Scalable Semi-Supervised Classifier Aggregation

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Abstract

We present and empirically evaluate an efficient algorithm that learns to predict using an ensemble of binary classifiers. It uses the structure of the ensemble predictions on unlabeled data to yield classification performance gains without making assumptions on the predictions or their origin, and does this as scalably as linear learning.

1 Introduction

Ensemble-based learning is a very successful approach to learning classifiers. The best known methods include boosting [1], bagging [2], and random forests [3]. The power of these methods has been clearly demonstrated in open large-scale learning competitions such as the Netflix Prize [5] and the ImageNet Challenge [4]. In general, these methods train a large number of “base” classifiers and then combine them using a (possibly weighted) majority vote. By aggregating over classifiers, ensemble methods reduce the variance of the predictions, and sometimes also reduce the bias [6].

Existing learning methods for ensembles rely solely on a labeled training set. In this paper we propose an ensemble method that uses a large unlabeled data set in addition to the labeled set. Our work is therefore at the intersection of semi-supervised learning [7, 8] and ensemble learning.

This paper is based on recent theoretical results of Balsubramani and Freund [9]. The main contribution of this paper is to extend and apply those results in the context of random forests [3] and perform experiments in which we show that, when the number of labeled examples is small, our algorithm’s performance is at least that of random forests, and often significantly better.

For the sake of completeness, we provide an intuitive introduction to the analysis given in [9]. How can unlabeled data help in the context of ensemble learning? Consider a simple example. Suppose the ensemble consists of six classifiers, partitioned into three “A” rules and three “B” rules. Suppose that the “A” rules each have error at most $\frac{1}{3}$ and the “B” rules each have error at most $\frac{1}{6}$.

If given only this information, we might take the majority vote over the six rules, possibly giving higher weights to the “B” rules because they have lower errors.

Suppose, however, that we are given the unlabeled information in Table 1. The columns of this table correspond to the six classifiers and the rows correspond to the six unlabeled examples. Each entry corresponds to the prediction of the given classifier on the given example. As we see, the main difference between the “A” rules and the “B” rules is that any two “A” rules disagree with probability $\frac{1}{3}$, whereas the “B” rules always agree. For this example, it can be seen that the only possible true labeling of the unlabeled data that is consistent with Table 1 and with the errors of the classifiers is that all the examples are labeled ‘+’. Equivalently, the majority vote over the “A” rules has zero error, performing significantly better than any of the base rules.

We assume that the upper bounds on the errors are, with high probability, true on the actual distribution. Such bounds can be derived using large deviation bounds or bootstrap-type methods.
In contrast, giving the “B” rules equal weight would result in an a rule with error $1/6$. Crucially, this superiority of taking a majority over the “A” rules can be inferred solely from the unlabeled examples in Table 1.

|       | A classifiers | B classifiers |
|-------|-------------|---------------|
| $x_1$ | -           | +             |
| $x_2$ | -           | +             |
| $x_3$ | +           | -             |
| $x_4$ | +           | +             |
| $x_5$ | +           | -             |
| $x_6$ | +           | -             |
| error | 1/3         | 1/3           |

Table 1: An example of the utility of unlabeled examples in ensemble learning

This is only a toy example, but it provides some insight into the ways in which unlabeled data can be useful:

- When combining classifiers, diversity is important. It can be better to combine less accurate rules that disagree with each other than to combine more accurate rules that tend to agree.
- The bounds on the errors of the rules can be seen as a set of constraints. A complementary set of constraints is provided by the unlabeled examples. These two sets of constraints can be combined to improve the accuracy of the ensemble classifier.

The above setup was recently introduced and analyzed in [9]. That paper characterizes the problem as a zero-sum game between a predictor and nature. It then characterizes the minimax solution of the game, which corresponds to an efficient algorithm for transductive learning.

In this paper, we build on the worst-case framework of [9] to devise an efficient and practical semi-supervised aggregation algorithm for random forests. To achieve this, we extend the framework to handle specialists – classifiers which only venture to predict on a subset of the data, and abstain from predicting on the rest. Specialists can be very useful in targeting regions of the data on which to precisely suggest a prediction.

The high-level idea of our algorithm is to artificially generate new specialists from the ensemble. We incorporate these, and the targeted information they carry, into the worst-case framework of [9]. The resulting aggregated predictor inherits the advantages of the original framework:

(A) **Efficient**: Learning reduces to solving a scalable $p$-dimensional convex optimization, and test-time prediction is as efficient and parallelizable as $p$-dimensional linear prediction.

