A Discriminative Model for Semi-Supervised Learning

Maria-Florina Balcan
School of Computer Science, Georgia Institute of Technology
Avrim Blum
Computer Science Department, Carnegie Mellon University

Supervised learning — that is, learning from labeled examples — is an area of Machine Learning that has reached substantial maturity. It has generated general-purpose and practically-successful algorithms and the foundations are quite well understood and captured by theoretical frameworks such as the PAC-learning model and the Statistical Learning theory framework. However, for many contemporary practical problems such as classifying web pages or detecting spam, there is often additional information available in the form of unlabeled data, which is often much cheaper and more plentiful than labeled data. As a consequence, there has recently been substantial interest in semi-supervised learning — using unlabeled data together with labeled data — since any useful information that reduces the amount of labeled data needed can be a significant benefit. Several techniques have been developed for doing this, along with experimental results on a variety of different learning problems. Unfortunately, the standard learning frameworks for reasoning about supervised learning do not capture the key aspects and the assumptions underlying these semi-supervised learning methods.

In this paper we describe an augmented version of the PAC model designed for semi-supervised learning, that can be used to reason about many of the different approaches taken over the past decade in the Machine Learning community. This model provides a unified framework for analyzing when and why unlabeled data can help, in which one can analyze both sample-complexity and algorithmic issues. The model can be viewed as an extension of the standard PAC model where, in addition to a concept class \( C \), one also proposes a compatibility notion: a type of compatibility that one believes the target concept should have with the underlying distribution of data. Unlabeled data is then potentially helpful in this setting because it allows one to estimate compatibility over the space of hypotheses, and to reduce the size of the search space from the whole set of hypotheses \( C \) down to those that, according to one’s assumptions, are a-priori reasonable with respect to the distribution. As we show, many of the assumptions underlying existing semi-supervised learning algorithms can be formulated in this framework.

After proposing the model, we then analyze sample-complexity issues in this setting: that is, how much of each type of data one should expect to need in order to learn well, and what the key quantities are that these numbers depend on. We also consider the algorithmic question of how to efficiently optimize for natural classes and compatibility notions, and provide several algorithmic results including an improved bound for Co-Training with linear separators when the distribution satisfies independence given the label.

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1. INTRODUCTION

In recent years there has been substantial and growing interest in using unlabeled data together with labeled data in machine learning. The motivation is clear: in many applications, unlabeled data can be much cheaper and much more plentiful than labeled data. If useful information can be extracted from unlabeled examples that allows for learning from fewer labeled examples, this can be a substantial benefit. A number of Semi-Supervised learning techniques have been developed for doing this, along with experimental results on a variety of different learning problems. These include label propagation for word-sense disambiguation [Yarowsky 1995], co-training for classifying web pages [Blum and Mitchell 1998] and improving visual detectors [Levin et al. 2003], transductive SVM [Joachims 1999] and EM [Nigam et al. 2000] for text classification, graph-based methods [Zhu et al. 2003c], and others. The problem of learning from labeled and unlabeled data has been the topic of several ICML workshops [Ghani et al. 2003; Amini et al. 2005] as well as a recent book [Chapelle et al. 2006] and survey article [Zhu 2006].

What makes unlabeled data so useful and what many of these methods exploit, is that for a wide variety of learning problems, the natural regularities of the problem involve not only the form of the function being learned by also how this function relates to the distribution of data. For example, in many problems one might expect the target function should cut through low density regions of the space, a property used by the transductive SVM algorithm [Joachims 1999]. In other problems one might expect the target to be self-consistent in some way, a property used by Co-training [Blum and Mitchell 1998]. Unlabeled data is potentially useful in these settings because it then allows one to reduce the search space to a set which is a-priori reasonable with respect to the underlying distribution.

Unfortunately, however, the underlying assumptions of these semi-supervised learning methods are not captured well by standard theoretical models. The main goal of this work is to propose a unified theoretical framework for semi-supervised learning, in which one can analyze when and why unlabeled data can help, and in which one can discuss both sample-complexity and algorithmic issues in a discriminative (PAC-model style) framework.

One difficulty from a theoretical point of view is that standard discriminative learning models do not allow one to specify relations that one believes the target should have with the underlying distribution. In particular, both in the PAC model [Valiant 1984; Blumer et al. 1989; Kearns and Vazirani 1994] and the Statistical Learning Theory framework [Vapnik 1998] there is intentionally a complete
disconnect between the data distribution $D$ and the target function $f$ being learned. The only prior belief is that $f$ belongs to some class $C$: even if the data distribution $D$ is known fully, any function $f \in C$ is still possible. For instance, in the PAC model, it is perfectly natural (and common) to talk about the problem of learning a concept class such as DNF formulas [Linial et al. 1989; Verbeurgt 1990] or an intersection of halfspaces [Baum 1990; Blum and Kannan 1997; Vempala 1997; Klivans et al. 2002] over the uniform distribution; but clearly in this case unlabeled data is useless — you can just generate it yourself. For learning over an unknown distribution, unlabeled data can help somewhat in the standard models (e.g., by allowing one to use distribution-specific algorithms and sample-complexity bounds [Benedek and Itai 1991; Kaarainen 2005]), but this does not seem to capture the power of unlabeled data in practical semi-supervised learning methods. In fact, a recent result of Simon [2009] shows that the information-theoretic advantage provided by knowing the distribution in the standard PAC model is extremely limited.

In generative models, one can easily talk theoretically about the use of unlabeled data, e.g., [Castelli and Cover 1995; 1996]. However, these results typically make strong assumptions that essentially imply that there is only one natural distinction to be made for a given (unlabeled) data distribution. For instance, a typical generative model would be that we assume positive examples are generated by one Gaussian, and negative examples are generated by another Gaussian. In this case, given enough unlabeled data, we could in principle recover the Gaussians and would need labeled data only to tell us which Gaussian is the positive one and which is the negative one.\footnote{[Castelli and Cover 1995; 1996] do not assume Gaussians in particular, but they do assume the distributions are distinguishable, which from this perspective has the same issue.} However, this is too strong an assumption for most real-world settings. Instead, we would like our model to allow for a distribution over data (e.g., documents we want to classify) where there are a number of plausible distinctions we might want to make. In addition, we would like a general framework that can be used to model many different uses of unlabeled data.

1.1 Overview of Our Model

In this paper, we present a PAC-style framework that bridges between these positions and can be used to help think about and analyze many of the ways unlabeled data is typically used. This framework extends the PAC learning model in a way that allows one to express not only the form of target function one is considering, but also relationships that one hopes the target function and underlying distribution will possess. We then analyze both sample-complexity issues—that is, how much of each type of data one should expect to need in order to learn well—as well as algorithmic results in this model. We derive bounds for both the realizable (PAC) and agnostic (statistical learning framework) settings.

We focus for most of this paper on inductive binary classification. In this setting, we assume that our data comes according to a fixed unknown distribution $D$ over an instance space $\mathcal{X}$, and is labeled by some unknown target function $c^* : \mathcal{X} \rightarrow \{0, 1\}$. As in the standard PAC model, in the “realizable case”, we make the assumption that the target is in a given class concept class $C$, whereas in the “agnostic case” we do not make this assumption and instead aim to compete with the best function.
in the given class $C$. A learning algorithm is given a set of labeled examples drawn i.i.d. from $D$ and labeled by $c^*$ as well as a (usually larger) set of unlabeled examples from $D$. The goal is to perform some optimization over these samples and to output a hypothesis that agrees with the target or with the best approximation of the target in $C$ over most of the distribution.

We now describe how we extend the PAC model to capture the kinds of beliefs and assumptions used in semi-supervised learning. The main idea is to augment the PAC notion of a concept class, which is a set of functions (such as linear separators or decision trees), with a notion of compatibility between a function and the data distribution that we hope the target function will satisfy. Rather than talking of “learning a concept class $C$,” we will talk of “learning a concept class $C$ under compatibility notion $\chi$.” For example, suppose we believe there should exist a low-error linear separator, and that furthermore, if the data happens to cluster, then this separator does not slice through the middle of any such clusters. Then we would want a compatibility notion that penalizes functions that do, in fact, slice through clusters. In this framework, the ability of unlabeled data to help depends on two quantities: first, the extent to which the target function indeed satisfies the given assumptions, and second, the extent to which the distribution allows this assumption to rule out alternative hypotheses. For instance, if the data does not cluster at all (say the underlying distribution is uniform in a ball), then all functions would equally satisfy this compatibility notion and the assumption is not useful. From a Bayesian perspective, one can think of this as a PAC model for a setting in which one’s prior is not just over functions, but also over how the function and underlying distribution relate to each other.

To make our model formal, we will need to ensure that the degree of compatibility be something that can be estimated from a finite sample. To do this, we will require that the compatibility notion $\chi$ in fact be a function from $C \times X$ to $[0, 1]$, where the compatibility of a hypothesis $h$ with the data distribution $D$ is then $E_{x \sim D}[\chi(h, x)]$. That is, we require that the degree of incompatibility be a kind of unlabeled loss function, and the incompatibility of a hypothesis $h$ with a data distribution $D$ is a quantity we can think of as an “unlabeled error rate” that measures how a-priori unreasonable we believe some proposed hypothesis to be. For instance, in the example above of a “margin-style” compatibility, we could define $\chi(f, x)$ to be an increasing function of the distance of $x$ to the separator $f$. In this case, the unlabeled error rate, $1 - \chi(f, D)$, is a measure of the probability mass close to the proposed separator. In co-training, where each example $x$ has two “views” ($x = (x_1, x_2)$), the underlying belief is that the true target $c^*$ can be decomposed into functions $\langle c^*_1, c^*_2 \rangle$ over each view such that for most examples, $c^*_1(x_1) = c^*_2(x_2)$. In this case, we can define $\chi(\langle f_1, f_2 \rangle, \langle x_1, x_2 \rangle) = 1$ if $f_1(x_1) = f_2(x_2)$, and 0 if $f_1(x_1) \neq f_2(x_2)$. Then the compatibility of a hypothesis $\langle f_1, f_2 \rangle$ with an underlying distribution $D$ is $Pr_{(x_1, x_2) \sim D}[f_1(x_1) = f_2(x_2)]$.

This framework allows us to analyze the ability of a finite unlabeled sample to reduce our dependence on labeled examples, as a function of (a) the compatibility of the target function (i.e., how correct we were in our assumption) and (b) various measures of the “helpfulness” of the distribution. In particular, in our model, we find that unlabeled data can help in several distinct ways.
(1) If the target function is highly compatible with $D$ and belongs to $C$, then if we have enough unlabeled data to estimate compatibility over all $f \in C$, we can in principle reduce the size of the search space from $C$ down to just those $f \in C$ whose estimated compatibility is high. For instance, if $D$ is “helpful”, then the set of such functions will be much smaller than the entire set $C$ and we thus need many fewer labeled examples to learn well. In the agnostic case, or even in realizable case if the number of labeled examples is severely limited, we can do (unlabeled)-data-dependent structural risk minimization to trade off labeled error and incompatibility.

(2) By providing an estimate of $D$, unlabeled data can allow us to use a more refined distribution-specific notion of “hypothesis space size” such as Annealed VC-entropy [Devroye et al. 1996], Rademacher complexities [Koltchinskii 2001; Bartlett and Mendelson 2002; Boucheron et al. 2005] or the size of the smallest $\epsilon$-cover [Benedek and Itai 1991], rather than VC-dimension [Blumer et al. 1989; Kearns and Vazirani 1994]. In fact, for many natural notions of compatibility we find that the sense in which unlabeled data reduces the “size” of the search space is best described in these distribution-specific measures.

(3) Finally, if the distribution is especially helpful, we may find that not only does the set of compatible $f \in C$ have a small $\epsilon$-cover, but also the elements of the cover are far apart. In that case, if we assume the target function is fully compatible, we may be able to learn from even fewer labeled examples than the $\Omega(1/\epsilon)$ needed just to verify a good hypothesis. For instance, as one application of this, we show that under the assumption of independence given the label, one can efficiently perform Co-Training of linear separators from a single labeled example!

A key feature of our framework is that it allows us to address the issue of how much unlabeled data we should expect to need. Roughly, the “VCdim/$\epsilon^2$” form of standard sample complexity bounds now becomes a bound on the number of unlabeled examples we need to uniformly estimate compatibilities. However, technically, the set whose VC-dimension we now care about is not $C$ but rather a set defined by both $C$ and $\chi$: that is, the overall complexity depends both on the complexity of $C$ and the complexity of the notion of compatibility (see Section 3.1.2). One consequence of our model is that if the target function and data distribution are both well behaved with respect to the compatibility notion, then the sample-size bounds we get for labeled data can substantially beat what one could hope to achieve through pure labeled-data bounds, and we illustrate this with a number of examples through the paper.

We can also talk about a transductive analog of our inductive model where one is given a fixed set $S$ of examples, of which some small random subset is labeled, and the goal is to predict well over $S$. In this case we again express the relationship we hope the target function has with the data through a compatibility notion $\chi$. However, since in this case the compatibility of a given hypothesis is completely determined by $S$ (which is known), we will not need to require that compatibility be an expectation over unlabeled examples. In this case unlabeled data can help in the ways (1) and (3) described above.
1.2 Summary of Main Results

The primary contributions of this paper are three-fold.

1. As described above, we develop a new discriminative (PAC-style) model for semi-supervised learning, that can be used to analyze when unlabeled data can help and how much unlabeled data is needed in order to gain its benefits, as well as the algorithmic problems involved.

2. We present a number of fundamental sample complexity bounds illustrating important and subtle issues in this framework. We present both uniform-convergence results—which apply to any algorithm that is able to find rules of low error and high compatibility—as well as $\epsilon$-cover-based bounds that apply to a more restricted class of algorithms but can be substantially tighter. Our main uniform convergence result (Theorem 11) applies to the fully-agnostic setting and shows how an algorithm that minimizes the sum of empirical error plus a regularization term that is a function of the compatibility structure, will achieve a near-optimal tradeoff and strong generalization guarantees. Our main $\epsilon$-cover bound (Theorem 13) then gives even tighter guarantees for algorithms that first use the unlabeled data to select a small subclass of hypotheses and then optimize over them. We also describe several natural, interesting cases where the set of highly compatible hypotheses is large but it has a small epsilon cover; in these cases, the $\epsilon$-cover-based bounds can apply even though with high probability there still exist high-error hypotheses in the class consistent with the labeled and unlabeled examples.

3. We present several algorithmic results in this model matching the various sample complexity bounds we provide. As a warmup we first present a simple algorithm for learning disjunctions in the fully realizable case for a simple compatibility notion (Theorem 17). Next, more interestingly we show how we can efficiently perform a near-optimal data dependent structural risk minimization in a natural transductive graph-based learning setting (Theorem 18). Finally, we present a new algorithm for Co-Training with linear separators (Theorems 20 and 21) that, if the distribution satisfies independence given the label assumption, requires only a single labeled example to learn to any desired error rate $\epsilon$ and is computationally efficient (i.e., achieves PAC guarantees). This substantially improves on the results of [Blum and Mitchell 1998] which required enough labeled examples to produce an initial weak hypothesis, and in the process we get also a simplification to the noisy halfspace learning algorithm of [Blum et al. 1998].

Our framework has helped analyze many existing semi-supervised learning methods used in practice and has guided the development of new semi-supervised learning algorithms and analyses. We discuss this further in Section 6.1.

1.3 Structure of this Paper

We begin by describing the general setting in which our results apply as well as several examples to illustrate our framework in Section 2. We then give results both for sample complexity (in principle, how much data is needed to learn) and efficient algorithms. In terms of sample-complexity, we start by discussing uniform
convergence results in Section 3.1. For clarity we begin with the case of finite hypothesis spaces in Section 3.1.1, and then discuss infinite hypothesis spaces in Section 3.1.2. We present results for the agnostic case, where one must trade off labeled error and incompatibility, in Section 3.1.3. In Section 3.2 we give results based on the notion of \( \epsilon \)-cover size, which can be substantially tighter though they require algorithms of a specific type (that first use the unlabeled data to choose a small set of “representative” hypotheses and then choose among the representatives based on the labeled data).

In Section 4, we give our algorithmic results. Our main results here are an efficient algorithm for graph-based structural risk minimization, nearly matching the sample complexity bounds in Section 3.1.3, and an efficient algorithm for Co-Training with linear separators, nearly matching the bounds of Section 3.2. In Section 5 we discuss transductive learning, connections with generative models and with other ways of using unlabeled data in machine learning, as well as the relationship between our model and the Luckiness Framework [Shawe-Taylor et al. 1998] developed in the context of supervised learning. Finally, in Section 6 we discuss some implications of our model and present our conclusions, as well a number of open problems.

2. A FORMAL FRAMEWORK

In this section we introduce general notation and terminology we use throughout the paper, and describe our model for semi-supervised learning. In particular, we formally define what we mean by a notion of compatibility and we illustrate it through a number of examples including margins and co-training.

We will focus on binary classification problems. We assume that our data comes according to a fixed unknown distribution \( D \) over an instance space \( \mathcal{X} \), and is labeled by some unknown target function \( c^* : \mathcal{X} \to \{0, 1\} \). A learning algorithm is given a set \( S_L \) of labeled examples drawn i.i.d. from \( D \) and labeled by \( c^* \) as well as a (usually larger) set \( S_U \) of unlabeled examples from \( D \). The goal is to perform some optimization over the samples \( S_L \) and \( S_U \) and to output a hypothesis that agrees with the target over most of the distribution. In particular, the error rate (also called “0 − 1 loss”) of a given hypothesis \( f \) is defined as

\[
\text{err}(f) = \text{err}_D(f) = \Pr_{x \sim D}[f(x) \neq c^*(x)].
\]

For any two hypotheses \( f_1, f_2 \), the distance with respect to \( D \) between \( f_1 \) and \( f_2 \) is defined as

\[
d(f_1, f_2) = d_D(f_1, f_2) = \Pr_{x \sim D}[f_1(x) \neq f_2(x)].
\]

We will use \( \hat{\text{err}}(f) \) to denote the empirical error rate of \( f \) on a given labeled sample (i.e., the fraction of mistakes on the sample) and \( \hat{d}(f_1, f_2) \) to denote the empirical distance between \( f_1 \) and \( f_2 \) on a given unlabeled sample (the fraction of the sample on which they disagree). As in the standard PAC model, a concept class or hypothesis space is a set of functions over the instance space \( \mathcal{X} \). In the “realizable case”, we make the assumption that the target is in a given class \( \mathcal{C} \), whereas in the “agnostic case” we do not make this assumption and instead aim to compete with the best function in the given class \( \mathcal{C} \).

