

# Meta-Prediction for Collective Classification

Luke K. McDowell<sup>1</sup>, Kalyan Moy Gupta<sup>2</sup>, and David W. Aha<sup>3</sup>

<sup>1</sup>Dept. of Computer Science; U.S. Naval Academy; Annapolis, MD 21402

<sup>2</sup>Knexus Research Corp.; Springfield, VA 22153

<sup>3</sup>Navy Center for Applied Research in Artificial Intelligence;

Naval Research Laboratory (Code 5514); Washington, DC 20375

lmcadowel@usna.edu | kalyan.gupta@knexusresearch.com | david.aha@nrl.navy.mil

## Abstract

When data instances are inter-related, as are nodes in a social network or hyperlink graph, algorithms for collective classification (CC) can significantly improve accuracy. Recently, an algorithm for CC named *Cautious ICA* ( $ICA_C$ ) was shown to improve accuracy compared to the popular ICA algorithm.  $ICA_C$  improves performance by initially favoring its more confident predictions during collective inference. In this paper, we introduce  $ICA_{MC}$ , a new algorithm that outperforms  $ICA_C$  when the attributes that describe each node are not highly predictive.  $ICA_{MC}$  learns a meta-classifier that identifies which node label predictions are most likely to be correct. We show that this approach significantly increases accuracy on a range of real and synthetic data sets. We also describe new features for the meta-classifier and demonstrate that a simple search can identify an effective feature set that increases accuracy.

## Introduction

In many classification tasks, the instances to be classified (such as web pages or people in a social network) are related in some way. *Collective classification* (CC) is a methodology that jointly classifies such instances (or *nodes*). CC algorithms can attain higher accuracies than non-collective methods when nodes are interrelated (Neville and Jensen 2000; Taskar, Abbeel, and Koller 2002). Several CC algorithms have been studied, including relaxation labeling (Chakrabarti, Dom, and Indyk 1998), the Iterative Classification Algorithm (ICA) (Sen *et al.* 2008), loopy belief propagation (LBP) (Taskar *et al.* 2002), and Gibbs sampling (Jensen, Neville, and Gallagher 2004).

We focus on ICA because it is a popular and computationally efficient algorithm that has good classification performance (Sen *et al.* 2008). It makes initial label predictions for each node  $v_i$ , then iteratively re-computes them based on the predictions for *every* node that links to  $v_i$ . Recently, a variant of ICA named *Cautious ICA* ( $ICA_C$ ) (McDowell *et al.* 2007, 2009) was shown to often attain higher accuracies than ICA.  $ICA_C$  is based on the

observation that, since some label predictions will be incorrect, ICA's use of *all* predictions may sometimes decrease accuracy. To counter this effect,  $ICA_C$  instead initially uses only *some* label predictions. By "cautiously" choosing only those predictions that appear more likely to be correct,  $ICA_C$  can increase accuracy vs. ICA.

In this paper, we introduce Meta-Cautious ICA ( $ICA_{MC}$ ), which is exactly like  $ICA_C$  except in how it selects the set of predicted labels to use during classification. In particular,  $ICA_{MC}$  learns a *meta-classifier* to predict the likelihood that a label prediction is correct. By carefully constructing a meta-training set from the original training set,  $ICA_{MC}$  can learn this classifier and use it to select more reliable predicted labels than  $ICA_C$ , increasing accuracy.

Our contributions are as follows. First, we present  $ICA_{MC}$ , a novel algorithm that can significantly increase accuracy compared to  $ICA_C$ , especially when the attributes that describe each node are not very predictive. Second, we introduce a technique to improve accuracy by generating more training examples for  $ICA_{MC}$ 's meta-classifier. Third, we describe new features for the meta-classifier and demonstrate that, while the most effective meta-features for  $ICA_{MC}$  are task-dependent, a simple search identifies an effective set that increases accuracy. Empirical evaluations using real and synthetic datasets support our claims.

We next review CC and the ICA and  $ICA_C$  algorithms. Then we introduce  $ICA_{MC}$ . Finally, we present our experimental evaluation and discuss future research issues.