(B) **Versatile/robust**: No assumptions about the structure or origin of the predictions or labels.

(C) **No introduced parameters**: The aggregation method is completely data-dependent.

(D) **Safe**: Accuracy guaranteed to be at least that of the best classifier in the ensemble.

We develop these ideas in the rest of this paper, reviewing the core worst-case setting of [9] in Section 2 and specifying how to incorporate specialists and the resulting learning algorithm in Section 3.

Then we perform an exploratory evaluation of the framework on data in Section 4. Though the framework of [9] and our extensions can be applied to any ensemble of arbitrary origin, in this paper we focus on random forests, which have been repeatedly demonstrated to have state-of-the-art, robust classification performance in a wide variety of situations [11]. We use a random forest as a base ensemble whose predictions we aggregate. But unlike conventional random forests, we do not simply take a majority vote over tree predictions, instead using a unlabeled-data-dependent aggregation strategy dictated by the worst-case framework we employ.

2 Preliminaries

A few definitions are required to discuss these issues concretely, following [9]. Write $[a]_+ = \max(0, a)$ and $[n] = \{1, 2, \ldots, n\}$. All vector inequalities are componentwise.
We first consider an ensemble \( \mathcal{H} = \{ h_1, \ldots, h_p \} \) and unlabeled data \( x_1, \ldots, x_n \) on which we wish to predict. As in \( \cite{9} \), the predictions and labels are allowed to be randomized, represented by values in \([-1, 1]\) instead of just the two values \( \{-1, 1\} \). The ensemble’s predictions on the unlabeled data are denoted by \( F \):

\[
F = \begin{pmatrix}
h_1(x_1) & h_1(x_2) & \cdots & h_1(x_n) \\
\vdots & \vdots & \ddots & \vdots \\
h_p(x_1) & h_p(x_2) & \cdots & h_p(x_n)
\end{pmatrix} \in [-1, 1]^{p \times n} \tag{1}
\]

We use vector notation for the rows and columns of \( F \): \( h_i = (h_i(x_1), \ldots, h_i(x_n))^\top \) and \( x_j = (h_1(x_j), \ldots, h_p(x_j))^\top \). The true labels on the test data \( T \) are represented by \( z = (z_1; \ldots; z_n) \in [-1, 1]^n \). The labels \( z \) are hidden from the predictor, but we assume the predictor has knowledge of a correlation vector \( b \in (0, 1]^n \) such that \( \frac{1}{n} \sum_{j} h_j(x_j) z_j \geq b_i \), i.e. \( \frac{1}{n} F z \geq b \). These \( p \) inequalities exactly represent upper bounds on individual classifier error rates, which can be estimated from the training set w.h.p. in a standard way also used by ERM \( \cite{9} \).

### 2.1 The Transductive Binary Classification Game

The idea of \( \cite{9} \) is to formulate the ensemble aggregation problem as a two-player zero-sum game between a predictor and an adversary. In this game, the predictor is the first player, who plays \( g = (g_1, g_2; \ldots; g_n) \), a randomized label \( g_i \in [-1, 1] \) for each example \( \{x_i\}_{i=1}^n \). The adversary then sets the labels \( z \in [-1, 1]^n \) under ensemble classifier error constraints defined by \( b \).\footnote{Since \( b \) is calculated from the training set, the problem is feasible w.h.p., which we assume.} The predictor’s goal is to minimize the worst-case expected classification error on the test data (w.r.t. the randomized labelings \( z \) and \( g \)), which is just \( \frac{1}{n} \left( 1 - \frac{1}{n} z^\top g \right) \). This is equivalently viewed as maximizing worst-case correlation \( \frac{1}{n} z^\top g \). To summarize concretely, we study the following game:

\[
V := \max_{g \in [-1,1]^n} \min_{z \in [-1,1]^n, \frac{1}{n} F z \geq b} \frac{1}{n} z^\top g \tag{2}
\]

The minimax theorem ((\( \cite{1} \), p.144) applies to the game (2). Therefore, there is an optimal strategy \( g^* \) such that \( \min_{z \in [-1,1]^n, \frac{1}{n} F z \geq b} \frac{1}{n} z^\top g^* \geq V \), guaranteeing worst-case prediction error \( \frac{1}{n} (1 - V) \) on the unlabeled data. This optimal strategy \( g^* \) is a simple function of a particular weighting over the \( p \) hypotheses – a non-negative \( p \)-vector.

