review articles

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**Tapping into the “folk knowledge” needed to advance machine learning applications.**

**by Pedro domingos**

**a few useful things to Know about machine Learning**

## ***Machine learning systeMs*** automatically learn programs from data. This is often a very attractive alternative to manually constructing them, and in the last decade the use of machine learning has spread rapidly throughout computer science and beyond.

Machine learning is used in Web search, spam filters, recommender systems, ad placement, credit scoring, fraud detection, stock trading, drug design, and many other applications. A recent report from the McKinsey Global Institute asserts that machine learning (a.k.a. data mining or predictive analytics) will be the driver of the next big wave of innovation.15 Several fine textbooks are available to interested practitioners and researchers (for example, Mitchell16 and Witten et al.24). However, much of the “folk knowledge” that

is needed to successfully develop machine learning applications is not readily available in them. As a result, many machine learning projects take much longer than necessary or wind up producing less-than-ideal results. Yet much of this folk knowledge is fairly easy to communicate. This is the purpose of this article.

**key insights**

**machine learning algorithms can figure out how to perform important tasks**

**by generalizing from examples. this is often feasible and cost-effective where manual programming is not. as more data becomes available, more ambitious problems can be tackled.**

**machine learning is widely used in computer science and other fields. however, developing successful machine learning applications requires a substantial amount of “black art” that is difficult to find in textbooks.**

**this article summarizes 12 key lessons that machine learning researchers and practitioners have learned. these include pitfalls to avoid, important issues to focus on, and answers to common questions.**



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Many different types of machine learning exist, but for illustration purposes I will focus on the most mature and widely used one: clas- sification. Nevertheless, the issues I will discuss apply across all of ma- chine learning. A *classifier* is a sys- tem that inputs (typically) a vector of discrete and/or continuous *fea- ture values* and outputs a single dis- crete value, the *class*. For example, a spam filter classifies email mes- sages into “spam” or “not spam,” and its input may be a Boolean vec- tor **x** = (*x*1,…,*xj*,…,*xd*), where *xj* = 1 if the *j*th word in the dictionary appears in the email and *xj* = 0 otherwise. A *learner* inputs a training set of ex- amples (**x***i*, *yi*), where **x***i* = (*xi*,*1* , . . . , *xi*,*d*) is an observed input and *yi* is the corresponding output, and outputs a classifier. The test of the learner is whether this classifier produces the correct output *yt* for future examples **x***t* (for example, whether the spam filter correctly classifies previously unseen email messages as spam or not spam).

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### Learning = Representation + Evaluation + optimization

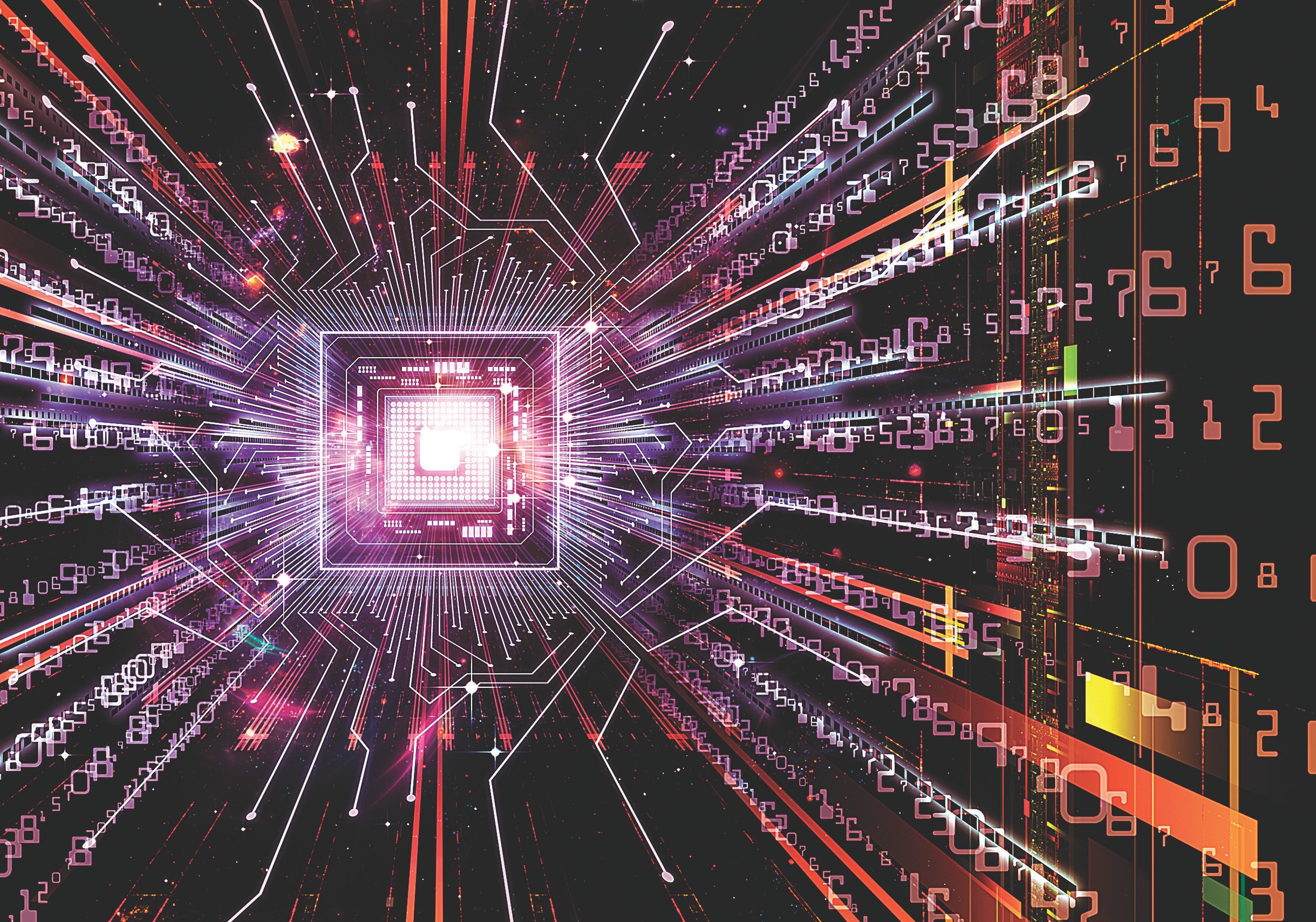
Suppose you have an application that you think machine learning might be good for. The first problem facing you is the bewildering variety of learning al- gorithms available. Which one to use? There are literally thousands available, and hundreds more are published each year. The key to not getting lost in this huge space is to realize that it consists of combinations of just three compo- nents. The components are:

