Learning Theory Analysis for Association Rules and Sequential Event Prediction

Cynthia Rudin
Sloan School of Management
Massachusetts Institute of Technology
77 Massachusetts Avenue
Cambridge, MA 02139, USA

Benjamin Letham
Operations Research Center
Massachusetts Institute of Technology
77 Massachusetts Avenue
Cambridge, MA 02139, USA

David Madigan
Department of Statistics
Columbia University
1255 Amsterdam Avenue
New York, NY 10027, USA

Editor: John Shawe-Taylor

Abstract

We present a theoretical analysis for prediction algorithms based on association rules. As part of this analysis, we introduce a problem for which rules are particularly natural, called “sequential event prediction.” In sequential event prediction, events in a sequence are revealed one by one, and the goal is to determine which event will next be revealed. The training set is a collection of past sequences of events. An example application is to predict which item will next be placed into a customer’s online shopping cart, given his/her past purchases. In the context of this problem, algorithms based on association rules have distinct advantages over classical statistical and machine learning methods: they look at correlations based on subsets of co-occurring past events (items a and b imply item c), they can be applied to the sequential event prediction problem in a natural way, they can potentially handle the “cold start” problem where the training set is small, and they yield interpretable predictions. In this work, we present two algorithms that incorporate association rules. These algorithms can be used both for sequential event prediction and for supervised classification, and they are simple enough that they can possibly be understood by users, customers, patients, managers, etc. We provide generalization guarantees on these algorithms based on algorithmic stability analysis from statistical learning theory. We include a discussion of the strict minimum support threshold often used in association rule mining, and introduce an “adjusted confidence” measure that provides a weaker minimum support condition that has advantages over the strict minimum support. The paper brings together ideas from statistical learning theory, association rule mining and Bayesian analysis.

Keywords: statistical learning theory, algorithmic stability, association rules, sequence prediction, associative classification

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1. Introduction

Consider the problem of predicting the next event within a current event sequence, given a “sequence database” of past event sequences to learn from. We might wish to do this, for instance, using data generated by a customer placing items into the virtual basket of an online grocery store such as NYC’s Fresh Direct, Peapod by Stop & Shop, or Roche Bros. The customer adds items one by one into the current basket, creating a sequence of events. The customer has identified him- or herself, so that all past orders are known. After each item selection, a confirmation screen contains a small list of recommendations for items that are not already in the basket. If the store can find patterns within the customer’s past purchases, it may be able to accurately recommend the next item that the customer will add to the basket. Another example is to predict each next symptom of a sick patient, given the patient’s past sequence of symptoms and treatments, and a database of the timelines of symptoms and treatments for other patients. We call the problem of predicting these sequentially revealed events based on past sequences of events “sequential event prediction.”

In these examples, a subset of past events (for instance, a set of ingredients for a particular recipe) can be useful in predicting the next event. In order to make predictions using subsets of past events, we employ association rules (Agrawal et al., 1993). An association rule in this setting is an implication \( a \rightarrow b \) (such as lettuce and carrots \( \rightarrow \) tomatoes), where \( a \) is a subset of items, and \( b \) is a single item. The association rule approach has the distinct advantage in being able to directly model underlying conditional probabilities \( P(b|a) \) eschewing the linearity assumptions underlying many classical supervised classification, regression, and ranking methods. Rules also yield predictive models that are interpretable, meaning that for the rule \( a \rightarrow b \), it is clear that \( b \) was recommended because \( a \) is satisfied.

The association rules approach makes predictions from subsets of co-occurring past events. Using subsets may make the estimation problem much easier, because it helps avoid problems with the curse of dimensionality. For instance \( P(\text{tomatoes} | \text{lettuce and carrots}) \) could be much easier to estimate than \( P(\text{tomatoes} | \text{lettuce, carrots, pears, potatoes, ketchup, eggs, bread, etc.}) \). This is precisely why learning algorithms created from rules can be helpful for the “cold start” problem in recommender systems, where predictions need to be made when there are not enough data available to accurately compute the full probability of a new item being purchased.

There are two main contributions in this work: a generalization analysis for association-rule-based algorithms, and a formal definition of the problem of sequential event prediction. An important part of the rule-based analysis is how a fundamental property of a rule, namely the “support,” is incorporated into the generalization bounds. The “support” of an itemset is the number of times that the itemset has appeared in the sequence database. For instance, the support of lettuce is the number of times lettuce has been purchased in the past. Typically in association rule mining, a strict minimum support threshold condition is placed on the support of itemsets within a rule, so that rules falling below the minimum support threshold are simply discarded. The idea of a condition on the support is not shared with other types of supervised learning algorithms, since they do not use subsets in the same way as when using rules. Thus a new aspect of generalization is explored in this analysis in that it handles predictions created from subsets of data. In classical supervised learning paradigms, bounds scale only with the sample size, and a large sample is necessary to create a generalization guarantee. In the context of association rules, the minimum support threshold forces predictions to be made only when there are enough data. Thus, in the association rules analysis, there are now two mechanisms for generalization: first a large sample, and second,
a minimum support. These are separate mechanisms, in the sense that it is possible to generalize with a somewhat small sample size and a large minimum support threshold, and it is also possible to generalize with a large sample size and no support threshold. We thus derive two types of bounds: large sample bounds, which scale with the sample size, and small sample bounds, which scale with the minimum support of rules. Using both large and small sample bounds (that is, the minimum of the two bounds) gives a complete picture. The large sample bounds are of order $O(\sqrt{1/m})$ as in classical analysis of supervised learning, where $m$ denotes the number of event sequences in the database, that is, the number of past baskets ordered by the online grocery store customer.

Most of our bounds are derived using a specific notion of algorithmic stability called “pointwise hypothesis stability.” The original notions of algorithmic stability were invented in the 1970’s and have been revitalized recently (Devroye and Wagner, 1979; Bousquet and Elisseeff, 2002), the main idea being that algorithms may be better able to generalize if they are insensitive to small changes in the training data such as the removal or change of one training example. The pointwise hypothesis stability specifically considers the average change in loss that will occur at one of the training examples if that example is removed from the training set. Our generalization analysis uses conditions on the minimum support of rules in order to bound the pointwise hypothesis stability.

There are two algorithms considered in this work. At the core of each algorithm is a method for rank-ordering association rules where the list of possible rules is generated using the customer’s past purchase history and subsets of items within the current basket. These algorithms build off of the rule mining literature that has been developing since the early 1990’s (Agrawal et al., 1993) by using an application-specific rule mining method as a subroutine. Our algorithms are interpretable in two different ways: the predictive model coming out of the algorithm is interpretable, and the whole algorithm for producing the predictive model is interpretable. In other words, the algorithms are straightforward enough that they can be understood by users, customers, patients, managers, etc. Further, the rules within the predictive model can provide a simple reason to the customer why an item might be relevant, or identify that a key ingredient is missing from a particular recipe. The rules provide “IF, THEN, ELSE” conditions, and yield models of the same form as those from the expert systems literature from the early days of artificial intelligence (Jackson, 1998). Many authors have emphasized the importance of interpretability and explanation in predictive modeling (see, for example, the work of Madigan et al., 1997).

The first of the two algorithms considered in this work uses a fixed minimum support threshold to exclude rules whose itemsets occur rarely. Then the remaining rules are ranked according to the “confidence,” which for rule $a \rightarrow b$ is the empirical probability that $b$ will be in the basket given that $a$ is in the basket. The right-hand sides of the highest ranked rules will be recommended by the algorithm. However, the use of a strict minimum support threshold is problematic for several well-known reasons, for instance it is known that important rules (“nuggets,” which are rare but strong rules) are often excluded by a minimum support threshold condition.

The other algorithm introduced in this work provides an alternative to the minimum support threshold, in that rules are ranked by an “adjusted” confidence, which is a simple Bayesian shrinkage estimator of the probability of a rule $P(b|a)$. The right-hand sides of rules with the highest adjusted confidence are recommended by the algorithm. For this algorithm, the generalization guarantee is smoothly controlled by a parameter $K$, which provides only a weak (less restrictive) minimum support condition. The key benefits of an algorithm based on the adjusted confidence are that: 1) it allows the possibility of choosing very accurate (high confidence) rules that have appeared very few times in the training set (low support), and 2) given two rules with the same or similar prediction
accuracy on the training set (confidence), the rule that appears more frequently (higher support) achieves a higher adjusted confidence and is thus preferred over the other rule.

All of the bounds are tied to the measure of quality (the loss function) used within the analysis. We would like to directly compare the performance of algorithms for various settings of the adjusted confidence’s $K$ parameter (and for the minimum support threshold $\theta$). It is problematic to have the loss defined using the same $K$ value as the algorithm, in that case we would be using a different method of evaluation for each setting of $K$, and we would not be able to directly compare performance across different settings of $K$. To allow a direct comparison, we select one reference value of the adjusted confidence, called $K_r$ (r for “reference”), and the loss depends on $K_r$ rather than on $K$. The bounds are written generally in terms of $K_r$. The special case $K_r = 0$ is where the algorithm is evaluated with respect to the confidence measure. The small sample bounds for the adjusted confidence algorithm have two terms: one that generally decreases with $K$ (as the support increases, there is better generalization) and the other that decreases as $K$ gets closer to $K_r$ (better generalization as the algorithm is closer to the way it is being measured). These two terms are thus agreeing if $K_r > K$ and competing if $K_r < K$. In practice, the choice of $K$ can be determined in several ways: $K$ can be manually determined (for instance by the customer), it can be set using side information as considered by McCormick et al. (2012), or it can be set via cross-validation on an extra hold-out set.

The novel elements of the paper include: 1) generalization analysis that incorporates the use of association rules, for both classification and sequential event prediction, 2) the algorithm based on adjusted confidence, where the adjusted confidence is a Bayesian version of the confidence, 3) the definition of a new supervised learning problem, namely sequential event prediction. The work falls in the intersection of several fields that are rarely connected: association rule mining and associative classification, supervised machine learning and generalization bounds from statistical learning theory, and Bayesian analysis.

In terms of applications, the definition of “sequential event prediction” was inspired by, but not restricted to, online grocery stores. Examples are Fresh Direct, Amazon.com grocery, and netgrocer.com. Many supermarket chains with local outlets also offer an online shop-and-delivery option, such as Peapod (paired with Stop & Shop and Giant). Other online retailers and recommendation engines may benefit from ranking algorithms that are transparent to the user like amazon.com’s “customers who purchased this also purchased that” recommender system. The same techniques used to solve the sequential event prediction problem could be used in medical applications to predict, for instance, the winners at each round of a tournament (e.g., the winners of games in a football season), or the next move of a video game player in order to design a more interesting game. The work of McCormick et al. (2012) contains a Bayesian algorithm, based on the analysis introduced in this paper, for predicting conditions of medical patients in a clinical trial. The work of Letham et al. (2013b) uses empirical risk miniimization to solve sequential event prediction problems dealing with email recipient recommendation, healthcare, and cooking.

Section 2 describes the two rule-based prediction algorithms, one based on a hard thresholding of the support (min support) and the other based on the soft thresholding (adjusted confidence). Section 3 formally defines sequential event prediction. Section 4 provides the generalization analysis, Section 5 contains proofs, and Section 6 provides experimental validation. Section 7 contains a summary of relevant literature. Appendix A discusses the suitability of regression approaches for solving the sequential event prediction problem. Appendix B provides additional experimental results. Appendix C contains an additional proof.
2. Derivation of Algorithms

We assume an interface similar to that of Fresh Direct, where users add items one by one into the basket. After each selection, a confirmation screen contains a handful of recommendations for items that are not already in the customer’s basket. The customer’s past orders are known.

The set of items is $X$, for instance $X = \{\text{apples, bananas, pears, etc}\}$; $X$ is the set of possible events. The customer has a past history of orders $S$ which is a collection of $m$ baskets, $S = \{z_i\}_{i=1,...,m}$, $z_i \subseteq X$; $S$ is the sequence database. The customer’s current basket is usually denoted by $B \subseteq X$; $B$ is the current sequence. An algorithm uses $B$ and $S$ to find rules $a \rightarrow b$, where $a$ is in the basket and $b$ is not in the basket. For instance, if salsa and guacamole are in the basket $B$ and also if salsa, guacamole and tortilla chips were often purchased together in $S$, then the rule (salsa and guacamole) $\rightarrow$ tortilla chips might be used to recommend tortilla chips.

The support of $a$, written $\text{Sup}(a)$ or $\#a$, is the number of times in the past the customer has ordered itemset $a$,

$$\text{Sup}(a) := \#a := \sum_{i=1}^{m} \mathbb{1}_{[a \subseteq z_i]}.$$

If $a = \emptyset$, meaning $a$ contains no items, then $\#a := \sum_{i=1}^{m} 1 = m$. The confidence of a rule $a \rightarrow b$ is denoted “Conf” or “$f_{S,0}$”:

$$\text{Conf}(a \rightarrow b) := f_{S,0}(a,b) := \frac{\#(a \cup b)}{\#a},$$

the fraction of times $b$ is purchased given that $a$ is purchased. It is an estimate of the conditional probability of $b$ given $a$. Ultimately an algorithm should order rules by conditional probability; however, the rules that possess the highest confidence values often have a left-hand side with small support, and their confidence values do not yield good estimates for the true conditional probabilities. Note that $a \cup b$ is the union of the set $a$ with item $b$ (the intersection is empty). In this work we introduce the “adjusted” confidence as a remedy for this problem: The adjusted confidence for rule $a \rightarrow b$ is:

$$f_{S,K}(a,b) := \frac{\#(a \cup b)}{\#a + K}.$$ 

The adjusted confidence for $K = 0$ is equivalent to the confidence.

The adjusted confidence is a particular Bayesian estimate of the confidence. Specifically, assuming a beta prior distribution for the confidence, the posterior mean is given by:

$$\hat{p} = \frac{L + \#(a \cup b)}{L + K + \#a},$$

where $L$ and $K$ denote the parameters of the beta prior distribution. The beta distribution is the “conjugate” prior distribution for a binomial likelihood. For the adjusted confidence we choose $L = 0$. This choice yields the benefits of the lower bounds derived in the remainder of this section, and the stability properties described later. The prior for the adjusted confidence tends to bias rules towards the bottom of the ranked list. Any rule achieving a high adjusted confidence must overcome this bias.

Other possible choices for $L$ and $K$ are meaningful. For instance we could choose the following:
• Collaborative filtering prior: have \( L/(L+K) \) represent the probability of purchasing item \( b \) given that item \( a \) was purchased, calculated over a subset of other customers. This biases estimates of the target user’s behavior towards the “average” user.

• Revenue management prior: choose \( L \) and \( K \) based on the item’s price, so more expensive items are more likely to be recommended.

• Time dependent prior: use only the customer’s most recent orders, and choose \( L \) and \( K \) to summarize the user’s behavior before this point.

A rule cannot have a high adjusted confidence unless it has a large enough confidence and also a large enough support on the left-hand side. To see this, consider the case when we take \( f_{S,K}(a,b) \) large, meaning for some \( \eta \), we have \( f_{S,K}(a,b) > \eta \), implying:

\[
\text{Conf}(a \rightarrow b) = \frac{f_{S,0}(a,b)}{\#a} > \eta \frac{\#a + K}{\#a} \geq \eta,
\]

\[
\text{Sup}(a) = \#a \geq (\#a + K) \frac{\#(a \cup b)}{\#a + K} > (\#a + K)\eta, \text{ implying } \text{Sup}(a) = \#a > \frac{\eta K}{1 - \eta}.
\]

And further, expression (1) implies:

\[
\text{Sup}(a \cup b) = \#(a \cup b) > \eta(\#a + K) > \eta K/(1 - \eta).
\]

Thus, rules attaining high values of adjusted confidence have a lower bound on confidence, and a lower bound on support of both the right and left-hand sides, which means a better estimate of the conditional probability. The bounds clearly do not provide any advantage when \( K = 0 \) and the confidence is used.

As \( K \) increases, rules with low support are heavily penalized, so they tend not to be at the top of the list. On the other hand, such rules might be chosen when all other rules have low confidence. That is an advantage of having no firm minimum support cutoff: “nuggets” that have fairly low support may filter to the top. Figure 1 illustrates this by showing the support of rules ordered by adjusted confidence, for two values of \( K \), using a transactional data set “T25I10D10KN200” from the IBM Quest Market-Basket Synthetic Data Generator (Agrawal and Srikant, 1994) which mimics a retail data set.\(^1\) We use all rules with either one or no items on the left and one item on the right (as produced for instance by GenRules, presented in Algorithm 1). On each scatter plot, each of the rules is represented by a point. The rules are ordered on the x-axis by adjusted confidence, and the support of the rule is indicated on the y-axis. As \( K \) increases, rules with the highest adjusted confidence are required to achieve a higher support, as can be seen from the gap in the lower left corner of the scatter plot for larger \( K \).

We now formally state the recommendation algorithms. Both algorithms use a subroutine for mining association rules to generate a set of candidate rules. GenRules (Algorithm 1) is one of the simplest such rule mining algorithms, which in practice should be replaced by a rule mining algorithm that retrieves rules tailored to the application. There is a vast literature on such algorithms since the field of association rule mining evolved on their development, e.g. Apriori (Agrawal et al., 1993). GenRules requires a set \( A \) which is the set of allowed left-hand sides of rules.

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\(^1\) The data set generated is T25I10D10KN200 that contains 10K transactions, 200 items, and where the average length of transactions is 25 and the average pattern length is 10.
Figure 1: Support vs. rank in adjusted confidence for $K = 0, 10, 50$. Rules with the highest adjusted confidence are on the left.

Algorithm 1: Subroutine GenRules.

**Input:** $(S, B, X)$, that is, past orders $S = \{z_i\}_{i=1,\ldots,m}$, current basket $B \subset X$, set of items $X$

**Output:** Set of all rules $\{a_j \rightarrow b_j\}_j$ where $b_j$ is a single item that is not in the basket $B$, and where $a_j$ is either a subset of items in the basket $B$, or else it is the empty set. Also the left-hand side $a_j$ must be allowed (meaning it is in $A$). That is, output rules $\{a_j \rightarrow b_j\}_j$ such that $b_j \in X \setminus B$ and $a_j \subseteq B \subset X$ with $a_j \in A$, or $a_j = \emptyset$.

2.1 Max Confidence, Min Support Algorithm

The max confidence, min support algorithm, shown as Algorithm 2, is based on the idea of eliminating rules whose itemsets occur rarely, which is commonly done in the rule-mining literature. For this algorithm, the rules are ranked by confidence, and rules that do not achieve a predetermined fixed minimum support threshold are completely omitted. The algorithm recommends the right-hand sides from the top ranked rules. Specifically, if $c$ items are to be recommended to the user, the algorithm picks the top ranked $c$ distinct items.

It is common that the minimum support threshold is imposed on the right and left side $\text{Sup}(a \cup b) \geq \theta$; however, as long as $\text{Sup}(a)$ is large, we can get a reasonable estimate of $P(b|a)$. In that sense, it is sufficient (and less restrictive) to impose the minimum support threshold on the left side: $\text{Sup}(a) \geq \theta$. Here $\theta$ is a number determined beforehand (for instance, the support of the left must be at least 5 items). In this work, we only have a required minimum support on the left side. As a technical note, we might worry about the minimum support threshold being so high that there are no rules that meet the threshold. This is actually not a major concern because of the minimum support being imposed only on the left-hand side: as long as $m \geq \theta$, all rules $\emptyset \rightarrow b$ meet the minimum support threshold.