## Collective Classification

Assume we are given a graph  $G = (V, E, X, Y, C)$ , where  $V$  is a set of nodes,  $E$  is a set of (possibly directed) edges, each  $x_i \in X$  is an attribute vector for node  $v_i \in V$ , each  $Y_i \in Y$  is a label variable for  $v_i$ , and  $C$  is the set of possible labels. We are also given a set of "known" label values  $Y^K$  for nodes  $V^K \subset V$ , so that  $Y^K = \{y_i \mid v_i \in V^K\}$ . Finally, assume that we are given a training graph  $G_{Tr}$ , which is defined similarly to  $G$  except that every node in  $G_{Tr}$  is a "known" node. Then the task is to infer  $Y^U = Y - Y^K$ , which are the values of  $Y_i$  for the nodes in  $G$  whose labels are unknown. For each node  $v_i$ , let  $y_i$  be the true label and  $\hat{y}_i$  be the predicted label.

For example, consider the task of predicting whether a web page belongs to a professor or a student. Conventional supervised learning approaches ignore the links and classify each page using attributes derived from its content (e.g., words present in the page). In contrast, a technique for *relational classification* explicitly uses the links to construct additional features for classification (e.g., for each page, include as features the words from hyperlinked pages). These relational features can increase classification accuracy, though not always (Chakrabarti *et al.* 1998). Alternatively, even greater (and usually more reliable) increases can occur when the *class labels* of the linked pages are used to derive relevant relational features (Jensen *et al.* 2004). However, using features based on these labels is challenging because some or all of these labels are initially unknown. Thus, their labels must first be predicted (without using relational features) and then re-predicted in some manner (using all features). This process of *jointly* inferring the labels of interrelated nodes is known as *collective classification* (CC).

We next describe two existing collective inference algorithms, ICA and ICA<sub>C</sub>, and then introduce ICA<sub>MC</sub>. Each algorithm relies on a given *node classifier* ( $M_{AR}$ ) that predicts a node’s label using both attributes and relations.

### ICA: Inference using *all* predicted labels

Figure 1 shows pseudocode for ICA, ICA<sub>C</sub>, and ICA<sub>MC</sub> (depending on *AlgType*). Step 1 is a “bootstrap” step that predicts the class label  $\hat{y}_i$  for each node in  $V^U$  using only attributes ( $conf_i$  records the confidence of this prediction, but ICA does not use it). ICA then iterates (step 2). During each iteration, it selects *all* available label predictions (step 3), computes the relational features’ values based on them (step 4), and then re-predicts the class label of each node using both attributes and relational features (step 5). Step 6 is ignored for ICA. After iterating, step 7 returns the final set of predicted class labels and their confidence values.

### ICA<sub>C</sub>: Inference using *some* predicted labels

In steps 3-4 of Figure 1, ICA assumes that the predicted node labels are all equally likely to be correct. When *AlgType* is instead ICA<sub>C</sub>, the inference becomes more cautious by only considering more confident predictions. Specifically, step 3 “commits” into  $Y'$  only the most confident  $m$  of the currently predicted labels; other labels are considered *missing* and are ignored. Step 4 computes the relational features using only the committed labels, and step 5 performs classification using this information. Step 3 gradually increases the fraction of predicted labels that are committed per iteration. Node label assignments committed in an iteration  $h$  are not necessarily committed again in future iterations (and may in fact change).