* **Representation.** A classifier must be represented in some formal lan- guage that the computer can handle. Conversely, choosing a representa- tion for a learner is tantamount to choosing the set of classifiers that it can possibly learn. This set is called the *hypothesis space* of the learner. If a classifier is not in the hypothesis space, it cannot be learned. A related question, that I address later, is how to represent the input, in other words, what features to use.
* **Evaluation.** An evaluation func- tion (also called *objective function*

or *scoring function*) is needed to dis- tinguish good classifiers from bad ones. The evaluation function used internally by the algorithm may dif- fer from the external one that we want the classifier to optimize, for ease of optimization and due to the issues I will discuss.

* **Optimization.** Finally, we need a method to search among the clas- sifiers in the language for the high- est-scoring one. The choice of op- timization technique is key to the efficiency of the learner, and also helps determine the classifier pro- duced if the evaluation function has more than one optimum. It is com- mon for new learners to start out using off-the-shelf optimizers, which are lat- er replaced by custom-designed ones.

The accompanying table shows common examples of each of these three components. For example, *k*- nearest neighbor classifies a test ex- ample by finding the *k* most similar training examples and predicting the majority class among them. Hyper- plane-based methods form a linear



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combination of the features per class and predict the class with the high- est-valued combination. Decision trees test one feature at each internal node, with one branch for each fea- ture value, and have class predictions at the leaves. Algorithm 1 (above) shows a bare-bones decision tree learner for Boolean domains, using information gain and greedy search.20 InfoGain(*xj*, *y*) is the mutual informa- tion between feature *xj* and the class *y*. MakeNode(*x*,*c*0,*c*1) returns a node that tests feature *x* and has *c*0 as the child for *x* = 0 and *c*1 as the child for *x* = 1.

Of course, not all combinations of one component from each column of the table make equal sense. For exam- ple, discrete representations naturally go with combinatorial optimization, and continuous ones with continu- ous optimization. Nevertheless, many learners have both discrete and con- tinuous components, and in fact the

day may not be far when every single possible combination has appeared in some learner!

Most textbooks are organized by representation, and it is easy to over- look the fact that the other compo- nents are equally important. There is no simple recipe for choosing each component, but I will touch on some of the key issues here. As we will see, some choices in a machine learning project may be even more important than the choice of learner.

### it’s Generalization that counts

The fundamental goal of machine learning is to generalize beyond the examples in the training set. This is because, no matter how much data we have, it is very unlikely that we will see those exact examples again at test time. (Notice that, if there are 100,000 words in the dictionary, the spam fil- ter described above has 2100,000 pos-

sible different inputs.) Doing well on the training set is easy (just memorize the examples). The most common mistake among machine learning be- ginners is to test on the training data and have the illusion of success. If the chosen classifier is then tested on new data, it is often no better than ran- dom guessing. So, if you hire someone to build a classifier, be sure to keep some of the data to yourself and test the classifier they give you on it. Con- versely, if you have been hired to build a classifier, set some of the data aside from the beginning, and only use it to test your chosen classifier at the very end, followed by learning your final classifier on the whole data.