**Definition 1 (Slack Function).** Let \( \sigma \geq 0^p \) be a weight vector over \( \mathcal{H} \) (not necessarily a distribution). The vector of **ensemble predictions** is \( F^\top \sigma = (x_1^\top \sigma, \ldots, x_n^\top \sigma) \), whose elements’ magnitudes are the **margins**. The prediction slack function is

\[
\gamma(\sigma, b) := \gamma(\sigma) := -b^\top \sigma + \frac{1}{n} \sum_{j=1}^n \left( |x_j^\top \sigma| - 1 \right)_+ \tag{3}
\]

The **optimal weight vector** \( \sigma^* \) is any minimizer of the slack function: \( \sigma^* \in \arg \min_{\sigma \geq 0^p} \gamma(\sigma) \).

The main result of \( \cite{9} \) uses these to describe the minimax equilibrium of the game (2).

**Theorem 2 (\( \cite{9} \)).** The minimax value of the game (2) is \( V = -\gamma(\sigma^*) \). The minimax optimal predictions are defined as follows: for each \( i \in [n] \),

\[
g_i^* := g_i(\sigma^*) = \begin{cases} 
    x_i^\top \sigma^* & |x_i^\top \sigma^*| < 1 \\
    \text{sgn}(x_i^\top \sigma^*) & \text{otherwise}
\end{cases}
\]

### 2.2 Interpretation

The above game-theoretic framework describes a way to learn in a transductive setting, in which test data (but not their labels) are known in advance. But it also applies to a standard setup in which all
the data are independently sampled from some unknown distribution; in such a case, $b$ is estimated from the training set, and contains all the label information used.

Theorem 2 therefore specifies a learning algorithm for aggregating the $p$ ensemble classifiers’ predictions: estimate $b$ from the training data, optimize the slack function $\gamma(\sigma)$ to find $\sigma^*$, and finally predict with $g^*$ as calculated from $\sigma^*$ and the test set. The resulting predictions are guaranteed to have low error, as measured by $V$.

The slack function (3) merits further scrutiny. Its first term depends only on the labeled data and not the unlabeled set, while the second term $\frac{1}{n} \sum_{j=1}^n \left( |x_j^\top \sigma| - 1 \right)_+$ incorporates only unlabeled information. These two terms trade off smoothly – as the problem setting becomes fully supervised and unlabeled information is absent, the first term dominates, and $\sigma^*$ tends to put all its weight on the best single classifier like ERM.

Indeed, this viewpoint suggests a (loose) interpretation of the second term as an unsupervised regularizer for the otherwise fully supervised optimization of the “average” error $b^\top \sigma$. It turns out that a change in the regularization factor corresponds to different constraints on the true labels $z$:

**Theorem 3 (9).** Let $V_{\alpha} := \max_{g \in [-1,1]^n} \min_{z \in [-\alpha, \alpha]^n} \frac{1}{n} \sum_{j=1}^n \left( x_j^\top g \right)$ for any $\alpha > 0$. Then $V_{\alpha} = \min \left[ -b^\top \sigma + \frac{\alpha}{n} \sum_{j=1}^n \left( x_j^\top \sigma - 1 \right)_+ \right].$

So the regularized optimization assumes each $z_j \in [-\alpha, \alpha]$. For $\alpha < 1$, this is equivalent to assuming the usual binary labels ($\alpha = 1$), and then adding uniform random label noise: flipping the label w.p. $\frac{1}{2} (1 - \alpha)$ on each of the $n$ examples independently. This encourages “clipping” of the ensemble predictions $x_j^\top \sigma^*$ to the $\sigma^*$-weighted majority vote predictions, as specified by $g^*$.

### 2.3 Advantages and Disadvantages

This formulation has several significant merits that would seem to recommend its use in practical situations. It is very efficient – once $b$ is estimated (a scalable task, given the labeled set), the slack function $\gamma$ is effectively an average over convex functions of i.i.d. unlabeled examples, and consequently is amenable to standard convex optimization techniques (9) like stochastic gradient descent (SGD) and variants. These only operate in $p$ dimensions, independent of $n$ (which is $\gg p$). The slack function is Lipschitz and well-behaved, resulting in stable approximate learning.

