

A review of machine learning applications in wildfire science and management

Piyush Jain, Sean C.P. Coogan, Sriram Ganapathi Subramanian, Mark Crowley, Steve Taylor, and Mike D. Flannigan

Abstract: Artificial intelligence has been applied in wildfire science and management since the 1990s, with early applications including neural networks and expert systems. Since then, the field has rapidly progressed congruently with the wide adoption of machine learning (ML) methods in the environmental sciences. Here, we present a scoping review of ML applications in wildfire science and management. Our overall objective is to improve awareness of ML methods among wildfire researchers and managers, as well as illustrate the diverse and challenging range of problems in wildfire science available to ML data scientists. To that end, we first present an overview of popular ML approaches used in wildfire science to date and then review the use of ML in wildfire science as broadly categorized into six problem domains, including (i) fuels characterization, fire detection, and mapping; (ii) fire weather and climate change; (iii) fire occurrence, susceptibility, and risk; (iv) fire behavior prediction; (v) fire effects; and (vi) fire management. Furthermore, we discuss the advantages and limitations of various ML approaches relating to data size, computational requirements, generalizability, and interpretability, as well as identify opportunities for future advances in the science and management of wildfires within a data science context. In total, to the end of 2019, we identified 300 relevant publications in which the most frequently used ML methods across problem domains included random forests, MaxEnt, artificial neural networks, decision trees, support vector machines, and genetic algorithms. As such, there exists opportunities to apply more current ML methods — including deep learning and agent-based learning — in the wildfire sciences, especially in instances involving very large multivariate datasets. We must recognize, however, that despite the ability of ML models to learn on their own, expertise in wildfire science is necessary to ensure realistic modelling of fire processes across multiple scales, while the complexity of some ML methods such as deep learning requires a dedicated and sophisticated knowledge of their application. Finally, we stress that the wildfire research and management communities play an active role in providing relevant, high-quality, and freely available wildfire data for use by practitioners of ML methods.

Key words: machine learning, wildfire science, fire management, wildland fire, support vector machine, artificial neural network, decision trees, Bayesian networks, reinforcement learning, deep learning.

Résumé : L'intelligence artificielle a été utilisée en science et en gestion des feux de forêt depuis les années 1990, les premières applications comprenant les réseaux neuronaux et les systèmes experts. Depuis lors, le domaine a rapidement progressé parallèlement à l'adoption des méthodes d'apprentissage machine (AM) en sciences de l'environnement. Les auteurs présentent ici une synthèse du cadrage des applications de l'AM en science et en gestion des feux de forêt. Leur objectif global consiste à améliorer la notoriété des méthodes d'AM auprès des chercheurs et des gestionnaires des feux de forêt, de même qu'à illustrer l'étendue vaste et complexe des problèmes en science des feux de forêt dont disposent les scientifiques spécialistes de données en AM. À cette fin, ils présentent d'abord un survol des approches populaires en AM utilisées en science des feux de forêt à ce jour et font ensuite la synthèse de l'utilisation de l'AM en science des feux de forêt, selon six grands domaines de problèmes dont (i) la caractérisation des carburants, la détection et la cartographie de l'incendie; (ii) la température de l'incendie et les changements climatiques; (iii) les circonstances, la susceptibilité et le risque d'incendie; (iv) la prédiction du comportement de l'incendie; (v) les effets de l'incendie; et (vi) la gestion de l'incendie. Par ailleurs, les auteurs discutent des avantages et des limites de différentes approches d'AM en lien avec la taille des données, les exigences de calcul, le potentiel de généralisation et d'interprétation et identifient également les possibilités d'avancées futures en science et gestion des feux de forêt dans le contexte de la science des données. Ils ont identifié au total 300 publications pertinentes jusqu'à la fin de 2019 qui comprennent les méthodes d'AM les plus fréquemment utilisées à travers les domaines de problèmes, dont les forêts aléatoires, MaxEnt, les réseaux de neurones artificiels, les arbres de décision, les séparateurs à vaste marge et les algorithmes génétiques. Il existe ainsi des possibilités d'appliquer davantage de méthodes actuelles d'AM — y compris l'apprentissage profond et l'apprentissage basé sur l'agent — en sciences des feux de forêt, particulièrement dans les cas impliquant de très grands ensembles de données multivariées. Ils

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reconnaissent cependant que, malgré la capacité des méthodes en AM d'apprendre par elles-mêmes, l'expertise en science des feux de forêt est nécessaire pour s'assurer d'une modélisation réaliste des processus des incendies à différentes échelles, alors que la complexité de certaines méthodes en AM telles que l'apprentissage profond, requiert une connaissance approfondie et spécifique de leur application. Finalement, ils soulignent que les communautés qui se consacrent à la recherche et à la gestion des feux de forêt jouent un rôle actif en fournissant des données pertinentes, de haute qualité et en libre accès à l'usage des praticiens des méthodes en AM. [Traduit par la Rédaction]

Mots-clés : apprentissage machine, science des feux de forêt, gestion des feux, feu de végétation, séparateurs à vaste marge, réseau de neurones artificiels, arbres de décision, réseaux bayésiens, apprentissage par renforcement, apprentissage profond.

1. Introduction

Wildland fire is a widespread and critical element of the Earth's system (Bond and Keeley 2005) and a continuous global feature that occurs in every month of the year. Presently, global annual area burned is estimated to be approximately 420 Mha (Giglio et al. 2018), which is greater in area than the country of India. Globally, most of the area burned by wildfires occurs in grasslands and savannas. Humans are responsible for starting over 90% of wildland fires, and lightning is responsible for almost all of the remaining ignitions. Wildland fires can result in significant impacts to humans, either directly through loss of life and destruction to communities or indirectly through smoke exposure. Moreover, as the climate warms, we are seeing increasing impacts from wildland fire (Coogan et al. 2019). Consequently, billions of dollars are spent every year on fire management activities aimed at mitigating or preventing wildfires' negative effects. Understanding and better predicting wildfires is therefore crucial in several important areas of wildfire management, including emergency response, ecosystem management, land-use planning, and climate adaptation to name a few.

Wildland fire itself is a complex process; its occurrence and behavior are the product of several interrelated factors, including ignition source, fuel composition, weather, and topography. Furthermore, fire activity can be examined across a vast range of scales — from ignition and combustion processes that occur at a scale of centimetres over a period of seconds to fire spread and growth over minutes to days from metres to kilometres. At larger extents, measures of fire frequency may be measured over years to millennia at regional, continental, and planetary scales (see Simard (1991) for a classification of fire-severity scales and Taylor et al. (2013) for a review of numerical and statistical models that have been used to characterize and predict fire activity at a range of scales). For example, combustion and fire behavior are fundamentally physicochemical processes that can be usefully represented in mechanistic (i.e., physics-based) models at relatively fine scales (Coen 2018); however, such models are often limited by both the ability to resolve relevant physical processes and the quality and availability of input data (Hoffman et al. 2016). Moreover, with the limitations associated with currently available computing power, it is not feasible to apply physical models to inform fire management and research across the larger and longer scales that are needed sometimes in near real time. Thus, wildfire science and management rely heavily on the development of empirical and statistical models for meso-, synoptic-, strategic-, and global-scale processes (Simard 1991), the utility of which are dependent on their ability to represent the often complex and non-linear relationships among the variables of interest, as well as by the quality and availability of data.

While the complexities of wildland fire often present challenges for modelling, significant advances have been made in wildfire monitoring and observation primarily due to the increasing availability and capability of remote-sensing technologies. Several satellites (e.g., NASA TERRA and AQUA, NOAA GOES), for instance, have onboard fire detection sensors (e.g., Advanced Very High Resolution Radiometer (AVHRR), Moderate Resolution Imaging Spectroradiometer (MODIS), and Visible Infrared Imaging Ra-

diometer Suite (VIIRS)). These sensors, along with those on other satellites (e.g., LANDSAT series), routinely monitor vegetation distributions and changes. Additionally, improvements in numerical weather prediction and climate models are simultaneously offering smaller spatial resolutions and longer lead forecast times (Bauer et al. 2015), which potentially offer improved predictability of extreme fire weather events. Such developments make a data-centric approach to wildfire modeling a natural evolution for many research problems given sufficient data. Consequently, there has been a growing interest in the use of machine learning (ML) methodologies in wildfire science and management in recent years.

Although no formal definition exists, we adopt the conventional interpretation of ML as the study of computer algorithms that can improve automatically through experience (Mitchell 1997). This approach is necessarily data-centric, with the performance of ML algorithms dependent on the quality and quantity of available data relevant to the task at hand. The field of ML has undergone an explosion of new algorithmic advances in recent years and is deeply connected to the broader field of artificial intelligence (AI). AI researchers aim to understand and synthesize intelligent agents that can act appropriately to their situation and objectives, adapt to changing environments, and learn from experience (Poole and Mackworth 2010). The motivations for using AI for forested ecosystem related research, including disturbances due to wildfire, insects, and disease, were discussed in an earlier paper (Schmoldt 2001), while Olden et al. (2008) further argued for the use of ML methods to model complex problems in ecology. The use of ML models in the environmental sciences has seen a rapid uptake in the last decade, as evidenced by recent reviews in the geosciences (Karpatne et al. 2017), forest ecology (Liu et al. 2018), extreme weather prediction (McGovern et al. 2017), flood forecasting (Mosavi et al. 2018), statistical downscaling (Vandal et al. 2019), remote sensing (Lary et al. 2016), and water resources (Shen 2018; Sun and Scanlon 2019). Two recent perspectives have also made compelling arguments for the application of deep learning (DL) in Earth system sciences (Reichstein et al. 2019) and for tackling climate change (Rolnick et al. 2019). To date, however, no such paper has synthesized the diversity of ML approaches used in the various challenges facing wildland fire science.

In this paper, we review the current state of literature on ML applications in wildfire science and management. Our overall objective is to improve awareness of ML methods among fire researchers and managers and illustrate the diverse and challenging problems in wildfire open to data scientists. This paper is organized as follows. In section 2, we discuss commonly used ML methods, focusing on those most commonly encountered in wildfire science. In section 3, we give an overview of the scoping review and literature search methodology employed in this paper. In this section, we also highlight the results of our literature search and examine the uptake of ML methods in wildfire science since the 1990s. In section 4, we review the relevant literature within six broadly categorized wildfire modeling domains: (i) fuels characterization, fire detection, and mapping; (ii) fire weather and climate change; (iii) fire probability and risk; (iv) fire behavior prediction; (v) fire effects; and (vi) fire management. In section 5, we discuss our findings and identify further opportunities for the

application of ML methods in wildfire science and management. Finally, in section 6, we offer conclusions. Thus, this review will serve to guide and inform both researchers and practitioners in the wildfire community looking to use ML methods, as well as provide ML researchers with the opportunity to identify possible applications in wildfire science and management.

2. Artificial intelligence and machine learning

Machine learning can be defined as a set of methods that “detect patterns in data, use the uncovered patterns to predict future data or other outcomes of interest” from Machine Learning: A Probabilistic Perspective, 2012 (Murphy 2012).

ML itself can be seen as a branch of AI or statistics, depending who you ask, that focuses on building predictive, descriptive, or actionable models for a given problem by using collected data, or incoming data, specific to that problem. ML methods learn directly from data and dispense with the need for a large number of expert rules or the need to model individual environmental variables with perfect accuracy. ML algorithms develop their own internal model of the underlying distributions when learning from data and thus need not be explicitly provided with physical properties of different parameters. For example, in the task of modeling wildland fire spread, the relevant physical properties include fuel composition, local weather, and topography. The current state-of-the-art method in wildfire prediction includes physics-based simulators that fire fighters and strategic planners rely on to take many critical decisions regarding allocation of scarce fire-fighting resources in the event of a wildfire (Sullivan 2007). These physics-based simulators, however, have certain critical limitations: they normally render very low accuracies, have a prediction bias in regions where they are designed to be used, and are often hard to design and implement due to the requirement of a large number of expert rules. Furthermore, modelling many complex environmental variables is often difficult due to large resource requirements and complex or heterogeneous data formats. ML algorithms, however, learn their own mappings between parametric rules directly from the data and do not require expert rules, which is particularly advantageous when the number of parameters is quite large and their physical properties are quite complex, as in the case of wildland fire. Therefore, an ML approach to wildfire response may help to avoid many of the limitations of physics-based simulators.

A major goal of this review is to provide an overview of the various ML methods utilized in wildfire science and management. Importantly, we also provide a generalized framework for guiding wildfire scientists interested in applying ML methods to specific problem domains in wildland fire research. This conceptual framework, derived from the approach in Murphy (2012) and modified to show examples relevant to wildland fire and management, is shown in Fig. 1. In general, ML methods can be identified as belonging to one of three types: supervised learning, unsupervised learning, or agent-based learning. We describe each of these below.

Supervised learning — In supervised ML, all problems can be seen as one of learning a parametrized function, often called a “model”, that maps inputs (i.e., predictor variables) to outputs (or “target variables”), both of which are known. The goal of supervised learning is to use an algorithm to learn the parameters of that function using available data. In fact, both linear and logistic regressions can be seen as very simple forms of supervised learning. The most popular ML methods fall into this category.

Unsupervised learning — If the target variables are not available, then ML problems are typically much harder to solve. In unsupervised learning, the canonical tasks are dimensionality reduction and clustering, with relationships or patterns being extracted from the data without any guidance as to the “correct” answer. Extracting embedded dimensions that minimize variance

or assigning data points to (labelled) classes that maximize some notion of natural proximity or other measures of similarity are examples of unsupervised ML tasks.

Agent-based learning — Between supervised and unsupervised learning is a group of ML methods in which learning happens by simulating behaviors and interactions of a single autonomous agent or a group of autonomous agents. These are general unsupervised methods that use incomplete information about the target variables (i.e., information is available for some instances but not for others), requiring generalizable models to be learned. A specific case in this space is reinforcement learning (Sutton and Barto 1998), which is used to model decision-making problems over time whereby critical parts of the environment can only be observed interactively through trial and error. This class of problems arises often in the real world and requires efficient learning and careful definition of values (or preferences) and exploration strategies.

In the next section, we present a brief introduction to commonly used ML methods from the aforementioned learning paradigms. We note that this list is not meant to be exhaustive and that some methods can accommodate both supervised and unsupervised learning tasks. It should be noted that the classification of a method as belonging to either ML or traditional statistics is often a question of taste. For the purpose of this review, and in the interests of economy, we have designated a number of methods as belonging to traditional statistics rather than ML. For a complete listing, see Tables 1 and 2.