Contamination of your classifier by test data can occur in insidious ways, for example, if you use test data to tune parameters and do a lot of tun- ing. (Machine learning algorithms have lots of knobs, and success of- ten comes from twiddling them a lot, so this is a real concern.) Of course, holding out data reduces the amount available for training. This can be mit- igated by doing cross-validation: ran- domly dividing your training data into (say) 10 subsets, holding out each one while training on the rest, testing each learned classifier on the examples it did not see, and averaging the results to see how well the particular param- eter setting does.

**LearnDt** (*TrainSet*)

**if** all examples in *TrainSet* have the same class *y*\* **then return** Makeleaf(*y*\*)

**if** no feature *xj* has InfoGain(*xj* ,*y*) > 0 **then** *y*\* ← Most frequent class in *TrainSet* **return** Makeleaf(*y*\*)

*x*\* ← argmax*xj* InfoGain(*xj*, *y*)

*TS*0 ← examples in *TrainSet* with *x*\* = 0

*TS*1 ← examples in *TrainSet* with *x*\* = 1

**return** Makenode(*x*\*, learnDt(*TS*0), learnDt(*TS*1))

**algorithm 1. Decision tree induction.**

|  |  |  |
| --- | --- | --- |
| **table 1. the three components of learning algorithms.** | | |
| **Representation** | **evaluation** | **optimization** |
| Instances | Accuracy/error rate | combinatorial optimization |
| K-nearest neighbor | Precision and recall | Greedy search |
| Support vector machines | Squared error | beam search |
| Hyperplanes | likelihood | branch-and-bound |
| naive bayes | Posterior probability | continuous optimization |
| logistic regression | Information gain | Unconstrained |
| Decision trees | K-l divergence | Gradient descent |
| Sets of rules | cost/Utility | conjugate gradient |
| Propositional rules | Margin | Quasi-newton methods |
| logic programs |  | constrained |
| neural networks |  | linear programming |
| Graphical models |  | Quadratic programming |
| bayesian networks |  |  |
| conditional random fields |  |  |

In the early days of machine learn- ing, the need to keep training and test data separate was not widely appreci- ated. This was partly because, if the learner has a very limited representa- tion (for example, hyperplanes), the difference between training and test error may not be large. But with very flexible classifiers (for example, deci- sion trees), or even with linear classifi- ers with a lot of features, strict separa- tion is mandatory.

Notice that generalization being the goal has an interesting conse- quence for machine learning. Unlike in most other optimization problems, we do not have access to the function we want to optimize! We have to use training error as a surrogate for test error, and this is fraught with dan- ger. (How to deal with it is addressed later.) On the positive side, since the objective function is only a proxy for the true goal, we may not need to fully

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optimize it; in fact, a local optimum returned by simple greedy search may be better than the global optimum.

**Data alone is not enough** Generalization being the goal has an- other major consequence: Data alone is not enough, no matter how much of it you have. Consider learning a Boolean function of (say) 100 vari- ables from a million examples. There are 2100 − 106 examples whose classes you do not know. How do you figure out what those classes are? In the ab- sence of further information, there is just no way to do this that beats flip- ping a coin. This observation was first made (in somewhat different form) by the philosopher David Hume over 200 years ago, but even today many mis- takes in machine learning stem from failing to appreciate it. Every learner must embody some knowledge or as- sumptions beyond the data it is given in order to generalize beyond it. This notion was formalized by Wolpert in his famous “no free lunch” theorems, according to which no learner can beat random guessing over all pos- sible functions to be learned.25

This seems like rather depressing news. How then can we ever hope to learn anything? Luckily, the functions we want to learn in the real world are *not* drawn uniformly from the set of all mathematically possible functions! In fact, very general assumptions—like smoothness, similar examples hav- ing similar classes, limited depen- dences, or limited complexity—are often enough to do very well, and this is a large part of why machine learn- ing has been so successful. Like de- duction, induction (what learners do) is a knowledge lever: it turns a small amount of input knowledge into a large amount of output knowledge. Induction is a vastly more powerful lever than deduction, requiring much less input knowledge to produce use- ful results, but it still needs more than zero input knowledge to work. And, as with any lever, the more we put in, the more we can get out.

**test-set accuracy (%)**

A corollary of this is that one of the key criteria for choosing a representa- tion is which kinds of knowledge are easily expressed in it. For example, if we have a lot of knowledge about what makes examples similar in our do-

main, instance-based methods may be a good choice. If we have knowl- edge about probabilistic dependen- cies, graphical models are a good fit. And if we have knowledge about what kinds of preconditions are required by each class, “IF . . . THEN . . .” rules may be the best option. The most useful learners in this regard are those that do not just have assumptions hard- wired into them, but allow us to state them explicitly, vary them widely, and incorporate them automatically into the learning (for example, using first- order logic21 or grammars6).

In retrospect, the need for knowl- edge in learning should not be sur- prising. Machine learning is not magic; it cannot get something from nothing. What it does is get more from less. Programming, like all en- gineering, is a lot of work: we have to build everything from scratch. Learn- ing is more like farming, which lets nature do most of the work. Farmers combine seeds with nutrients to grow crops. Learners combine knowledge with data to grow programs.

