

Machine Learning: New Ideas and Tools in Environmental Science and Engineering

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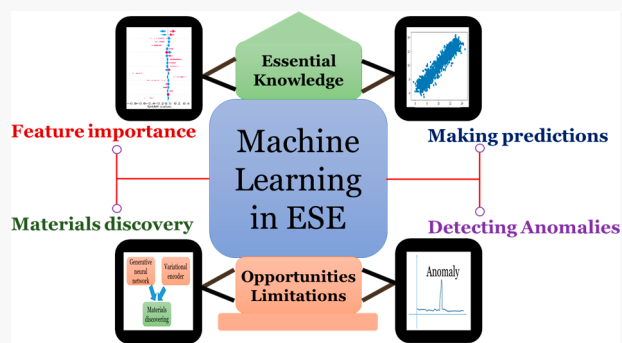
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ABSTRACT: The rapid increase in both the quantity and complexity of data that are being generated daily in the field of environmental science and engineering (ESE) demands accompanied advancement in data analytics. Advanced data analysis approaches, such as machine learning (ML), have become indispensable tools for revealing hidden patterns or deducing correlations for which conventional analytical methods face limitations or challenges. However, ML concepts and practices have not been widely utilized by researchers in ESE. This feature explores the potential of ML to revolutionize data analysis and modeling in the ESE field, and covers the essential knowledge needed for such applications. First, we use five examples to illustrate how ML addresses complex ESE problems. We then summarize four major types of applications of ML in ESE: making predictions; extracting feature importance; detecting anomalies; and discovering new materials or chemicals. Next, we introduce the essential knowledge required and current shortcomings in ML applications in ESE, with a focus on three important but often overlooked components when applying ML: correct model development, proper model interpretation, and sound applicability analysis. Finally, we discuss challenges and future opportunities in the application of ML tools in ESE to highlight the potential of ML in this field.

KEYWORDS: applicability domain, artificial intelligence, best practices, feature importance, machine learning modeling, model applications, model interpretation, predictive modeling



1. INTRODUCTION

The rapid advancement in environmental analytical tools and monitoring technologies has led to a corresponding explosive expansion in both the quantity and complexity in data generation, which demands more advanced and powerful computational and data analytical approaches beyond traditional statistical tools. Data analytical approaches that have less dependence on prior knowledge, such as machine learning (ML), have shown promise in solving complex data patterns or formats because of their powerful fitting abilities. As a result, the past decade has witnessed a rapid growth of ML, especially deep learning, in a variety of applications, such as image classification and machine translation. These tools are revolutionizing many scientific fields, from chemistry,¹ material sciences,² and biomedicine,³ to quantum physics.⁴ Researchers in the broad field of environmental science and engineering (ESE) have also adopted ML enthusiastically, as demonstrated by the explosive growth in the number of publications (5855 between 1990 and 2020) on the applications of ML in ESE (Figure 1). These applications cover broad areas, including assessing environmental risks, evaluating the health of water

and wastewater infrastructure, optimizing treatment technologies, identifying and characterizing pollution sources, and performing life cycle analysis, among others.

The definition of ML is that “ML algorithms build a model based on sample data, known as ‘training data’, to make predictions or decisions without being explicitly programmed to do so”.⁵ Example ML algorithms include random forest, support vector machines, and artificial neural networks.^{6–8} Deep learning is one class of ML, in which “deep” refers to the multilayered neural network structures,⁹ such as recurrent neural networks and convolution neural networks.^{10–12} Every ML algorithm can be decomposed into three key components: the structure of the algorithm, such as random forest and deep



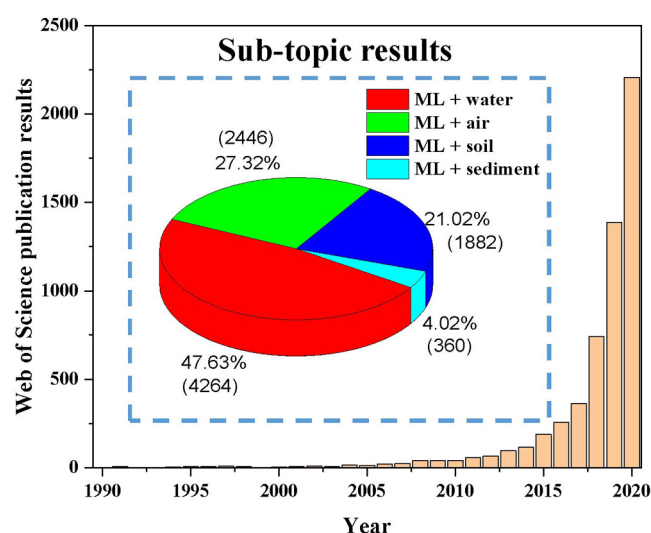


Figure 1. Number of publications in ML applications in ESE based on the Web of Science (access date 1/28/2021) with the keyword “machine learning” combined with the categories of environmental science, water resources, public environmental occupational health, environmental engineering, and environmental studies. The inset shows the subtopic results from 1990 to 2020 with the keywords specified.

neural networks; the goal to achieve, such as prediction accuracy and squared error; and the training method to achieve the goal, such as stochastic gradient descent.¹³ One major advantage of ML algorithms is that they can easily identify trends or patterns in data without human intervention. Their predictive performance can be continually improved with more available data. Multidimensional and multivariety data can also be handled by ML even in dynamic or uncertain environments.^{14–17}

Many complex ESE problems can be addressed by ML. Five specific examples are explained in section 2, and a comprehensive summary is included in section 3. A challenge, and opportunity, is that even as ML algorithms are increasingly applied to ESE problems, these algorithms are being continuously improved and new ones are developed. ESE practitioners passively benefit from increasingly advanced ML methods, as they allow more challenging problems to be addressed. ESE practitioners can also make an active contribution to the development of more advanced ML methods by identifying and communicating the encountered limitations to ML researchers.