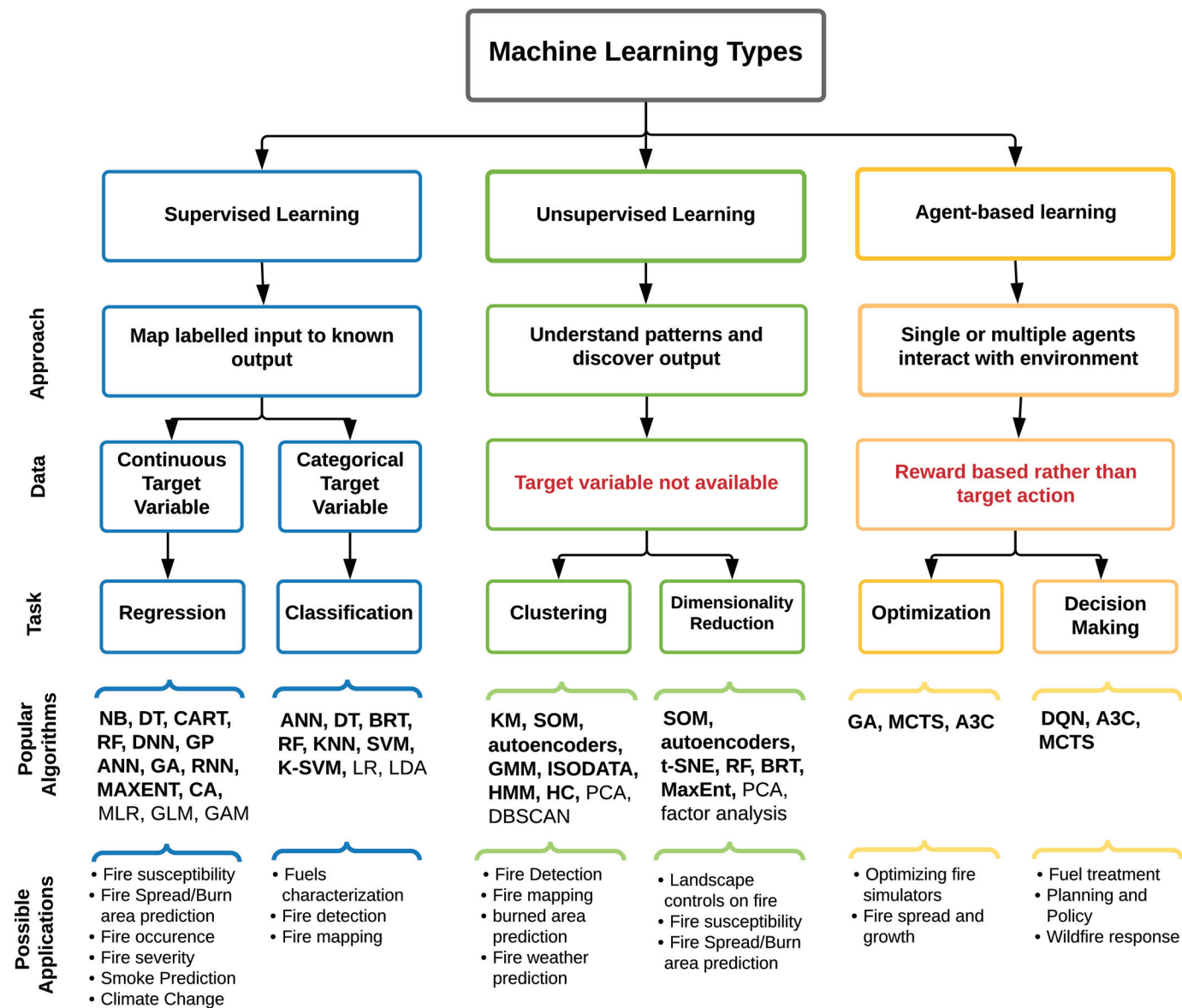
2.1. Decision trees

Decision trees (DTs) (Breiman et al. 1984) belong to the class of supervised learning algorithms and are another example of a universal function approximator, although in their basic form such universality is difficult to achieve. DTs can be used for both classification and regression problems. A DT is a set of if-then-else rules with multiple branches joined by decision nodes and terminated by leaf nodes. The decision node is where the tree splits into different branches, with each branch corresponding to the particular decision being taken by the algorithm, and leaf nodes represent the model output. This could be a label for a classification problem or a continuous value in the case of a regression problem. A large set of decision nodes is used in this way to build the DT. The objective of DTs is to accurately capture the relationships between input and outputs using the smallest possible tree that avoids overfitting. C4.5 (Quinlan 1993) and classification and regression trees (CART; Breiman et al. 1984) are examples of common single DT algorithms. Note that while the term CART is also used as an umbrella term for single-tree methods, we use DT here to refer to all such methods. The majority of DT applications are ensemble DT (EDT) models that use multiple trees in parallel (i.e., bootstrap aggregation or bagging) or sequentially (i.e., boosting) to arrive at a final model. In this way, EDTs make use of many weak learners to form a strong learner while being robust to overfitting. EDTs are well described in many ML-AI textbooks and are widely available as implemented libraries.

2.1.1. Random forests

A random forest (RF) (Breiman 2001) is an ensemble model composed of a many individually trained DTs and is the most popular implementation of a bagged DT. Each component DT in an RF model makes a classification decision whereby the class with the maximum number of votes is determined to be the final classification for the input data. RFs can also be used for regression with the final output being determined by averaging over the individual-tree outputs. The underlying principle of the RF algorithm is that a random subset of features is selected at each node of each tree; the samples for training each component tree are selected using bagging, which resamples (with replacement) the

Fig. 1. A diagram showing the main machine learning (ML) types, types of data, and modeling tasks in relation to popular algorithms and potential applications in wildfire science and management. Note that the algorithms shown in bold type are core ML methods, whereas those algorithms not in bold type are often not considered ML methods (see Tables 1 and 2).



original set of data points. The high performance of this algorithm is achieved by minimizing correlation between trees while reducing model variance so that a large number of different trees provides greater accuracy than individual trees; however, this improved performance comes at the cost of an increase in bias and loss of interpretability (although variable importance can still be inferred through permutation tests).

2.1.2. Boosted ensembles

Boosting describes a strategy in which one combines a set of weak learners, usually DTs, to make a strong learner using a sequential additive model. Each successive model improves on the previous one by taking into account the model errors from the previous model, which can be done in more than one way. For example, the adaptive boosting algorithm, known as AdaBoost (Freund and Shapire 1995), works by increasing the weight of observations that were previously misclassified. This can, in principle, reduce the classification error leading to a high level of precision (Hastie et al. 2009).

Another very popular implementation for ensemble boosted trees is the gradient boosting machine (GBM), which makes use of the fact that each DT model represents a function that can be differentiated with respect to its parameters, i.e., how much a change in the parameters will change the output of the function. GBMs sequentially build an ensemble of multiple weak learners by following a simple gradient that points in the opposite direction to weakest results of the current combined model (Friedman 2001).

The details for the GBM algorithm are as follows. Denoting the target output as Y and given a tree-based ensemble model, represented as a function $T_i(X) \rightarrow Y$, after already adding i weak learners, the "perfect" function for weak learner $i + 1$ would be $h(x) = T_i(x) - Y$, which exactly corrects the previous model (i.e., $T_{(i+1)}(x) = T_i(x) + h(x) = Y$). In practice, we can only approach this perfect update by performing functional gradient descent in which we use an approximation of the true residual (i.e., loss function) at each step. In our case, this approximation is simply the sum of the residuals

Table 1. Table of abbreviations and definitions for common machine learning algorithms referred to in the text.

Abbreviation	Definition
A3C	Asynchronous advantage actor-critic
AdaBoost	Adaptive boosting
ANFIS	Adaptive neuro fuzzy inference system
ANN	Artificial neural networks
ADP	Approximate dynamic programming (also known as reinforcement learning)
Bag	Bagged decision trees
BN	Bayesian networks
BRT	Boosted regression trees (also known as gradient boosted machine)
BULC	Bayesian updating of land cover
CART	Classification and regression tree
CNN	Convolutional neural network
DNN	Deep neural network
DQN	Deep Q-network
DT	Decision trees (including CART, J48, and jRip)
EDT	Ensemble decision trees (including bagging and boosting)
ELM	Extreme machine learning (i.e., feedforward network)
GA	Genetic algorithms (also known as evolutionary algorithms)
GBM	Gradient boosted machine (also known as boosted regression trees, including XGBoost, AdaBoost, and LogitBoost)
GMM	Gaussian mixture models
GP	Gaussian processes
HCL	Hard competitive learning
HMM	Hidden Markov models
ISODATA	Iterative self-organizing DATA algorithm
KNN	<i>k</i> nearest neighbor
KM	<i>k</i> -means clustering
LB	LogitBoost (including AdaBoost)
LSTM	Long short-term memory
MaxEnt	Maximum entropy
MCMC	Markov chain Monte Carlo
MCTS	Monte Carlo tree search
MLP	Multilayer perceptron
MDP	Markov decision process
NB	Naive Bayes
NFM	Neuro-fuzzy models
PSO	Particle swarm optimization
RF	Random forest
RL	Reinforcement learning
RNN	Recurrent neural network
SGB	Stochastic gradient boosting
SOM	Self-organizing maps
SVM	Support vector machines
t-SNE	<i>t</i> -distributed stochastic neighbor embedding

Table 2. Table of abbreviations and definitions for common data analysis algorithms usually considered as foundational to, or outside of, machine learning itself.

Abbreviation	Definition
DBSCAN	Density-based spatial clustering of applications with noise
GAM	Generalized additive model
GLM	Generalized linear model
KLR	Kernel logistic regression
LDA	Linear discriminant analysis
LR	Logistic regression
MARS	Multivariate adaptive regression splines
MLR	Multiple linear regression
PCA	Principal component analysis
SLR	Simple linear regression

from each weak learner DT , $L(Y, T(X)) = \sum_i Y - T_i(X)$. GBM explicitly uses the gradient $\nabla_T L(Y, T_i(X))$ of the loss function of each tree to fit a new tree and add it to the ensemble.

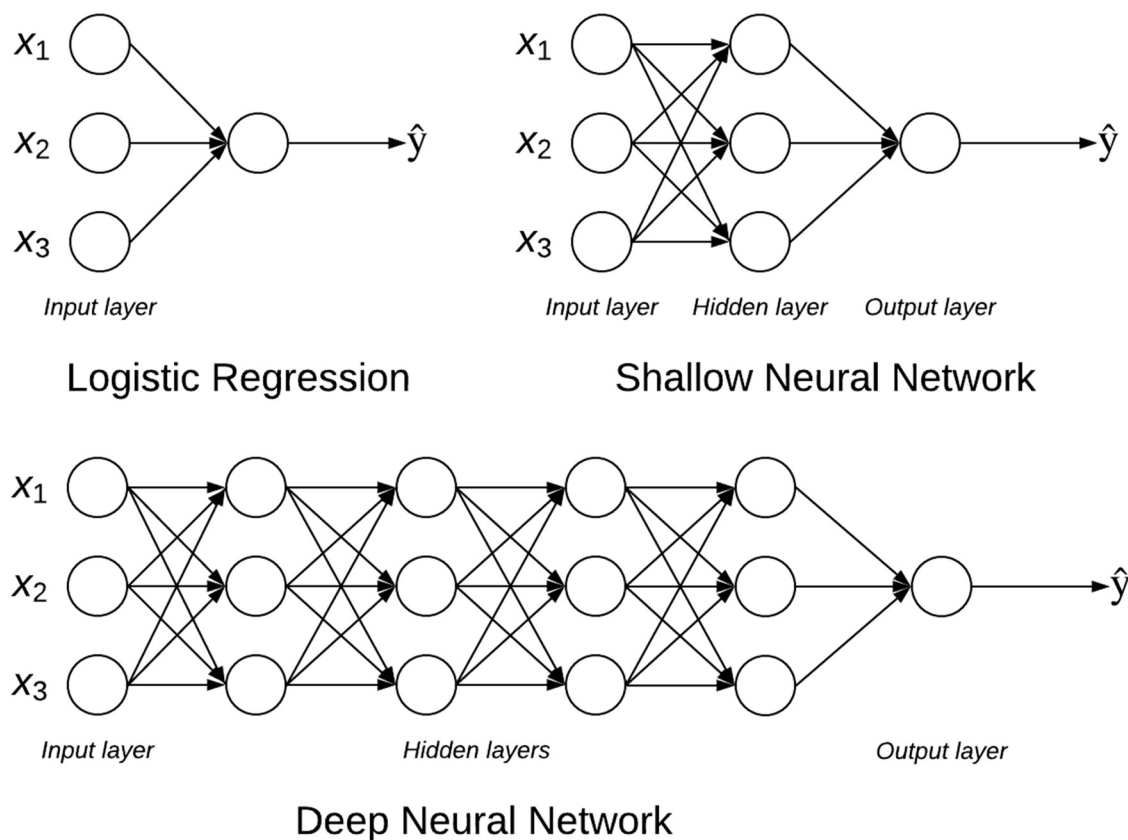
In a number of domains and particularly in the context of ecological modeling, GBM is often referred to as boosted regression

trees (BRTs) (Elith et al. 2008). For consistency with most of the literature reviewed in this paper, we henceforth use the latter term. It should be noted that while deep neural networks (DNNs) and EDT methods are both universal function approximators, EDTs are more easily interpretable and faster to learn with less data than DNNs. There are, however, fewer and fewer cases in which tree-based methods can be shown to provide superior performance on any particular metric when DNNs are trained properly with enough data (see, for example, Korotcov et al. 2017).

2.2. Support vector machines

Another category of supervised learning includes support vector machines (SVMs) (Hearst et al. 1998) and related kernel-based methods. SVM is a classifier that determines the hyperplane (decision boundary) in an n -dimensional space separating the boundary of each class, for data in n dimensions. SVM finds the optimal hyperplane in such a way that the distance between the nearest point of each class and the decision boundary is maximized. If the data can be separated by a line, then the hyperplane is defined to be of the form $w^T x + b = 0$, where the w is the weight vector, x is the input vector, and b is the bias. The distance of the hyperplane to the closest data point d , called a support vector, is defined as the

Fig. 2. Logistic regression can be seen as a basic building block for neural networks, with no hidden layer and a sigmoid activation function. Classic shallow neural networks (also known as multilayer perceptrons) have at least one hidden layer and can have a variety of activation functions. Deep neural networks essentially have a much larger number of hidden layers and use additional regularization and optimization methods to enhance training.



margin of separation. The objective is to find the optimal hyperplane that minimizes the margin. If they are not linearly separable, kernel SVM methods such as radial basis functions (RBF) first apply a set of transformations to the data to a higher dimensional space where finding this hyperplane would be easier. SVMs have been widely used for both classification and regression problems, although recently developed DL algorithms have proven to be more efficient than SVMs given a large amount of training data; however, for problems with limited training samples, SVMs might give better performances than DL-based classifiers.

2.3. Artificial neural networks and deep learning

The basic unit of an artificial neural network (ANN) is a neuron (also called a perceptron or logistic unit). A neuron is inspired by the functioning of neurons in mammalian brains in that it can learn simple associations, but in reality, it is much simpler than its biological counterpart. A neuron has a set of inputs that are combined linearly through multiplication with weights associated with the input. The final weighted sum forms the output signal, which is then passed through a (generally) nonlinear activation function. Examples of activation functions include sigmoid, tanh, and the rectified linear unit (ReLU). This nonlinearity is important for general learning as it creates an abrupt cutoff (or threshold) between positive and negative signals. The weights on each connection represent the function parameters, which are fit using supervised learning by optimizing the threshold so that it reaches a maximally distinguishing value.

In practice, even simple ANNs, often called multilayered perceptrons (MLP), combine many neuron units in parallel, each processing the same input with independent weights. In addition, a second layer of hidden neuron units can be added to allow more

degrees of freedom to fit general functions (see Fig. 2). MLPs are capable of solving simple classification and regression problems. For instance, if the task is one of classification, then the output is the predicted class for the input data, whereas in the case of a regression task, the output is the regressed value for the input data. Deep learning (LeCun et al. 2015) refers to using deep neural networks (DNNs), which are ANNs with multiple hidden layers (nominally more than three) and include convolutional neural networks (CNNs), popularized in image analysis, and recurrent neural networks (RNNs), which can be used to model dynamic temporal phenomena. The architecture of DNNs can vary in connectivity between nodes, the number of layers employed, the types of activation functions used, and many other types of hyperparameters. Nodes within a single layer can be fully connected or connected with some form of convolutional layer (e.g., CNNs), recurrent units (e.g., RNNs), or other sparse connectivity. The only requirement of all these connectivity structures and activation functions is that they are differentiable.

Regardless of the architecture, the most common process of training an ANN involves processing input data fed through the network layers and activation functions to produce an output. In the supervised setting, this output is then compared with the known true output (i.e., labelled training data), resulting in an error measurement (loss or cost function) used to evaluate model performance. The error for DNNs is commonly calculated as a cross-entropy loss between the predicted output label and the true output label. As every part of the network is mathematically differentiable, we can compute a gradient for the entire network. This gradient is used to calculate the proportional change in each network weight needed to produce an infinitesimal increase in

the likelihood of the network producing the same output for the most recent output. The gradient is then weighted by the computed error, and thereafter, all the weights are updated in sequence using a backpropagation algorithm (Hecht-Nielsen 1992).

