Tree-based Partitioning of Data for Association Rule Mining

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Abstract. The most computationally demanding aspect of association rule mining is the identification and counting of support of the frequent sets of items that occur together sufficiently often to be the basis of potentially interesting rules. The task increases in difficulty with the scale of the data and also with its density. The greatest challenge is posed by data that is too large to be contained in primary memory, especially when high data density and/or low support thresholds give rise to very large numbers of candidates that must be counted. In this paper we consider strategies for partitioning the data to deal effectively with such cases. We describe a partitioning approach which organizes the data into tree structures that can be processed independently. We present experimental results that show the method scales well for increasing dimensions of data, and performs significantly better than alternatives, especially when dealing with dense data and low support thresholds.

Keywords: Association Rules, Partial Support, Data Structures, Set-Enumeration Tree

1. Introduction

An association rule (Agrawal et al, 1993) is an implication of the form $A \rightarrow B$, relating disjoint sets of database attributes, which is interpreted to mean “if the set of attribute-values $A$ is found together in a database record, then it is likely that the set $B$ will be present also”. Association Rule Mining involves the discovery, in a tabular database, of all such rules that satisfy defined threshold requirements. Of these requirements, the most fundamental concerns frequency: a rule is likely to be applicable only if the relationship it describes occurs sufficiently often in the data. The support for the rule $A \rightarrow B$ is the number (or

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proportion) of database records within which the set of attribute-values \( A \cup B \) is found. The frequent sets are those sets for which the support exceeds some threshold value. Association Rule Mining requires that all frequent sets are identified, and their support determined, so that other properties of rules such as confidence and lift \( (\text{Berry, 1997}) \) can be calculated.

It is recognised that identifying the frequent sets is the most computationally demanding aspect of Association Rule Mining. The problem arises because the number of possible sets is exponential in the number of possible attribute-values. Continuously-valued attributes can be dealt with by discretization, and, for convenience of processing, most methods convert multiple-valued attributes into a number of binary attributes, or items, each of which can be said to be present or absent in each record. For most real data, the number \( n \) of such items is likely to be such that counting the support of all \( 2^n \) sets of items (itemsets) is infeasible. For this reason, almost all methods attempt to count the support only of candidate itemsets that are identified as possible frequent sets. It is, of course, not possible to completely determine the candidate itemsets in advance, and it will therefore be necessary to consider many itemsets that are not in fact frequent.

In general, algorithms for finding frequent sets involve one or (usually) several passes of the source data, in each of which the support for some set of candidate itemsets is counted. The performance of these methods, clearly, depends both on the size of the original database, typically millions or billions of records, and on the number of candidate itemsets being considered. The number of possible candidates increases with increasing density of data (greater number of items present in a record) and with decreasing support thresholds. In applications such as medical epidemiology, we may be searching for rules that associate rather rare items within quite densely-populated data, and in these cases the low support-thresholds required may lead to very large candidate sets. These factors motivate a continuing search for efficient algorithms.

Performance will be affected, especially, if the magnitudes involved make it impossible for the algorithm to proceed entirely within primary memory. In these cases, some strategy for partitioning the data may be required to enable algorithmic stages to be carried out on primary-memory-resident data. Effective partitioning will reduce the number of accesses to secondary memory. In this paper we examine methods of partitioning to limit the total primary memory requirement, including that required both for the source data and for the candidate sets. We consider both ‘horizontal’ partitioning, which divides the source data into sets of records, and ‘vertical’ partitioning, which partitions records into sets of items. We describe a new method of vertical partitioning that exploits tree structures we have previously developed for Association Rule Mining. Experimental results are presented that show this method offers significantly better performance than horizontal partitioning.

2. Background

Most methods for finding frequent sets are based to a greater or lesser extent on the “Apriori” algorithm \( (\text{Agrawal and Srikant, 1994}) \). Apriori performs repeated passes of the database, successively computing support-counts for sets of single items, pairs, triplets, and so on. At the end of each pass, sets that fail to reach the required support threshold are eliminated, and candidates for the next pass
are constructed as supersets of the remaining (frequent) sets. Since no set can be frequent which has an infrequent subset, this procedure guarantees that all frequent sets will be found.