Despite the initial success of ML in ESE, many concerns remain. First, ESE researchers may be eager to use ML in their research but lack the knowledge of how to properly employ it, which may lead to incorrect applications of ML to their data sets. Second, increased data volume and complexity has enabled the use of more advanced ML algorithms, such as deep neural networks, capable of capturing sophisticated nonlinear relationships. However, these types of models are often of a “black box” nature; therefore, model interpretation is vital for investigating whether ML model predictions are consistent with the fundamental principles of the domain science. Although there is a growing field of model interpretability,¹⁸ such interpretation is still commonly neglected in ESE.^{19–22} Alternatively, emerging approaches to building neural networks that enforce physical symmetries or directly implement differential equations that parallel chemical kinetic differential

equations,^{23,24} have not yet been adopted in ESE. Third, applicability domain (AD) analysis of ML models is yet to be practiced by researchers in ESE after model development,^{19–22} other than development of quantitative structure–activity relationships (QSARs).²⁵ To introduce ML to the general ESE researcher’s toolbox, this feature aims to discuss the current status, essential knowledge, shortcomings, challenges, and future opportunities of ML in ESE to highlight the potential of ML in the ESE field.

2. HOW DOES ML ADDRESS ESE PROBLEMS?

ESE applications often utilize supervised ML approaches, which are essentially interpolation—given a big enough library of inputs (independent variables or “features”) matched with outputs (dependent variables or “outcomes”), these algorithms after training can take a new input and predict its corresponding output. Where this differs from traditional statistical tools is the ability to treat a large number of features that have weak or nonlinear relationships with the outcomes. ML can be more effective than traditional statistical tools in handling various data formats, such as text, images, and graphs, where the important information is not contained in a single input variable, nor are the important variables known ahead of time, but instead where some previously unknown combination of features are needed to determine the outcome. These unique properties of ML are especially suitable for solving complex environmental problems with rich sets of input features.²⁶ We, here, take five specific examples from different environmental fields to illustrate how ML can address complex environmental problems. We will have a comprehensive discussion of current ML applications in ESE in section 3.

2.1. Prediction of Particulate Matter (PM_{2.5}). When predicting daily to yearly variations in PM_{2.5} for a region, the major drivers include meteorological conditions, such as air temperature, dew point temperature, visibility, pressure, potential evaporation, downward longwave radiation flux, downward shortwave radiation flux, relative humidity, and wind; and land-use variables, such as limited access highway, highway, local roads, and forest cover.²⁷ In addition, there are regional differences associated with meteorological conditions and daily variations in PM_{2.5} levels. These factors interact with each other to form complex relationships, which are challenging to process using traditional statistical tools but can be effectively handled by ML when sufficient training data are given to learn their relationships. For example, whereas multiple linear regression of PM_{2.5} prediction from these factors only achieves an R^2 of 0.60, an artificial neural network using the same features and data can achieve a higher R^2 of 0.74.²⁸

2.2. Prediction of Water Resource Availability. Climate change—increasing average temperature, uneven shifting of precipitation patterns, and frequent occurrence of extreme climate events like floods and droughts—can substantially impact the predictability and availability of water resources across regions. To better manage and sustain future water resources, it is essential to develop decision support tools that can handle these variations and uncertainties, which arise due to either interactions between natural and human systems or from the variability of climate. Addressing such challenges requires insights into past and future patterns, yet this insight can be difficult to develop based on traditional statistical approaches. In contrast, ML allows effective analysis and

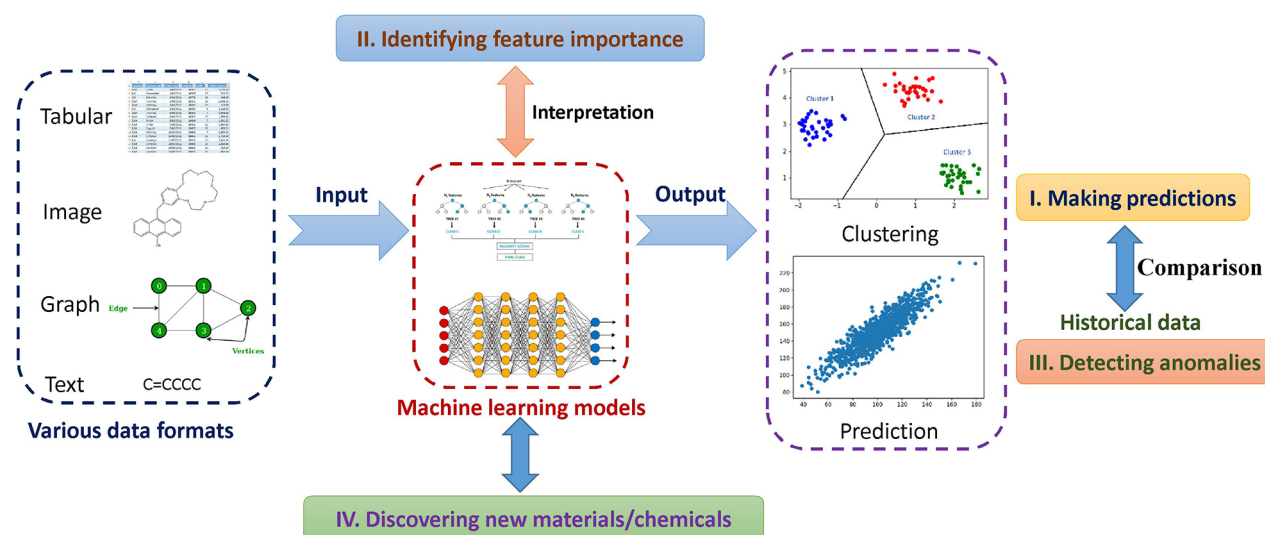


Figure 2. Four common applications of ML in ESE. Different data formats can be used for inputs in ML to develop models for (I) prediction purposes. By interpreting the ML models, the importance of features (II) can be obtained. By comparing the prediction with the historical data, (III) anomaly detection can be achieved. ML can also be used to (IV) discover new materials.

prediction of future water availability through processing data related to climate change and water system interactions.²⁹