ANNs can also be configured for unsupervised learning tasks. For example, self-organizing maps (SOMs) are a form of ANN adapted for dealing with spatial data and have therefore found widespread use in the atmospheric sciences (Skific and Francis 2012). A SOM is a form of unsupervised learning that consists of a two-dimensional array of nodes as the input layer, representing, for example, a gridded atmospheric variable at a single time. The algorithm clusters similar atmospheric patterns together, resulting in a dimensionality reduction of the input data. More recently, unsupervised learning methods from DL such as autoencoder networks are starting to replace SOMs in the environmental sciences (Shen 2018).

2.4. Bayesian methods

2.4.1. Bayesian networks

Bayesian networks (Bayes net, belief network; BN) are a popular tool in many applied domains because they provide an intuitive graphical language for specifying the probabilistic relationships between variables, as well as the tools for calculating the resulting probabilities (Pearl 1988). The basis of BNs is Bayes' theorem, which relates the conditional and marginal probabilities of random variables. BNs can be treated as an ML task if one is trying to automatically fit the parameters of the model from data or, even more challenging, to learn the best graphical structure that should be used to represent a dataset. BNs have close ties to causal reasoning, but it is important to remember that the relationships encoded in a BN are inherently correlational rather than causal. BNs are acyclic graphs, consisting of nodes and arrows (or arcs), defining a probability distribution over variables \mathcal{U} . The set of parents of a node (variable) X , denoted π_X , are all nodes with directed arcs going into X . BNs provide compact representation of conditional distributions as $p(X_i | X_1, \dots, X_{i-1}) = p(X_i | \pi_{X_i})$, where X_1, \dots, X_{i-1} are arranged to be all of the ancestors of X_i other than its direct parents. Each node X is associated with a conditional probability table over X and its parents defining $p(X | \pi_X)$. If a node has no parents, a prior distribution is specified for $p(X)$. The joint probability distribution of the network is then specified by the chain rule $P(\mathcal{U}) = \prod_{X \in \mathcal{U}} p(X | \pi_X)$.

2.4.2. Naïve Bayes

A special case of a BN is the naïve Bayes (NB) classifier, which assumes conditional independence between input features, allowing the likelihood function to be constructed by a simple multiplication of the conditional probability of each input variable conditional on the output. Therefore, while NB is fast and straightforward to implement, prediction accuracy can be low for problems in which the assumption of conditional independence does not hold.

2.4.3. Maximum entropy

Maximum entropy (MaxEnt), originally introduced by Phillips et al. (2006), is a presence-only framework that fits a spatial probability distribution by maximizing entropy, consistent with existing knowledge. MaxEnt can be considered a Bayesian method as it is compatible with an application of Bayes theorem as existing knowledge is equivalent to specifying a prior distribution. MaxEnt has found widespread use in landscape ecology species distribution modeling (Elith et al. 2011), where prior knowledge consists of occurrence observations for the species of interest.

2.5. Reward-based methods

2.5.1. Genetic algorithms

Genetic algorithms (GA) are heuristic algorithms inspired by Darwin's theory of evolution (natural selection) and belong to a more general class of evolutionary algorithms (Mitchell 1996). GAs are often used to generate solutions to search and optimization problems by using biologically motivated operators such as mutation, crossover, and selection. In general, GAs involve several steps. The first step involves creating an initial population of potential solutions, with each solution encoded as a chromosome. In the second step, a fitness function appropriate to the problem is defined, which returns a fitness score determining how likely an individual is to be chosen for reproduction. The third step requires the selection of pairs of individuals, denoted as parents. In the fourth step, a new population of finite individuals is created by generating two new offspring from each set of parents using crossover whereby a new chromosome is created by some random selection process from each parent's chromosomes. In the final step called mutation, a small sample of the new population is chosen, and a small perturbation is made to the parameters to maintain diversity. The entire process is repeated many times until the desired results are satisfactory (based on the fitness function) or some measure of convergence is reached.

2.5.2. Reinforcement learning

Reinforcement learning (RL) represents a very different learning paradigm to supervised or unsupervised learning. In RL, an agent (or actor) interacts with its environment and learns a desired behavior (set of actions) to maximize some reward. RL is a solution to a Markov decision process (MDP) in which the transition probabilities are not explicitly known but need to be learned. This type of learning is well suited to problems of decision-making such as required for automated control (e.g., robotics) or for system optimization (e.g., management policies). Various RL algorithms include Monte Carlo tree search (MCTS), Q-learning, and actor-critic algorithms. For an introduction to RL, see Sutton and Barto (2018).

2.6. Clustering methods

Clustering is the process of splitting a set of points into groups such that each point in a group is more similar to its own group than to any other group. There are different ways in which clustering can be done; for example, the K-means clustering (KM) algorithm (MacQueen 1967), based on a centroid model, is perhaps the most well-known clustering algorithm. In KM, the notion of similarity is based on closeness to the centroid of each cluster. KM is an iterative process in which the centroid of a group and points belonging to a group are updated at each step. The KM algorithm consists of five steps: (i) specify the number of clusters; (ii) randomly assign each data point to a cluster; (iii) calculate the centroids of each cluster; (iv) reassign the points to the nearest centroids; and (v) recompute the cluster centroids. Steps iv and v repeat until no further changes are possible. Although KM is the most widely used clustering algorithm, several other clustering algorithms exist, for example, agglomerative hierarchical clustering (HC), Gaussian mixture models (GMMs), and iterative self-organizing Data (ISODATA).

2.7. Other methods

2.7.1. k-Nearest neighbor

The k-nearest neighbor (KNN) algorithm is a simple but very effective supervised classification algorithm based on the intuitive premise that similar data points are in close proximity according to some metric (Altman 1992). Specifically, a KNN calculates the similarity of data points to each other using the Euclidean distance between the k nearest data points. The optimal value of k can be found experimentally over a range of values using the

classification error. KNN is widely used in applications in which a search query is performed such that the results should be similar to another pre-existing entity. Examples of this include finding similar images to a specified image and recommender systems. Another popular application of KNN is outlier (or anomaly) detection whereby the points (in a multidimensional space) farthest away from their nearest neighbors may be classified as outliers.

2.7.2. Neuro-fuzzy models

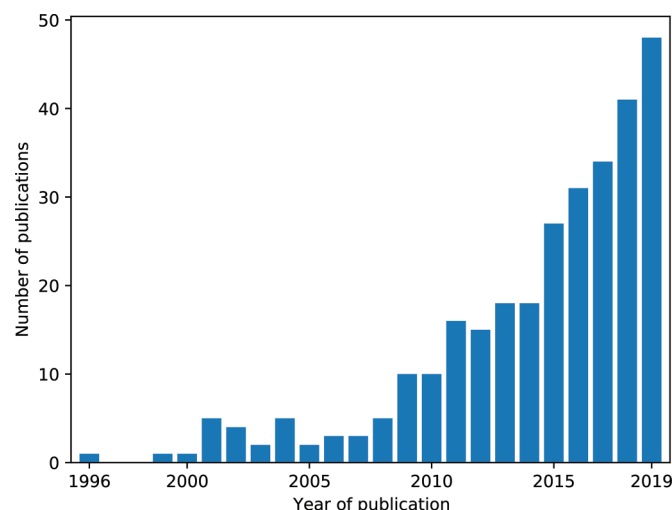
Fuzzy logic is an approach for encoding expert human knowledge into a system by defining logical rules about how different classes overlap and interact without being constrained to “all-or-nothing” notions of set inclusion or probability of occurrence. Although early implementations of fuzzy logic systems depended on setting rules manually and therefore are not considered ML, using fuzzy rules as inputs or extracting them from ML methods is often described as “neuro-fuzzy” methods. For example, the adaptive neuro-fuzzy inference system (ANFIS) (Jang 1993) fuses fuzzy logical rules with an ANN approach, while trying to maintain the benefits of both. ANFIS is a universal function approximator like ANNs; however, as this algorithm originated in the 1990s, it precedes the recent DL revolution so is not necessarily appropriate for very large data problems with complex patterns arising in high-dimensional spaces. Alternatively, human-acquired fuzzy rules can be integrated into ANNs learning; however, it is not guaranteed that the resulting trained neural network will still be interpretable. It should be noted that fuzzy rules and fuzzy logic are not a major direction of research within the core ML community.

3. Literature search and scoping review

The combination of ML and wildfire science and management comprises a diverse range of topics in a relatively nascent field of multidisciplinary research. Thus, we employed a scoping review methodology (Arksey and O'Malley 2005; Levac et al. 2010) for this paper. The goal of a scoping review is to characterize the existing literature in a particular field of study, particularly when a topic has yet to be extensively reviewed and the related concepts are complex and heterogeneous (Pham et al. 2014). Furthermore, scoping reviews can be particularly useful for summarizing and disseminating research findings and for identifying research gaps in the published literature. A critical review of methodological advances and limitations and comparison with other methods is left for future work. We performed a literature search using the Google Scholar and Scopus databases and the key words “wildfire” or “wildland fire” or “forest fire” or “bushfire” in combination with “machine learning” or “random forest” or “decision trees” or “regression trees” or “support vector machine” or “maximum entropy” or “neural network” or “deep learning” or “reinforcement learning”. We also used the Fire Research Institute's online database (<http://fireresearchinstitute.org>) using the following search terms: “Artificial Intelligence”; “Machine Learning”; “Random Forests”; “Expert Systems”; and “Support Vector Machines”. Furthermore, we obtained papers from references cited within papers that we had obtained using the aforementioned literature databases.

After performing our literature search, we identified a total of 300 publications relevant to the topic of ML applications in wildfire science and management (see Supplementary material¹ for a full bibliography). Furthermore, a search of the Scopus database revealed a dramatic increase in the number of wildfire and ML articles published in recent years (see Fig. 3). After identifying publications for review, we further applied the following criteria to exclude nonrelevant or unsuitable publications: (i) conference

Fig. 3. Number of publications by year for 300 publications on topic of machine learning and wildfire science and management as identified in this review.



submissions in which a journal publication describing the same work was available, (ii) conference posters, (iii) articles in which the methodology and results were not adequately described to conduct an assessment of the study, (iv) articles not available to us either by open access or by subscription, and (v) studies that did not present new methodologies or results.

4. Wildfire applications

In summary, we found a total of 300 journal papers or conference proceedings on the topic of ML applications in wildfire science and management, published up to the end of 2019 (Fig. 4). We found that the problem domains with the highest application of ML methods were fire occurrence, susceptibility, and risk (127 papers) followed by fuels characterization, fire detection, and mapping (66 papers), fire behavior prediction (43 papers), fire effects (35 papers), fire weather and climate change (20 papers), and fire management (16 papers). Within fire occurrence, susceptibility, and risk, the subdomains with the most papers were fire-susceptibility mapping (71 papers) and landscape controls on fire (101 papers). Note that some papers appear in multiple problem domains or subdomains. Refer to Table 3 and the Supplementary material¹ for a breakdown of each problem subdomain and the ML methods used, as well as study areas considered.

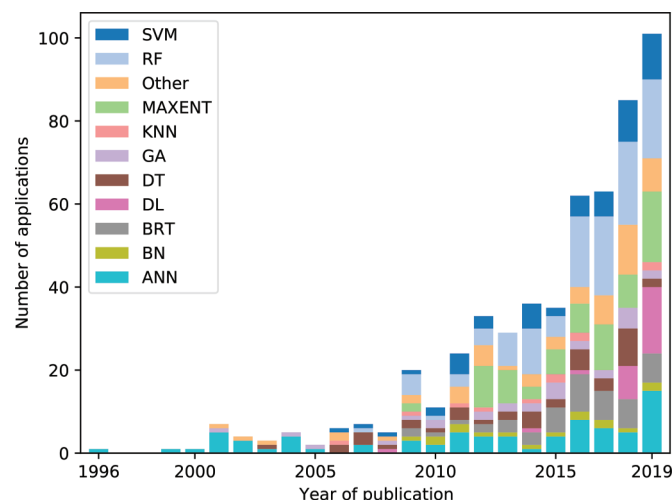
4.1. Fuels characterization, fire detection, and mapping

4.1.1. Fuels characterization

Fires ignite in a few fuel particles; subsequent heat transfer between particles through conduction, radiation, and convection and the resulting fire behavior (fuel consumption, spread rate, intensity) is influenced by properties of the live and dead vegetative fuels, including moisture content, biomass, and vertical and horizontal distribution. Fuel properties are a required input in all fire behavior models, whether it be a simple categorical vegetation type, as in the Canadian Fire Behavior Prediction (FBP) System, or as physical quantities in three-dimensional space (e.g., see FIRETEC model; Linn et al. 2002). Research to predict fuel properties has been carried out at two different scales: (i) regression applications to predict quantities such as the crown biomass of single trees from more easily measured variables such as height and diameter, and (ii) classification applications to map fuel type

¹Supplementary material is available with the article through the journal Web site at <http://nrcresearchpress.com/doi/suppl/10.1139/er-2020-0019>.

Fig. 4. Number of machine learning (ML) applications by category and year for 300 publications on topics of ML and wildfire science and management as identified in this review. See Table 1 for the definition of abbreviations.



descriptors or fuel quantities over a landscape from visual interpretation of air photographs or by interpretation of the spectral properties of remote sensing imagery. Relatively few studies, however, have employed ML to wildfire fuel prediction, leaving the potential for substantially more research in this area.

In an early study, Riaño et al. (2005) used an ANN to predict and map the equivalent water thickness and dry matter content of wet and dry leaf samples from 49 species of broadleaf plants using reflectance and transmittance values in the Ispra region of Italy. Pierce et al. (2012) used RF to classify important canopy fuel variables (e.g., canopy cover, canopy height, canopy base height, and canopy bulk density) related to wildland fire in Lassen Volcanic National Park, California, using field measurements, topographic data, and NDVI to produce forest canopy fuel maps. Likewise, Riley et al. (2014) used RF with Landfire and biophysical variables to perform fuel classification and mapping in eastern Oregon. The authors of the aforementioned study achieved relatively high overall modelling accuracy, for example, 97% for forest height, 86% for forest cover, and 84% for existing vegetation group (i.e., fuel type). López-Serrano et al. (2016) compared the performance of three common ML methods (SVM, KNN, and RF) and multiple linear regression in estimating aboveground biomass in the Sierra Madre Occidental, Mexico. The authors reported the advantages and limitations of each method and concluded that the nonparametric ML methods had an advantage over multiple linear regression for biomass estimation. García et al. (2011) used SVM to classify LiDAR and multispectral data to map fuel types in Spain. Chirici et al. (2013) compared the use of CART, RF, and stochastic gradient boosting (SGB), an ensemble tree method that uses both boosting and bagging, for mapping forest fuel types in Italy and found that SGB had the highest overall accuracy.

4.1.2. Fire detection

Detecting wildfires as soon as possible after they have ignited, and therefore while they are still relatively small, is critical to facilitating a quick and effective response. Traditionally, fires have mainly been detected by human observers by distinguishing smoke in the field of view directly from a fire tower or from a video feed from a tower, aircraft, or from the ground. All of these methods can be limited by spatial or temporal coverage, human error, the presence of smoke from other fires, and hours of daylight. Automated detection of heat signatures or smoke in infrared (IR) or optical images can extend the spatial and temporal coverage of detection, the detection efficiency in smoky condi-

tions, and remove bias associated with human observation. The analytical task is a classification problem that is quite well suited to ML methods.

For example, Arrue et al. (2000) used ANNs and IR image processing (in combination with visual imagery and meteorological and geographic data used in a decision function using fuzzy logic) to identify true wildfires. Several researchers have similarly employed ANNs for fire detection (Al-Rawi et al. 2001; Angayarkkani and Radhakrishnan 2010; Fernandes et al. 2004a, 2004b; X. Li et al. 2015; Soliman et al. 2010; Utkin et al. 2002; Sayad et al. 2019). In addition, Liu et al. (2015) used ANNs on wireless sensor networks to build a fire detection system in which multicriteria detection was used on multiple attributes (e.g., flame, heat, light, and radiation) to detect and raise alarms. Other ML methods used in fire detection systems include SVM to automatically detect wildfires from videoframes (Zhao et al. 2011), GA for multi-objective optimization of a LiDAR-based fire detection system (Cordoba et al. 2004), BN in a vision-based early fire detection system (Ko et al. 2010), ANFIS (Angayarkkani and Radhakrishnan 2011; Wang et al. 2011), and KM (Srinivasa et al. 2008).

