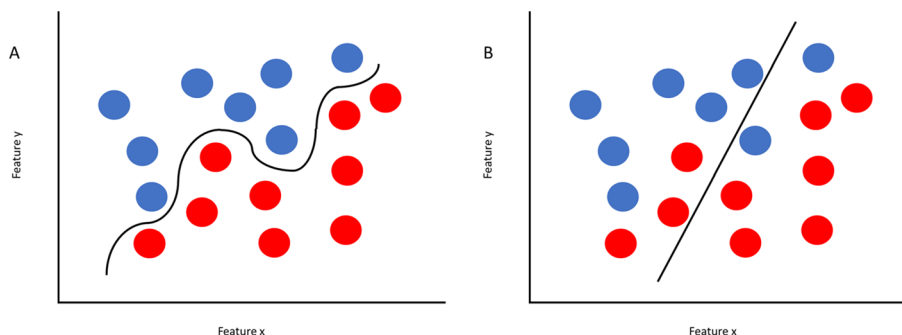


Fig. 1 Plots depicting (A) overfitting and (B) underfitting of a dataset.

spectrum of a mixed soil/microplastic sample (a data point) may have a large number of visible peaks (predictors), which the PLS can use to identify the types of plastics within it. In a classification problem, it may produce a response variable to indicate whether polymers are visible and measurable in the soil. PLS looks at the available information and transforms the data into what are known as latent variables, which are essentially summaries of the data that capture the most important information from the predictors. It will group these latent variables together and reduce the dimensionality of the data, focusing only on the most important and predictive elements. The important elements are selected by determining which combinations of predictors have the highest covariance with the response variables (if the prevalence of one predictor increases, so does the likelihood of identifying a microplastic). Once these elements have been determined, the model can use them to produce the response variable defined by the classifier.<sup>59</sup>

### 3.2 Support vector machine

Support Vector Machines (SVMs) are classification tools that attempt to separate data into classes by examining features of that data. This type of model was developed to focus on binary classifications (yes/no systems), informing the user whether an object is or is not what they are looking for. In the case of MPs,

this can be used to determine if a particle in a microscope image is an MP, or simply leftover debris from the extraction process.

An SVM works by plotting the data set based on two or more features (these make up the axes of your plot). Features in the case of MPs may refer to physical dimensions like size or aspect ratio, or for spectroscopic data sets the features may be spectral peaks known to correlate with a given functional group. The data set is then plotted on this set of feature axes (Fig. 2). The next step is for the algorithm to create a boundary line between through the data set called a hyperplane. As there are a number of hyperplanes that can be drawn through a dataset, SVM maximizes the margin between different classes, making it clear that there is a true distinction between each side of the hyperplane. One advantage SVM has over similar models like PLS is that the hyperplane can be drawn through three-dimensional space using kernel functions, improving the classification by adding extra features. Looking at our example in Fig. 2, the side of the hyperplane that the data is found on will determine if the object is or is not a MP. For many cases, it may be obvious whether the data shows an MP or not. Many data points will not be so clear, and that's why having the algorithm find the correct hyperplane is so important and why having an appropriate training set can greatly improve the results of your analysis. SVMs tend to be used more for classification problems than regression problems.<sup>60</sup>

### 3.3 K-nearest neighbour (KNN)

KNN models also look to classify a data point based on its similarity and proximity to known datasets. For example, a known dataset has been classified into two groups based on a set of features, and the model has placed each data point into one of two separate groups (Fig. 3). As a new data point is introduced into the model, it will be sorted based on the assigned features. But what if it falls between classifications? In the case of K-Nearest Neighbour, its classification will be based upon its proximity to one of the other classifications (generally based on a Euclidean distance function). The distance values will be sorted from shortest distance to longest distance. A  $K$  value is then selected. This refers to the selected population size (starting with the smallest distance), which will be averaged. High  $K$  values lead to underfitting, meaning that the model is too simple to represent the true relationship between the input

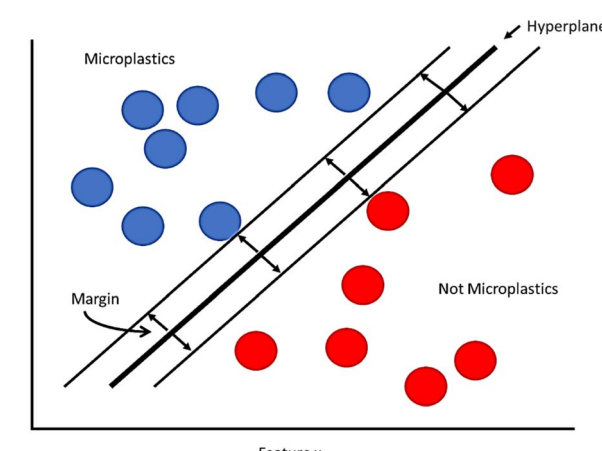


Fig. 2 Visualization of a support vector machine Model.