We now formally describe what we mean by a notion of compatibility. A notion of compatibility is a mapping from a hypothesis \( f \) and a distribution \( D \) to \([0, 1]\)
indicating how “compatible” \( f \) is with \( D \). In order for this to be estimable from a finite sample, we require that compatibility be an expectation over individual examples. Specifically, we define:

**Definition 1.** A legal notion of compatibility is a function \( \chi : \mathcal{C} \times \mathcal{X} \rightarrow [0, 1] \) where we (overloading notation) define \( \chi(f, D) = \mathbb{E}_{x \sim D}[\chi(f, x)] \). Given a sample \( S \), we define \( \chi(f, S) \) to be the empirical average of \( \chi \) over the sample.

**Note:** One could also allow compatibility functions over \( k \)-tuples of examples, in which case our (unlabeled) sample-complexity bounds would simply increase by a factor of \( k \).

**Definition 2.** Given compatibility notion \( \chi \), the incompatibility of \( f \) with \( D \) is \( 1 - \chi(f, D) \). We will also call this its unlabeled error rate, \( \text{err}_{\text{unl}}(f) \), when \( \chi \) and \( D \) are clear from context. For a given sample \( S \), we use \( \hat{\text{err}}_{\text{unl}}(f) = 1 - \chi(f, S) \) to denote the empirical average over \( S \).

Finally, we need a notation for the set of functions whose incompatibility is at most some given value \( \tau \).

**Definition 3.** Given value \( \tau \), we define \( \mathcal{C}_{D, \chi}(\tau) = \{ f \in \mathcal{C} : \text{err}_{\text{unl}}(f) \leq \tau \} \). So, e.g., \( \mathcal{C}_{D, \chi}(1) = \mathcal{C} \). Similarly, for a sample \( S \), we define \( \mathcal{C}_{S, \chi}(\tau) = \{ f \in \mathcal{C} : \hat{\text{err}}_{\text{unl}}(f) \leq \tau \} \).

**The transductive case:** In transductive learning, one is given a fixed set \( S \) of examples, of which some small random subset is labeled, and the goal is to predict well over \( S \). That is, we know which examples we will be tested on up front, and so we can view this as a case of learning from a known distribution (the uniform distribution over \( S \)). Since in this case the compatibility of a given hypothesis is completely determined by \( S \) (which is known), we will not need to require that compatibility be an expectation over unlabeled examples. For this setting or for the setting in which \( D \) is actually known in advance, we can drop this requirement entirely and allow any notion of compatibility \( \chi(f, D) \) to be legal.

For convenience, we summarize the notation used throughout the paper in Appendix A. We now give several examples to illustrate this framework:

**Example 1.** Suppose examples are points in \( \mathbb{R}^d \) and \( \mathcal{C} \) is the class of linear separators. A natural belief in this setting is that data should be “well-separated”: not only should the target function separate the positive and negative examples, but it should do so by some reasonable margin \( \gamma \). This is the assumption used by Transductive SVM, also called Semi-Supervised SVM (S3VM) [Joachims 1999; Bie

\[ \chi(f, D) = \mathbb{E}_{(x_1, \ldots, x_k) \sim D^k}[\chi(f, x_1, \ldots, x_k)] \]. In order to estimate this quantity, we simply draw a sample of size \( n \) from \( D \), and partition it into \( n/k \) groups of size \( k \), namely \( (x_i, \ldots, x_i) \) for \( i = 1, \ldots, n/k \); our estimate is then simply the average over those groups, i.e., \( \sum_{i=1}^{n/k} \chi(f, x_i, \ldots, x_i) \).
and Cristianini 2003; Chapelle and Zien 2005]. In this case, if we are given \( \gamma \) up front, we could define \( \chi(f, x) = 1 \) if \( x \) is farther than distance \( \gamma \) from the hyperplane defined by \( f \), and \( \chi(f, x) = 0 \) otherwise. So, the incompatibility of \( f \) with \( D \) is the probability mass within distance \( \gamma \) of the hyperplane \( f \cdot x = 0 \). Alternatively, if we do not want to commit to a specific \( \gamma \) in advance, we could define \( \chi(f, x) \) to be a smooth function of the distance of \( x \) to the separator, as done in [Chapelle and Zien 2005]. Note that in contrast, defining compatibility of a hypothesis based on the largest \( \gamma \) such that \( D \) has probability mass exactly zero within distance \( \gamma \) of the separator would not fit our model: it cannot be written as an expectation over individual examples and indeed would not be a good definition since one cannot distinguish “zero” from “exponentially close to zero” from a small sample of unlabeled data.

Example 2. In co-training [Blum and Mitchell 1998], we assume examples \( x \) each contain two “views”: \( x = (x_1, x_2) \), and our goal is to learn a pair of functions \( \langle f_1, f_2 \rangle \), one on each view. For instance, if our goal is to classify web pages, we might use \( x_1 \) to represent the words on the page itself and \( x_2 \) to represent the words attached to links pointing to this page from other pages. The hope underlying co-training is that the two parts of the example are generally consistent, which then allows the algorithm to bootstrap from unlabeled data. For example, iterative co-training uses a small amount of labeled data to learn some initial information (e.g., if a link with the words “my advisor” points to a page then that page is probably a faculty member’s home page). Then, when it finds an unlabeled example where one side is confident (e.g., the link says “my advisor”), it uses that to label the example for training over the other view. In regularized co-training, one attempts to directly optimize a weighted combination of accuracy on labeled data and agreement over unlabeled data. These approaches have been used for a variety of learning problems, including named entity classification [Collins and Singer 1999], text classification [Nigam and Ghani 2000; Ghani 2001], natural language processing [Pierce and Cardie 2001], large scale document classification [Park and Zhang 2003], and visual detectors [Levin et al. 2003]. As mentioned in Section 1, the assumptions underlying this method fit naturally into our framework. In particular, we can define the incompatibility of some hypothesis \( \langle f_1, f_2 \rangle \) with distribution \( D \) as \( \Pr_{(x_1, x_2) \sim D}[f_1(x_1) \neq f_2(x_2)] \). Similar notions are given in subsequent work of [Rosenberg and Bartlett 2007; Sridharan and Kakade 2008] for other types of learning problems (e.g. regression) and for other loss functions.

Example 3. In transductive graph-based methods, we are given a set of unlabeled examples connected in a graph \( G \), where the interpretation of an edge is that we believe the two endpoints of the edge should have the same label. Given a few labeled vertices, various graph-based methods then attempt to use them to infer labels for the remaining points. If we are willing to view \( D \) as a distribution over edges (a uniform distribution if \( G \) is unweighted), then as in co-training we can define the incompatibility of some hypothesis \( f \) as the probability mass of edges that are cut by \( f \), which then motivates various cut-based algorithms. For instance, if we require \( f \) to be boolean, then the mincut method of [Blum and Chawla 2001] finds the most-compatible hypothesis consistent with the labeled data; if we allow \( f \) to be fractional and define \( 1 - \chi(f, (x_1, x_2)) = (f(x_1) - f(x_2))^2 \), then the algorithm
of [Zhu et al. 2003c] finds the most-compatible consistent hypothesis. If we do not wish to view $D$ as a distribution over edges, we could have $D$ be a distribution over vertices and broaden Definition 1 to allow for $\chi$ to be a function over pairs of examples. In fact, as mentioned in section 2 (right after definition 1), since we have perfect knowledge of $D$ in this setting we can allow any compatibility function $\chi(f, D)$ to be legal. We discuss connections to other graph-based methods in Section 5.1.

Example 4. As a special case of co-training, suppose examples are pairs of points in $R^d$, $C$ is the class of linear separators, and we believe the two points in each pair should both be on the same side of the target function. (So, this is a version of co-training where we require $f_1 = f_2$.) The motivation is that we want to use pairwise information as in Example 3, but we also want to use the features of each data point. For instance, in the word-sense disambiguation problem studied by [Yarowsky 1995], the goal is to determine which of several dictionary definitions is intended for some target word in a piece of text (e.g., is “plant” being used to indicate a tree or a factory?). The local context around each word can be viewed as placing it into $R^d$, but the edges correspond to a completely different type of information: the belief that if a word appears twice in the same document, it is probably being used in the same sense both times. In this setting, we could use the same compatibility function as in Example 3, but rather than having the concept class $C$ be all possible functions, we restrict $C$ to just linear separators.

Example 5. In a related setting to co-training considered by [Leskes 2005], examples are single points in $X$ but we have a pair of hypothesis spaces $\langle C_1, C_2 \rangle$ (or more generally a $k$-tuple $\langle C_1, \ldots, C_k \rangle$), and the goal is to find a pair of hypotheses $\langle f_1, f_2 \rangle \in C_1 \times C_2$ with low error over labeled data and that agree over the distribution. For instance, if data is sufficiently “well-separated”, one might expect there to exist both a good linear separator and a good decision tree, and one would like to use this assumption to reduce the need for labeled data. In this case one could define compatibility of $\langle f_1, f_2 \rangle$ with $D$ as $\Pr_{x \sim D}[f_1(x) = f_2(x)]$, or the similar notions given in [Leskes 2005; Shawe-Taylor 2006].

3. SAMPLE COMPLEXITY RESULTS

We now present several sample-complexity bounds that can be derived in this framework, showing how unlabeled data, together with a suitable compatibility notion, can reduce the need for labeled examples. Our key focus is to understand the fundamental quantities that determine these bounds, in order to better understand what it is about the learning problem one can hope to leverage from unlabeled data.

The high-level structure of all of these results is as follows. First, given enough unlabeled data (where “enough” will be a function of some measure of the complexity of $C$ and possibly of $\chi$ as well), we can uniformly estimate the true compatibilities of all functions in $C$ using their empirical compatibilities over the sample. Then, by using this quantity to give a preference ordering over the functions in $C$, in the realizable case we can reduce “$C$” down to “the set of functions in $C$ whose compatibility is not much larger than the true target function” in bounds for the number of labeled examples needed for learning. We also show how we can do (unlabeled)-data-dependent structural risk minimization to trade off labeled er-
error and incompatibility – this is extremely useful both in the agnostic case and in the realizable case when the number of labeled examples is severely limited. The specific bounds differ in terms of the exact complexity measures used (and a few other issues) and we provide examples illustrating when and how certain complexity measures can be significantly more powerful than others. Moreover, one can prove fallback properties of these procedures — the number of labeled examples required is never much worse than the number of labeled examples required by a standard supervised learning algorithm\footnote{See for example Corollary 12 and the discussion following Corollary 6.}. However, if the assumptions happen to be right, one can significantly benefit by using the unlabeled data.

Since it is immediate to derive their analogues in the simpler transductive setting, we write all our generic bounds in the inductive setting, though some of the examples we consider are transductive.

3.1 Uniform Convergence Bounds

We begin with uniform convergence bounds (later in Section 3.2 we give tighter $\epsilon$-cover bounds that apply to algorithms of a particular form). For clarity, we begin with the case of finite hypothesis spaces where we measure the “size” of a set of functions by just the number of functions in the set. We then discuss several issues that arise when considering infinite hypothesis spaces, such as what is an appropriate measure for the “size” of the set of compatible functions, and the need to account for the complexity of the compatibility notion itself. Note that in the standard PAC model, one typically talks of either the realizable case, where we assume that the target function $c^*$ belongs to $\mathcal{C}$, or the agnostic case where we allow any target function $c^*$ [Kearns and Vazirani 1994]. In our setting, we have the additional issue of unlabeled error rate, and can either make an a-priori assumption that the target function’s unlabeled error is low, or else provide a bound in which our sample size (or error rate) depends on whatever its unlabeled error happens to be. We begin in Sections 3.1.1 and 3.1.2 with bounds for the the setting in which we assume $c^* \in \mathcal{C}$, and then in Section 3.1.3 we consider the agnostic case where we remove this assumption.

3.1.1 Finite hypothesis spaces. We first give a bound for the “doubly realizable” case where we assume $c^* \in \mathcal{C}$ and $err_{unl}(c^*) = 0$.

**Theorem 4.** If $c^* \in \mathcal{C}$ and $err_{unl}(c^*) = 0$, then $m_u$ unlabeled examples and $m_l$ labeled examples are sufficient to learn to error $\epsilon$ with probability $1 - \delta$, where

$$m_u = \frac{1}{\epsilon} \left[ \ln |\mathcal{C}| + \ln \frac{2}{\delta} \right] \quad \text{and} \quad m_l = \frac{1}{\epsilon} \left[ \ln |\mathcal{C}_{D,\chi}(\epsilon)| + \ln \frac{2}{\delta} \right].$$

In particular, with probability at least $1 - \delta$, all $f \in \mathcal{C}$ with $\tilde{err}(f) = 0$ and $\tilde{err}_{unl}(f) = 0$ have $err(f) \leq \epsilon$.

**Proof.** The probability that a given hypothesis $f$ with $err_{unl}(f) > \epsilon$ has $\tilde{err}_{unl}(f) = 0$ is at most $(1 - \epsilon)^{m_u} < \frac{\delta}{2|\mathcal{C}|}$ for the given value of $m_u$. Therefore, by the union bound, the number of unlabeled examples is sufficient to ensure that with probability $1 - \frac{\delta}{2}$, only hypotheses in $\mathcal{C}_{D,\chi}(\epsilon)$ have $\tilde{err}_{unl}(f) = 0$. The number of
labeled examples then similarly ensures that with probability $1 - \frac{\delta}{2}$, none of those whose true error is at least $\epsilon$ have an empirical error of 0, yielding the theorem.

**Interpretation:** If the target function indeed is perfectly correct and compatible, then Theorem 4 gives sufficient conditions on the number of examples needed to ensure that an algorithm that optimizes both quantities over the observed data will, in fact, achieve a PAC guarantee. In section 4.1 we describe an efficient algorithm for learning disjunctions under a natural notion of compatibility matching the bounds in this theorem.

We can think of Theorem 4 as bounding the number of labeled examples we need as a function of the “helpfulness” of the distribution $D$ with respect to our notion of compatibility. That is, in our context, a helpful distribution is one in which $C_{D,\chi}(\epsilon)$ is small, and so we do not need much labeled data to identify a good function among them. We can get a similar bound in the situation when the target function is not fully compatible:

**Theorem 5.** If $c^* \in C$ and $err_{\text{unl}}(c^*) = t$, then $m_u$ unlabeled examples and $m_l$ labeled examples are sufficient to learn to error $\epsilon$ with probability $1 - \delta$, for

$$m_u = \frac{2}{\epsilon^2} \left[ \ln |C| + \ln \frac{4}{\delta} \right] \quad \text{and} \quad m_l = \frac{1}{\epsilon} \left[ \ln |C_{D,\chi}(t + 2\epsilon)| + \ln \frac{2}{\delta} \right].$$

In particular, with probability at least $1 - \delta$, the $f \in C$ that optimizes $err_{\text{unl}}(f)$ subject to $\hat{err}(f) = 0$ has $err(f) \leq \epsilon$.

Alternatively, given the above number of unlabeled examples $m_u$, for any number of labeled examples $m_l$, with probability at least $1 - \delta$, the $f \in C$ that optimizes $\hat{err}_{\text{unl}}(f)$ subject to $\hat{err}(f) = 0$ has

$$err(f) \leq \frac{1}{m_l} \left[ \ln |C_{D,\chi}(err_{\text{unl}}(c^*) + 2\epsilon)| + \ln \frac{2}{\delta} \right]. \tag{1}$$

**Proof.** By Hoeffding bounds, $m_u$ is sufficiently large so that with probability at least $1 - \delta/2$, all $f \in C$ have $|\hat{err}_{\text{unl}}(f) - err_{\text{unl}}(f)| \leq \epsilon$. Thus we have:

$$\{f \in C : \hat{err}_{\text{unl}}(f) \leq t + \epsilon\} \subseteq C_{D,\chi}(t + 2\epsilon).$$

For the first implication, the given bound on $m_l$ is sufficient so that with probability at least $1 - \delta$, all $f \in C$ with $\hat{err}(f) = 0$ and $\hat{err}_{\text{unl}}(f) \leq t + \epsilon$ have $err(f) \leq \epsilon$; furthermore, $\hat{err}_{\text{unl}}(c^*) \leq t + \epsilon$, so such a function $f$ exists. Therefore, with probability at least $1 - \delta$, the $f \in C$ that optimizes $\hat{err}_{\text{unl}}(f)$ subject to $\hat{err}(f) = 0$ has $err(f) \leq \epsilon$, as desired. For the second implication, inequality (1) follows immediately by solving for the labeled estimation-error as a function of $m_l$.

**Interpretation:** Theorem 5 has several implications. Specifically:

1. If we can optimize the (empirical) unlabeled error rate subject to having zero empirical labeled error, then to achieve low true error it suffices to draw a number of labeled examples that depends logarithmically on the number of functions in $C$ whose unlabeled error rate is at most $2\epsilon$ greater than that of the target $c^*$.  

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(2) Alternatively, for any given number of labeled examples \(m_l\), we can provide a bound (given in equation 1) on our error rate that again depends logarithmically on the number of such functions.

(3) If we have a desired maximum error rate \(\epsilon\) and do not know the value of \(err_{unl}(c^*)\) but have the ability to draw additional labeled examples as needed, then we can simply do a standard “doubling trick” on \(m_l\). On each round, we check if the hypothesis \(f\) found indeed has sufficiently low empirical unlabeled error rate, and we spread the “\(\delta\)” parameter across the different runs. See, e.g., Corollary 9 in Section 3.1.2.