2.3. Data Collection and Interpretation Across Water Facilities. An enormous amount and wide spectrum of data in water utilities have been collected from in-place supervisory control and data acquisition systems, including flow rate, temperature, dissolved oxygen concentration, turbidity, and chlorine content, to name a few, as well as data from ex situ laboratory information management systems and computerized maintenance management systems. Although these rich and complex data sets have been around for years, the gathered data sets from treatment facilities are often incomplete and not connected to each other. The current approach to data collection, interpretation, and utilization is not suitable for rapid identification of malfunction, swift control and adjustment under transient fluctuations, or efficient decision-making regarding facility operations. This is because these traditional models are mainly based on statistics,^{30–34} which are valid only for a limited operating range and cannot capture the time-varying or nonlinear behavior of dynamic systems. In contrast, ML models can adapt to fast-changing situations and, because they do not rely on predetermined rules, they can use varied, dynamic data to update themselves for better predictions.^{35,36}

2.4. Modeling of Biochemical Wastewater Treatment Systems. Most wastewater treatment plants (WWTPs) use activated sludge and other processes to remove contaminants, primarily including organic carbon, nitrogen, and phosphorus. There can be tens of thousands of different microbial species present in each system. The abundance of different organisms, as well as the time-varying and highly nonlinear characteristics of the WWTP process, makes the relationship with process performance a difficult one to investigate. Deterministic models based on the fundamental biokinetics for activated sludge processes are not particularly practical due to the complex nature of biological reactions, highly multivariable aspects of treatment plants, the expertise required to calibrate activated sludge models, and system-specific calibration.³⁷ The influent wastewater flow and composition also vary over time and follow dynamic patterns. However, ML techniques could predict sludge bulking in wastewater treatment plants with

improved accuracies without the burden of calibration.³⁸ ML carries the potential to characterize the relationships between microbial communities and parameters of the process based on the fusion of sequencing data and model outputs. For example, Zhu et al. compared the abilities of nine single or hybrid models to predict the next day's influent wastewater flow rate.³⁹ Compared to principal component regression (adjusted $R^2 \approx 0.66$; mean absolute percentage error $\approx 18.2\%$) and partial least-squares (0.70; 17.5%), the hybrid model using an artificial neural network (0.83; 13.3%) achieved the best results.

2.5. Prediction and Identification of Endocrine-Disrupting Chemicals (EDCs). Currently, there are over one hundred thousand chemicals available on the market, many of which lack toxicity data, including data on their endocrine disrupting activity.⁴⁰ In screening and predicting potential EDCs, many traditional tools, such as read-cross, are based on the concept that structurally similar compounds carry similar toxicity and these tools have been successfully applied.^{41,42} However, focusing on a limited set of chemicals or only structurally similar compounds makes it difficult to apply these traditional tools to tens of thousands of untested chemicals. Additionally, because of the complex molecular mechanisms behind endocrine disruption, many traditional tools that rely solely on structural information usually experience “activity cliffs”, that is, a group of seemingly structurally similar chemicals have different endocrine disrupting activities.^{43,44} These problems stress the unsuitability of traditional data analysis methods, and researchers have tried many approaches to combine ML tools with big data (including various biological, physicochemical information, etc.) to solve them. For instance, a computational method was developed to automatically extract useful biologically active characteristic fragments of the estrogen receptor alpha (ER α) and the androgen receptor (AR) based on more than 4000 chemicals and to built chemical fragment–in vitro relationships (118 for ER α and 99 for AR) to illustrate new molecular mechanisms of EDCs.⁴⁵ Pathway activity prediction, such as the AR-mediated pathway, used 1746 compounds from 11 high-throughput in vitro screening assays, considered multiple

points (receptor binding, coregulator recruitment, gene transcription, and protein production) and multiple cell types, and showed extremely high prediction accuracy.^{46,47} Furthermore, combined with 16 different ML methods, pathway activity information was used to build 91 predictive models for EDCs' activity on the AR. The final consensus model showed a highly predictive performance on 87 914 chemicals.⁴⁷ Similar research also can be found in an ER study that used 19 different ML methods and 32 464 chemical data sets to build 48 models for ER activity prediction.⁴⁸

3. CURRENT STATUS OF ML APPLICATIONS IN ESE

Figure 2 shows four common applications of ML in ESE: making predictions, identifying feature importance, detecting anomalies, and discovering new materials or chemicals. Table 1 provides more detailed examples for each of the applications.

3.1. Making Predictions. Overall, making predictions is one of the most popular applications of ML in ESE. This application can be implemented through either regression or classification modeling. The key assumption is that the distribution of training examples provided to the algorithm is representative of the examples that the model will be asked to predict. In this sense, it does not matter if the problem is time-dependent or -independent, as long as "past" observations are predictive of "future" queries. In addition, if the predictions are related to the location or sites, they can be seen as geographic space-based predictions, such as predicting arsenic concentrations in groundwater at different sites.⁴⁹ Most prediction applications are in the realm of supervised learning, in which the sample outputs are labeled. For example, ML methods have been extensively employed to predict changes of various wastewater variables, such as nitrogen, phosphorus, solids, chemical oxygen demand (COD), biochemical oxygen demand (BOD), and future flow rate;^{39,50–53} or atmospheric pollutants, such as PM_{2.5} and carbonaceous aerosols.^{54–56}

In addition to supervised learning, unsupervised ML approaches have also emerged recently. In unsupervised methods, the computer algorithm works on its own to find patterns in the data, that is, in an "unsupervised" fashion without intervention from the user. Most applications of unsupervised ML have been used to automatically categorize data into separate groups or "clusters" that have similar characteristics. In the context of environmental studies, there have been recent unsupervised ML applications, such as *t*-distributed stochastic neighbor embedding (*t*-SNE) or *k*-means clustering, on categorizing the carbon–fluorine bond dissociation energies of per- and polyfluoroalkyl substances (PFAS) to understand bond dissociation energies.^{57,58} These algorithms allow the visualization of high-dimensional data as two-dimensional "clusters" where data points grouped within a cluster share similar characteristics with each other. It is important to reiterate that these clusters were automatically chosen by these unsupervised ML algorithms, without human intervention. As such, these results demonstrate that these algorithms can be useful tools for automatically classifying and rationalizing chemical trends in environmental contaminants that would otherwise have been difficult to detect manually.