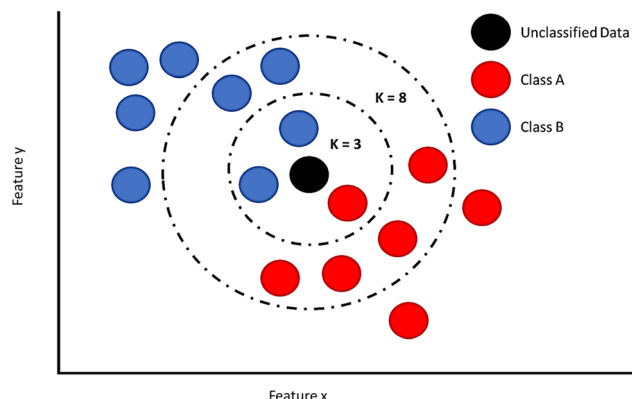


Fig. 3 Visualization of a K-nearest neighbours model.

(new data point) and output (classification). Low  $K$  values can lead to overfitting, as the model becomes too sensitive to variations in the data, making it difficult to properly present the trend of the data, especially when it varies from the training data. Therefore, a Goldilocks zone needs to be found for  $k$ .<sup>61</sup>

#### 3.4 Random forest (RF)

A Random Forest (RF) is a machine learning algorithm that builds on the concept of a decision tree (Fig. 4). A decision tree is a method of determining an output based on a series of binary choices. For example, if one is trying to determine what type of plastic is under investigation, there are a series of choices to be made that can help make that determination? Does the IR spectrum have a C–H stretch at  $2900\text{ cm}^{-1}$ ? If no, then it's probably not PS. Does it have an N–H stretch at  $2930\text{ cm}^{-1}$ ? If yes, then it may be a nylon. In ML, an RF model will take a number of decision trees and run them simultaneously. This is done to minimize overfitting to a training set, or reduce bias that is caused by errors built into each individual tree. For a classification RF model, the classification is based on the majority vote of the individual decision trees. For a regression analysis, the output is the average output of all the individual decision trees. In this way, RF models help to generalize the output and produce more accurate results.<sup>62</sup>

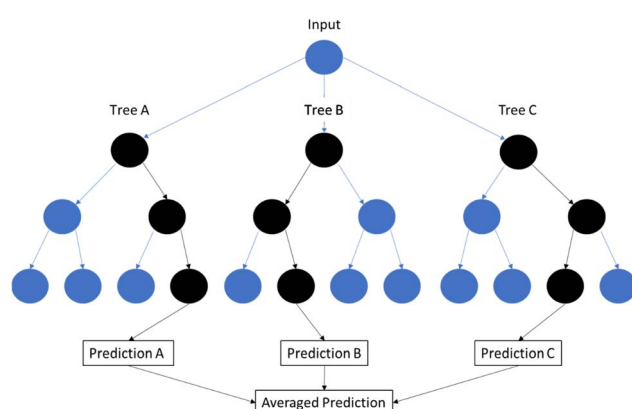


Fig. 4 Visualization of a random forest model.

#### 3.5 Neural networks (NN)

NNs are designed to act like neurons in the brain, taking an input value and weighing it against known information to determine the correct output.<sup>63</sup> NNs use a series of weighted nodes to evaluate the input information. The internal layers, known as hidden layers, are made up of a series of nodes that evaluate one characteristic of the input (Fig. 5). Is this particle round? Is it blue? Does it fluoresce? The model then compares the input data against the training data, and passes the information onto the next layer. Each of these decisions have an associated weight based on how important that characteristic is to determining the output. More weight may be put on a particle having a blue or green colour *versus* whether a particle is round or square. Once the data has passed through a specified set of hidden layers, it reaches the output layer where it makes its final determination based on the appraisal of the data. For a classification model, the output layer may be a single node that makes a binary decision; yes, this is a MP or no, it is not. It's important to note that NNs move in the forward direction (called a feed forward network). Newer models<sup>64</sup> can institute what is known as a recurrent neural network, where the output of some nodes can be used to affect later inputs into the same nodes, acting as an internal learning process. Other neural network models continue to appear as the field matures.