CNNs (i.e., deep learning), which are able to extract features and patterns from spatial images and are finding widespread use in object detection tasks, have recently been applied to the problem of fire detection. Several of these applications trained the models on terrestrial-based images of fire and (or) smoke (Zhang et al. 2016; B. Zhang et al. 2018; Q.X. Zhang 2018; Yuan et al. 2018; Akhloufi et al. 2018; Barmoutis et al. 2019; Jakubowski et al. 2019; Sousa et al. 2019; X. Li et al. 2018, T. Li et al. 2019; Muhammad et al. 2018; Wang et al. 2019). Of particular note, Q.X. Zhang et al. (2018) found that CNNs outperformed a SVM-based method, and Barmoutis et al. (2019) found that a faster region-based CNN outperformed another CNN based on YOLO ("you only look once"). Yuan et al. (2018) used CNN combined with optical flow to include time-dependent information. X. Li et al. (2018) similarly used a three-dimensional CNN to incorporate both spatial and temporal information and so were able to treat smoke detection as a segmentation problem for video images. Another approach by Cao et al. (2019) used convolutional layers as part of a long short-term memory (LSTM) neural network for smoke detection from a sequence of images (i.e., video feed). They found that the LSTM method achieved 97.8% accuracy, a 4.4% improvement over a single image based DL method.

Perhaps of greater utility for fire management were fire/smoke detection models trained on either unmanned aerial vehicle (UAV) images (Zhao et al. 2018; Alexandrov et al. 2019) or satellite imagery including GOES-16 (Phan and Nguyen 2019) and MODIS (Ba et al. 2019). Zhao et al. (2018) compared SVM, ANN, and three CNN models and found that their 15-layer CNN performed best, with an accuracy of 98%. By comparison, the SVM-based method, which was unable to extract spatial features, only had an accuracy of 43%. Alexandrov et al. (2019) found that YOLO was both faster and more accurate than a region-based CNN method in contrast to Barmoutis et al. (2019).

4.1.3. Fire perimeter and severity mapping

Fire maps have two management applications: (i) accurate maps of the location of the active fire perimeter are important for daily planning of suppression activities and (or) evacuations, including modeling fire growth; and (ii) maps of the final burn perimeter and fire severity are important for assessing and predicting the economic and ecological impacts of wildland fire and for recovery planning. Historically, fire perimeters were sketch-mapped from the air, from a ground or aerial GPS or other traverse, or by air-photo interpretation. Developing methods for mapping fire perimeters and burn severity (a measure of above- and below-ground biomass loss due to fire) from remote sensing imagery has been an area of active research since the advent of remote sensing in the 1970s and is mainly concerned with classifying active fire areas

Table 3. Summary of machine learning (ML) methods applied to different problem domains in wildfire science and management.

Section	Domain	NFM	SVM	KM	GA	BN	BRT	ANN	DT	RF	KNN	MAXENT	DL	NB	Other
1.1	Fuels characterization	—	2	—	—	—	1	1	1	4	1	—	—	—	—
1.2	Fire detection	2	3	1	1	1	—	12	—	—	—	—	18	—	3
1.3	Fire perimeter and severity mapping	1	12	1	2	—	1	6	1	4	2	1	—	—	6
2.1	Fire-weather prediction	—	—	1	—	—	—	—	—	1	—	—	—	—	3
2.2	Lightning prediction	—	—	—	—	—	—	—	1	2	—	—	—	—	—
2.3	Climate change	—	1	—	—	—	6	2	2	5	—	7	—	—	—
3.1	Fire occurrence prediction	—	3	—	—	1	—	7	1	5	1	2	—	1	4
3.2	Landscape-scale burned area prediction	—	1	1	1	—	—	1	1	2	—	1	1	—	1
3.3	Fire susceptibility mapping	2	12	1	3	2	8	16	9	26	—	27	1	2	3
3.4	Landscape controls on fire	2	10	1	3	2	19	11	15	40	1	30	1	1	2
4.1	Fire spread and growth	—	—	—	13	2	—	4	—	1	1	—	3	—	2
4.2	Burned area and fire-severity prediction	—	7	—	1	1	3	10	7	6	3	—	2	1	5
5.1	Soil erosion and deposits	—	—	1	—	—	—	1	1	—	—	1	—	—	—
5.2	Smoke and particulate levels	—	2	—	—	—	3	3	—	5	2	—	—	—	2
5.3	Post-fire regeneration and ecology	—	1	—	1	1	6	1	2	10	—	2	—	1	—
5.4	Socioeconomic effects	—	—	—	—	1	—	—	—	—	—	—	—	—	—
6.1	Planning and policy	—	—	—	1	1	—	—	—	2	—	—	—	—	2
6.2	Fuel treatment	—	—	—	1	1	—	—	—	—	—	—	—	—	1
6.3	Wildfire preparedness and response	—	—	—	1	2	1	1	—	—	—	1	1	—	1
6.4	Social factors	—	—	—	—	1	—	—	—	—	—	—	—	—	—

Note: Abbreviations for the ML methods are defined in Table 1. Note that in some cases, a paper may use more than one ML method and (or) appear in multiple problem domains.

from inactive or nonburned areas, burned from unburned areas (for extinguished fires), or fire-severity measures such as the normalized burn ratio (Lutes et al. 2006).

In early studies using ML methods for fire mapping, Al-Rawi et al. (2001) and Al-Rawi et al. (2002) used ANNs (specifically, the supervised ART-II neural network) for burn-scar mapping and fire detection. Pu and Gong (2004) compared logistic regression (LR) with ANN for burn-scar mapping using Landsat images; both methods achieved high accuracy (>97%). Interestingly, however, the authors found that LR was more efficient for their relatively limited dataset. The authors in Zammit et al. (2006) performed burned-area mapping for two large fires that occurred in France using satellite images and three ML algorithms, including SVM, KNN, and the KM algorithm; overall SVM had the best performance. Likewise, Dragozi et al. (2011) compared the use of SVM against a nearest neighbor method for burned-area mapping in Greece and found better performance with SVM. In fact, a number of studies (Alonso-Benito et al. 2008; Cao et al. 2009; Petropoulos et al. 2010, 2011; Zhao et al. 2015; Pereira et al. 2017; Branham et al. 2017; Hamilton et al. 2017) have successfully used SVM for burn-scar mapping using satellite data. Mitrakis et al. (2012) performed burned-area mapping in the Mediterranean region using a variety of ML algorithms, including a fuzzy neuron classifier (FNC), ANN, SVM, and AdaBoost, and found that while all methods displayed similar accuracy, the FNC performed slightly better. Dragozi et al. (2014) applied SVM and a feature selection method (based on fuzzy logic) to IKONOS imagery for burned-area mapping in Greece. Another approach to burned-area mapping in the Mediterranean used an ANN and MODIS hotspot data (Gómez and Martín 2011). Pereira et al. (2017) used a one-class SVM, which requires only positive training data (i.e., burned pixels), for burn-scar mapping, which may offer a more sample-efficient approach than general SVMs; the one-class SVM approach may be useful when good wild-fire training datasets are difficult to obtain. In Mithal et al. (2018), the authors developed a three-stage framework for burned-area mapping using MODIS data and ANNs. Crowley et al. (2019) used Bayesian updating of landcover (BULC) to merge burned-area classifications from three remote sensing sources (Landsat-8, Sentinel-2, and MODIS). Celik (2010) used GA for change detection in satellite images, while Sunar and Özkan (2001) used the interactive ISODATA and ANN to map burned areas.

In addition to burned-area mapping, ML methods have been used for burn-severity mapping, including GA (Brumby et al.

2001), MaxEnt (Quintano et al. 2019), bagged DTs (Sá et al. 2003), and others. For instance, Hultquist et al. (2014) used three popular ML approaches (Gaussian process regression (GPR) (Rasmussen and Williams 2006), RF, and SVM) for burn-severity assessment in the Big Sur ecoregion, California. RF gave the best overall performance and had lower sensitivity to different combinations of variables. All ML methods, however, performed better than conventional multiple regression techniques. Likewise, Hultquist et al. (2014) compared the use of GPR, RF, and SVM for burn-severity assessment and found that RF displayed the best performance. Another recent paper by Collins et al. (2018) investigated the applicability of RF for fire-severity mapping and discussed the advantages and limitations of RF for different fire and land conditions.

One recent paper by Langford et al. (2019) used a five-layer DNN for mapping fires in Interior Alaska with a number of MODIS-derived variables (e.g., NDVI and surface reflectance). They found that a validation-loss (VL) weight selection strategy for the unbalanced dataset (i.e., the no-fire class appeared much more frequently than fire) allowed them to achieve better accuracy compared with a XGBoost method; however, without the VL approach, XGBoost outperformed the DNN, highlighting the need for methods to deal with unbalanced datasets in fire mapping.

4.2. Fire weather and climate change

4.2.1. Fire weather prediction

Fire weather is a critical factor in determining whether a fire will start, how fast it will spread, and where it will spread. Fire weather observations are commonly obtained from surface weather station networks operated by meteorological services or fire management agencies. Weather observations may be interpolated from these point locations to a grid over the domain of interest, which may include diverse topographical conditions; the interpolation task is a regression problem. Weather observations may subsequently be used in the calculation of meteorologically based fire danger indices such as the Canadian Fire Weather Index (FWI) System (Van Wagner 1987). Future fire weather conditions and danger indices are commonly forecast using the output from numerical weather prediction (NWP) models (e.g., the European Forest Fire Information System; San-Miguel-Ayanz et al. 2012); however, errors in the calculation of fire danger indices that have a memory (such as the moisture indices of the FWI System) can

accumulate in such projections. It is noteworthy that surface fire danger measures may be correlated with large-scale weather and climatic patterns.

To date, there have been relatively few papers that address fire weather and danger prediction using ML. The first effort (Crimmins 2006) used SOMs to explore the synoptic climatology of extreme fire weather in the southwestern United States (US). Crimmins found three key patterns representing southwesterly flow and large geopotential height gradients that were associated with over 80% of the extreme fire weather days as determined by a fire weather index. Nauslar et al. (2019) used SOMs to determine the timing of the North American Monsoon that plays a major role on the length of the active fire season in the southwestern US. Lagerquist et al. (2017) also used SOMs to predict extreme fire weather in northern Alberta, Canada. Fire weather was defined by using extreme values of the Fine Fuel Moisture Code (FFMC), Initial Spread Index (ISI), and the Fire Weather Index (FWI), all components of the FWI System (Van Wagner 1987). Good performance was achieved with the FFMC and the ISI, and this approach has the potential to be used in near real time, allowing input into fire management decision systems. Other efforts have used a combination of conventional and ML approaches to interpolate meteorological fire danger in Australia (Sanabria et al. 2013).

4.2.2. Lightning prediction

Lightning is the second most common cause of wildfires (behind human causes); thus, predicting the location and timing of future storms and strikes is of great importance to predicting fire occurrence. Electronic lightning detection systems have been deployed in many parts of the world for several decades and have accrued rich strike location–time datasets. Lightning prediction models have employed these data to derive regression relationships with atmospheric conditions and stability indices that can be forecast with NWP. Ensemble forecasts of lightning using RF is a viable modelling approach for Alberta, Canada (Blouin et al. 2016). Bates et al. (2017) used two ML methods (CART and RF) and three statistical methods to classify wet and dry thunderstorms (lightning associated with dry thunderstorms are more likely to start fires) in Australia.

4.2.3. Climate change

Transfer modeling, whereby a model produced for one study region and (or) distribution of environmental conditions is applied to other cases (Phillips et al. 2006), is a common approach in climate change science. Model transferability should be considered when using ML methods to estimate projected quantities due to climate change or other environmental changes. With regards to climate change, transfer modeling is essentially an extrapolation task. Previous studies in the context of species distribution modeling have indicated that ML approaches may be suitable for transfer modeling under future climate scenarios. For example, Heikkinen et al. (2012) indicated that MaxEnt and generalized boosting methods (GBM) have better transferability than either ANN and RF and that the relatively poor transferability of RF may be due to overfitting.

There are several publications on wildfires and climate change that use ML approaches. Amatulli et al. (2013) found that multivariate adaptive regression splines (MARS) were better predictors of future monthly area burned for five European countries compared with multiple linear regression and RF. Parks et al. (2016) projected fire severity for future time periods in the western US using BRT. Young et al. (2017) similarly used BRT to project future fire intervals in Alaska and found up to a fourfold increase in (30 year) fire occurrence probability by 2100. Several authors used MaxEnt to project future fire probability globally (Moritz et al. 2012), for Mediterranean ecosystems (Batllori et al. 2013), in southwestern China (S. Li et al. 2017), in the US Pacific Northwest (Davis et al. 2017), and for southcentral US (Stroh et al. 2018). An alterna-

tive approach for projecting future potential burn probability was employed by Stralberg et al. (2018), who used RF to determine future vegetation distributions as inputs to ensemble Burn-P3 simulations. Another interesting paper of note was by Boulanger et al. (2018), who built a consensus model with two different predictor datasets and five different regression methods (generalized linear models, RF, BRT, CART, and MARS) to make projections of future area burned in Canada. The consensus model can be used to quantify uncertainty in future area burned estimates. The authors noted that model uncertainty for future periods (>200%) can be higher than that of different climate models under different carbon-forcing scenarios. This highlights the need for further work in the application of ML methods for projecting future fire danger under climate change.

4.3. Fire occurrence, susceptibility, and risk

Papers in this domain include prediction of fire occurrence and area burned (at a landscape or seasonal scale), mapping of fire susceptibility (or similar definitions of risk), and analysis of landscape or environmental controls on fire.

4.3.1. Fire occurrence prediction

Predictions of the number and location of fire starts in the upcoming day(s) are important for preparedness planning, that is, the acquisition of resources, including the relocation of mobile resources and readiness for expected fire activity. The origins of fire occurrence prediction (FOP) models go back almost 100 years (Nadeem et al. 2020). FOP models typically use regression methods to relate the response variable (fire reports or hotspots) to weather, lightning, and other covariates for a geographic unit or as a spatial probability. The seminal work of Brillinger and others in developing the spatiotemporal FOP framework is reviewed in Taylor et al. (2013). The most commonly used ML method in studies predicting fire occurrence were ANNs. As early as 1996, Vega-Garcia et al. (1996) used an ANN for human-caused wildfire prediction in Alberta, Canada, correctly predicting 85% of no-fire observations and 78% of fire observations. Not long after, Alonso-Betanzos et al. (2002) and Alonso-Betanzos et al. (2003) used ANN to predict a daily fire occurrence risk index using temperature, humidity, rainfall, and fire history as part of a larger system for real-time wildfire management system in the Galicia region of Spain. Vasilakos et al. (2007) used separate ANNs for three different indices representing fire weather (Fire Weather Index, FWI), hazard (Fire Hazard Index, FHI), and risk (Fire Risk Index) to create a composite fire ignition index (FII) for estimating the probability of wildfire occurrence on the Greek island of Lesbos. Sakr et al. (2010) used meteorological variables in a SVM to create a daily fire risk index corresponding to the number of fires that could potentially occur on a particular day. Sakr et al. (2011) then compared the use of SVM and ANN for fire occurrence prediction based only on relative humidity and cumulative precipitation up to the specific day. While Sakr et al. (2011) reported low errors for the number of fires predicted by both the SVM and ANN models, ANN models outperformed SVM; however, the SVM performed better on binary classification of fire or no fire. It is important to note, however, that ANNs encompass a wide range of possible network architectures. In an Australian study, Dutta et al. (2013) compared the use of 10 different types of ANN models for estimating monthly fire occurrence from climate data and found that an Elman RNN performed the best.