ICA<sub>C</sub> requires a confidence measure ( $conf_i$  in Figure 1) to rank the current label predictions. As with prior work (Neville and Jensen 2000, McDowell *et al.* 2007), we set  $conf_i$  to be the posterior probability of the most likely class

```

ICA_classify(V,E,X,YK,MAR,MA,MM,n,AlgType,Φ) =
// V=nodes; E=edges; X=attr. vectors; YK=labels of known nodes
// MAR=node classifier (uses attr. & relations); MA= classifier
// that uses attr. only; MM=meta-classifier (predicts correctness)
// n=# iters; AlgType=ICA, ICAC, or ICAMC; Φ=est. class distr.
1 for each node  $v_i \in V^U$  do // Bootstrap
   ( $\hat{y}_i, conf_i$ )  $\leftarrow$  MA( $x_i$ )
2 for h = 0 to n do
3 // Select node labels for computing relational feat. values
  if (AlgType = ICA) // Use all labels: Known or predicted
    Y'  $\leftarrow$  YK  $\cup$  { $\hat{y}_i \mid v_i \in V^U$ }
  else // ICAC or ICAMC: Use known and m
    m  $\leftarrow$  |VU| * (h/n) // most confident predicted labels
    Y'  $\leftarrow$  YK  $\cup$  { $\hat{y}_i \mid v_i \in V^U \wedge \text{rank}(conf_i) \leq m$ }
4 for each node  $v_i \in V^U$  do // Use labels selected above
    $f_i \leftarrow \text{calcRelatFeats}(V,E,Y')$  // to compute feat. values
5 for each node  $v_i \in V^U$  do // Re-predict labels using
   ( $\hat{y}_i, conf_i$ )  $\leftarrow$  MAR( $x_i, f_i$ ) // attributes and features
6 if (AlgType = ICAMC) // Compute meta-features;
   ZU  $\leftarrow$  {( $\hat{y}_i, conf_i$ )  $\mid v_i \in V^U$ } // use to re-estimate conf.
   for each node  $v_i \in V^U$  do // values for each node
      $mf_i \leftarrow \text{calcMetaFeatures}(i,V,E,X,Y',Y^K,Z^U,M_A,\Phi)$ 
      $conf_i \leftarrow M_M(mf_i)$ 
7 // Return most likely label (and conf. estimate) for each node
return { ( $\hat{y}_i, conf_i$ )  $\mid v_i \in V^U$  }

```

**Figure 1:** Pseudocode for ICA, ICA<sub>C</sub>, or ICA<sub>MC</sub>. Based on prior work (McDowell *et al.* 2009), we set n=10 iterations.

for each node  $v_i$ . This is computed by the node classifier  $M_{AR}$  based on the attributes and relational features of  $v_i$ .

ICA<sub>C</sub> performs well on a variety of real and synthetic data, and attains higher accuracies than ICA and similar accuracies as more time-consuming algorithms such as Gibbs sampling or LBP (McDowell *et al.* 2009). However, ICA<sub>C</sub>’s ability to select the “best” predicted labels depends entirely on the confidence value estimates from the node classifier. Accuracy may decrease if a misclassified node is nonetheless assigned a high confidence value.

### Improving ICA<sub>C</sub> with Meta-Caution

To address this potential problem with ICA<sub>C</sub>, we created ICA<sub>MC</sub>. They are identical except that ICA<sub>MC</sub> uses a separate “meta classifier” to predict how likely each prediction  $\hat{y}_i$  is to be correct. Below we describe ICA<sub>MC</sub>’s use of this meta-classifier, methods for generating its training data, and methods for constructing its features.

### ICA<sub>MC</sub>: Inference using *predicted correct* labels

Figure 1 shows that ICA<sub>MC</sub> changes ICA<sub>C</sub> only in step 6. In particular, after using the node classifier to predict the label  $\hat{y}_i$  (and associated confidence  $conf_i$ ) for every node, ICA<sub>MC</sub> computes the meta-feature values and then uses the meta-classifier  $M_M$  to predict how likely  $\hat{y}_i$  is to be correct. These predictions serve as the new confidence values that are then used in Step 3 of the next iteration to select the

committed set  $Y'$ . If the meta-classifier's confidence predictions more accurately identify those nodes whose labels are correctly predicted (compared to  $ICA_C$ 's simple confidence values), then accuracy should increase.

### Generating meta-training data

Learning the meta-classifier requires constructing appropriate meta-training data, which we represent as a set of vectors. Figure 2 shows the pseudocode for this task, whose algorithm employs a holdout graph (a subset of the training set) with nodes  $V$ , edges  $E$ , attributes  $X$ , and true labels  $Y$ . For each of  $T$  trials, step 3 randomly selects  $lp\%$  of the nodes to be *known*; this value is chosen to replicate the fraction of known labels that are present in the test set. It then executes  $ICA_C$  on the graph, given the known nodes (step 4). This yields the set  $Z^U$ , which contains the label predictions and associated confidence values for each node in  $V^U$ . Using these and the expected class distribution  $\Phi$  (from the training set), it then generates a meta-training vector per node (steps 5-7). This vector includes eight meta-features (described later) and a Boolean value that indicates whether prediction  $\hat{y}_i$  is correct. This training data is later used to learn the meta-classifier that predicts the correctness of the  $\hat{y}_i$  estimates given the values of the meta-features.