The thresholded confidence is denoted by $\bar{f}_{S,\theta}$:

$$\bar{f}_{S,\theta}(a, b) := f_{S,0}(a, b) \text{ if } \#a \geq \theta, \text{ and } \bar{f}_{S,\theta}(a, b) := 0 \text{ otherwise.}$$
2.2 Adjusted Confidence Algorithm

The adjusted confidence algorithm is shown as Algorithm 3. A chosen value of $K$ is used to compute the adjusted confidence for each rule, and rules are then ranked according to adjusted confidence.

The definition of the adjusted confidence makes an implicit assumption that the order in which items were placed into previous baskets is irrelevant. It is easy to include a dependence on the order by defining a “directed” version of the adjusted confidence, and calculations can be adapted accordingly. The numerator of the adjusted confidence becomes the number of past orders where $a$ is placed in the basket before $b$.

$$f_{S,K}^{(directed)}(a,b) = \frac{\#\{(a \cup b) : b \text{ follows } a\}}{\#a + K}.$$  

2.3 Rule Selection

In classical supervised machine learning problems, like classification and regression, designing features is one of the main engineering challenges. In association rule modeling, the analogous challenge is designing the allowed sets of items for the left and right sides of rules. For instance, we can choose to capture only positive correlations, as if customers were purchasing items from several independent recipes. The present work considers mainly positive correlations, for the purpose of exposition and to keep things simple. Beyond this, it is easily possible to capture negative corre-
lations between items by creating “negation” items, such as \( \neg b \). As an example of using negation rules in the ice cream category, we impose that for vanilla to be on the right, both chocolate and strawberry need to be on the left, in either their usual form or negated. Of these, the rule that is used corresponds to the current basket. In that case, \( \neg \)chocolate, \( \neg \)strawberry \( \rightarrow \) vanilla could have a high score in order to recommend vanilla when chocolate and strawberry are not in the basket, whereas chocolate, \( \neg \)strawberry \( \rightarrow \) vanilla might have a low score, conveying that since chocolate is already in the basket that vanilla should not be recommended. Alternatively, we could create a negation item \( \neg \)ice cream indicating that the basket contains no ice cream presently, so sprinkles + \( \neg \)ice cream \( \rightarrow \) vanilla could have a high score.

We can also use negation items on the right, where if there is a rule \( a \rightarrow \neg b \) that receives a higher score (confidence or adjusted confidence) than any other rules recommending \( b \), we can choose not to recommend \( b \). Rules can be designed to capture higher level correlations in specific regimes, for instance the allowed set \( A \) can contain up to three items in one product category, but only two items in another. It is not practical in general to exhaustively enumerate and use all possible rules in a rule modeling algorithm due to problems with computational complexity. The key is to find a small but good set of rules, for instance the set of rules containing exhaustively all subsets of 1, 2, or 3 items on the left; or perhaps use the top rules that come out of the Apriori algorithm (Agrawal et al., 1993). In Section 7 we provide citations to surveys on association rule mining and associative classification that discuss this important issue of rule-construction and rule-engineering.

2.4 Modeling Assumption

The general modeling assumption that we make with the two algorithms above can be written as follows, where current basket \( B \) is composed of items \( b_1, \ldots, b_t \), and \( X_i \) is the random variable governing whether item \( i \) will be placed into the basket next:

\[
\arg\max_{i=1, \ldots, m} P(X_i = 1 | X_{b_1} = 1, X_{b_2} = 1, \ldots, X_{b_t} = 1) = \arg\max_{i=1, \ldots, m} \max_{a \subseteq \{b_1, \ldots, b_t\}} P(X_i = 1 | X_{a_1} = 1, X_{a_2} = 1, \ldots).
\]

This expression states that the most likely item to be added next into the basket can be identified using a subset of items in the basket, denoted \( a \). That subset is restricted to fall into a class \( A \) which is chosen based on the application at hand and the ease in which that subset can be searched. The set \( A \) determines the hypothesis space for learning, and it would be chosen differently as we move from the small sample regime to the large sample regime, so that the right side of this expression would eventually look just like the left side when the sample is large.

The choice of \( A \) can help with the problem of “curse of dimensionality” by allowing us to look at small subsets on the left. A similar example to the one in the introduction is \( P( \text{machine will break} | \text{a particular part is old} ) \) could be much easier to estimate accurately than the full probability \( P( \text{machine will break} | \text{part 1 did poorly at last inspection, part 2 is very old, part 3 is new, part 4 is ok, part 612 is ok, etc.} ) \). The large dimensionality would likely be a problem when estimating the full probability. Further, the approximation also could actually be sufficient to estimate the full probability. We note that there are circumstances in which it is natural to only consider positive correlations. In the example of equipment failure, for instance, individual component failures would always increase the risk of overall failure. More typically, however, consideration of both positive and negative correlations will be important.
Our modeling assumption aligns with sequential event prediction, where only part of a sequence is available to make a prediction at time \( t \). This is a case where standard linear modeling approaches do not naturally apply, since one would need to make a linear combination of terms, some of which are unrealized. We discuss this more in Appendix A.

3. Definition of Sequential Event Prediction

For simplicity in notation, at each time the algorithm recommends only one item, \( c = 1 \). A basket \( z \) consists of an ordered (permuted) set of items, \( z \in 2^X \times \Pi \), where \( 2^X \) is the set of all subsets of \( X \), and \( \Pi \) is the set of permutations over at most \( |X| \) elements. We have a training set of \( m \) baskets \( S = \{z_i\}_{i=1}^m \) that are the customer’s past orders. Denote \( z \sim \mathcal{D} \) to mean that basket \( z \) is drawn randomly (iid) according to distribution \( \mathcal{D} \) over the space of possible items in baskets and permutations over those items, \( 2^X \times \Pi \). The \( t^{th} \) item added to the basket is written \( z_{t,i} \), where the dot is just a placeholder for the generic basket \( z \). The \( t^{th} \) element of the \( t^{th} \) basket in the training set is written \( z_{t,i} \). We define the number of items in basket \( z \) by \( T_z \), that is, \( T_z := |z| \). We introduce a generic scoring function \( f_S : (a, b) \mapsto \mathbb{R} \) where \( a \) is a subset of items and \( b \) is a single item. The input \( a \) to the score is \( \{z_1, \ldots, z_{t,i}\} \) or is a subset of \( \{z_1, \ldots, z_{t,i}\} \). For now we let \( a \) be the full set \( \{z_1, \ldots, z_{t,i}\} \). The input \( b \) is an item that is not already in the basket, \( b \in X \setminus \{z_1, \ldots, z_{t,i}\} \). The scoring function \( f_S \) comes from an algorithm that takes data set \( S \) as input. We can consider \( f_S \) to be parameterized, and the algorithm will learn the parameters of \( f_S \) from \( S \).

If the score \( f_S(\{z_1, \ldots, z_{t,i}\}, b) \) is larger than that of \( f_S(\{z_1, \ldots, z_{t,i}\}, z_{t,i+1}) \), it means that the algorithm recommended the wrong item. The loss function below counts the proportion of times this happens for each basket.

\[
\ell_{0-1}(f_S, z) := \frac{1}{T_z} \sum_{i=0}^{T_z-1} \begin{cases} 1 & \text{if } f_S(\{z_1, \ldots, z_{t,i}\}, z_{t,i+1}) - \max_{b \in X \setminus \{z_1, \ldots, z_{t,i}\}} f_S(\{z_1, \ldots, z_{t,i}\}, b) \leq 0 \\ 0 & \text{otherwise.} \end{cases}
\]

(Note that if \( z \) contains all items in \( X \), then the recommendation for the last item is deterministic, so we would not count it towards the loss.) The true error for sequential event prediction is an expectation of the loss with respect to \( \mathcal{D} \), and is again a random variable since the training set \( S \) is random.

\[
\text{TrueErr}(f_S) := \mathbb{E}_{z \sim \mathcal{D}} \ell_{0-1}(f_S, z).
\]

The empirical risk is the average loss with respect to \( S \):

\[
\text{EmpErr}(f_S) := \frac{1}{m} \sum_{i=1}^{m} \ell_{0-1}(f_S, z_i).
\]

The loss is bounded (by 1), the baskets are chosen independently, and the empirical risk is an average of iid random variables and the true risk is the expectation. Thus, the problem fits into the traditional scope of statistical learning, and the loss can be used within concentration arguments to obtain generalization bounds.

In the analysis below, we build the full algorithm for constructing \( f_S \) into the notation. The algorithms above are simple enough that they can be encoded within the same line of notation. To do this we will say that \( f_S \) acts on the the subset of \( \{z_1, \ldots, z_{t,i}\} \) within \( A \) that has the maximum
score. For instance, if we are using the adjusted confidence algorithm,

\[
f_S(\{z_1, \ldots, z_J\}, b) := \max_{a \in A, a \subseteq \{z_1, \ldots, z_J\}} f_{S,K}(a, b).
\]

The 0-1 loss is not smooth, so we will often use a smooth convex upper bound for the loss within the bounds. Specifically, for the way we have defined sequential event prediction, if any item has a higher score than the next item added, the algorithm incurs an error. (Even if that item is added later on, the algorithm incurs an error at this timestep.) To measure the size of that error, we can use the 0-1 loss, indicating whether or not our algorithm gave the highest score to the next item added. However, the 0-1 loss does not capture how close our algorithm was to correctly predicting the next item, though this information might be useful in determining how well the algorithm will generalize. We approximate the 0-1 loss using a modified loss that decays linearly near the discontinuity. This modified loss allows us to consider differences in adjusted confidence, not just whether one is larger than another:

\[
|(\text{adjusted conf. of highest-scoring-correct rule}) - (\text{adjusted conf. of highest-scoring-incorrect rule})|.
\]

However, as discussed in the introduction, if we adjust the loss function’s \( K \) value to match the adjusted confidence \( K \) value, then we cannot fairly compare the algorithm’s performance using two different values of \( K \). An illustration of this point is that for large \( K \), all adjusted confidence values are \( \ll 1 \), and for small \( K \), the adjusted confidence can be \( \approx 1 \); differences in adjusted confidence for small \( K \) cannot be directly compared to those for large \( K \). Since we want to directly compare performance as \( K \) is adjusted, we fix an evaluation measure that is separate from the choice of \( K \). Specifically, we use the difference in adjusted confidence values with respect to a reference \( K_r \):

\[
|(\{\text{adjusted conf.}\}_K, \text{of highest-scoring-correct rule}_K) - (\{\text{adjusted conf.}\}_K, \text{of highest-scoring-incorrect rule}_K)|.
\]

The reference \( K_r \) is a parameter of the loss function, whereas \( K \) is a parameter of an algorithm. We set \( K_r = 0 \) to measure loss using the difference in confidence, and \( K = 0 \) for an algorithm that chooses rules according to the confidence. As \( K \) gets farther from \( K_r \), the algorithm is more distant from the way it is being evaluated, which leads to worse generalization. Note that for \( K_r = K \), the 0-1 loss is the same as the sign of (2).

A similar loss will be used in classification, where we incur an error if the adjusted confidence of the incorrect label is higher than that of the correct label.

4. Generalization

Our goal in this section is to provide a foundation for supervised learning with association rules, and also a foundation for sequential event prediction. We will consider several quantities that may be important in the learning process: \( m \), \( K \) or \( \theta \), the size of the set of possible itemsets \( |A| \), and the probability of the least probable itemsets and items.

As part of this section, we establish bounds for vanilla supervised binary classification with rules. Specifically we consider “max-score” association rule classifiers. For a given example, a max-score classifier assigns a score to the label +1 and a score to the label -1, and chooses the label
corresponding to the higher of the two scores. Max-score association rule classifiers are a special
type of “associative classifier” (Liu et al., 1998) and are also a type of “decision list” (Rivest, 1987).
The result in 4.2 is a uniform bound based on the VC dimension of the set of max-score classifiers.
This bound does not depend explicitly on $K$, which we hypothesize is an important quantity for the
learning process.

In order to understand how $K$ might affect learning, we use algorithmic stability analysis. This
approach originated in the 1970’s (Rogers and Wagner, 1978; Devroeye and Wagner, 1979) and
was revitalized by Bousquet and Elisseeff (2002). Stability bounds depend on how the space of
functions is searched by the algorithm (rather than the size of the function space), so it often yields
more insightful bounds. These bounds are still not often directly useful due to large multiplicative
constants (in our case a factor of 6), but they capture more closely the scalability relationship of
a particular algorithm with respect to important quantities in the learning process. The calculation
required for an algorithmic stability bound is to show that the empirical error will not dramatically
change by altering or removing one of the training examples and re-running the algorithm. There
are many different ways to measure the stability of an algorithm; most of the bounds presented here
use a specific type of algorithmic stability (pointwise hypothesis stability) so that the bounds scale
correctly with the number of training examples $m$.

Section 4.1 presents a basic stability bound for sequential event prediction. Section 4.2 presents
a uniform VC bound for classification with max-score classifiers. Section 4.3 provides notation.
Section 4.4 presents another basic stability bound for sequential event prediction, for a rule-based
loss function. We then focus on stability bounds for the rule-based algorithms provided in Section 2.
Specifically, Section 4.5 provides stability bounds for the large sample asymptotic regime (for both
sequential event prediction and classification). Then we consider the new small $m$ regime in Section
4.6, starting with stability bounds that formally show that minimum support thresholds can lead to
to better generalization (for both sequential event prediction and classification). From there, we present
small sample bounds for the adjusted confidence algorithm, for classification and (separately) for
sequential event prediction.

We note that the space of possible baskets (up to a maximum size) is a combinatorially large,
discrete space. Because the space is discrete, all probability estimates converge to the true proba-
bilities, which means that an algorithm that is statistically consistent can be obtained by estimating
$p(b|B)$ directly for the current basket $B$. If $m$ is large, prediction is easy. The difficult part is when
we have only enough data to well estimate conditionals that are much smaller, $P(b|a), a \subset B$. That
is the problem we are concerned with. Consistency does not imply anything about generalization
bounds for the finite sample case.

4.1 General Stability Bound for Sequential Event Prediction

In this section we provide a basic stability-based bound for sequential event prediction, by analogy
with Theorem 17 of Bousquet and Elisseeff (2002) (B&E).

We define a sequential event prediction algorithm producing $f_S$ to have strong sequential event
prediction stability $\beta$ (by analogy with B&E Definition 15) if the following holds:

$$\forall S \in D^m, \forall i \in \{1, \ldots, m\} \quad \| \max_{t=0, \ldots, T-1} | f_S(\{z_1, \ldots, z_T\}, z_{t+1}) - f_{S,i}(\{z_1, \ldots, z_t\}, z_{t+1}) \|_\infty \leq \beta,$$
where the $\infty$-norm is over baskets. A definition we will use from B&E is as follows: an algorithm producing function $f_S$ with uniform stability $\beta'$ obeys:

$$\forall S, \forall i \in \{1, \ldots, m\}, \|\ell(f_S, \cdot) - \ell(f_{S\setminus i}, \cdot)\|_\infty \leq \beta'.$$

Let us define a modified loss function. Let symbol $\Delta$ temporarily denote $f_S(\{z_1, \ldots, z_J\}, z_{J+1}) - \max_{b \in X \setminus \{z_1, \ldots, z_J\}} f_S(\{z_1, \ldots, z_J\}, b)$ in the expression below. The loss is:

$$\ell(f_S, z) := \frac{1}{T} \sum_{t=0}^{T-1} \left\{ \begin{array}{ll} 1 & \text{if } \Delta \leq 0 \\ 1 - \frac{1}{\gamma} \Delta & \text{if } 0 \leq \Delta \leq \gamma \\ 0 & \text{if } \Delta \geq \gamma. \end{array} \right.$$

The empirical error and leave-one-out error defined for this loss are:

$$\text{EmpErr}_\gamma(f_S, z_i) := \frac{1}{m} \sum_{i=1}^{m} \ell(f_S, z_i),$$

$$\text{LooErr}_\gamma(f_S, z_i) := \frac{1}{m} \sum_{i=1}^{m} \ell(f_{S\setminus i}, z_i).$$

**Lemma 1** A sequential event prediction algorithm producing $f_S$ with strong sequential event prediction stability $\beta$ has uniform stability $2\beta'/\gamma$ with respect to the loss function $\ell_\gamma$.

**Proof**

$$|\ell_\gamma(f_S, z) - \ell_\gamma(f_{S\setminus i}, z)| \leq \frac{1}{T} \sum_{t=0}^{T-1} \frac{1}{\gamma} \left[ f_S(\{z_1, \ldots, z_J\}, z_{J+1}) - \max_{b \in X \setminus \{z_1, \ldots, z_J\}} f_S(\{z_1, \ldots, z_J\}, b) \right]$$

$$- \left[ f_{S\setminus i}(\{z_1, \ldots, z_J\}, z_{J+1}) - \max_{b \in X \setminus \{z_1, \ldots, z_J\}} f_{S\setminus i}(\{z_1, \ldots, z_J\}, b) \right]$$

$$\leq \frac{1}{\gamma} \sum_{t=0}^{T-1} \left[ f_S(\{z_1, \ldots, z_J\}, z_{J+1}) - f_{S\setminus i}(\{z_1, \ldots, z_J\}, z_{J+1}) \right] +$$

$$\left| \max_{b \in X \setminus \{z_1, \ldots, z_J\}} f_S(\{z_1, \ldots, z_J\}, b) - \max_{b \in X \setminus \{z_1, \ldots, z_J\}} f_{S\setminus i}(\{z_1, \ldots, z_J\}, b) \right|$$

$$\leq \frac{1}{\gamma} 2\beta'.$$

The first inequality uses the Lipschitz property of the loss, as well as an upper bound from moving the absolute values inside the sum. The third inequality uses the strong stability with respect to $f_S$. 

The following theorem is analogous to Theorem 17 in B&E, for sequential event prediction. The proof is a direct application of Theorem 12 of B&E to the sequential event prediction loss, combined with Lemma 1.
Theorem 2 Let \( f_S \) be a sequential event prediction algorithm with sequential event stability \( \beta \). Then for all \( \gamma > 0 \) and any \( m \geq 1 \) and any \( \delta \in (0,1) \) with probability at least \( 1 - \delta \) over the random draw of sample \( S \),

\[
\text{TrueErr}(f_S) \leq \text{EmpErr}_\gamma(f_S) + \frac{4\beta}{\gamma} + \left(8m \frac{\beta}{\gamma} + 1\right) \sqrt{\frac{\ln(1/\delta)}{2m}}
\]

and with probability at least \( 1 - \delta \) over the random draw of sample \( S \),

\[
\text{TrueErr}(f_S) \leq \text{LooErr}_\gamma(f_S) + \frac{4\beta}{\gamma} + \left(8m \frac{\beta}{\gamma} + 1\right) \sqrt{\frac{\ln(1/\delta)}{2m}}.
\]

As with classification algorithms, the type of stability one would need to apply these bounds can be quite difficult to achieve, as it requires that the change in the model is small for any training set when any example is removed. This is particularly difficult to achieve when the sample size is somewhat small. For the association rule bounds, we know that uniform stability is not possible for many algorithms that perform well. However, there are some algorithms that do exhibit stronger stability, as we will discuss.