After 2012, RF became the more popular method for predicting fire occurrence among the papers reviewed here. Stojanova et al. (2012) evaluated several ML methods for predicting fire outbreaks using geographical, remotely sensed, and meteorological data in Slovenia, including single classifier methods (i.e., KNN, naïve Bayes, DT (using the J48 and JRIP algorithms), LR, SVM, and BN), and ensemble methods (AdaBoost, DT with bagging, and RF). The ensemble methods DT with bagging and RF displayed the best

predictive performance, with bagging having higher precision and RF having better recall. Vecín-Arias et al. (2016) found that RF performed slightly better than LR for predicting lightning fire occurrence in the Iberian Peninsula, based on topography, vegetation, meteorology, and lightning characteristics. Similarly, Cao et al. (2017) found that a cost-sensitive RF analysis outperformed GLM and ANN models for predicting wildfire ignition susceptibility. In recent noncomparative studies, Yu et al. (2017) used RF to predict fire risk ratings in Cambodia using publicly available remotely sensed products, while Van Beusekom et al. (2018) used RF to predict fire occurrence in Puerto Rico and found precipitation to be the most important predictor. The MaxEnt method has also been used for fire occurrence prediction (De Angelis et al. 2015; Chen et al. 2015). For example, De Angelis et al. (2015) used MaxEnt to evaluate different meteorological variables and fire indices (e.g., the Canadian FWI) for daily fire risk forecasting in the mountainous Canton Ticino region of Switzerland. The authors of that study found that combinations of such variables increased predictive power for identifying daily meteorological conditions for wildfires. Dutta et al. (2016) use a two-stage ML approach (ensemble of unsupervised deep belief neural networks with conventional supervised ensemble machine learning (DBNet)) to predict bush-fire hotspot incidence on a weekly time scale. In the first unsupervised DL phase, Dutta et al. (2016) used DBNet to generate simple features from environmental and climatic surfaces. In the second supervised ensemble classification stage, features extracted from the first stage were fed as training inputs to 10 ML classifiers (i.e., conventional supervised binary tree, linear discriminant analyzer, naïve Bayes, KNN, bagging tree, AdaBoost, gentle boosting tree, random under-sampling boosting tree, subspace discriminant, and subspace KNN) to establish the best classifier for bush fire hotspot estimation. The authors found that bagging and the conventional KNN classifier were the two best classifiers, with 94.5% and 91.8% accuracy, respectively.

4.3.2. Landscape-scale burned-area prediction

The use of ML methods in studies of burned-area prediction has only occurred relatively recently compared with other wildfire domains, yet such studies have incorporated a variety of ML methods. For example, Cheng and Wang (2008) used an RNN to forecast annual average area burned in Canada, while Archibald et al. (2009) used RF to evaluate the relative importance of human and climatic drivers of burnt area in southern Africa. Arnold et al. (2014) used hard competitive learning (HCL) to identify clusters of unique pre-fire antecedent climate conditions in the interior western US, which they then used to construct fire danger models based on MaxEnt.

Mayr et al. (2018) evaluated five common statistical and ML methods for predicting burned area and fire occurrence in Namibia, including GLM, MARS, regression trees from recursive partitioning (RPART), RF, and SVMs for regression (SVR). The RF model performed best for predicting burned area and fire occurrence; however, adjusted R^2 values were slightly higher for RPART and SVR in both cases. Likewise, de Bem et al. (2018) compared the use of LR and ANN for modelling burned area in Brazil. Both LR and ANN showed similar performance; however, the ANN had better accuracy when identifying nonburned areas but displayed lower accuracy when classifying burned areas.

4.3.3. Fire-susceptibility mapping

A considerable number of references (71) used various ML algorithms to map wildfire susceptibility, corresponding to either the spatial probability or density of fire occurrence (or other measures of fire risk such as burn severity), although other terms such as fire vulnerability and risk have also been used. The general approach was to build a spatial fire-susceptibility model using either remotely sensed or agency-reported fire data with some combination of landscape, climate, structural, and anthropogenic vari-

ables as explanatory variables. In general, the various modeling approaches used either a presence-only framework (e.g., MaxEnt) or a presence-absence framework (e.g., BRT or RF).

Early attempts at fire-susceptibility mapping used CART (Amatulli et al. 2006; Amatulli and Camia 2007; Lozano et al. 2008). Amatulli and Camia (2007) compared fire density maps in central Italy using CART and MARS and found that while CART was more accurate, MARS led to a smoother density model. More recent work has used ensemble-based classifiers such as RF and BRT or ANNs (see Table S.3.3 in Supplementary material¹ for a full list). Several of these papers also compared ML and non-ML methods for fire-susceptibility mapping and, in general, found superior performance from the ML methods. Specifically, Adab (2017) mapped fire hazard in northeastern Iran and found that ANN performed better than binary logistic regression (BLR) with an area under the curve (AUC) of 87% compared with 81% for BLR. Bisquert et al. (2012) found that ANN outperformed LR for mapping fire risk in the northwestern Spain. Goldarag et al. (2016) also compared ANN and linear regression for fire-susceptibility mapping in northern Iran and found that ANN had much better accuracy (93.49%) than linear regression (65.76%). Guo et al. (2016a, 2016b) compared RF and LR for fire-susceptibility mapping in China and found that RF led to better performance. Oliveira et al. (2012) compared RF and LR for fire density mapping in Mediterranean Europe and found that RF outperformed linear regression. De Vasconcelos et al. (2001) found that ANN had better classification accuracy than LR for ignition probability maps in parts of Portugal.

Referring to Table 3 and section S.3.3 of Supplementary material,¹ a frequently used ML method for fire-susceptibility mapping was MaxEnt. In particular, Vilar et al. (2016) found that MaxEnt performed better than GLM for fire-susceptibility mapping in central Spain with respect to sensitivity (i.e., true positive rate) and commission error (i.e., false positive rate), even though the AUC was lower. Of further note, Duane et al. (2015) partitioned their fire data into topography-driven, wind-driven, and convection-driven fires in Catalonia and mapped the fire susceptibility for each fire type.

Other ML methods used for regional fire-susceptibility mapping include BNs (Bashari et al. 2016; Dlamini 2011) and novel hybrid methods such as neuro-fuzzy systems (Jaafari et al. 2019; Bui et al. 2017). Bashari et al. (2016) noted that BNs may be useful because they allow probabilities to be updated when new observations become available. SVM was also used by a number of authors as a benchmark for other ML methods (Ghorbanzadeh et al. 2019b; Gigović et al. 2019; Hong et al. 2018; Jaafari and Pourghasemi 2019; Thach et al. 2018; Rodrigues and De la Riva 2014; Sachdeva et al. 2018; Tehrany et al. 2018; Bui et al. 2017; van Breugel et al. 2016; Zhang et al. 2019), but as we discuss later, it did not perform as well as other methods with which it was being compared.

There were two applications of ML for mapping global fire susceptibility, including Moritz et al. (2012) using MaxEnt and Luo et al. (2013) using RF. Both of these papers found that at a global scale, precipitation was one of the most important predictors of fire risk.

The majority of papers considered thus far used the entire study period (typically four or more years) to map fire susceptibility, thereby neglecting the temporal aspect of fire risk; however, a few authors have considered various temporal factors to map fire susceptibility. Martín et al. (2019) included seasonality and holidays as explanatory variables for fire probability in northeastern Spain. Vacciano et al. (2018) predicted fire susceptibility separately for the winter and summer seasons. Several papers produced maps of fire susceptibility in the eastern US by month of year (Peters et al. 2013; Peters and Iverson 2017). Parisien et al. (2014) examined differences in annual fire-susceptibility maps and a 31-year climatology for the US, highlighting the role of climate variability as a driver of fire occurrence. In particular, they found that the 90th percentile of the Canadian FWI was the dominant factor for an-

nual fire risk but not for climatological fire risk. Cao et al. (2017) considered a 10-day resolution (corresponding to the available fire data) for fire risk mapping, which makes their approach similar to fire occurrence prediction.

In addition to fire-susceptibility mapping, a few papers focused on other aspects of fire risk, including mapping probability of burn-severity classes (Holden et al. 2009; Parks et al. 2018; Tracy et al. 2018). Parks et al. (2018) additionally considered the role of fuel treatments on fire probability, which has obvious implications for fire management. Additionally, Ghorbanzadeh et al. (2019a) combined fire-susceptibility maps with vulnerability and infrastructure indicators to produce a fire hazard map.

A number of papers directly compared three or more ML (and sometimes non-ML) methods for fire-susceptibility mapping. Here we highlight some of these papers that elucidate the performance and advantages–disadvantages of various ML methods. Cao et al. (2017) found that a cost-sensitive RF model outperformed a standard RF model, ANN, as well as probit and LR. Ghorbanzadeh et al. (2019a) compared ANN, SVM, and RF and found the best performance with RF. Gigović et al. (2019) compared SVM and RF for fire-susceptibility mapping in combination with Bayesian averaging to generate ensemble models. They found that the ensemble model led to marginal improvement (AUC = 0.848) over SVM (AUC = 0.834) and RF (AUC = 0.844). For mapping both wildfire ignitions and potential natural vegetation in Ethiopia, van Breugel et al. (2016) also considered ensemble models consisting of a weighted combination of ML methods (RF, SVM, BRT, MaxEnt, ANN, and CART) and non-ML methods (GLM and MARS) and concluded that the ensemble member performed best over a number of metrics. In that paper, however, RF showed the best overall performance of all methods, including the ensemble model.

Jaafari et al. (2018) compared five DT-based classifiers for wildfire-susceptibility mapping in Iran. Here, the alternating decision tree (ADT) classifier achieved the highest performance (94.3% accuracy) in both training and validation sets. Thach et al. (2018) compared SVM, RF, and an MLP neural network for forest fire danger mapping in the region of Thuan Chau in Vietnam. They found that the performances of all models were comparable, although MLP had the highest AUC values. Interestingly Pourtaghi et al. (2016) found that a generalized additive model (GAM) outperformed RF and BRT for fire-susceptibility mapping in the Golestan province in Iran. This was one of the few examples that we found in which a non-ML method outperformed ML methods. Rodrigues and De la Riva (2014) compared RF, BRT, SVM, and LR for fire-susceptibility mapping and found that RF led to the highest accuracy, as well as the most parsimonious model. Tehrany et al. (2018) compared a LogitBoost ensemble-based DT (LEDT) algorithm with SVM, RF, and kernel logistic regression (KLR) for fire-susceptibility mapping in the Lao Cai region of Vietnam and found the best performance with LEDT, closely followed by RF. Finally, of particular note, Zhang et al. (2019) compared CNN, RF, SVM, ANN, and KLR for fire-susceptibility mapping in the Yunnan province of China. This was the only application of DL that we could find for fire-susceptibility mapping. The authors found that a CNN outperformed the other algorithms, with an overall accuracy of 87.92% compared with RF (84.36%), SVM (80.04%), MLP (78.47%), and KLR (81.23%). They noted that the benefit of CNNs is that they incorporate spatial correlations so that they can learn spatial features; however, the downside is that DL models are not as easily interpreted as other ML methods (e.g., RF and BRT).

4.3.4. Landscape controls on fire

Many of the ML methods used in fire-susceptibility mapping have also been used to examine landscape controls—the relative importance of weather, vegetation, topography, and structural and anthropogenic variables, on fire activity—which may facilitate hypothesis formation and testing or model building. From Table 3, the most commonly used methods in this section were

MaxEnt, RF, BRT, and ANN. These methods all allow for the determination of variable importance (i.e., the relative influence of predictor variables in a given model on a response variable). A commonly used method to ascertain variable importance is through the use of partial-dependence plots (Hastie et al. 2009). This method works by averaging over models that exclude the predictor variable of interest, with the resulting reduction in AUC (or other performance metrics) representing the marginal effect of the variable on the response. Partial-dependence plots have the advantage of being able to be applied to a wide range of ML methods. A related method for determining variable importance, often used for RFs, is a permutation test that involves random permutation of each predictor variable (Strobl et al. 2007). Another model-dependent approach used for ANN is the use of partial derivatives (of the activation functions of hidden and output nodes) as outlined by Vasilakos et al. (2009). It should be noted that while many other methods for model interpretation and variable dependence exist, a discussion of these methods is outside the scope of this paper.

In general, the drivers of fire occurrence or area burned varied greatly by the study area considered (including the size of area) and the methods used. Consistent with other work on “top down” and “bottom up” drivers of fire activity at large scales, climate variables were often determined to be the main drivers of fire activity, whereas at smaller scales, anthropogenic or structural factors exerted a larger influence. Here we discuss some of the papers that highlight the diversity of results for different study areas and spatial scales (global, country, ecoregion, urban) but refer the reader to section S.3.4 of Supplementary material¹ for a full listing of papers in this section. Note that many of the papers listed under section S.3.4 also belong to the fire-susceptibility mapping section and have already been discussed there.

Aldersley et al. (2011) considered drivers of monthly area burned at global and regional scales using both regression trees and RF. They found that climate factors (high temperature, moderate precipitation, and dry spells) were the most important drivers at the global scale, although at the regional scale, the models exhibited higher variability due to the influence of anthropogenic factors. At a continental scale, Mansuy et al. (2019) used MaxEnt to show that climate variables were the dominant controls (over landscape and human factors) on area burned for most ecoregions for both protected areas and outside these areas, although anthropogenic factors exerted a stronger influence in some regions such as the Tropical Wet Forests ecoregion. Masrur et al. (2018) used RF to investigate controls on circumpolar Arctic fire and found that June surface temperature anomalies were the most important variable for determining the likelihood of wildfire occurrence on an annual scale. Chingono and Mbohwa (2015) used MaxEnt to model fire occurrences in southern Africa where most fires are human-caused and found that vegetation (i.e., dry mass productivity and NDVI) was the main driver of biomass burning. Curt et al. (2015) used BRT to examine drivers of fire in New Caledonia. Interestingly, they found that human factors (such as distance to villages, cities, or roads) were dominant influences for predicting fire ignitions, whereas vegetation and weather factors were most important for area burned. Curt et al. (2016) modeled fire probabilities by different fire ignition causes (lightning, intentional, accidental, professional negligence, and personal negligence) in southeastern France. They found that socioeconomic factors (e.g., housing and road density) were the dominant factors for ignitions and area burned for human-caused fires. Fernandes et al. (2016) used BRT to examine large fires in Portugal and found that high pyrodiversity (i.e., spatial structure due to fire recurrence) and low landscape fuel connectivity were important drivers of area burned. Leys et al. (2017) used RF to find the drivers that determine sedimentary charcoal counts to reconstruct grassfire history in the US Great Plains. Not surprisingly, they found that fire regime characteristics (e.g., area burned and fire frequency) were the

most important variables and concluded that charcoal records can therefore be used to reconstruct fire histories. [L.M. Li et al. \(2009\)](#) used ANNs to show that wildfire probability was strongly influenced by population density in Japan, with a peak determined by the interplay of positive and negative effects of human presence. This relationship, however, becomes more complex when weather parameters and forest cover percentage are added to the model. [Liu et al. \(2013\)](#) used BRT to study factors influencing fire size in the Great Xingan Mountains in northeastern China. Their method included a “moving window” resampling technique that allowed them to look at the relative influence of variables at different spatial scales. They showed that the most dominant factors influencing fire size were fuel and topography for small fires, but fire weather became the dominant factor for larger fires. For regions of high population density, anthropogenic or structural factors are often dominant for fire susceptibility. For example, [Molina et al. \(2019\)](#) used MaxEnt to show that distance to roads, settlements, or powerlines were the dominant factors for fire occurrence probability in the Andalusia region in southern Spain. MaxEnt has also been used for estimating spatial fire probability under different scenarios such as future projections of housing development and private land conservation ([Syphard et al. 2016](#)). One study in China using RF found that mean spring temperature was the most important variable for fire occurrence, whereas forest stock was most important for area burned ([Ying et al. 2018](#)).