3.2. Identifying Feature Importance. This refers to the techniques of assigning scores to input features or independent predictors to evaluate their relative importance to the outcome. This is also referred to as "model interpretability," "feature extraction," or the study of "latent spaces" in the ML literature.¹⁸ For example, air pollution, as a complex global

environmental issue, is affected by many factors. ML techniques such as support vector machines, neural networks, and feature extraction methods can be especially useful to determine the most significant factors for the modeling of particulate matter.²⁷ The prerequisite for model interpretation is that the obtained ML model has satisfactory predictive performance or ML has already correctly "learned" the underlying relationship between the features and the outcome. Thus, interpretation reveals the implicit knowledge "learned" by the ML model and indicates whether the model is based on a correct "understanding" of the underlying mechanisms, which is important for validating the model. This process may also yield new knowledge. For example, Mori et al. interpreted ML models to provide evidence-based information on how stressors and ecologically important environmental factors interact and drive ecological processes and microbial biomass.⁹⁸ Such information cannot be obtained without performing model interpretation. Similarly, there are several attempts to interpret ML models to correlate factors with the chemical activity of EDCs, including unveiling features that make EDCs chemically active, determining the type of activity on the ER α or the AR, and how these features exert their functions.⁴⁵ Using ML techniques, researchers also found that the octanol–water partitioning coefficient ($\log K_{ow}$) plays a dominant role in regulating plant uptake of organic contaminants, while their molecular weight plays a secondary role.⁹⁹ In recent years, data-driven analytics such as ML have become key tools for discovery in public health and ESE research to find hidden patterns and causal relationships and to identify key features, that is, chemical exposure or other social economic parameters, that are linked with health outcomes.^{84,100}

3.3. Anomaly Detection. This refers to the identification of historical or current abnormal events to avoid irregularities or unreliable operations; this is widely used to prevent credit card fraud.¹⁰¹ The basic principle is that new observations are compared to the learned distribution of (mostly)-normal historical data to determine statistically improbable deviations. For instance, anomaly detection has been used to identify burst locations of pipes and detect contamination events in water distribution networks,^{90–92} with the former contributing to less water loss and the latter important for reducing public health risks. ML can also be used to make predictions for a future event by comparing those predictions with current data to identify potential outliers and then calculate the probability of future contamination events.

3.4. Discovering Materials and Chemicals. Discovering materials and chemicals based on ML is another rapidly growing application in ESE, such as designing environmentally friendly adsorbents and catalysts. Take biopolymers as an example, researchers are developing biodegradable polymer materials that can functionally replace plastics made from fossil feedstocks, thus reducing plastic pollution in the environment. Toward this goal, they apply an innovative ML approach to bring together two adaptive codesign loops: (i) a chemical loop, where the structure and function of possible chemical combinations of the polymer backbone and side chains are explored for their predicted properties^{96,102} and (ii) a synthetic biology adaptive design loop, in which the biosynthetic pathways involved in polymer production and the roles of their component genes and proteins, are investigated and targeted for improvement by genetic engineering methods. Another example is to develop novel adsorbents, where two

Table 1. Example Applications of ML in ESE

application	media	example	input	output	performance	refs
making predictions	air	PM _{2.5} formation	meteorological conditions, land-use variables, etc.	PM _{2.5} concentration	better than multiple linear models	27, 28, 59
	water	prediction of carbonaceous aerosols	air quality variables and meteorological conditions	organic carbon, elemental carbon concentrations	better than generalized additive models	56
	water	QSARs for predicting the reactivity of contaminants	molecular fingerprints or molecular descriptors	reaction rate constants	better than multiple linear regression	60–63
		single- and multisolute adsorption modeling	descriptors for chemicals and adsorbents	adsorbed amounts or coefficients	outperformed multilinear regression models	64, 65
		deep learning and systematic understanding of urban water management infrastructure	water quantity and quality	energy consumption, chemical use, and environmental impacts	understood systems complexity, identified unnecessary energy/chemical inputs, and reduced unwanted effects	66
		predicting water quality parameters for wastewater effluent	various operating parameters such as mixed liquor suspended solids	water quality characteristics	outperformed activated sludge models	67–69
		intelligent monitoring of membrane fouling in water and wastewater treatment	operating parameters and influent water characteristics	membrane permeability	introduced an integrated approach to control membrane fouling	70, 71
	soil/sediment	monitoring of water or wastewater quality and risk management	water/wastewater quality variables	monitoring variables	superior over multiple linear regression model	51
		prediction and early warning of combined sewer and facility overflow	rainfall data in conjunction with meteorological conditions	future combined sewage flow rate	better than classical, statistical methods	39
		predicting arsenic concentrations in groundwater	various climate and soil parameters, geology, and topography	probability of arsenic in groundwater exceeding the World Health Organization (WHO) guideline	better than statistical models	49
		prediction of plant uptake efficiency	physicochemical properties of chemicals	concentration factors	improved accuracy compared to mechanistic models	72
		neural network predictive models for cadmium and cerium uptake by plants	plant properties	cadmium and cerium concentrations	showed high accuracy and low error	73
toxicity		QSARs for predicting toxic end points of high concern, e.g., bioconcentration factor, skin sensitization, carcinogenicity, mutagenicity, developmental toxicity	"SMILES" of chemicals, molecular descriptors, and physicochemical properties, relevant experimental results	qualitative and quantitative activity prediction results	improved the prediction accuracy and innovatively predicted some in vivo toxicity end points with complex mechanisms that traditional tools could not predict	74–77
		identifying potential endocrine-disrupting chemicals (EDCs)	1D, 2D, and 3D molecular descriptors, physicochemical properties, in vitro and in vivo assays	EDC activity	active pollutants can be prioritized for regulators	47, 48, 78–81
identifying feature importance	air	understanding the relationship between pollutant concentrations and influencing factors	meteorological and land-use variables	PM _{2.5}	identified the most important variables to PM _{2.5} prediction	27
	water	understanding the relationship between factors and the EDCs' chemical activity (chemical features)	physicochemical properties of chemicals, high throughput screening of in vitro data (e.g., ToxCast from U.S.EPA), and toxicity data in vivo	active/inactive and the types of toxic activity	showed high prediction accuracy, broke the limitation of the "black box", and identified structural features	45, 47, 82
	soil/sediment	understanding the relationship between factors and uptake/transport of organic contaminants	physicochemical properties of chemicals	uptake efficiency	introduced new relationships for plant uptake of contaminants	83
environmental health	environmental	identifying key causal factors that predict environmental and human health outcomes	environmental exposure, water quality parameters, social economic factors	public health outcome (i.e., premature birth rate)	improved the identification of multiple factors that contribute to and predict health outcome	84, 85
	others	understanding the molecular mechanisms between environmental pollutants and nuclear receptors	computational information, such as molecular docking and classical molecular dynamics simulations	ligand–receptor interactions	found key residues and binding modes, built new virtual screening models	86, 87
		mechanism profiling of cellular stress response, hepatotoxicity caused by oxidative stress	in vitro assay models, time-series toxicogenomic data and big data	ranking in vitro–in vivo correlations; most relevant bioassay(s) related to hepatotoxicity, identifying key molecular toxicity mechanisms	fully explored the source-to outcome continuum of modern experimental toxicology	88, 89
		reducing animal experiments and exploring the relationship between in vitro bioassays and in vivo acute toxicity end points	high throughput screening in vitro data and in vivo acute toxicity data	nonanimal models for extrapolating in vitro to in vivo	built chemical fragment–in vitro–in vivo relationships	43

