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Practical PAC Learning

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Abstract

We present new strategies for "probably approximately correct" (pac) learning that use fewer training examples than previous approaches. The idea is to observe training examples one-at-a-time and decide "on-line" when to return a hypothesis, rather than collect a large fixed-size training sample. This yields sequential learning procedures that pac-learn by observing a small random number of examples. We provide theoretical bounds on the expected training sample size of our procedure — but establish its efficiency primarily by a series of experiments which show sequential learning actually uses many times fewer training examples in practice. These results demonstrate that paclearning can be far more efficiently achieved in practice than previously thought.

1 Introduction

We consider the standard problem of learning an accurate classifier from examples: given a target classification scheme $c: X \to Y$ defined on a domain X, we are interested in observing a sequence of training examples $\langle \langle x_1, c(x_1) \rangle, ..., \langle x_t, c(x_t) \rangle \rangle$ and producing a hypothesis $h: X \to Y$ that agrees with c on as much of the domain as possible. Here we adopt the standard batch training protocol, where after a finite number of training examples the learner must produce a hypothesis h, which is then tested ad infinitum on subsequent training examples.

In practice, domain objects can be represented in many different ways (e.g., boolean or real-valued vectors, or structured descriptions like strings, graphs, terms, etc.), and so too can hypotheses (e.g., decision trees, neural networks, nearest neighbor classifiers, etc.). However, regardless of the specific representation used, the central question is always how best to extrapolate the classifications of a few domain objects to an accurate classification scheme over the entire domain.

Motivation: Classification learning is by far the most studied in machine learning research. The immense interest in this problem arises from the fact that classification itself is an important subtask in many appli-

cations — in fact, comprising the central function of most expert systems [Clancey, 1985]. The importance of learning in this context is that we often lack the requisite knowledge needed to specify an appropriate classifier, and yet have access to many correctly classified examples. In such situations, we can attempt to exploit the wealth of available data to overcome inadequate prior knowledge — and hence, use learning as an effective classifier synthesis tool. In fact, there are numerous examples where learning systems have produced classifiers that outperform the best available "hand-coded" systems, e.g., [le Cun, et al., 1989; Weiss and Kulikowski, 1991].

Although empirical research tends to examine the performance properties of particular hypothesis guessing strategies on specific domains, the underlying goal of classification learning research is to uncover whatever general principles might underly the effective extrapolation of training object classifications to entire domains. However, it has often been observed that there really is no such thing as a "general purpose" extrapolation strategy [Schaffer, 1994] — a particular strategy performs well on a specific application only by fortuitous predisposition: it just happens to "guess right" on unseen domain objects, whether by prior knowledge or luck. To guarantee success, one must supply prior constraints.

The current trend towards theoretical analysis in machine learning represents a fundamental shift in emphasis away from discovering "universal" extrapolation strategies, towards explicitly acknowledging the role played by prior constraints in yielding successful extrapolation. The role of a theoretical analysis is not to prescribe prior knowledge/constraints, but to determine the best that can be achieved given whatever is known beforehand.

1.1 Pac-learning theory

The most influential analysis of classification learning is the theory of "probably approximately correct" (pac) learning introduced by Valiant [1984]. Rather than speculate about the mechanisms that might underly "general purpose" classification learning, Valiant's idea was to characterize those situations where successful learning could be *provably* achieved, and where it is demonstrably impossible.

Problem: Pac-learning theory adopts an "i.i.d. random examples" model of the learning situation, which

assumes domain objects are independently generated by a fixed distribution P_X and labelled according to a fixed target concept $c: X \to \{0,1\}$. Under this model, the error of a hypothesis $h: X \to \{0,1\}$ with respect to c and P_X is given by $P_X\{h(x) \neq c(x)\}$. Here we consider the difficulty of meeting the so-called pac-criterion: producing a hypothesis h with error at most ϵ , with probability at least $1-\delta$, for specified accuracy and reliability parameters ϵ and δ . Of course, the difficulty of achieving this criterion depends on how much we know about c and P_X beforehand. Pac-learning theory adopts a model of prior knowledge where we assume the target concept c belongs to some known class C, but nothing is known about the domain distribution P_X . Given this model, we naturally consider what can be achieved in the "worst case, distribution-free" sense:

Definition 1 (Pac-learning problem) A learner L solves the pac-learning problem $\langle X, C, \epsilon, \delta \rangle$ (or "pac (ϵ, δ) -learns C") if, for any c in C and P_X , L produces a hypothesis h such that $P_X\{h(x) \neq c(x)\} \leq \epsilon$ with probability at least $1-\delta$ (over possible training sequences).