#### 3.6 Choosing the best algorithm

This section has shown that there a vast number of ML algorithms available for researchers to use, and breadth of the field is a feature not a bug. In line with Wolpert's "No Free Lunch Theorem",<sup>65</sup> these is no single machine learning approach that performs best across all possible problems. Therefore, to find the best model, one must simply test several and find the best

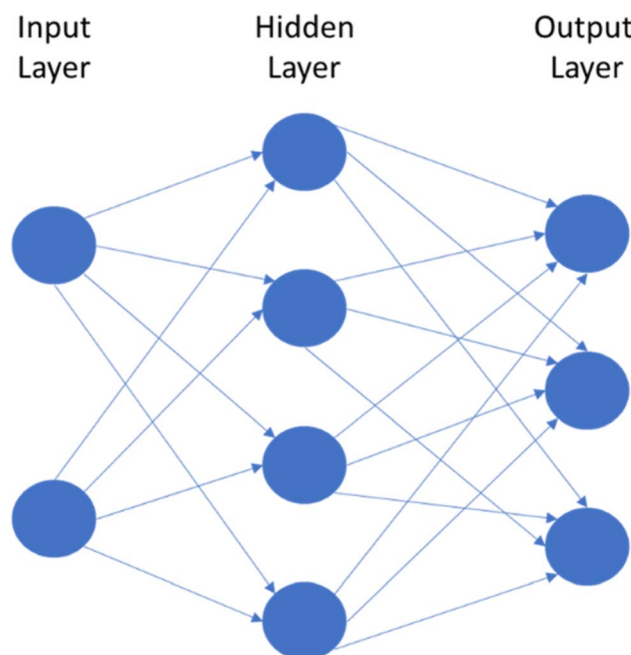


Fig. 5 Visualization of a neural network.



Table 2 Comparison of ML Algorithms (Adapted from Kotsiantis, 2007<sup>66</sup>)

	Decision trees	Neural networks	KNN	SVM
Accuracy in general	**	***	**	****
Speed of learning	***	*	****	*
Speed of classification	****	****	*	**
Tolerance to missing values	***	*	**	****
Tolerance to irrelevant attributes	***	*	**	****
Tolerance to highly interdependent attributes	**	***	*	***
Dealing with discrete/binary/continuous attributes	****	***	***	**
Tolerance to noise	**	**	*	**
Dealing with danger of overfitting	**	*	***	**
Attempts for incremental learning	**	***	***	**
Explanation ability	****	*	**	*
Model parameter handling	***	*	***	*

for yourself. Thankfully, Kotsiantis<sup>66</sup> has put together an excellent review of a number of models that demonstrates their pros and cons (see Table 2). If overall accuracy is a priority, maybe an SVM is the best model, even if it learns at a slow pace. Maybe classification speed matters most, so a NN is the ideal model. Overall, Kotsiantis provides us an excellent starting point, but there is no substitute for hard work.

## 4 Application of ML techniques to MPs in various matrices

### 4.1 Identification of MPs

MP identification typically relies on spectroscopic techniques (such as IR or Raman spectroscopy), which analyze sample composition. IR and Raman spectroscopy offer distinct advantages, providing unique fingerprints of the targeted specimen. When combined with microscopic techniques, they enable individual particle analysis, enhancing characterization capabilities. These techniques are also non-destructive, allowing for further analysis of the particles by other means, such as TGA or Py-GCMS. On the other hand, this analysis is quite time-consuming, and identifying uncommon substances (that may have undergone degradation) can be tricky if they fall outside of known libraries. This is why ML has begun to be paired with these techniques to both speed up and improve the accuracy of identification.