4.2 Classification with Association Rules: A Uniform Bound

In the classification problem, each basket receives a single label that is one of two possible labels \( \{+1, -1\} \). This contrasts with sequential event prediction where there is a sequence of labels, one for each item in the basket as it arrives. For classification, we represent basket \( x \) as a binary vector, where entry \( j \) is 1 if item \( j \) is in the basket. We sample baskets with labels, \( z = (x, y) \), where \( x \in 2^X \) is a set of items (or, equivalently, a binary feature vector) and \( y \in \{-1, 1\} \) is the corresponding label. Each labeled basket \( z \) is chosen randomly (iid) from a fixed (but unknown) probability distribution \( D \) over baskets and labels. Given a training set \( S \) of \( m \) labeled baskets, we wish to construct a classifier that can assign the correct label to new, unlabeled baskets. We begin by defining a scoring function \( g : A \times \{-1, 1\} \rightarrow \mathbb{R} \) that assigns a score \( g(a, y) \) to a rule \( a \rightarrow y \).

The set of left-hand sides \( A \) can be any collection of itemsets so long as every \( x \in 2^X \) contains at least one \( a \in A \). We define a valid scoring function as one where \( \forall a \in A, g(a, 1) \neq g(a, -1) \) and \( \forall a_1, a_2 \in A, \max_{y \in \{-1, 1\}} g(a_1, y) \neq \max_{y \in \{-1, 1\}} g(a_2, y) \), that is, there are no ties. The validity requirement will be discussed in the following paragraph. Define \( G \) to be the class of all valid scoring functions. We now define a class of decision functions that use a valid scoring function \( g \in G \) to provide a label to a basket \( x, f_g : 2^X \rightarrow \{-1, 1\} \). The decision function assigns the label corresponding to the highest scoring rule whose left-hand side is contained in \( x \). Specifically,

\[
f_g(x) = \arg\max_{y \in \{-1, 1\}} \max_{a \in A, a \subseteq x} g(a, y).
\]

We call such a classifier a “max-score association rule classifier” (or “decision list”) because it uses the association rule with the maximum score to perform the classification. Let \( \mathcal{F}_{\text{maxscore}} \) be the class of all max-score association rule classifiers:

\[
\mathcal{F}_{\text{maxscore}} := \{f_g : g \in G\}.
\]

We will bound the VC dimension of class \( \mathcal{F}_{\text{maxscore}} \). By definition, the VC dimension is the size of the largest set of baskets to which arbitrary labels can be assigned using some \( f_g \in \mathcal{F}_{\text{maxscore}} \); it is the size of the largest set that can be shattered.

The argmax in (3) is unique because \( g \) is valid, thus there are no ties. If ties are allowed but broken randomly, arbitrary labels can be realized with some probability, for example by taking
$g(a, y) = 0$ for all $a$ and $y$. In this case the VC dimension can be considered to be infinite, which motivates our definition of a valid scoring function. This problem actually happens with any classification problem where function $f(x) = 0 \forall x$ is within the hypothesis space, thereby allowing all points to sit on the decision boundary. Our definition of validity is equivalent to one in which ties are allowed but are broken deterministically using a pre-determined ordering on the rules. In practice, ties are generally broken in a deterministic way by the computer, so the inclusion of the function $f = 0$ is not problematic.

The true error of the max-score association rule classifier is the expected misclassification error:

$$\text{TrueErrClass}(f_g) := \mathbb{E}_{(x, y) \sim \mathcal{D}}[1[f_g(x) \neq y]].$$

The empirical error is the average misclassification error over a training set of $m$ baskets:

$$\text{EmpErrClass}(f_g) := \frac{1}{m} \sum_{i=1}^{m} [1[f_g(x_i) \neq y_i]].$$

The main result of this subsection is the following theorem, which indicates that the size of the allowed set of left-hand sides may influence generalization.

**Theorem 3 (VC Dimension for Classification)**

The VC dimension $h$ of the set of max-score classifiers is equal to the size of the allowed set of left-hand sides of rules:

$$\text{VCdim}(\mathcal{F}_{\text{maxscore}}) := h := |A|.$$  

From this theorem, classical results such as those of Vapnik (1999, Equations 20 and 21) can be directly applied to obtain a generalization bound:

**Corollary 4 (Uniform Generalization Bound for Classification)**

With probability at least $1 - \delta$ the following holds simultaneously for all $f_g \in \mathcal{F}_{\text{maxscore}}$:

$$\text{TrueErrClass}(f_g) \leq \text{EmpErrClass}(f_g) + \frac{\varepsilon}{2} \left(1 + \sqrt{1 + \frac{4\text{EmpErrClass}(f_g)}{\varepsilon}}\right),$$

where $\varepsilon = 4 \frac{|A|}{m} \left(\frac{\ln \frac{2m}{|A|} + 1}{|A|}\right) - \ln \delta$.

**Note 1 (on uniform bounds):** The result of Theorem 3 holds generally, well beyond the simple adjusted confidence or max confidence, min support algorithms. Those two algorithms correspond to specific choices of the scoring function $g$: the adjusted confidence algorithm takes $g(a, y) = f_{S,K}(a, y)$, and the max confidence, min support algorithm takes $g(a, y) = f_{S,\theta}(a, y)$. We could use other strategies to choose $g$, for example, choosing $f_g \in \mathcal{F}$ to minimize an empirical risk (similar to what we do in Letham et al., 2013c).

**Note 2 (on replacing itemsets with general boolean operators):** Although in this paper we restrict our attention to left-hand sides that are sets of items (e.g., “apples and oranges”), association rules can be constructed using the boolean operators AND, OR, and NOT (e.g., “apples or oranges but not bananas”). In this case, the left-hand sides of rules are not contained in $x$, rather they are true with respect to $x$. By replacing “contained in $x$” with “true with respect to $x$” in the first half of the
Recall that we sample $z \mid s$ sides, drawn from.) Because of this, if we are willing to redefine $A$ with zero probability. (This depends on the unknown probability distribution that the examples are classification and then for sequential event prediction. We will now introduce the notation that will be used for the algorithmic stability bounds, first for $\text{4.3 Notation for Algorithmic Stability Bounds}$

$$\text{TrueErrClass}(f_g) \leq \text{EmpErrClass}(f_g) + \sqrt{\frac{1}{2m} \left( \ln(2|F_{\text{maxscore}}|) + \ln \frac{1}{\delta} \right)}.$$ 

The value of $|F_{\text{maxscore}}|$ is at most $2^{|A|}$. This is because there are $|A|$ ways to determine $\max_{a \in A, a \subseteq x} g(a, y)$, and there are 2 ways to determine the argmax over $y$. The bound then depends on $\sqrt{|A|}$ (as classical VC bounds would also give, using Theorem 3), but not log $|A|$. Note that the bound is meaningful when $|A| < m$ so that $2^{|A|} < 2^m$.

Note 3 (dependence on $|A|$): It is possible that many of the possible left-hand sides in $|A|$ are realized with zero probability. (This depends on the unknown probability distribution that the examples are drawn from.) Because of this, if we are willing to redefine $A$ to include only realizable left-hand sides, $|A|$ can be replaced in the bound by $|A|$, where $A = \{a \in A : P_e(a \subseteq x > 0)\}$ are the itemsets that have some probability of being chosen.

### 4.3 Notation for Algorithmic Stability Bounds

We will now introduce the notation that will be used for the algorithmic stability bounds, first for classification and then for sequential event prediction.

#### 4.3.1 Notation for Classification Bounds

Recall that we sample $z = (x, y)$ where $x \in 2^X$ is a set of items and $y \in \{-1, 1\}$ is the corresponding label. Each $z$ is sampled randomly (iid) according to a distribution $\mathcal{D}$ over the space $2^X \times \{-1, 1\}$. The adjusted confidence algorithm uses the training set $S$ of $m$ iid baskets to compute the adjusted confidences $f_{S,K}$ and find a rule that will be used to label the basket. We use $z = (x, y)$ to refer to a general labeled basket, and $z_i = (x_i, y_i)$ to refer specifically to the $i$th labeled basket in the training set. We define a highest-scoring-correct rule for $x$ as a rule with the highest adjusted confidence that predicts the correct label $y$. The left-hand side of a highest-scoring-correct rule obeys:

$$a_{SxK}^+ \in \arg\max_{a \subseteq x, a \in A} f_{S,K}(a, y) = \arg\max_{a \subseteq x, a \in A} \frac{#(a \cup y)}{#a + K},$$

where $K \geq 0$. If more than one rule is tied for the maximum adjusted confidence, one can now be chosen randomly. If the true label $y$ is not found in the training set, then the confidence of all rules with $y$ on the right-hand side will be 0, and we take $\emptyset \rightarrow y$ as the maximizing rule. We define a highest-scoring incorrect rule for $x$ as a rule with the highest adjusted confidence that predicts the incorrect label $-y$, so the left-hand side obeys:

$$a_{SxK}^- \in \arg\max_{a \subseteq x, a \in A} f_{S,K}(a, -y) = \arg\max_{a \subseteq x, a \in A} \frac{#(a \cup -y)}{#a + K}.$$
Again, if the label $-y$ is not found in the training set, we take $\emptyset \rightarrow -y$ as the maximizing rule. Otherwise, ties are broken randomly.

A misclassification error is made for labeled basket $z$ when the highest-scoring-correct rule, $a_\delta^+ \rightarrow y$, has a lower adjusted confidence than the highest-scoring incorrect rule $a_\delta^- \rightarrow -y$. As discussed earlier, we will measure this difference in adjusted confidence values with respect to a reference $K_r$ in order to allow comparisons with different values of $K$. We will take $K_r \geq 0$. This leads to the definition of the 0-1 loss for classification:

$$
\ell_{0-1,K_r}(f_{S,K}, z) := \begin{cases} 
1 & \text{if } f_{S,K}(a_\delta^+ \rightarrow y) - f_{S,K}(a_\delta^- \rightarrow -y) \leq 0 \\
0 & \text{otherwise.}
\end{cases}
$$

The term $f_{S,K}(a_\delta^+ \rightarrow y) - f_{S,K}(a_\delta^- \rightarrow -y)$ is the “margin” of example $z$ (that is, the gap in score between the predictions for the two classes, see also Shen and Wang, 2007).

We will now define the true error which, when $K = K_r$, is a specific case of TrueErrClass defined in (4). (The function $g$ is chosen using the data set, and it is $f_{S,K}$.) The true error is an expectation of a loss function with respect to $D$, and is a random variable since the training set $S$ is random, $S \sim D^m$.

$$
\text{TrueErrClass}(f_{S,K}, K_r) := \mathbb{E}_{z \sim D} \ell_{0-1,K_r}(f_{S,K}, z).
$$

We approximate the true error using a different loss $\ell_{0-1,K_r}$ that is a continuous upper bound on the 0-1 loss $\ell_{0-1,K_r}$. It is defined with respect to $K_r$ and another real-valued parameter $\gamma > 0$ as follows:

$$
\ell_{\gamma,K_r}(f_{S,K}, z) := c_\gamma(f_{S,K}(a_\delta^+ \rightarrow y) - f_{S,K}(a_\delta^- \rightarrow -y)),
$$

where $c_\gamma : \mathbb{R} \rightarrow [0, 1]$,

$$
c_\gamma(y) = \begin{cases} 
1 & \text{for } y \leq 0 \\
1 - y/\gamma & \text{for } 0 \leq y \leq \gamma \\
0 & \text{for } y \geq \gamma.
\end{cases}
$$

As $\gamma$ approaches 0, loss $c_\gamma$ approaches the standard 0-1 loss. Also, $\ell_{0-1,K_r}(f_{S,K}, z) \leq \ell_{\gamma,K_r}(f_{S,K}, z)$.

We define TrueErrClass$_\gamma$ using this loss:

$$
\text{TrueErrClass}_\gamma(f_{S,K}, K_r) := \mathbb{E}_{z \sim D} \ell_{\gamma,K_r}(f_{S,K}, z),
$$

where TrueErrClass $\leq$ TrueErrClass$_\gamma$. The generalization bounds for classification will bound TrueErrClass by considering the difference between TrueErrClass$_\gamma$ and its empirical counterpart that we will soon define. For training basket $x_i$, the left-hand side of a highest-scoring-correct rule obeys:

$$
a_\delta^+ \subseteq x_i \in \arg\max_{a \subseteq x_i, a \in A} f_{S,K}(a, y_i),
$$

and the left-hand side of a highest-scoring-incorrect rule obeys:

$$
a_\delta^- \subseteq x_i \in \arg\max_{a \subseteq x_i, a \in A} f_{S,K}(a, -y_i).
$$

The empirical error is an average of the loss over the baskets:

$$
\text{EmpErrClass}_\gamma(f_{S,K}, K_r) := \frac{1}{m} \sum_{i=1}^{m} \ell_{\gamma,K_r}(f_{S,K}, z_i).
$$

3457
For the max confidence, min support algorithm, we substitute $\theta$ where $K$ appears in the notation. For instance, for general labeled basket $z = (x, y)$, we analogously define:

$$a_{\text{max}}^+ \in \arg\max_{a \subseteq x, a \in A} f_{S, \theta}(a, y),$$
$$a_{\text{max}}^- \in \arg\max_{a \subseteq x, a \in A} f_{S, \theta}(a, -y),$$

$$\ell_{0-1}(\bar{a}_{\text{max}}^+, z) = \begin{cases} 1 & \text{if } f_{S, K}(a_{\text{max}}^+, y) - f_{S, K}(a_{\text{max}}^-, y) \leq 0 \\ 0 & \text{otherwise} \end{cases},$$

and TrueErrClass$(\bar{f}_{S, \theta}, K_r)$ and TrueErrClass$(\bar{f}_{S, \theta}, K_r)$ are defined analogously as expectations of the losses, and EmpErrClass$(\bar{f}_{S, \theta}, K_r)$ is again an average of the loss over the training baskets.

### 4.3.2 Notation for Sequential Event Prediction Bounds

The notation and the bounds for sequential event prediction are similar to those of classification, the main differences being an additional index $t$ to denote the different time steps, and a set of possible incorrect recommendations in the place of the single incorrect label $-y$. As defined in Section 3, a basket $z$ consists of an ordered (permuted) set of items, $z \in 2^X \times \Pi$, where $2^X$ is the set of all subsets of $X$, and $\Pi$ is the set of permutations over at most $|X|$ elements. We have a training set of $m$ baskets $S = \{z_i\}_{1 \leq i \leq m}$ that are the customer’s past orders. Denote $z \sim D$ to mean that basket $z$ is drawn randomly (iid) according to distribution $D$ over the space of possible items in baskets and permutations over those items, $2^X \times \Pi$. The $t^{th}$ element added to the basket is written $z_{t}$, where the dot is just a placeholder for the generic basket $z$. The $t^{th}$ element of the $i^{th}$ basket in the training set is written $z_{i,t}$. We define the number of items in basket $z$ by $T_z$, that is, $T_z := |z|$.

For sequential event prediction, a highest-scoring-correct rule is a highest scoring rule that has the next item $z_{t+1}$ on the right. The left-hand side $a_{\text{max}}^+$ of a highest-scoring-correct rule obey:

$$a_{\text{max}}^+ \in \arg\max_{a \subseteq \{z_1, \ldots, z_t\}, a \in A} f_{S, K}(a, z_{t+1}).$$

If $z_{t+1}$ has never been purchased, the adjusted confidence for all rules $a \rightarrow z_{t+1}$ is 0, and we choose the maximizing rule to be $\emptyset \rightarrow z_{t+1}$. Also at time 0 when the basket is empty, the maximizing rule is $\emptyset \rightarrow z_{t+1}$.

The algorithm incurs an error when it recommends an incorrect item. A highest-scoring-incorrect rule is a highest scoring rule that does not have $z_{t+1}$ on the right. It is denoted $a_{\text{max}}^- \rightarrow b_{\text{max}}^+$, and obeys:

$$[a_{\text{max}}^-, b_{\text{max}}^+] \in \arg\max_{a \subseteq \{z_1, \ldots, z_t\}, a \in A, \ b \in X} f_{S, K}(a, b).$$

If there is more than one highest-scoring rule, one is chosen at random (with the exception that all incorrect rules are tied at zero adjusted confidence, in which case the left side is taken as $\emptyset$ and the right side is chosen randomly). At time $t = 0$, the left side is again $\emptyset$. The adjusted confidence algorithm determines $a_{\text{max}}^+, a_{\text{max}}^-$, and $b_{\text{max}}^+$, whereas nature chooses $z_{t+1}$.

---

2. Even though we define an order for the basket for this discussion of prediction, we are still using the undirected adjusted confidence to make recommendations rather than the directed version introduced in Section 2. The results can be trivially extended to the directed case.
The empirical error is defined as:

\[ \ell_{0-1,K_r}(f_{S,K},z) := \frac{1}{T_z} \sum_{i=0}^{T_z-1} \begin{cases} 1 & \text{if } f_{S,K}(a_{S,t,K}^+,z_{i+1}) - f_{S,K}(a_{S,t,K},b_{S,t,K}^z) \leq 0 \\ 0 & \text{otherwise}. \end{cases} \]

The true error for sequential event prediction is an expectation of the loss:

\[ \text{TrueErr}(f_{S,K},K_r) := \mathbb{E}_{z \sim \mathcal{D}} \ell_{0-1,K_r}(f_{S,K},z). \]

We create an upper bound for the true error by using a different loss \( \ell_{\gamma,K_r} \) that is a continuous upper bound on the 0-1 loss \( \ell_{0-1,K_r} \). It is defined analogously to classification, with respect to \( K_r \) and \( c_{s \gamma} \):

\[ \ell_{\gamma,K_r}(f_{S,K},z) := \frac{1}{T_z} \sum_{i=0}^{T_z-1} c_{s \gamma}(f_{S,K}(a_{S,t,K}^+,z_{i+1}) - f_{S,K}(a_{S,t,K},b_{S,t,K}^z)). \]

It is true that \( \ell_{0-1,K_r}(f_{S,K},z) \leq \ell_{\gamma,K_r}(f_{S,K},z) \). We define \( \text{TrueErr}_{s \gamma} \):

\[ \text{TrueErr}_{s \gamma}(f_{S,K},K_r) := \mathbb{E}_{z \sim \mathcal{D}} \ell_{\gamma,K_r}(f_{S,K},z), \]

where \( \text{TrueErr} \leq \text{TrueErr}_{\gamma} \). The first set of results for sequential event prediction below bound \( \text{TrueErr} \) by considering the difference between \( \text{TrueErr}_{\gamma} \) and its empirical counterpart that we will soon define.

For the specific training basket \( z_i \), the left-hand side \( a_{S,t,K}^+ \) of a highest-scoring-correct rule at time \( t \) obeys:

\[ a_{S,t,K}^+ \in \arg \max_{a \subseteq \{z_1, \ldots, z_t\}, a \in A} f_{S,K}(a, z_{i+1}), \]

similarly, a highest-scoring-incorrect rule for \( z_i \) at time \( t \) has:

\[ [a_{S,t,K}^+, b_{S,t,K}^z] \in \arg \max_{a \subseteq \{z_1, \ldots, z_t\}, a \in A} f_{S,K}(a, b). \]

The empirical error is defined as:

\[ \text{EmpErr}_{s \gamma}(f_{S,K},K_r) := \frac{1}{m} \sum_{i=1}^{m} \ell_{\gamma,K_r}(f_{S,K},z_i). \]

For the max confidence, min support algorithm, we again substitute \( \theta \) where \( K \) appears in the notation. For example, we define:

\[ a_{S,t,K}^+ \in \arg \max_{a \subseteq \{z_1, \ldots, z_t\}, a \in A} f_{S,\theta}(a, z_{i+1}), \]

\[ [a_{S,t,K}^+, b_{S,t,K}^\theta] \in \arg \max_{a \subseteq \{z_1, \ldots, z_t\}, a \in A} f_{S,\theta}(a, b), \]

\[ \ell_{0-1,K_r}(f_{S,\theta},z) := \frac{1}{T_z} \sum_{i=0}^{T_z-1} \begin{cases} 1 & \text{if } f_{S,K}(a_{S,t,K}^+,z_{i+1}) - f_{S,K}(a_{S,t,K},b_{S,t,K}^\theta) \leq 0 \\ 0 & \text{otherwise}. \end{cases} \]

\[ \ell_{\gamma,K_r}(f_{S,\theta},z) := \frac{1}{T_z} \sum_{i=0}^{T_z-1} c_{s \gamma}(f_{S,K}(a_{S,t,K}^+,z_{i+1}) - f_{S,K}(a_{S,t,K},b_{S,t,K}^\theta)). \]
TrueErr($\tilde{f}_{S, \theta, K_t}$) and TrueErr$_{(t)}$($\tilde{f}_{S, \theta, K_t}$) are expectations of the losses, and EmpErr$_{(t)}$($\tilde{f}_{S, \theta, K_t}$) is an average of the loss over the training baskets.