Finally, before going to infinite hypothesis spaces, we give a simple Occam-style version of the above bounds for this setting. Given a sample \(S\), let us define \(\text{desc}_S(f) = \ln |C_{S,\chi}(\hat{err}_{unl}(f))|\). That is, \(\text{desc}_S(f)\) is the description length of \(f\) (in “nats”) if we sort hypotheses by their empirical compatibility and output the index of \(f\) in this ordering. Similarly, define \(\epsilon\cdot\text{desc}_D(f) = \ln |C_{D,\chi}(err_{unl}(f) + \epsilon)|\). This is an upper-bound on the description length of \(f\) if we sort hypotheses by an \(\epsilon\)-approximation to their true compatibility. Then we immediately get a bound as follows:

**Corollary 6.** For any set \(S\) of unlabeled data, given \(m_l\) labeled examples, with probability at least \(1 - \delta\), all \(f \in \mathcal{C}\) satisfying \(\hat{err}(f) = 0\) and \(\text{desc}_S(f) \leq \epsilon m_l\) have \(err(f) \leq \epsilon\). Furthermore, if \(|S| \geq \frac{1}{\epsilon^2}\ln |\mathcal{C}| + \ln \frac{2}{\delta}\), then with probability at least \(1 - \delta\), all \(f \in \mathcal{C}\) satisfy \(\text{desc}_S(f) \leq \epsilon \cdot \text{desc}_D(f)\).

**Interpretation:** The point of this bound is that an algorithm can use observable quantities (the “empirical description length” of the hypothesis produced) to determine if it can be confident that its true error rate is low. Furthermore, if we have enough unlabeled data, the observable quantities will be no worse than if we were learning a slightly less compatible function using an infinite-size unlabeled sample.

Note that if we begin with a non-distribution-dependent ordering of hypotheses, inducing some description length \(\text{desc}(f)\), and our compatibility assumptions turn out to be wrong, then it could well be that \(\text{desc}_D(c^*) > \text{desc}(c^*)\). In this case our use of unlabeled data would end up hurting rather than helping. However, notice that by merely interleaving the initial ordering and the ordering produced by \(S\), we get a new description length \(\text{desc}_{\text{new}}(f)\) such that

\[
\text{desc}_{\text{new}}(f) \leq 1 + \min(\text{desc}(f), \text{desc}_S(f)).
\]

Thus, up to an additive constant, we can get the best of both orderings.

Also, if we have the ability to purchase additional labeled examples until the function produced is sufficiently “short” compared to the amount of data, then we can perform the usual stratification and be confident whenever we find a consistent function \(f\) such that

\[
\text{desc}_S(f) \leq \epsilon m_l - \ln \left(\frac{m_l (m_l + 1)}{\delta}\right),
\]

where \(m_l\) is the number of labeled examples seen so far.

### 3.1.2 Infinite hypothesis spaces.

To reduce notation, we will assume in the rest of this paper that \(\chi(f, x) \in \{0, 1\}\) so that \(\chi(f, D) = \Pr_{x \sim D}[\chi(f, x) = 1]\). However,
all our sample complexity results can be easily extended to the general case. Before giving our main theorems, we first discuss some subtle issues that come up when dealing with infinite hypothesis spaces.

The first issue that arises is that in order to achieve uniform convergence of unlabeled error rates, the set whose complexity we care about is not $\mathcal{C}$ but rather the potentially more complex set $\chi(\mathcal{C}) = \{\chi_f : f \in \mathcal{C}\}$ where

$$\chi_f : \mathcal{X} \rightarrow \{0, 1\} \text{ is defined as } \chi_f(x) = \chi(f, x).$$

For instance, suppose examples are just points on the line, and $\mathcal{C} = \{f_a(x) : f_a(x) = 1 \text{ if } x \leq a\}$. In this case, $\text{VCdim}(\mathcal{C}) = 1$. However, we could imagine a compatibility function such that $\chi(f_a, x)$ depends on some complicated relationship between the real numbers $a$ and $x$. In this case, $\text{VCdim}(\chi(\mathcal{C}))$ is much larger, and we would need many more unlabeled examples to estimate compatibility over all of $\mathcal{C}$. Of course, with $\text{VCdim}(\mathcal{C}) = 1$ we do not need to uniformly estimate compatibilities in order to learn from a small number of labeled examples.\(^5\) However in Appendix B we present an explicit lower bound where indeed having an unlabeled sample size depending only on $\text{VCdim}(\mathcal{C})$ and $1/\epsilon$ results in a substantially larger number of labeled examples needed for uniform convergence of labeled error rates. See Theorem 22 in Appendix B.

A second issue is that we need an appropriate measure for the “size” of the set of surviving functions. VC-dimension tends not to be a good choice: for instance, if we consider the case of Example 1 (margins), then even if data is concentrated in two well-separated “blobs”, the set of compatible separators still has as large a VC-dimension as the entire class even though they are all very similar with respect to $D$ (see, e.g., Figure 1 after Theorem 8 below). Instead, it is better to consider distribution dependent complexity measures such as annealed VC-entropy [Devroye et al. 1996] or Rademacher averages [Koltchinskii 2001; Bartlett and Mendelson 2002; Boucheron et al. 2005]. For this we introduce some notation. Specifically, for any $\mathcal{C}$, we denote by $\mathcal{C}[m, D]$ the expected number of splits of $m$ points (drawn i.i.d.) from $D$ using concepts in $\mathcal{C}$. Also, for a given (fixed) $S \subseteq \mathcal{X}$, we will denote by $\mathcal{S}$ the uniform distribution over $S$, and by $\mathcal{C}[m, \mathcal{S}]$ the expected number of splits of $m$ points from $\mathcal{S}$ using concepts in $\mathcal{C}$. The following is the analog of Theorem 5 for the infinite case.

**Theorem 7.** If $c^* \in \mathcal{C}$ and $\text{err}_{unl}(c^*) = t$, then $m_u$ unlabeled examples and $m_l$ labeled examples are sufficient to learn to error $\epsilon$ with probability $1 - \delta$, for

$$m_u = O\left(\frac{\text{VCdim}(\chi(\mathcal{C}))}{\epsilon^2} \ln \frac{1}{\epsilon} + \frac{1}{\epsilon^2} \ln \frac{2}{\delta}\right)$$

and

$$m_l = \frac{2}{\epsilon} \left[\ln \left(2\mathcal{C}_{D,\chi}(t+2\epsilon)[2m_l, D]\right) + \ln \frac{4}{\delta}\right],$$

where recall $\mathcal{C}_{D,\chi}(t+2\epsilon)[2m_l, D]$ is the expected number of splits of $2m_l$ points drawn from $D$ using concepts in $\mathcal{C}$ of unlabeled error rate $\leq t + 2\epsilon$. In particular, with

\(^5\)If by “small” we mean $O(1/\epsilon)$. Later, in Section 3.2, we present results where unlabeled examples can reduce the labeled sample size below even this level.
probability at least 1 − δ, the f ∈ C that optimizes ε̂err\text{unl}(f) subject to ε̂err(f) = 0 has err(f) ≤ ε.

PROOF. Let S be the set of m_u unlabeled examples. By standard VC-dimension bounds (e.g., see Theorem 24 in Appendix D) the number of unlabeled examples given is sufficient to ensure that with probability at least 1 − δ, we have

\[ \Pr_{x \sim \mathcal{S}}[\chi_f(x) = 1] - \Pr_{x \sim \mathcal{D}}[\chi_f(x) = 1] \leq \epsilon \quad \text{for all } \chi_f \in \chi(C). \]

Since \( \chi_f(x) = \chi(f, x) \), this implies that we have

\[ |\varepsilon\hat{r}_{\text{unl}}(f) - \varepsilon r_{\text{unl}}(f)| \leq \epsilon \quad \text{for all } f \in C. \]

So, the set of hypotheses with \( \varepsilon\hat{r}_{\text{unl}}(f) \leq t + \epsilon \) is contained in \( C_{D, \chi}(t + 2\epsilon) \).

The bound on the number of labeled examples now follows directly from known concentration results using the expected number of partitions instead of the maximum in the standard VC-dimension bounds (e.g., see Theorem 25 in Appendix D). This bound ensures that with probability 1 − δ, none of the functions \( f \in C_{D, \chi}(t + 2\epsilon) \) with \( \text{err}(f) \geq \epsilon \) have \( \varepsilon\hat{r}(f) = 0 \).

The above two arguments together imply that with probability 1 − δ, all \( f \in C \) with \( \varepsilon\hat{r}(f) = 0 \) and \( \varepsilon\hat{r}_{\text{unl}}(f) \leq t + \epsilon \) have \( \text{err}(f) \leq \epsilon \), and furthermore \( c^* \) has \( \varepsilon\hat{r}_{\text{unl}}(c^*) \leq t + \epsilon \). This in turn implies that with probability at least 1 − δ, the \( f \in C \) that optimizes \( \varepsilon\hat{r}_{\text{unl}}(f) \) subject to \( \varepsilon\hat{r}(f) = 0 \) has \( \text{err}(f) \leq \epsilon \) as desired. □

We can also give a bound where we specify the number of labeled examples as a function of the unlabeled sample; this is useful because we can imagine our learning algorithm performing some calculations over the unlabeled data and then deciding how many labeled examples to purchase.

**Theorem 8.** If \( c^* \in C \) and \( \text{err}_{\text{unl}}(c^*) = t \), then an unlabeled sample \( S \) of size

\[ O\left( \frac{\max[\text{VCdim}(C), \text{VCdim}(\chi(C))]}{\epsilon^2} \ln \frac{1}{\epsilon} + \frac{1}{\epsilon^2} \ln \frac{2}{\delta} \right) \]

is sufficient so that if we label \( m_l \) examples drawn uniformly at random from \( S \), where

\[ m_l > \frac{4}{\epsilon} \left[ \ln(2C_{S, \chi}(t + \epsilon)[2m_l, \mathcal{S}]) + \ln \frac{4}{\delta} \right] \]

then with probability at least 1 − δ, the \( f \in C \) that optimizes \( \varepsilon\hat{r}_{\text{unl}}(f) \) subject to \( \varepsilon\hat{r}(f) = 0 \) has \( \text{err}(f) \leq \epsilon \).

PROOF. Standard VC-bounds (in the same form as for Theorem 7) imply that the number of labeled examples \( m_l \) is sufficient to guarantee the conclusion of the theorem with “\( \text{err}(f) \)” replaced by “\( \text{err}_{\mathcal{S}}(f) \)” (the error with respect to \( \mathcal{S} \)) and “\( \epsilon \)” replaced with “\( \epsilon/2 \)”. The number of unlabeled examples is enough to ensure that, with probability \( \geq 1 - \delta \), for all \( f \in C \), \( |\text{err}(f) - \text{err}_{\mathcal{S}}(f)| \leq \epsilon/2 \). Combining these two statements yields the theorem. □

Note that if we assume \( \text{err}_{\text{unl}}(c^*) = 0 \), then we can use the set \( C_{S, \chi}(0) \) instead of \( C_{S, \chi}(t + \epsilon) \) in the formula giving the number of labeled examples in Theorem 8.

**Note:** Notice that for the setting of Example 1, in the worst case (over distributions \( D \)) this will essentially recover the standard margin sample-complexity bounds.
for the number of labeled examples. In particular, \( C_{S,\chi}(0) \) contains only those separators that split \( S \) with margin \( \geq \gamma \), and therefore, \( s = \left| C_{S,\chi}(0)[2m_l, S] \right| \) is no greater than the maximum number of ways of splitting \( 2m_l \) points with margin \( \gamma \). However, if the distribution is helpful, then the bounds can be much better because there may be many fewer ways of splitting \( S \) with margin \( \gamma \). For instance, in the case of two well-separated “blobs” illustrated in Figure 1, if \( S \) is large enough, we would have just \( s = 4 \).

Fig. 1. Linear separators with a margin-based notion of compatibility. If the distribution is uniform over two well-separated “blobs” and the unlabeled set \( S \) is sufficiently large, the set \( C_{S,\chi}(0) \) contains only four different partitions of \( S \), shown in the figure as \( f_1, f_2, f_3, \) and \( f_4 \). Therefore, Theorem 8 implies that we only need \( O(1/\epsilon) \) labeled examples to learn well.

Theorem 8 immediately implies the following stratified version, which applies to the case in which one repeatedly draws labeled examples until that number is sufficient to justify the most-compatible hypothesis found.

**Corollary 9.** An unlabeled sample \( S \) of size

\[
O \left( \frac{\max\{ VCdim(\mathcal{C}), VCdim(\chi(\mathcal{C}))\}}{\epsilon^2} \ln \frac{1}{\epsilon} + \frac{1}{\epsilon^2} \ln \frac{2}{\delta} \right)
\]

is sufficient so that with probability \( \geq 1 - \delta \) we have that simultaneously for every \( k \geq 0 \) the following is true: if we label \( m_k \) examples drawn uniformly at random from \( S \), where

\[
m_k > \frac{4}{\epsilon} \left[ \ln (2C_{S,\chi}((k+1)\epsilon)[2m_k, S]) + \ln \frac{4(k+1)(k+2)}{\delta} \right]
\]

then all \( f \in \mathcal{C} \) with \( \hat{err}(f) = 0 \) and \( \hat{err}_{unl}(f) \leq (k+1)\epsilon \) have \( err(f) \leq \epsilon \).

**Interpretation:** This corollary is an analog of Theorem 6 and it justifies a stratification based on the estimated unlabeled error rates. That is, beginning with \( k = 0 \), one draws the specified number of examples and checks to see if a sufficiently compatible hypothesis can be found. If so, one halts with success, and if not, one increments \( k \) and tries again. Since \( k \leq \frac{1}{\epsilon} \), we clearly have a fallback property: the number of labeled examples required is never much worse than the number of labeled examples required by a standard supervised learning algorithm.
If one does not have the ability to draw additional labeled examples, then we can
fix $m_l$ and instead stratify over estimation error as in [Bartlett et al. 1999]. We
discuss this further in our agnostic bounds in Section 3.1.3 below.

3.1.3 The agnostic case. The bounds given so far have been based on the
assumption that the target function belongs to $\mathcal{C}$ (so that we can assume there will
exist $f \in \mathcal{C}$ with $\hat{e}(f) = 0$). One can also derive analogous results for the agnostic
(unrealizable) case, where we do not make that assumption. We first present one
immediate bound of this form, and then show how we can use it in order to trade
off labeled and unlabeled error in a near-optimal way. We also discuss the relation
of this to a common regularization technique used in semi-supervised learning. As
we will see, the differences between these two point to certain potential pitfalls in
the standard regularization approach.

Theorem 10. Let $f_t^* = \arg\min_{f \in \mathcal{C}} [\text{err}(f) | \text{err}_{unl}(f) \leq t]$. Then an unlabeled
sample $S$ of size

$$\mathcal{O} \left( \frac{\max[\text{VCdim}(\mathcal{C}), \text{VCdim}(\chi(\mathcal{C}))]}{\epsilon^2} \ln \frac{1}{\epsilon} + \frac{1}{\epsilon^2} \ln \frac{1}{\delta} \right)$$

and a labeled sample of size

$$m_l \geq \frac{8}{\epsilon^2} \left[ \ln \left( C_{D,\chi}(t+2\epsilon|2m_l, D) \right) + \ln \frac{8}{\delta} \right]$$

is sufficient so that with probability $\geq 1 - \delta$, the $f \in \mathcal{C}$ that optimizes $\hat{e}(f)$ subject
to $\hat{e}(f) \leq t + \epsilon$ has

$$\text{err}(f) \leq \text{err}(f_t^*) + \epsilon + \sqrt{\ln(4/\delta)/(2m_l)} \leq \text{err}(f_t^*) + 2\epsilon.$$  

Proof. The given unlabeled sample size implies that with probability $1 - \delta/4$, all $f \in \mathcal{C}$ have $|\hat{e} - e| \leq \epsilon$, which also implies that $\hat{e}(f_t^*) \leq t + \epsilon$. The labeled sample size, using standard VC bounds (e.g, Theorem 26 in the
Appendix D) imply that with probability at least $1 - \delta/2$, all $f \in C_{D,\chi}(t+2\epsilon)$ have
$\text{err}(f) - \hat{e}(f) \leq \epsilon$. Finally, by Hoeffding bounds, with probability at least $1 - \delta/4$ we have

$$\text{err}(f_t^*) \leq \hat{e}(f_t^*) + \sqrt{\ln(4/\delta)/(2m_l)}.$$ 

Therefore, with probability at least $1 - \delta$, the $f \in \mathcal{C}$ that optimizes $\hat{e}(f)$ subject
to $\hat{e}(f) \leq t + \epsilon$ has

$$\text{err}(f) \leq \hat{e}(f) + \epsilon \leq \hat{e}(f_t^*) + \epsilon \leq \text{err}(f_t^*) + \epsilon + \sqrt{\ln(4/\delta)/(2m_l)} \leq \text{err}(f_t^*) + 2\epsilon,$$

as desired.  \qed

Interpretation: Given a value $t$, Theorem 10 bounds the number of labeled ex-
amples needed to achieve error at most $\epsilon$ larger than that of the best function $f_t^*$ of
unlabeled error rate at most $t$. Alternatively, one can also state Theorem 10 in the
form more commonly used in statistical learning theory: given any number of la-
bled examples $m_l$ and given $t > 0$, Theorem 10 implies that with high probability,
the function $f$ that optimizes $\hat{\epsilon}(f)$ subject to $\hat{\epsilon}(f) \leq t + \epsilon$ satisfies

$$err(f) \leq \hat{\epsilon}(f) + \epsilon_t \leq err(f^*) + \epsilon_t + \sqrt{\frac{\ln(4/\delta)}{2m_l}}$$

where

$$\epsilon_t = \sqrt{\frac{8}{m_l} \ln \left( 8C_{D,\chi(t+2\epsilon)|2m_l, D}/\delta \right)}.$$ 

Note that as usual, there is an inherent tradeoff here between the quality of the comparison function $f^*$, which improves as $t$ increases, and the estimation error $\epsilon_t$, which gets worse as $t$ increases. Ideally, one would like to achieve a bound of

$$\min_t [err(f^*_t) + \epsilon_t] + \sqrt{\ln(4/\delta)/(2m_l)};$$

i.e., as if the optimal value of $t$ were known in advance. We can perform nearly as well as this bound by (1) performing a stratification over $t$ (so that the bound holds simultaneously for all values of $t$) and (2) using an estimate $\hat{\epsilon}_t$ of $\epsilon_t$ that we can calculate from the unlabeled sample and therefore use in the optimization. In particular, letting $f_t = \arg\min_{f \in C} [\hat{\epsilon}(f') : \hat{\epsilon}(f') \leq t]$, we will output $f = \arg\min_{f \in C} [\hat{\epsilon}(f(t)) + \hat{\epsilon}_t]$.