Moreover, test-time prediction is extremely efficient, because it only requires the $p$-dimensional weighting $\sigma^*$ and can be computed example-by-example on the test set using only a dot product in $\mathbb{R}^p$. The form of $g^*$ and its dependence on $\sigma^*$ facilitates interpretation as well, as it resembles familiar objects: sigmoid link functions for linear classifiers.

Other advantages of this method also bear mention: it makes no assumptions on the structure of $\mathcal{H}$ or $F$, is robust by nature of its worst-case setting, and adds no input parameters that need tuning. These benefits are notable because they will be inherited by our extension of the framework that constitutes our main contribution.

However, this algorithm’s practical performance can still be mediocre on real data, which is often easier to predict than an adversarial setup would have us believe. As a result, we seek to add more information in the form of constraints on the adversarial setup, to narrow the gap between it and reality.

### 3 Learning with Specialists

To address this issue, we examine a scenario in which our prediction is binary, but each classifier in the ensemble can abstain on any subset of the examples instead of predicting $\pm 1$. It is a specialist that predicts only over a subset of the input, and we think of its abstain/participate decision being randomized in the same way as in the previous section. In this section, we extend the framework of Section 2.1 to arbitrary specialists, and discuss the semi-supervised learning algorithm that results.
In our formulation, suppose that for a classifier $i \in [p]$ and an example $x$, the classifier decides to abstain with probability $1 - \gamma_i(x)$. But if the decision is to participate, the classifier predicts $h_i(x) \in [-1, 1]$ as previously. Our only assumption on $\{\gamma_i(x_1), \ldots, \gamma_i(x_n)\}$ is the reasonable one that $\sum_{j=1}^n \gamma_i(x_j) > 0$, so classifier $i$ is not a worthless specialist that abstains everywhere.

The constraint on classifier $i$ is now not on its overall correlation with $z$ on the entire test set, but on the correlation with $z$ restricted to the examples on which it participates. In other words, for some $[\rho_i]_i \in [0, 1]$,\footnote{Here $[\cdot]_i$ represents the $i$th element of a vector.}

$$\sum_{j=1}^n \left( \frac{\gamma_i(x_j)}{\sum_{k=1}^n \gamma_i(x_k)} \right) h_i(x_j) z_j \geq [\rho_i]_i$$

(4)

Define $\rho_i(x_j) := \frac{\gamma_i(x_j)}{\sum_{k=1}^n \gamma_i(x_k)}$, a distribution over $j \in [n]$ for convenience. Now redefine our unlabeled data matrix as follows:

$$S = \begin{pmatrix} \rho_1(x_1) h_1(x_1) & \rho_1(x_2) h_1(x_2) & \cdots & \rho_1(x_n) h_1(x_n) \\ \vdots & \vdots & \ddots & \vdots \\ \rho_p(x_1) h_p(x_1) & \rho_p(x_2) h_p(x_2) & \cdots & \rho_p(x_n) h_p(x_n) \end{pmatrix}$$

(5)

so that the constraints (4) can be written as $\frac{1}{n} S z \geq b_S$, analogous to the initial prediction game (2).

To summarize, our specialist ensemble aggregation game is stated as

$$V_S := \min_{z \in [-1, 1]^n} \max_{z \in [0, 1]^n} \frac{1}{n} z^T g$$

(6)

We can immediately solve this game from Thm. 2 with $(S, b_S)$ simply taking the place of $(F, b)$.

**Theorem 4** (Solution of the Specialist Aggregation Game). The awake ensemble prediction w.r.t. weighting $\sigma \geq 0^p$ on example $x_i$ is $[S^T \sigma]_i = n \sum_{j=1}^p \rho_j(x_i) h_j(x_i) \sigma_j$. The slack function is now

$$\gamma_S(\sigma) := \frac{1}{n} \sum_{j=1}^n \left( [S^T \sigma]_j - 1 \right)_+ - b_S^T \sigma$$

(7)

The minimax value of this game is $V_S = \max_{\sigma \geq 0^p} -\gamma_S(\sigma) = -\gamma_S(\sigma_S^*)$. The minimax optimal predictions are defined as follows: for all $i \in [n]$,

$$[g_S^*]_i = \begin{cases} [S^T \sigma_S^*]_i & \text{sgn}([S^T \sigma_S^*]_i) < 1 \\ \top & \text{otherwise} \end{cases}$$