One of the inherent performance weaknesses of Apriori is that it requires the source data to be scanned repeatedly; in principle, the number of passes required is one greater than the size of the largest frequent set. This is especially a problem if, as is likely to be the case in many applications, the source data cannot be contained in primary memory. An early refinement of the method attempted to reduce this cost by partitioning the data into a number of equal-sized segments that can be so contained. The “Partition” algorithm (Savasere et al, 1995) applies the Apriori procedure to each data segment in turn, retaining the segment in primary storage throughout its repeated passes. For each segment, thus, a set of locally frequent itemsets is determined, each of which reaches the proportionate threshold of support in that segment. A second pass of the complete database is required to establish which of the locally frequent sets are (globally) frequent. Similar thinking motivated the strategy introduced by Toivonen (1996). Here, a random sample of the source data, small enough to contain in primary memory, is first processed using the Apriori procedure, with a modified support threshold and other modifications designed to make it likely that all the globally frequent sets will be identified in the sample. The sets thus found become candidates for a single full pass of the source data to verify this.

The drawback of both these approaches highlights the second weakness of Apriori: that the number of candidates whose support is to be counted may become very large, especially when the data is such that the frequent sets may contain many items (the “long pattern” problem (Agrawal et al, 2000)). If, for example, there is just one set of 20 items that reaches the threshold of support, then the method inescapably requires the support for all the $2^{20}$ subsets of this set to be counted. In the Partition algorithm, this is exacerbated because there may be many more sets that are locally frequent in some partition, even though they are not globally frequent. In Toivonen’s method, also, it is necessary for the initial processing of the sample to identify an enlarged candidate set, to give a reasonable probability that all the actual frequent sets will be included. Both these methods also require all candidates to be retained in primary memory (for efficient processing) during the final database pass.

Other methods (Bayardo, 1998; Bayardo, 1999; Agrawal et al, 2000; Zaki et al, 1997) aim to identify maximal frequent sets without first examining all their subsets. These algorithms may cope better with densely-populated databases and long patterns than the others described, but again usually involve multiple database passes. The DepthProject (Agrawal et al, 2000) algorithm bypasses the problem by explicitly targeting memory-resident data. The method of Zaki et al. (1997) is of interest, as it introduces a different kind of partitioning. In this, candidate sets are partitioned into clusters which can be processed independently. The problem with the method is that, especially when dealing with dense data and low support thresholds, expensive pre-processing is required before effective clustering can be identified. The partitioning by equivalence class, however, is relevant to the methods we will describe.

Our methods begin by performing a single pass of the database to perform a partial summation of the support totals. These partial counts are stored in a tree structure that we call the $P$-tree, which enumerates itemsets counted in lexicographic order. The term $P$-tree has been used elsewhere (e.g. (Huang et al, 2002)) to describe other structures: here we use it to denote the structure we
first introduced in (Goulbourne et al., 2000), representing a tree of Partial support counts. The P-tree contains all the sets of items present as distinct records in the database, plus some additional sets that are leading subsets of these.

To illustrate this, consider a database with items \{a, b, c, d, e\}, and 20 records:

\{abce, abde, abd, abde, abe, ace, ade, b, bde, bce, bd, be, cd, cde, ce, d, de, e\}

(Not necessarily in this order). For convenience, we will use the notation \text{abd}, for example, to denote the set of items \{a, b, d\}. Figure 1 shows the P-tree that would be constructed. The counts stored at each node are incomplete support-totals, representing support derived from the set and its succeeding supersets in the tree.

We then apply to this structure an algorithm, Apriori-TPF, which completes the summation of the final support counts, storing the results in a second set-enumeration tree (the T-tree, of Total support counts), ordered in the opposite way to the P-tree. The T-tree finally contains all frequent sets with their complete support-counts. The algorithm used, essentially a form of Apriori that makes use of the partial counting that has already been done, is described in (Coenen et al., 2001a), where we also explain the rationale of the approach and its advantages. Experimental results reported in (Coenen et al., 2001a) demonstrate significant performance gains in comparison with Apriori, and also some improvements over the FP-growth (Han et al., 2001) algorithm, which uses somewhat similar structures and has some similar properties. The FP-tree used in (Han et al., 2001) is a more pointer-rich structure than the P-tree, leading to greater difficulties in dealing with non-memory-resident data, although strategies for this have been proposed, which will be discussed further below. The CATS tree, an extension of the FP-tree proposed in (Cheung and Zaiane, 2003), also assumes no limitation on main memory capacity. In this paper we consider implementations of Apriori-TPF in cases when it will be impossible to contain all the data required in main memory, requiring some strategy for partitioning this.