Specifically, given a set $S$ of unlabeled examples and $m_l$ labeled examples, let

$$\hat{\epsilon}_t = \hat{\epsilon}_t (S, m_l) = \sqrt{\frac{24}{m_l} \ln (8C_{S,\chi}(t)[m_l, S])},$$

where we define $C_{S,\chi}(t)[m_l, S]$ to be the number of different partitions of the first $m_l$ points in $S$ using functions in $C_{S,\chi}(t)$, i.e., using functions of empirical unlabeled error at most $t$ (we assume $|S| \geq m_l$). Then we have the following theorem.

**Theorem 11.** Let $f^*_t = \arg\min_{f \in C} [\hat{\epsilon}(f') : \hat{\epsilon}(f') \leq t]$ and define $\hat{\epsilon}(f') = \hat{\epsilon}_t'$ for $t' = \hat{\epsilon}(f')$. Then, given $m_l$ labeled examples, with probability at least $1 - \delta$, the function

$$f = \arg\min_{f \in C} [\hat{\epsilon}(f') + \hat{\epsilon}(f')]$$

satisfies the guarantee that

$$err(f) \leq \min_t [err(f^*_t) + \hat{\epsilon}(f^*_t)] + 5 \sqrt{\frac{\ln(8/\delta)}{m_l}}.$$ 

**Proof.** First we argue that with probability at least $1 - \delta/2$, for all $f' \in C$ we have

$$err(f') \leq \hat{\epsilon}(f') + \hat{\epsilon}(f') + 4 \sqrt{\frac{\ln(8/\delta)}{m_l}}.$$ 

In particular, define $C_0 = C_{S,\chi}(0)$ and inductively for $k > 0$ define $C_k = C_{S,\chi}(t_k)$ for $t_k$ such that $C_k[m_l, S] = 8C_{k-1}[m_l, S]$. (If necessary, arbitrarily order the functions with empirical unlabeled error exactly $t_k$ and choose a prefix such that the size condition holds.) Also, we may assume without loss of generality that $C_0[m_l, S] \geq 1$. 

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Then, using bounds of [Boucheron et al. 2000] (see also Appendix D), we have that with probability at least \(1 - \delta/2^{k+2}\), all \(f' \in C_k \setminus C_{k-1}\) satisfy:

\[
err(f') \leq \hat{err}(f') + \sqrt{\frac{6}{m_l} \ln(C_k[|m_l, S|])} + 4 \sqrt{\frac{1}{m_l} \ln(2^{k+3}/\delta)}
\]

\[
\leq \hat{err}(f') + \sqrt{\frac{6}{m_l} \ln(C_k[|m_l, S|])} + 4 \sqrt{\frac{1}{m_l} \ln(2^k)} + 4 \sqrt{\frac{1}{m_l} \ln(8/\delta)}
\]

\[
\leq \hat{err}(f') + 2 \sqrt{\frac{6}{m_l} \ln(C_k[|m_l, S|])} + 4 \sqrt{\frac{1}{m_l} \ln(8/\delta)}
\]

\[
\leq \hat{err}(f') + \hat{err}(f) + 4 \sqrt{\frac{1}{m_l} \ln(8/\delta)}.
\]

Now, let \(f^* = \arg\min_{f^*} [\hat{err}(f^*) + \hat{err}(f)]\). By Hoeffding bounds, with probability at least \(1 - \delta/2\) we have \(\hat{err}(f^*) \leq err(f^*) + \sqrt{\ln(2/\delta)}/(2m_l)\). Also, by construction we have \(\hat{err}(f) + \hat{err}(f^*) \leq \hat{err}(f^*) + \hat{err}(f^*)\). Therefore with probability at least \(1 - \delta\) we have:

\[
err(f) \leq \hat{err}(f) + \hat{err}(f^*) + \hat{err}(f) + 4 \sqrt{\ln(8/\delta)/m_l}
\]

\[
\leq \hat{err}(f^*) + \hat{err}(f) + 4 \sqrt{\ln(8/\delta)/m_l}
\]

\[
\leq err(f^*) + \hat{err}(f^*) + 4 \sqrt{\ln(8/\delta)/m_l}
\]

as desired. \(\square\)

The above result bounds the error of the function \(f\) produced in terms of the empirical unlabeled error rate of \(f^*\). If our unlabeled sample \(S\) is sufficiently large to estimate all unlabeled error rates to \(\pm \epsilon\), then with high probability we have \(\hat{err}_{unl}(f^*_t) \leq t + \epsilon\), so \(\hat{err}(f_t) \leq \hat{err}(f^*_t)\), and moreover \(C_{S,t}(t + \epsilon) \subseteq C_{S,t}(t + 2\epsilon)\). So, our error term \(\hat{err}(f_t)\) is at most \(\sqrt{\frac{2}{m_l} \ln \left(8C_{S,t}(t + 2\epsilon)[m_l, S]\right)}\). Recall that our ideal error term \(\epsilon_t\) for the case that \(t\) was given to the algorithm in advance, factoring out the dependence on \(\delta\), was \(\sqrt{\frac{2}{m_l} \ln \left(8C_{S,t}(t + 2\epsilon)[2m_l, D]\right)}\). [Boucheron et al. 2000] show that for any class \(\mathcal{C}\), the quantity \(\ln(C[m, S])\) is tightly concentrated about \(\ln(C[m, D])\) (see also Theorem 29 in the Appendix E), so up to multiplicative constants, these two bounds are quite close.

**Interpretation and use of unlabeled error rate as a regularizer:** The above theorem suggests to optimize the sum of the empirical labeled error rate plus an estimation-error bound (regularization function) based on the unlabeled error rate. A common related approach used in practice in machine learning (e.g., [Chapelle et al. 2006]) is to just directly optimize the sum of the two kinds of error: i.e., to find \(\arg\min_f [\hat{err}(f) + \hat{err}_{unl}(f)]\). However, this is not generically justified in our framework, because the labeled and unlabeled error rates are really of different "types". In particular, depending on the concept class and notion of compatibility, a small change in unlabeled error rate could substantially change the size of the
For example, suppose all functions in $C$ have unlabeled error rate $0.6$, except for two: function $f_0$ has unlabeled error rate $0$ and labeled error rate $1/2$, and function $f_{0.5}$ has unlabeled error rate $0.5$ and labeled error rate $1/10$. Suppose also that $C$ is sufficiently large that with high probability it contains some functions $f$ that drastically overfit, giving $\hat{err}(f) = 0$ even though their true error is close to $1/2$. In this case, we would like our algorithm to pick out $f_{0.5}$ (since its labeled error rate is fairly low, and we cannot trust the functions of unlabeled error $0.6$). However, even if we use a regularization parameter $\lambda$, there is no way to make $f_{0.5} = \arg\min_f [\hat{err}(f) + \lambda err_{unl}(f)]$: in particular, one cannot have $1/10 + 0.5\lambda \leq \min[1/2 + 0\lambda, 0 + 0.6\lambda]$. So, in this case, this approach will not have the desired behavior.

**Application:** A natural setting where we can efficiently apply the bounds of Theorem 11 is that of learning linear separators in a constant-dimensional space under a margin-based notion of compatibility. Given an unlabeled sample $S$, there are only a polynomial number of partitions of $S$ induced by the class $C$ of linear separators, and for each one we can efficiently find the separator of highest compatibility. This in turn implies that by sampling, we can efficiently estimate the penalty term $\hat{\epsilon}_t$ for any given $t$, and therefore (again because $C$ induces only a polynomial number of distinct partitions of $S$) we can efficiently perform the argmin computation in Theorem 11. In Section 4.2 we demonstrate a more involved use of the bounds of Theorem 11 in a natural graph-based learning setting, where $C$ induces an exponential number of partitions of the dataset. Here, we are able to find a good upper bound on the penalty term $\hat{\epsilon}(f)$ which can be easily computed and has a nice functional form. Moreover, because of this form, we can efficiently perform the argmin computation solving for the hypothesis of smallest empirical error plus penalty, without needing to use a surrogate loss for the empirical error term as is often done in cases of large hypothesis spaces. See Section 4.2.

**Fallback Guarantees:** We end this section with a corollary pointing out that if we have a purely supervised regularization function, we can combine that with the compatibility-based bound of Theorem 11 to perform nearly as well as the best of the two.

**Corollary 12.** Suppose $\lambda(f)$ is a regularization function such that given $m_l$ labeled examples, with probability at least $1 - \delta$ the hypothesis

$$f = \arg\min_{f'} [\hat{err}(f') + \lambda(f')]$$

satisfies

$$err(f) \leq \min_{f'} [err(f') + \lambda(f')] + g(m_l, \delta)$$

for some function $g$. Also let $h(m, \delta) = 5\sqrt{\ln(8/\delta)/m}$. Then with probability at least $1 - 2\delta$, the hypothesis

$$f = \arg\min_{f'} [\hat{err}(f') + \min[\hat{\epsilon}(f') + h(m_l, \delta), \lambda(f') + g(m_l, \delta)]]$$

satisfies $err(f) \leq \min_{f'} [err(f') + \min[\hat{\epsilon}(f') + h(m_l, \delta), \lambda(f') + g(m_l, \delta)]]$.\[^6\]  

\[^6\] On the other hand, for certain compatibility notions and under certain natural assumptions, one can use unlabeled error rate directly, e.g., see e.g., [Sridharan and Kakade 2008].
Note that one could also derive bounds based on other distribution dependent complexity measures and using other concentration results (see e.g. [Boucheron et al. 2005]).

3.2 \( \epsilon \)-Cover-based Bounds

The results in the previous section are uniform convergence bounds: they provide guarantees for any algorithm that optimizes over the observed data. In this section, we consider stronger bounds based on \( \epsilon \)-covers that apply to algorithms that behave in a specific way: they first use the unlabeled examples to choose a “representative” set of compatible hypotheses, and then use the labeled sample to choose among these. Bounds based on \( \epsilon \)-covers exist in the classical PAC setting, but in our framework these bounds and algorithms of this type are especially natural, and the bounds are often much lower than what can be achieved via uniform convergence. For simplicity, we restrict ourselves in this section to the realizable case. However one can combine ideas in Section 3.1.3 with ideas in this section in order to derive bounds in the agnostic case as well. We first present our generic bounds. In Section 3.2.1 we discuss natural settings in which they can be especially useful, and in Section 3.2.2 we present even tighter bounds for co-training.

Recall that a set \( C \subseteq 2^X \) is an \( \epsilon \)-cover for \( C \) with respect to \( D \) if for every \( f \in C \) there is a \( f' \in C \) which is \( \epsilon \)-close to \( f \). That is, \( \Pr_{x \sim D}(f(x) \neq f'(x)) \leq \epsilon \).

**Theorem 13.** Assume \( c^* \in C \) and let \( p \) be the size of a minimum \( \epsilon \)-cover for \( C \). Let \( m_u \) unlabeled examples and \( m_l \) labeled examples

\[
m_u = O\left( \frac{\max\{VCDim(C), VCDim(\chi(C))\}}{\epsilon^2} \ln \frac{1}{\epsilon} + \frac{1}{\epsilon^2} \ln \frac{2}{\delta} \right) \quad \text{and} \quad m_l = O\left( \frac{1}{\epsilon} \ln \frac{p}{\delta} \right),
\]

we can with probability \( 1 - \delta \) identify a hypothesis \( f \in C \) with \( err(f) \leq 6\epsilon \).

**Proof.** Let \( t = err_{unl}(c^*) \). Now, given the unlabeled sample \( S_U \), define \( C' \subseteq C \) as follows: for every labeling of \( S_U \) that is consistent with some \( f \in C \), choose a hypothesis in \( C \) for which \( err_{unl}(f) \) is smallest among all the hypotheses corresponding to that labeling. Next, we obtain \( C_\epsilon \) by eliminating from \( C' \) those hypotheses \( f \) with the property that \( err_{unl}(f) > t + \epsilon \). We then apply a greedy procedure on \( C_\epsilon \) to obtain \( G_\epsilon = \{g_1, \ldots, g_s\} \), as follows:

1. Initialize \( C_\epsilon^i = C_\epsilon \) and \( i = 1 \).
2. Using the unlabeled sample \( S_U \), determine \( C_\epsilon^{i+1} \) by deleting from \( C_\epsilon^i \) those hypotheses \( f \) with the property that \( \hat{d}(f, i) < 3\epsilon \).
3. If \( C_\epsilon^{i+1} = \emptyset \) then set \( s = i \) and stop; else, increase \( i \) by 1 and goto 1.

We now show that with high probability, \( G_\epsilon \) is a 5\( \epsilon \)-cover of \( \mathcal{C}_{D, \chi}(t) \) with respect to \( D \) and has size at most \( p \). First, our bound on \( m_u \) is sufficient to ensure that with probability \( 1 - \frac{2}{\delta} \), we have (a) \( |\hat{d}(f, g) - d(f, g)| \leq \epsilon \) for all \( f, g \in C \) and (b) \( |err_{unl}(f) - err_{unl}(g)| \leq \epsilon \) for all \( f, g \in C \). Let us assume in the remainder that this (a) and (b) are indeed satisfied. Now, (a) implies that any two functions in \( C \) that agree on \( S_U \) have distance at most \( \epsilon \), and therefore \( C' \) is an \( \epsilon \)-cover of \( C \). Using
(b), this in turn implies that $C_\epsilon$ is an $\epsilon$-cover for $C_{D,\chi}(t)$. By construction, $G_\epsilon$ is a 3$\epsilon$-cover of $C_\epsilon$ with respect to distribution $\mathbb{S}_D$, and thus (using (a)) $G_\epsilon$ is a 4$\epsilon$-cover of $C_\epsilon$ with respect to $D$, which implies that $G_\epsilon$ is a 5$\epsilon$-cover of $C_{D,\chi}(t)$ with respect to $D$.

We now argue that $G_\epsilon$ has size at most $p$. Fix some optimal $\epsilon$-cover $\{f_1, \ldots, f_p\}$ of $C_{D,\chi}(err_{unl}(\epsilon^* + 2\epsilon))$. Consider function $g_i$ and suppose that $g_i$ is covered by $f_{\sigma(i)}$. Then the set of functions deleted in step (2) of the procedure include those functions $f$ satisfying $d(g_i, f) < 2\epsilon$ which by triangle inequality includes those satisfying $d(f_{\sigma(i)}, f) \leq \epsilon$. Therefore, the set of functions deleted include those covered by $f_{\sigma(i)}$ and so for all $j > i$, $\sigma(j) \neq \sigma(i)$; in particular, $\sigma$ is 1-1. This implies that $G_\epsilon$ has size at most $p$.

Finally, to learn $\epsilon^*$ we simply output the function $f \in G_\epsilon$ of lowest empirical error over the labeled sample. By Chernoff bounds, the number of labeled examples is enough to ensure that with probability $\geq 1 - \frac{\delta}{\epsilon}$ the empirical optimum hypothesis in $G_\epsilon$ has true error at most $6\epsilon$. This implies that overall, with probability $\geq 1 - \delta$, we find a hypothesis of error at most $6\epsilon$. □

Note that Theorem 13 relies on knowing a good upper bound on $err_{unl}(\epsilon^*)$. If we do not have such an upper bound, then one can perform a stratification as in Sections 3.1.2 and 3.1.3. For example, if we have a desired maximum error rate $\epsilon$ and we do not know a good upper bound for $err_{unl}(\epsilon^*)$ but we have the ability to draw additional labeled examples as needed, then we can simply run the procedure in Theorem 13 for various values of $p$, testing on each round to see if the hypothesis $f$ found indeed has zero empirical labeled error rate. One can show that $m_l = O\left(\frac{1}{\epsilon^2}\ln \frac{1}{\delta}\right)$ labeled examples are sufficient in total for all the “validation” steps. If the number of labeled examples $m_l$ is fixed, then one can also perform a stratification over the target error $\epsilon$.

### 3.2.1 Some illustrative examples

To illustrate the power of $\epsilon$-cover bounds, we now present two examples where these bounds allow for learning from significantly fewer labeled examples than is possible using uniform convergence.

