An Effective Process for Finding Frequent Sequential Traversal Patterns on Varying Weight Range

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Summary
Many frequent sequential traversal pattern mining algorithms have been developed which mine the set of frequent subsequences traversal pattern satisfying a minimum support constraint in a session database. However, previous frequent sequential traversal pattern mining algorithms give equal weightage to sequential traversal patterns while the pages in sequential traversal patterns have different importance and have different weightage.

Another main problem in most of the frequent sequential traversal pattern mining algorithms is that they produce a large number of sequential traversal patterns when a minimum support is lowered and they do not provide alternative ways to adjust the number of sequential traversal patterns other than increasing the minimum support. In this paper, we propose a frequent sequential traversal pattern mining algorithm with dynamic weight constraint.

Our main approach is to support the weight constraints into the sequential traversal pattern while maintaining the downward closure property with weight range. A weight range is defined to maintain the downward closure property and pages are given different range and traversal sequences assign a minimum and maximum weight. Here we are accessing pattern utility graph for dynamically approach. In scanning a session database, a maximum and minimum weight in the session database is used to prune infrequent sequential traversal subsequence by doing downward closure property can be maintained. Our method produces a few but important sequential traversal patterns in session databases with a low minimum support, by adjusting a weight range of pages and sequence.

Keywords:
Sequential traversal pattern mining, Weight constraint, Web usage mining, Data mining

1. Introduction
The World Wide Web is an immense source of data that can come either from the Web content, represented by the billions of pages publicly available, or from the Web usage, represented by the log information daily collected by all the servers around the world. Web Mining is that area of Data Mining which deals with the extraction of interesting knowledge from the World Wide Web [1].

More precisely, Web Content Mining is that part of Web Mining which focuses on the raw information available in Web pages; source data mainly consist of textual data in Web pages (e.g., words, but also tags); typical applications are content-based categorization and content-based ranking of Web pages [2]. Web Structure Mining is that part of Web Mining which focuses on the structure of Web sites; source data mainly consist of the structural information present in Web pages (e.g., links to other pages); typical applications are link-based categorization of Web pages, ranking of Web pages through a combination of content and structure [3], and reverse engineering of Web site models. Web Usage Mining is that part of Web Mining which deals with the extraction of knowledge from server log files; source data mainly consist of the (textual) logs that are collected when users access Web servers and might be represented in standard formats (e.g., Common Log Format, Extended Log Format, LogML)[5]; typical applications are those based on user modeling techniques, such as Web personalization, adaptive Web sites, and user modeling.

Srivastava et al. [6] systematically discuss the development of WUM and classify the content of WUM. Zhang and Liang [7] show the importance of data preprocessing in Web Usage Mining and present an algorithm called "USIA" which boasts high efficiency. Wang and Meinel [8] point out that user behaviors recovery and pattern definition play more important roles in web mining than other applications so they give a new insight on behavior recovery and complicated pattern definition. As current Web Usage Mining applications rely exclusively on the server log files, Guo et al. [9] propose a system that integrates Web page clustering into log file association mining and use the cluster labels as Web page content indicators in the hope of mining novel and interesting association rules from the combined data source.

2. Problem Definition And Related Work

2.1 Problem definition
Let P = {P1, P2... Pn} be a unique set of pages. A session S is an ordered list of itemsets, denoted as (s1, s2, .., sm),
SPADE and PrefixSpan were two of the fastest algorithms. Bitmap representation of each sequence. Before SPAM, traversal of the search space combined with a vertical format was proposed which improves the mining process. The main idea of PrefixSpan is to examine only the prefix subsequences and project only their corresponding suffix subsequences into projected databases. In each projected database, sequential patterns are grown by exploring only the prefix of the sequence. A sequence database, $D = \{S_1, S_2, .., S_n\}$, is a set of tuples $(sid, s)$, where $sid$ is a sequence identifier and $S$ is an input sequence. A sequence $a = (X_1, X_2, .., X_n)$ is called a subsequence of another sequence $b = (Y_1, Y_2, .., Y_m)$ $(n \leq m)$, and $b$ is called a super sequence of $a$ if there exist an integer $1 \leq i_1 < ... < i_n < m$ such that $X_1 \subseteq Y_{i_1}$, $..., X_n \subseteq Y_{i_n}$. A tuple $(sid, S)$ is said to contain a sequence $Sa$ if $S$ is a super sequence of $Sa$. The support of a sequence $Sa$ in a sequence database $D$ is the number of tuples in SDB that contains $Sa$. Given a support threshold, $\text{min}_\text{sup}$, a sequence $Sa$ is a frequent sequence in the sequence database if the support of the sequence $Sa$ is no less than a minimum support threshold. The problem of sequential pattern mining is to find the complete set of sequential patterns in the database with a support constraint.