**4.1.1 Raman Spectroscopy.** Raman Spectroscopy has been viewed as a favorable candidate to pair with ML, given its ability to produce distinct spectra for each object under investigation. This allows the ML model to compare against known spectra (used as training data) in order to classify the new unknown spectrum. Lei *et al.*<sup>66</sup> have used Raman in combination with RF and KNN models to identify MP polymers purchased from commercial sources. Luo *et al.*<sup>67</sup> also worked with purchased polymers, but then mixed them into various water samples (lake, river, sea, tap, and ultrafiltered) to add natural impurities. These solutions (mixed at 10, 1, and 0.1 ppm) were then filtered, measured by Raman spectroscopy, and run through a feed forward NN (also known as a Sparse Auto-Encoder or SAE), an SVM, and a back-propagation NN (BPNN) for comparison. The SAE was able to identify the polymers within the water mixtures

with 99% accuracy, besting the SVM (94%) and the BPNN (81%). These studies are important to show ideal case studies, but these don't provide the whole story. Environmental and soil samples are subject to weathering from the sun, wind, and water, as well as from their interactions with humans, causing changes in the Raman spectrum. In order to approach the problem of weathering, Ramanna *et al.*<sup>68</sup> chose to train on Raman datasets of unweathered polymers, such as the Spectral Library of Plastic Particles (or SLoPP)<sup>69</sup> and Mendeley's Raman Database of microplastics weathered under natural environments,<sup>70</sup> and then tested the model on their weathered datasets, such as SLoPP-e. While some pre-processing in the form of normalization and discretization of the data was needed, an RF model was able to correctly determine the identity of the weathered polymers with an accuracy of >90%. Ren *et al.*<sup>71</sup> also chose to test environmentally degraded samples, applying natural and UV light to a series of commercial plastics. It was found that numerous Raman bands saw their intensity change due to oxidative stress on surface functional groups and carbon chains by the UV light. As training data is often based on pristine samples, these light-induced changes in the spectra will differ from the training data. Using a convolutional NN, Luo was still able to identify these weathered samples by their Raman spectra, although a decrease in accuracy from 96% to 95% was noted as a result of these intensity changes.

The above examples were performed using synthetic or pristine polymers, free from environmental matrices. However, when dealing with soil, marine sediments, peat, or wastewater biosolid, classification becomes significantly more challenging. One approach to tackle this is to pair the ML with human expertise. Weber *et al.*<sup>72</sup> collected sediment samples from a series of wastewater treatment plants in Germany, and extracted the MPs *via* density separation in sodium polytungstate before using hydrogen peroxide and sodium hypochlorite to tackle the organic components still dwelling amongst the MPs. They would then generate more than sixty thousand Raman spectra from the samples, which were painstakingly classified by humans based on a set of criteria. This resulted in a high rate of false positives in classification of MPs, though it had a higher overall precision (higher percent of MPs were categorized and classified). Next, the data was input into



a deep-learning NN, where it was found to have higher rate of true positives in classification of MPs than the human only methodology, but had a precision rate lower than the human approach. They then chose to combine the human and machine, first applying the ML algorithm to collect all possible MP signals, and then using the human expert to remove the false positives. Along with a major reduction in the time needed to perform the analysis, this combined method led to a recall of >98% (compared to human only value of roughly 80%) and a precision of around 97% (compared to machine-only 90%). These results show that ML can act as a powerful complementary technique to current classification methods.

What happens when the MPs are still mixed within the matrix? Li *et al.*<sup>73</sup> chose to combine Raman imaging, Raman spectroscopy, and a PLS method in order to examine MPs found in river sand without extraction or digestion of mineral and organic matter. A volume maximizer (AVmax)<sup>74</sup> is used to unmix spectral signals and maximize features of interest before using an unconstrained form of PLS to predict which components in an image are MPs, along with their identity. Li was able to discern PE that was mixed into river sand samples, without any physical preprocessing or filtering of the samples.