**Graph-based learning:** Consider the setting of graph-based algorithms (e.g., Example 3). In particular, the input is a graph $G$ where each node is an example and $C$ is the class of all boolean functions over the nodes of $G$. Let us define the incompatibility of a hypothesis to be the fraction of edges in $G$ cut by it. Suppose now that the graph $G$ consists of two cliques of $n/2$ vertices, connected together by $en^2/4$ edges. Suppose the target function $\epsilon^*$ labels one of the cliques as positive and one as negative, so the target function indeed has unlabeled error rate less than $\epsilon$. Now, given any set $S_L$ of $m_l < en/4$ labeled examples, there is always a highly-compatible hypothesis consistent with $S_L$ that just separates the positive points in $S_L$ from the entire rest of the graph: the number of edges cut will be at most $nm_l < en^2/4$. However, such a hypothesis has true error nearly 1/2 since it has

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7 Specifically, note that as we increase $t$ (our current estimate for the unlabeled error rate of the target function), the associated $p$ (which is an integer) increases in discrete jumps, $p_1, p_2, \ldots$. We can then simply spread the “$\delta$” parameter across the different runs, in particular run $i$ would use $\delta/(i + 1)$. Since $p_i \geq i$, this implies that $m_l = O\left(\frac{1}{\epsilon^2}\ln \frac{1}{\delta}\right)$ labeled examples are sufficient for all the “validation” steps.
less than $\epsilon n/4$ positive examples. So, we do not yet have uniform convergence over the space of highly compatible hypotheses, since this hypothesis has zero empirical error but high true error. Indeed, this illustrates an overfitting problem that can occur with a direct minimum-cut approach to learning [Blum and Chawla 2001; Joachims 2003; Blum et al. 2004]. On the other hand, the set of functions of unlabeled error rate less than $\epsilon$ has a small $\epsilon$-cover: in particular, any partition of $G$ that cuts less than $\epsilon n^2/4$ edges must be $\epsilon$-close to (a) the all-positive function, (b) the all-negative function, (c) the target function $c^*$, or (d) the complement of the target function $1 - c^*$. So, $\epsilon$-cover bounds act as if the concept class had only 4 functions and so by Theorem 13 we need only $O(\frac{1}{\epsilon^2} \ln \frac{1}{\delta})$ labeled examples to learn well.\(^8\) (In fact, since the functions in the cover are all far from each other, we really need only $O(\ln \frac{1}{\delta})$ examples. This issue is explored further in Theorem 15).

**Simple co-training:** For another case where $\epsilon$-cover bounds can beat uniform-convergence bounds, imagine examples are pairs of points in \{0,1\}^d, $C$ is the class of linear separators, and compatibility is determined by whether both points are on the same side of the separator (i.e., the case of Example 4). Now suppose for simplicity that the target function just splits the hypercube on the first coordinate, and the distribution is uniform over pairs having the same first coordinate (so the target is fully compatible). We then have the following.

**Theorem 14.** Given poly($d$) unlabeled examples $S_U$ and $\frac{1}{\epsilon} \log_2 d$ labeled examples $S_L$, with high probability there will exist functions of true error $1/2 - 2^{-\frac{1}{2} \sqrt{d}}$ that are consistent with $S_L$ and compatible with $S_U$.

**Proof.** Let $V$ be the set of all variables (not including $x_1$) that (a) appear in every positive example of $S_L$ and (b) appear in no negative example of $S_L$. In other words, these are variables $x_i$ such that the function $f(x) = x_i$ correctly classifies all examples in $S_L$. Over the draw of $S_L$, each variable has a $(1/2)^{|S_L|} = 1/\sqrt{d}$ chance of belonging to $V$, so the expected size of $V$ is $(d-1)/\sqrt{d}$ and so by Chernoff bounds, with high probability $V$ has size at least $\frac{1}{2} \sqrt{d}$. Now, consider the hypothesis corresponding to the conjunction of all variables in $V$. This correctly classifies the examples in $S_L$, and with probability at least $1 - 2|S_U|2^{-|V|}$ it classifies every other example in $S_U$ negative because each example in $S_U$ has only a $1/2^{|V|}$ chance of satisfying every variable in $V$. Since $|S_U| = \text{poly}(d)$, this means that with high probability this conjunction is compatible with $S_U$ and consistent with $S_L$, even though its true error is at least $1/2 - 2^{-\frac{1}{2} \sqrt{d}}$. \(\square\)

So, given only a set $S_U$ of poly($d$) unlabeled examples and a set $S_L$ of $\frac{1}{\epsilon} \log_2 d$ labeled examples we would not want to use a uniform convergence based algorithm since we do not yet have uniform convergence. In contrast, the cover-size of the set of functions compatible with $S_U$ is constant, so $\epsilon$-cover based bounds again allow learning from just only $O(\frac{1}{\epsilon^2} \ln \frac{1}{\delta})$ labeled examples (Theorem 13). In fact as we show in Theorem 15 we only need $O\left(\log_2 \frac{1}{\delta} \right)$ labeled examples in this case.

\(^8\)Effectively, $\epsilon$-cover bounds allow one to rule out a hypothesis that, say, just separates the positive points in $S_L$ from the rest of the graph by noting that this hypothesis is very close (with respect to $D$) to the all-negative hypothesis, and that hypothesis has a high labeled-error rate.
3.2.2 Learning from even fewer labeled examples. In some cases, unlabeled data can allow us to learn from even fewer labeled examples than given by Theorem 13. In particular, consider a co-training setting where the target \( c^* \) is fully compatible and \( D \) satisfies the property that the two views \( x_1 \) and \( x_2 \) are conditionally independent given the label \( c^*(x_1, x_2) \). As shown by [Blum and Mitchell 1998], one can boost any weak hypothesis from unlabeled data in this setting (assuming one has enough labeled data to produce a weak hypothesis). Related sample complexity results are given in [Dasgupta et al. 2001]. In fact, we can use the notion of \( \epsilon \)-covers to show that we can learn from just a single labeled example. Specifically, for any concept classes \( C_1 \) and \( C_2 \), we have:

**Theorem 15.** Assume that \( \text{err}(c^*) = \text{err}_{\text{unl}}(c^*) = 0 \) and \( D \) satisfies independence given the label. Then for any \( \tau \leq \epsilon/4 \), using \( m_u \) unlabeled examples and \( m_l \) labeled examples we can find a hypothesis that with probability \( 1 - \delta \) has error at most \( \epsilon \), for

\[
m_u = O\left(\frac{1}{\tau} \left[(V\text{Cdim}(C_1) + V\text{Cdim}(C_2)) \ln \frac{1}{\tau} + \ln \frac{2}{\delta}\right]\right) \quad \text{and} \quad m_l = O\left(\log \frac{1}{\delta}\right).
\]

**Proof.** We will assume for simplicity the setting of Example 3, where \( c^* = c^*_1 = c^*_2 \) and also \( D_1 = D_2 = D \) (the general case is handled similarly, but just requires more notation).

We start by characterizing the hypotheses with low unlabeled error rate. Recall that \( \chi(f, D) = \text{Pr}_{(x_1, x_2) \sim D}[f(x_1) = f(x_2)] \), and for concreteness assume \( f \) predicts using \( x_1 \) if \( f(x_1) \neq f(x_2) \). Consider \( f \in C \) with \( \text{err}_{\text{unl}}(f) \leq \tau \) and let us define \( p_+ = \text{Pr}_{x \in D}[c^*(x) = 0] \), \( p_+ = \text{Pr}_{x \in D}[c^*(x) = 1] \) and for \( i, j \in \{0, 1\} \) define \( p_{ij} = \text{Pr}_{x \in D}[f(x) = i, c^*(x) = j] \). We clearly have \( \text{err}(f) = p_{10} + p_{11} \). From \( \text{err}_{\text{unl}}(f) = \text{Pr}_{(x_1, x_2) \sim D}[f(x_1) \neq f(x_2)] \leq \tau \), using the independence given the label of \( D \), we get

\[
\frac{2p_{10}p_{00}}{p_{10} + p_{00}} + \frac{2p_{11}}{p_{01} + p_{11}} \leq \tau.
\]

In particular, the fact that \( \frac{2p_{10}p_{00}}{p_{10} + p_{00}} \leq \tau \) implies that we cannot have both \( p_{10} > \tau \) and \( p_{00} > \tau \), and the fact that \( \frac{2p_{11}}{p_{01} + p_{11}} \leq \tau \) implies that we cannot have both \( p_{01} > \tau \) and \( p_{11} > \tau \). Therefore, any hypothesis \( f \) with \( \text{err}_{\text{unl}}(f) \leq \tau \) falls in one of the following categories:

1. \( f \) is “close to \( c^* \)”: \( p_{10} \leq \tau \) and \( p_{01} \leq \tau \); so \( \text{err}(f) \leq 2\tau \).
2. \( f \) is “close to \( \overline{c^*} \)”: \( p_{00} \leq \tau \) and \( p_{11} \leq \tau \); so \( \text{err}(f) \geq 1 - 2\tau \).
3. \( f \) “almost always predicts negative”: \( p_{10} \leq \tau \) and \( p_{11} \leq \tau \); so \( \text{Pr}[f(x) = 0] \geq 1 - 2\tau \).
4. \( f \) “almost always predicts positive”: \( p_{00} \leq \tau \) and \( p_{01} \leq \tau \); so \( \text{Pr}[f(x) = 0] \leq 2\tau \).

Let \( f_1 \) be the constant positive function and \( f_0 \) be the constant negative function. Note that our bound on \( m_u \) is sufficient to ensure that with probability \( \geq 1 - \frac{\delta}{2} \), we have (a) \( |\hat{d}(f, g) - d(f, g)| \leq \tau \) for all \( f, g \in C \) and (b) all \( f \in C \) with \( \text{err}_{\text{unl}}(f) = 0 \) satisfy \( \text{err}_{\text{unl}}(f) \leq \tau \). Let us assume in the remainder that this (a) and (b) are indeed satisfied. By our previous analysis, there are at most four kinds of hypotheses...
consistent with unlabeled data: those close to $c^*$, those close to its complement $c^\perp$, those close to $f_0$, and those close to $f_1$. Furthermore, $c^*$, $c^\perp$, $f_0$, and $f_1$ are compatible with the unlabeled data.

So, algorithmically, we first check to see if there exists a hypothesis $g \in \mathcal{C}$ with $\hat{\text{err}}_{\text{unl}}(g) = 0$ such that $d(f_1, g) \geq 3\tau$ and $\hat{d}(f_0, g) \geq 3\tau$. If such a hypothesis $g$ exists, then it must satisfy either case (1) or (2) above. Therefore, we know that one of $\{g, f\}$ is $2\tau$-close to $c^*$. If not, we must have $p_+ \leq 4\tau$ or $p_- \leq 4\tau$, in which case we know that one of $\{f_0, f_1\}$ is $4\tau$-close to $c^*$. So, either way we have a set of two functions, opposite to each other, one of which is at least $4\tau$-close to $c^*$. We finally use $O(\log \frac{1}{\tau})$ labeled examples to pick one of these to output, namely the one with lowest empirical labeled error. Lemma 16 below then implies that with probability $1 - \delta$ the function we output has error at most $4\tau \leq \epsilon$. □

**Lemma 16.** Consider $\tau < \frac{1}{8}$. Let $C_\tau = \{f, f\}$ be a subset of $\mathcal{C}$ containing two opposite hypotheses with the property that one of them is $\tau$-close to $c^*$. Then, $m_l > 6 \log \left(\frac{1}{\delta} \right) \left(\frac{1}{\tau}\right)$ labeled examples are sufficient so that with probability $\geq 1 - \delta$, the concept in $C_\tau$ that is $\tau$-close to $c^*$ in fact has lower empirical error.

**Proof.** See Appendix E. □

In particular, by reducing $\tau$ to $\text{poly}(\delta)$ in Theorem 15, we can reduce the number of labeled examples needed $m_l$ to one. Note however that we will need polynomially more unlabeled examples.

In fact, the result in Theorem 15 can be extended to the case that $D^+$ and $D^-$ merely satisfy constant expansion rather than full independence given the label, see [Balcan et al. 2004].

**Note:** Theorem 15 illustrates that if data is especially well behaved with respect to the compatibility notion, then our bounds on labeled data can be extremely good. In Section 4.3, we show for the case of linear separators and independence given the label, we can give efficient algorithms, achieving the bounds in Theorem 15 in terms of labeled examples by a polynomial time algorithm. Note, however, that both these bounds rely heavily on the assumption that the target is fully compatible. If the assumption is more of a “hope” than a belief, then one would need an additional sample of $1/\epsilon$ labeled examples just to validate the hypothesis produced.

4. **ALGORITHMIC RESULTS**

In this section we give several examples of efficient algorithms in our model that are able to take advantage of unlabeled data and learn using sample sizes comparable to those described in Section 3. We begin in Section 4.1 by considering the problem of learning disjunctions in the doubly realizable case for a simple compatibility notion, presenting an algorithm achieving the sample-size bounds in Section 3.1.1. Next in Section 4.2 we give a more interesting example showing how we can efficiently perform data dependent structural risk minimization in a natural transductive graph-based learning setting, illustrating the bounds of Section 3.1.3. Finally, in section 4.3 we present a new efficient algorithm for Co-Training with linear separators that, if the distribution satisfies independence given the label, requires only a single labeled example to learn to any desired error rate $\epsilon$, matching the $\epsilon$-cover labeled sample bounds in Section 3.2.
Note that our focus is on achieving a low-error hypothesis (also called minimizing 0-1 loss). Another common practice in machine learning (both for supervised and semi-supervised learning) is to instead try to minimize a surrogate convex loss that is easier to optimize [Chapelle et al. 2006]. While this does simplify the computational problem, it does not in general solve the true goal of achieving low error since in general the minimizer of the surrogate loss can have a large 0-1 loss.

4.1 An efficient semi-supervised algorithm for disjunctions

We give here a simple example to illustrate the bounds in Section 3.1.1, and for which we can give a polynomial-time algorithm that takes advantage of them. Let the instance space $\mathcal{X} = \{0,1\}^d$, and for $x \in \mathcal{X}$, let $\text{vars}(x)$ be the set of variables set to 1 by $x$. Let $\mathcal{C}$ be the class of monotone disjunctions (e.g., $x_1 \lor x_3 \lor x_6$), and for $f \in \mathcal{C}$, let $\text{vars}(f)$ be the set of variables disjoined by $f$. Now, suppose we say an example $x$ is compatible with function $f$ if either $\text{vars}(x) \subseteq \text{vars}(f)$ or else $\text{vars}(x) \cap \text{vars}(f) = \emptyset$. This is a very strong notion of “margin”: it says, in essence, that every variable is either a positive indicator or a negative indicator, and no example should contain both positive and negative indicators.

We assume the “doubly realizable” case where we assume $c^* \in \mathcal{C}$ and $\text{err}_{unl}(c^*) = 0$. Given this setup, we can give a simple learning algorithm for this pair $(\mathcal{C}, \chi)$ with sample size bounds matching those in Theorem 4. Specifically, we prove:

**Theorem 17.** The class $\mathcal{C}$ of monotone disjunctions is learnable in polynomial time under the compatibility notion defined above with sample size bounds matching those in Theorem 4.

**Proof.** We begin by using our unlabeled data to construct a graph on $d$ vertices (one per variable), putting an edge between two vertices $i$ and $j$ if there is any example $x$ in our unlabeled sample with $i, j \in \text{vars}(x)$. We now use our labeled data to label the components. If the target function is fully compatible, then no component will get multiple labels (if some component does get multiple labels, we halt with failure). Finally, we produce the hypothesis $f$ such that $\text{vars}(f)$ is the union of the positively-labeled components. This is fully compatible with the unlabeled data and has zero error on the labeled data, so by Theorem 4, if the sizes of the data sets are as given in the bounds, with high probability the hypothesis produced will have error at most $\epsilon$. 

Notice that if we want to view the algorithm as “purchasing” labeled data, then we can simply examine the graph, count the number of connected components $k$, and then request $\frac{1}{\epsilon}[k \ln 2 + \ln \frac{2}{\delta}]$ labeled examples. (Here, $2^k = |\mathcal{C}_{S,\chi}(0)|$.) By the proof of Theorem 4, with high probability $2^k \leq |\mathcal{C}_{D,\chi}(\epsilon)|$, so we are purchasing no more than the number of labeled examples in the theorem statement.

Also, it is interesting to see the difference between a “helpful” and “non-helpful” distribution for this problem. An especially non-helpful distribution would be the uniform distribution over all examples $x$ with $|\text{vars}(x)| = 1$, in which there are $d$ components. In this case, unlabeled data does not help at all, and one still needs $\Omega(d)$ labeled examples (or, even $\Omega\left(\frac{d}{\epsilon}\right)$ if the distribution is non-uniform as in the lower bounds of [Ehrenfeucht et al. 1989]). On the other hand, a helpful distribution is one such that with high probability the number of components is small, such as the case of features appearing independently given the label.
For the semi-agnostic case, where \( c^* \in C \) but \( \text{err}_{\text{unl}}(c^*) > 0 \), the natural approach would be to solve for the consistent hypothesis of minimum unlabeled error over \( S \). Unfortunately, as shown in Appendix C, this problem is NP-hard. A different problem where one can efficiently solve for the most-compatible consistent hypothesis is graph-based learning with a cut-size compatibility notion. Indeed, this is what is done by the min-cut learning algorithm of [Blum and Chawla 2001]. In fact, for this problem one can even do a full data-dependent structural risk minimization, and we describe this in Section 4.2 below.

4.2 Efficient structural risk minimization for graph-based learning

We show here how we can apply the bounds of Theorem 11 in a natural graph-based learning setting to yield an efficient algorithm for structural risk minimization. Interestingly, in this setting not only do we have a good upper bound on the penalty term \( \hat{\epsilon}(f) \) with a nice functional form, but also because of this form, we can efficiently solve for the hypothesis of smallest empirical error plus penalty, without needing to use a surrogate loss for the empirical error term. This allows us to efficiently achieve an agnostic PAC guarantee, getting error nearly as good as the best function of any given compatibility level, with a gap that degrades gracefully with the degree of incompatibility and the number of labeled examples.

We assume our input is an unweighted graph \( G \) over the dataset \( S \). This is a transductive problem (see also Section 5.1): \( S \) is fixed and our goal is to do well with respect to the uniform distribution over \( S \). Our hypothesis class \( C \) consists of all \( 2^{|S|} \) labelings of \( S \), and we consider a cut-size notion of compatibility. Specifically, we define \( \text{err}_{\text{unl}}(f) \) to be the fraction of edges in \( G \) whose endpoints are assigned different labels by \( f \) (i.e., the fraction of edges cut by \( f \)). For ease of discussion, we assume that \( G \) is connected, so the only hypotheses of perfect compatibility are the all-positive and all-negative labelings. Given a random labeled subset of \( S \), our goal in applying Theorem 11 is to identify the hypothesis \( f \) minimizing the sum of its empirical labeled error \( \hat{\text{err}}(f) \) and the regularization term \( \hat{\epsilon}(f) \). We show here how for this problem we can produce a good bound on \( \hat{\epsilon}(f) \), and moreover we can identify the desired hypothesis efficiently, yielding an efficient well-motivated regularized semi-supervised learning algorithm. Specifically, we use Theorem 11 to prove the following result. Let \( t_{\text{min}} \) be the fraction of edges cut by the minimum nontrivial cut in \( G \).