For example, we might be interested in solving the problem $\langle X=I\!\!R^{10},C=$ halfspace, $\epsilon=0.01,\delta=0.05\rangle$, where domain objects are described by 10 real-valued attributes, the target concept is known to be some linear-halfspace of $I\!\!R^{10}$, and we wish to produce a hypothesis with 1% error with probability at least 95%. Our goal is to solve these learning problems as efficiently as possible — i.e., using a minimum of data and computational resources. The primary focus of this paper is on improving the data-efficiency of pac-learners, rather than their computational-efficiency.

Results: Some of the most important technical results of pac-learning theory concern the data resources needed to solve pac-learning problems. Intuitively, it should take more training examples to pac-learn a complex concept class than a simple one, since it is harder to disambiguate possible target concepts from a complex class. The question is: how can one measure the representational complexity of a concept class C so as to precisely determine the number of training examples needed to $pac(\epsilon, \delta)$ -learn C? It turns out that just such a measure is provided by the Vapnik-Chervonenkis (VC) dimension of C: Ehrenfeucht et al. [1989] have shown that, for any concept class C with vc(C) = d, the minimum number of training examples needed by any learner to $pac(\epsilon, \delta)$ learn C is at least $t_{EHKV}(C,\epsilon,\delta) = \max\left\{\frac{d-1}{32\epsilon}, \frac{1-\epsilon}{\epsilon}\ln\frac{1}{\delta}\right\}$. Furthermore, there is a simple $\mathit{fixed-sample-size}$ learning procedure, F, that always meets this lower bound to within constant and log factors, and hence learns with near-optimal data-efficiency; see Figure 1. In particular, Blumer et al. [1989] have shown that for any² conProcedure F (C, ϵ, δ)

Collect $T_{\mathbf{F}}(C, \epsilon, \delta)$ training examples, sufficient to eliminate all ϵ -bad concepts from C with prob. at least $1-\delta$. Return any $h \in C$ that correctly classifies every example.

Figure 1: Procedure F

cept class C, $T_{BEHW}(C,\epsilon,\delta) = \max\left\{\frac{8d}{\epsilon}\log_2\frac{13}{\epsilon}, \frac{4}{\epsilon}\log_2\frac{2}{\delta}\right\}$ random training examples are sufficient to ensure F pac (ϵ,δ) -learns C, where $d=\mathrm{vc}(C)$. (This result has since been improved by Shawe-Taylor et~al.~[1993] to $T_{STAB}(C,\epsilon,\delta) = \frac{1}{\epsilon(1-\sqrt{\epsilon})}\left(2d\ln\frac{6}{\epsilon}+\ln\frac{2}{\delta}\right)$.) Overall, these are powerful results as they characterize the necessary and sufficient training sample sizes needed to pac-learn any concept class C in terms of a "tight" linear function of its VCdimension.

1.2 Issue

However, despite these impressive results, pac-learning theory has arguably had little direct impact on the actual practice of machine learning. Why? Beyond criticisms of certain modelling assumptions (e.g., noise-free examples, bivalent classifications, etc.— which actually have been addressed the pac-framework, cf. [Haussler, 1992]), the most prevalent criticism of pac-learning theory is that the actual numbers of training examples it demands are far too large to be practical.