We set  $T=10$  to conduct ten trials with different *known* nodes each time. The goal is to reduce the bias that might otherwise occur due to the particular selection of  $Y^K$  in step 3. We later compare this with the one-trial approach ( $T=1$ ).

### Generating meta-features from meta-training data

$ICA_{MC}$  needs useful meta-features to predict when the node classifier has *correctly* classified a node. The constructed features are based on two key premises. First, we assume that the data exhibits *relational autocorrelation* (correlation of class labels among interrelated nodes, Jensen *et al.*, 2004) for use by the node classifier. Thus, each node's predicted label will be influenced by the predictions of its neighboring labels. Second, since  $ICA_{MC}$  (like  $ICA_C$ ) exploits only some of the predicted labels during each iteration, not all neighbor labels will affect the prediction for  $v_i$ . We assume that the accuracy of prediction  $\hat{y}_i$  for iteration  $j$  is affected only by the neighbors of  $v_i$  that were included in the committed set  $Y$  during that same iteration. Let  $N_i$  refer to the set of such neighbors for  $v_i$ .

Based on these two premises and additional intuitions described below, we designed eight features for this initial study of  $ICA_{MC}$ . The first three features are based on ones used by Bilgic and Getoor (2008) for a related problem that is discussed later. Future work should examine these choices and others in more detail.

Suppose the CC algorithm predicts  $\hat{y}_i$  to be the label for node  $v_i$ , with confidence  $conf_i$ . Then  $v_i$ 's features are:

1. *Local score*: The CC algorithm's predictions should differ from those of an attribute-only classifier (e.g.,  $M_A$  in Figure 1), or there is no point in executing CC. However, if  $M_A$  and the node classifier  $M_{AR}$  agree on a prediction, then it is more likely to be correct. This