#### 4.1.2 Infrared-based spectroscopy and microscopy.

Infrared spectroscopy has also found a home in the study of microplastics for many of the same reasons as Raman. Its quick analysis time and distinct fingerprint-like spectra makes it an ideal technique to combine with ML. The easiest technique is to create a binary classification model; are there MPs present? Yes, or no? ML has been combined with NNs in the past for the classification of pristine polymers,<sup>75</sup> but, as with Raman, mixing with various aqueous matrices and soil samples has become a more pressing focus. Zou *et al.*<sup>76</sup> set out to determine if coloured and colourless plastics could be visually confirmed sitting on a background of compost and quartz sand using a combination of near-IR (NIR) hyperspectral imaging and a PLS ML technique. Hyperspectral images would be taken of the plastics placed on top of the environmental background (but not mixed inside). The PLS model was trained on a series of known plastics on the compost background. For coloured plastics, identification of plastics was performed with high accuracy (>95%). For colourless plastics, the accuracy was lower, but overall identification could be performed with 80–90% accuracy depending on the polymer. Shan *et al.*<sup>77</sup> also used a NIR-based hyperspectral imaging technique to identify microplastics extracted from soil using an SVM, with a particular emphasis on assessing the difficulty of identifying white coloured PE *vs.* black coloured PE. These researchers found that their model performed better when analyzing white coloured PE, suggesting shadows in the imagery caused misidentifications by the SVM model. Paul *et al.*<sup>78</sup> would go a step further, using NIR to examine a series of Australian soil samples spiked with cryo-milled PE, PP, PS, and polyethylene terephthalate (PET) polymers in known quantities. PLS and SVM models were trained on NIR spectra of MP-soil mixtures, MP-free soils, and real-world samples that had some amount of MP in it. They were able to predict whether a sample had MPs or not with relative accuracy, but only above

a concentration threshold (approximately 1%). It also struggled with multiple polymer types present. Chen *et al.*<sup>79</sup> also examined soil samples containing spikes of PE and polyamide (PA) in various concentration, comparing three types of hyperspectral imaging (visible-NIR, InGaAs, and Mercury Cadmium Telluride (MCT) based systems). Using SVM and PLS to help classify the data, Chen determined that InGaAs and MCT performed best in identifying the presence PE and PA in a soil sample, even at concentrations as low as 1.6%. It was postulated that visible-NIR hyperspectral imaging performed worse as its spectral range was more colour sensitive than InGaAs or MCT, suggesting that further research is needed towards the application of these short-wave infrared systems.

The addition of a polymer spike adds certainty to the analysis, but in real world samples the concentration of MPs is expected to be much lower. In the case of marine and ocean samples, large volumes of water need to be processed to analyze measurable quantities of MPs. Tian *et al.*<sup>80</sup> collected large volumes of river water samples from the German Rhine and Meus rivers in order to classify the weathered MPs found in the water. After extensive sieving, chemical treatment, and density filtration, the resulting particulate was examined using LDIR, which uses a Quantum Cascade Laser as its infrared source. Using both a KNN model and Decision Tree model, the researchers were able to classify the unknown polymers with 89.9% accuracy (KNN) and 77.1% accuracy (decision tree). They then went one step further, using a non-supervised model (Density-Based Spatial Clustering of Applications with Noise, or DBSCAN) to help determine the remaining unknowns (which are likely eroded and weathered versions of the known polymers). DBSCAN, as an unsupervised technique, will simply group together data points with similar features. If one happens to know what some of those data points are, and they are clustered with a series of unknowns, it is reasonable to believe some of those unknowns fall into the same category. While this technique cannot truly confirm the identity of the unknown data points, it allows a rough estimate of which of the remaining points may be MPs, and which are simply outliers.

Water samples and soil samples are unlikely to have similar weathering processes, so it's important to examine soil as well. Hufnagl *et al.*<sup>81</sup> chose to examine MPs within soil, wastewater treatment plant outlet, deep sediment, and compost samples, which required the soil to be sieved and separated using mechanical and chemical techniques. Once the MPs were isolated, they used an RF model was used to identify polymers from focal plane array-based micro-Fourier transform infrared (FPA- $\mu$ FTIR) imaging (Fig. 6). This IR technique creates chemical images by recording thousands of IR spectra. The RF model would then predict the identity of the particles by comparing the measured spectra against known reference or training data. This technique was able to distinguish up to 20 different polymers on a 1000 × 1000-pixel image in less than 10 minutes, while also measuring length, width, aspect ratio, area, and orientation. This technique was shown to work in a wide variety of matrices, including sediment, soil, compost, and sewage sludge.