Example: Consider the $\langle X = \mathbb{R}^{10}, C = \mathsf{halfspaces}, \epsilon =$ $0.01, \delta = 0.05$ problem mentioned earlier. Noting that $\mathrm{vc}(C) = 11$, we can simply use T_{BEHW} to determine a sufficient sample size for Procedure F. But here we find T_{BEHW} demands 91,030 training examples! (Even the improved T_{STAB} demands 15, 981 examples in this case.) This seems like an outrageous number given the apparently modest parameter settings. Moreover, these results compare poorly to the empirical "rule of thumb" that, for a concept class defined by w free parameters, roughly $T_{thumb} = \frac{w}{\epsilon}$ training examples are needed to achieve an error of ϵ [Baum and Haussler, 1989]. Applied here, T_{thumb} demands only 1, 100 training examples — an order of magnitude fewer than T_{STAB} . (Of course, this rule of thumb comes with no guarantees, but it does give an indication of how many training examples practitioners would deem "reasonable" for this problem.) Furthermore, T_{BEHW} and T_{STAB} are orders of magnitude larger than the best known lower bound t_{EHKV} , which demands only 32 training examples in this case! See Table 1 in Section 3 for a direct comparison.

This shows that, although the theoretical upper and lower bounds are tight up to constant and log factors, they give results that are orders of magnitude apart in practice. This has drastic consequences for the applicability of the theory, since in practice it is often training data, not computation time, that is the critical resource. I.e., cutting the training sample size in half would be a significant improvement in most applications, even if this came with a slight increase in overall computation time.

[Blumer, et al., 1989], which we will assume throughout.

¹The VCdimension measures how "fine grained" C is by the maximum number of domain objects C can independently label [Vapnik and Chervonenkis, 1971]. This is an abstract combinatorial measure which applies to arbitrary domains and concept classes. Moreover, it often gives intuitive results (e.g., the class of halfspace concepts on \mathbb{R}^n is defined by n+1 "free parameters" and also has a VCdimension of n+1).

²C must satisfy certain (benign) measurability constraints

The apparent inefficiency of pac-learning has lead to much speculation about the sources of difficulty. The predominant "folk wisdom" is that the large sample sizes follow from the worst case nature of the pacguarantees [Haussler, 1990] — that is, the worst case bounds are inherently unreasonable because they must take into account "pathological" domain distributions and target concepts which force large training sample sizes (moreover, the argument continues, these pathological situations do not arise in "typical" applications). In fact, this belief motivates much research that makes distributional assumptions in order to improve dataefficiency, e.g., Benedek and Itai, 1988; Aha, et al., 1991; Bartlett and Williamson, 1991]. However, notice that this line of reasoning is actually quite weak: First of all, no-one can demonstrate that these "pathological" situations really exist (for this would be tantamount to improving the lower bound t_{EHKV}). Secondly, it is clear from the previous example that the current bounds are loose, and can likely be substantially improved — e.g., T_{STAB} and t_{EHKV} differ by roughly a factor of $64 \ln \frac{6}{\epsilon}$.

Approach: In this paper we investigate an alternative view: perhaps the simplistic (collect; find) learning procedure F is not particularly data-efficient. This raises the obvious question of whether alternative learning strategies might be more data-efficient than F. Here we investigate sequential learning procedures that observe training examples one-at-a-time and autonomously decide "online" when to stop training and return a hypothesis. The idea is that we should be able to detect situations where an accurate hypothesis can be reliably returned, even before the sufficient sample size bounds have been reached (e.g., we might detect that C has been reduced to a single possible target). The hope is that, in this way, we can significantly reduce the number of training examples observed, while still meeting the exact same pac-criterion as before: namely, that an ϵ -accurate hypothesis be returned with probability at least $1-\delta$ for any target concept $c \in C$ and distribution P_x . An underlying assumption here is that we are willing to incur a slight computational penalty to obtain a significant improvement in data-efficiency. This is motivated by the fact that training data is usually the most critical resource in practical learning applications.

The remainder of this paper develops a few simple sequential learning procedures that (i) are correct paclearners, (ii) are provably data-efficient, and (iii) use many times fewer training examples in empirical case studies.

2 Sequential pac-learning

A sequential learner L consists of a stopping rule T_L , that maps training sequences to stopping times, and a hypothesizer H_L , that maps finite training sequences to hypotheses. Our basic strategy for constructing sequential pac-learners is to take an arbitrary consistent hypothesizer H for C (which produces hypotheses $h \in C$ that correctly classify every observed training example), collect H's hypotheses, and test these against subsequent training examples until one proves to have sufficiently

Procedure R (C, ϵ, δ, H)

FIX a sequence $\{\delta_i\}_1^{\infty}$ such that $\sum \delta_i = \delta$. Call H to obtain an initial hypothesis h_0 .