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generateMetaVectors(V,E,X,Y,MAR,MA,n,T,lp,Φ) =
// V=nodes, E=edges, X=attribute vectors, Y=node labels
// MAR = node classifier (uses attrs. & relats), MA = classifier
// (attrs. only), n=# ICAC iters., T=# randomized trials to use
// lp=labeled proportion, Φ = expected class distribution
1 MetaTrainVecs ← ∅
2 for j=1 to T do
3 // Randomly select some nodes to be "known"
  YK ← randomSelectSomeNodes(V, Y, lp) // Randomize
  VU ← {vi | ∃yi ∈ Y-YK} // Nodes used for prediction
4 // Run ICAC to predict labels and compute confidences
  ZU ← ICA_classify(V,E,X,YK,MAR,MA,∅,n,ICAC,Φ)
5 for each vi ∈ VU do // Calc. and store meta-feature vectors
6   mfi ← calcMetaFeatures(i,V,E,X,Y,YK,ZU,MA,Φ)
7   MetaTrainVecs ← MetaTrainVecs ∪ mfi
8 return MetaTrainVecs // return all vectors of meta-features

```

**Figure 2:** Pseudocode to generate training vectors for the meta classifier used by  $ICA_{MC}$ .

heuristic is captured by using, for each  $v_i$ ,  $M_A$ 's confidence value for the  $\hat{y}_i$  that was predicted by  $M_{AR}$ . "Known" nodes are assumed to be fully correct (score of 1), though this could be reduced to account for possible noise:

$$lf_i = \begin{cases} P(Y_i = \hat{y}_i | x_i) & v_i \notin V^K \\ 1 & v_i \in V^K \end{cases}$$

2. *Relational score*: If a node is surrounded by nodes whose predictions are more likely (e.g., have high  $lf$  scores), then its prediction is also more likely:

$$rf_i = \frac{1}{|N_i|} \sum_{v_j \in N_i} lf_j$$

3. *Global score*: Let  $Prior(c)$  be the fraction of training nodes with class label  $c$ , and  $Posterior(c)$  be the fraction of test set labels predicted as  $c$  by the CC algorithm. If  $Posterior(c)$  is much higher than  $Prior(c)$ , then many nodes with predicted label  $c$  may be incorrect. Thus, the global score measures whether class  $y_i$  is over or under-represented in the posterior distribution:

$$gf_i = \frac{1 + Prior(\hat{y}_i) - Posterior(\hat{y}_i)}{2}$$

4. *Node confidence*: If the node classifier is confident in some prediction  $\hat{y}_i$  (high posterior probability), then this suggests that  $\hat{y}_i$  is more likely to be correct:

$$cf_i = \begin{cases} conf_i & v_i \notin V^K \\ 1 & v_i \in V^K \end{cases}$$

If only this feature is used,  $ICA_{MC}$  devolves to  $ICA_C$ .

5. *Neighbor confidence*: As with the relational score, more confident neighbor predictions suggest that a node's prediction is more likely to be correct:

$$nf_i = \frac{1}{|N_i|} \sum_{v_j \in N_i} cf_j$$

6. *Neighbor agreement*: If most of node  $v_i$ 's neighbors have the same predicted label, this may indicate that  $\hat{y}_i$  is more likely to be correct. Let  $\text{count}_1(N_i)$  and  $\text{count}_2(N_i)$  indicate the count of the two most frequent label predictions in  $N_i$ . If the former value is large and the latter is small, then neighbor agreement is high:

$$\text{naf}_i = \frac{1}{|N_i|} (\text{count}_1(N_i) - \text{count}_2(N_i))$$

7. *Known neighbors*: Having many “known” neighbors increases the chances that a node’s prediction is correct:

$$\text{knt}_i = |N_i \cap V^K|$$

8. *Known vicinity*: A node’s prediction may also be influenced by known nodes that are linked to it by one or more intervening nodes. We use a simple measure that favors direct known neighbors, then counts (with reduced weight) any known nodes reached via one additional node  $v'$ :

$$\text{kvt}_i = |N_i \cap V^K| + \frac{1}{2} |v_j \mid \exists v' \in (N_i \cap N_j)|$$

Each of these eight features may not be useful for every dataset. However,  $\text{ICA}_{\text{MC}}$  needs only *some* of the features to be useful – the meta-classifier (we use logistic regression) will learn appropriate parameters for each feature based on their predictive accuracy on the meta-training data. Also, features that provide no benefit are discarded by the feature search process described later.

## Evaluation

**Hypotheses.** By default,  $\text{ICA}_{\text{MC}}$  uses feature search and ten randomized training data trials. This  $\text{ICA}_{\text{MC}}$  attains higher accuracies than  $\text{ICA}_{\text{C}}$  (Hypothesis #1),  $\text{ICA}_{\text{MC}}$  without such trials (#2),  $\text{ICA}_{\text{MC}}$  without feature search (#3), and  $\text{ICA}_{\text{MC}}$  with just the three features used by Bilgic and Getoor (#4).

**Data Sets.** We used the following data sets (see Table 1):

1. *Cora* (see Sen *et al.* 2008): A collection of machine learning papers categorized into seven classes.
2. *CiteSeer* (see Sen *et al.* 2008): A collection of research papers drawn from the CiteSeer collection.