### overfitting has many faces

What if the knowledge and data we have are not sufficient to completely determine the correct classifier? Then we run the risk of just hallucinating a classifier (or parts of it) that is not grounded in reality, and is simply en- coding random quirks in the data. This problem is called *overfitting*, and is the bugbear of machine learning. When your learner outputs a classi- fier that is 100% accurate on the train- ing data but only 50% accurate on test data, when in fact it could have output

**figure 2. naïve Bayes can outperform a state-of-the-art rule learner (c4.5rules) even when the true classifier is a set of rules.**

Bayes

C4.5

80

75

70

65

60

55

50

10

100

1000

**number of examples**

10000

one that is 75% accurate on both, it has overfit.

Everyone in machine learning knows about overfitting, but it comes in many forms that are not immedi- ately obvious. One way to understand overfitting is by decomposing gener- alization error into *bias* and *variance*.9 Bias is a learner’s tendency to con- sistently learn the same wrong thing. Variance is the tendency to learn ran- dom things irrespective of the real sig- nal. Figure 1 illustrates this by an anal- ogy with throwing darts at a board. A linear learner has high bias, because when the frontier between two classes is not a hyperplane the learner is un- able to induce it. Decision trees do not have this problem because they can represent any Boolean function, but on the other hand they can suffer from high variance: decision trees learned on different training sets generated by the same phenomenon are often very different, when in fact they should be



low bias

High bias

High variance

low variance

**figure 1. Bias and variance in dart-throwing.**

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the same. Similar reasoning applies to the choice of optimization meth- od: beam search has lower bias than greedy search, but higher variance, be- cause it tries more hypotheses. Thus, contrary to intuition, a more powerful learner is not necessarily better than a less powerful one.

Figure 2 illustrates this.a Even though the true classifier is a set of rules, with up to 1,000 examples na- ive Bayes is more accurate than a rule learner. This happens despite naive Bayes’s false assumption that the frontier is linear! Situations like this are common in machine learn- ing: strong false assumptions can be better than weak true ones, because a learner with the latter needs more data to avoid overfitting.

Cross-validation can help to com- bat overfitting, for example by using it to choose the best size of decision tree to learn. But it is no panacea, since if we use it to make too many parameter choices it can itself start to overfit.17

Besides cross-validation, there are many methods to combat overfit- ting. The most popular one is adding a *regularization term* to the evaluation function. This can, for example, pe- nalize classifiers with more structure, thereby favoring smaller ones with less room to overfit. Another option is to perform a statistical significance test like chi-square before adding new structure, to decide whether the dis- tribution of the class really is differ- ent with and without this structure. These techniques are particularly use- ful when data is very scarce. Neverthe- less, you should be skeptical of claims that a particular technique “solves” the overfitting problem. It is easy to avoid overfitting (variance) by falling into the opposite error of underfitting (bias). Simultaneously avoiding both requires learning a perfect classifier, and short of knowing it in advance there is no single technique that will always do best (no free lunch).

A common misconception about overfitting is that it is caused by noise,

a Training examples consist of 64 Boolean fea- tures and a Boolean class computed from them according to a set of “IF . . . THEN . . .” rules. The curves are the average of 100 runs with different randomly generated sets of rules. Error bars are two standard deviations. See Domingos and Pazzani10 for details.

like training examples labeled with the wrong class. This can indeed ag- gravate overfitting, by making the learner draw a capricious frontier to keep those examples on what it thinks is the right side. But severe overfitting can occur even in the absence of noise. For instance, suppose we learn a Bool- ean classifier that is just the disjunc- tion of the examples labeled “true” in the training set. (In other words, the classifier is a Boolean formula in disjunctive normal form, where each term is the conjunction of the feature values of one specific training exam- ple.) This classifier gets all the training examples right and every positive test example wrong, regardless of whether the training data is noisy or not.

The problem of *multiple testing*13 is closely related to overfitting. Standard statistical tests assume that only one hypothesis is being tested, but mod- ern learners can easily test millions before they are done. As a result what looks significant may in fact not be. For example, a mutual fund that beats the market 10 years in a row looks very impressive, until you realize that, if there are 1,000 funds and each has a 50% chance of beating the market on any given year, it is quite likely that one will succeed all 10 times just by luck. This problem can be combatted by correcting the significance tests to take the number of hypotheses into account, but this can also lead to un- derfitting. A better approach is to con- trol the fraction of falsely accepted non-null hypotheses, known as the *false discovery rate*.3

**intuition Fails in high Dimensions** After overfitting, the biggest problem in machine learning is the *curse of dimensionality*. This expression was coined by Bellman in 1961 to refer to the fact that many algorithms that work fine in low dimensions become intractable when the input is high- dimensional. But in machine learn- ing it refers to much more. General- izing correctly becomes exponentially harder as the dimensionality (number of features) of the examples grows, be- cause a fixed-size training set covers a dwindling fraction of the input space. Even with a moderate dimension of 100 and a huge training set of a trillion examples, the latter covers only a frac-

tion of about 10−18 of the input space. This is what makes machine learning both necessary and hard.