SEQUENTIALLY observe training examples $\langle x_t, y_t \rangle$, t = 1, 2, ...:

If current hypothesis h_i makes a mistake, call H to obtain a consistent h_{i+1} (drop h_i , begin testing h_{i+1}).

RETURN current hypothesis h_i if it correctly classifies $\frac{1}{\epsilon} \ln \frac{1}{\delta_i}$ consecutive training examples.

Figure 2: Procedure R

small error. The main challenge is finding an appropriate stopping rule that guarantees the pac-criterion, while observing as few training examples as possible.

Note that, in general, a sequential learner observes a random, rather than fixed, number of training examples. Thus, to compare the data-efficiency of our approach with previous techniques, we must compare a distribution of sample sizes to a fixed number. There are a number of ways one could do this, but we focus on what is arguably the most natural measure: comparing the average (i.e., expected) training sample size of a sequential learner with the fixed sample size demanded by previous approaches to solve the same pac-learning problem.

Obvious approach: Perhaps the most obvious strategy for sequential pac-learning is based on the idea of repeated significance testing: test a series of hypotheses generated by H until one correctly classifies a sufficient number of consecutive training examples; see Procedure R in Figure 2.³ Although this is a plausible approach (which, in fact, works well in practice), it is hard to prove reasonable bounds on R's expected sample size. The problem is that R rejects "good enough" hypotheses with high probability, and yet takes a long time to do so (i.e., R rejects hypotheses of error ϵ with probability $1-\delta$, but this takes $\frac{1}{\epsilon}$ expected time). Thus, if H produces a series of "borderline" hypotheses, R will take a long time to terminate (expected time about $\frac{1}{\epsilon\delta}$, which is not very good). Fortunately, there is a better approach.

Better approach: Here we introduce a novel learning procedure, S (Figure 3), which is also based on repeated significance testing, but avoids the apparent inefficiency of R's "survival testing" approach. S is based on two ideas: First, instead of throwing away H's hypotheses after a single mistake, S saves hypotheses and continues testing them until one proves to have small error. Second, S identifies accurate hypotheses by using a sequential probability ratio test (sprt) [Wald, 1947] to test each candidate "on-line" (in parallel); Figure 4. Thus, S never rejects a potentially acceptable hypothesis, and quickly identifies any sufficiently accurate candidate.

Procedure S is a correct pac-learner in the exact same sense as F: The key property of S is that its call to sprt

³Variants of Procedure R have been proposed by many authors in the past [Linial, et al., 1991; Oblow, 1992], primarily to achieve "nonuniform" pac-learning. However, the goals of nonuniform pac-learning fundamentally differ from what we are trying to accomplish here (see Footnote 6).

Procedure S (C, ϵ, δ, H)

FIX a sequence $\{\delta_i = \frac{6\delta}{\pi^2 i^2}\}_1^{\infty}$ and a constant $\kappa > 1$. Initialize a list of hypotheses with h_0 , obtained by calling H.

SEQUENTIALLY observe training examples $\langle x_t, y_t \rangle$, t = 1, 2, ...:

If the most recent hypothesis h_i makes a mistake, call H to add a new, consistent hypothesis h_{i+1} to the list.

TEST all hypotheses in the list (in parallel) by calling $\operatorname{sprt}(h_i(x) \neq c(x), \frac{\epsilon}{5}, \epsilon, \delta_i, 0)$

for each h_i (when generated).

RETURN the first generated hypothesis h_i sprt accepts.

Figure 3: Procedure S

eventually accepts any $\frac{\epsilon}{\kappa}$ -good hypothesis with probability 1 (wp1), but only accepts an ϵ -bad hypothesis h_i with probability at most δ_i . This implies that S eventually halts wp1, and returns an ϵ -good hypothesis with probability at least $1-\delta$, for any target concept $c \in C$ and domain distribution P_X (thus, achieving the exact same worst case pac-guarantees as F). This property also allows us to prove a reasonable upper bound on the average number of training examples S observes for any target concept $c \in C$ and domain distribution P_X .