### 2.2 Related work

GSP [18] mines sequential patterns based on an Apriori like approach by generating all candidate sequences. This is inefficient and ineffective. To overcome this problem, the database projection growth based approach, FreeSpan [15], was developed. Although FreeSpan outperforms the Apriori based GSP algorithm, FreeSpan may generate any substring combination in a sequence. The projection in FreeSpan must keep all sequences in the original sequence database without length reduction. PrefixSpan [16], a more efficient pattern growth algorithm was proposed which improves the mining process. The main idea of PrefixSpan is to examine only the prefix subsequences and project only their corresponding suffix subsequences into projected databases. In each projected database, sequential patterns are grown by exploring only local frequent patterns.

In SPADE [25], a vertical id-list data format was presented and the frequent sequence enumeration was performed by a simple join on id lists. SPADE can be considered as an extension of vertical format based frequent pattern mining. SPAM [4] utilizes depth first traversal of the search space combined with a vertical bitmap representation of each sequence. Before SPAM, SPADE and PrefixSpan were two of the fastest algorithms. According to performance evaluations [4], SPAM outperforms SPADE on most datasets and PrefixSpan outperforms SPAM slightly on very small datasets. Except for this case, SPAM outperforms PrefixSpan.

### 3. Proposed Work

In this section, we suggest an efficient sequential traversal pattern mining algorithm in which the main approach is to apply weight constraints into the frequent sequential traversal tree while maintaining the downward closure property. We discuss our algorithm in detail and show actual examples for sequential traversal pattern mining with weight constraint.

**Definition 3.1 Weight Range**: A weight of a web page is a non-negative real number that shows the importance of each web page. The weight of each web page is assigned to reflect the importance of each web page in the session database.

**Definition 3.2 Traversal sequence with weight**: We can use the term, traversal sequence with weight to represent a set of sequential traversal patterns with weight.

**Definition 3.3 Average Weight of traversal**: We can use the term, average weight of subsequence is the sum of weight all pages in traversal divided by total number of pages in sequence.

**Definition 3.4 Minimum and Maximum weight of subsequence**: Here we define the maximum and minimum weight of traversal is average weight. If the weight of sequence come under the maximum and minimum weight range than given sequence is frequent otherwise infrequent.

### 3.1 Sequential traversal pattern with weight constraint

In this paper, pages of traversals are assigned with weights to show their importance. For example, when users traverse web site, they may have different interest in each page, and therefore stay for different times. Web pages can be assigned with a weight standing for the user stay time, frequency of pages, content of pages and type of web site. This paper generalizes the mining problem to the case where pages of traversals are given such weights showing their importance. The weights are taken into account in the measurement of support, the ratio of traversals which contains a candidate pattern. If a page of traversal has a weight which doesn’t between the weight ranges then it is removed from session of user and treated as an outlier, and can not consider for the support. For example, when users visit web site, they may traverse through a page very fast to another page, or do another
work for a long time during web site visit. This type of page visit is not useful and consider as an outlier because the page is not attentively read by the user.

Table 1: A sequence database as a running example.

<table>
<thead>
<tr>
<th>Sid</th>
<th>Traversal Sequence</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>P2 P1 P3 P4 P1 P5</td>
<td>0.2,0.3,0.12,0.34,0.6,0.3</td>
</tr>
<tr>
<td>S2</td>
<td>P1 P2 P4 P3 P4 P2</td>
<td>0.12,0.5,0.91,0.12,0.4,0.26</td>
</tr>
<tr>
<td>S3</td>
<td>P1 P2 P1 P3 P6 P7</td>
<td>0.6,0.2,0.32,0.56,0.45,0.7</td>
</tr>
<tr>
<td>S4</td>
<td>P2 P3 P6 P5 P1 P4</td>
<td>0.5,0.56,0.32,0.23,0.7,0.54</td>
</tr>
</tbody>
</table>

Table 2: The example of page with weight range

<table>
<thead>
<tr>
<th>S.No.</th>
<th>Page</th>
<th>Support</th>
<th>Weight Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>P1</td>
<td>4</td>
<td>0.12 – 0.56</td>
</tr>
<tr>
<td>2</td>
<td>P2</td>
<td>4</td>
<td>0.45 – 0.67</td>
</tr>
<tr>
<td>3</td>
<td>P3</td>
<td>4</td>
<td>0.23 – 0.67</td>
</tr>
<tr>
<td>4</td>
<td>P4</td>
<td>3</td>
<td>0.12 – 0.45</td>
</tr>
<tr>
<td>5</td>
<td>P5</td>
<td>2</td>
<td>0.34 – 0.67</td>
</tr>
<tr>
<td>6</td>
<td>P6</td>
<td>2</td>
<td>0.24 – 0.8</td>
</tr>
<tr>
<td>7</td>
<td>P7</td>
<td>0</td>
<td>0.12 – 0.56</td>
</tr>
</tbody>
</table>

In this section, we propose the concept of sequential traversal patterns with weight constraint, and show their importance.