### 4.4 General Stability Bound for Sequential Event Prediction with Rule-Based Loss

This section contains a stability bound for sequential event prediction, by analogy with Theorem 17 of Bousquet and Elisseeff (2002), using the loss we just defined, which involves rules. We need to define what is meant by a rule-based sequential event prediction algorithm. To keep this definition general, we define an algorithm $\text{Alg}$ to take as input a data set $S$, basket $z$, and item $b^*$ (where $b^*$ is the desired output for basket $z$), and have the algorithm output: (i) the left hand side of the algorithm’s chosen rule to predict $b^*$, which we call $a_{S, z, b^*, \text{Alg}}^+$, (ii) the algorithm’s chosen rule that predicts an item other than $b^*$, which is called $a_{S, z, b^*, \text{Alg}}^-$.

We define $\text{Alg} : S, z, b^* \mapsto a_{S, z, b^*, \text{Alg}}^+; a_{S, z, b^*, \text{Alg}}^-; b_{S, z, b^*, \text{Alg}}$ to have uniform rule stability $\beta$ for sequential event prediction with respect to $K_t$ if:

$$\forall S, \forall z, \forall b^*, \text{ we have } |f_{S, K_t}(a_{S, z, b^*, \text{Alg}}^+, b^*) - f_{S, K_t}(a_{S, z, b^*, \text{Alg}}^-, b^*)| \leq \beta$$

That is, the algorithm is stable whenever (i) the adjusted confidence of the rules used to predict both $b^*$ is not affected much by the removal of one training example, and (ii) when the adjusted confidence of the rule to predict something other than $b^*$ is not affected much by the removal of one training example. We can then show:

**Lemma 5** A rule-based sequential event prediction algorithm with uniform rule stability $\beta$ has uniform stability $2\beta/\gamma$ with respect to the loss function $\ell_{t,K_t}$.

**Proof**

$$|\ell_{t,K_t}(\text{Alg}(S, \cdot, \cdot), z) - \ell_{t,K_t}(\text{Alg}(S^{i/}, \cdot, \cdot), z)|$$

$$= \left| \frac{1}{T_z} \sum_{t=0}^{T_z-1} c_t \left( f_{S, K_t}(a_{S, z, 1 \ldots j, z_{j+1}, \text{Alg}}, z_{j+1}) ight. ight.$$

$$- f_{S, K_t}(a_{S, z, 1 \ldots j, z_{j+1}, \text{Alg}}^-, z_{j+1}) 
- c_t \left( f_{S, K_t}(a_{S, z, 1 \ldots j, z_{j+1}, \text{Alg}}^+, z_{j+1}) ight. 
- f_{S, K_t}(a_{S, z, 1 \ldots j, z_{j+1}, \text{Alg}}^-, z_{j+1}) 
\left. \left. \right| \right|$$

$$\leq \frac{1}{2} \frac{2\beta}{\gamma}.$$

3460
In the first inequality, we used the Lipschitz property of the loss, and properties of absolute values. In the second inequality, we used the definition of uniform rule stability for both absolute value terms with \( b^* \) being \( z_{,j+1} \), and basket \( z \) being \( z_{,1} \ldots z_{,j} \).

Adapting the definitions in the previous subsection to \( Alg \) (rather than \( f_S \)), the following theorem is analogous to Theorem 17 in B&E, for the rule-based loss \( \ell_{\gamma,K} \) for sequential event prediction. The proof is an application of Theorem 12 of B&E to the rule-based sequential event prediction loss, combined with Lemma 5.

**Theorem 6** Let \( Alg \) be a sequential event prediction algorithm with uniform rule stability \( \beta \) for sequential event stability. Then for all \( \gamma > 0 \) and any \( m \geq 1 \) and any \( \delta \in (0, 1) \) with probability at least \( 1 - \delta \) over the random draw of sample \( S \),

\[
\text{TrueErr}(Alg, K_r) \leq \text{EmpErr}_\gamma(Alg, K_r) + \frac{4\beta}{\gamma} + \left( 8m\frac{\beta}{\gamma} + 1 \right) \sqrt{\frac{\ln(1/\delta)}{2m}}
\]

and with probability at least \( 1 - \delta \) over the random draw of sample \( S \),

\[
\text{TrueErr}(Alg, K_r) \leq \text{LooErr}_\gamma(Alg, K_r) + \frac{4\beta}{\gamma} + \left( 8m\frac{\beta}{\gamma} + 1 \right) \sqrt{\frac{\ln(1/\delta)}{2m}}.
\]

We now focus our attention back to the rule-based algorithms from Section 2, and derive a variety of bounds for these algorithms.

**4.5 Generalization Analysis for Large \( m \)**

The choice of minimum support threshold \( \theta \) or the choice of parameter \( K \) matters mainly in the regime where \( m \) is small. For the max confidence, min support algorithm, when \( m \) is large, then all (realizable) itemsets have appeared more times than the minimum support threshold with high probability. For the adjusted confidence algorithm, when \( m \) is large, prediction ability is guaranteed as follows.

**Theorem 7** (Generalization Bound for Adjusted Confidence Algorithm, Large \( m \))

For set of rules \( A, K \geq 0, K_r \geq 0 \), with probability at least \( 1 - \delta \) (with respect to training set \( S \sim \mathcal{D}^m \)),

\[
\text{TrueErr}(f_{S,K}, K_r) \leq \text{EmpErr}_\gamma(f_{S,K}, K_r) + \sqrt{\frac{1}{\delta} \left[ \frac{1}{2m} + 6\beta \right]} + O \left( \frac{1}{m^2} \right),
\]

where \( \beta = \frac{2|A|}{\gamma} \left[ \frac{1}{(m-1)p_{\min A} + K} + \frac{|K_r - K|}{m-1} \right] + O \left( \frac{1}{m^2} \right) \),

and where \( A = \{ a \in A : P_z(a \subseteq z) > 0 \} \) are the itemsets that have some probability of being chosen. Out of these, any itemset that is the least likely to be chosen has probability \( p_{\min A} \):

\[
p_{\min A} := \min_{a \in A} P_{z \sim \mathcal{D}}(a \subseteq z).
\]
As a corollary, the same result holds for classification, replacing $\text{TrueErr}(f_{S,K}, K_r)$ with $\text{TrueErrClass}(f_{S,K}, K_r)$ and $\text{EmpErr}_\gamma(f_{S,K}, K_r)$ with $\text{EmpErrClass}_\gamma(f_{S,K}, K_r)$.

A special case is where $K_r = K = 0$: the algorithm chooses the rule with maximum confidence, and accuracy is then judged by the difference in confidence values between the highest-scoring-incorrect rule and the highest-scoring-correct rule. The bound reduces to:

**Corollary 8 (Generalization Bound for Maximum Confidence Setting, Large $m$)**

With probability at least $1 - \delta$ (with respect to $S \sim D_m$),

$$\text{TrueErr}(f_{S,0}, 0) \leq \text{EmpErr}_\gamma(f_{S,0}, 0) + \sqrt{\frac{1}{\delta} \left[ \frac{1}{2m} + \frac{12|A|}{\gamma(m-1)p_{\min A}} \right]} + O\left(\frac{1}{m^2}\right).$$

Again the result holds for classification with appropriate substitutions. The use of the pointwise hypothesis stability within this proof is the key to providing a decay of order $\sqrt{1/m}$. Now that this bound is established, we move to the small sample case, where the minimum support is the force that provides generalization.

### 4.6 Generalization Analysis for Small $m$

The first small sample result is a general bound for the max confidence, min support algorithm, which holds for both classification and sequential event prediction. The max confidence, min support algorithm has uniform stability, which is a stronger kind of stability than pointwise hypothesis stability. This result strengthens the one in the conference version of this work (Rudin et al., 2011), where we used the bound for pointwise hypothesis stability; uniform stability implies pointwise hypothesis stability, so the result in the conference version follows automatically.

**Theorem 9 (Generalization Bound for Max Confidence, Min Support)**

For $\theta \geq 1$, $K_r \geq 0$, with probability at least $1 - \delta$ (with respect to $S \sim D_m$), $m > \theta$,

$$\text{TrueErr}(\tilde{f}_{S,\theta}, K_r) \leq \text{EmpErr}_\gamma(\tilde{f}_{S,\theta}, K_r) + 2\beta + (4m\beta + 1)\sqrt{\frac{\ln 1/\delta}{2m}}$$

where $\beta = \frac{2}{\gamma} \left[ \frac{1}{\theta} + K_r \left( \frac{1}{\theta + K_r} \right) \left( 1 + \frac{1}{\theta} \right) \right].$

Note that $|A|$ does not appear in the bound. For classification, $\text{TrueErr}(\tilde{f}_{S,\theta}, K_r)$ is replaced by $\text{TrueErrClass}(\tilde{f}_{S,\theta}, K_r)$ and $\text{EmpErr}_\gamma(\tilde{f}_{S,\theta}, K_r)$ is replaced by $\text{EmpErrClass}_\gamma(\tilde{f}_{S,\theta}, K_r)$. Figure 2 shows $\beta$ as a function of $\theta$ for several different values of $K_r$. The special case of interest is when $K_r = 0$, so that the loss is judged with respect to differences in confidence, as follows:

**Corollary 10 (Generalization Bound for Max Confidence, Min Support, $K_r = 0$)**

For $\theta \geq 1$, with probability at least $1 - \delta$ (with respect to $S \sim D_m$), $m > \theta$,

$$\text{TrueErr}(\tilde{f}_{S,\theta}, 0) \leq \text{EmpErr}_\gamma(\tilde{f}_{S,\theta}, 0) + \frac{4}{\gamma\theta} + \left( \frac{8m}{\gamma\theta} + 1 \right)\sqrt{\frac{\ln 1/\delta}{2m}}.$$
Figure 2: $\beta$ vs. $\theta$ from Theorem 9, with $\gamma = 1$. The different curves are different values of $K_r = 0, 1, 5, 10, 50$ from bottom to top.

It is common to use a minimum support threshold that is a fraction of $m$, for instance, $\theta = 0.1 \times m$. In that case, the bound again scales with $\sqrt{(1/m)}$. Note that there is no generalization guarantee when $\theta = 0$; the minimum support threshold enables generalization in the small $m$ case.

Now we discuss the adjusted confidence algorithm for small $m$ setting. We present separate small sample bounds for classification and sequential event prediction.

**Theorem 11** (Generalization Bound for Adjusted Confidence Algorithm, Small $m$, For Classification Only) For $K > 0, K_r \geq 0$, with probability at least $1 - \delta$,

$$ \text{TrueErrClass}(f_{S,K}, K_r) \leq \text{EmpErrClass}_\gamma(f_{S,K}, K_r) + \sqrt{\frac{1}{2m} \left[ \frac{1}{2} \beta + \sqrt{\beta} \right]} $$

where

$$ \beta = \frac{2}{\gamma K} \left( 1 - \frac{(m-1)p_{y,\min}}{m + K} \right) + \frac{2}{\gamma} |K_r - K| \mathbb{E}_{\xi \sim \text{Bin}(m-1, p_{y,\min})} \left[ \frac{1}{K} \left( \frac{m}{m+K} + \frac{1}{K} \left( 1 - \frac{\xi}{m+K} \right) \right) + K_r \left( \frac{1}{m+K} \right) \right] $$

where $p_{y,\min} = \min(P(y = 1), P(y = -1))$ is the probability of the less popular label.

Again, $|A|$ does not appear in the bound, and generalization is provided by $K$, and the difference between $K$ and $K_r$; the interpretation will be further discussed after we state the small sample bound for sequential event prediction.

In the proof of the following theorem, if we were to use the definitions established in Section 4.3.2, the bound does not simplify beyond a certain point and is difficult to read at an intuitive level. From that bound, it would not be easy to see what are the important quantities for the learning process, and how they scale. In what follows, we redefine the loss function slightly, so that it approximates a 0-1 loss from below instead of from above. This provides a concise and intuitive bound.
Define a highest-scoring rule $a^*_{S,tK} \to b^*_{S,tK}$ as a rule that achieves the maximum adjusted confidence, over all of the possible rules. It will either be equal to $a^+_{S,tK} \to z_{t,j+1}$ or $a^*_{S,tK} \to b^*_{S,tK}$, depending on which has the larger adjusted confidence:

$$\left[a^*_{S,tK}, b^*_{S,tK}\right] \subseteq \arg\max_{a \in \{z_{t-1,j-1}, \ldots, z_{t,j} \} \in A^*, b \in \mathcal{X} \in \{z_{t-1,j-1}, \ldots, z_{t,j}\}} f_{S,K}(a, b).$$

Note that $b^*_{S,tK}$ can be equal to $z_{t,j+1}$ whereas $b^*_{S,tK}$ cannot. The notation for $a^*_{S,tK}$ and $b^*_{S,tK}$ is similar, and the new loss is:

$$\ell^\text{new}_{0-1,K} (f_{S,K}, z) := \frac{1}{T_z} \sum_{t=0}^{T_z-1} \begin{cases} 1 & \text{if } f_{S,K}(a^+_{S,tK}, z_{t,j+1}) - f_{S,K}(a^*_{S,tK}, b^*_{S,tK}) < 0 \\ 0 & \text{otherwise.} \end{cases}$$

By definition, the difference $f_{S,K}(a^+_{S,tK}, z_{t,j+1}) - f_{S,K}(a^*_{S,tK}, b^*_{S,tK})$ can never be strictly positive. The continuous approximation is:

$$\ell^\text{new}_{\gamma,K} (f_{S,K}, z) := \frac{1}{T_z} \sum_{t=0}^{T_z-1} c^\text{new}_\gamma(f_{S,K}(a^+_{S,tK}, z_{t,j+1}) - f_{S,K}(a^*_{S,tK}, b^*_{S,tK})), \text{ where}$$

$$c^\text{new}_\gamma(y) = \begin{cases} 1 & \text{for } y \leq -\gamma \\ -y/\gamma & \text{for } -\gamma < y \leq 0 \\ 0 & \text{for } y \geq 0. \end{cases}$$

As $\gamma$ approaches 0, the $c_\gamma$ loss approaches the 0-1 loss. We define $\text{TrueErr}^\text{new}_\gamma (f_{S,K}, K_r)$ and $\text{EmpErr}^\text{new}_\gamma (f_{S,K}, K_r)$ using this loss: $\text{TrueErr}^\text{new}_\gamma (f_{S,K}, K_r) := \mathbb{E}_{z \sim D} \ell^\text{new}_{\gamma,K} (f_{S,K}, z)$, and $\text{EmpErr}^\text{new}_\gamma (f_{S,K}, K_r) := \frac{1}{m} \sum_{i=1}^{m} \ell^\text{new}_{\gamma,K} (f_{S,K}, z_i)$.

The minimum support threshold condition we used in Theorem 9 is replaced by a weaker condition on the support. This weaker condition has the benefit of allowing more rules to be used in order to achieve a better empirical error; however, it is more difficult to get a generalization guarantee. This support condition is derived from the fact that the adjusted confidence of the highest-scoring rule $a^*_{S,tK} \to b^*_{S,tK}$ exceeds that of the highest-scoring-correct rule $a^+_{S,tK} \to z_{t,j+1}$, which exceeds that of the marginal rule $\varnothing \to z_{t,j+1}$:

$$\frac{\#a^*_{S,tK} \backslash K}{a^*_{S,tK} + K} \geq \frac{\#(a^*_{S,tK} \cup b^*_{S,tK})}{a^*_{S,tK} + K} \geq \frac{\#(a^+_{S,tK} \cup z_{t,j+1})}{a^+_{S,tK} + K} \geq \frac{\#z_{t,j+1}}{m + K}. \quad (5)$$

This leads to a lower bound on the support $\#a^*_{S,tK}$:

$$\#a^*_{S,tK} \geq K \left(\frac{\#z_{t,j+1}}{m + K - \#z_{t,j+1}}\right). \quad (6)$$

This is not a hard minimum support threshold, yet since the support generally increases as $K$ increases, the bound will give a better guarantee for large $K$. Note that in the original notation, we would replace the condition (5) with $\frac{\#a^*_{S,tK} \backslash K}{a^*_{S,tK} + K} \geq \frac{\#(a^*_{S,tK} \cup b^*_{S,tK})}{a^*_{S,tK} + K} \geq \frac{\#z_{t,j+1}}{m + K}$ and proceed with analogous steps in the proof.

**Theorem 12** (Generalization Bound for Adjusted Confidence Algorithm, Small $m$) For $K > 0, K_r \geq 0$, with probability at least $1 - \delta$,

$$\text{TrueErr}^\text{new}_\gamma (f_{S,K}, K_r) \leq \text{EmpErr}^\text{new}_\gamma (f_{S,K}, K_r) + \sqrt{\frac{1}{\delta} \left[ \frac{1}{2m} + 6\beta \right]}$$

where
Figure 3: $\beta$ and $\beta_{\text{Approx}}$ vs. $K$, where $K_r = 10$, $p_{\text{min}} = 0.3$, $m = 20$, $\gamma = 1$.

\[
\beta = \frac{2}{\gamma K} \left(1 - \frac{(m-1)p_{\text{min}}}{m+K}\right) + \frac{2}{\gamma} |K_r - K| \mathbb{E}_{\zeta \sim \text{Bin}(m-1,p_{\text{min}})} \left(\frac{1}{K_{m+K-\zeta-1}} + K_r \left(\frac{m}{m+K} + \frac{1}{K} \left(1 - \frac{\zeta}{m+K}\right)\right)\right),
\]

and where $Q = \{x \in X : P_{\zeta,D}(x \in z) > 0\}$ are the items that have some probability of being chosen by the customer. Out of these, any item that is the least likely to be chosen has probability $p_{\text{min}} := \min_{x \in Q} P_{\zeta,D}(x \in z)$.

The stability $\beta$ has two main terms. The first term decreases generally as $1/K$. The second term arises from the error in measuring loss with $K_r$ rather than $K$. In order to interpret $\beta$, consider the following approximation to the expectation in the bound, which assumes that $m$ is large and that $m \gg K \gg 0$, and that $\zeta \approx mp_{\text{min}}$:

\[
\beta \approx \frac{2}{\gamma K} \left(1 - \frac{(m-1)p_{\text{min}}}{m+K}\right) + \frac{2}{\gamma} |K_r - K| \frac{1}{K_{1-p_{\text{min}}}} + K_r.
\]  

Intuitively, if either $K$ is close to $K_r$ or $p_{\text{min}}$ is large (close to 1) then this term becomes small. Figure 3 shows an example plot of $\beta$ and the approximation using (7), which we denote by $\beta_{\text{Approx}}$.