In the no-specialists case, the vector $\rho_i$ is the uniform distribution $(\frac{1}{n}, \ldots, \frac{1}{n})$ for any $i \in [p]$, and the problem reduces to the prediction game (2). As in the original prediction game, the minimax equilibrium depends on the data only through the ensemble predictions, but these are now of a different form. Each example is now weighted proportionally to $\rho_j(x_i)$. So on any given example $x_i$, only hypotheses which participate on it will be counted; and those that specialize the most narrowly, and participate on the fewest other examples, will have more influence on the eventual prediction $g_i$, ceteris paribus.

### 3.1 Creating Specialists for an Algorithm

We can now present the main ensemble aggregation method of this paper, which creates specialists from the ensemble, adding them as additional constraints (rows of $S$). The algorithm, HEDGECLIPPER, is given in Fig. 1 and instantiates our specialist learning framework with a random forest $\mathcal{F}$. As an initial exploration of the framework here, since the goal is better classification performance, random forests are an appropriate base ensemble because they are known to exhibit state-of-the-art performance $\mathcal{F}$. Their well-known advantages also include scalability, robustness (to corrupt data and parameter choices), and interpretability; each of these benefits is shared by our aggregation algorithm, which consequently inherits them all.
Furthermore, decision trees are a natural fit as the ensemble classifiers because they are inherently hierarchical. Intuitively (and indeed formally too [12]), they act like nearest-neighbor (NN) predictors w.r.t. a distance that is “adaptive” to the data. So each tree in a random forest represents a somewhat different, nonparametric partition of the data space into regions in which one of the labels ±1 dominates. Each such region corresponds exactly to a leaf of the tree.

The idea of HEDGECLIPPER is simply to consider each leaf in the forest as a specialist, which predicts only on the data falling into it. By the NN intuition above, these specialists can be viewed as predicting on data that is near them, where the supervised training of the tree attempts to define the purest possible partitioning of the space. A pure partitioning results in many specialists with \(|b_S| \approx 1\), each of which contributes to the awake ensemble prediction w.r.t. \(\sigma^*\) over its domain, to influence it towards the correct label (inasmuch as \(|b_S|\) is high).

Though the idea is complex in concept for a large forest with many arbitrarily overlapping leaves from different trees, it fits the worst-case specialist framework of the previous sections. So the algorithm is still essentially linear learning with convex optimization, as we have described.

**Algorithm 1** HEDGECLIPPER

| Input: Labeled set \(L\), unlabeled set \(U\) |
| 1: Using \(L\), grow trees \(T = \{T_1, \ldots, T_p\}\) (regularized; see Sec. 3.2) |
| 2: Using \(L\), estimate \(b_S\) on \(T\) and its leaves |
| 3: Using \(U\), (approximately) optimize \(\gamma\) to estimate \(\sigma^*_S\) |
| Output: The estimated weighting \(\sigma^*_S\), for use at test time |

Figure 1: At left is algorithm HEDGECLIPPER. At right is a schematic of how the forest structure is related to the unlabeled data matrix \(S\), with a given example \(x\) highlighted. The two colors in the matrix represent \(\pm 1\) predictions, and white cells abstentions.

**3.2 Discussion**

Trees in random forests have thousands of leaves in practice. As we are advocating adding so many extra specialists to the ensemble for the optimization, it is natural to ask whether this erodes some of the advantages we have claimed earlier.

Computationally, it does not. When \(\rho_j(x_i) = 0\), i.e. classifier \(j\) abstains always on \(x_i\), then the value of \(h_j(x_i)\) is irrelevant. So storing \(S\) in a sparse matrix format is natural in our setup, with the accompanying efficiency gain in computing \(S^\top \sigma\) while learning \(\sigma^*\) and predicting with it. This turns out to be crucial to efficiency – each tree induces a partitioning of the data, so the set of rows corresponding to any tree contains \(n\) nonzero entries in total. This is seen in Fig. 1.

Statistically, the situation is more complex. On one hand, there is no danger of overfitting in the traditional sense, regardless of how many specialists are added. Each additional specialist can only shrink the constraint set that the adversary must follow in the game \(\gamma\). It only adds information about \(z\), and therefore the performance \(V_S\) can only improve, if the game is solved exactly.