Table 1. continued

application	media	example	input	output	performance	refs
detecting anomalies	water	identifying contamination events	water quality indicators, for example, chlorine, electric conductivity, pH, temperature, total organic carbon, turbidity	probability of contamination	improved the identification of the source of fecal pollution in water	90, 91
		identifying the burst location of pipes	flow, pressure, or demand	probability of burst	deep learning models can successfully predict the bursts in a real-life network	92
		identifying abnormal events in wastewater treatment processes	wastewater influent and process variables	classified wastewater variables	successfully detected instrument and process anomalies	93
chemical assessment		QSAR models for governments, industry, and other stakeholders to fill gaps in (eco)toxicity data needed for assessing chemical hazards	chemical structural information and environmental information	structural characteristics and potential mechanism or mode of action of a target chemical	many QSAR models have been widely accepted, such as the OECD (Organization for Economic Co-operation and Development) QSAR Toolbox	94
	air	discovering new adsorption materials for CO ₂	edges, vertices, topologies of metal–organic frameworks (MOFs)	new MOF structures	discovered MOFs are strongly competitive against some of the best-performing MOFs/zeolites ever reported	95
water		discovering and designing new bioplastics materials, with desired durability and faster degradability in the environment than conventional plastics	polymer property data from literature and experiments, polymer features based on composition, configuration, topology, and polarity	prediction of target properties (e.g., Glass Transition Temperature) from candidate polymers	relationships determined between polymer chemical compositions and target properties; enabled the prediction of thermomechanical performance	96
	others	drug discovery and development of environmentally friendly chemicals; enabling new environmental technology	systematic and comprehensive high-dimensional data, single cell technology data	new environmentally friendly compounds; determine microbial community diversity and sampling size	challenges lie primarily with the lack of interpretability and repeatability of ML-generated results	97

different strategies have been used: the generative adversarial network (e.g., to discover new zeolite structures)¹⁰³ and the variational autoencoder (e.g., to obtain new structures for a metal–organic framework).¹⁰⁴ Such research typically has three steps: (1) train a model based on one of the ML algorithms, (2) generate new structures using the generator or decoder, and (3) validate the new structures by experiments or molecular simulation.

4. ESSENTIAL KNOWLEDGE REQUIREMENT AND CURRENT SHORTCOMINGS IN ML APPLICATIONS IN ESE

Although ML has shown great potential for solving ESE issues, inexperience with ML may lead to unsatisfactory performance or inappropriate applications of ML tools. This problem has been recognized by the closely affiliated material science and chemistry communities, which have felt the need to address “best practices” for ML.^{105,106} For example, applying cross-validation to the entire data set is one of the most common mistakes made by inexperienced researchers. Cross-validation should be applied to the training data set to screen ML algorithms and tune the hyperparameters,¹⁰⁷ and a completely independent test data set should be reserved to test the predictive performance of the obtained model.^{108,109} Researchers may confuse the validation data set with the test data set.¹¹⁰ Not doing this properly may have serious overfitting risks.¹¹¹ Seeking collaborations between researchers in the two fields can resolve these issues, but ESE researchers should acquire basic knowledge to fully benefit from the advantages of ML. For example, feature engineering (e.g., which features to use) and hyperparameters (e.g., the depth of trees in a random-forest model or the number of layers in a deep neural network) play important roles in the design of ML models, but often draw upon domain expertise. Therefore, effective collaboration will only be possible if ESE researchers understand fundamental ML concepts.

Figure 3 shows a typical workflow of ML model development and highlights key steps that should be carefully followed to deliver meaningful results when applying ML to ESE. The benefits of ML tools can only be realized if they are correctly used. For example, a common assumption in ML applications is that a larger volume of data will always generate better predictive performance of the ML model. However, more “representative” data rather than “big” data are more important for obtaining robust, powerful ML models. The term “representative” refers to the diversity of data. If a data set has too many similar data points, that is, they are close to each other in mathematical space, even though the sample size is large the model will not necessarily have the desired predictive performance. For instance, collecting chemicals with diverse chemical structures is preferred for developing widely applicable QSAR models. Once ML is deemed appropriate for a given application, collecting a sufficient amount of “representative” data is vital to the subsequent model development. It is also important that all key input features are considered in the input, as stated in a recent article that “if a well-trained neural network provided with sufficient data cannot achieve the desired accuracy, it is likely that the data provided is not entirely relevant to the task being predicted.”¹¹² The same statement may be true for other ML algorithms. Therefore, a deep understanding of the ESE application is crucial for developing correct and meaningful ML models.