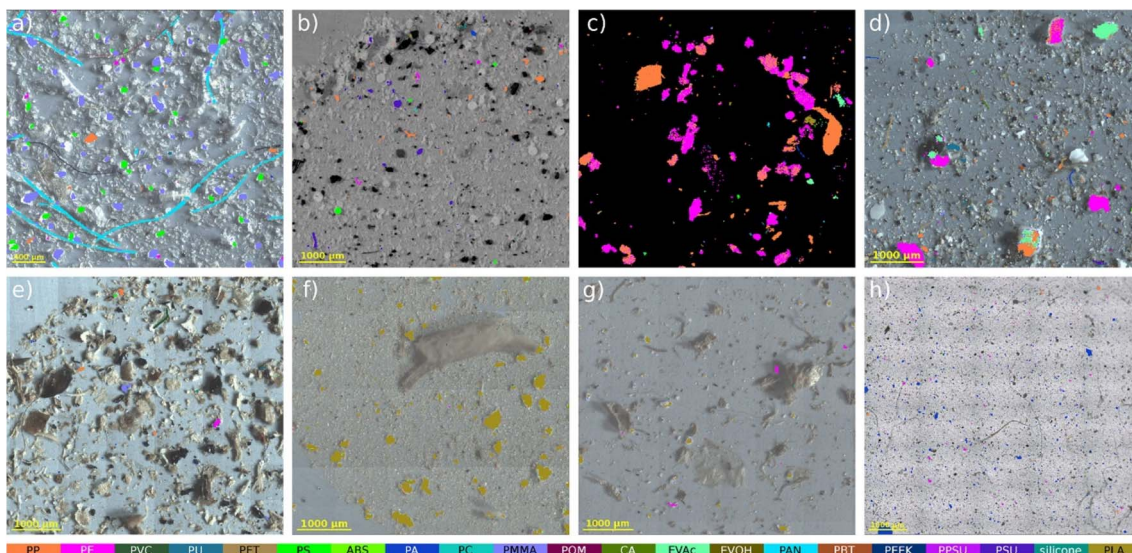


Fig. 6 Application examples for different matrices. (a–c). Plankton samples (d) wastewater treatment plant outlet, (e) deep sediment sample, (f) soil sample, (g) compost sample, and (h) sea salt sample. Adapted with permission under a Creative Commons Attribution 3.0 Unported License from Hufnagl *et al.*<sup>81</sup> Copyright 2022, The Royal Society of Chemistry.

## 4.2 Quantification of MPs

Quantifying the number of MPs in a sample can be just as important as knowing which kind of MPs are in the sample. The ability to determine the number of MPs in soil or water can help monitor the effectiveness of pollution mitigation efforts, and inform researchers whether the situation is improving or deteriorating. By combining ML with spectroscopic techniques, such as IR, Raman, and NIR, ML can be used to determine if a sample reaches a certain threshold of MP concentration, and can be even used to estimate the number of MPs in the given sample.

Quantification of the number of particles (without identification), thus far, has seen most of its progress focused on simpler samples, such as pure microplastics<sup>82</sup> or water samples that have very little in the way of matrix that obscures and obfuscates the MPs.<sup>83</sup> Tan *et al.*<sup>84</sup> chose an in-silica approach to count the number of MPs in a sample using mass measurements obtained from existing datasets, including those taken from beach sands, seawater, and seabirds. They computationally “sieved” the particles into different size ranges and determine the total particle weight in each size range. Using particle density and size measurements for rubber particles, films, beads, fibres, and organic components in the known datasets, they could train an RF model and a linear regression model (known as a Kernel Ridge Regression)<sup>85</sup> to estimate the number of MP particles on the basis of the aggregate particle weight measurements. This allowed the researchers to predict the number of MPs in a single population or in mixed MP samples. Results showed the model would perform better than humans for large and homogeneous mixtures, and that organic material that can be often difficult to remove was not a major source of error, as it represented such a small component of the mixtures overall mass.

Moving on from simulation and testing, more complex samples and matrices would become the focus of study. Using Australian soil samples spiked with PET and low-density PE, Ng *et al.*<sup>86</sup> combined NIR spectroscopy with a NN to classify samples based on the percentage of microplastics found within the sample. They were able to sort soil samples into batches that were less than 1% MPs, between 1% and 3% MPs, and above 3% MPs based on the NIR images with relative accuracy, although it had trouble separating situations of no MP concentration and low MPs concentration. Wu *et al.*<sup>87</sup> also went the spiked-polymer route, adding PE, PS, PP, and PVC to waste incineration ash. The samples were examined with NIR, and care was taken to exclude spectral information from the ash using background subtractions from pure ash samples. The application of an SVM model was able to predict the quantity of plastics within the ash with a greater than 89% accuracy for each of the polymers. Each of these studies show that spectroscopic techniques combined with ML are able to parse through matrices in order to complete quantification experiments, particularly when the concentration is high, as one would expect from a spiked sample.