Theorem 1 For $\epsilon > 0$, $\delta > 0$, and any (well behaved) concept class C with vc(C) = d: using a consistent hypothesizer H for C and any constant $\kappa > 1$, Procedure S observes an average training sample size of at most

$$ET_{\mathbb{S}}(C,\epsilon,\delta) \leq \left(\frac{\kappa}{\kappa-1-\ln\kappa}\right) \frac{1}{\epsilon} \left(\left[2.12\kappa d+3\right] \ln \frac{14\kappa}{\epsilon} + \ln \frac{1}{\delta}\right)$$

for any target concept $c \in C$ and distribution P_x .

Although this is a crude bound, it is interesting to note that it scales the same as T_{BEHW} and T_{STAB} . Moreover, this bound actually beats T_{BEHW} and T_{STAB} for small values of δ [Schuurmans, 1995]. However, as shown below, S actually performs much better in practice than any bounds we can prove about its performance. Since this is not a possibility for fixed-sample-sized approaches, we expect S to perform much better than T_{BEHW} and T_{STAB} in practical applications.

Before demonstrating S's advantage in empirical tests, we first note that there are inherent limits to the data-efficiency even of sequential learning.

Theorem 2 For sufficiently small ϵ and δ , and any concept class C with $\operatorname{vc}(C) = d \geq 2$: any learner that always observes (for any fixed $c \in C$ and P_X) an average training sample size less than

$$t_{avg}(C,\epsilon,\delta) = \max \big\{ \frac{d-1}{480\epsilon},\, \frac{1-\delta}{4\epsilon} \big\}$$

cannot meet the $pac(\epsilon, \delta)$ -criterion for all $c \in C$ and P_X . Notice that this lower bound scales the same as t_{EHKV} in terms of ϵ and VC(C) — which shows that no new concept classes become pac-learnable merely by considering a sequential over fixed-sample-size approach.

Procedure sprt $(\phi(x), a, r, \delta_{acc}, \delta_{rej})$

For boolean random variable $\phi(x)$, test

 H_{acc} : $P_X\{\phi(x)=1\} \le a$ vs. H_{rej} : $P_X\{\phi(x)=1\} \ge r$, with:

– probability of deciding H_{acc} given H_{rej} bounded by δ_{acc} ,

- probability of deciding H_{rej} given H_{acc} bounded by δ_{rej} .

SEQUENTIALLY observe $\phi_t = \phi(x_t)$, t = 1, 2, ...; monitoring:

$$S_t = \sum_{j=1}^t \phi_j \ln \frac{a}{r} + (1 - \phi_j) \ln \frac{1-a}{1-r}.$$

RETURN "accept" if ever $S_t \ge \ln 1/\delta_{acc}$.

RETURN "reject" if ever $S_t \leq \ln \delta_{rej}$.

Figure 4: Procedure sprt

3 Empirical efficiency

Although the theoretical advantage we can demonstrate for S is only slight, we expect S to perform $much\ better$ in practice than any bounds we can prove about its performance. This is because S's actual data-efficiency in any particular case study is determined by the specific case at hand, and not by the worst case situation (or, worse yet, what we can prove about the worst case situation). In fact, in empirical studies, S proves to be far more efficient than any bounds we can prove about its performance, and many times more efficient than T_{BEHW} or T_{STAB} . This is easily demonstrated by a simple example.

Illustration: We tested Procedure S on the problem $\langle X = \mathbb{R}^n, C = \text{halfspaces}, \epsilon = 0.01, \delta = 0.05 \rangle$ with the following setup: Training objects were generated according to a uniform distribution on $[-1, 1]^n$ and labelled by a fixed target halfspace (defined by a "diagonal" hyperplane passing through the origin 0^n with norm directed towards $\mathbf{1}^n$). The constant κ was set to 3.14619 (so that $\frac{\kappa}{\kappa-1-\ln\kappa} = \kappa$), and we supplied S with a hypothesizer H that finds consistent halfspace concepts.⁵ We ran Procedure S 100 times for n = 10 and obtained the results shown in Table 1. Notice that S's average training sample size of 3, 402 is about 5 times smaller than T_{STAB} , 27 times smaller than T_{BEHW} , and only about 3 times larger than T_{thumb} . It is important to emphasize that S obtains these empirical sample size improvements while maintaining the exact same worst case pac-guarantees as before (that an ϵ -accurate hypothesis is returned with probability at least $1-\delta$). These results are in fact representative over the entire range of parameter settings: S's empirical advantage actually *improves* for increased problem dimension n (Figure 5), and is maintained at higher accuracy and reliability levels [Schuurmans, 1995]. Overall, S appears to be pac-learning with near-practical data-efficiency in this example.