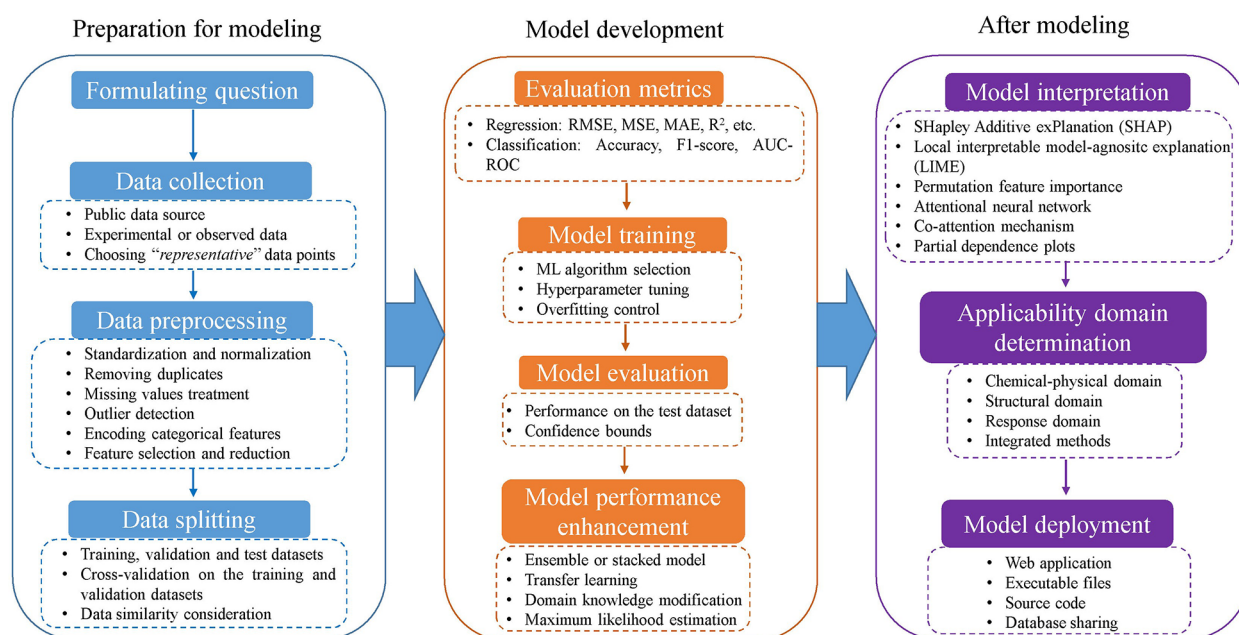


Figure 3. Typical workflow for the development of ML models, where RMSE, MSE, MAE, and R^2 refer to the root mean squared error, mean squared error, mean absolute error, and coefficient determination, while AUC-ROC means the area under the curve–receiver operating characteristic curve. For more details about specific ML techniques listed in Figure 3, please refer to the following references: data preprocessing, data splitting, and model development;^{113,114} model interpretation, including SHAP,¹¹⁵ LIME,¹¹⁶ permutation feature importance,⁶ attentional neural network,¹¹⁷ coattention mechanism,¹¹⁸ and partial dependence plots;¹¹⁹ and applicability domain determination.²⁵

Once a data set is ready, proper data preprocessing and splitting are necessary to avoid possible data leakage. Data leakage refers to cases where information used in model testing is inadvertently present during the model training process, so that the estimated prediction performance on the test set overestimates the true predictive performance of the model.¹²⁰ For example, duplicate samples may result in the same sample in both the training and test data sets, and the model may simply memorize these data without generalizing to new data points. Data leakage problems can often be subtle. To avoid data leakage, data standardization or normalization should be conducted based on the training data set and then applied to the test data set. For data splitting, the test data set should always be set aside and never be used for model development to ensure the independence of the test data set in evaluating the model performance. Cross-validation should be applied to the training data set rather than the entire data set, and the purpose is not to deliver a final predictive model but to evaluate the unbiased generalization ability of the ML algorithms. Feature selection can improve prediction accuracy, reduce model training time, and reduce the overfitting risk. Better model performance is commonly observed when an effective feature selection method is applied.^{39,50}

During model training, different evaluation metrics are used in different applications to evaluate the model performance; but in general, multiple metrics should be employed for a given model. For an imbalanced data set in which the number of positive cases is more than the number of negative cases, accuracy cannot be a reliable evaluation metric. Instead, metrics such as the F1 score,¹²¹ AUC (area under the curve)–ROC (receiver operating characteristics),¹²² or Matthews correlation coefficient (MCC) should be employed.¹²³ Tuning hyperparameters is required to calibrate a successful model.^{107,124} The optimum hyperparameters are determined to be the ones that minimize the errors on the validation data

set. After model development, the predictive performance of the model is evaluated on the test data set, which should never be used in model training or hyperparameter selection. This is consistent with a real application where the query sample is “new” and not merely identical with a previously observed training example. The model performance might be further improved by developing ensemble or stacked models, applying transfer learning, or performing domain knowledge modification. For example, Xiao et al. has developed an ensemble model that combines random forest, a generalized additive model, and extreme gradient boosting to predict historical $PM_{2.5}$ concentrations in China based on satellite data.²² However, it is not guaranteed that ensemble models are always better than single models, for example, when predicting the concentrations of ultrafine particles in the air.¹⁹

Model interpretation is an indispensable step after model development because it examines if the model predictions are consistent with the domain science (i.e., ground truth). Certain ML algorithms (e.g., ensemble methods, neural networks) are less interpretable than traditional statistical models.^{125,126} It will be problematic if a model with apparently high predictive performance makes predictions that are physically impossible; therefore, it is sometimes necessary to adjust the model and “teach” the model the correct domain science to obtain results that make sense. For example, based on the interpretation of the results of an ML model, one may need to remove the incorrectly identified features or adjust their weights and bias and retrain the model. Additional benefits related to model interpretation include (1) validating the reliability of the model, that is, if the model makes predictions aligned with domain knowledge; (2) unveiling implicit knowledge or even new knowledge that the model reveals; (3) informing feature engineering, such as selecting the most useful features; and (4) informing data collection, such as selecting data containing the desirable features during data collection.