One can observe that if $K_r > K$, then both terms tend to improve (decrease) with increasing $K$. When $K_r < K$, then the two terms can compete as $K$ increases.

4.7 Summary of Bounds

We have provided probabilistic guarantees on performance that show the following: 1) For large $m$, the association rule-based algorithms have a performance guarantee of the same order as other bounds for supervised learning. 2) For small $m$, the minimum support threshold guarantees generalization (at the expense of possibly removing important rules). 3) The adjusted confidence provides a weaker support threshold, allowing important rules to be used, while still being able to generalize. 4) All generalization guarantees depend on the way the goodness of the algorithm is measured (the choice of $K_r$ in the loss function). 5) Important quantities in the learning process may include: $|A|$ or $|\mathcal{A}|$, $K$ or $\theta$, $p_{\text{min},A}$ or $p_{\text{min}}$ (or $p_{\gamma,\text{min}}$).
5. Proofs

In this section, we prove all results from Section 4.

**Proof (Of Theorem 3)** First we show that \( h \leq |A| \). To do this, we must show that for any collection of baskets \( x_1, \ldots, x_N \), \( N > |A| \), there exists a corresponding set of labels \( y_1, \ldots, y_N \) that cannot be realized by any max-score association rule classifier. For each \( x_i \), we introduce a vector \( \bar{x}_i \) of length \( |A| \), where each element corresponds to an \( a \in A \). The element of \( \bar{x}_i \) corresponding to \( a \) is 1 if \( a \subseteq x_i \) and 0 otherwise. Each vector \( \bar{x}_i \) is an element of \( \mathbb{R}^{|A|} \), so the collection of vectors \( \bar{x}_1, \ldots, \bar{x}_N \) must be linearly dependent if \( N > |A| \). By linear dependence and the fact that every \( \bar{x}_i \) is non-zero and non-negative, there must exist coefficients \( c_i \) and disjoint, non-empty sets \( M_0 \) and \( M_1 \) such that:

\[
\sum_{i \in M_0} c_i \bar{x}_i = \sum_{i \in M_1} c_i \bar{x}_i, \quad c_i > 0. \tag{8}
\]

Define \( A_0 = \{ a \in A : a \subseteq x_i \text{ for some } i \in M_0 \} \) and \( A_1 = \{ a \in A : a \subseteq x_i \text{ for some } i \in M_1 \} \). If \( a \subseteq x_i \), for some \( i \in M_0 \), then the corresponding element of \( \bar{x}_i \) will be 1 and the same element in the left part of (8) will be strictly positive. Then, (8) implies that \( a \subseteq x_j \) for some \( j \in M_1 \). Thus, \( A_0 \subseteq A_1 \), and the reverse argument shows \( A_1 \subseteq A_0 \), so \( A_0 = A_1 \). There exists a left-hand side with maximum score, \( a^* = \arg \max_{a \in A_0} \max_{y \in \{-1, 1\}} g(a, y) = \arg \max_{a \in A_1} \max_{y \in \{-1, 1\}} g(a, y) \). The label assigned to \( x_i \), where \( i \) is in \( M_0 \) or \( M_1 \) and \( x_i \) contains itemset \( a^* \), is \( y^* = \arg \max_{y \in \{-1, 1\}} g(a^*, y) \). Thus for at least one \( i \in M_0 \) and at least one \( j \in M_1 \), \( f_{\text{maxscore}}(x_i) = y^* = f_{\text{maxscore}}(x_j) \). Set \( y_i = -1 \) for all \( i \in M_0 \) and \( y_i = 1 \) for all \( i \in M_1 \) and this set of labels cannot be realized, which shows that \( h \leq |A| \).

We now show that this upper bound can be achieved by providing a set of \( |A| \) baskets and finding elements of \( F_{\text{maxscore}} \) that can assign them arbitrary labels. Specifically, we list the elements of \( A \) as \( a_1, \ldots, a_{|A|} \) and take \( x_i = a_i \), for \( i = 1, \ldots, |A| \). Thus each basket is one of the left-hand sides from the allowed set. The elements of \( A \) are not all the same size, and some elements of \( A \) may contain other elements; this could cause problems when we are constructing a max-score classifier that uniquely assigns a given label to each basket. To get around this, we will place the elements of \( A \) in order of increasing size. The possible sizes of elements of \( A \) are denoted \( l_1, \ldots, l_L \), so that \( l_1 < l_2 < \ldots < l_L \). We arrange the elements of \( A \) into sets based on their sizes: \( S_k = \{ i : |a_i| = l_k \}, \quad k = 1, 2, \ldots, L \). We are now ready to construct a classifier \( f_{\text{maxscore}} \) so that, given an arbitrary set of labels \( \{ y_i \} \), it can label the \( x_i \)'s according to the \( y_i \)'s. For all \( i \in S_1 \), we set \( g(a_i, y_i) = c_1 \), any positive number, and \( g(a_i, -y_i) = 0 \). Thus, for the corresponding \( x_i \), \( f_{\text{maxscore}}(x_i) = y_i \). Similarly, for all \( i \in S_2 \), we set \( g(a_i, y_i) = c_2, \quad c_2 > c_1, \) and \( g(a_i, -y_i) = 0 \). For any \( i \in S_2 \), it may be that there exists some \( j \in S_1 \) such that \( a_j \subset x_i \). However, because \( c_2 > c_1 \), the rule with the maximum score will be “\( a_j \rightarrow y_i \)” and \( x_i \) is labeled as desired. In general, for any \( i \in S_k \), we set \( g(a_i, y_i) = c_k \), where \( c_{k-1} < c_k < c_{k+1} \) and \( g(a_i, -y_i) = 0 \) to get \( f_{\text{maxscore}}(x_i) = y_i \). Because this set of \( |A| \) examples can be arbitrarily labeled using elements of \( F_{\text{maxscore}} \), we have \( h \geq |A| \), which combined with the previous result shows that \( h = |A| \).

The remaining theorems are based on the algorithmic stability bounds of Bousquet and Elisseeff (2002) (B&E). Many of the proofs that we provide for classification are essentially identical to those for sequential event prediction. In these cases, the proofs are given for sequential event prediction, and afterwards the translation to classification is outlined. The proofs follow this outline: first, we show how differences in adjusted confidence values with respect to \( K_r \) can be translated into differences with respect to \( K \) (Lemma 15). Then we bound the difference in adjusted confidence values (Lemma 16) in terms of the support. Various lower bounds on the support are used to obtain...
stability for each of the separate cases: large $m$ (Theorem 7), small $m$ for the max confidence, min support algorithm (Theorem 9, which uses uniform stability), small $m$ for classification with the adjusted confidence algorithm (Theorem 11), and small $m$ for sequential event prediction with the adjusted confidence algorithm (Theorem 12).

Following notation of Bousquet and Elisseeff (2002), the input space and output space are $X$ and $Y$. Their training set is $S \in \tilde{Z}^m$, $S = \{\bar{x}_i = (x_i, y_i), \ldots, \bar{x}_m = (x_m, y_m)\}$. An algorithm is a function $A$ from $\tilde{Z}^m$ into $\mathcal{Y} \subset Y^X$ which maps a learning set $S$ onto a function $A_S$ from $X$ to $Y$. The loss is $\ell(f, \bar{z}) = c(f(x), y)$, where $c : Y \times Y \to \mathbb{R}$. $S^{\setminus i}$ means to exclude the $i^{th}$ example $\bar{z}_i$. B&E assume that $Y \subset \mathbb{R}$ but we believe this assumption is unnecessary. In any case, $Y$ is empty for sequential event prediction. An algorithm $A$ has pointwise hypothesis stability $\beta$ with respect to the loss function $\ell$ if the following holds:

$$\forall i \in \{1, \ldots, m\}, \mathbb{E}_{S \sim \text{emp}}[|\ell(A_S, \bar{z}_i) - \ell(A_{S^{\setminus i}}, \bar{z}_i)|] \leq \beta.$$ 

An algorithm $A$ has uniform stability $\beta$ with respect to the loss function $\ell$ if the following holds:

$$\forall S \in Z^m, \forall i \in \{1, \ldots, m\}, ||\ell(A_S, \cdot) - \ell(A_{S^{\setminus i}}, \cdot)||_\infty \leq \beta.$$ 

The empirical error is defined by:

$$R_{\text{emp}}(A, S) := \frac{1}{m} \sum_{i=1}^{m} \ell(A_S, \bar{z}_i)$$

and the true error is:

$$R(A, S) := \mathbb{E}_z[\ell(A_S, \bar{z})].$$

We will use the following results that are based on ideas of Devroye and Wagner (1979).

**Theorem 13** (B&E Pointwise Hypothesis Stability Bound) (Bousquet and Elisseeff, 2002, Theorem 11, first part)

For any learning algorithm $A$ with pointwise hypothesis stability $\beta$ with respect to a loss function $\ell$, such that the value of $\ell$ is at most $M$, we have with probability $1 - \delta$,

$$R(A, S) \leq R_{\text{emp}}(A, S) + \sqrt{\frac{M^2 + 12Mm\beta}{2m\delta}}.$$

**Theorem 14** (B&E Uniform Stability Bound) (Bousquet and Elisseeff, 2002, Theorem 12, first part)

For any learning algorithm $A$ with uniform stability $\beta$ with respect to a loss function $\ell$, such that the value of $\ell$ is at most $M$, we have with probability $1 - \delta$ over a random draw of $S$,

$$R \leq R_{\text{emp}} + 2\beta + (4m\beta + M)\sqrt{\ln \frac{1}{\delta}} \frac{1}{2m}.$$

Translating B&E’s notation to the adjusted confidence setting for sequential event prediction, $\bar{z}_i = x_i = z_i$, with $z_i \in 2^X \times \Pi$. For our problem, $f(x_i)$ is the value of the loss and the $y_i$’s are not defined. In other words, $\ell(A_S, \bar{z}_i) = c(f(x_i), y_i) = f(x_i)$ which in our notation is equal to $\ell_{Y, K_r}(f_S, \bar{z}_i)$. For the max confidence, min support setting, $\ell(A_S, \bar{z}_i)$ translates to $\ell_{Y, K_r}(f_S, \bar{z}_i)$. The adjusted confidence is bounded by 1 so $M = 1$.

The following lemma allows us to convert differences in adjusted confidence with respect to $K_r$ into differences with respect to $K$. 3467
Lemma 15 (Conversion of Adjusted Confidence) For $K \geq 0, K_r \geq 0, 0 \leq s_1 \leq S_1, 0 \leq s_2 \leq S_2$

\[
\frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \leq \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \left( 1 + \frac{|K_r - K|}{S_1 + K} \right) \left( \frac{1}{S_2 + K} \right) \left[ \frac{|K_r - K|}{S_1 + K} + \frac{|K_r - K|}{S_2 + K} \left( \frac{s_2}{S_2 + K} \right) \right]
\]

where $\tilde{S} = \min(S_1, S_2)$.

Proof

\[
\left| \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \right| = \left| \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} + (-K_r + K) \left[ \frac{s_1}{S_1 + K} \left( \frac{1}{S_1 + K} \right) - \frac{s_2}{S_2 + K} \left( \frac{1}{S_2 + K} \right) \right] \right| \\
\leq \left| \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \right| + |K_r - K| \left| \frac{s_1}{S_1 + K} \left( \frac{1}{S_1 + K} \right) - \frac{s_2}{S_2 + K} \left( \frac{1}{S_2 + K} \right) \right|.
\]

Taking just the second absolute value term:

\[
\left| \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \right| = \left| \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \right| = \left| \frac{s_1}{S_1 + K} \left( \frac{1}{S_1 + K} \right) - \frac{s_2}{S_2 + K} \left( \frac{1}{S_2 + K} \right) \right| \leq \left| \frac{s_1}{S_1 + K} - \frac{s_2}{S_2 + K} \right| + 1 \left( \frac{1}{S_1 + K} - \frac{1}{S_2 + K} \right)
\]

Putting this back into (9) yields the statement.

The next results bound the difference in the highest adjusted confidence values when the basket $z_i$ is removed from $S$. We require some additional notation in order to exclude basket $i$. Denote $# / i a$ to be the number of times $a$ has appeared in $S / i$, that is, $# / i a = \sum_{t \neq i} 1_{[a \in z_t]}$. For sequential event prediction, the left-hand side of a highest-scoring-correct rule for a general basket $z$ on $S / i$ obeys:

\[
a^{+}_{S / i, K} \in \arg\max_{a \in \{z_1, \ldots, z_i\}, a \in A} f_{S / i, K}(a, z_{i+1}) = \arg\max_{a \in \{z_1, \ldots, z_i\}, a \in A} \frac{# / i (a \cup z_{i+1})}{# / i a + K}.
\]

A highest-scoring-incorrect rule for basket $z$ on $S / i$ obeys:

\[
[a_{S / i, K}^{+}, b_{S / i, K}^{+}] \in \arg\max_{a \in \{z_1, \ldots, z_i\}, a \in A, b \in X_{\{z_1, \ldots, z_i, a\}}} f_{S / i, K}(a, b) = \arg\max_{a \in \{z_1, \ldots, z_i\}, a \in A, b \in X_{\{z_1, \ldots, z_i, a\}}} \frac{# / i (a \cup b)}{# / i a + K}.
\]

In Lemma 16 below, we bound the difference in adjusted confidence of a general basket $z$ when $z_i$ is removed from the training set, in the sequential event prediction setting.
Lemma 16 (Difference in Adjusted Confidence)
Define $\hat{a}_z := \min(\#a_{S^zK}, \#a_{S^zK}^+)$ and $\hat{a}_z := \min(\#a_{S^zK}, \#a_{S^zK}^+)$. Then,

(I) \[ |f_{S,K}(a_{S^zK}^+, b_{S^zK}) - f_{S,K}(a_{S^zK}^+, b_{S^zK}^-)| \leq \frac{1}{\hat{a}_z + K}, \]

(II) \[ |f_{S,K}(a_{S^zK}, z_{j+1}) - f_{S,K}(a_{S^zK}, z_{j+1})| \leq \frac{1}{\hat{a}_z + K}. \]

Proof Any itemset $a$ is either in $z_i$ or not, thus $\#a^i_a \geq \#a - 1$ and $\#a^i_a \leq \#a$. Also the number of times we see $a \cup b$ is less than or equal to the number of times we see $a$. These observations lead to the following inequalities that will be used throughout the proof:

\[
\begin{align*}
\#a_{S^zK}^+ \cup b_{S^zK}^- & \geq \#a_{S^zK}^+ b_{S^zK}^- - 1, \\
\#a_{S^zK}^+ & \leq \#a_{S^zK}^+, \\
\#(a_{S^zK}^+ \cup b_{S^zK}^-) & \geq \#(a_{S^zK}^+ \cup b_{S^zK}^-), \\
\#a_{S^zK}^+ & \leq \#a_{S^zK}^+ + 1, \\
\#(a_{S^zK}^+ \cup z_{j+1}) & \geq \#(a_{S^zK}^+ \cup z_{j+1}) - 1, \\
\#a_{S^zK}^+ & \leq \#a_{S^zK}^+ + 1, \\
\#(a_{S^zK}^+ \cup z_{j+1}) & \leq \#(a_{S^zK}^+ \cup z_{j+1}) - 1.
\end{align*}
\]

To prove (I) we provide upper bounds for both $f_{S,K}(a_{S^zK}^+, b_{S^zK}) - f_{S,K}(a_{S^zK}^+, b_{S^zK}^-)$ and $f_{S,K}(a_{S^zK}^+, b_{S^zK}^-) - f_{S,K}(a_{S^zK}^+, b_{S^zK}^-)$. Using that for basket $z$ the adjusted confidence of the highest-scoring-incorrect rule on $S^i$, $a_{S^zK}^+ \rightarrow b_{S^zK}^-$, exceeds that of another incorrect rule $a_{S^zK}^\rightarrow b_{S^zK}^-$, and using inequalities (10) and (11),

\[
\frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} \geq \frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} \geq \frac{\#a_{S^zK}^+ \cup b_{S^zK}^- - 1}{\#a_{S^zK}^+ + K}.
\]

Using the inequality above:

\[
\begin{align*}
f_{S,K}(a_{S^zK}^+, b_{S^zK}) - f_{S,K}(a_{S^zK}^+, b_{S^zK}^-) & = \frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} - \frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} \\
& \leq \frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} - \frac{\#a_{S^zK}^+ \cup b_{S^zK}^- - 1}{\#a_{S^zK}^+ + K} = \frac{1}{\#a_{S^zK}^+ + K}.
\end{align*}
\]

Considering the other direction, using that the highest-scoring-incorrect rule under $S$ has higher adjusted confidence than the rule $a_{S^zK}^\rightarrow b_{S^zK}^-$ and inequalities (12) and (13):

\[
\frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} \geq \frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + K} \geq \frac{\#a_{S^zK}^+ \cup b_{S^zK}^-}{\#a_{S^zK}^+ + 1 + K}.
\]
Using this, and inequality (14),
\[
\begin{align*}
\ell_{\gamma,K}(f_{S,K},z) - \ell_{\gamma,K}(f_{S',\gamma,K},z) & = \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \left| c_t \left( f_{S,K}(a_{S,t+1}^+,b_{S,t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right) \\
& - \left( f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right) \right| \\
& \leq \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \left| f_{S,K}(a_{S,t+1}^+,b_{S,t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right| \\
& \leq \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \left| f_{S,K}(a_{S,t+1}^+,b_{S,t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right| + \left| f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right| \\
& =: \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \text{term}_1 + \text{term}_2.
\end{align*}
\]

Together with (20) this proves (I). The proof of part (II) is identical, using \(a_{S,t+1}^+\) and \(a_{S',t+1}^+\) in the place of \(a_{S,t+1}^-\) and \(a_{S',t+1}^-\), in the place of \(b_{S,t+1}^\gamma\) and \(b_{S',t+1}^\gamma\), and inequalities (15)-(19).

The following lemma is the backbone for our stability computations. The upper bound in this lemma depends only on the supports of the relevant rules. Recall that \(\hat{a} \in \min(\#a_{S,t+1}^+,\#a_{S',t+1}^+)\) and \(\hat{a} \in \min(\#a_{S,t+1}^+,\#a_{S',t+1}^+)\).