3. Strategies for Partitioning

3.1. Horizontal partitioning

The natural implementation of Apriori, when source data cannot be contained in primary memory, requires all the data to be read from secondary memory in each pass. The equivalent for Apriori-TPF, because the first stage of the method involves the construction of a P-tree, requires a partitioning of the data into segments of manageable size. We will refer to this form of partitioning, in which each segment contains a number of complete database records, as 'horizontal' partitioning (HP), or segmentation. We first take each segment of data separately and create for it a P-tree that is then stored in secondary memory. The stored P-trees are then treated as a composite structure from which we compute the final support totals for all the frequent sets, storing these in a single T-tree. Each pass of Apriori-TPF requires each of the P-trees to be read in turn from secondary memory. The method creates a final T-tree in primary memory, which contains all the frequent sets and their support-counts.
3.2. Vertical partitioning

The drawback of the simple approach outlined above is that it replicates the two weaknesses of the Apriori methodology. As with Apriori, all the source data (now in the form of P-trees) must be re-read from secondary memory in each pass. The second problem is that the entire T-tree, which finally contains all the frequent sets, must be contained in primary memory while counting proceeds. As we have noted, this tree may itself become very large, especially when long frequent patterns are encountered. Even if the tree is not too large to be contained in primary memory, a large set of candidates leads to slower counting, in Apriori-TFP, just as for Apriori.

A possible alternative way of partitioning the data is to divide the set of items under consideration into subsets, each of which defines a vertical partition of the data set. The problem with this is that, in general, the sets for which support is to be counted contain items from several partitions. The P-tree structure offers another form of vertical partitioning, into subtrees that represent equivalence classes of the items represented. In this case, again, it is still not possible to compute the support for a set by considering only the subtree in which it is located. Although succeeding supersets of a set S in the P-tree are located in the subtree rooted at S, predecessor supersets are scattered throughout the preceding part of the P-tree. For example, consider the support for the set bd in the data used for Figure 1. In the subtree rooted at b, we find a partial support total for bd, which includes the total for its superset bde. To complete the support count for bd, however, we must add in the counts recorded for its preceding supersets.
$bde$, $abd$ (incorporating $abde$) and $abde$, the latter two of which are in the subtree headed by $a$.

The problem can be overcome by a different partitioning of the $P$-tree structure. Our Tree Partitioning (TP) method begins by dividing the ordered set of items into subsequences. For example, for the data used in Figure 1, we might define 3 sequences of items, $\{a, b\}$, $\{c, d\}$ and $\{e\}$, labelled 1, 2, 3 respectively. For each sequence we define a Partition-$P$-tree ($PP$-tree), labelled $PP_1$, $PP_2$ and $PP_3$. The construction of these is a slight modification of the original method. The first partition-tree, $PP_1$, is a proper $P$-tree that counts the partial support for the power set of $\{a, b\}$. $PP_2$, however, counts all those sets that include a member of $\{c, d\}$ in a tree that includes just these items and their predecessors. The third tree, $PP_3$, will count all sets that include any member of $\{e\}$. The three trees obtained, from our example, are illustrated in Figure 2. The $PP$-trees are, in effect, overlapping partitions of the $P$-tree of Figure 1, with some restructuring resulting from the omission of nodes when they are not needed.

The effect of this is that the total support for any set $S$ can now be obtained from the $PP$-tree corresponding to the last item within $S$; for example, we now find all the counts contributing to the support of $bd$ are included in $PP_2$. The drawback is that the later trees in the sequence are of increasing size; in particular, $PP_3$ in our example is almost as large as the original $P$-tree. We can overcome this, however, by a suitable reordering of the items. In descending order of their frequency in the data, the items of our example are $e$, $d$, $b$, $c$ and $a$. Using the same data as for Figures 1 and 2, we will construct $PP$-trees using this ordering, for the sets of items $\{e, d\}$, $\{b, c\}$ and $\{a\}$ respectively.