3. *WebKB* (see Neville and Jensen 2007): A collection of web pages from four computer science departments.
4. *Synthetic*: We generate synthetic data using Sen *et al.*'s (2008) graph generator. Similar to their defaults, we use a degree of homophily of 0.7 and a link density of 0.4.

**Table 1:** Data sets summary

Characteristics	Cora	CiteSeer	WebKB	Syn.
Total nodes	2708	3312	1541	n.a.
Avg. # nodes per test set	400	400	385	250
Avg. links per node	2.7	2.7	6	3.3
Class labels	7	6	6	5
Non-rel. features avail.	1433	3703	100	10
Non-rel. features used	10	10	10	10
Relational features used	2	2	3	1
Folds	5	5	4	25

**Feature Representation.** Our node representation includes relational features and non-relational attributes, as described below.

*Non-relational (content) attributes:* The real datasets are all textual. We use a bag-of-words representation for the textual content of each node, where the feature corresponding to a word is assigned *true* if it occurs in the node and *false* otherwise.

Our version of the WebKB dataset has 100 words available. For Cora and CiteSeer, we used information gain to select the 100 highest-scoring words, based on McDowell *et al.* (2007), which reported that using more did not improve performance. Our focus is on the case where relatively few attributes are available (or the attributes are not very predictive) as may occur in large real-world networks (c.f., Macskassy and Provost 2007, Gallagher *et al.* 2008). Thus, for most of our experiments we randomly select 10 of the 100 available words to use as attributes. We also briefly discuss results when using 100 attributes. For the synthetic data, ten binary attributes are generated using the technique described by McDowell *et al.* (2009). This model has a parameter *ap* (attribute predictiveness) that ranges from 0.0 to 1.0; it indicates how strongly predictive the attributes are of the class label. We evaluate *ap* using the values {0.2, 0.4, 0.6}.

*Relational features:* Each relational feature value is a multiset. For instance, a possible feature value is {3 A, 2 B, 1 missing}, which indicates that a node links to 3 other nodes whose predicted label is A, 2 nodes whose prediction is B, and 1 node labeled *missing*. During inference, each label in the multiset (excluding *missing* labels) is separately used to update the probability that a node has label *c*. This is the “independent value” approach that was introduced by Neville *et al.* (2003), used by Neville and Jensen (2007), and shown to be superior to “count” or “proportion” features by McDowell *et al.* (2009). See Neville *et al.* (2003) for more details.

For Cora and CiteSeer, we compute a “multiset” feature using only incoming links, and a separate such feature using only outgoing links. For WebKB, we also compute one such feature using “co-citation” links (a co-citation link exists between nodes *i* and *j* if some node *k* links to both of them). For the synthetic data, the links are undirected, so there is a single relational feature.

**Classifiers.** For the node classifier, we used a naïve Bayes classifier. McDowell *et al.* (2009) reported that, using multiset features, it attained higher accuracies than did alternatives such as logistic regression. For the meta-classifier, we used logistic regression, as did Bilgic and Getoor (2008). Future work should consider other choices.

**Test Procedure.** We conducted an n-fold cross-validation study for each tested algorithm. For WebKB, we treated each of the four schools as a separate fold. For Cora and CiteSeer, we created five disjoint test sets by using “similarity-driven snowball sampling” (McDowell *et al.* 2009). This is similar to the approach of Sen *et al.* (2008).

For all 3 datasets we tested on one graph, trained on two others, and used the remaining two (one for WebKB) as a holdout set for learning the meta-classifier and performing the meta-feature search.

For the synthetic data, we performed 25 separate trials. For each trial we generated three disjoint graphs: one test set, one training set, and one holdout set.

We randomly selected  $lp=10\%$  of each test set to form  $V^K$  (nodes with known labels). This is a “sparsely labeled” task, which is common in real data (Gallagher *et al.* 2008).

To search for which of the eight meta-features to use with  $ICA_{MC}$ , we use the simple, greedy Backwards Sequential Elimination (BSE) algorithm (Kittler, 1986). It evaluates accuracy on the holdout set with  $ICA_{MC}$ , then recursively eliminates any meta-feature whose removal increases accuracy. To increase robustness, accuracy is averaged over ten executions of  $ICA_{MC}$ , each time using a different set of initial “known” labels (as done for  $T=10$  in Figure 2). The final set of meta-features is used for testing.