Example: In session S1 the weight of P2 is 0.2 and the support is 4. The weight range for P2 is from 0.45 to 0.67. So, when we construct the frequent sequential traversal pattern tree P2 is eliminated from session.

3.2 Frequent Sequential Traversal Pattern Tree with weight constraint

In this section, a data structure called FSTP-tree is constructed. FSTP-Tree is a data structure, which must satisfy the following conditions. Firstly, it consists one root “null”, a set of item prefix subtrees as the children of the root, and a frequent-page head table. Secondly, every page in the page prefix subtree contains three fields: the name of the page, the support of the page, and a link to the next same page. Thirdly, every page in the frequent-page table contains three fields: the name of the page, the weight range and a link to the first node in the tree which denotes this page. The following algorithm to build the FSTP-tree:

**Algorithm 1 (FSTP-tree Building: Building a FSTP-tree of the SDB)**

**Input:** A session database SDB, weights of pages and a minimum support

**Output:** corresponding FSTP-tree

**Method:**

1) Scan the whole SDB and find frequent pages from SDB based on support and weight range assign to the page. Here we add only those pages that come under the weight range and contribute to the support and those not come in given range consider as outlier and not contribute to support.
2) Create the root of the FSTP-tree, and label it NULL.
3) Scan the whole SDB for the second time. For each session in the SDB, we only preserve the pages which are frequent and have a weight in given weight range, and hold the traversal sequences of pages. The different branches of same prefix can be merged.

3.3 FSTPMW Algorithm

The divide-and-conquer strategy is used for finding frequent sequential traversal patterns.

To handle the ordered problem, the FSTPMW uses a merging method. Each frequent ordered pattern whose first page is P1 must be contained in one or more session. The merging process in fact is rebuilding a smaller FSTP-tree. This time, the relative sessions all contains P1 as the first web page.

The complete algorithm given as:

**Algorithm 2 (FSTPMW: Mining frequent sequential traversal pattern)**

**Input:** FSTP-tree

**Output:** frequent sequential traversal pattern

**Method:** call FSTPMW (Weight range for each page, support, Minimum & Maximum Weight Range)

**Procedure** FSTPMW (FSTPtreeRootNode node,String prefix )

```java
{    for each node x in the corresponding page head table do
        if x.support less than minimum support then
            calculate the average weight of prefix
            if minimum weight<=average weight<=maximum weight
            {output prefix;}
            return;
        else if i.subs.count == 0 then
            prefix = prefix + i.content;
            calculate the average weight of prefix
            if minimum weight<=average weight<=maximum weight
            {output prefix;}
            return;
        else
            call CombineTree(i);
        for each node j in i.subs do
```
call FSTPMW (j, prefix+i);
end for
end if
end for

4. Analysis and Performance evaluation

In this section, we present our performance study over various datasets. We report our experimental results on the performance of FSTPMW in comparison with a recently developed algorithm; WSpan [26], which is the fastest algorithm for mining sequential patterns. The main purpose of this experiment is to demonstrate how effectively the sequential traversal patterns with weight constraint can be generated by incorporating a weight page, weight of sequence with a support. First, we show how the number of sequential traversal patterns can be adjusted through user assign weights, the efficiency in terms of runtime of the FSTPMW algorithm, and the quality of sequential traversal patterns. Second, we show that FSTPMW has good scalability against the number of sequence transactions in the datasets.

4.1 Environmental results. Comparison of FSTPMW and WSpan

In this performance test, we focused on the efficiency of using a weight range. Our experiment shows that in most cases, FSTPMW outperforms WSpan. First, we evaluate the performance on the kosarak dataset.

4.2 Further extension

FSTPMW basically focuses on sequential pattern mining with weight constraint uses a weight range to adjust the number of sequential traversal patterns. Frequent sequential traversal pattern mining can be extended by considering levels of support and/or weight of sequential traversal patterns. There are many areas in which items have different importance and patterns with a similar level of support and/or weight are more meaningful. For example, the concept of strong support and/or weight affinity can be applied in DNA analysis. We can give importance to specific DNA patterns and find interesting DNA patterns. Our algorithm can be extended by considering the levels of support and/or weight of sequential traversal patterns. By not only giving a balance between the two measures of support and/or weight, but also considering both support and/or weight affinity between items within patterns, more valuable sequential traversal patterns can be generated.

5. Conclusion

Many studies exist on mining sequential frequent patterns. One of the main limitations of the traditional approach for mining sequential traversal patterns is that all items are
treated uniformly, while each page of web site has different importance. Moreover, previous sequential traversal pattern mining generates a very large number of subsequence as the minimum support becomes lower. In this paper, we developed FSTPMW which focused on frequent sequential traversal pattern mining with weight constraint. A weight range is used to adjust the number of sequential patterns. The extensive performance analysis shows that FSTPMW is efficient and scalable in mining sequential traversal patterns.

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