The results are shown in Figure 3. Now, because the less frequent items appear later in the sequence, the trees become successively more sparse, so that $PP_3$ now has only 13 nodes, compared with the 23 of $PP_3$ in Figure 2. In fact, our previous work has shown (Coenen and Leng, 2001b) that ordering items in this way leads to a smaller $P$-tree and faster operation of Apriori-TP. The additional advantage for partitioning is that the $PP$-trees become more compact and more equal in size. The total support-count for $bd$ (now ordered as $db$) is again to be found within $PP_2$, but now requires the addition of only 2 counts ($db + edb$).

### 3.3. Counting total support using $PP$-trees

The form of partitioning we have described offers us a way of dividing the source data into a number of $PP$-trees each of which may then be processed independently. With a sufficiently large data set, it will of course still not be possible to construct the $PP$-trees within primary memory. We can, however, combine this approach with a (horizontal) segmentation of the original data into segments small enough to allow the corresponding $PP$-trees to be contained in primary store.

The overall method is as follows. For clarity, we will use the term segment when we refer to the horizontal division of the data into sets of records, and partition when we refer to the vertical division into sets of items and the corresponding tree structures:

1. Obtain an (at least approximate) ordering of the frequency of items.
2. Using this ordering, choose an appropriate partitioning of the items into \( n \) sequences \( 1, 2, 3, \ldots \) etc.

3. Divide the source data into \( m \) segments.

4. For each segment of data, construct \( n \) PP-trees in primary memory, storing finally to disk. This construction phase involves just one pass of the source data.

5. For partition 1, read the PP1 trees for all segments into memory, and apply the Apriori-TPF algorithm to build a \( T \)-tree that finds the final frequent sets in the partition. This stage requires the PP1 trees for each segment of data to be read once only. The \( T \)-tree remains in memory throughout, finally being stored to disk.

6. Repeat step 5 for partitions 2, 3, \ldots, \( n \).

The method offers two speed advantages over simple horizontal segmentation. First, we have now effectively reduced the number of disk passes to 2: one (step 4) to construct the PP-trees, and a second pass (of the stored trees) to complete the counting (steps 5 and 6). The second advantage is that we are now, at each stage, dealing with smaller tree structures, leading to faster traversal and counting.
3.4. Comparison with other methods

As we have noted above, our P-tree structure, first presented in (Goulbourne et al., 2000), has many properties in common with the FP-tree structure developed independently and contemporaneously by Han et. al. (2001). The principal differences are two. First, the nodes of the FP-tree correspond to individual items, whereas in the P-tree a sequence of items which is partially closed (i.e. which has no leading subsequence with greater support in the tree) will be stored as a single tree node. Thus, for example, two transactions \{a, b, c, d, e\} and \{a, b, c, x, y\}, which share a common prefix \{a, b, c\}, would require in all 7 nodes in the FP-tree.

In the P-tree, conversely, only 3 nodes would necessarily be created: a parent for \{a, b, c\}, and child nodes for \{d, e\} and \{x, y\}. The second difference is that, in order to implement the FP-growth algorithm, the FP-tree must store pointers at each node to link all nodes representing the same item, and also to link a node to its parent and child nodes. The Apriori-TPP algorithm, however, treats the P-tree essentially as a set of nodes which can be processed in any order. This makes it possible, once the tree has been constructed, to store it in a tabular form in which no pointers are required.

Both these differences lead to a more compact tree structure. Furthermore, the absence of pointers allows us easily to use horizontal partitioning in order to build a succession of P-trees, each of which is vertically partitioned into PP-trees, as described above. It is then straightforward to collect all the PP-trees representing a single partition from their separate segments, and use them in any order to construct the final T-tree for that partition.
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Partitioning the FP-tree is necessarily more complex. The partitioning we describe, as illustrated in Figure 3, is essentially similar to that obtained by the construction of conditional databases described in (Han et al, 2001) and (Pei et al, 2000). In (Han et al, 2004) two strategies are proposed for dealing with an FP-tree too large for primary storage. In the first of these, parallel projection, the original database is partitioned into a set of projected databases, one for each item. Each projected database contains only transactions in which the item is present with some predecessors in the item ordering. Thus, each projected database essentially represents the same information as would be contained in a corresponding PP-tree in our method. The second method described, partition projection, would (using our example to illustrate) first construct the a-conditional database corresponding to PP3, and after building the FP-tree for this, would copy relevant transactions (e.g. \{e, d, b, ca\}), into the next (c-conditional) database, as \{e, d, b, c\}. This reduces the total size of the projected databases, at a cost of some additional processing.