**Tested Algorithms.** We tested ICA,  $ICA_C$ , and  $ICA_{MC}$ . In addition, to assess the utility of  $ICA_{MC}$ ’s design decisions, we also tested three of its ablated variants:

1. “1 trial instead of 10”: this uses only one randomized trial to collect meta-training data (i.e.,  $T=1$  in Figure 2) and only one evaluation trial for the meta-feature search.
2. “No meta-feature search”: This skips search and uses all eight meta-features that were previously described.
3. “Only Bilgic meta-feats”: This uses just features #1, #2, and #3 – the set used by Bilgic and Getoor (2008).

**Performance Measure.** We compared all the algorithms on their average classification error rate on the test sets.

**Analysis.** We performed independent analyses for each prediction task and joint analyses by pooling the observations, either for all the real data sets or for all the synthetic data conditions shown. Our analysis uses one-tailed paired t-tests accepted at the 95% confidence level.

**Results.** Table 2 displays the classification error rates averaged over all the folds for each algorithm. For each (data set, algorithm) pair, the best result is shown in bold.

*Result 1:  $ICA_{MC}$  significantly outperforms  $ICA_C$  and ICA when attribute predictiveness is low:* Comparing  $ICA_{MC}$  with  $ICA_C$ , we find that  $ICA_{MC}$  reduces classification error by 2.3-8.0% for the real data, and 1.9-6.9% for the synthetic data. This improvement is significant in every case ( $p < .03$  for the real data and  $p < .045$  for the synthetic data). In addition, the pooled analyses found significant gains for both the real and synthetic data. Therefore, we accept Hypothesis #1.

For the synthetic data, the gains clearly decrease as attribute predictiveness ( $ap$ ) increases. This is consistent with the results of McDowell *et al.* (2009), who report that the cautious use of relational information is more important for CC algorithms when  $ap$  and/or the number of attributes is small. Since  $ICA_{MC}$  is even more cautious than  $ICA_C$ ,  $ICA_{MC}$  has larger gains over  $ICA_C$  when  $ap$  is small (the same relative trend exists between  $ICA_C$  and the non-

**Table 2:** Average % classification error rate

Core Algorithms	“Real” datasets			Synthetic data		
	Cora	CS	Web KB	$ap=.2$	$ap=.4$	$ap=.6$
ICA	51.5 <sup>†</sup>	61.0 <sup>†</sup>	60.3 <sup>†</sup>	53.3 <sup>†</sup>	35.9 <sup>†</sup>	22.6 <sup>†</sup>
$ICA_C$	36.2 <sup>†</sup>	37.6 <sup>†</sup>	32.5 <sup>†</sup>	38.8 <sup>†</sup>	27.8 <sup>†</sup>	18.3 <sup>†</sup>
$ICA_{MC}$	<b>31.3</b>	<b>35.3</b>	<b>24.5</b>	<b>31.9</b>	25.0	16.4
Gain*	4.9	2.3	8.0	6.9	2.8	1.9
<b>Variants of <math>ICA_{MC}</math></b>						
1 trial instead of 10	35.4 <sup>†</sup>	35.8	30.0 <sup>†</sup>	36.4 <sup>†</sup>	27.5 <sup>†</sup>	17.2
No meta-feat. search	35.9 <sup>†</sup>	37.6 <sup>†</sup>	31.5 <sup>†</sup>	33.5	<b>24.9</b>	<b>16.2</b>
Only Bilgic meta-feats	42.1 <sup>†</sup>	47.1 <sup>†</sup>	26.4	37.3 <sup>†</sup>	27.8 <sup>†</sup>	18.2 <sup>†</sup>

<sup>†</sup> indicates significantly worse behavior than  $ICA_{MC}$ .

\* indicates gain from meta-caution ( $ICA_C - ICA_{MC}$ )

cautious ICA). Nonetheless,  $ICA_{MC}$  continues to provide a small gain even when  $ap$  is high – a gain of 0.9% when  $ap=0.8$  (results not shown).

For the real data,  $ICA_{MC}$  provides gains for all three datasets, where the largest gain is with WebKB. WebKB has more complex and numerous linking patterns (Macskassy and Provost 2007). For this reason,  $ICA_{MC}$ ’s careful selection of which neighboring labels to use for prediction may be especially important with WebKB.

We repeated these experiments with real data using 2, 5, or 20 attributes (instead of 10) and found similar results. In every case pooled analyses found a significant gain for  $ICA_{MC}$  over  $ICA_C$  (average gains ranging from 3.2- 6.9%), with the largest gains occurring with WebKB. As with the synthetic data, these gains diminish when the attributes are more predictive. For instance, when 100 attributes are used the gains of  $ICA_{MC}$  remained but were small (0.2-1.0%) and statistically insignificant. These results suggest that  $ICA_{MC}$  is especially helpful when the attributes alone are not very predictive, and at least does no harm otherwise.