Thus far, model interpretation has been largely overlooked in ESE. An example from the biomedical field demonstrates the necessity of model interpretation. McCloskey et al. used the attribution method to interpret their model and discovered that a deep neural network still learns spurious binding logic, despite its perfect classification accuracy on the protein–ligand binding data set.¹²⁷ Zhong et al. recently used the Shapley method to interpret how a deep neural network model and an extreme gradient boosting model made predictions on rate constants toward hydroxyl radicals ($\log k_{\text{OH}^\bullet}$).⁶⁰ They found that the former model assigned equal contributions of the same atom groups to $\log k_{\text{OH}^\bullet}$ no matter which compounds this atom group was in, which contradicts the chemical principle that the reactivity of a structural functional group is affected by its chemical environment. By comparison, the latter model correctly considered these contributions differently. Moreover, there are many complex ESE issues for which identifying the dominant drivers is extremely difficult. In these cases, if a successful model is obtained, one may be able to extract some implicit new knowledge that is otherwise unrecognized (see section 3.2 for more details).

Applicability domain (AD) is a concept from QSARs. It defines a domain in which the activity of a new compound can be reliably predicted based on the calibrated QSAR.²⁵ Recently this concept has been applied to materials science problems.¹²⁸ After model development, verifying the reliability of a prediction is the key idea of applicability domain analysis. However, studies seldom conduct AD analyses except for those applying QSAR models.²⁵ We propose expanding this concept to other ML models after they have been successfully developed and validated. There is no unified method to determine the AD of a model because diverse data types are used, such as image, text, and tabular data; and several measures may be applicable, for example, chemical-physical or structural domains for QSAR models.²⁵ Nevertheless, AD should be defined with a clear mathematic form, such as the similarity calculation for QSAR models.²⁵

Model deployment means how your model is shared with others. Commonly used model deployment approaches include sharing source code, providing executable files, and web applications. Sharing source code allows others to reuse or modify the models but requires expertise in coding; web applications or executable files provide ready-to-use tools to make predictions but limit the ability of other researchers to modify or augment the tools. Deploying ML models in more than one way may help ML models reach more users.

5. CHALLENGES

As an emerging tool for ESE, ML faces many challenges in applying it successfully. The following challenges are listed based on their order in the model development process, from data collection to model application.

5.1. Data Scarcity and Quality. The first challenge is how to effectively collect valid, high-quality data. For instance, since the outbreak of the COVID-19 pandemic, over 20 000 peer-reviewed papers have been published in a few months, among which many efforts have been launched to detect this virus in raw and treated wastewater.^{129–134} Ironically, the almost unlimited accessibility of data makes it hard to obtain the most relevant and qualified data. No generally agreed upon guidelines are available to sample, transport, store, and analyze the wastewater samples, leading to variations in the data quality. These low-quality data cannot be easily used for

modeling. On the other hand, ML requires a large sample size to build robust predictive models and make accurate predictions.^{135–137} Unfortunately, there is often a lack of mature databases for numerous environmental applications, and the data are often either scattered or not available in the literature. For example, researchers have systematically summarized the modes of action (MOAs) for thyroid hormone receptors (TRs)-related endocrine disruptions.¹³⁸ Developing high-throughput screening (HTS) in vitro assays based on MOAs can quickly generate a large volume of valid and reliable data for ML modeling. However, until recently, few in vitro assays were available to examine MOAs. For instance, EDCs can interact with one of 26 molecular processes to induce TRs-related endocrine disruptions, but only 14 out of the 26 (53.8%) processes have assays that have demonstrated reliability and are available for screening endocrine disrupting potentials.¹³⁸ Even when there are data, different researchers often conduct experiments under different conditions of water quality, soil/sediment properties, catalyst or adsorbent type and loading, etc., which creates discrepancies among the collected data. To develop predictive models that are truly robust and widely applicable, we should first build a large, consistent source data set.

5.2. Overfitting. Overfitting means that a model shows excellent predictive performance on the training samples but fails to accurately predict for new samples. This is especially a problem for more complex ML algorithms which contain large numbers of learned parameters. Concerns about overfitting and a lack of interpretability hinder, for example, the further optimization of automatic control processes in water/wastewater systems via ML algorithms compared with first-principle theoretical models.¹³⁹ Detecting overfitting remains a challenge, but methods that can reduce the overfitting risk include feature selection,¹⁴⁰ data augmentation,¹⁴¹ cross-validation,¹⁰⁸ regularization,¹⁴² model simplification/choice,¹⁴³ dropout,¹⁴⁴ and early stopping.¹⁴⁵