More seriously, the similarity- based reasoning that machine learn- ing algorithms depend on (explicitly or implicitly) breaks down in high di- mensions. Consider a nearest neigh- bor classifier with Hamming distance

as the similarity measure, and sup- pose the class is just *x*1 ∧ *x*2. If there are no other features, this is an easy problem. But if there are 98 irrelevant

features *x*3,..., *x*100, the noise from them completely swamps the signal in *x*1 and *x*2, and nearest neighbor effec- tively makes random predictions.

Even more disturbing is that near- est neighbor still has a problem even if all 100 features are relevant! This is because in high dimensions all examples look alike. Suppose, for instance, that examples are laid out on a regular grid, and consider a test example *xt*. If the grid is *d*-dimen- sional, *xt*’s 2*d* nearest examples are all at the same distance from it. So as the dimensionality increases, more and more examples become nearest neighbors of *xt*, until the choice of nearest neighbor (and therefore of class) is effectively random.

This is only one instance of a more general problem with high dimen- sions: our intuitions, which come from a three-dimensional world, of- ten do not apply in high-dimensional ones. In high dimensions, most of the mass of a multivariate Gaussian dis- tribution is not near the mean, but in an increasingly distant “shell” around it; and most of the volume of a high- dimensional orange is in the skin, not the pulp. If a constant number of ex- amples is distributed uniformly in a high-dimensional hypercube, beyond some dimensionality most examples are closer to a face of the hypercube than to their nearest neighbor. And if we approximate a hypersphere by in- scribing it in a hypercube, in high di- mensions almost all the volume of the hypercube is outside the hypersphere. This is bad news for machine learning, where shapes of one type are often ap- proximated by shapes of another.

Building a classifier in two or three dimensions is easy; we can find a rea- sonable frontier between examples of different classes just by visual in-

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spection. (It has even been said that if people could see in high dimensions machine learning would not be neces- sary.) But in high dimensions it is dif- ficult to understand what is happen- ing. This in turn makes it difficult to design a good classifier. Naively, one might think that gathering more fea- tures never hurts, since at worst they provide no new information about the class. But in fact their benefits may be outweighed by the curse of dimen- sionality.

Fortunately, there is an effect that partly counteracts the curse, which might be called the “blessing of non- uniformity.” In most applications examples are not spread uniformly throughout the instance space, but are concentrated on or near a lower- dimensional manifold. For example, *k*-nearest neighbor works quite well for handwritten digit recognition even though images of digits have one dimension per pixel, because the space of digit images is much smaller than the space of all possible images. Learners can implicitly take advan- tage of this lower effective dimension, or algorithms for explicitly reducing the dimensionality can be used (for example, Tenenbaum22).

### theoretical Guarantees are not What they seem

Machine learning papers are full of theoretical guarantees. The most com- mon type is a bound on the number of examples needed to ensure good gen- eralization. What should you make of these guarantees? First of all, it is re- markable that they are even possible. Induction is traditionally contrasted with deduction: in deduction you can guarantee that the conclusions are correct; in induction all bets are off. Or such was the conventional wisdom for many centuries. One of the major developments of recent decades has been the realization that in fact we can have guarantees on the results of in- duction, particularly if we are willing to settle for probabilistic guarantees.

The basic argument is remarkably simple.5 Let’s say a classifier is bad if its true error rate is greater than ε. Then the probability that a bad clas- sifier is consistent with *n* random, in- dependent training examples is less than (1 − ε)*n*. Let *b* be the number of

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bad classifiers in the learner’s hypoth- esis space *H*. The probability that at least one of them is consistent is less than *b*(1 − ε)*n*, by the union bound. As- suming the learner always returns a consistent classifier, the probability that this classifier is bad is then less than |*H*|(1 − ε)*n*, where we have used the fact that *b* ≤ |*H*|. So if we want this

probability to be less than δ, it suffices to make *n* > ln(δ/|*H*|)/ ln(1 − ε) ≥ 1/ε (ln

|*H*| + ln 1/δ).

Unfortunately, guarantees of this

type have to be taken with a large grain of salt. This is because the bounds ob- tained in this way are usually extreme- ly loose. The wonderful feature of the bound above is that the required num- ber of examples only grows logarith-

mically with |*H*| and 1/δ. Unfortunate- ly, most interesting hypothesis spaces

are *doubly* exponential in the number of features *d*, which still leaves us needing a number of examples expo- nential in *d*. For example, consider the space of Boolean functions of *d* Boolean variables. If there are *e* pos- sible different examples, there are 2*e* possible different functions, so since there are 2*d* possible examples, the total number of functions is 22*d*. And even for hypothesis spaces that are “merely” exponential, the bound is still very loose, because the union bound is very pessimistic. For exam- ple, if there are 100 Boolean features and the hypothesis space is decision trees with up to 10 levels, to guarantee

δ = ε = 1% in the bound above we need half a million examples. But in prac-

tice a small fraction of this suffices for accurate learning.