Spiked samples have a known number of MPs to measure, but this is not the case with environmental samples. Lorenzo-Navarro *et al.*<sup>88</sup> collected beach sands from the Canary Islands archipelago. Even after pre-treatment *via* mechanical sieving and density-based separation techniques, they were unable to completely separate polymer pellets and fibres from tar and organic particles. In order to properly count the MPs, they had to first introduce a classifier that would separate the pellets and the fibres from the tar and organic material based off microscope imagery. A series of ML models were then employed to contrast and compare, including a KNN, an RF, and an SVM. Fibres and pellets proved easy to classify based on geometric properties (fibres have high aspect ratios, pellets are spherical), while the remaining components would require more

properties (colour, texture, *etc.*) to be further classified. Once the models could identify the particles by class, it could then carry out the job of counting. SVM was found to be the best of the techniques, with a recall rate of 88%.

While soil itself is a complex matrix, this also means that it has more features that can be analyzed and measured. Tran *et al.*<sup>89</sup> used this to their advantage, as they went about predicting MP quantities in peatland sediments based on soil pH, salinity, and composition, amongst others. Using peatland soil from the three industrial regions around the Mekong Delta of Vietnam, they measured 7 different physicochemical properties for 300 samples, and, after performing sieving and density filtration, counted the number of MPs found in each sample. This information was used as training and testing data for RF and SVM models (although the large differences in the region made mixing the data from different areas too monumental of a task). Their results showed that both pH and total organic carbon (TOC) had a positive correlation with the number of MPs in a sample from a given area. Salinity and electrical conductivity were found to have a negative correlation. Overall, the correlations were not considered strong enough to stand on their own (the models couldn't predict samples with high numbers of MPs due to the weak correlations), but it has merit as a complimentary tool to the many spectroscopic and microscopic tools seen thus far.

### 4.3 Methodological comparison

The studies discussed above show that ML techniques can be applied to MPs derived from myriad environmental sources, whether provided as pristine polymer spikes or weathered and eroded samples found *in situ*. Each of these studies remains reliant on the ability to extract the MPs from the matrix diligently enough for spectroscopic and microscopic techniques to be applied. New techniques, such as those demonstrated by Ng *et al.*<sup>86</sup> and Tran *et al.*,<sup>89</sup> allow for analysis to occur with very little preprocessing of the matrix, or by using the matrix to improve the analysis. By focusing on these kinds of techniques, the difficult extraction step can be minimized, allowing for more reliable and reproducible data between groups.

This work has previously discussed the important decision of which ML model to select for the task and a quick perusal of the above studies show how Wolpert's "No Free Lunch Theorem" rings true. Table 3 shows a comparison of all the studies in this work, with a particular emphasis on the ML models used in each study and their final accuracy determination. Almost all studies used multiple ML models and, while each study had one model that worked best, no particular model stands head and shoulders above the rest across the board. When it comes to applying ML tools in this field, it is simply a necessity to roll up one's sleeves and try several algorithms to find the best one. It should be noted that Weber

Table 3 Comparison of ML studies

Study	Analysis technique	Matrix	Goal	ML technique	Accuracy of technique
Lei <i>et al.</i> <sup>66</sup>	Raman	Commercial plastics	Classification	RF KNN	>95% >95%
Luo <i>et al.</i> <sup>67</sup>	Raman	Water samples	Classification	NN SVM BPNN	99% 94% 81%
Ramanna <i>et al.</i> <sup>68</sup>	Raman	Weathered plastics	Classification	RF NN SVM DT KNN	93.81% 71.13% 73.19% 69.07% 73.19%
Ren <i>et al.</i> <sup>71</sup>	Raman	Weathered plastics	Classification	CNN	95%
Weber <i>et al.</i> <sup>72</sup>	Raman	Wastewater sediments	Classification	NN/human	98%
Li <i>et al.</i> <sup>73</sup>	Raman	River sand	Classification	PLS	>94%
Zou <i>et al.</i> <sup>76</sup>	NIR	Sand	Classification	PLS	>95% (coloured) >80% (colourless)
Shan <i>et al.</i> <sup>77</sup>	NIR	Soil	Classification	SVM	76% (black) 77% (white)
Paul <i>et al.</i> <sup>78</sup>	NIR	Soil	Classification	SVM PLS	92% 83%
Chen <i>et al.</i> <sup>79</sup>	SWIR	Soil	Classification	PLS	>95% (InGaAs) >92% (MCT) >95% (InGaAs) >92% (MCT)
Tian <i>et al.</i> <sup>80</sup>	LDIR	River water	Classification	KNN DT	89.9% 77.1%
Hufnagl <i>et al.</i> <sup>81</sup>	$\mu$ FTIR	Soil	Classification	RF	>90%
Ng <i>et al.</i> <sup>86</sup>	Vis-NIR	Soil	Quantification	CNN	78.5%
Wu <i>et al.</i> <sup>87</sup>	NIR	Waste incineration ash	Quantification	SVM	89%
Lorenzo-Navarro <i>et al.</i> <sup>88</sup>	Microscopy	Beach sand	Quantification	KNN RF SVM	72.1% 81.9% 88%