**Theorem 18.** If \( G \) has a partition of true error \( \alpha \) and cutting a \( t \) fraction of edges, then there exists an efficient algorithm that with probability at least \( 1 - \delta \) produces a hypothesis \( f \) of error at most \( \alpha + O \left( \sqrt{\frac{1 + t/t_{\text{min}}}{m_l}} \ln(m_l/\delta) \right) \).

**Proof.** Algorithmically, to use Theorem 11 we have two tasks. The first is to calculate or determine a good upper bound on the regularization term \( \hat{\epsilon}(f) \), and the second is to efficiently find \( \arg\min_f [\hat{\text{err}}(f) + \hat{\epsilon}(f)] \), where \( \hat{\epsilon}(f) \) is our computed upper-bound on \( \hat{\epsilon}(f) \). (Note that if we required \( \hat{\text{err}}(f) = 0 \), i.e. perfect behavior on the given labeled data, we could easily compute the hypothesis minimizing \( \hat{\epsilon}(f) \) by collapsing the labeled positive and negative examples into a source and sink respectively and solving for the minimum cut. However, we instead want to perform a tradeoff.)
For the first task, we can use results of Kleinberg et al. [Kleinberg 2000; Kleinberg et al. 2004] showing that for any connected graph $G$, the VC-dimension of the set of partitions of unlabeled error rate at most $t$ is $O(1 + t/t\text{min})$ where $t\text{min}$ is the fraction of edges cut by the minimum nontrivial cut in $G$.\footnote{Typically one would state this using $t$ and $t\text{min}$ as the number of edges cut rather than the fraction, but we will normalize both here (which doesn’t affect their ratio) to be consistent with the terminology used throughout this paper.} Therefore, by Sauer’s lemma this implies that

$$\ln(C_{S\lambda}(t)[m_i, S]) = \ln(m_i^{O(1+t/t\text{min})}) = O((1 + t/t\text{min})\ln m_i).$$

Notice that this is a graph-dependent quantity since $t\text{min}$ is a function of the graph $G$. We define $\bar{e}(f) = O\left(\sqrt{(1 + t/t\text{min})\ln(m_i)/m_i}\right)$ to be the quantity $\bar{e}(f)$ produced using this upper bound.

For the second task, we need to compute the function $f$ minimizing empirical error plus our estimation error bound. For this we can use the following approach. First, connect a source $v_s$ to all labeled positive examples via edges of weight 1, a sink $v_t$ to all labeled negative examples via edges of weight 1, and assign each edge in $G$ a weight of $\lambda$. If we now find the minimum $v_s-v_t$ cut, this will produce the function $f_\lambda$ minimizing the number of labeled mistakes plus $\lambda$ times the number of unlabeled mistakes (edges cut in $G$). Unfortunately, our estimation error bound $\bar{e}(f)$ is not a linear function of the number of unlabeled mistakes so we cannot directly convert it to a scalar $\lambda$. However, it is a concave function, which means that the argmin of $\bar{e}r(f) + \lambda \bar{e}(f)$ must be the minimizer of $\bar{e}r(f) + \lambda^* err_{\text{val}}(f)$ for some value $\lambda^*$ (see Lemma 19 below). Therefore, we can scan through values of $\lambda$ to determine all distinct functions $f_\lambda$ produced (there are at most $m_i + 1$ such functions since each will have at least one more labeled mistake than the previous) and then output the argmin among these.

Thus, we can use this approach to efficiently solve for our desired hypothesis. This completes the proof. □

**Lemma 19.** Let $S = \{(x_1, y_1), \ldots, (x_n, y_n)\} \subset R^2$, and let $g(y)$ be a concave increasing function. Then the point $(x_i, y_i) \in S$ minimizing $x + g(y)$ must be the point in $S$ minimizing $x + \lambda y$ for some $\lambda > 0$.

**Proof.** Let $S' \subseteq S$ be the set of points in $S$ that minimize $x + \lambda y$ for some $\lambda > 0$. Then $S'$ is exactly the points $(x_k, y_k)$ on the convex hull of $S$ such that for all $j \neq k$, either $x_j > x_k$ or $y_j > y_k$. Now, suppose $(x_i, y_i)$ is the point in $S$ minimizing $x + g(y)$. Then clearly there cannot be $(x_j, y_j) \neq (x_i, y_i)$ such that $x_j < x_i$ and $y_j > y_i$ because then $x_j + g(y_j) < x_i + g(y_i)$ since $g$ is an increasing function. Furthermore, $(x_i, y_i)$ must be on the convex hull of $S$. In particular, suppose not. Then there must be some $(x_j, y_j), (x_k, y_k) \in S$ with $x_i = \alpha x_j + (1 - \alpha)x_k$ for $\alpha \in [0, 1]$ but $y_i > \alpha y_j + (1 - \alpha)y_k$. In that case we have

$$x_i + g(y_i) > \alpha x_j + (1 - \alpha)x_k + g(\alpha y_j + (1 - \alpha)y_k)$$

$$\geq \alpha x_j + (1 - \alpha)x_k + \alpha g(y_j) + (1 - \alpha)g(y_k)$$

$$\geq \min(x_j + g(y_j), x_k + g(y_k)).$$

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where the first “≥” follows from concavity of \( g \). This in turn contradiction the definition of \((x_i, y_i)\). Therefore, \((x_i, y_i)\) must minimize \(x + \lambda y\) for some \(\lambda > 0\).

4.3 Co-training with linear separators: an efficient cover based algorithm

In this section we consider the problem of learning the class \( C \) of linear separators under the Co-training compatibility notion, in the doubly-realizable case. For simplicity we focus first on the case of Example 4: the target function is a linear separator in \( \mathbb{R}^d \) and each example is a pair of points, both of which are assumed to be on the same side of the separator (i.e., an example is a line-segment that does not cross the target hyperplane). We then show how our results can be extended to the more general setting.

Unfortunately, unlike the previous examples, the consistency problem here even with double realizability is NP-hard: given a graph \( G \) embedded in \( \mathbb{R}^d \) with two distinguished points \( s \) and \( t \), it is NP-hard to find the linear separator with \( s \) on one side and \( t \) on the other that cuts the minimum number of edges, even if the minimum is zero [Flaxman 2003]. For this reason, we will make an additional assumption, that the two points in an example are each drawn independently given the label. That is, there is a single distribution \( \bar{D} \) over \( \mathbb{R}^d \), and with some probability \( p_+ \), two points are drawn i.i.d. from \( \bar{D}_+ \) (\( \bar{D} \) restricted to the positive side of the target function) and with probability \( 1 - p_+ \), the two are drawn i.i.d from \( \bar{D}_- \) (\( \bar{D} \) restricted to the negative side of the target function). Note that our sample complexity results in Section 3.2 extend to weaker assumptions such as distributional expansion introduced by [Balcan et al. 2004], but we need true independence for our algorithmic results. [Blum and Mitchell 1998] also give positive algorithmic results for co-training when (a) the two views of an example are drawn independently given the label (which we are assuming now), (b) the underlying function is learnable via Statistical Query algorithms\(^{10}\) (which is true for linear separators [Blum et al. 1998]), and (c) we have enough labeled data to produce a weakly-useful hypothesis (defined below) on one of the views to begin with. We give here an improvement over that result by showing how we can run the algorithm in [Blum and Mitchell 1998] with only a single labeled example, thus obtaining an efficient algorithm in our model. It is worth noticing that in the process, we also somewhat simplify the results of [Blum et al. 1998] on efficiently learning linear separators with noise without a margin assumption.

For the analysis below, we need the following definition. A weakly-useful predictor is a function \( f \) such that for some \( \beta \) that is at least inverse polynomial in the input size we have:

\[
\Pr[f(x) = 1|c^*(x) = 1] > \Pr[f(x) = 1|c^*(x) = 0] + \beta.
\]

(2)

It is equivalent to the usual notion of a “weak hypothesis” [Kearns and Vazirani 1994] when the target function is balanced, but requires the hypothesis give more information when the target function is unbalanced [Blum and Mitchell 1998]. Also, we will assume for convenience that the target separator passes through the origin, and let us denote the separator by \( c^* \cdot x = 0 \).

\(^{10}\)For a detailed description of the Statistical Query model see [Kearns 1998] and [Kearns and Vazirani 1994].
We now describe an efficient algorithm to learn to any desired error rate $\epsilon$ in this setting from just a single labeled example. For clarity, we first describe an algorithm whose running time depends polynomially on both the dimension $d$ and $1/\gamma$, where $\gamma$ is a soft margin of separation between positive and negative examples. Formally, in this case we assume that at least some non-negligible probability mass of examples $x$ satisfy

$$\frac{|x \cdot c^*|}{|x||c^*|} \geq \gamma.$$ 

I.e., they have distance at least $\gamma$ to the separating hyperplane $x \cdot c^* = 0$ after normalization. This is a common type of assumption in machine learning (in fact, often one makes the much stronger assumption that nearly all probability mass is on examples $x$ satisfying this condition). We then show how one can replace the dependence on $1/\gamma$ with instead a polynomial dependence on the number of bits of precision $b$ in the data, using the Outlier Removal Lemma of [Blum et al. 1998] and [Dunagan and Vempala 2001].

**Algorithm 1** Co-training with Linear Separators. The Soft Margin Case.

**Input:** $\epsilon$, $\delta$, $T$, a set $S_L$ of $m_l$ labeled examples drawn i.i.d from $D$, a set $S_U$ of $m_u$ unlabeled examples drawn i.i.d from $D$.

Let $h_p$ be the all-positive function. Let $h_n$ be the all-negative function.

Let $\tau = \epsilon/6$, $\epsilon_1 = \tau/4$.

(1) For $i = 1, \ldots, T$ do
- Choose a random halfspace $f_i$ going through the origin.
- Feed $f_i$, $S_U$, the error parameters $\epsilon_1$, and the confidence parameter $\delta/6$ into the bootstrapping procedure of [Blum and Mitchell 1998] to produce hypothesis $h_i$.

(2) Let $h$ be argmin$_{h_i} \{\hat{e}r_{ unl}(h_i)|d(h, h_p) \geq 3\tau, \hat{d}(h, h_n) \geq 3\tau\}$. If $\hat{e}r_{ unl}(h_i) \geq 3\epsilon_1$, then let $h = h_p$.

(3) Use $S_L$ to output either $h$ or $\bar{h}$: output the hypothesis with lowest empirical error on the set $S_L$.

**Theorem 20.** Assume that at least an $\alpha$ probability mass of examples $x$ have margin $\frac{|x \cdot c^*|}{|x||c^*|} \geq \gamma$ with respect to the target separator $c^*$. There is a polynomial-time algorithm (polynomial in $d$, $1/\gamma$, $1/\alpha$, $1/\epsilon$, and $1/\delta$) to learn a linear separator under the above assumptions to error rate $\epsilon$ with probability $1-\delta$, from a polynomial number of unlabeled examples and a single labeled example.

**Proof.** Let $\epsilon$ and $\delta$ be the desired accuracy and confidence parameters. Let $T = O\left(\frac{1}{\alpha^2 \ln \left(\frac{1}{\epsilon}\right)}\right)$, $m_u = \text{poly}(1/\gamma, 1/\alpha, 1/\epsilon, 1/\delta, d)$, and $m_l = 1$. We run Algorithm 1 with the inputs $\epsilon$, $\delta$, $T$, $S_L$, $S_U$, and $m_l = 1$. Let $\tau = \epsilon/6$, $\epsilon_1 = \tau/4$.

In order to prove the desired result, we start with a few facts.
We first note that our bound on \( m_u \) is sufficient to ensure that with probability 
\[ 1 - \frac{\delta}{3}, \] we have (a) \( |\hat{d}(f, g) - d(f, g)| \leq \tau \) for all \( f, g \in C \) and (b) all \( f \in C \) have 
\[ |err_{\text{unl}}(f) - err_{\text{unl}}(\hat{f})| \leq \epsilon_1. \]

We now argue that if at least an \( \alpha \) probability mass of examples \( x \) have margin 
\[ \frac{|x \cdot c^*|}{\gamma} \geq \gamma \] with respect to the target separator \( c^* \), then a random halfspace has at least a \( \text{poly}(\alpha, \gamma) \) probability of being a weakly-useful predictor. (Note that [Blum et al. 1998] uses the Perceptron algorithm to get weak learning; here, we need something simpler since we need to save our labeled example to the very end.)

Specifically, consider a point \( x \) of margin \( \gamma_x \geq \gamma \). By definition, the margin is the cosine of the angle between \( x \) and \( c^* \), and therefore the angle between \( x \) and \( c^* \) is

\[ \pi/2 - \cos^{-1}(\gamma_x) \leq \pi/2 - \gamma. \]

Now, imagine that we draw \( f \) at random subject to \( f \cdot c^* \geq 0 \) (half of the \( f \)'s will have this property) and define \( f(x) = \text{sign}(f \cdot x) \). Then,

\[ \Pr_f(f(x) \neq c^*(x)|f \cdot c^* \geq 0) \leq (\pi/2 - \gamma)/\pi = 1/2 - \gamma/\pi. \]

Moreover, if \( x \) does not have margin \( \gamma \) then at the very least we have \( \Pr_f(f(x) \neq c^*(x)|f \cdot c^* \geq 0) \leq 1/2. \)

Now define distribution \( D^* = \frac{1}{3}D_+ + \frac{1}{3}D_- \); that is \( D^* \) is the distribution \( D \) but balanced to 50% positive and 50% negative. With respect to \( D^* \) at least an \( \alpha/2 \) probability mass of the examples have margin at least \( \gamma \), and therefore:

\[ E_f[err_{D^*}(f)|f \cdot c^* \geq 0] \leq 1/2 - (\alpha/2)(\gamma/\pi). \]

Since \( err(f) \) is a bounded quantity, by Markov inequality this means that at least an \( \Omega(\alpha \gamma) \) probability mass of functions \( f \) must satisfy

\[ err_{D^*}(f) \leq \frac{1}{2} - \frac{\alpha \gamma}{4 \pi} \]

which in turn implies that they must be useful weakly predictors with respect to \( D \) as defined in Equation (2) with \( \beta = \frac{\alpha \gamma}{4 \pi} \).

The second part of the argument is as follows. Note that in Step(1) of our algorithm we repeat the following process for \( T \) iterations: pick a random \( f_i \), and plug it into the bootstrapping theorem of [Blum and Mitchell 1998] (which, given a distribution over unlabeled pairs \( (x'_1, x'_2) \), will use \( f_i(x'_1) \) as a noisy label of \( x'_2 \), feeding the result into a Statistical Query algorithm). Since \( T = O \left( \frac{1}{\alpha \gamma} \ln \left( \frac{1}{\delta} \right) \right) \), using the above observation about random halfspaces being weak predictors, we obtain that with high probability at least \( 1 - \delta/6 \), at least one of the random hypothesis \( f_i \) was a weakly-useful predictor; and since \( m_u = \text{poly}(1/\gamma, 1/\alpha, 1/\epsilon, 1/\delta, d) \) we also have the associated hypothesis \( h_i \) output by the bootstrapping procedure of [Blum and Mitchell 1998] will with probability at least \( 1 - \delta/6 \) satisfy \( err(h_i) \leq \epsilon_1 \). This implies that with high probability at least \( 1 - 2\delta/3 \), at least one of the hypothesis \( h_i \) we find in Step 1 has true labeled error at most \( \epsilon_1 \). For the rest of the hypotheses we find in Step 1, we have no guarantees.

We now observe the following. First of all, any function \( f \) with small \( err(f) \) must have small \( err_{\text{unl}}(f) \); in particular,

\[ err_{\text{unl}}(f) = \Pr(f(x_1) \neq f(x_2)) \leq 2err(f). \]
This implies that with high probability at least $1 - 2\delta/3$, at least one of the hypotheses $h_i$ we find in Step 1 has true unlabeled error at most $2\epsilon_1$, and therefore empirical unlabeled error at most $3\epsilon_1$. Secondly, because of the assumption of independence given the label, as shown in Theorem 15, with high probability the only functions with unlabeled error at most $\tau$ are functions $2\tau$-close to $c^*$, $2\tau$-close to $-c^*$, $2\tau$-close to the “all positive” function, or $2\tau$-close to the “all negative” function.

In Step (2) we first examine all the hypotheses produced in Step 1, and we pick the hypothesis $h$ with the smallest empirical unlabeled error rate subject to being empirically at least $3\tau$-far from the “all-positive” or “all-negative” functions. If the empirical error rate of this hypothesis $h$ is at most $3\epsilon_1$, we know that its true unlabeled error rate is at most $3\epsilon_1 \leq \tau$, which further implies that either $h$ or $-h$ is $2\tau$ close to $c^*$. However, if the empirical unlabeled error rate of $h$ is greater than $3\epsilon_1$, then we know that the target must be $4\tau$-close to the all-positive or all-negative function so we simply choose $h = “all positive”$ (this is true since the unlabeled sample was large enough so that $|d(f, g) - d(f, g)| \leq \tau$).

So, we have argued that with probability at least $1 - 2\delta/3$ either $h$ or $-h$ is $4\tau$-close to $c^*$. We can now just use $O\left(\log(\frac{1}{\epsilon})\left(\frac{1}{\tau}\right)\right)$ labeled examples to determine which case is which (Lemma 16). This quantity is at most 1 and our error rate is at most $\epsilon$ if we set $\tau \leq \epsilon/4$ and $\tau$ sufficiently small compared to $\delta$. This completes the proof. \(\square\)

The above algorithm assumes one can efficiently pick a random unit-length vector in $R^d$, but the argument easily goes through even if we do this to only $O(ln 1/\gamma)$ bits of precision.