Some authors examined controls on fire severity using high resolution data for a single large fire. For example, several studies used RF to examine controls on burn severity for the 2013 Rim fire in the Sierra Nevada ([Lydersen et al. 2014, 2017](#); [Kane et al. 2015](#)). At smaller spatial scales, fire weather was the most important variable for fire severity, whereas fuel treatments were most important at larger spatial scales ([Lydersen et al. 2017](#)). A similar study by [Harris and Taylor \(2017\)](#) showed that previous fire severity was an important factor influencing fire severity for the Rim fire. For the 2005 Riba de Saelices fire, [Viedma et al. \(2015\)](#) looked at factors contributing to burn severity using a BRT model and found that burning conditions (including fire weather variables) were more important than stand structure and topography. For burn severity, these papers all used the relative differenced normalized burn ratio (RdNBR) metric, derived from Landsat satellite images, which allowed spatial modeling at high resolutions (e.g., 30 m × 30 m). In addition to the more commonly used ML methods, one paper by [Wu et al. \(2015\)](#) used KNN to identify spatially homogeneous fire environment zones by clustering climate, vegetation, topography, and human activity related variables. They then used CART to examine variable importance for each of three fire environment zones in south-eastern China. For landscape controls on fire, there were few studies comparing multiple ML methods. One such study by [Nelson et al. \(2017\)](#) compared CART, BRT, and RF for classifying different fire size classes in British Columbia, Canada. For both central and periphery regions, they found the best performing model to be BRT followed by CART and RF. For example, in the central region, BRT achieved a classification accuracy of 88% compared with 82.9% and 49.6% for the CART and RF models, respectively. It is not clear from the study why RF performed poorly, although it was noted that variable importance differed appreciably among the three models.

4.4. Fire behavior prediction

In general, fire behavior includes physical processes and characteristics at a variety of scales, including combustion rate, flaming, smouldering residence time, fuel consumption, flame height, and flame depth; however, the papers in this section deal mainly with larger scale processes and characteristics such as the prediction of fire spread rates, fire growth, burned area, and fire severity, conditional on the occurrence (ignition) of one, or more, wildfires. Here, our emphasis is on prognostic applications, in contrast to the problem domain in section 4.1 (Fuels characteriza-

tion, fire detection, and mapping) in which we focused on diagnostic applications.

4.4.1. Fire spread and growth

Predicting the spread of a wildland fire is an important task for fire management agencies, particularly to aid in the deployment of suppression resources or to anticipate evacuations one or more days in advance. Thus, a large number of models have been developed using different approaches. In a series of reviews, [Sullivan \(2009a, 2009b, 2009c\)](#) described fire spread models that he classified as being of physical or quasi-physical nature or empirical or quasi-empirical nature, as well as mathematical analogues and simulation models. Many fire growth simulation models convert one-dimensional empirical or quasi-empirical spread rate models to two dimensions and then propagate a fire perimeter across a modelled landscape.

A wide range of ML methods have been applied to predict fire growth. For example, [Markuzon and Kolitz \(2009\)](#) tested several classifiers (RF, BNs, and KNN) to estimate if a fire would become large either one or two days following its observation; they found that each of the tested methods performed similarly, with RF correctly classifying large fires at a rate over 75%, albeit with a number of false positives. [Vakalis et al. \(2004\)](#) used an ANN in combination with a fuzzy logic model to estimate the rate of spread in the mountainous region of Attica in Greece. A number of papers used GAs to optimize input parameters to a physics or empirically based fire simulator to improve fire spread predictions ([Abdalhaq et al. 2005](#); [Rodriguez et al. 2008, 2009](#); [Artés et al. 2014, 2016](#); [Carrillo et al. 2016](#); [Denham et al. 2012](#); [Cencerrado et al. 2012, 2013, 2014](#); [Artés et al. 2017](#); [Denham and Laneri 2018](#)). For example, [Cencerrado et al. \(2014\)](#) developed a framework based on GAs to shorten the time needed to run deterministic fire spread simulations. They tested the framework using the FARSITE ([Finney 2004](#)) fire spread simulator with different input scenarios sampled from distributions of vegetation models, wind speed-direction, and dead-live fuel moisture content. The algorithm used a fitness function that discarded the most time-intensive simulations but did not lead to an appreciable decrease in the accuracy of the simulations. Such an approach is potentially useful for fire management where it is desirable to predict fire behavior as far in advance as possible so that the information can be enacted upon. This approach may greatly reduce overall simulation time by reducing the input parameter space as also noted by [Artés et al. \(2016\)](#) and [Denham et al. \(2012\)](#) or through parallelization of simulation runs for stochastic approaches ([Artés et al. 2017](#); [Denham and Laneri 2018](#)). A different goal was considered by [Ascoli et al. \(2015\)](#), who used a GA to optimize fuel models in southern Europe by calibrating the model with respect to rate of spread observations.

[Kozik et al. \(2013\)](#) presented a fire spread model that used a novel ANN implementation that incorporated a Kalman filter for data assimilation that could potentially be run in real time, the resulting model more closely resembling that of complex cellular automata (CA) than a traditional ANN. The same authors later implemented this model and simulated fire growth under various scenarios with different wind speeds and directions, although a direct comparison with real fire data was not possible ([Kozik et al. 2014](#)).

[Zheng et al. \(2017\)](#) simulated fire spread by integrating a CA model with an extreme learning machine (ELM; a type of feed-forward ANN). Transition rules for the CA were determined by the ELM trained with data from historical fires, as well as vegetation, topographic, and meteorological data. Likewise, [Chetehouna et al. \(2015\)](#) used ANNs to predict fire behavior, including rate of spread and flame height and angle. In contrast, [Subramanian and Crowley \(2017\)](#) formulated the problem of fire spread prediction as a Markov decision process in which they proposed solutions based on both a classic RL algorithm and a deep RL algorithm; the

authors found that the DL approach improved on the traditional approach when tested on two large fires in Alberta, Canada. The authors further developed this work to compare five widely used reinforcement learning algorithms (Subramanian and Crowley 2018) and found that the asynchronous advantage actor-critic (A3C) and Monte Carlo tree search (MCTS) algorithms achieved the best accuracy. Meanwhile, Khakzad (2019) developed a fire spread model to predict the risk of fire spread in wildland-industrial interfaces, using dynamic Bayesian networks (DBN) in combination with a deterministic fire spread model. The Canadian Fire Behavior Prediction (FBP) System, which uses meteorological and fuel conditions data as inputs, determined the fire spread probabilities from one node to another in the aforementioned DBN.

More recently, Hodges and Lattimer (2019) trained a (deep-learning) CNN to predict fire spread using environmental variables (topography and weather- and fuel-related variables). Outputs of the CNN were spatial grids corresponding to the probability that the burn map reached a pixel and the probability that the burn map did not reach a pixel. Their method achieved a mean precision of 89% and mean sensitivity of 80% with reference 6 hourly burn maps computed using the physics-based FARSITE simulator. Radke et al. (2019) also used a similar approach to predict daily fire spread for the 2016 Beaver Creek fire in Colorado.

4.4.2. Burned-area and fire-severity prediction

There are a number of papers that focus on using ML approaches to directly predict the final area burned from a wildfire. Cortez and Morais (2007) compared multiple regression and four different ML methods (DT, RF, ANN, and SVM) to predict area burned using fire and weather (i.e., temperature, precipitation, relative humidity, and wind speed) data from the Montesinho Natural Park in northeastern Portugal and found that SVM displayed the best performance. A number of publications subsequently used the data from Cortez and Morais (2007) to predict area burned using various ML methods, including ANN (Safi and Bouroumi 2013; Storer and Green 2016), GAs (Castelli et al. 2015), both ANN and SVM (Al Janabi et al. 2018), and DTs (Alberg 2015; H. Li et al. 2018). Notably, Castelli et al. (2015) found that a GA variant outperformed other ML methods including SVM. Xie and Shi (2014) used a similar set of input variables with SVM to predict burned area in the area around Guangzhou City in China. In addition to these studies, Toujani et al. (2018) used hidden Markov models (HMM) to predict burned area in northwestern Tunisia, where the spatiotemporal factors used as inputs to the model were initially clustered using SOMs. Liang et al. (2019) compared back-propagation ANNs, RNN, and LSTM ANNs to predict wildfire scale, a quantity related to area burned and fire duration, in Alberta, Canada. They found that the highest accuracy (90.9%) was achieved with LSTM.

Most recently, Xie and Peng (2019) compared a number of ML methods for estimating area burned (regression) and binary classification of fire sizes (>5 ha) in Montesinho Natural Park, Portugal. For the regression task, they found that a tuned RF algorithm performed better than a standard RF, tuned and standard gradient boosted machines, tuned and standard GLMs, and DL. For the classification problem, they found that extreme gradient boosting and DL had a higher accuracy than CART, RF, SVM, ANN, and LR.

By attempting to predict membership of burned-area size classes, a number of papers were able to recast the problem of burned-area prediction as a classification problem. For example, Yu et al. (2011) used a combination of SOMs and back-propagation ANNs to classify forest fires into size categories based on meteorological variables. This approach gave Yu et al. (2011) better accuracy (90%) when compared with a rules-based method (82%). Özbayoglu and Bozer (2012) estimated burned-area size classes using geographical and meteorological data using three different ML methods: (i) MLP; (ii) radial basis function networks (RBFN); and (iii) SVM. Overall, the best-performing method was MLP, which achieved a 65% success rate

using humidity and wind speed as predictors. Zwirgmaier et al. (2013) used a BN to predict area-burned classes using historical fire data, fire weather data, fire behavior indices, land cover, and topographic data. Shidik and Mustofa (2014) used a hybrid model (fuzzy C-means and back-propagation ANN) to estimate fire size classes using data from Cortez and Morais (2007); the hybrid model performed best, with an accuracy of 97.50% when compared with naïve Bayes (55.5%), DT (86.5%), RF (73.1%), KNN (85.5%), and SVM (90.3%). Mitsopoulos and Mallinis (2017) compared BRT, RF, and LR to predict three burned-area classes for fires in Greece. They found that RF led to the best performance of the three tested methods and that fire suppression and weather were the two most important explanatory variables. Coffield et al. (2019) compared CART, RF, ANN, KNN, and gradient boosting to predict three burned-area classes at time of ignition in Alaska. They found that a parsimonious model using CART with vapor pressure deficit (VPD) provided the best performance of the models and variables considered.

We found only one study that used ML to predict fire behavior related to fire severity, which is important in the context of fire ecology, suggesting that there are opportunities to apply ML in this domain of wildfire science. In that paper, Zald and Dunn (2018) used RF to determine that the most important predictor of fire severity was daily fire weather, followed by stand age and ownership, with less predictability given by topographic features.

4.5. Fire effects

Fire effects prediction studies have largely used regression-based approaches to relate costs, losses, or other impacts (e.g., soils, post-fire ecology, wildlife, and socioeconomic factors) to physical measures of fire severity and exposure. Importantly, this category also includes wildfire smoke and particulate modelling (but not smoke detection, which was previously discussed in section 4.1.2, Fire detection).

4.5.1. Soil erosion and deposits

Mallinis et al. (2009) modelled potential post-fire soil erosion risk following a large intensive wildfire in the Mediterranean area using CART and KM algorithms. In that paper, before wildfire, 55% of the study area was classified as having severe or heavy erosion potential compared with 90% after fire, with an overall classification accuracy of 86%. Meanwhile, Buckland et al. (2019) used ANNs to examine the relationships between sand deposition in semi-arid grasslands and wildfire occurrence, land use, and climatic conditions. The authors then predicted soil erosion levels in the future given climate change assumptions.

4.5.2. Smoke and particulate levels

Smoke emitted from wildfires can seriously lower air quality with adverse effects on the health of both humans and animals, as well as other impacts. Thus, it is not surprising that ML methods have been used to understand the dynamics of smoke from wildland fire. For example, Yao et al. (2018b) used RF to predict the minimum height of forest fire smoke using data from the CALIPSO satellite. More commonly, ML methods have also been used to estimate population exposure to fine particulate matter (e.g., PM_{2.5}: atmospheric particulate matter with diameter less than 2.5 µm), which can be useful for epidemiological studies and for informing public health actions. One such study by Yao et al. (2018a) also used RF to estimate hourly concentrations of PM_{2.5} in British Columbia, Canada. Zou et al. (2019) compared RF, BRT, and MLR to estimate regional PM_{2.5} concentrations in the Pacific Northwest and found that RF performed much better than the other algorithms. In another very broad study covering several datasets and ML methods, Reid et al. (2015) estimated spatial distributions of PM_{2.5} concentrations during the 2008 northern California wildfires. The authors of the aforementioned study used 29 predictor variables and compared 11 different statistical mod-

els, including RF, BRT, SVM, and KNN. Overall, the BRT and RF models displayed the best performance. Emissions other than particulate matter have also been modelled using ML, as [Lozhkin et al. \(2016\)](#) used an ANN to predict carbon monoxide concentrations emitted from a peat fire in Siberia, Russia. In another study, the authors used 10 different statistical and ML methods and 21 covariates (including weather, geography, land use, and atmospheric chemistry) to predict ozone exposures before and after wildfire events ([Watson et al. 2019](#)). Here, gradient boosting gave the best results with respect to both root mean square error and R^2 values, followed by RF and SVM. In a different application related to smoke, [Fuentes et al. \(2019\)](#) used ANNs to detect smoke in several different grape varieties used for wine making.

4.5.3. Post-fire regeneration, succession, and ecology

The study of post-fire regeneration is an important aspect of understanding forest and ecosystem responses and resilience to wildfire disturbances, with important ecological and economic consequences. RF, for example, has been a popular ML method for understanding the important variables driving post-fire regeneration ([João et al. 2018](#); [Vijayakumar et al. 2016](#)). Burn severity is an important metric for understanding the impacts of wildfire on vegetation and post-fire regeneration, soils, and potential successional shifts in forest composition and, as such, has been included in many ML studies in this section, including [Barrett et al. \(2011\)](#), [Cai et al. \(2013\)](#), [Cardil et al. \(2019\)](#), [Chapin et al. \(2014\)](#), [Divya and Vijayalakshmi \(2016\)](#), [Fairman et al. \(2017\)](#), [Han et al. \(2015\)](#), [Johnstone et al. \(2010\)](#); [Liu and Yang 2014](#); [Martín-Alcón and Coll 2016](#), [Sherrill and Romme \(2012\)](#), and [Thompson and Spies \(2010\)](#). For instance, [Cardil et al. \(2019\)](#) used BRT to demonstrate that remotely sensed data (i.e., the RdNBR index) can provide an acceptable assessment of fire-induced impacts (i.e., burn severity) on forest vegetation, while [Fairman et al. \(2017\)](#) used RF to identify the variables most important in explaining plot-level mortality and regeneration of *Eucalyptus pauciflora* in Victoria, Australia, affected by high-severity wildfires and subsequent re-burns. [Debouk et al. \(2013\)](#) assessed post-fire vegetation regeneration status using field measurements, a canopy height model, and Lidar (i.e., three-dimensional laser scanning) data with a simple ANN. Post-fire regeneration also has important implications for the successional trajectories of forested areas, and a few studies have examined this using ML approaches ([Barrett et al. 2011](#); [Cai et al. 2013](#); [Johnstone et al. 2010](#)). For example, [Barrett et al. \(2011\)](#) used RF to model fire severity from which they made an assessment of the area susceptible to a shift from coniferous to deciduous forest cover in the Alaskan boreal forest, while [Cai et al. \(2013\)](#) used BRT to assess the influence of environmental variables and burn severity on the composition and density of post-fire tree recruitment and thus the trajectory of succession in northeastern China. In other studies not directly related to post-fire regeneration, [Hermosilla et al. \(2015\)](#) used RF to attribute annual forest change to one of four categories, including wildfire, in Saskatchewan, Canada, while [Jung et al. \(2013\)](#) used GA and RF to estimate the basal area of post-fire residual spruce (*Picea obovate*) and fir (*Abies sibirica*) stands in central Siberia using remotely sensed data. [Magadzire et al. \(2019\)](#) used MaxEnt to demonstrate that fire return interval and species life history traits affected the distribution of plant species in South Africa. ML has also been used to examine fire effects on the hydrological cycle, as [Poon and Kinoshita \(2018\)](#) used SVM to estimate both pre- and post-wildfire evapotranspiration using remotely sensed variables.