et al.'s<sup>72</sup> combination of a neural network with a human "expert" led to not only high classification accuracy, but also minimized false positive and false negatives, suggesting that removing the human element from machine learning remains unwise.

## 5 Beyond spectroscopy

While this work has placed an emphasis on the synthesis of spectroscopic techniques and ML, it is important to note that a number of other techniques have been used to study MPs, particularly thermometric techniques such as Thermogravimetric Analysis (TGA)<sup>90,91</sup> and Pyrolysis Gas Chromatography Mass Spectrometry (Py-GCMS).<sup>92-94</sup> There has recently been a push to utilize ML alongside these techniques. Chowdhury *et al.* used a NN alongside TGA to predict the heat deterioration of PET particles.<sup>95</sup> Christian *et al.* tested a number of ML algorithms to examine PET degradation amongst modified and aged PET particles.<sup>96</sup> Zhang *et al.* tested RF and SVM models to predict the presence of PE and PP based on decomposition data produced by Py-GCMS.<sup>97</sup> Lastly, Forbes *et al.* was able to characterize MPs as pristine, weathered, or in mixtures by combining Py-GCMS with a series of ML models.<sup>98</sup> Each of these studies show how ML can work in conjunction with thermal methods to analyze MPs. What each of these studies lack is the application of these techniques to MPs in environmental samples. Particularly with soil, the field remains very much in its infancy, with direct application of ML on MPs in soil samples yet to be studied at this time.<sup>99</sup>

## 6 Limitations and conclusions

The use of ML to study soil samples has shown amazing promise in analyzing MPs in complex matrices, but limitations still remain. All ML models require large sets of data for training and validation, which can be difficult to obtain. Many of the above examples were required to pull data from multiple databases (often derived from different sources and methods), or even produce synthetic data to be trained upon. These datasets are also not easily accessible. Many datasets may be found behind paywalls or held back for proprietary reasons, making it difficult to get a strong dataset to train on. The field is desperately calling out for more easily accessible datasets, and, even more ideally, a single open access database where it can all be archived and made available for future researchers.

Another major limitation is the complexity of the matrices themselves. Extracting MPs from different matrices involves multiple steps to remove the mineral and organic content, and often doesn't see complete separation or full recovery of the MPs. Remove too much of the soil, and one doesn't get an accurate measurement of MPs. Remove too little, and the MPs get buried in the noise. Add in the fact that soil or water varies heavily between regions in composition and usage, and it becomes difficult to apply a one-sized fits all approach to analysis. This may be solved through the creation of new standards and reference materials created from individual soil/MP mixtures, which may speed up and improve the accuracy of

the training and testing procedure. There remains an opening for metrologists to add their own expertise to the field.

A final limitation of ML comes from the reproducibility of ML models, as sample preparation (choosing one chemical degradant over another), the type of spectrometer (such as variations in laser power and intensity), and even the data acquisition method (selection of range of interest, duration of laser application) can lead to variations in the data that is fed into the models. In order to combat this, full transparency of data and methodology will be key to demonstrating why and how models may vary. Both raw and processed data should be made available publicly, as well as any assumptions regarding why each data point was or was not included in the training/testing dataset. An advantage of making all the data available is that large databases can be built to create training data, improving future models. The ML code itself needs to be made readily available, so that the code can be evaluated and tested for errors and inaccuracies and even improved upon for future iterations. If the data and code are available for all to see, then reproducibility can be much improved, or, in the cases where there is poor reproducibility, the source of the issue can be laid bare for all to see.

Despite these limitations, the future remains open to optimism. As more data is produced, the available pool of training data increases, improving the reliability of the models. As new techniques and models come online, research can dig deeper into their data and finally gain a full picture of the MPs scattered throughout our world.

## Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

## Conflicts of interest

The author declares that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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