We now extend the result to the case that we make no margin assumption.

**Theorem 21.** There is a polynomial-time algorithm (in $d$, $b$, $1/\epsilon$, and $1/\delta$), where $d$ is the dimension of the space and $b$ is the number of bits per example) to learn a linear separator under the agreement notion of compatibility if the distribution $D$ satisfies independence given the label, from a polynomial number of unlabeled examples and a single labeled example.

**Proof.** We begin by drawing a large unlabeled sample $S$ (of size polynomial in $d$ and $b$). We then compute a linear transformation $T$ that when applied to $S$ has the property that for any hyperplane $w \cdot x = 0$, at least a $1/poly(d, b)$ fraction of $T(S)$ has margin at least $1/poly(d, b)$. We can do this via the Outlier Removal Lemma of [Blum et al. 1998] and [Dunagan and Vempala 2001]. Specifically, the Outlier Removal Lemma states that given a set of points $S$, one can algorithmically remove an $\epsilon'$ fraction of $S$ and ensure that for the remaining set $S'$, for any vector $w$, we have

$$\max_{x \in S'}(w \cdot x)^2 \leq poly(d, b, 1/\epsilon')E_{z \in S'}[(w \cdot x)^2],$$

where $b$ is the number of bits needed to describe the input points. Given such a set $S'$, one can then use its eigenvectors to compute a standard linear transformation (also described in [Blum et al. 1998]) $T : R^d \rightarrow R^{d'}$, where $d' \leq d$ is the dimension of the subspace spanned by $S'$, such that in the transformed space, for all unit-length $w$, we have $E_{z \in T(S')}(w \cdot x)^2] = 1$. In particular, since the maximum of $(w \cdot x)^2$ is
bounded, this implies that for any vector \( w \in \mathbb{R}^d \), at least an \( \alpha \) fraction of points \( x \in T(S') \) have margin at least \( \alpha \) for some \( \alpha \geq 1/poly(b, d, 1/\epsilon') \).

Now, choose \( \epsilon' = \epsilon/4 \), and let \( D' \) be the distribution \( \tilde{D} \) restricted to the space spanned by \( S' \). By VC-dimension bounds, \( |S| = \tilde{O}(d/\alpha) \) is sufficient so that with high probability, (a) \( D' \) has probability mass at least \( 1 - \epsilon/2 \), and (b) the vector \( T(c^*) \) has at least an \( \alpha/2 \) probability mass of \( T(D') \) at margin \( \geq \alpha \). Thus, the linear transformation \( T \) converts the distribution \( D' \) into one satisfying the conditions needed for Theorem 20, and any hypothesis produced with error \( \leq \epsilon/2 \) on \( D' \) will have error at most \( \epsilon \) on \( D \). So, we simply apply \( T \) to \( D' \) and run the algorithm for Theorem 20 to produce a low-error linear separator.

**Note:** We can easily extend our algorithm to the standard co-training setting (where \( c_1^* \) can be different from \( c_2^* \)) as follows: we repeat the procedure in a symmetric fashion, and then just try all combinations of pairs of functions returned to find one of small unlabeled error rate, not close to “all positive”, or “all negative”. Finally we use \( O \left( \log(\frac{1}{\epsilon}) \left( \frac{1}{\delta} \right) \right) \) labeled examples to produce a low error hypothesis (and here we use only one part of the example and only one of the functions in the pair).

5. RELATED MODELS

In this section we further discuss how to incorporate several existing transductive semi-supervised learning approaches in our model, describe connections with generative models and other ways of using unlabeled data in Machine Learning, and discuss the relationship between our model and the luckiness framework of [Shawe-Taylor et al. 1998].

5.1 Transductive Semi-Supervised Learning

We show how a number of existing transductive graph-based learning algorithms can be modeled in our framework. In the following we assume that there is weighted graph \( G \) defined over \( S \), which is given a-priori and encodes the prior knowledge. We denote by \( W \) the weighted adjacency matrix of \( G \) and by \( C \) the set of all binary functions over \( S \).

**Minimum Cut** Suppose for \( f \in C \) we define the incompatibility of \( f \) to be the weight of the cut in \( G \) determined by \( f \). This is the implicit notion of compatibility considered in [Blum and Chawla 2001], and as discussed in Section 4.2, the minimum-cut algorithm used by [Blum and Chawla 2001] finds the most compatible hypothesis that is correct on the labeled data, solving the problem in the semi-agnostic case.

**Normalized Cut** For \( f \in C \) define \( size(f) \) to be the weight of the cut in \( G \) determined by \( f \), and let \( neg(f) \) and \( pos(f) \) be the number of points in \( S \) on which \( f \) predicts negative and positive, respectively. For the normalized cut setting of [Joachims 2003] we can define the incompatibility of \( f \in C_S \) to be \( \frac{size(f)}{neg(f) \cdot pos(f)} \). This is the penalty function used in [Joachims 2003], and again, in the semi-realizable case, algorithmically the goal would be to find a highly compatible hypothesis that is correct on the labeled data. Unfortunately, the corresponding optimization problem is in this case is NP-hard. Still, several approximate solutions have been considered,
leading to different semi-supervised learning algorithms. For instance, Joachims [2003] considers a spectral relaxation that leads to the “SGT algorithm”; another relaxation based on semidefinite programming is considered in [Bie and Cristianini 2004].

Harmonic Function We can also model the algorithms introduced in [Zhu et al. 2003c; 2003a] as follows. If we consider \( f \) to be a probabilistic prediction function defined over \( S \), then we can define the incompatibility of \( f \) to be

\[
\sum_{i,j} w_{i,j} (f(i) - f(j))^2 = f^T L f,
\]

where \( L \) is the un-normalized Laplacian of \( G \). Similarly we can model the algorithm introduced by Zhao et al. [Zhou et al. 2004] by using an incompatibility of \( f \) given by \( ||f||_K = f^T K f \) where \( K \) is a kernel derived from the graph (see for instance [Zhu et al. 2003b]).

5.2 Connections to Generative Models

It is also interesting to consider how generative models can be fit into our model. As mentioned in Section 1, a typical assumption in a generative setting is that \( D \) is a mixture with the probability density function

\[
p(x|\theta) = p_0 \cdot p_0(x|\theta_0) + p_1 \cdot p_1(x|\theta_1)
\]

(see for instance [Ratsaby and Venkatesh 1995; Castelli and Cover 1995; 1996]). In other words, the labeled examples are generated according to the following mechanism: a label \( y \in \{0, 1\} \) is drawn according to the distribution of classes \( \{p_0, p_1\} \) and then a corresponding random feature vector is drawn according to the class-conditional density \( p_y \). The assumption typically used is that the mixture is identifiable. Identifiability ensures that the Bayes optimal decision border \( \{x : p_0 \cdot p_0(x|\theta_0) = p_1 \cdot p_1(x|\theta_1)\} \) can be deduced if \( p(x|\theta) \) is known, and therefore one can construct an estimate of the Bayes border by using \( p(x|\hat{\theta}) \) instead of \( p(x|\theta) \). Essentially once the decision border is estimated, a small labeled sample suffices to learn (with high confidence and small error) the appropriate class labels associated with the two disjoint regions generated by the estimate of the Bayes decision border. To see how we can incorporate this setting in our model, consider for illustration the setting in [Ratsaby and Venkatesh 1995]; there they assume that \( p_0 = p_1 \), and that the class conditional densities are \( d \)-dimensional Gaussians with unit covariance and unknown mean vectors \( \theta_i \in \mathbb{R}^d \). The algorithm used is the following: the unknown parameter vector \( \theta = (\theta_0, \theta_1) \) is estimated from unlabeled data using a maximum likelihood estimate; this determines a hypothesis which is a linear separator that passes through the point \( (\hat{\theta}_0 + \hat{\theta}_1)/2 \) and is orthogonal to the vector \( \hat{\theta}_1 - \hat{\theta}_0 \); finally each of the two decision regions separated by the hyperplane is labeled according to the majority of the labeled examples in the region. Given this setting, a natural notion of compatibility we can consider is the expected log-likelihood function (where the expectation is taken with respect to the unknown distribution specified by \( \theta \)). Specifically, we can identify a legal hypothesis \( f_\theta \) with
the set of parameters $\theta = (\theta_0, \theta_1)$ that determine it, and then we can define

$$\chi(f_{\theta}, D) = E_{x \in D} \log(p(x|\theta)).$$

[Ratsaby and Venkatesh 1995] show that if the unlabeled sample is large enough, then all hypotheses specified by parameters $\theta$ which are close enough to $\theta$, will have the property that their empirical compatibilities will be close enough to their true compatibilities. This then implies (together with other observations about Gaussian mixtures) that the maximum likelihood estimate will be close enough to $\theta$, up to permutations. (This actually motivates $\chi$ as a good compatibility function in our model.)

More generally, we can deal with other parametric families using the same compatibility notion; however, we will need to impose constraints on the distributions allowed in order to ensure that the compatibility is actually well defined (the expected log-likelihood is bounded).

As mentioned in Section 1, this kind of generative setting is really at the extreme of our model. The assumption that the distribution that generates the data is truly a mixture implies that if we knew the distribution, then there are only two possible concepts left (and this makes the unlabeled data extremely useful).

5.3 Connections to the Luckiness Framework

It is worth noticing that there is a strong connection between our approach and the luckiness framework [Shawe-Taylor et al. 1998; Mendelson and Philips 2003]. In both cases, the idea is to define an ordering of hypotheses that depends on the data, in the hope that we will be “lucky” and find that the target function appears early in the ordering. There are two main differences, however. The first is that the luckiness framework (because it was designed for supervised learning only) uses labeled data both for estimating compatibility and for learning: this is a more difficult task, and as a result our bounds on labeled data can be significantly better. For instance, in Example 4 described in Section 2, for any non-degenerate distribution, a dataset of $d^2$ pairs can with probability 1 be completely shattered by fully-compatible hypotheses, so the luckiness framework does not help. In contrast, with a larger (unlabeled) sample, one can potentially reduce the space of compatible functions quite significantly, and learn from $o(d)$ or even $O(1)$ labeled examples depending on the distribution — see Section 3.2 and Section 4. Secondly, the luckiness framework talks about compatibility between a hypothesis and a sample, whereas we define compatibility with respect to a distribution. This allows us to talk about the amount of unlabeled data needed to estimate true compatibility. There are also a number of differences at the technical level of the definitions.

5.4 Relationship to Other Ways of Using Unlabeled Data for Learning

It is well known that when learning under an unknown distribution, unlabeled data might help somewhat even in the standard discriminative models by allowing one to use both distribution-specific algorithms [Benedek and Itai 1991], [Kuariainen 2005], [Sokolovska et al. 2008] and/or tighter data dependent sample-complexity bounds [Bartlett and Mendelson 2002; Koltchinskii 2001]. However in many of these methods one chooses a class of functions or a prior over functions before performing the inference. This does not capture the power of unlabeled data in many practical
semi-supervised learning methods, where typically one has some idea about what the structure of the data tells about the target function, and where the choice of prior can be made more precise after seeing the unlabeled data [Blum and Mitchell 1998; Joachims 1999; Leskes 2005; Rosenberg and Bartlett 2007]. Our focus in this work has been to provide a unified discriminative framework for reasoning about usefulness of unlabeled data in such settings in which one can analyze both sample complexity and algorithmic results.

Another use of unlabeled data that has been studied is for model selection [Schuurmans and Southey 2002]. Here, unlabeled data is used to measure the distance between hypotheses produced at different complexity levels, and serves as a form of “sanity check” to indicate when clear overfitting has occurred. Note that as opposed to our model, in this setting again, the hierarchy (or sequence of nested subclasses) is fixed before the inference and it does not adjust based on the unlabeled data, and in this sense the use of unlabeled data is weaker.

Finally, one other extremely popular setting studied for using unlabeled data in concept learning is that of active learning. As in semi-supervised learning, in active learning the algorithm initially sees only the unlabeled portion of a pool of examples drawn from some underlying distribution. However, in active learning, the algorithm then gets to choose which examples in the pool to have labeled for it rather than just getting labels for a random subset of examples. There have recently been a number of papers on algorithmic results (both theoretical and empirical) and analysis frameworks for active learning – see e.g. [Freund et al. 1993; Tong and Koller. 2001; Dasgupta 2005; Kääriäinen 2006; Balcan et al. 2007; Balcan et al. 2008; Balcan et al. 2009; Dasgupta et al. 2007]. It is worth noting that as opposed to the semi-supervised learning setting, and similarly to the classical supervised learning discriminative settings (PAC and Statistical Learning Theory) the only prior belief about the learning problem in the existing active learning frameworks is that the target function (or a good approximation of it) belongs to a given concept class. For simple concept classes such as linear separators on the line one can achieve an exponential improvement (over the usual supervised learning setting) in the labeled data sample complexity, under no additional assumptions about the learning problem [Cohen et al. 1994; Dasgupta 2005; Balcan et al. 2009]; in general, however, for more complicated concept classes, the speed-ups achievable in the active learning setting depend on the match between the distribution over example-label pairs and the hypothesis class, and often on the target hypothesis in the class [Balcan et al. 2009; Hanneke 2007; Balcan et al. 2008; Friedman 2009; Koltchinskii 2009]. In terms of labeled data savings, in the active learning literature the focus has been in reducing the $1/\epsilon$ term while for most of this paper, with a few exceptions (Theorem 15 and Theorem 20) we have focused on reducing the capacity term.

6. CONCLUSIONS

The formulation of the PAC learning model by Valiant [1984] and the Statistical Learning Theory framework by Vapnik [1982] were instrumental in the development of machine learning and the design and analysis of algorithms for supervised learning. Many modern learning problems, however, call for semi-supervised methods
that can take advantage of large quantities of unlabeled data that are often available, and while a large number of algorithms have been explored, there has been no unifying theoretical framework. In this paper, we develop such a framework that captures many of the ways unlabeled data is typically used, and the fundamental assumptions underlying these approaches. This framework allows one to analyze when and why unlabeled data can help and what the basic quantities are that these data bounds depend on. The high level implication of our analysis is that unlabeled data is useful if (a) we have a good notion of compatibility so that the target function indeed has a low unlabeled error rate, (b) the distribution $D$ is helpful in the sense that not too many other hypotheses also have a low unlabeled error rate, and (c) we have enough unlabeled data to estimate unlabeled error rates well. We then make these statements precise through a series of sample-complexity results, giving bounds as well as identifying the key quantities of interest. In addition, we give several efficient algorithms for learning in this framework. One consequence of our model is that if the target function and data distribution are both well behaved with respect to the compatibility notion, then the sample-size bounds we get can substantially beat what one could hope to achieve using labeled data alone, and we have illustrated this with a number of examples through the paper.

6.1 Subsequent Work

Following the conference version of this paper, several authors have used our framework for reasoning about semi-supervised learning, as well as for developing new algorithms and analyses of semi-supervised learning. For example [Shawe-Taylor 2006; Rosenberg and Bartlett 2007; Ganchev et al. 2008] use it in the context of agreement-based multi-view learning for either classification with specific convex loss functions (e.g., hinge loss) or for regression. Sridharan and Kakade [2008] use our framework in order to provide a general analysis of multi-view learning for a variety of loss functions and learning tasks (classification and regression) along with characterizations of suitable notions of compatibility functions. Parts of this work appear as a book chapter in [Chapelle et al. 2006] and as stated in the introduction of that book, our framework can be used to obtain bounds for a number of the semi-supervised learning methods used in the other chapters.

6.2 Open Problems and Future Directions

Our framework sets up the foundations for a broad range of theoretically fundamental and practically relevant questions, both specific and high-level.

One broad category of such questions is for what natural classes $C$ and compatibility notions $\chi$ can one provide an efficient algorithm that learns the pair $(C, \chi)$ with running time and sample sizes polynomial in the bounds of Theorem 4? For example, a natural question of this form is: can one generalize the algorithm of Section 4.1 to allow for irrelevant variables that are neither positive nor negative indicators? That is, suppose we define a “two-sided disjunction” $h$ to be a pair of disjunctions $(h_+, h_-)$ where $h$ is compatible with $D$ iff for all examples $x$, $h_+(x) = -h_-(x)$ (and let us define $h(x) = h_+(x)$). Can we efficiently learn the class of two-sided disjunctions under this notion of compatibility?

Alternatively, as a different generalization of the problem analyzed in Section 4.1, suppose that again every variable is either a positive or negative indicator, but we
relax the “margin” condition. In particular, suppose we require that every example $x$ either contain at least 60% of the positive indicators and at most 40% of the negative indicators (for positive examples) or vice versa (for negative examples). Can this class be learned efficiently with bounds comparable to those from Theorem 4? Along somewhat different lines, can one generalize the cover-based algorithm given for Co-Training with linear separators, to assume some condition weaker than independence given the label, while maintaining computational efficiency? More broadly, it would be interesting to further extend the development of efficient cover-based algorithms for other interesting concept classes and notions of compatibility.

A particularly interesting consequence of this sample complexity analysis in Section 3.1.3 is that it suggests a new broad class of semi-supervised learning procedures which could greatly improve on the existing (more heuristically justified) regularization based semi-supervised learning procedures. We have exemplified the use of this analysis in the context of graph-based learning algorithms with a cut-size compatibility notion, but the question remains open for other learning problems. An interesting open question here is to design computationally efficient algorithms for learning using theoretically-justified regularization terms for other natural notions of compatibility.

It would also be interesting to analyze an active semi-supervised learning setting. As mentioned in Section 5.4, in the active learning model the only prior typically used is that the target function (or a good approximation to it) belong to a certain concept class, but the algorithm is allowed to query unlabeled examples of its own choice. It would be interesting to combine this interactive capability with the type of assumptions employed by semi-supervised learning, and derive a general theory that captures improvements in labeled data sample complexity both in terms of the $1/\varepsilon$ and of the capacity term.

More broadly, our framework can also be viewed as falling under the general area of learning with data-dependent hypothesis spaces, and it would be interesting to analyze this for settings such as learning with weakly-labeled data or additional information from world knowledge as in [Gabrilovich and Markovitch 2005].