However, for learning we are only concerned with approximately optimizing \(\gamma_S(\sigma)\) and solving the game. This presents several statistical challenges. Standard optimization methods do not converge as well in high ambient dimension, even given the structure of our problem. In addition, random forests practically perform best when each tree is grown to overfit. In our case, on any sizable test set, small leaves would cause some entries of \(S\) to have large magnitude, \(\gg 1\). This can foil an algorithm like HEDGECLIPPER by causing it to vary wildly during the optimization, particularly since those leaves’ \(|b_S|\) values are only roughly estimated.

From an optimization perspective, some of these issues can be addressed by e.g. (pseudo)-second-order methods [13], whose effect would be interesting to explore in future work. Our implementation opts for another approach – to grow trees constrained to have a nontrivial minimum weight per leaf. Of course, there are many other ways to handle this, including using the tree structure beyond the
leaves; we just aim to conduct an exploratory evaluation here, as several of these areas remain ripe for future research.

4 Experimental Evaluation

We now turn to evaluating HEDGECLIPPER on publicly available datasets. Our implementation uses minibatch SGD to optimize (6), runs in Python on top of the popular open-source learning package scikit-learn, and runs out-of-core (with an $n$-independent memory footprint), taking advantage of the scalability of our formulation. The datasets are drawn from UCI/LibSVM as well as data mining sites like Kaggle. No further preprocessing was done on the data. We refer to “Base RF” as the forest of constrained trees from which our implementation draws its specialists. We restrict the training data available to the algorithm, using mostly supervised datasets because these far outnumber medium/large-scale public semi-supervised datasets. Unused labeled examples are combined with the test examples (and the extra unlabeled set, if any is provided) to form the set of unlabeled data used by the algorithm. Further information and discussion on the protocol is in the appendix.

Class-imbalanced and noisy sets are included to demonstrate the aforementioned practical advantages of HEDGECLIPPER. Therefore, AUC is an appropriate measure of performance, and these results are in Table 2. Results are averaged over 10 runs, each drawing a different random subsample of labeled data. The best results according to a paired t-test are in bold.

We find that the use of unlabeled data is sufficient to achieve improvements over even traditionally overfitted RFs in many cases. Notably, in most cases there is a significant benefit given by unlabeled data in our formulation, as compared to the base RF used. The boosting-type methods also perform fairly well, as we discuss in the next section.

![Figure 2: Class-conditional “awake ensemble prediction” ($\mathbf{x}^\top \sigma^*$) distributions, on SUSY. Rows (top to bottom): $\{1K, 10K, 100K\}$ labels. Columns (left to right): $\alpha = \{1.0, 0.3, 3.0\}$, and the base RF.](image)

The awake ensemble prediction values $\mathbf{x}^\top \sigma$ on the unlabeled set are a natural way to visualize and explore the operation of the algorithm on the data, in an analogous way to the margin distribution in boosting [6]. One representative sample is in Fig. 2 on SUSY, a dataset with many (5M) examples, roughly evenly split between $\pm 1$. These plots demonstrate that our algorithm produces much more peaked class-conditional ensemble prediction distributions than random forests, suggesting “margin-based” learning applications. Changing $\alpha$ alters the aggressiveness of the clipping, inducing a more or less peaked distribution. The other datasets without dramatic label imbalance show strikingly similar qualitative behavior in these respects, and these plots help choose an appropriate $\alpha$ in practice (see appendix).

Datasets with very low dimension (compared to $n$) seem to exhibit little to no significant improvement from our method; we believe this is because the distinct feature splits found by the random forest are few in number, and it is the diversity in ensemble predictions that enables HEDGECLIPPER to clip (weighted majority vote) dramatically and achieve its performance gains.

5 Related and Future Work

This paper’s framework and algorithms are superficially reminiscent of boosting, another paradigm that uses voting behavior to aggregate an ensemble and has a game-theoretic intuition tied to linear programming [1] [15]. There has been some work on semi-supervised versions of boosting [16],

\[\text{It is possible to make this footprint independent of } d \text{ as well by hashing features [14], which is outside the scope of this manuscript.}\]
but it departs from this principled structure and has little in common with our approach. Classical boosting algorithms like AdaBoost [17] make no attempt to use unlabeled data. It is an interesting open problem to incorporate boosting ideas into our formulation, particularly since the two boosting-type methods acquit themselves well in our results, and can pack information parsimoniously into many fewer ensemble classifiers than random forests.