A number of other researchers have made use of the FP-tree or variants thereof. The CFR-tree described in (Liu et al, 2003) stores frequent closed itemsets in a form that facilitates subsequent query processing. The construction algorithm is similar to that used for FP-tree construction, using conditional databases as described in (Han et al, 2001) to partition the data and produce separate trees for query processing. The focus of this work is on the form of a structure that can be re-used efficiently, rather than on the efficiency of the construction algorithm. Reusability is also a feature of our P-tree structures, which retain all relevant information from the original data as well as performing part of the support-counting. In (Huang et al, 2002) a structure is described, also (coincidentally) called a P-tree, which is quite similar to our P-tree, but (like the FP-tree) stores only one item at each node. The P-tree of (Huang et al, 2002) shares with ours the property of retaining all the information from the database needed for counting of support at any threshold, rather than just counts of frequent sets. The approach described in (Huang et al, 2002) constructs FP-trees from the P-tree rather than from the original data; results presented show that this offers significant performance gains when multiple FP-trees are required. The partitioning strategy described for dealing with large databases is essentially horizontal, dividing the data into segments for each of which a P-tree is constructed. The method produces a single overall FP-tree, however, for which further partitioning might become necessary.

The COFI-trees proposed in (El-Hajj and Zaiane, 2003a) also create subtrees that can be processed independently, but require an initial construction of an FP-tree that must be retained in primary memory for efficient processing. In (El-Hajj and Zaiane, 2003b), a method is described for building COFI-trees from an inverted database structure called “Inverted Matrix”. In this structure, each item is represented by a row of the matrix which lists all transactions in which the item occurs. Each element of the list contains a pointer to the next item in the transaction. Constructing the Inverted Matrix requires two passes of the original database and will, in general, lead to an expansion in size because of the need to store a pointer with every item occurrence. The COFI-trees constructed from this have similar properties to the conditional FP-trees produced by database projection, as in (Han et al, 2001). However, to produce the COFI-trees it is necessary to mine the large inverted matrix, and the pointers between rows of this imply that no simple partitioning of this is possible. The approach seems
to work well with very sparse databases, but for dense data it seems likely that following the links through a disk-resident matrix will be a costly overhead.

In general, all methods that use FP-tree-like structures require to employ projection of conditional databases in some manner in order to avoid the problem of dealing with a single large FP-tree or its equivalent in primary memory. In our experiments, described below, we compare the performance of Apriori-TP using our Tree Partitioning method with that of FP-growth using database projection.

4. Results

We first consider some general performance properties of our method. To investigate performance, we have used synthetic data sets constructed using the QUEST generator described in (Agrawal and Srikant, 1994). The programs were written in standard C++ and run under the Linux operating system. We performed these experiments on an AMD Athlon workstation with a clock rate of 1.3 GHz, 256 Kb of cache, and 512 Mb of RAM. The data was stored on an NFS server (1Gb filestore).

We first need to establish that the method scales acceptably; that is, that the partitioning strategy successfully constrains the maximum requirement for primary memory, without leading to unacceptable execution times. For this purpose we generated data sets with parameters T10, 5.500: i.e 500 items, with an average record-length of 10 items and an expected maximal frequent pattern size of 5. We divided the data into segments of 50,000 records, and within each segment generated 500 partitions, i.e. a PP-tree for each item. In this and all other experiments, the tree partitioning is naive: after ordering the items, we partition into sequences of equal length (in this case, 1). In fact, our experiments seem to show that increasing the degree of (vertical) partitioning always reduces the primary memory requirement (as would be expected), and also almost always reduces execution time. The latter, less obvious result arises because the increased time taken to construct a greater number of PP-trees (step 4 of the algorithm outlined above) is usually more than compensated by the faster processing of smaller T-trees (steps 5 and 6).

Figure 4 shows the overall time to generate frequent sets, with a support threshold of 0.01%, for datasets of increasing size, i.e. 1, 2, 3, 4 and 5 segments. The figure shows the performance of the Tree-partitioning method (TP) in comparison with the simple method involving horizontal partitioning (segmentation) only (HP). The times illustrated include both the time to construct the P-trees (HP method) or PP-trees (TP method) and to execute the Apriori-TPF algorithm. As can be seen, Tree Partitioning offers substantially better performance than horizontal partitioning, and its performance scales linearly with the size of the dataset.