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REFERENCES


\footnote{Note that a number of weaker conditions are known to produce good sample bounds if computational considerations are ignored [Balcan et al. 2004].}


on The Continuum from Labeled to Unlabeled Data in Machine Learning. Washington, DC, USA, 912–912.

A. DIRECTORY OF NOTATION

For convenience, we summarize here the notation used throughout the paper.

<table>
<thead>
<tr>
<th>Notation</th>
<th>Definition and interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>( C )</td>
<td>concept class</td>
</tr>
<tr>
<td>( D )</td>
<td>distribution over unlabeled examples</td>
</tr>
<tr>
<td>( S )</td>
<td>a sample, assumed drawn i.i.d. from ( D )</td>
</tr>
<tr>
<td>( S )</td>
<td>for a sample ( S ), ( S ) is the uniform distribution over ( S )</td>
</tr>
<tr>
<td>( c^* )</td>
<td>the target function</td>
</tr>
<tr>
<td>( \text{err}(f) )</td>
<td>the zero-one error of ( f ), ( \text{err}(f) = \Pr_{x \sim D}[f(x) \neq c^*(x)] )</td>
</tr>
<tr>
<td>( \hat{\text{err}}(f) )</td>
<td>the empirical zero-one error of ( f )</td>
</tr>
<tr>
<td>( \chi )</td>
<td>compatibility notion, ( \chi : C \times X \rightarrow [0, 1] )</td>
</tr>
<tr>
<td>( \chi(f, D) )</td>
<td>the compatibility of ( f ) w.r.t. the underlying distribution ( D )</td>
</tr>
<tr>
<td>( \chi(f, S) )</td>
<td>the compatibility of ( f ) w.r.t ( S )</td>
</tr>
<tr>
<td>( \text{err}_{\text{unl}}(f) )</td>
<td>the unlabeled error rate of ( f ), i.e., ( \text{err}_{\text{unl}}(f) = 1 - \chi(f, D) )</td>
</tr>
<tr>
<td>( \hat{\text{err}}_{\text{unl}}(f) )</td>
<td>the empirical unlabeled error rate of ( f ), i.e., ( \hat{\text{err}}_{\text{unl}}(f) = 1 - \chi(f, S) )</td>
</tr>
<tr>
<td>( C_{D,\chi}(\tau) )</td>
<td>the subset of functions in ( C ) whose incompatibility is at most ( \tau ); i.e., ( C_{D,\chi}(\tau) = { f \in C : \text{err}_{\text{unl}}(f) \leq \tau } )</td>
</tr>
<tr>
<td>( C_{S,\chi}(\tau) )</td>
<td>the subset of functions in ( C ) whose empirical incompatibility is at most ( \tau ); i.e., ( C_{S,\chi}(\tau) = { f \in C : \hat{\text{err}}_{\text{unl}}(f) \leq \tau } )</td>
</tr>
<tr>
<td>( \chi(C) )</td>
<td>the class of functions induced by both ( C ) and ( \chi ) i.e., ( \chi(C) = { \chi_f : f \in C } ) where we define ( \chi_f(x) = \chi(f, x) )</td>
</tr>
<tr>
<td>( C[m, D] )</td>
<td>the expected number of splits of ( m ) points (drawn i.i.d.) from ( D ) using concepts in ( C )</td>
</tr>
<tr>
<td>( C[m, S] )</td>
<td>the expected number of splits of ( m ) points from ( S ) using concepts in ( C )</td>
</tr>
<tr>
<td>( C[m, S] )</td>
<td>the number of different partitions of the first ( m ) points in ( S ) using concepts in ( C )</td>
</tr>
<tr>
<td>( C_{D,\chi}(\tau)[m, D] )</td>
<td>the expected number of splits of ( m ) points drawn i.i.d. from ( D ) using concepts in ( C_{D,\chi}(\tau) )</td>
</tr>
<tr>
<td>( C_{S,\chi}(\tau)[m, S] )</td>
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<td>the number of different partitions of the first ( m ) points in ( S ) using functions in ( C_{S,\chi}(\tau) )</td>
</tr>
</tbody>
</table>

B. UNIFORM CONVERGENCE OF UNLABELED ERROR RATES: A LOWER BOUND

As mentioned in Section 3.1.2, in order to achieve uniform convergence of unlabeled error rates, we need to consider the complexity of the set \( \chi(C) \) and not just the set
C. In this section we present an explicit lower bound giving a class C, compatibility notion χ and distribution D where this lack of uniform convergence of unlabeled error rates indeed results in a substantially larger number of labeled examples being needed for uniform convergence of labeled error rates. Specifically, we prove the following theorem.

**Theorem 22.** There exists a class C, compatibility notion χ, distribution D and target c∗ ∈ C_{D,χ}(0) such that given only poly(VCdim(C), 1/ε) unlabeled examples and o(VCdim(C)/ε) labeled examples, with high probability there exist hypotheses in C of zero empirical labeled error and zero empirical unlabeled error and yet with true error ≥ 2ε. On the other hand, C_{D,χ}(ε)[m, D] = 1, so given sufficient unlabeled examples as in Theorem 7, with high probability all hypotheses in C of zero empirical unlabeled error have true error 0.

**Proof.** We will present a class C, a compatibility notion χ, and a distribution D such that (a) the target function belongs to C_{D,χ}(0), (b) VCdim(χ(C)) is arbitrarily higher than VCdim(C), which itself is arbitrarily higher than ln(C_{D,χ}(ε)[m, D]), and (c) given only poly(VCdim(C), 1/ε) unlabeled examples and o(VCdim(C)/ε) labeled examples, with high probability there exist hypotheses in C of zero empirical labeled error and zero empirical unlabeled error and yet with high true error.

We first need a few preliminaries. Let A_1, A_2, ..., A_N be a collection of subsets of (0, 1) with the following properties: (1) every point z ∈ {0, 1} belongs to half of the sets A_j, and (2) with high probability, the union of a random collection of o(√N) of the sets A_j does not cover all of (0, 1). For example, we can do this by defining A_j to be the set of points in z ∈ {0, 1} such that bit ⌊j/2⌋ in the binary representation of z is equal to j − 1 mod 2; that is, A_1 = (0, 1/2), A_2 = [1/2, 1), A_3 = (0, 1/4) ∪ [1/2, 3/4), A_4 = [1/4, 1/2) ∪ [3/4, 1), and so on. Note that this satisfies property (2) since with high probability a random collection of o(√N) of the sets will not include two sets for the same bit ⌊j/2⌋.

Now, let C be the class of homogeneous linear separators in R^d (i.e., of the form w · x > 0) where w_i ∈ (-1, 1) for all i. We define D to be the distribution that puts 1 − 4ε probability mass on the origin and spreads the remaining 4ε probability mass equally over all points x such that all coordinates of x are equal to 0 except for one x_i ∈ (1, 2); for example, points such as (0, 0, 1, 27, 0). This is much like the typical example used for VC-dimension lower bounds, but with the non-origin points spread out a bit. Finally, we define our compatibility notion as follows:

- If all w_i ≤ 0 or if x is the origin, then χ(w, x) = 1.
- Else, let x_k be the first non-zero coordinate of x (for x in the support of D there is only one) and let w_l be the first positive coordinate of w. So, x_k ∈ (1 + (j − 1)/N, 1 + j/N] for some 1 ≤ j ≤ N. If w_i ∈ A_j then χ(w, x) = 0, else χ(w, x) = 1.

In other words, if not all coordinates in w are negative, then a random non-origin example x will index a random set A_j and thereby have a 50% chance of causing w to make an unlabeled mistake.

The target function has all w_i ≤ 0 (i.e., is all-negative over the support of D), which is perfectly compatible. We now show that this C, χ, and D satisfy the conditions of the theorem.
Let us fix $\epsilon$ and $d = VCdim(C)$. First, for any given polynomial $p$ we can set $N$ sufficiently large so that $p(VCdim(C), 1/\epsilon) = o(\sqrt{N})$. Now notice that in the above example, any hypothesis that is not all negative over the support of $D$ (and therefore has some $w_i \in (0, 1)$) has incompatibility exactly $2\epsilon$ since there is $4\epsilon$ probability mass on examples $x$ that are not the origin, and given that $x$ is not the origin, all sets $A_j$ are equally likely (so, by property (1) of the sets $A_j$, there is a 50% chance that $w_i \in A_j$). Thus, given sufficiently many unlabeled examples as in Theorem 7, we can with high probability reduce $C$ down to a single compatible way of labeling the support of $D$. On the other hand, by property (2) of the sets $A_j$, if we have $o(VCdim(C)/\epsilon)$ unlabeled examples, then with high probability there is some point $z \in (0, 1)$ which is not in the union of the sets $A_j$ chosen so far. This means that, by setting any desired positive weight to $z$, the set $C_{S, \chi}(0)$ contains all ways of labeling the $d$ line segments in the support of $D$. Thus, given $o(VCdim(C)/\epsilon)$ labeled examples, with high probability there remains a hypothesis of zero empirical labeled error and zero empirical unlabeled error and yet with true error $\geq (4\epsilon)/2 = 2\epsilon$.

C. HARDNESS OF COMPATIBILITY OPTIMIZATION

We show here that for the setting of Section 4.1, finding the hypothesis $f \in C$ of minimum empirical unlabeled error subject to having zero empirical labeled error is NP-hard. Recall that here $C$ is the class of disjunctions over $\{0, 1\}^d$, and an example $x$ is compatible with hypothesis $f$ if $\text{vars}(x) \subseteq \text{vars}(f)$ or $\text{vars}(x) \cap \text{vars}(f) = \phi$.

Theorem 23. For the compatibility notion as given in Section 4.1, it is NP-hard to find the disjunction of minimum empirical unlabeled error subject to having zero empirical labeled error.

Proof. We reduce from vertex cover. Given an instance $G = (V, E)$ of vertex cover on $n = d - 1$ vertices, create the following instance of the learning problem. Make one variable for each vertex in $G$ plus one new variable $d$. For each edge $(i, j) \in E$, create a labeled positive example with ones in positions $i$ and $j$. For each $i \in V$, create $n^2$ identical unlabeled examples with ones in positions $i$ and $d$. Finally create a single labeled negative example with a single one in position $d$.

Any disjunction consistent with all the labeled examples must be a vertex cover of $G$ because it has to correctly classify all the positive labeled examples, and it cannot contain variable $d$ since otherwise it would make a mistake on the negative example. Considering now the unlabeled error, each variable in the disjunction creates $n^2$ unlabeled mistakes on the examples with ones in positions $i$ and $d$, plus it makes some number (at most $|E| < n^2$ total) of unlabeled mistakes on the examples corresponding to edges in $G$. Therefore, compatibility is optimized when the disjunction is as small as possible, and so the most-compatible disjunction consistent with the labeled examples is exactly the minimum vertex cover of $G$.

D. STANDARD RESULTS

We state here a few known generalization bounds and concentration results used in our proofs. We start with several classic results. See, e.g., [Devroye et al. 1996].

Theorem 24. Suppose that $C$ is a set of functions from $X$ to $\{0, 1\}$ with finite VC-dimension $V \geq 1$. For any distribution $D$ over $X$, any target function (not
necessarily in \( C \), and any \( \epsilon, \delta > 0 \), if we draw a sample from \( D \) of size

\[
m(\epsilon, \delta; V) = \frac{64}{\epsilon^2} \left( 2V \ln \left( \frac{12}{\epsilon} \right) + \ln \left( \frac{4}{\delta} \right) \right),
\]

then with probability at least \( 1 - \delta \), we have \( |\text{err}(h) - \hat{\text{err}}(h)| \leq \epsilon \) for all \( f \in C \).

**Theorem 25.** Suppose that \( C \) is a set of functions from \( X \) to \( \{0, 1\} \) with finite VC-dimension \( V \geq 1 \). For any probability distribution \( D \) over \( X \), any target function \( c^* \), we have

\[
\Pr_S \left[ \sup_{f \in C, \hat{\text{err}}(f) = 0} |\text{err}(f) - \hat{\text{err}}(f)| \geq \epsilon \right] \leq 2C[2m, D]e^{-m\epsilon^2/2}.
\]

So, for any \( \epsilon, \delta > 0 \), if we draw a sample from \( D \) of size

\[
m \geq \frac{2}{\epsilon} \left( 2 \ln(C[2m, D]) + \ln \left( \frac{2}{\delta} \right) \right),
\]

then with probability at least \( 1 - \delta \), we have that all functions \( f \) with \( \hat{\text{err}}(f) = 0 \) satisfy \( \text{err}(f) \leq \epsilon \).

**Theorem 26.** Suppose that \( C \) is a set of functions from \( X \) to \( \{0, 1\} \) with finite VC-dimension \( V \geq 1 \). For any probability distribution \( D \) over \( X \), any target function \( c^* \), we have

\[
\Pr_S \left[ \sup_{f \in C} |\text{err}(f) - \hat{\text{err}}(f)| \geq \epsilon \right] \leq 4C[2m, D]e^{-m\epsilon^2/8}.
\]

So, for any \( \epsilon, \delta > 0 \), if we draw from \( D \) a sample satisfying

\[
m \geq \frac{8}{\epsilon^2} \left( \ln(C[m, D]) + \ln \left( \frac{4}{\delta} \right) \right),
\]

then with probability at least \( 1 - \delta \) all functions \( f \) satisfy \( \text{err}(f) - \hat{\text{err}}(f) \leq \epsilon \).

We now state a result from [Boucheron et al. 2000]. Let \( C[S] \) denote the number of ways of labeling set \( S \) using functions in \( C \).

**Theorem 27.** Suppose that \( C \) is a set of functions from \( X \) to \( \{0, 1\} \). Let \( D \) be an arbitrary, but fixed probability distribution over \( X \). For any target function and for any i.i.d. sample of \( S \) of size \( m \) from \( D \), let \( f_m \) be the function that minimizes the empirical error over \( S \). Then for any \( \delta > 0 \), the probability that

\[
\text{err}(f_m) \leq \hat{\text{err}}(f_m) + \sqrt{\frac{6 \ln C[S]}{m}} + 4 \sqrt{\frac{\ln(2/\delta)}{m}}
\]

is greater than \( 1 - \delta \).

In fact the above statement is true even if in the right-hand side we use \( C[S'] \) instead of \( C[S] \) where \( S' \) is another i.i.d sample of size \( m \) drawn from \( D \). Using the notation introduced in section 3.1.3, since \( |S| = m \), we can also use \( C[m, S] \) instead of \( C[S] \) in the above statement.
THEOREM 28. For any class of functions we have:
\[
\Pr_S[\ln(C[S]) \geq E[\ln(C[S])] + \alpha] \leq \exp\left[-\frac{\alpha^2}{2E[\ln(C[S])] + 2\alpha/3}\right].
\] (3)

Also,
\[
E[\ln C[S]] \leq \ln E[C[S]] \leq (e - 1)E[\ln C[S]].
\] (4)

E. ADDITIONAL PROOFS

With the notations from Appendix D and in section 3.1.3 we have:

**THEOREM 29.** For any class of functions we have:

\[
\Pr_S[\ln(C[m, S]) \geq 2\ln C[m, D] + \alpha] \leq e^{-\alpha}.
\] (5)

**PROOF.** Inequality (3) implies that:

\[
\Pr_S[\ln(C[S]) \geq 2E[\ln(C[S])] + \alpha] \leq \exp\left[-\frac{(\alpha + E[\ln(C[S)])^2}{2E[\ln(C[S])] + 2(E[\ln(C[S])] + \alpha)/3}\right].
\]

Since \(\frac{(\alpha + E)^2}{2E[\ln(C[S])] + 2\alpha/3} \geq \alpha\) for any \(\alpha \geq 0\) we get

\[
\Pr_S[\ln(C[S]) \geq 2E[\ln(C[S])] + \alpha] \leq e^{-\alpha}.
\]

Combining this together with the following fact (implied by Inequality (4))

\[
\Pr_S[\ln(C[S]) \geq 2\ln E[C[S]] + \alpha] \leq \Pr_S[\ln(C[S]) \geq 2E[\ln(C[S])] + \alpha],
\]

we get

\[
\Pr_S[\ln(C[S]) \geq 2\ln E[C[S]] + \alpha] \leq e^{-\alpha}.
\]

Using the facts that for \(S\) of size \(m\), \(C[m, S] = C[S]\) and \(E_S[C[m, S]] = C[m, D]\), we get the desired result. \(\square\)

We finally prove here a lemma used in Section 3.2.2.

**LEMMA 16.** Consider \(\tau < \frac{1}{\delta}\). Let \(C_\tau = \{f, \overline{f}\}\) be a subset of \(C\) containing two opposite hypotheses with the property that one of them is \(\tau\)-close to \(c^*\). Then, \(m_l > 6\log_2(\frac{1}{\delta})\) labeled examples are sufficient so that with probability \(\geq 1 - \delta\), the concept in \(C_\tau\) that is \(\tau\)-close to \(c^*\) in fact has lower empirical error.

**PROOF.** We need to show that if \(m_l > 6\log_2(\frac{1}{\delta})\), then

\[
\sum_{k=0}^{m_l} \binom{m_l}{k} \tau^{m_l-k} (1-\tau)^k \leq \delta.
\]

Since \(\tau < \frac{1}{\delta}\) we have:

\[
\sum_{k=0}^{m_l} \binom{m_l}{k} \tau^{m_l-k} (1-\tau)^k \leq \sum_{k=0}^{m_l} \binom{m_l}{k} \tau^{m_l-k} = \tau^{m_l} \sum_{k=0}^{m_l} \binom{m_l}{k} \tau^{m_l-k}
\]

and so \(S \leq (\sqrt{\tau} \cdot 2)^{m_l}\). For \(\tau < \frac{1}{\delta}\) and \(m_l > 6\log_2(\frac{1}{\delta}) = 6\log_2(\frac{1}{\delta})\) it’s easy to see that \((\sqrt{\tau} \cdot 2)^{m_l} < \delta\), which implies the desired result. \(\square\)