*Result 2:  $ICA_{MC}$  with randomized trials and meta-feature search outperforms simpler variants:* The bottom of Table 2 shows results with the variants of  $ICA_{MC}$  that do not use multiple randomized trials or do less or no meta-feature search.  $ICA_{MC}$  outperforms the “1 trial instead of 10” and “Only Bilgic meta-feats” variants, often significantly, and pooled analyses find that  $ICA_{MC}$  outperforms both, for the real and for the synthetic data. Thus, we accept Hypotheses #2 and #4.  $ICA_{MC}$  also significantly outperforms the variant that uses all eight meta-features (“No meta-feat. search”) for the real data, but not for the synthetic data (perhaps because simpler, undirected linking patterns were used in the synthetic data). Thus, we reject Hypothesis #3.

Despite the rejection of one hypothesis,  $ICA_{MC}$  always outperformed all three variants (or lagged by at most 0.2%) and significantly outperformed all three variants on the real datasets. Some of the variants that simplify  $ICA_{MC}$ ’s search process sometimes performed notably worse than even  $ICA_C$ . Together, these results suggest that the complete  $ICA_{MC}$ , with randomized trials and feature search, is the most robust performer.

## Discussion

ICA<sub>MC</sub> increased accuracy compared to ICA and ICA<sub>C</sub>. However, *why* does ICA<sub>MC</sub>'s meta-classifier more effectively identify reliable predictions than does ICA<sub>C</sub>'s node classifier? First, the meta-classifier's task is simpler: choosing between two values (correct or incorrect) vs. between all possible class labels. Second, the meta-classifier can use additional information, such as the number of known labels, which has no obvious utility for predicting a particular label, but does help estimate the *correctness* of the resultant prediction. Finally, using two different classifiers helps to reduce the bias due to using the Naïve Bayes node classifier alone.

Meta-feature search often significantly increased ICA<sub>MC</sub>'s accuracy. However, is the same set of features almost always chosen? On average, the "global score" and "node confidence" features were selected most often, and "known neighbor" least often. This varied substantially, however, with some features selected 90% of the time for one dataset and never for another. These results, combined with the results from Table 2, suggest that search is essential to make ICA<sub>MC</sub> robust across different data, even if the default set of meta-features is further refined.

We are not aware of any other work that uses a meta-classifier to improve the operation of a CC inference algorithm, although Bilgic and Getoor (2008) did use a similar predictor to identify the *least likely* CC label predictions (in order to "purchase" the correct labels for them). In contrast, we seek the *most likely* predictions (to favor them for inference). They considered three features for this different task, which our search algorithm selected for ICA<sub>MC</sub> 62%, 67%, and 91% of the time, respectively. Thus, their features are also useful for our task, although the results of the previous section show that using *only* those features leads to very poor performance for ICA<sub>MC</sub>.

Compared to ICA<sub>C</sub>, ICA<sub>MC</sub> requires additional computation: to execute ICA<sub>C</sub> when collecting meta-training data, to execute ICA<sub>MC</sub> for feature selection, and to train the meta-classifier for each combination of meta-features that are considered. However, in many real-world graphs each node links to at most  $k$  other nodes, in which case each of these steps is linear in the number of nodes. In addition, once the meta-classifier is learned, ICA<sub>MC</sub> requires little additional time for inference compared to ICA<sub>C</sub> (i.e., it needs only one additional execution of the meta-classifier per iteration).

## Conclusion

We demonstrated that Meta-Cautious ICA (ICA<sub>MC</sub>) significantly outperforms ICA<sub>C</sub> for many tasks. Moreover, we showed that aspects of ICA<sub>MC</sub> – in particular, its use of multiple randomized training data trials and its use of search for selecting meta-features – were essential to achieving performance that was robust across a range of datasets. Since ICA<sub>C</sub> has already been shown to be a very effective CC algorithm, these results suggest that ICA<sub>MC</sub> should be seriously considered for CC applications,

particularly when attributes alone do not yield high predictive accuracy.

Further work is needed to confirm our results using other datasets, meta-features, and classifiers, and to consider how meta-caution might be extended to other CC algorithms. In addition, we intend to consider techniques for further reducing the time complexity of ICA<sub>MC</sub> compared to ICA<sub>C</sub>.

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