Further, we have to be careful about what a bound like this means. For instance, it does not say that, if your learner returned a hypothesis consistent with a particular training set, then this hypothesis probably generalizes well. What it says is that, given a large enough training set, with high probability your learner will ei- ther return a hypothesis that general- izes well or be unable to find a consis- tent hypothesis. The bound also says nothing about how to select a good hypothesis space. It only tells us that, if the hypothesis space contains the true classifier, then the probability that the learner outputs a bad classi- fier decreases with training set size.

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If we shrink the hypothesis space, the bound improves, but the chances that it contains the true classifier shrink also. (There are bounds for the case where the true classifier is not in the hypothesis space, but similar consid- erations apply to them.)

Another common type of theoreti- cal guarantee is asymptotic: given in- finite data, the learner is guaranteed to output the correct classifier. This is reassuring, but it would be rash to choose one learner over another be- cause of its asymptotic guarantees. In practice, we are seldom in the asymp- totic regime (also known as “asymp- topia”). And, because of the bias-vari- ance trade-off I discussed earlier, if learner A is better than learner B given infinite data, B is often better than A given finite data.

The main role of theoretical guar- antees in machine learning is not as a criterion for practical decisions, but as a source of understanding and driving force for algorithm design. In this capacity, they are quite useful; in- deed, the close interplay of theory and practice is one of the main reasons machine learning has made so much progress over the years. But caveat emptor: learning is a complex phe- nomenon, and just because a learner has a theoretical justification and works in practice does not mean the former is the reason for the latter.

### feature engineering is the Key

At the end of the day, some machine learning projects succeed and some fail. What makes the difference? Eas- ily the most important factor is the features used. Learning is easy if you have many independent features that each correlate well with the class. On the other hand, if the class is a very complex function of the features, you may not be able to learn it. Often, the raw data is not in a form that is ame- nable to learning, but you can con- struct features from it that are. This is typically where most of the effort in a machine learning project goes. It is often also one of the most interesting parts, where intuition, creativity and “black art” are as important as the technical stuff.

First-timers are often surprised by how little time in a machine learning project is spent actually doing ma-

# a dumb algorithm with lots and lots of data beats

**a clever one with modest amounts of it.**

chine learning. But it makes sense if you consider how time-consuming it is to gather data, integrate it, clean it and preprocess it, and how much trial and error can go into feature design. Also, machine learning is not a one- shot process of building a dataset and running a learner, but rather an itera- tive process of running the learner, analyzing the results, modifying the data and/or the learner, and repeat- ing. Learning is often the quickest part of this, but that is because we have already mastered it pretty well! Feature engineering is more diffi- cult because it is domain-specific, while learners can be largely general purpose. However, there is no sharp frontier between the two, and this is another reason the most useful learn- ers are those that facilitate incorpo- rating knowledge.

Of course, one of the holy grails of machine learning is to automate more and more of the feature engi- neering process. One way this is often done today is by automatically gener- ating large numbers of candidate fea- tures and selecting the best by (say) their information gain with respect to the class. But bear in mind that features that look irrelevant in isola- tion may be relevant in combination. For example, if the class is an XOR of *k* input features, each of them by it- self carries no information about the class. (If you want to annoy machine learners, bring up XOR.) On the other hand, running a learner with a very large number of features to find out which ones are useful in combination may be too time-consuming, or cause overfitting. So there is ultimately no replacement for the smarts you put into feature engineering.

### more Data Beats

**a cleverer algorithm**

Suppose you have constructed the best set of features you can, but the classifiers you receive are still not ac- curate enough. What can you do now? There are two main choices: design a better learning algorithm, or gather more data (more examples, and pos- sibly more raw features, subject to the curse of dimensionality). Machine learning researchers are mainly con- cerned with the former, but pragmati- cally the quickest path to success is

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often to just get more data. As a rule of thumb, a dumb algorithm with lots and lots of data beats a clever one with modest amounts of it. (After all, ma- chine learning is all about letting data do the heavy lifting.)

This does bring up another prob- lem, however: scalability. In most of computer science, the two main lim- ited resources are time and memory. In machine learning, there is a third one: training data. Which one is the bottleneck has changed from decade to decade. In the 1980s it tended to be data. Today it is often time. Enor- mous mountains of data are avail- able, but there is not enough time to process it, so it goes unused. This leads to a paradox: even though in principle more data means that more complex classifiers can be learned, in practice simpler classifiers wind up being used, because complex ones take too long to learn. Part of the an- swer is to come up with fast ways to learn complex classifiers, and indeed there has been remarkable progress in this direction (for example, Hulten and Domingos11).

Part of the reason using cleverer algorithms has a smaller payoff than you might expect is that, to a first ap- proximation, they all do the same. This is surprising when you consider representations as different as, say, sets of rules and neural networks. But in fact propositional rules are readily encoded as neural networks, and sim- ilar relationships hold between other representations. All learners essen- tially work by grouping nearby exam- ples into the same class; the key dif- ference is in the meaning of “nearby.” With nonuniformly distributed data, learners can produce widely different frontiers while still making the same predictions in the regions that matter (those with a substantial number of training examples, and therefore also where most test examples are likely to appear). This also helps explain why powerful learners can be unstable but still accurate. Figure 3 illustrates this in 2D; the effect is much stronger in high dimensions.