Interestingly, S also outperforms the simplistic procedure R on this problem. Figure 6 shows that, R performs nearly as well as S on easy problems (low dimension, accuracy, reliability), but S's advantage grows significantly as these parameters are scaled up.

⁴Provided $VC(C) < \infty$ (details omitted). Proofs of all results mentioned in this paper (and more) are outlined in [Schuurmans and Greiner, 1995]. Complete details appear in [Schuurmans, 1995].

⁵Specifically, we used the BFGS secant optimization procedure [Dennis and Schnabel, 1983] with a "relaxation" objective function [Duda and Hart, 1973].

Explanations: These results demonstrate a clear advantage for sequential over fixed-sample-size learning: we solve the exact same pac-learning problem using far fewer training examples in this case. Of course these preceding results are anecdotal, and it is tempting to explain away the advantage as a mere artifact of the specific experimental setup. However, we have found that these experimental results are, in fact, quite robust.

First, the previous experiment only tested a single domain distribution (uniform), which could happen to be a particularly "easy" one for S. To counter this claim, we repeated the experiment with various domain distributions to see if any could seriously affect S's performance. In particular, we considered three different transformations of the uniform $[-1,1]^n$ distribution: spherical (nonlinear compression towards origin), pyramidal (compression from opposite corners towards hyperplane), and accretive (translation towards discrete points in $\{-1,1\}^n$). Surprisingly, none of these transformations had any noticeable effect on S's performance [Schuurmans, 1995]; as demonstrated in Figure 7 for the pyramidal case.

A second reason for S's advantage might be that the specific target concept (diagonal halfspace) is a particularly "easy" one for $S-i.e.,\ H$ could somehow be biased to guess similar hypotheses. However, this is easily shown not to be the case: We repeated the original experiment on 10 different target halfspaces, each successively closer to "axis-parallel," and found that none of these made any appreciable difference; Figure 8.

Third, it could be the case that the class of halfspaces concepts happens to be "easy" among classes with comparable VCdimension. This turns out to be partly true: We have been able to construct alternative concept classes which force S to observe slightly more training examples; see Figure 9. However, we have yet to devise any concept class (with the same VCdimension) that can even double S's original performance on halfspaces. In fact, S's performance often improves for different concept classes (particularly finite ones). Overall, it appears that halfspaces is not a remarkably hard or easy class for a given VCdimension.

Another explanation of S's advantage over F is that T_{STAB} might possibly be a gross overestimate of the true worst case situation (which seems likely, given the gap between T_{STAB} and t_{EHKV}). Of course, this means that any current advantage enjoyed by S could potentially be overcome by future improvements to T_{STAB} — but notice that we can enjoy S's improved performance immediately, without having to wait for theoreticians to improve the bounds. (Ensuring the correctness of F requires one to prove some bound is sufficient; this is not a requirement for Procedure S since its correctness is completely decoupled from its efficiency.)

A final explanation of S's advantage is that sequential learning might be *inherently* more efficient than fixed-sample-size learning. Clearly, since the sequential approach *generalizes* the fixed-sample-size approach, it can be no worse than F. The question is how substantial an advantage can be obtained in principle? This is left largely unanswered by our empirical results, and remains an interesting open topic for future research.