5.3. Bias of ML Models. In a technical sense, bias refers to systematic distortions of data sets and algorithms that result in undesirable outcomes. Data set biases occur when the training data are not representative of the planned use case and can arise when training data are inadvertently contaminated with the desired outcome information or when the training data are missing relevant examples. Data contamination problems are a more subtle form of the data leakage discussed above, and can be difficult to identify without domain expertise. For example, ML models trained to predict the severity of pneumonia from patient X-rays can inadvertently learn to distinguish the type of machine used to acquire the image (for example, the sickest patients are scanned with a portable X-ray machine, whereas healthier patients can be moved to a more sophisticated machine), rather than the information in the intended image itself.¹⁴⁶ Missing data is a problem in scientific communities, as peer-reviewed publications only report the most promising positive results, omitting negative results and outliers, which are crucial for ML model performance.¹⁴⁷ Data can also be missing because of choices by human scientists, such as habits in using particular reagents, reaction conditions, or sampling plans, or neglecting to collect data that violate a currently accepted theory. These types of anthropogenic biases in data sets also degrade ML performance.¹⁴⁸ Algorithmic bias occurs when the inherent structure of the model or loss function does not correspond to the desired use case. For example, the application of a linear classifier to data with nonlinear concave

or convex boundaries cannot succeed; it is necessary to choose a different type of ML model capable of describing the complex relationship embedded in the data. Similarly, the use of an inappropriate success metric (discussed above) during the training process will result in unintended predictions. The timely identification of the possible bias in an ML model is crucial for its applications (beyond that of even environmental issues). The issue of bias can be mitigated by improving the interpretability of the ML model, where domain knowledge can be incorporated to institute judgment on its validity. A practical strategy to identify bias in ML models is via convening an ensemble of ML models, which compares the results of different ML models on the same set of problems. This comparison will help identify the consistency in ML model performance and allow bias associated with a particular ML model to be identified.

5.4. Other Underlying Concerns. Besides the above-mentioned challenges, there are still some broader concerns that deserve to be further discussed. (1) We should not over trust or overestimate ML tools. It is always necessary to verify the findings either experimentally or based on the domain knowledge or experience. (2) Traditional statistical tools may be more appropriate than ML in some cases, such as when there are small sample sizes. (3) Not every ESE problem can be solved by ML tools directly. How to elegantly convert such ESE problems to ones that can be addressed by ML requires artful design.

6. FUTURE OPPORTUNITIES AND OUTLOOK

Although challenges lie ahead, there are still many opportunities, as described below.

6.1. Balance Model Fidelity and Interpretability. Environmental systems are characterized by complex interactions of various parameters and processes. Conventional modeling is often based on significant simplifications and assumptions relying on the domain knowledge of human experts. Despite many challenges to adopting ML in ESE, ML offers a unique opportunity to precisely predict the input and output relationships for complicated systems without a priori hypotheses. ML models are designed to make the most accurate predictions possible by understanding the complex relationships embedded in data. However, the complexity of environmental problems leads to an added layer of challenge regarding interpretation and bias for ML. The interpretation of the modeling results is often difficult as ML models are like “black box” networks in which the impact of inputs on the output cannot be understood easily. Integration of an ML model with a traditional mechanistic model may retain their respective merits and can be used to solve comprehensive problems. We encourage ESE researchers to think creatively within their domains and use ML when appropriate.

6.2. Data Sharing. Data sharing should be a consensus in ESE because a significant amount of time is spent on collecting and cleaning data before model development. It is essential to build an open-access data-sharing community in ESE where we can help increase the size and diversity of data by adding more experimental or observed data points, similar to ImageNet (<http://www.image-net.org/>). In this way, more accurate models with larger applicability domains can be developed to better serve ESE. It is desirable for the obtained databases and source codes for the models to be released to the public through platforms, such as GitHub (<http://github.com/>), a widely used web-based service for code sharing, review, and

management of open-source projects. Repositories such as DLHub (<http://dlhub.org>) not only archive data and program code, but also make it easy to use the models without the need to install software. We recommend that journal editors and peer-reviewers require that papers relying upon ML provide complete data sets and analysis source code in an electronic format (either in public repositories or as supporting information) as a necessary condition for publication, unless there are compelling privacy or legal nondisclosure reasons that preclude data-sharing.

6.3. Data Collection from Trusted Sources. Similar to other research areas, it is challenging to be inclusive of all valid publications in ESE and extract necessary information from different sources. Data can be obtained from stakeholders (e.g., water and wastewater utilities, government databases, etc.) or directly measured using instruments. Another trusted source of big data is the vast amount of literature available from scientific articles and reports. These traditional data collections are usually based on structured and homogeneous data, while textual data from the literature consider unstructured and structured texts with different formats and types using linguistic and statistical techniques. Literature text mining is a new area for ML and a void in ESE. Using ML methods to analyze ESE literature data effectively can reveal trends and patterns and underlying knowledge in complex and ever-changing environmental systems. For example, Zhu et al. proposed novel methods that analyze the complex publication data across many domains using deep text preprocessing,¹⁴⁹ such as acronym and abbreviation detection, chemical expression identification, and synonym combination, which has been beneficial in extracting the true value of big literature data. We recommend the creation and adoption of formal data and metadata schemas by the ESE community to facilitate data reuse.

6.4. Applications of ML Models. Beyond providing accurate predictions, one next step would be to create additional knowledge; for example, new materials or applications. On the one hand, ML models can help “discover” new information from the existing data. For example, one research group identified eight antibacterial compounds that are structurally distant from known antibiotics from the ZINC15 database (>107 million chemicals) based on an ML model.¹⁵⁰ On the other hand, ML can help create new chemical spaces beyond manually defined heuristics, which enables exploration of a vast landscape of possible material chemistries and properties that would not be possible to explore with an Edisonian trial and error approach.¹⁵¹ The flexibility of adjustment and the capability of pseudoreal time calculation and validation under varying conditions uniquely enable ML to have precise control of diverse ESE systems. This use of ML thus facilitates discovery and innovation.

6.5. Educational Opportunities. To take advantage of the rapid development of ML and unparalleled computational power, it is imperative that the next generation of ESE practitioners are prepared to properly utilize these tools. We recommend the creation of interdisciplinary certificates, specializations, and degrees in, for example, *Environmental Data Science*, which will train the next generation of environmental engineers and scientists at the interface of computer, data, and environmental sciences to prepare them for these challenges. Environmental students in this track may take several essential computer and data science courses, such as *Data Structures*, *Data Science*, *Intro to AI*, *Machine Learning*,

and Programming. Computer and data science students may take courses from the ESE curriculum to receive a certificate or minor in ESE.

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Notes

The authors declare no competing financial interest.

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