As a rule, it pays to try the simplest learners first (for example, naïve Bayes before logistic regression, *k*-nearest neighbor before support vector ma- chines). More sophisticated learn-

ers are seductive, but they are usually harder to use, because they have more knobs you need to turn to get good re- sults, and because their internals are more opaque.

Learners can be divided into two major types: those whose representa- tion has a fixed size, like linear classi- fiers, and those whose representation can grow with the data, like decision trees. (The latter are sometimes called nonparametric learners, but this is somewhat unfortunate, since they usually wind up learning many more parameters than parametric ones.) Fixed-size learners can only take ad- vantage of so much data. (Notice how the accuracy of naive Bayes asymptotes at around 70% in Figure 2.) Variable- size learners can in principle learn any function given sufficient data, but in practice they may not, because of limi- tations of the algorithm (for example, greedy search falls into local optima) or computational cost. Also, because of the curse of dimensionality, no ex- isting amount of data may be enough. For these reasons, clever algorithms— those that make the most of the data and computing resources available— often pay off in the end, provided you are willing to put in the effort. There is no sharp frontier between design- ing learners and learning classifiers; rather, any given piece of knowledge could be encoded in the learner or learned from data. So machine learn- ing projects often wind up having a significant component of learner de- sign, and practitioners need to have some expertise in it.12

In the end, the biggest bottleneck is not data or CPU cycles, but human



D. Tree

SVM

knn

n. bayes

**Figure 3. very different frontiers can yield similar predictions. (+ and – are training examples of two classes.)**

cycles. In research papers, learners are typically compared on measures of accuracy and computational cost. But human effort saved and insight gained, although harder to measure, are often more important. This favors learners that produce human-under- standable output (for example, rule sets). And the organizations that make the most of machine learning are those that have in place an infrastruc- ture that makes experimenting with many different learners, data sources, and learning problems easy and effi- cient, and where there is a close col- laboration between machine learning experts and application domain ones.

**Learn many models, not Just one** In the early days of machine learn- ing, everyone had a favorite learner, together with some a priori reasons to believe in its superiority. Most ef- fort went into trying many variations of it and selecting the best one. Then systematic empirical comparisons showed that the best learner varies from application to application, and systems containing many different learners started to appear. Effort now went into trying many variations of many learners, and still selecting just the best one. But then researchers noticed that, if instead of selecting the best variation found, we combine many variations, the results are bet- ter—often much better—and at little extra effort for the user.

Creating such *model ensembles* is now standard.1 In the simplest tech- nique, called *bagging*, we simply gen- erate random variations of the train- ing set by resampling, learn a classifier on each, and combine the results by voting. This works because it greatly reduces variance while only slightly increasing bias. In *boosting*, training examples have weights, and these are varied so that each new classifier fo- cuses on the examples the previous ones tended to get wrong. In *stacking*, the outputs of individual classifiers become the inputs of a “higher-level” learner that figures out how best to combine them.

Many other techniques exist, and the trend is toward larger and larger ensembles. In the Netflix prize, teams from all over the world competed to build the best video recommender

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system (http://netflixprize.com). As the competition progressed, teams found they obtained the best results by combining their learners with oth- er teams’, and merged into larger and larger teams. The winner and runner- up were both stacked ensembles of over 100 learners, and combining the two ensembles further improved the results. Doubtless we will see even larger ones in the future.

Model ensembles should not be confused with Bayesian model av- eraging (BMA)—the theoretically optimal approach to learning.4 In BMA, predictions on new examples are made by averaging the individual predictions of *all* classifiers in the hypothesis space, weighted by how well the classifiers explain the train- ing data and how much we believe in them a priori. Despite their su- perficial similarities, ensembles and BMA are very different. Ensembles change the hypothesis space (for ex- ample, from single decision trees to linear combinations of them), and can take a wide variety of forms. BMA assigns weights to the hypotheses in the original space according to a fixed formula. BMA weights are extremely different from those produced by (say) bagging or boosting: the latter are fairly even, while the former are extremely skewed, to the point where the single highest-weight classifier usually dominates, making BMA ef- fectively equivalent to just selecting it.8 A practical consequence of this is that, while model ensembles are a key part of the machine learning toolkit, BMA is seldom worth the trouble.

### Simplicity Does not imply Accuracy

Occam’s razor famously states that entities should not be multiplied be- yond necessity. In machine learning, this is often taken to mean that, given two classifiers with the same training error, the simpler of the two will likely have the lowest test error. Purported proofs of this claim appear regularly in the literature, but in fact there are many counterexamples to it, and the “no free lunch” theorems imply it can- not be true.