For $\langle X = \mathbb{R}^{10}, C = \text{halfspaces}, \epsilon = 0.01, \delta = 0.05 \rangle$:

${\bf Sufficient:}$	T_{BEHW}	=	91,030
Improved:	$T_{\scriptscriptstyle STAB}$	=	15,981
Folklore:	T_{thumb}	\approx	1, 100
Necessary:	$t_{\scriptscriptstyle EHKV}$	=	32

After 100 trials, Procedure S used:

${ m avg} \; T_{ m S}$	=	3,402
$\max T_{S}$	=	5,155
$\min T_{\mathbf{S}}^{T}$	=	2,267

Table 1: A direct comparison of training sample sizes for the pac-learning problem $\langle \mathbb{R}^{10}, \mathsf{halfspaces}, \epsilon = 0.01, \delta = 0.05 \rangle$.

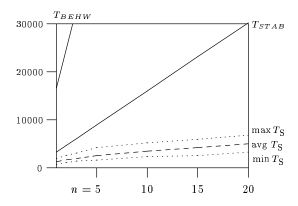


Figure 5: Scaling in input dimension n. Number of training examples observed for $\langle \mathbb{R}^n$, halfspaces, $\epsilon = 0.01$, $\delta = 0.05 \rangle$ with n = 1, 2, 3, 5, 10, 15, 20. (Results of 100 runs each.)

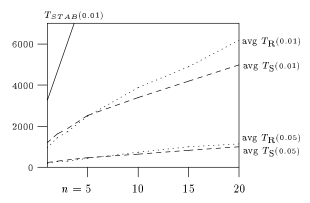


Figure 6: Comparing S versus R. Number of training examples observed for $\langle I\!\!R^n, {\sf halfspaces}, \epsilon, \delta = 0.05 \rangle$ with n=1,2,3,5,10,15,20 and $\epsilon=0.01,~0.05.~ \left(T_{STAB}({\tt 0.05})\right)$ and T_{BEHW} not shown.)

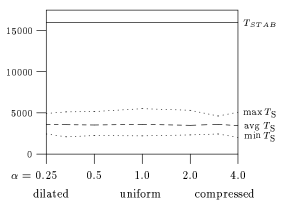


Figure 7: Comparing different domain distributions. Results for $\langle R^{10}$, halfspaces, $\epsilon=0.01$, $\delta=0.05\rangle$ under *pyramidal* transformations of the uniform[-1,1]¹⁰ distribution. *X-axis*: power factor of transformed dot products.

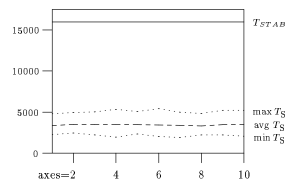


Figure 8: Comparing different target concepts. Results for $\langle I\!\!R^{10}, {\sf halfspaces}, \epsilon = 0.01, \delta = 0.05 \rangle$ with "diagonal" target concepts depending on r = 1, 2, ..., 10 relevant axes.

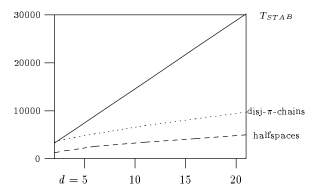


Figure 9: Comparing different concept classes with matching VCdimensions. Average $T_{\rm S}$ for $\langle R^n, C_i, \epsilon=0.01, \delta=0.05\rangle$ with C_1 = halfspaces, C_2 = disj- π -chains, and VC(C_i) = 2, 3, 4, 6, 11, 16, 21. (The class of disj- π -chains is defined by $\frac{1}{d\epsilon}$ copies of a d-dimensional "product chain" of concepts, where the concepts in different copies are mutually exclusive [Schuurmans, 1995]. This class has VCdimension d.)

Advantages: Despite the empirical nature of these results, sequential learning holds many clear advantages over fixed-sample-size learning for solving pac-learning problems: First, the sequential approach decouples the actual data-efficiency of a pac-learner from the precise bounds we can prove about its performance a priori. Thus, the actual data-efficiency of a sequential learner depends on the specific case at hand, not on what we can prove about the worst case situation. Consequently, the sequential approach automatically takes advantage of beneficial situations like "easy" target concepts and domain distributions [Oblow, 1992], or a "good" hypothesizer that makes lucky guesses — without the system designer having to explicitly notice that these beneficial situations exist a priori! More importantly, the true worst case data-efficiency of sequential learning depends on the true worst case convergence properties of the concept class, not on the particular bounds we happen to be able to prove at the time (i.e., if bad concepts are eliminated sooner than proven bounds, then S automatically stops sooner). So, in effect, we are able to exploit the optimal worst case bounds right now, even though we are unable to prove exactly what they are.