There is a long-recognized connection between transductive and semi-supervised learning, and our method bridges these two settings. Popular methods such as the transductive SVM [18] and graph-based or nearest-neighbor algorithms, which dominate the semi-supervised literature [8], have shown promise largely in data-poor regimes because they face major scalability challenges. Our focus on ensemble aggregation instead allows us to keep a computationally inexpensive linear formulation and avoid considering the underlying feature space of the data.

Learning with specialists has been studied in an adversarial online setting [19], as in the work of Freund et al. [20]. Though that paper’s setting and focus is different from ours, the optimal algorithms it derives also depend on each specialist’s average error on the examples on which it is awake.

Finally, we re-emphasize the generality of our formulation, which leaves many interesting questions yet to be explored. The specialists we form are not restricted to being trees; there are other ways of dividing the data like clustering methods. Indeed, the composition of the ensemble can be heterogeneous and even incorporate other semi-supervised methods. Our method is complementary to myriad classification algorithms, and we hope to stimulate inquiry into the many research avenues this opens.

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A Additional Information on Experiments

A.1 Datasets

Information on the datasets used:

| Dataset     | Data sizes $m/n$ | Dim. $d$ | Comments                  |
|-------------|-----------------|----------|---------------------------|
| kagg-prot   | 3751/1776       | 12       | Kaggle challenge [21]     |
| ssl-text    | 1500            | 11960    | [7]                       |
| kagg-cred   | 150000          | 10       | Kaggle challenge [22]     |
| ala         | 1605/30956      | 123      | LibSVM                    |
| wla         | 2477/47272      | 300      | LibSVM                    |
| covtype     | 581012          | 54       | LibSVM                    |
| ssl-secstr  | 83679 (unlab-   | 315      | [7]                       |
|             | eled:1189472)   |          |                           |
| SUSY        | 50000000        | 18       | UCI                       |
| epsilon     | 5000000         | 2000     | PASCAL Large Scale Learning Challenge 2008 |
| webspam-uni | 3500000        | 254      | LibSVM                    |

All data from the challenges (e.g. kagg-cred) lacked test labels, so the results reported are averaged over 10 random splits of the training data.

A.2 Algorithms

In all cases, the random forests were grown with default parameters for the feature and data splits (bootstrap data sample of size $n$, and $\sim \sqrt{d}$ features considered per split). Varying these fundamentally changes the induced diversity of the trees/partitions, but exploration of such aspects is left to future work. All the comparator algorithms were also run with default parameters – in many cases like RFs, they are fairly insensitive to parameter choice.

To overcome the statistical issues discussed in Sec. 3.2 we found we needed to enforce some regularization on the tree used. We chose to impose a constraint on the minimum number of training examples in any leaf of the tree. This constraint was imposed as a parameter to grow the forest; thereafter, we could use all resulting leaves as specialists. To avoid any leaf specialist weights being too large but still collect as many leaf specialists as possible, we set the minimum number of examples per leaf to 10 with $\geq 1K$ labeled examples, and to 4 otherwise.

We also tried an alternative way of avoiding small specialists: to simply grow an unregularized forest and then filter out leaves, selecting only large enough leaves as specialists. This generally performed comparably or worse, consistent with the intuition that the diversity in unregularized tree predictions often manifests largely on small leaves.

A notable issue we encountered is the setting of the “noise” rescaling factor $\alpha$. We found HEDGECLIPPER to be relatively insensitive to the precise choice of $\alpha$, so it essentially sufficed for our experiments to try three choices: $\{0.3, 1.0, 3.0\}$. The last is $> 1.0$, and therefore does not have an interpretation in terms of uniform label noise, but it is certainly a valid computational tool.

Which of these three $\alpha$s to choose? Generally, we found that choosing $\alpha = 0.3$ does not hurt performance, because our performance goals are often met best by separating the class-conditional peaks as much as possible. This is dangerous for more class-unbalanced datasets like kagg-cred, however, for which the default $\alpha = 1.0$ works best. A useful heuristic which we used to choose $\alpha$ is simply to look at the class-conditional awake ensemble prediction distribution as plotted in Fig. 2; the distribution can be roughly estimated and plotted on the fly, and we can quickly ascertain sensible choices of parameters like $\alpha$. 