Considering the potential impacts of wildfires on wildlife, it is perhaps surprising that relatively few of such studies have adopted ML approaches; however, ML methods have been used to predict the impacts of wildfire and other drivers on species distributions and arthropod communities. [Hradsky et al. \(2017\)](#), for example, used nonparametric BNs to describe and quantify the drivers of faunal distributions in wildfire-affected landscapes in

southeastern Australia. Similarly, [Reside et al. \(2012\)](#) used MaxEnt to model bird species distributions in response to fire regime shifts in northern Australia, which is an important aspect of conservation planning in the region. ML has also been used to look at the effects of wildfire on fauna at the community level, as [Luo et al. \(2017\)](#) used DTs, association rule mining, and AdaBoost to examine the effects of fire disturbance on spider communities in Cangshan Mountain, China.

4.5.4. Socioeconomic effects

ML methods have been little used to model socioeconomic impacts of fire to date. We found one study in which BNs were used to predict the economic impacts of wildfires in Greece from 2006–2010 due to housing losses ([Papakosta et al. 2017](#)). The authors did this by first defining a causal relationship between the participating variables and then using BNs to estimate housing damages. It is worth noting that the problem of detecting these causal relationships from data is a difficult task and remains an active area of research in AI.

4.6. Fire management

The goal of contemporary fire management is to have the appropriate amount of fire on the landscape, which may be accomplished through the management of vegetation, including prescribed burning, the management of human activities (prevention), and fire suppression. Fire management is a form of risk management that seeks to maximize fire benefits and minimize costs and losses ([Finney 2005](#)). Fire management decisions have a wide range of scales, including long-term strategic decisions about the acquisition and location of resources or the application of vegetation management in large regions, medium-term tactical decisions about the acquisition of additional resources, relocation, or release of resources during the fire season, and short-term real-time operational decisions about the deployment and utilization of resources on individual incidents. Fire preparedness and response is a supply chain with a hierarchical dependence. [Taylor \(2020\)](#) describes 20 common decision types in fire management and maps the spatiotemporal dimensions of their decision spaces.

Fire management models can be predictive, for example, the probability of initial attack success, or prescriptive as in maximizing or minimizing an objective function (e.g., optimal helicopter routing to minimize travel time in crew deployment). While advances have been made in the domain of wildfire management using ML techniques, there have been relatively few studies in this area compared with other wildfire problem domains. Thus, there appears to be great potential for ML to be applied to wildfire management problems, which may lead to novel and innovative approaches in the future.

4.6.1. Planning and policy

An important area of fire management is planning and policy, and various ML methods have been applied to address pertinent challenges. For example, [Bao et al. \(2015\)](#) used GA, which are useful for solving multi-objective optimization problems, to optimize watchtower locations for forest fire monitoring. [Bradley et al. \(2016\)](#) used RF to investigate the relationship between the protected status of forest in the western US and burn severity. Likewise, [Ruffault and Mouillot \(2015\)](#) also used BRTs to assess the impact of fire policy introduced in the 1980s on fire activity in southern France and the relationships between fire and weather, and [Penman et al. \(2011\)](#) used BNs to build a framework to simultaneously assess the relative merits of multiple management strategies in Wollemi National Park, NSW, Australia. [McGregor et al. \(2016\)](#) used MDP and a model-free Monte Carlo method to create fast-running simulations (based on the FARSITE simulator) and interactive visualizations of forest futures over 100 years based on alternate high-level suppression policies. [McGregor et al. \(2017\)](#) demonstrated ways in which a variety of ML and optimiza-

tion methods could be used to create an interactive approximate simulation tool for fire managers. The authors of the aforementioned study utilized a modified version of the FARSITE fire-spread simulator, which was augmented to run thousands of simulation trajectories while including new models of lightning-strike occurrences, fire duration, and a forest vegetation simulator. [McGregor et al. \(2017\)](#) also clearly showed how DTs could be used to analyze a hierarchy of decision thresholds for deciding whether or not to suppress a fire; their hierarchy splits on fuel levels, then intensity estimations, and finally weather predictors to arrive at a generalizable policy.

4.6.2. Fuel treatment

ML methods have also been used to model the effects of fuel treatments to mitigate wildfire risk. For example, [Penman et al. \(2014\)](#) used a BN to examine the relative risk reduction of using prescribed burns on the landscape versus within the 500 m interface zone adjacent to houses in the Sydney Basin, Australia. [Lauer et al. \(2017\)](#) used approximate dynamic programming (also known as RL) to determine the optimal timing and location of fuel treatments and timber harvest for a fire-threatened landscape in Oregon, US, with the objective of maximizing wealth through timber management. Similarly, [Arca et al. \(2015\)](#) used GA for multi-objective optimization of fuel treatments.

4.6.3. Wildfire preparedness and response

Wildfire preparedness and response issues have also been examined using ML techniques. [Costafreda Aumedes et al. \(2015\)](#) used ANNs to model the relationships among daily fire load, fire duration, fire type, fire size, and response time, as well as personnel and terrestrial and (or) aerial units deployed for individual wildfires in Spain. Most of the models in [Costafreda Aumedes et al. \(2015\)](#) highlighted the positive correlation of burned area and fire duration with the number of resources assigned to each fire, and some highlighted the negative influence of daily fire load. In another study, [Penman et al. \(2015\)](#) used BNs to assess the relative influence of preventative and suppression management strategies on the probability of house loss in the Sydney Basin, Australia. [O'Connor et al. \(2017\)](#) used BRT to develop a predictive model of fire control locations in the northern Rocky Mountains, US, based on the likelihood of final fire perimeters, while [Homchaudhuri et al. \(2010\)](#) used GAs to optimize fire-line generation. [Rodrigues et al. \(2019\)](#) modelled the probability that wildfire will escape initial attack using a RF model trained with fire location, detection time, arrival time, weather, fuel types, and available resources data. Important variables in [Rodrigues et al. \(2019\)](#) included fire weather and simultaneity of events. [Julian and Kochenderfer \(2018\)](#) used two different RL algorithms to develop a system for autonomous control of one or more aircraft to monitor active wildfires.

4.6.4. Social factors

Recently, the use of ML in fire management has grown to encompass more novel aspects of fire management, even including the investigation of criminal motives related to arson. [Delgado et al. \(2018\)](#) used BNs to characterize wildfire arsonists in Spain, thereby identifying five motivational archetypes (i.e., slight negligence, gross negligence, impulsiveness, profit, and revenge).

5. Discussion

ML methods have seen a spectacular evolution in development, accuracy, computational efficiency, and application in many fields since the 1990s. It is therefore not surprising that ML has been helpful in providing new insights into several critical sustainability and social challenges in the 21st century ([Gomes 2009](#); [Sullivan et al. 2014](#); [Butler 2017](#)). The recent uptake and success of ML methods has been driven in large part by ongoing advances in computational power and technology. For example, the recent

use of bandwidth-optimized graphics processing units (GPUs) takes advantage of parallel processing for simultaneous execution of computationally expensive tasks, which has facilitated a wider use of computationally demanding but more accurate methods such as DNNs. The advantages of powerful but efficient ML methods are therefore widely anticipated as being useful in wildfire science and management.

However, despite some early papers suggesting that data-driven techniques would be useful in forest fire management ([Latham 1987](#); [Kourtz 1990, 1993](#)), our review has shown that there was relatively slow adoption of ML-based research in wildfire science up to the 2000s compared with other fields, followed by a sharp increase in publication rate in the last decade. In the early 2000s, data-mining techniques were quite popular, and classic ML methods such as DTs, RF, and bagging and boosting techniques began to appear in the wildfire science literature (e.g., [Stojanova et al. 2006](#)). In fact, some researchers started using simple feed-forward ANNs for small-scale applications as early as the mid-1990s and early 2000s (e.g., [McCormick et al. \(1999\)](#); [Al-Rawi et al. \(2002\)](#)). In the last three decades, almost all major ML methods have been used in some way in wildfire applications, although some more computationally demanding methods such as SOMs and cellular automata have only been actively experimented with in the last decade ([Toujani et al. 2018](#); [Zheng et al. 2017](#)). Furthermore, the recent development of DL algorithms, with a particular focus on extracting spatial features from images, has led to a sharp rise in the application of DL for wildfire applications in the last decade. It is evident, however, from our review that while an increasing number of ML methodologies have been used across a variety of fire research domains over the past 30 years, this research is unevenly distributed among ML algorithms, research domains, and tasks and has had limited application in fire management.

Many fire science and management questions can be framed within a fire risk context. [Xi et al. \(2019\)](#) discussed the advantages of adopting a risk framework with regard to statistical modeling of wildfires. There the risk components of hazard, vulnerability, and exposure are replaced by fire probability, fire behavior, and fire effects, respectively. Most fire management activities can be framed as risk controls to mitigate these components of risk. Traditionally, methods used in wildfire fire science to address these various questions have included physical modeling (e.g., [Sullivan 2009a, 2009b, 2009c](#)), statistical methods (e.g., [Taylor et al. 2013](#); [Xi et al. 2019](#)), simulation modeling (e.g., [Keane et al. 2004](#)), and operations research methods (e.g., [Martell 2015](#); [Minas et al. 2012](#)).

In simple terms, any analytical study begins with one or more of four questions: what happened?; why did it happen?; what will happen?; or what to do? Corresponding data-driven approaches to address these questions are called descriptive, diagnostic, predictive, and prescriptive analytics, respectively. The type of analytical approach adopted then circumscribes the types of methodological approaches (e.g., regression, classification, clustering, dimensionality reduction, decision-making) and sets of possible algorithms appropriate to the analysis.

In our review, we found that studies incorporating ML methods in wildland fire science were predominantly associated with descriptive or diagnostic analytics, reflecting the large body of work on fire detection and mapping using classification methods and on fire-susceptibility mapping and landscape controls on fire using regression approaches. In many cases, the ML methods identified in our review are an alternative to statistical methods used for clustering and regression. While the aforementioned tasks are undoubtedly very important for understanding wildland fire, we found much less work associated with predictive or prescriptive analytics such as fire occurrence prediction (predictive), fire behavior prediction (predictive), and fire management (prescriptive). This may be because (i) specific domain knowledge is required to frame fire management problems, (ii) fire manage-

ment data are often not publicly available, need a lot of work to transform into an easily analyzable form, or do not exist at the scale of the problem, and (iii) some fire management problems are not suited or cannot be fully addressed by ML approaches. We note that much of the work on fire risk in the fire-susceptibility and mapping domain used historical fire and environmental data to map fire susceptibility; therefore, while that work aims to inform future fire risk, it cannot be considered to be predictive analytics, except, for example, when it was used in combination with climate change projections. It appears then that, in general, wildfire science research is currently more closely aligned with descriptive and diagnostic analytics, whereas wildfire management goals are aligned with predictive and prescriptive analytics. This fundamental difference identifies new opportunities for research in fire management, which we discuss later in this paper.

In the remainder of the paper, we examine some considerations for the use of ML methods, including data considerations, model selection and accuracy, implementation challenges, interpretation, opportunities, and implications for fire management.

5.1. Data considerations

ML is a data-centric modeling paradigm concerned with finding patterns in data. Importantly, data scientists need to determine, often in collaboration with fire managers or domain experts, whether there are suitable and sufficient data for a given modeling task. Some of the criteria for suitable data include whether (i) the predictands and covariates are or can be wrangled into the same temporal and spatial scale, (ii) the observations are a representative sample of the full range of conditions that may occur in application of a model to future observations, and (iii) whether the data are at a spatiotemporal scale appropriate to the fire science or management question. The first of these criteria can be relaxed in some ML models such as ANNs and DNNs in which inputs and outputs can be at different spatial or temporal scales for appropriately designed network architectures, although data normalization may still be required. The second criterion also addresses the important question of whether enough data exists for training a given algorithm for a given problem. In general, this question depends on the nature of the problem, complexity of the underlying model, data uncertainty, and many other factors (for a further discussion of data requirements for ML, see Roh et al. 2018). In any case, many complex problems require a substantive data wrangling effort to acquire, perform quality assurance on, and fuse data into sampling units at the appropriate spatiotemporal scale. An example of this is in daily fire occurrence prediction in which observations of a variety of features (e.g., continuous measures such as fire arrival time and location or lightning strike times and locations) are discretized into three-dimensional (e.g., longitude, latitude, and day) cells called voxels. Another important consideration for the collection and use of data in ML is selection bias. A form of spatial selection bias called preferential sampling occurs when sampling occurs preferentially in locations where one expects a certain response (Diggle et al. 2010). For example, preferential sampling may occur in air monitoring, because sensors may be placed in locations where poor air quality is expected (Shaddick and Zidek 2014). In general, preferential sampling or other selection biases may be avoided altogether by selecting an appropriate sampling strategy at the experimental design phase or, when this is not possible, by taking it into account in model evaluation (Zadrozny 2004).

For the problem domain fire detection and mapping, most applications of ML used some form of imagery (e.g., remotely sensed satellite images or terrestrial photographs). In particular, many papers used satellite data (e.g., Landsat, MODIS) to determine vegetation differences before and after a fire and so were able to map area burned. For fire detection, many applications considered either remotely sensed data for hotspot or smoke detection or photographs of wildfires (used as inputs to an image classification

problem). For fire weather and climate change, the three main sources of data were weather station observations, climate re-analyses (modelled data that include historical observations), or GCMs for future climate projections. Re-analyses and GCMs are typically highly dimensional large gridded spatiotemporal datasets that require careful feature selection and (or) dimensional reduction for ML applications. Fire occurrence prediction, susceptibility, and risk applications use a large number of different environmental variables as predictors, but almost all use fire locations and associated temporal information as predictands. Fire data itself is usually collated from fire management agencies in the form of georeferenced points or perimeter data, along with reported dates, ignition cause, and other related variables. Care should be taken using such data because changes in reporting standards or accuracy may lead to data inhomogeneity. As well as fire locations and perimeters, fire severity is an attribute of much interest to fire scientists. Fire severity is often determined from remotely sensed data and represented using variables such as the differenced normalized burn ratio (dNBR) and variants or through field sampling; however, remotely sensed estimates of burn severity should be considered as proxies as they have low skill in some ecosystems. Other fire ecology research historically relies on in situ field sampling, although many of the ML applications attempt to resolve features of interest using remotely sensed data. Smoke data can also be derived from remotely sensed imagery or from air quality sensors (e.g., PM_{2.5}).

Continued advances in remote sensing, as well as the quality and availability of remotely sensed data products, in weather and climate modeling have led to increased availability of large spatiotemporal datasets, which presents both an opportunity and challenge for the application of ML methods in wildfire research and management. The era of “big data” has seen the development of cloud computing platforms to provide the computing and data storage facilities to deal with these large datasets. For example, in our review, we found two papers (Crowley et al. 2019; Quintero et al. 2019) that used Google Earth Engine, which integrates geospatial datasets with a coding environment (Gorelick et al. 2017). In any case, data processing and management plays an important role in the use of large geospatial datasets.