Computation: We also note that Procedure S only introduces reasonable computational overhead over Procedure F; and in fact, is often more computationally-efficient than R! Although, at first glance, S appears to be extremely space-inefficient, this rarely amounts to a significant expense in practical applications. The point is that, in practice, it is the task of finding consistent hypotheses (calling H) that takes most of the work—storing hypotheses once found (updating statistics, etc.) does not require much overhead in comparison. Consequently, R is often slower than S (even though it uses less space) simply because R tends to call H more often.

4 Additional results

Special cases: We have obtained even stronger results in slightly restricted settings [Schuurmans and Greiner, 1995]. For example, a variant of Procedure S can serve as a sequential "mistake bounded to pac" conversion procedure that is provably more efficient than Littlestone's fixed-sample-size procedure [Littlestone, 1989] (and which uses $30 \ times$ fewer training examples in empirical tests). We also obtain stronger improvements for the case of distribution specific pac-learning (where we assume the learner $knows\ P_x$, but not the target concept $c \in C$). Notice that a sequential approach is still possible in this case; and, in fact, a variant of Procedure S can pac-learn concept spaces (C, P_x) using 5 times fewer training examples than the best known fixed-sample-size procedure developed in [Benedek and Itai, 1988].

Range of applicability: Beyond improving data-efficiency, sequential learning is also applicable to a much wider range of pac-learning problems than fixed-sample-size learning. For example, Procedure S can be directly applied to "nearest neighbor" and "decision-tree" hypothesizers (like CART [Breiman, et al., 1984]) which implicitly consider concept classes of infinite VCdimension. No fixed-sample-size bound can ever be sufficient

in these cases, and yet Procedure S can be applied to pac-learn these classes "as is." The only catch is that we can no longer place a *uniform* upper bound on S's expected training sample size.⁶

5 Conclusion

Research directions: There are numerous directions for future research. First, since our empirical results address "artificial" learning problems, it would be interesting to test these procedures on "real world" data sets (e.g., as contained in the UCI repository of machine learning databases) to verify that the same empirical advantages can be realized there. Another important research direction is to extend our techniques to deal with classification noise, which remains the main barrier between the results presented here and real applications. Finally, one can also consider a slightly different learning scenario which perhaps has more practical applications than pac-learning: rather than first fixing the accuracy and reliability parameters and then determining a sufficient sample size, it is much more natural to take a fixed sample size, fix a reliability parameter, and produce an estimate of the accuracy achieved by the learner's final hypothesis. In this regard, we are currently investigating a variant of Procedure S which produces hypotheses with small (but reliable!) error estimates.

Contributions: We have described a novel pac-learning procedure, S, that uses far fewer training examples than previous approaches. Procedure S is, in effect, generic test procedure that can pac-learn arbitrary concept classes C (with finite VCdimension), provided only that we can supply a hypothesizer H that produces consistent concepts from C. This procedure introduces little computational overhead, and yet substantially reduces the number of training examples needed to pac-learn in practice — as demonstrated in numerous case studies where S used $many\ times$ fewer training examples than the previous best known approaches, while still maintaining the $exact\ same$ worst case pac-guarantees.

In a way, these results exploit the empirical advantage demonstrated by practical learning algorithms over the theoretical bounds, to improve the efficiency of paclearning. Overall, our results show how pac-learning can be far more efficiently achieved in practice than previously thought — countering the claim that pac-learning can never be feasibly achieved in real applications.

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⁶ It is important not to confuse the idea of sequential with nonuniform pac-learning [Linial, et al., 1991; Oblow, 1992]. Although nonuniform pac-learning procedures also use "on-line" stopping rules very similar to R, they do not share the same theoretical advantages shown for S. Sequential pac-learning seeks to obtain a uniform improvement in data-efficiency for all cases permitted by our prior knowledge, whereas nonuniform pac-learning sacrifices data-efficiency in some situations to obtain an improvement in others. These two concerns are in fact orthogonal.