Importantly, this performance is achieved within conservative requirements for primary storage. In the second phase of the TP method (steps 5 and 6 of the algorithm outlined above) it is necessary to contain in memory all the PP-trees for one partition, and the corresponding T-tree containing the frequent sets in that partition. In the experiment of Figure 4, this led to a maximum memory requirement for the TP method that varied from 1.38 Mb (1 segment) to 1.6Mb (5 segments). In general, larger data sets, requiring greater horizontal segmentation, lead to some increase in the combined size of the PP-trees, but this is relatively slight. By contrast, the HP method requires the P-tree for one
data segment and the whole of the \( T \)-tree to be contained in primary store, leading to a maximum memory requirement of between 116 and 128 Mb in the case illustrated.

The combined sizes of the \( PP \)-trees for any one segment are, of course, greater than the size of a corresponding \( P \)-tree. In the experiment of Figure 4, the sum of the sizes of the \( PP \)-trees for any one segment was about 15.56 Mb (varying little between segments), compared to a \( P \)-tree size of about 1.85 Mb. This was not the dominant store requirement in the case we have illustrated, but in other cases could be a constraint during the construction of the \( PP \)-trees (step 4 of the method, above). If this is so, the problem can easily be overcome by imposing a greater degree of horizontal segmentation. Our experiments show that increasing the number of segments has little effect on execution times, while reducing memory requirements during the \( PP \)-tree construction. Figure 5 shows the results of experiments with increasing segmentation of \( T10.I5.N500.D50000 \), again with a support threshold of 0.01%. Having first established the linear scaling of the method, in the results presented above, this experiment for convenience used this relatively small database, but in order to replicate the problems of dealing with large databases we imposed a requirement that only one segment can be retained in primary store at one time.

In this case we imposed a vertical partitioning of 50 items/partition (10 partitions in all), while varying the number of segments. Figure 5 shows that the overall execution time (for both methods) increases only slightly with increasing segmentation. The total memory requirement to contain all the \( PP \)-trees for any one segment decreases, as one would expect, from 7.8 Mb (1 segment) to a maximum of 0.22 Mb (50 segments).

The method scales well with increasing number of items, also. Figure 6 shows the execution times for the TP method for \( T10.I5.D50000 \), with a support threshold of 0.01, a vertical partitioning of 10 items/partition, and horizontal partitioning of 10,000 records per segment (i.e., 5 segments). The maximum

\[ \text{Time (sec)} \]

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure4}
\caption{Execution times for T10.I5.N500 (0.01% support)}
\end{figure}
memory requirement in this case also remained small (between 6 and 15Mb, with no general upward trend). Conversely, the simple HP method has a rapidly increasing memory requirement in this case because of the greater size of the unpartitioned T-tree.

The performance of the method when dealing with more dense datasets is shown in figures 7 and 8. In these experiments, we generated databases with D=250000, N=500, varying the T and I parameters. Here we compared the TP method both with HP and with a method based on the "Negative Border" approach of (Toivonen, 1996) (labelled NB in the figures). In the latter, P-trees are first constructed for all segments, as for the HP method. Then, all the proportionately-frequent sets in the first segment of data are found, using a support threshold reduced to 2/3 of the original, and also retaining the negative border of the sets found, i.e. those sets which, although not themselves frequent,
have no infrequent subsets. The frequent sets, with their negative border, are stored in a $T$-tree which is kept in store to complete the counts for these sets for the remaining segments. The reduced support threshold, and the inclusion of the negative border, make it very likely that all the finally frequent sets will be included in this tree, in which case the method requires the disk-stored $P$-trees to be read once only.

The figures show the results with a support threshold of 0.1%, using 5 segments, and 10 items/partition. With this support threshold (higher than in the previous experiments) there is little difference in the total execution times for the T10.5 data; in this case, the faster computation of the frequent sets by the TP method is offset by the longer time taken to construct the $PP$-trees. With more dense data, however, the latter factor becomes increasingly significant, and the advantage of the TP method becomes increasingly apparent. This is principally because of the much smaller candidate sets that are involved. This becomes apparent from the comparison of maximal memory requirements, shown in Figure 8. This reflects the growing size of the candidate sets (and hence the $T$-tree) as the data density increases, leading both to larger memory requirements and to longer times to find candidates. The problem is particularly acute with the ’Negative Border’ method. This works well with relatively sparse data, but at high density the reduced support threshold and the inclusion of the negative border lead to very large candidate sets.