5.2. Model selection and accuracy

Given a wildfire science question or management problem and available relevant data, a critical question to ask is what is the most appropriate modeling tool to address the problem? Is it a standard statistical model (e.g., linear regression or LR), a physical model (e.g., FIRETEC or other fire simulator), a ML model, or a combination of approaches? Moreover, which specific algorithm will yield the most accurate classification or regression? Given the heterogeneity of research questions, study areas, and datasets considered in the papers reviewed here, it is not possible to comprehensively answer these questions with respect to ML approaches. Even when multiple studies used the same dataset (Cortez and Morais 2007; Safi and Bouroumi 2013; Storer and Green 2016; Castelli et al. 2015; Al Janabi et al. 2018; Alberg 2015; H. Li et al. 2018), the different research questions considered meant that a direct comparison of ML methods was not possible between research studies. However, a number of individual studies did make comparisons between multiple ML methods or between ML and statistical methods for a given wildfire modeling problem and dataset. Here we highlight some of their findings to provide some guidance with respect to model selection. In our review, we found 29 papers comparing ML and statistical methods, and in the majority of these cases, ML methods were found to be more accurate than traditional statistical methods (e.g., GLMs) or displayed similar performance (Pu and Gong 2004; Bates et al. 2017; de Bem et al. 2018). In only one study on climate change by Amatulli et al. (2013), MARS was found to be superior to RF for their analytical task. A sizable number of the comparative studies

(14) involved classification problems that used LR as a benchmark method against ANN or ensemble tree methods. For studies comparing multiple ML methods, there was considerable variation in the choice of most accurate method; however, in general, ensemble methods tended to outperform single classifier methods (e.g., Stojanova et al. 2012; Dutta et al. 2016; Mayr et al. 2018; Nelson et al. 2017; Reid et al. 2015; Watson et al. 2019), except in one case in which the most accurate model (CART) was also the most parsimonious (Coffield et al. 2019). A few more recent papers also highlighted the advantages of DL over other methods. In particular, for fire detection, Zhang et al. (2018) compared CNNs with SVM and found that CNNs were more accurate, while Zhao et al. (2018) similarly found CNNs superior to SVMs and ANNs. For fire-susceptibility mapping, Zhang et al. (2019) found CNNs to be more accurate than RF, SVMs, and ANNs. For time series forecasting problems, Liang et al. (2019) found that LSTMs outperformed ANNs. Finally, Cao et al. (2019) found that using an LSTM combined with a CNN led to better fire detection performance from video compared with CNNs alone.

In any case, more rigorous intermodel comparisons are needed to reveal under which conditions and in what sense are particular methods more accurate, as well as to establish procedures for evaluating accuracy. ML methods are also prone to overfitting, so it is important to evaluate models with robust test datasets using appropriate cross-validation strategies. For example, the naïve application of cross-validation to data that have spatial or spatiotemporal dependencies may lead to overly optimistic evaluations (Roberts et al. 2017). In general, one also desires to minimize errors associated with either under- or over-specification of the model, a problem known as the bias-variance trade-off (Geman et al. 1992); however, several recent advances have been made to reduce overfitting in ML models, e.g., regularization techniques in DNNs (Kukačka et al. 2017). Moreover, when interpreting comparisons between ML and statistical methods, we should be cognizant that just as some ML methods require expert knowledge, the accuracy of statistical methods can also vary with the skill of the practitioner. Thompson and Calkin (2011) also emphasize the need for identifying sources of uncertainty in modeling so that they can be better managed.

5.3. Implementation challenges

Beyond data and model selection, two important considerations for model specification are feature selection and spatial autocorrelation. Knowledge of the problem domain is extremely important in identifying a set of candidate features. However, while many ML methods are not limited by the number of features, more variables do not necessarily make for a more accurate, interpretable, or easily implemented model (Schoenberg 2016; Breiman 2001) and can lead to overfitting and increased computational time. Two different ML methods to enable selection of a reduced and more optimal set of features include GAs and particle swarm optimization (PSO). Sachdeva et al. (2018) used a GA to select input features for BRT and found that this method gave the best accuracy compared with ANN, RF, SVM, SVM with PSO (PSO-SVM), DTs, LR, and NB. Hong et al. (2018) employed a similar approach for fire-susceptibility mapping and found that this led to improvements for both SVM and RF compared with their non-optimized counterparts. Tracy et al. (2018) used a novel random subset feature selection algorithm for feature selection, which they found led to higher AUC values and lower model complexity. Jaafari et al. (2019) used a NFM combined with the imperialist competitive algorithm (a variant of GA) for feature selection, which led to very high model accuracy (0.99) in their study. Bui et al. (2017) used PSO to choose inputs to a neural fuzzy model and found that this improved results. Zhang et al. (2019) also considered the information gain ratio for feature selection. As noted in Moritz et al. (2012) and Mayr et al. (2018), one should also take spatial autocorrelation into account when modeling fire probabil-

ities spatially. In general, the presence of spatial autocorrelation violates the assumption of independence for parametric models, which can degrade model performance. One approach to deal with autocorrelation requires subsampling to remove any spatial autocorrelation (Moritz et al. 2012). It is also often necessary to subsample from non-fire locations due to class imbalance between ignitions and non-ignitions (e.g., Cao et al. 2017; Zhang et al. 2019). Song et al. (2017) considered spatial econometric models and found that a spatial autocorrelation model worked better than RF, although Kim et al. (2019) noted that RF may be robust to spatial autocorrelation with large samples. In contrast to many ML methods, a strength of CNN is its ability to exploit spatial correlation in the data to enable the extraction of spatial features.

5.4. Interpretation

A major obstacle for the adoption of ML methods to fire modeling tasks is the perceived lack of interpretability or explainability of such methods, which are often considered to be “black box” models. Users (in this case, fire fighters and managers) need to trust ML model predictions and so have the confidence and justification to apply these models, particularly when proposed solutions are considered novel. Model interpretability should therefore be an important aspect of model development if models are to be selected and deployed in fire management operations. Model interpretability varies significantly across the different types of ML. For example, conventional thinking is that tree-based methods are more interpretable than neural network methods. This is because a single DT classifier can be rendered as a flow chart corresponding to if-then-else statements, whereas an ANN represents a nonlinear function approximated through a series of nonlinear activations. However, because they combine multiple trees in an optimized way, ensemble tree classifiers are less interpretable than single tree classifiers. On the other hand, BNs are one example of an ML technique in which good explanations for results can be inferred due to their graphical representation; however, full Bayesian learning on large-scale data is very computationally expensive, which may have limited early applications; however, as computational power has increased, we have seen an increase in the popularity of BNs in wildfire science and management applications (e.g., Penman et al. 2015; Papakosta et al. 2017).

DL-based architectures are widely considered to be among the least interpretable ML models, despite that they can achieve very accurate function approximation (Chakraborty et al. 2017). In fact, this is demonstrative of the well-known trade-off between prediction accuracy and interpretability (for an in-depth discussion, see Kuhn and Johnson 2013). The ML community, however, recognizes the problem of interpretability and work is underway to develop procedures that allow for greater interpretability of ML methods, including for DL (see, for example, McGovern et al. 2019) or model-agnostic approaches (Ribeiro et al. 2016). Runge et al. (2019) further argue that casual inference methods should be used in conjunction with predictive models to improve our understanding of physical systems. Finally, it is worth noting that assessing variable importance (see section 4.3.4) for a given model can play a role in model interpretation.

5.5. Opportunities

Our review highlights a number of potential opportunities in wildfire science and management for ML applications where ML has not yet been applied or is underutilized. Here we examine ML advances in other areas of environmental science that have analogous problems in wildland fire science and that may be useful for identifying further ML applications. For instance, J. Li et al. (2011) compared ML algorithms for spatial interpolation and found that a RF model combined with geostatistical methods yielded good results; a similar method could be used to improve interpolation of fire weather observations from weather stations and so enhance fire danger monitoring. Rasp and Lerch (2018) showed that

ANNs could improve weather forecasts by post-processing ensemble forecasts, an approach that could similarly be applied to improve short-term forecasts of fire weather. Belayneh et al. (2014) used ANNs and SVMs combined with wavelet transforms for long-term drought forecasting in Ethiopia; such methods could also be useful for forecasting drought in the context of fire danger potential. In the context of numerical weather prediction, Cohen et al. (2019) found better predictability using ML methods than dynamical models for subseasonal to seasonal weather forecasting, suggesting similar applications for long-term fire weather forecasting. McGovern et al. (2017) discussed how AI techniques can be leveraged to improve decision-making around high-impact weather. More recently, Reichstein et al. (2019) have further argued for the use of DL in the environmental sciences, citing its potential to extract spatiotemporal features from large geospatial datasets. Kussul et al. (2017) used CNNs to classify land cover and crop types and found that CNNs improved the results over standard ANN models; a similar approach could be used for fuels classification, which is an important input to fire behavior prediction models. Shi et al. (2016) also used CNNs to detect clouds in remotely sensed imagery and were able to differentiate between thin and thick clouds. A similar approach could be used for smoke detection, which is important for fire detection, as well as in determining the presence of false negatives in hotspot data (due to smoke or cloud obscuration). Finally, recent proposals have called for hybrid models that combine process-based models and ML methods (Reichstein et al. 2019). For example, ML models may replace user-specified parameterizations in numerical weather prediction models (Brenowitz and Bretherton 2018). Other recent approaches use ML methods to determine the solutions to nonlinear partial differential equations (Raissi and Karniadakis 2018; Raissi et al. 2019). Such methods could find future applications in improving fire behavior prediction models based on computationally expensive physics-based fire simulators in coupled fire-atmosphere models or in smoke dispersion modeling. In any case, the applications of ML that we have outlined are meant for illustrative purposes and are not meant to represent an exhaustive list of all possible applications.

5.6. Implications for fire management

We believe that ML has been underutilized in fire management, particularly with respect to problems belonging to either predictive or prescriptive analytics. Fire management comprises a set of risk control measures that are often cast in the framework of the emergency response phases: prevention, mitigation, preparedness, response, recovery, and review (Tymstra et al. 2019). In terms of financial expenditure, by far the largest percentage is spent in the response phase (Stocks and Martell 2016). In practice, fire management is largely determined by the need to manage resources in response to active or expected wildfires, typically for lead times of days to weeks, or to manage vegetative fuels. This suggests the opportunity for increased research in areas of fire weather prediction, fire occurrence prediction, and fire behavior prediction, as well as optimizing fire operations and fuel treatments. The identification of these areas, as well as the fact that wildfire is both a spatial and temporal process, further reiterate the need for ML applications for time series forecasting.

In this review, there were few papers that used time series ML methods for forecasting problems, suggesting an opportunity for further work in this area. In particular, RNNs were used for fire behavior prediction (Cheng and Wang 2008; Kozik et al. 2013, 2014) and fire occurrence prediction (Dutta et al. 2013). The most common variants of RNNs are LSTM networks (Hochreiter and Schmidhuber 1997), which have been used for burned-area prediction (Liang et al. 2019) and fire detection (Cao et al. 2019). Because these methods implicitly model dynamical processes, they should lead to improved forecasting models compared with standard ANNs. For example, Gensler et al. (2017) used LSTMs to forecast

solar power and Kim et al. (2017) used CNNs combined with LSTM for forecasting precipitation. We anticipate that these methods could also be employed for fire weather, fire occurrence, and fire behavior prediction.

We note that there are a number of operational research and management science methods used in fire management research, including queuing, optimization, and simulation of complex system dynamics (e.g., Martell 2015), in which ML algorithms do not seem to provide an obvious alternative, e.g., planning models to simulate the interactions between fire management resource configurations and fire dynamics reviewed by Mavsar et al. (2013). From our review, a few papers used agent-based learning methods for fire management. In particular, RL was used for optimizing fuel treatments (Lauer et al. 2017) or for autonomous control of aircraft for fire monitoring (Julian and Kochenderfer 2018). GAs were used for generating optimal fire lines for active fires (Homchaudhuri et al. 2010) and for reducing the time for fire simulation (Cencerrado et al. 2014). However, more work is needed to identify where ML methods could contribute to tactical, operational, or strategic fire management decision-making.

An important challenge for the fire research and management communities is enabling the transition of potentially useful ML models to fire management operations. Although we have identified several papers emphasizing that their ML models could be deployed in fire management operations (Artés et al. 2016; Alonso-Betanzos et al. 2002; Iliadis 2005; Stojanova et al. 2012; Davis et al. 1989, 1986; Liu et al. 2015), it can be difficult to assess whether and how a study has been adopted by, or influenced by, fire management agencies. This challenge is often exacerbated by a lack of resources and (or) funding, as well as the different priorities and institutional cultures of researchers and fire managers. One possible solution to this problem would be the formation of working groups dedicated to enabling this transition, preferably at the research proposal phase. In general, enabling operational ML methods will require tighter integration and greater collaboration between the research and management communities, particularly with regards to project design, data compilation and variable selection, implementation, and interpretation. It is worth noting, however, that this is not a problem unique to ML; it is a long-standing and common issue in many areas of fire research and other applied science disciplines in which continuous effort is required to maintain communications and relationships between researchers and practitioners.

Finally, we would like to stress that we believe that the wildfire research and management communities should play an active role in providing relevant, high-quality, and freely available wildfire data for use by practitioners of ML methods. For example, burned area and fire weather data made available by Cortez and Morais (2007) was subsequently used by a number of authors in their work. It is imperative that the quality of data collected by management agencies be as robust as possible, as the results of any modelling process are dependent on the data used for analysis. It is worth considering how new data on, for example, hourly fire growth or the daily use of fire management resources could be used in ML methods to yield better predictions or management recommendations — using new tools to answer new questions may require better or more complete data. Conversely, we must recognize that despite ML models being able to learn on their own, expertise in wildfire science is necessary to ensure realistic modelling of wildfire processes, while the complexity of some ML methods (e.g., DL) requires a dedicated and sophisticated knowledge of their application (we note that many of the most popular ML methods used in this study such as RF, MaxEnt, and DTs are fairly easy to implement). The observation that no single ML algorithm is superior for all classes of problem, an idea encapsulated by the “no free lunch” theorem (Wolpert 1996), further reinforces the need for domain-specific knowledge. Thus, the proper implementation of ML in wildfire science is a challenging endeavor,

often requiring multidisciplinary teams and (or) interdisciplinary specialists to effectively produce meaningful results.

5.7. A word of caution

ML holds tremendous potential for a number of wildfire science and management problem domains. As indicated in this review, much work has already been undertaken in a number of areas, although further work is clearly needed for fire management specific problems. Despite this potential, ML should not be considered a panacea for all fire research areas. ML is best suited to problems in which there are sufficient high-quality data, and this is not always the case. For example, for problems related to fire management policy, data are needed at large spatiotemporal scales (i.e., ecosystem-administrative spatial units at time scales of decades or even centuries), and such data may simply not yet exist in current inventories. At the other extreme, data are needed at very fine spatiotemporal scales for fire spread and behavior modeling, including high-resolution fuel maps and surface weather variables, which are often not available at the required scale and are difficult to acquire even in an experimental context. Another limitation of ML may occur when one attempts to make predictions where no analog exists in the observed data, such as may be the case with climate change prediction.

6. Conclusions

Our review shows that the application of ML methods in wildfire science and management has been steadily increasing since their first use in the 1990s, across core problem domains using a wide range of ML methods. The bulk of work undertaken thus far has used traditional methods such as RF, BRT, MaxEnt, SVM, and ANNs, partly due to the ease of application and partly due to their simple interpretability in many cases; however, problem domains associated with predictive (e.g., predicted fire behavior) or prescriptive (e.g., optimizing fire management decisions) analytics have seen much less work with ML methods. We therefore suggest that opportunities exist for both the wildfire community and ML practitioners to apply ML methods in these areas. Moreover, the increasing availability of large spatiotemporal datasets from, for example, climate models or remote sensing may be amenable to the use of DL methods, which can efficiently extract spatial or temporal features from data. Another major opportunity is the application of agent-based learning to fire management operations, although many other opportunities exist. We must recognize, however, that despite ML models being able to learn on their own, expertise in wildfire science is necessary to ensure realistic modelling of wildfire processes across multiple scales, while the complexity of some ML methods (e.g., DL) requires a dedicated and sophisticated knowledge of their application. Furthermore, a major obstacle for the adoption of ML methods to fire modeling tasks is the perceived lack of interpretability of such methods, which are often considered to be black box models. The ML community, however, recognizes this problem and work is underway to develop methods that allow for greater interpretability of ML methods (see, for example, McGovern et al. 2019). Data-driven approaches are by definition data-dependent — if the fire management community wants to more fully exploit powerful ML methods, we need to consider data as a valuable resource and examine what further information on fire events or operations is needed to apply ML approaches to management problems. In closing, wildland fire science is a diverse multifaceted discipline that requires a multipronged approach, a challenge made greater by the need to mitigate and adapt to current and future fire regimes.

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