We saw one counterexample previ- ously: model ensembles. The gener- alization error of a boosted ensemble

# Just because a function can be represented does not mean

**it can be learned.**

continues to improve by adding clas- sifiers even after the training error has reached zero. Another counterexam- ple is support vector machines, which can effectively have an infinite num- ber of parameters without overfitting. Conversely, the function sign(sin(*ax*)) can discriminate an arbitrarily large, arbitrarily labeled set of points on the *x* axis, even though it has only one pa- rameter.23 Thus, contrary to intuition, there is no necessary connection be- tween the number of parameters of a model and its tendency to overfit.

A more sophisticated view instead equates complexity with the size of the hypothesis space, on the basis that smaller spaces allow hypotheses to be represented by shorter codes. Bounds like the one in the section on theoreti- cal guarantees might then be viewed as implying that shorter hypotheses generalize better. This can be further refined by assigning shorter codes to the hypotheses in the space we have some a priori preference for. But viewing this as “proof” of a trade-off between accuracy and simplicity is circular reasoning: we made the hy- potheses we prefer simpler by design, and if they are accurate it is because our preferences are accurate, not be- cause the hypotheses are “simple” in the representation we chose.

A further complication arises from the fact that few learners search their hypothesis space exhaustively. A learner with a larger hypothesis space that tries fewer hypotheses from it is less likely to overfit than one that tries more hypotheses from a smaller space. As Pearl18 points out, the size of the hypothesis space is only a rough guide to what really matters for relat- ing training and test error: the proce- dure by which a hypothesis is chosen. Domingos7 surveys the main argu- ments and evidence on the issue of Occam’s razor in machine learning. The conclusion is that simpler hy- potheses should be preferred because simplicity is a virtue in its own right, not because of a hypothetical connec- tion with accuracy. This is probably

what Occam meant in the first place.

### Representable Does not imply Learnable

Essentially all representations used in variable-size learners have associated

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theorems of the form “Every function can be represented, or approximated arbitrarily closely, using this repre- sentation.” Reassured by this, fans of the representation often proceed to ignore all others. However, just be- cause a function can be represented does not mean it can be learned. For example, standard decision tree learn- ers cannot learn trees with more leaves than there are training examples. In continuous spaces, representing even simple functions using a fixed set of primitives often requires an infinite number of components. Further, if the hypothesis space has many local optima of the evaluation function, as is often the case, the learner may not find the true function even if it is rep- resentable. Given finite data, time and memory, standard learners can learn only a tiny subset of all possible func- tions, and these subsets are different for learners with different represen- tations. Therefore the key question is not “Can it be represented?” to which the answer is often trivial, but “Can it be learned?” And it pays to try different learners (and possibly combine them). Some representations are exponen- tially more compact than others for some functions. As a result, they may also require exponentially less data to learn those functions. Many learners work by forming linear combinations of simple basis functions. For exam- ple, support vector machines form combinations of kernels centered at some of the training examples (the support vectors). Representing parity of *n* bits in this way requires 2*n* basis functions. But using a representation with more layers (that is, more steps between input and output), parity can be encoded in a linear-size classifier. Finding methods to learn these deeper representations is one of the major re- search frontiers in machine learning.2

### Correlation Does not imply Causation

The point that correlation does not imply causation is made so often that it is perhaps not worth belaboring. But, even though learners of the kind we have been discussing can only learn correlations, their results are often treated as representing causal relations. Isn’t this wrong? If so, then why do people do it?

More often than not, the goal of learning predictive models is to use them as guides to action. If we find that beer and diapers are often bought together at the supermar- ket, then perhaps putting beer next to the diaper section will increase sales. (This is a famous example in the world of data mining.) But short of actually doing the experiment it is difficult to tell. Machine learning is usually applied to *observational* data, where the predictive variables are not under the control of the learner, as opposed to *experimental* data, where they are. Some learning algorithms can potentially extract causal infor- mation from observational data, but their applicability is rather restrict- ed.19 On the other hand, correlation is a sign of a potential causal connec- tion, and we can use it as a guide to further investigation (for example, trying to understand what the causal chain might be).

Many researchers believe that cau- sality is only a convenient fiction. For example, there is no notion of causal- ity in physical laws. Whether or not causality really exists is a deep philo- sophical question with no definitive answer in sight, but there are two practical points for machine learn- ers. First, whether or not we call them “causal,” we would like to predict the effects of our actions, not just corre- lations between observable variables. Second, if you can obtain experimen- tal data (for example by randomly as- signing visitors to different versions of a Web site), then by all means do so.14

### Conclusion

Like any discipline, machine learn- ing has a lot of “folk wisdom” that can be difficult to come by, but is crucial for success. This article summarized some of the most salient items. Of course, it is only a complement to the more conventional study of machine learning. Check out [http://www.](http://www/) cs.washington.edu/homes/pedrod/ class for a complete online machine learning course that combines formal and informal aspects. There is also a treasure trove of machine learning lectures at http://www.videolectures. net. A good open source machine learning toolkit is Weka.24

Happy learning!

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