In Figures 9 and 10 we compare the performance of the three methods for different support thresholds. These results again relate to the T10.5.5.N500.D250000 data, for support thresholds decreasing from 1.0 through to 0.01. In these experiments, we divided the data into 5 segments, and for the TP method used 500 partitions (1 item/partition). Here again, as can be seen from Figure 9, the overhead of constructing the multiple $PP$-trees for the TP method leads to relatively poor execution times when the support threshold is high; in this case, the NB method is fastest. As the support threshold is reduced, however, the increasing cost of servicing a growing candidate set leads to rapidly increasing memory requirements and execution times for the alternative methods, whereas the TP method scales much better.

Finally, we compare our method with an implementation of $FP$-$growth$, using the $FP$-tree structure of (Han et al, 2001). For this purpose, we implemented both Apriori-TP, using our TP method, and an $FP$-$growth$ algorithm in Java, with the aim of obtaining as fair a comparison as possible. In this experiment, we used the dataset T20.110.N500.D500K which was divided into 5 equal segments. For $FP$-$growth$, we processed each segment in turn to generate conditional databases for each of the 500 items, using the parallel projection method described in (Han et al, 2004). The conditional database-segments were then combined in order to build an overall conditional-$FP$-$tree$ for each item in turn, to which the $FP$-$growth$ algorithm was applied to produce frequent sets. For Apriori-TP, each segment is used to construct 500 $PP$-trees, as described above. The experiments were in this case run on a 1.2 GHz Intel Celeron CPU with 512 MBytes of RAM.

The overall execution times for the two methods are shown in Figure 11, for support thresholds from 1.0 down to 0.1. As can be seen, on this data Apriori-TP with tree partitioning outperforms a comparable implementation of $FP$-$growth$ partitioned by parallel projection. We believe the advantage of Apriori-TP in this case arises from the relatively efficient iterative processing of the simple $P$-tree structures, in comparison with the recursive generation of multiple
5. Conclusions

Because Data Mining is principally concerned with obtaining information from data of very large dimensions, it is important that methods used should scale effectively to deal with the most extreme cases. In this paper we have examined ways of partitioning data for Association Rule Mining. Our aim has been to identify methods that will enable efficient counting of frequent sets in cases where the data is much too large to be contained in primary memory, and also where the density of the data means that the number of candidates to be considered becomes very large. Our starting point was a method which makes use of an initial preprocessing of the data into a tree structure (the \textit{F}-tree) which incorpo-

\textit{FP}-trees required by \textit{FP-growth}. This is especially expensive with relatively dense data, because of the greater depth of recursion required.
rates a partial counting of support totals. In previous work we have shown this method to offer significant performance advantages. Here, we have investigated ways of applying the approach in cases that require the data to be partitioned for primary memory use. We have, in particular, described a method that involves a partitioning of the tree structures involved to enable separate subtrees to be processed independently. The advantage of this approach is that it allows both the original data to be partitioned into more manageable subsets, and also partitions the candidate sets to be counted. The latter results in both lower memory requirements and also faster counting.

The experimental results we have reported here show that the Tree Partitioning method described is extremely effective in limiting the maximal memory requirements of the algorithm, while its execution time scales only slowly and linearly with increasing data dimensions. Its overall performance, both in execu-
tion time and especially in memory requirements, is significantly better than that obtained from either simple data segmentation or from other methods considered. The advantage increases with increasing density of data and with reduced thresholds of support - i.e. for the cases that are in general most challenging for association rule mining. Furthermore, a relatively high proportion of the time required by the method is taken up in the preprocessing stage during which the PP-trees are constructed. Because this stage is independent of the later stages, in many applications it could be accepted as a one-off data preparation cost. In this case, the gain over other methods becomes even more marked. Note also that the P-tree construction, and the partitioning thereof, is essentially generic: it leads to no loss of relevant information, and so could be used as the first stage of other quite different algorithms for completing the support-counts. A further advantage, not examined here, is that the independent processing of subtrees can be carried out in parallel.

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