**Lemma 17 (Large Support Implies Stability)**

\[
\ell_{\gamma,K}(f_{S,K},z) - \ell_{\gamma,K}(f_{S',\gamma,K},z) \leq \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \left| \frac{1}{\hat{a} + K} + |K_r - K| \left( \frac{m}{\hat{a} + K} + \frac{1}{\hat{a} + K} \right) \right|.
\]

**Proof**

\[
\ell_{\gamma,K}(f_{S,K},z) - \ell_{\gamma,K}(f_{S',\gamma,K},z) = \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} c_t \left( f_{S,K}(a_{S,t+1}^+,b_{S,t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right)
\]

\[
\leq \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \left| f_{S,K}(a_{S,t+1}^+,b_{S,t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right| + \left| f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) - f_{S',\gamma,K}(a_{S',t+1}^+,b_{S',t+1}) \right|.
\]

\[
=: \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \text{term}_1 + \text{term}_2.
\]

3470
The first inequality above used that $c_\gamma$ is $1/\gamma$-Lipschitz. Consider an upper bound for term$_1$ as follows from Lemma 15:

$$\text{term}_1 = |f_{S,K}(a_{S\cap K}^-,b_{S\cap K}^-) - f_{S,K}(a_{S\cap K}^+,b_{S\cap K}^+)|$$

$$\leq |f_{S,K}(a_{S\cap K}^-,b_{S\cap K}^-) - f_{S,K}(a_{S\cap K}^+,b_{S\cap K}^+)| \left(1 + \frac{|K_r - K|}{\#a_{S\cap K}^- + K_r}\right)$$

$$+ \frac{|K_r - K|}{\min(\#a_{S\cap K}^-,\#a_{S\cap K}^+) + K_r} \frac{\#a_{S\cap K}^+}{\#a_{S\cap K}^- + K}$$

$$\leq \frac{|K_r - K|}{\min(\#a_{S\cap K}^-,\#a_{S\cap K}^+) + K_r} \frac{\#a_{S\cap K}^+}{\#a_{S\cap K}^- + K}.$$

Now incorporating Lemma 16 and that $\frac{\#(a_{S\cap K}^- \cup b_{S\cap K}^+)}{\#a_{S\cap K}^-} \leq m-1+K \leq \frac{m-1+K}{m+K},$

$$\text{term}_1 \leq \frac{1}{\bar{a}_z + K} \left(1 + \frac{|K_r - K|}{\bar{a}_z + K_r}\right) + \frac{|K_r - K|}{\bar{a}_z + K_r} \frac{m}{m+K}$$

$$= \frac{1}{\bar{a}_z + K} + \frac{|K_r - K|}{\bar{a}_z + K_r} \left[\frac{1}{m+K} \left(\frac{m}{m+K} \right) + \frac{1}{\bar{a}_z + K_r}\right].$$

The same steps can be followed exactly for term$_2$.  

The following lemma is used for the proof for the large sample bound.

**Lemma 18** (Asymptotic Expectation of $1/(\#a + K)$) For any itemset $a \in A$ and any $K \geq 0$,

$$\mathbb{E}_{S \sim D} \frac{1}{\#a + K} \leq \frac{1}{mp_a + K} + O \left(\frac{1}{m^2}\right),$$

where $p_a$ is the probability that a random basket contains $a$, that is, $p_a = P_{S \sim D}(a \subseteq z)$.

Since $\#a$ is binomially distributed, $\#a \sim \text{Binomial}(m, p_a)$, the proof of this lemma can be found by directly applying Lemma 21 in Appendix C.

We now give the proof of pointwise hypothesis stability for the large sample bound. We are interested in the change in adjusted confidence of specific basket $z_i$ when that same basket is removed from the training set, that is on $S'$. Because Lemma 17 holds for any $z$, it also holds for $z_i$, where $a_{z_i} := \min(\#a_{S \cap z_i K}^-,\#a_{S \cap z_i K}^+) \cup \#a_{S \cap z_i K}^-$ and $\bar{a}_{z_i} := \min(\#a_{S \cap z_i K}^+,\#a_{S \cap z_i K}^+) \cup \#a_{S \cap z_i K}^-$.

**Proof (Of Theorem 7)** First, note that:

$$\frac{1}{a_{z_i} + K_r} = \min(\#a_{S \cap z_i K}^- \cup a_{S \cap z_i K}^+) + K_r$$

$$\leq \min(\#a_{S \cap z_i K}^-,\#a_{S \cap z_i K}^+) + K_r \leq \sum_{a \in A} \frac{1}{\#a + K_r}.$$
By the same reasoning, similar upper bounds hold for \(1/(\hat{a}_{z_i} + K), 1/(\hat{a}_{z_i} + \bar{K}),\) and \(1/(\hat{a}_{z_i} + \bar{K}).\) Starting from Lemma 17 using specific basket \(z_i\) and incorporating these bounds on each fraction,  
\[
|\ell_{\mathcal{L}_K}(f_{S,K}, z_i) - \ell_{\mathcal{L}_K}(f_{S^{i-1}, K}, z_i)|
\leq \frac{2}{\gamma T_z} \sum_{t=0}^{T_z-1} \left[ \sum_{a \in \mathcal{A}} \frac{1}{\#a + K} + |K_r - K| \left( \sum_{a \in \mathcal{A}} \frac{1}{\#a + K_r} \right) \left( \frac{m}{m + K} + \sum_{a \in \mathcal{A}} \frac{1}{\#a + K} \right) \right].  
\tag{21}
\]
We have also that for any \(K_r,\) using that \(p_{\max} \leq p_a\) for all \(a \in \mathcal{A},\) and Lemma 18:
\[
\mathbb{E}_{S^{i-1} \sim D^{i-1}} \frac{1}{|\mathcal{A}|} \sum_{a \in \mathcal{A}} \frac{1}{\#a + K_r} \leq \frac{|\mathcal{A}|}{(m-1)p_{\max} + K_r} + O \left( \frac{1}{m^2} \right).  
\tag{22}
\]
Thus from (21) and (22), for any \(1 \leq i \leq m,\)
\[
\mathbb{E}_{S \sim D^m}|\ell_{\mathcal{L}_K}(f_{S,K}, z_i) - \ell_{\mathcal{L}_K}(f_{S^{i-1}, K}, z_i)|
\leq \frac{2}{\gamma T_z} \sum_{t=0}^{T_z-1} \mathbb{E}_{S^{i-1} \sim D^{i-1}} \left[ \sum_{a \in \mathcal{A}} \frac{1}{\#a + K} + |K_r - K| \left( \sum_{a \in \mathcal{A}} \frac{1}{\#a + K_r} \right) \left( \frac{m}{m + K} + \sum_{a \in \mathcal{A}} \frac{1}{\#a + K} \right) \right] + |K_r - K| \left( \sum_{a \in \mathcal{A}} \frac{1}{\#a + K_r} \right) \left( \frac{m}{m + K} \right) + O \left( \frac{1}{m^2} \right) =: \beta,
\]
where in the second inequality, we moved the \((\sum_{a \in \mathcal{A}} 1/(\#a + K_r))(\sum_{a \in \mathcal{A}} 1/(\#a + K))\) terms into the \(O \left( \frac{1}{m^2} \right).\) To see this, one can take a Taylor expansion around the mean for all of the terms similar to \(\frac{1}{\#a + K}\) as follows:
\[
\frac{1}{\#a + K} \approx \frac{1}{mp_a + K} - \frac{(#a - mp_a)}{(mp_a + K)^2} + \frac{(#a - mp_a)^2}{(mp_a + K)^3} + \ldots.
\]
When these terms are multiplied together, the result is always \(O \left( \frac{1}{m^2} \right).\) Thus, the algorithm has pointwise hypothesis stability \(\beta.\) Using \(\beta\) within the B&E theorem yields the result.

**Proof (Of Theorem 9)**

Starting from Lemma 17, we will use the minimum support threshold to provide the upper bound for the reciprocal of the support of rules. All of the steps used to derive Lemma 17 are valid for the max confidence, min support setting, only the notation needs to be changed. We define \(\hat{a}_{z,\theta} := \min(\#a_{S^i,\theta}, \#a'_{S^i,\theta}),\) and now define also \(\hat{a}_{z,\theta} := \min(\#a_{S^i,\theta}, \#a'_{S^i,\theta}).\) Lemma 17 provides for \(\tilde{f}_{S,\theta}\) and using \(K = 0:\)
\[
|\ell_{\mathcal{L}_K}(f_{S,\theta}, z) - \ell_{\mathcal{L}_K}(f_{S^{i-1}, \theta}, z)|
\leq \frac{1}{\gamma T_z} \sum_{t=0}^{T_z-1} \left[ \frac{1}{\hat{a}_{z,\theta} + K_r} \left( \frac{1}{\hat{a}_{z,\theta} + K_r} \left( \frac{1}{\hat{a}_{z,\theta}} \right) \right) + \frac{1}{\hat{a}_{z,\theta} + K_r} \left( \frac{1}{\hat{a}_{z,\theta}} \right) \right].
\]

3472
The requirement of a minimum support threshold ensures that for any particular item \( b \), the highest scoring rule with \( b \) on the right must have support at least \( \theta \), that is:

\[
\max_{a \subseteq \{z_1, \ldots, z_m\}, a \in A} f_{S, \theta}(a, b)
\]

includes only itemsets with support at least \( \theta \). If \( b \) has never been ordered, \( \max_a f_{S, \theta}(a, b) = 0 \) and we choose the maximizing rule to be \( \emptyset \rightarrow b \), with support \( m > m - 1 \geq \theta \). By this reasoning, all of the rules we use have support at least \( \theta \): \( \#a_{S^{i+1} \theta}^+ \geq \theta \), \( \#a_{S^{i+1} \theta}^- \geq \theta \), and \( \#a_{S^{i+1} \theta}^0 \geq \theta \). Thus, \( a_{\theta, \theta} \geq \theta \) and also \( a_{\varepsilon, \theta} \geq \theta \). Using this in the previous expression:

\[
\left| \ell_{T, K}(\hat{f}_{S, \theta}, x) - \ell_{T, K}(\hat{f}_{S^{i+1}, \theta}, x) \right| \\
\leq \frac{2}{\gamma} \frac{1}{T_i} \sum_{i=0}^{T_i-1} \left[ \frac{1}{\theta + K_r} \left( \frac{1}{\theta + K_r} \right) \left( 1 + \frac{1}{\theta} \right) \right] = \frac{2}{\gamma} \frac{1}{\theta + K_r} \left( \frac{1}{\theta + K_r} \right) \left( 1 + \frac{1}{\theta} \right) =: \beta.
\]

This expression holds for all \( S \) and all \( z \). It is thus an upper bound on the uniform stability. Using \( \beta \) within the B&E theorem yields the result. \( \square \)

The proofs of Theorems 7 and 9 for classification are essentially identical to those provided above for sequential event prediction. The left-hand side of a highest-scoring-correct rule for general basket \( x \) on \( S^{i+1} \) obeys:

\[
a_{S^{i+1} \theta}^+ \in \arg\max_{a \subseteq \{a \in A\}} f_{S^{i+1}, K}(a, y) = \arg\max_{a \subseteq \{a \in A\}} \frac{\#(a \cup y)}{\#(a + K^+)}.
\]

And the left-hand side of a highest-scoring-incorrect rule for \( x \) on \( S^{i+1} \) obeys:

\[
a_{S^{i+1} \theta}^- \in \arg\max_{a \subseteq \{a \in A\}} f_{S^{i+1}, K}(a, -y) = \arg\max_{a \subseteq \{a \in A\}} \frac{\#(a \cup -y)}{\#(a + K^+)}.
\]

We further define \( \hat{a}_x = \min(\#a_{S^{i+1} \theta}^+, \#a_{S^{i+1} \theta}^0) \) and \( \hat{a}_y := \min(\#a_{S^{i+1} \theta}^+, \#a_{S^{i+1} \theta}^0) \), and \( \hat{a}_x \) and \( \hat{a}_y \) as the analogous quantities for specific basket \( x_i \). Lemma 16, Lemma 17, and the proof of Theorem 7 all hold for classification by making the following substitutions in notation: \( \hat{a}_x \) and \( \hat{a}_y \) for \( a_x \) and \( a_y \); \( \hat{a}_x \) and \( \hat{a}_y \) for \( \hat{a}_x \) and \( \hat{a}_y \); \( \hat{a}_x \) and \( \hat{a}_y \) for \( \hat{a}_x \) and \( \hat{a}_y \); \( \hat{a}_x \) and \( \hat{a}_y \) for \( \hat{a}_x \) and \( \hat{a}_y \); \( \hat{a}_x \) and \( \hat{a}_y \) for \( \hat{a}_x \) and \( \hat{a}_y \); and \( \hat{a}_x \) and \( \hat{a}_y \) for \( \hat{a}_x \) and \( \hat{a}_y \). For Theorem 9, we again replace \( K \) with \( \theta \) in the notation to define \( \hat{a}_{x, \theta} = \min(\#a_{S^{i+1} \theta}^+, \#a_{S^{i+1} \theta}^+) \) and \( \hat{a}_{x, \theta} := \min(\#a_{S^{i+1} \theta}^+, \#a_{S^{i+1} \theta}^+) \), and then substitute \( \hat{a}_{x, \theta} \) and \( \hat{a}_{x, \theta} \) for \( \hat{a}_{x, \theta} \) and \( \hat{a}_{x, \theta} \) in the proof of the theorem.

The next lemma is specific to classification and is used for the small sample bound for the adjusted confidence algorithm.
Lemma 19 (Support Thresholds for Adjusted Confidence, Classification)
For specific basket \( x_i \), it is true that:

\[
\frac{1}{\hat{\alpha}_{x_i} + K} \leq \alpha_{K}, \text{ where } \alpha_{K} = \frac{m + K - \#^{i}(-y_i)}{K(\#^{i}(-y_i)) + K_r(m + K - \#^{i}(-y_i))};
\]

\[
\frac{1}{\hat{\alpha}_{x_i} + K} \leq \alpha_{K}, \text{ where } \alpha_{K} = \frac{1}{K} \left( 1 - \frac{\#^{i}(-y_i)}{m + K} \right);
\]

\[
\frac{1}{\hat{\alpha}_{x_i} + K} \leq \alpha_{K}, \text{ where } \alpha_{K} = \frac{m + K - \#^{i}y_i}{K(\#^{i}y_i) + K_r(m + K - \#^{i}y_i)}; \text{ and,}
\]

\[
\frac{1}{\hat{\alpha}_{x_i} + K} \leq \alpha_{K}, \text{ where } \alpha_{K} = \frac{1}{K} \left( 1 - \frac{\#^{i}y_i}{m + K} \right).
\]

Proof First we use the fact that on \( S \), the adjusted confidence of the highest-scoring-incorrect rule for \( x_i, a_{Sx_iK} \rightarrow y_i \), exceeds that of the rule \( \varnothing \rightarrow y_i \):

\[
\frac{\#a_{Sx_iK}}{\#a_{Sx_iK} + K} \geq \frac{\#(a_{Sx_iK} \cup -y_i)}{\#a_{Sx_iK} + K} \geq \frac{\#(-y_i)}{m + K} = \frac{\#^{i}(-y_i)}{m + K},
\]

where in the last step we used that basket \( x_i \) does not have label \(-y_i\). Rearranging,

\[
\frac{\#a_{Sx_iK}}{\#a_{Sx_iK} + K} \geq \hat{\sigma} \text{ where } \hat{\sigma} := K \left( \frac{\#^{i}(-y_i)}{m + K - \#^{i}(-y_i)} \right).
\]

Similarly, the adjusted confidence of the highest-scoring-incorrect rule for \( x_i \) with data set \( S^{i} \), \( a_{S^{i}x_iK} \rightarrow -y_i \), exceeds that of the rule \( \varnothing \rightarrow -y_i \), thus:

\[
\frac{\#^{i}a_{S^{i}x_iK}}{\#^{i}a_{S^{i}x_iK} + K} \geq \frac{\#^{i}(a_{S^{i}x_iK} \cup -y_i)}{\#^{i}a_{S^{i}x_iK} + K} \geq \frac{\#^{i}(-y_i)}{m - 1 + K} \geq \frac{\#^{i}(-y_i)}{m + K}.
\]

Rearranging, we find that \( \#^{i}a_{S^{i}x_iK} \geq \hat{\sigma} \). Thus, \( \tilde{\alpha}_{x_i} = \min(\#a_{Sx_iK}, \#^{i}a_{S^{i}x_iK}) \geq \hat{\sigma} \). We can derive a similar bound for \( \tilde{\alpha}_{x_i} \), beginning with \( a_{Sx_iK}^{+} \):

\[
\frac{\#a_{Sx_iK}^{+}}{\#a_{Sx_iK}^{+} + K} \geq \frac{\#(a_{Sx_iK}^{+} \cup y_i)}{\#a_{Sx_iK}^{+} + K} \geq \frac{\#y_i}{m + K} = \frac{\#^{i}y_i + 1}{m + K} > \frac{\#^{i}y_i}{m + K}.
\]

The first equality uses that basket \( x_i \) has label \( y_i \). Rearranging,

\[
\frac{\#a_{Sx_iK}^{+}}{\#a_{Sx_iK}^{+} + K} > \hat{\sigma} \text{ where } \hat{\sigma} := K \left( \frac{\#^{i}y_i}{m + K - \#^{i}y_i} \right).
\]

Similarly for \( \#^{i}a_{S^{i}x_iK}^{+} \):

\[
\frac{\#^{i}a_{S^{i}x_iK}^{+}}{\#^{i}a_{S^{i}x_iK}^{+} + K} \geq \frac{\#^{i}(a_{S^{i}x_iK}^{+} \cup y_i)}{\#^{i}a_{S^{i}x_iK}^{+} + K} \geq \frac{\#^{i}y_i}{m - 1 + K} \geq \frac{\#^{i}y_i}{m + K}.
\]
Rearranging, we find \( \#^1a_{S_x} \geq \hat{\sigma} \). Thus \( \hat{\sigma}_i = \min(\#^1a_{S_x}, \#^1a_{S_x}) \geq \hat{\sigma} \). These lower bounds on the supports are now used to create upper bounds for the reciprocals:

\[
\frac{1}{\bar{\sigma}_i + K} \leq \frac{1}{\bar{\sigma} + K} = \hat{\sigma}_i \text{ and } \frac{1}{\bar{\sigma}_i + K} \leq \frac{1}{\bar{\sigma} + K} = \hat{\sigma}_i.
\]

The bounds for \( \frac{1}{\bar{\sigma}_i + K} \) and \( \frac{1}{\bar{\sigma}_i + K} \) are obtained in a similar way using \( \hat{\sigma} \).

The proof of the small sample bound for classification follows directly from this lemma.

**Proof (Of Theorem 11)**

From Lemma 17, adapted for classification,

\[
|e_{\gamma,K_r}(f_{S,K}, z_i) - e_{\gamma,K_r}(f_{S^i,K}, z_i)| \\
\leq \frac{1}{\gamma} \left[ \frac{1}{\bar{\sigma}_i + K} + |K_r - K| \left[ \frac{1}{\bar{\sigma}_i + K_r} \left( \frac{m}{m + K} + \frac{1}{\bar{\sigma}_i + K} \right) \right] + \frac{1}{\bar{\sigma}_i + K} + |K_r - K| \left[ \frac{1}{\bar{\sigma}_i + K} \left( \frac{m}{m + K} + \frac{1}{\bar{\sigma}_i + K} \right) \right] \right].
\]

Combining this and Lemma 19, we have:

\[
|e_{\gamma,K_r}(f_{S,K}, z_i) - e_{\gamma,K_r}(f_{S^i,K}, z_i)| \\
\leq \frac{1}{\gamma} \left[ \hat{\sigma}_i + |K_r - K| \hat{\sigma}_i \left( \frac{m}{m + K} + \hat{\sigma}_i \right) + \hat{\sigma}_i + |K_r - K| \hat{\sigma}_i \left( \frac{m}{m + K} + \hat{\sigma}_i \right) \right] \\
= \frac{1}{\gamma} \left( \hat{\sigma}_i + \hat{\sigma}_i \right) + \frac{1}{\gamma} |K_r - K| \left[ \hat{\sigma}_i \left( \frac{m}{m + K} + \hat{\sigma}_i \right) + \hat{\sigma}_i \left( \frac{m}{m + K} + \hat{\sigma}_i \right) \right].
\]

We now provide an upper bound on the expectation of this quantity, beginning with the first term:

\[
\mathbb{E}_{z_1, \ldots, z_m} \frac{1}{\gamma} (\hat{\sigma}_i + \hat{\sigma}_i) = \mathbb{E}_{z_1, \ldots, z_m} \frac{1}{\gamma} \left[ \left( 1 - \frac{\#^1(-y_i)}{m + K} \right) + \left( 1 - \frac{\#^1y_i}{m + K} \right) \right] \\
= \frac{1}{\gamma} \left[ 2 - \frac{(m - 1)p_{-y_i}}{m + K} - \frac{(m - 1)p_{y_i}}{m + K} \right] \\
\leq \frac{2}{\gamma} \left[ 1 - \frac{(m - 1)p_{y_{\min}}}{m + K} \right].
\]

Here we used the fact that the mean of the binomial distribution \( \text{Bin}(m - 1, p_{y_i}) \) is \( (m - 1)p_{y_i} \), and we use a lower bound for \( p_{y_i} \) and \( p_{-y_i} \), namely \( p_{y_{\min}} = \min(P(y = 1), P(y = -1)) \) the minimum...
probability of a randomly chosen basket having any particular label. For the second term,
\[
\mathbb{E}_{z_1, \ldots, z_m} \mathbb{I}_{K_r - K} \left[ \hat{\alpha}_{K_r} \left( \frac{m}{m + K} + \hat{\alpha}_K \right) + \hat{\alpha}_{K_r} \left( \frac{m}{m + K} + \hat{\alpha}_K \right) \right]
\]
\[
= \frac{1}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(-y_i)}{m + K - \#^i(-y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(-y_i)}{m + K} \right) \right) \right]
\]
\[
+ \frac{1}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(y_i)}{m + K - \#^i(y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(y_i)}{m + K} \right) \right) \right]
\]
\[
= \frac{1}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(y_i)}{m + K - \#^i(y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(y_i)}{m + K} \right) \right) \right]
\]
\[
+ \frac{1}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(-y_i)}{m + K - \#^i(-y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(-y_i)}{m + K} \right) \right) \right]
\]
\[
= \frac{1}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(y_i)}{m + K - \#^i(y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(y_i)}{m + K} \right) \right) \right]
\]
Since the function \( F(\zeta) \) is decreasing as \( \zeta \) increases, then an upper bound is produced by using the distribution \( \text{Bin}(m - 1, p_{y_{\min}}) \):
\[
\mathbb{E}_{z_1, \ldots, z_m} \frac{1}{\gamma} |K_r - K| \left[ \hat{\alpha}_{K_r} \left( \frac{m}{m + K} + \hat{\alpha}_K \right) + \hat{\alpha}_{K_r} \left( \frac{m}{m + K} + \hat{\alpha}_K \right) \right]
\]
\[
\leq \frac{2}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(y_i)}{m + K - \#^i(y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(y_i)}{m + K} \right) \right) \right]
\]
\[
= \frac{2}{\gamma} |K_r - K| \mathbb{E}_{z_1, \ldots, z_m} \left[ \frac{1}{K} \left( \frac{\#^i(y_i)}{m + K - \#^i(y_i)} + K_r \right) \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\#^i(y_i)}{m + K} \right) \right) \right] .
\]

The following lemma is similar to the previous lemma, but specific to sequential event prediction. It uses the support guarantee for the adjusted confidence algorithm (6) in order to bound the terms of Lemma 17, which holds with the same proof when the loss \( \ell_{\gamma,K} \) is changed to the new loss \( \ell_{\gamma,K}^{new} \) and superscript “*” is replaced by “**”. We define the analogy to \( a_{z_{\min}} \) as \( a_{z_{\min}}^* := \min(a_{S_{t+1}K}, a_{S_{t+1}K}^*) \). The result below will immediately yield a proof of Theorem 12.

**Lemma 20 (Support Thresholds for Adjusted Confidence, Sequential Event Prediction)**

For specific basket \( z_i \), define:
\[
\alpha_{K_r} := \frac{m + K - \#z_{i,t+1}}{K(\#z_{i,t+1} - 1) + K_r(m + K - \#z_{i,t+1})} \quad \text{and} \quad \alpha_K := \frac{1}{K} \left( 1 - \frac{\#z_{i,t+1} - 1}{m + K} \right).
\]

It is true that:
\[
\frac{1}{a_{z_i}^* + K_r} \leq \alpha_{K_r}, \quad \frac{1}{a_{z_i}^* + K} \leq \alpha_K, \quad \frac{1}{a_{z_i} + K_r} \leq \alpha_{K_r}, \quad \text{and} \quad \frac{1}{a_{z_i} + K} \leq \alpha_K.
\]
\textbf{Proof} Starting with (5), we know that \( a^*_{t_{ij}K} > \sigma \), where
\[
\sigma := K \left( \frac{\#z_{i, t+1} - 1}{m + K - \#z_{i, t+1}} \right).
\]

We use the same type of argument as in (5), incorporating the fact that on \( S^i \), the adjusted confidence of the highest scoring rule \( a^*_{S^i z_{ij}K} \rightarrow b^*_{S^i z_{ij}K} \) exceeds that of the highest-scoring-correct rule \( a^+_{S^i z_{ij}K} \rightarrow z_{i, t+1} \), which exceeds that of the rule \( \varnothing \rightarrow z_{i, t+1} \),
\[
\frac{\#a^+_{S^i z_{ij}K}}{\#a^*_{S^i z_{ij}K}} + K \geq \frac{\#(a^*_{S^i z_{ij}K} \cup b^*_{S^i z_{ij}K})}{\#a^*_{S^i z_{ij}K} + K} \geq \frac{\#(a^+_{S^i z_{ij}K} \cup z_{i, t+1})}{\#a^+_{S^i z_{ij}K} + K} \geq \frac{\#z_{i, t+1} - 1}{m + K}.
\]

Rearranging, we find that \( \#a^+_{S^i z_{ij}K} > \sigma \). Similarly for \( #a^+_{S^i z_{ij}K} \),
\[
\frac{\#a^+_{S^i z_{ij}K}}{\#a^*_S z_{ij}K} + K \geq \frac{\#(a^*_S z_{ij}K \cup z_{i, t+1})}{\#a^*_S z_{ij}K + K} \geq \frac{\#z_{i, t+1}}{m + K}.
\]

so \( #a^+_{S^i z_{ij}K} \geq K \left( \frac{\#z_{i, t+1}}{m + K - \#z_{i, t+1}} \right) > \sigma \). And again for \( #a^+_{S^i z_{ij}K} \) using (23),
\[
\frac{\#a^+_{S^i z_{ij}K}}{\#a^*_{S^i z_{ij}K}} + K \geq \frac{\#(a^*_{S^i z_{ij}K} \cup z_{i, t+1})}{\#a^*_{S^i z_{ij}K} + K} \geq \frac{\#z_{i, t+1} - 1}{m + K}.
\]

so \( #a^+_{S^i z_{ij}K} \geq \sigma \). We now have \( \hat{a}^*_{z_{ij}} = \min(\#a^*_{S^i z_{ij}K}, #a^+_{S^i z_{ij}K}) \geq \sigma \), and also \( \hat{a}_{z_{ij}} = \min(\#a^+_{S^i z_{ij}K}, #a^+_{S^i z_{ij}K}) \geq \sigma \). Since \( \sigma \) is a lower bound on all the supports, it can be used to create an upper bound for the reciprocals, as follows, using \( \hat{a}^*_{z_{ij}} \) as an example:
\[
\frac{1}{\hat{a}^*_{z_{ij}} + K} \leq \frac{1}{\sigma + K} = \alpha_{K^r} \text{ and } \frac{1}{\hat{a}_{z_{ij}} + K} \leq \frac{1}{\sigma + K} = \alpha_{K^r}.
\]

\textbf{Proof (Of Theorem 12)} First, all of the steps in the proof of Lemma 17 hold when we replace the loss \( \ell_{Y, K} \) with the new loss \( \ell^\text{new}_{Y, K} \). Replace \( c_Y \) with \( c^\text{new}_Y \), and \( a_{z_{ij}} \) by \( \hat{a}^*_{z_{ij}} \), so we obtain:
\[
|\ell^\text{new}_{Y, K}(f_{S, K, z_{ij}}) - \ell^\text{new}_{Y, K}(f_{S^i, K, z_{ij}})|
\leq \frac{1}{\gamma T_{z_{ij}}} \sum_{t=0}^{T_{z_{ij}} - 1} \left[ \frac{1}{\hat{a}^*_{z_{ij}} + K} + |K_r - K| \left[ \frac{1}{\hat{a}^*_{z_{ij}} + K} \left( \frac{m}{m + K + \frac{1}{\hat{a}^*_{z_{ij}} + K}} \right) \right] \right].
\]

Combining this and Lemma 20, we have:
\[
|\ell^\text{new}_{Y, K}(f_{S, K, z_{ij}}) - \ell^\text{new}_{Y, K}(f_{S^i, K, z_{ij}})| \leq \frac{2}{\gamma T_{z_{ij}}} \sum_{t=0}^{T_{z_{ij}} - 1} \alpha_K + |K_r - K| \alpha_K \left( \frac{m}{m + K + \alpha_K} \right).
\]
To calculate the stability, we need an upper bound on the expectation of this quantity. Let us first create an upper bound for the expectation of the first term, $\frac{2}{\gamma} \frac{1}{T_z} \sum_{t=0}^{T_z-1} \alpha_K$:

$$
\mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \frac{1}{\gamma} \frac{1}{T_z} \sum_{t=0}^{T_z-1} \alpha_K \right] = \mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \frac{1}{\gamma} \frac{1}{T_z} \sum_{t=0}^{T_z-1} \frac{1}{K} \left( 1 - \frac{\# z_{i,t+1} - 1}{m + K} \right) \right] \\
= \mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \frac{1}{\gamma} \frac{1}{T_z} \sum_{t=0}^{T_z-1} \frac{1}{K} \left( 1 - \frac{\# z_{i,t+1} - 1}{m + K} \right) \right] \\
= \mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \frac{1}{\gamma} \frac{1}{T_z} \sum_{t=0}^{T_z-1} \frac{1}{K} \left( 1 - \frac{(m-1)p_{z_{i,t+1}}}{m + K} \right) \right] \\
\leq \mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \frac{1}{\gamma} \frac{1}{T_z} \sum_{t=0}^{T_z-1} \frac{1}{K} \left( 1 - \frac{(m-1)p_{\min}}{m + K} \right) \right] = \frac{2}{\gamma} \frac{1}{K} \left( 1 - \frac{(m-1)p_{\min}}{m + K} \right).
$$

The first line above uses the definition of $\alpha_K$, the second line uses the fact that each basket is chosen independently, the third line uses that $z_{i,t+1}$ is always contained in $z_i$ and also uses the fact that the mean of the binomial distribution $\text{Bin}(m-1, p_{z_{i,t+1}})$ is $(m-1)p_{z_{i,t+1}}$. The fourth line uses that $p_{z_{i,t+1}}$ has the lower bound $p_{\min}$, which no longer depends on $z_i$.

We repeat this outline for the second term:

$$
\mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \left| K_r - K \right| \frac{1}{T_z} \sum_{t=0}^{T_z-1} \alpha_K \left( \frac{m}{m + K} + \alpha_K \right) \right] \\
= \mathbb{E}_{z_1, \ldots, z_m} \left[ 2 \left| K_r - K \right| \frac{1}{T_z} \sum_{t=0}^{T_z-1} \frac{1}{K} \left( 1 - \frac{\# z_{i,t+1} - 1}{m + K - \# z_{i,t+1}} \right) + K_r \right] \left( \frac{m}{m + K} + 1 \right) \left( 1 - \frac{\# z_{i,t+1} - 1}{m + K} \right) \\
= \frac{2}{\gamma} \left| K_r - K \right| \mathbb{E}_{z_1, \ldots, z_m} \left[ \sum_{t=0}^{T_z-1} \mathbb{E}_{z_1, \ldots, z_{i-1}, z_{i+1}, \ldots, z_m} \left[ \frac{1}{K} \left( 1 - \frac{\# z_{i,t+1} - 1}{m + K - \# z_{i,t+1}} \right) + K_r \right] \right] \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\# z_{i,t+1} - 1}{m + K} \right) \right) \\
= \frac{2}{\gamma} \left| K_r - K \right| \mathbb{E}_{z_1, \ldots, z_m} \left[ \sum_{t=0}^{T_z-1} \mathbb{E}_{\zeta \sim \text{Bin}(m-1, p_{z_{i,t+1}})} \frac{1}{K} \left( 1 - \frac{\zeta + 1}{m + K - \zeta - 1} \right) + K_r \right] \left( \frac{m}{m + K} + \frac{1}{K} \left( 1 - \frac{\zeta + 1}{m + K} \right) \right) \\
= \frac{2}{\gamma} \left| K_r - K \right| \mathbb{E}_{z_1, \ldots, z_m} \left[ \sum_{t=0}^{T_z-1} \mathbb{E}_{\zeta \sim \text{Bin}(m-1, p_{z_{i,t+1}})} F(\zeta) \right].
$$
Algorithm 4: Subroutine GenRules, simplest version that considers only “marginal” rules.

**Input:** \((S, B, X)\), that is, past orders \(S = \{z_i\}_{i=1}^m, z_i \subseteq X\), current basket \(B \subseteq X\), set of items \(X\)

**Output:** Set of all rules where \(a_j\) is an item in the basket \(B\) (or the empty set) and \(b_j\) is not in \(B\). That is, rules \(\{a_j \rightarrow b_j\}_j\) such that \(b_j \in X \setminus B\) and either \(a_j \in B\) or \(a_j = \emptyset\).

Since the function \(F\) is decreasing as \(\zeta\) increases, then an upper bound is produced by using the distribution \(\text{Bin}(m-1, p_{\text{min}})\). Namely,

\[
\mathbb{E}_{z_1, \ldots, z_m} \left[ |K_r - K| \frac{1}{T_{z_i}} \sum_{t=0}^{T_{z_i} - 1} \alpha_{K_r} \left( \frac{m}{m + K} + \alpha_K \right) \right] \\
\leq \mathbb{E}_{z_1, \ldots, z_m} \left[ |K_r - K| \frac{1}{T_{z_i}} \sum_{t=0}^{T_{z_i} - 1} \mathbb{E}_{\zeta \sim \text{Bin}(m-1, p_{\text{min}})} F(\zeta) \right] \\
= \mathbb{E}_{z_1, \ldots, z_m} \left[ |K_r - K| \frac{1}{m + K} \left( \frac{m}{m + K - \zeta - 1} \right) + K_r \left( \frac{m}{m + K} + \frac{1}{K_r} \left( 1 - \frac{\zeta}{m + K} \right) \right) \right].
\]

In all of the theorems and proofs, the empirical loss and true loss are defined only for the case where the algorithm only recommends one item \((c = 1)\). It is possible to use a vector norm to generalize to larger \(c\).

6. Experiments

All data sets chosen for these experiments are publicly available from the UCI machine learning repository (Bache and Lichman, 2013), and from the IBM Quest Market-Basket Synthetic Data Generator (Agrawal and Srikant, 1994). To obtain formatted market-basket data, categorical data were converted into binary features (one feature per category). Each feature represents an item, and each example represents a basket. The feature value \((0\) or \(1)\) indicates the presence of an item. Training baskets and test baskets were chosen randomly without replacement from the full data set. Since these data do not come naturally with a time ordering, items in the basket were randomly permuted to attain an order. At each iteration, rules were formed from one item or the empty item on the left, and one item on the right (See GenRules in Figure 4). Recommendations of one item were made using the following 15 algorithms: highest support, highest confidence, highest adjusted confidence for eight \(K\) levels, max confidence, min support algorithm for five support threshold levels \(\theta\). All 15 algorithms were evaluated by the average fraction of correct recommendations \(\text{AvgCorrect}\) per basket. As recommendations were made, it was common to have ties where multiple items were equally good to recommend, in which case the tie was broken at random; AvgCorrect is similar to \(\ell_{0.1_K}\) except for this way of dealing with ties.

The parameters of the experiment are: number of training baskets \((20\) in all cases\), number of test baskets \((100\) in all cases\), values of \(K\) for the adjusted confidence algorithm \((0.0001, 0.001, 0.01, 0.1, 1, 5, 10, 15)\), and values of \(\theta\) for the max confidence, min support algorithm \((1, 2, 3, 5, 15)\).
10). Note that two of these algorithms are the same: the max confidence algorithm is the same as the max confidence, min support algorithm for $\theta=1$. Data sets are: Car Evaluation (25 items, 1728 baskets), Chess King-Rook vs. King-Pawn, (75 items, 3196 baskets), MONK’s problems (19 items, 1711 baskets) Mushroom (119 items, 8124 baskets), Nursery (32 items, 12960 baskets), Plants (70 items, 34781 baskets), T20I18D10KN22CR50 (22 items, 10000 baskets).

Each experiment (training, test, evaluation for all 15 algorithms) was performed 100 times, (totaling $100 \times 100 \times 15 = 150,000$ test basket evaluations per data set, for each of 7 data sets). In Figures 4 and 5, the distribution of AvgCorrect values for data sets Chess and Monk are shown via boxplot, along with the mean and standard deviation of AvgCorrect values. Bold indicates that the mean is not significantly different from that of the algorithm with the largest mean value; that is, bold indicates the highest scores. The boxplots and means for the other data sets are shown in Figures 7 through 11 in Appendix B.

Figure 6 summarizes the results of all of the experiments by totaling the number of data sets for which each algorithm achieved one of the highest scores. The best performing algorithms were $K = 0.01$ and $K = 0.1$, both algorithms achieving one of the top scores for 6 out of 7 of the data sets. The single data set for which these algorithms did not achieve one the best scores was the very dense data set T20I18D10KN22CR50, where the algorithms requiring a higher support (the max support algorithm, and also the adjusted confidence algorithm for $K = 5$, 10, and 15) achieved the highest AvgCorrect score. In that case, the $K = 0.01$ and $K = 0.1$ algorithms still performed better than the max confidence, min support algorithms for the parameters we tried.

The adjusted confidence algorithm with a very small $K$ is similar to using the max confidence algorithm, except that whenever there is a tie, the tie is broken in favor of the rule with largest support. It seems that in most of the data sets we chose, this type of algorithm performed the best, which indicates two things. First, that for some data sets, increasing $K$ too much can have the same effect as a too-large minimum support threshold, in that large values of $K$ could potentially remove the best rules, leading to too much bias, and where the algorithm cannot explain enough of the variance in the data. Second, when comparing rules, it is important not to break ties at random as in the max confidence, min support algorithm, but instead to use the support of the rules. Another observation is that the performance levels of the adjusted confidence algorithm vary less than those of the max confidence, min support algorithm. In other words, our experiments indicate that a less-than-perfect choice of $K$ for the adjusted confidence algorithm is likely to perform better than a less-than-perfect choice of $\theta$ for the max confidence, min support algorithm.

7. Related Work

We provide background on related works within several fields: association rule mining and associative classification, decision lists, recommender systems, and Bayesian analysis. There is also a body of literature on pattern mining in sequences, but not in the sequential event prediction setting defined here. This type of work generally considers the order in which items are added, and often uses a Markov assumption (see, for instance, Ayres et al., 2002; Berchtold and Raftery, 2002), whereas in our work, subsets of items are used to predict the next item, possibly without regard to the order in which they occurred, and a Markov assumption can be false. There is also work relating statistics to pattern mining and sequence mining, (e.g., Chernoff bounds for the confidence, Jacquemont et al., 2009). Our work also relates to multi-class classification, since there is a multi-class classification step at each point in time $t$ of each sequence. For a recent work on generalization bounds in
Algorithm | mean ± standard dev.  
--- | ---  
Support | 0.0813 ± 0.0046  
Confidence | 0.0764 ± 0.0053  
K=0.0001 | 0.0831 ± 0.0045  
K=0.001 | 0.0832 ± 0.0048  
K=0.01 | 0.0835 ± 0.0041  
K=0.1 | 0.0834 ± 0.0049  
K=1 | 0.0835 ± 0.0043  
K=5 | 0.0821 ± 0.0049  
K=10 | 0.0821 ± 0.004  
K=15 | 0.0816 ± 0.0049  
θ=1 | 0.0759 ± 0.0049  
θ=2 | 0.0767 ± 0.0045  
θ=3 | 0.078 ± 0.0049  
θ=5 | 0.0794 ± 0.0052  
θ=10 | 0.0813 ± 0.0046  

Figure 4: *Left:* Boxplots of AvgCorrect values for Chess data set. *Right:* Means and standard deviations for Chess data set.

Algorithm | mean ± standard dev.  
--- | ---  
Support | 0.0943 ± 0.0126  
Confidence | 0.1103 ± 0.0145  
K=0.0001 | 0.1108 ± 0.0137  
K=0.001 | 0.1109 ± 0.0147  
K=0.01 | 0.1104 ± 0.0149  
K=0.1 | 0.11 ± 0.0151  
K=1 | 0.1081 ± 0.0148  
K=5 | 0.0992 ± 0.0138  
K=10 | 0.0947 ± 0.0133  
K=15 | 0.0948 ± 0.012  
θ=1 | 0.1098 ± 0.0138  
θ=2 | 0.1095 ± 0.0146  
θ=3 | 0.1092 ± 0.0146  
θ=5 | 0.1054 ± 0.0143  
θ=10 | 0.0944 ± 0.0129  

Figure 5: *Left:* Boxplots of AvgCorrect values for MONK’s problems data set. *Right:* Means and standard deviations for MONK’s problems data set.

multi-class classification see Shen and Wang (2007). Remember that in multi-class classification, each example is a feature vector, whereas in sequential event prediction, each example is an event sequence. Related work on generalization bounds includes those on algorithmic stability (Devroye and Wagner, 1979; Bousquet and Elisseeff, 2002).

### 7.1 Mining Association Rules

Association rule mining has proven successful for many applications, including market basket analysis (cross selling, product placement, affinity promotion, see also Kohavi et al., 2004), mining gene expression data (Jiang and Gruenwald, 2005), and weblog analysis (Huang et al., 2002).
majority of literature on association rule mining concerns the design of efficient algorithms to address the time-and-memory-consuming task of mining rules within very large databases. Discovering rules is usually a two-step process. First, itemsets are mined that meet a predetermined minimum support threshold. Then using this set, rules are formed and the strength of the rules is assessed using “interestingness” measures, such as the confidence. Many “interestingness” measures have been proposed in the literature (see Tan et al., 2002; Geng and Hamilton, 2007; McGarry, 2005). It is clearly possible to use the adjusted confidence as an interestingness measure for database exploration. In that setting, the adjusted confidence would provide a ranking of rules in terms of their ability to predict, including both “common sense rules” and “nuggets.”

Although association rule mining has proven successful for many applications, it is well-known that the usefulness of association rules and their impact on even a wider range of practical applications remains limited due to problems arising from the minimum support threshold: first, the large number of rules mined can be intractable to domain experts who analyze rules and act on them, unless the minimum support threshold is set to a large value; second, the heuristic choice of the minimum support threshold tends to over-prune the search space of association rules, disregarding “nuggets” which can be very useful in many applications. Most prior work relies on the strong requirement of the minimum support threshold; some exceptions include the works of Li et al. (1999); Koh (2008) and DuMouchel and Pregibon (2001). Some recent work (Cohen et al., 2001; Wang et al., 2001) attempts to avoid the support measure altogether. In our work, the use of the adjusted confidence eliminates the need for the minimum support threshold.

When a set of rules is used to form a classifier, this is called “associative classification” (see, for instance, Liu et al., 1998; Thabtah, 2007; Vanhoof and Depaire, 2010).

### 7.2 Decision Lists

A decision list is an ordered set of association rules that forms a classifier (Rivest, 1987). Usually decision lists are formed the same way as decision trees are formed, which is by greedily splitting

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Number of data sets</th>
</tr>
</thead>
<tbody>
<tr>
<td>Support</td>
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</tr>
<tr>
<td>Confidence</td>
<td>1</td>
</tr>
<tr>
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</tr>
<tr>
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<td>5</td>
</tr>
<tr>
<td>$K=0.01$</td>
<td>6</td>
</tr>
<tr>
<td>$K=0.1$</td>
<td>6</td>
</tr>
<tr>
<td>$K=1$</td>
<td>2</td>
</tr>
<tr>
<td>$K=5$</td>
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<td>2</td>
</tr>
<tr>
<td>$K=15$</td>
<td>2</td>
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<tr>
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<td>0</td>
</tr>
<tr>
<td>$\theta=10$</td>
<td>1</td>
</tr>
</tbody>
</table>

Figure 6: Summary of experiments: For each algorithm, the number of data sets where it performed comparably with the best algorithm.
on each nodes to form the tree, and then pruning (as in for instance, Li et al., 2001; Yin and Han, 2003; Simon et al., 2011; Marchand and Sokolova, 2005). However, it is possible to mine a set of rules, and order them to produce a classifier, as in the associative classification literature.

The work of Anthony (2004) contains a generalization bound for decision lists, but each rule in the list requires a linear combination, which is problematic in the sequential setting by the reasoning in Appendix A. (Similarly, there are many papers using a set of pre-computed rules as features for supervised learning, where a linear combination of rules is constructed, rather than a decision list; one recent example is by Friedman and Popescu 2008.)

In recent work, we have been learning the ordering of rules to form decision lists (Letham et al., 2013c).

### 7.3 Recommender Systems

Association rule mining has proven to be particularly useful for finding “goes with” relationships between items purchased simultaneously. Lin et al. (2002) also construct a recommender system using rules, having a minimum confidence threshold and then an adjustable minimum support threshold. Their scoring system is essentially based on support × confidence, which is not an estimate of $P(b|a)$ for rule $a \rightarrow b$. Lawrence et al. (2001) provide a recommender system for a grocery store, but the setting differs entirely from ours in that they always recommend items that have never been previously purchased.

In other work, we designed a Bayesian framework that estimates $K$ for the adjusted confidence by “borrowing strength” across both users and items (McCormick et al., 2012). We are also looking at different approaches to the sequential event prediction problem, where we allow the predictions to alter the sequence in which items are placed into the basket (Letham et al., 2013b). This work uses a supervised learning framework for sequential event prediction.

We also note that a recommender system based on a weighted version of the adjusted confidence won third place in the ECML Discovery Challenge in 2013 (Letham, 2013).

Often, item-based collaborative filtering is used for problems that are actually sequential event prediction problems. There are several problems in applying standard item-based collaborative filtering techniques in sequential event prediction, the first one being that standard item-based collaborative filtering requires us to compute a similarity measure between all “co-rated” items. The similarity measure is often symmetric between two items, there is no distinguishing between $P(a|b)$ and $P(b|a)$. Even if item $b$ is always found when $a$ is found, $P(b|a) = 1$, is it possible for $b$ not to be recommended when $a$ is present, even with more than sufficient data to see the pattern. Further, for an incomplete basket, we do not have the ratings for all “co-rated” items, since there is no natural way to differentiate between items that have not yet been purchased in this transaction, and items that will not be purchased in this transaction, as both have a “rating” of 0 at time $t$. Thus, the only ratings that are available are ratings of “1” indicating that an item is in the basket. In other words, where the association rule approach we present here is intrinsically sequential, it is unnatural to force item-based collaborative filtering into a sequential framework. In general, item-based collaborative filtering is not based in a typical machine learning setting, in that it is not based on either loss minimization or probabilistic modeling (as the association rule approach is). The work of Letham et al. (2013b) also shows experimentally that item-based collaborative filtering can be worse than the max-confidence association rule approach.
7.4 Bayesian Analysis

DuMouchel and Pregibon (2001, “D&P”) present a Bayesian approach to the identification of interesting itemsets. While not a rule mining algorithm per se, the approach could be extended to produce rules. D&P consider the ratio of observed itemset frequencies to baseline frequencies computed under a particular independence model. A prior distribution over the collection of such ratios results in shrinkage estimates for the true ratios. The amount of shrinkage depends on the observed frequency and tends to be more pronounced for less frequent itemsets. Our approach differs from D&P in several key regards. Most importantly we focus directly on Bayesian estimation for rules rather than itemsets. Second, D&P use an empirical Bayes approach to choose the prior hyperparameters. Since our approach requires just a single hyperparameter, \( K \), we instead let the user choose an appropriate value (the value might be determined by cross validation or empirical Bayes). Finally, D&P perform a stratified analysis; one interesting future direction for our proposed approach would be to incorporate stratification.

Breese et al. (1998) present a number of different algorithms for collaborative filtering, including two Bayesian approaches. One of their Bayesian approaches clusters users while the other constructs a Bayesian network. Condliff et al. (1999) present a hierarchical Bayesian approach to collaborative filtering that “borrows strength” across users. Neither Breese et al. nor Condliff et al. focus on repeated purchases but both present ideas and techniques that may have relevance to future versions of our approach, especially the borrowing strength ideas.

Our recent work (McCormick et al., 2012; Letham et al., 2013c) uses Bayesian analysis to order rules into decision lists.

8. Conclusion

This work synthesizes tools from several fields to analyze the use of association rules in a new supervised learning framework. This analysis is necessarily different from that of classical supervised learning analysis; as we have discussed, association rules provide two mechanisms for generalization: first a large sample, and second, a minimum support of rules. We considered two simple algorithms based on association rules: a max confidence, min support algorithm, and the Bayesian adjusted confidence algorithm. Both algorithms have a parameter that creates a bound on the support, regulating a tradeoff between accuracy on the training set and generalization ability. We have also demonstrated that the adjusted confidence introduced here has some advantages over the minimum support threshold that is commonly considered in association rule mining: it allows rare rules to be used while still encouraging generalization, and among rules with similar confidence, it prefers those with larger support.

Acknowledgments

We would like to express thanks to Gene Kogan for helpful discussions and inspiration. Support for this project was provided by the National Science Foundation under grant IIS-1053407.
Appendix A. Regression and the Sequential Event Prediction Problem

By using association rules to model conditional probabilities for the sequential event prediction problem, we make a general assumption about the Markov chains governing our application, namely that a subset of knowledge about the current state can be used to predict the most likely future state. In this section we will address the suitability of two natural regression approaches that do not make this assumption. Let \( X_i \) be an indicator variable that is 1 if item \( i \) is in the current basket and 0 otherwise.

A.1 First Regression Method

Apply regression (e.g., logistic regression) to create a model for each item separately. Consider the model for the last item (item \( m \)), where the predictor variables will be \( X_i \) for \( i \in \{1, \ldots, m-1\} \), and \( X_m \) will be the response variable. This model would provide:

\[
P(X_m = 1|X_1 = x_1, \ldots, X_{m-1} = x_{m-1}) = \frac{1}{1 + \exp(f)}
\]

where \( f = \sum_{i=1}^{m-1} \lambda_i x_i + \lambda_{0,m} \), with each \( x_i \in \{0, 1\} \).

Because the data are being revealed sequentially, the correct application of this technique is not straightforward. Only a partial basket is available when predictions need to be made. It is incorrect to substitute the current state of the basket directly into the formula above. For instance, if the current basket contains items 1 and 2, so \( X_1 = 1 \) and \( X_2 = 1 \), it is incorrect to write \( P(X_m = 1|X_1 = 1, X_2 = 1) = \frac{1}{1 + \exp(f)} \), where \( f = \lambda_1 + \lambda_2 + \lambda_{0,m} \). This statement would be equivalent to the expression:

\[
P(X_m = 1|X_1 = 1, X_2 = 1) = P(X_m = 1|X_1 = 1, X_2 = 1, X_3 = 0, \ldots, X_{m-1} = 0),
\]

which is clearly false in general. It is not that, for instance, \( X_3 = 0 \), it is simply that \( X_3 \) is not yet realized.

On the other hand, it is possible to integrate in order to obtain conditional probability estimates:

\[
P(X_m = 1|X_1 = 1, X_2 = 1) = \sum_{x_3=\{0,1\}, \ldots, x_{m-1}=\{0,1\}} P(X_m = 1|X_1 = 1, X_2 = 1, X_3 = x_3, \ldots, X_{m-1} = x_{m-1} = x_m) \times P(X_3 = x_3, \ldots, X_{m-1} = x_{m-1} = x_m),
\]

where estimates of \( P(X_3 = x_3, \ldots, X_{m-1} = x_{m-1}) \) would need to be made also for every one of the \( 2^{m-3} \) combinations of \( x_3, \ldots, x_{m-1} \). Thus, this approach would rely on a large number of uncertain estimates (given limited data, and even moderately large \( m \)), each introducing errors into the final estimate. This is in contrast to the association rule approaches where a class of conditional probabilities are directly estimated. Further, the regression method provided above would not be able to be explained easily to customers or managers. In most circumstances, it would also require a large amount of computation between recommendations. Finally, it is not clear how to incorporate the order in which items are placed into the basket within this type of model, whereas it is trivial to incorporate this into the association rule techniques as discussed in Section 2.2.
### A.2 Second Regression Method

Apply regression methods (e.g., logistic regression) for each item and at each timestep, in total $m \times T$ regression models, where $T$ is the size of the largest possible basket. This would give a direct way to incorporate time into the predictions. If the current basket contains $t$ items, one would use only the models constructed using the first $t$ items in each basket to predict the next item to be added. However, this would be making an entirely different assumption than the one given by the rule-mining approach. The rule-mining approach uses time only implicitly, and purchase patterns are counted the same regardless of the exact time within the transaction when the pattern occurred. In contrast, this regression approach would ignore all items added after time $t$ in previous baskets. If apples were always followed by oranges, but in the past apples and oranges were always added after timestep $t$, then this approach would fail to recommend oranges when apples are added before timestep $t$. Further, the models for each timestep $t$ must be constructed from baskets at least as large as $t$. This means that for very large baskets, there would only be a few past baskets that could be used to construct the models. Further, if the current basket is larger than any of the past baskets, the models would be trivial, since none of the past baskets can be used to construct them.

It may indeed be possible to use regression approaches for the sequential event prediction problem, but given the discussion above, it is not clear how this should be accomplished. We explore other ways to solve the sequential event prediction problem using supervised ranking techniques in another work (Letham et al., 2013b).

### Appendix B. Additional Experimental Results

See Figures 7 - 11.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>mean ± standard dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Support</td>
<td>0.115 ± 0.0176</td>
</tr>
<tr>
<td>Conf.</td>
<td>0.1125 ± 0.0143</td>
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<tr>
<td>$K=0.0001$</td>
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</tr>
<tr>
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<tr>
<td>$K=0.01$</td>
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</tr>
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<td>0.1161 ± 0.0143</td>
</tr>
<tr>
<td>$\theta=10$</td>
<td>0.1205 ± 0.0191</td>
</tr>
</tbody>
</table>

Figure 7: *Left*: Boxplots of AvgCorrect values for Cars data set. *Right*: Means and standard deviations for Cars data set.
### Appendix C. Lemma 21

**Lemma 21** For $t \sim \text{Binomial}(m, p)$ and $K \geq 0$,

$$
\mathbb{E} \left[ \frac{1}{K + t} \right] = \frac{1}{K + mp} + O \left( \frac{1}{m^2} \right).
$$

The proof of this lemma for $K = 0$ is provided by Rempala (2003). The proof of this lemma for $K > 0$ comes from Letham et al. (2013a), which we provide here for completeness. The proof of the lemma uses the following result.
Lemma 22 Let \( X \sim \text{Binomial}(m, p) \) and let \( \mu_k = \mathbb{E} \left[ (X - \mathbb{E}[X])^k \right] \) be the \( k \)th central moment. For integer \( k \geq 1 \), \( \mu_{2k} \) and \( \mu_{2k+1} \) are \( O(m^k) \).

Proof We will use induction. For \( k = 1 \), the central moments are well known (e.g., Johnson et al., 2005): \( \mu_2 = mp(1-p) \) and \( \mu_3 = mp(1-p)(1-2p) \), which are both \( O(m) \). We rely on the following recursion formula (Johnson et al., 2005; Romanovsky, 1923):

\[
\mu_{s+1} = p(1-p) \left( \frac{d\mu_s}{dp} + ms\mu_{s-1} \right). 
\]

Figure 10: Left: Boxplots of AvgCorrect values for Plants data set. Right: Means and standard deviations for Plants data set.

Figure 11: Left: Boxplots of AvgCorrect values for T20I18D10KN22CR50 data set. Right: Means and standard deviations for T20I18D10KN22CR50 data set.
Because $\mu_2$ and $\mu_3$ are polynomials in $p$, their derivatives will also be polynomials in $p$. This recursion makes it clear that for all $s$, $\mu_s$ is a polynomial in $p$ whose coefficients include terms involving $m$.

For the inductive step, suppose that the result holds for $k = s$. That is, $\mu_{2s}$ and $\mu_{2s+1}$ are $O(m^s)$. Then, by (24),
\[
\mu_{2(s+1)} = p(1-p) \left( \frac{d\mu_{2s+1}}{dp} + (2s+1)m\mu_{2s} \right).
\]
Differentiating $\mu_{2s+1}$ with respect to $p$ yields a term that is $O(m^s)$. The term $(2s+1)m\mu_{2s}$ is $O(m^{s+1})$, and thus $\mu_{2(s+1)}$ is $O(m^{s+1})$. Also,
\[
\mu_{2(s+1)+1} = p(1-p) \left( \frac{d\mu_{2(s+1)+1}}{dp} + 2(s+1)m\mu_{2s+1} \right).
\]
Here $\frac{d\mu_{2(s+1)}{dp}$ is $O(m^{s+1})$ and $2(s+1)m\mu_{2s+1}$ is $O(m^{s+1})$, and thus $\mu_{2(s+1)+1}$ is $O(m^{s+1})$.

This shows that if the result holds for $k = s$ then it must also hold for $k = s + 1$ which completes the proof.

We can now prove Lemma 21.

**Proof** (Of Lemma 21) We expand $\frac{1}{K+X}$ at $X = mp$:
\[
\mathbb{E} \left[ \frac{1}{K+X} \right] = \mathbb{E} \left[ \sum_{i=0}^{\infty} (-1)^i \frac{(X - mp)^i}{(K+mp)^{i+1}} \right]
\]
\[
= \sum_{i=0}^{\infty} (-1)^i \mathbb{E} \left[ \frac{(X - mp)^i}{(K+mp)^{i+1}} \right]
\]
\[
= \frac{1}{K+mp} \sum_{i=2}^{\infty} (-1)^i \frac{\mu_i}{(K+mp)^{i+1}} \tag{25}
\]
where $\mu_i$ is the $i$th central moment and we recognize that $\mu_1 = 0$. By Lemma 22,
\[
\frac{\mu_i}{(K+mp)^{i+1}} = O \left( \frac{m^{\frac{1}{2}i}}{O(m^i)} \right) = O \left( m^{\frac{1}{2}i-1} \right).
\]
The alternating sum in (25) can be split into two sums:
\[
\sum_{i=2}^{\infty} (-1)^i \frac{\mu_i}{(K+mp)^{i+1}} = \sum_{i=2}^{\infty} O \left( m^{\frac{1}{2}i-1} \right) = \sum_{i=2}^{\infty} O \left( \frac{1}{m^i} \right) + \sum_{i=3}^{\infty} O \left( \frac{1}{m^i} \right).
\]
These are, for $m$ large enough, bounded by a geometric series that converges to $O \left( \frac{1}{m^n} \right)$. ■

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