Efficient Searches for Similar Subsequences of Different Lengths in Sequence Databases *

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Abstract

We propose an indexing technique for fast retrieval of similar s ubsequences us ing t ime w arping di stances. A time warping d istance is a more suitable similarity measure t han t he Eucl idean di stance in many applications, where sequences may be of different lengths or different sampling rates. Our indexing technique uses a disk-based suffix tree as an index s tructure and employs lower-bound distance functions to filter out dis similar subsequences without false dismissals. To make the index structure compact and thus acceler ate the query processing, we convert sequences of continuous values to sequences of discrete values via a categorization method and store only a s ubset of suffixes whose first values are different from their p receding values. The exp erimental results reveal that our proposed technique can be a few orders of magnitude faster than sequential scanning.

1. Introduction

Similarity searches in sequence databases are important in many application d omains, such as inf ormation retrieval, d ata m ining, a nd c lustering. D etecting sto cks that have similar gro wth p atterns and f inding p atients whose lung lesions have similar evolution c haracteristics are a few ex amples of sim ilarity qu eries. Although sequential scanning can be used to answer these queries, it may require an en ormous processin g tim e ov er larg e sequence databases. Recently, several indexing techniques [1,5,10,22] have been proposed to speed up the processing of similarity queries.

Most of the previous techniques [1,10,22] for similarity searches use the Euclidean distance metric as a sim ilarity measure. However, in many applications, the sampling rates and the lengths of sequences may be different, making it d ifficult or im possible to use the Euclidean distance as a similarity measure. In the area of speech

recognition [15], this problem has been approached using a similarity m easure, called a tim e w arping distance [3,15], which allows sequences to be stretched or compressed along the time axis. Under time warping, any element of a sequ ence can be m atched to on e or m ore neighboring elements of another sequence. As an example [16]. let us consider two sequences, S_1 <20,20,21,21,20,20,23,23> and S ₂ = <20,21,20,23>where the sequence S_1 is the closing price of a stock taken every day and S₂ is the closing price of another stock taken e very o ther d ay. S₁ and S₂ cannot be compared directly because the sequence S_1 is longer than S_2 . The Euclidean distance betw een S₂ and any subsequence of length four of S_1 is greater than 1.41. However, if we duplicate every element of S2 using time warping, we find that the two sequences are identical.

In the matching of similar sequences, it is important to prevent the occu rrence of *false d ismissals* [1]. A false dismissal occurs when a sequ ence sim ilar to a query sequence is n ot in cluded in the an swer set. Indexing techniques th at assume the *triangular inequality* may produce f alse dismissals when the distance function not satisfying the triangular inequality is use d as a similarity measure [22]. Un fortunately, a time warping distance does not satisfy the triangular ine quality, which c an b e simply proved by a counter example [22]. T his propert y makes spatial access m ethods b ased on the triangular inequality unsuitable for similarity searches with a time warping distance.

In the area of string matching, a suffix tree [17] has been extensively used as an index structure to find the substrings that are exactly matched to the given q uery string. A suffix tree may be a good candidate for an index structure with a time warping distance because it does n ot assume an yg eometry or an yu nderlying distan ce functions. However, for a suffix tree to be used as an index structure f or sim ilarity search es, th e f ollowing problems h ave t o be addres sed: 1) A suffix tree is designed to find the exactly matched substrings. Its exact search algorithm needs to be extended to f ind similarly matched subsequences. 2) In general, a suffix tree is built

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from sequences whose elem ents tak e th e v alues f rom finite alphabets. However, sequences we consider in this paper are comprised of elem ents of con tinuous real values. A systematic method to co nvert co ntinuous element values to discrete values is required.

In this paper we propose a new indexing technique for the fast retrieval o f sim ilar sub sequences o f d ifferent lengths or different sampling rates. Our technique uses a time warping distance as a sim ilarity measure and a diskbased suffix tree as an index structure. To make the index structure com pact, we con vert sequences of con tinuous values to sequences of discrete values via a categorization method and store only a subset of suffixes whose first values are dif ferent f rom th eir im mediately preceding values. When the query sequence, Q, is given, a suffix tree is traversed and time warping distances between Q and subsequences con tained in a su ffix tree are computed. Because subsequences con tained in a su ffix tree are of discrete values, their exact distances from Q cannot be obtained. In stead, low er-bound distan ce functions are employed to estimate the exact distance; so our proposed technique guarantees no false dismissals.

This paper is organized as f ollows. In Section 2 we provide a brief overview of the related work on sequence matching problems. In Section 3, we give the definition and the property of a tim e w arping distance. Section 4 introduces the construction method and the similarity search algorithm of a disk-based suffix tree. We apply the ideas of a categorization and a sparse suffix tree in Section 5 and Section 6, respectively. Experimental results are given in Section 7.

2. Related work

Several approaches f or f ast retrieval o f similar sequences have recently been proposed. In [1], sequences are converted in to the f requency domain by a Discrete Fourier Transform and are subsequently mapped into multi-dimensional points that are m anaged by an R^{*}-tree; this technique w as extend ed to lo cate similar subsequences [10]. Since the approaches of [1,10] use the Euclidean distance metric as a sim ilarity m easure, sequences of different lengths or different sampling rates cannot be matched.

Sequence m atching that allo ws transf ormations is proposed in [11,16]. In [11], s equences are grouped into equivalent classes accordin g to the eir n ormal f orms. Though normal forms are invariant to shape-based transformations such as scaling and shifting, they do not handle the compressions or the stretches of element values along the time axis. The authors of [16] propose a class of transformations that c an be used in a query language to express sim ilarity with an R- tree index. They handle moving average and global time scaling, but not time warping.

The access m ethods of [5,14,21,22] perm it the matching of sequences of different lengths. [5] presents a modified version of an edit distance, considering two sequences matching if a majority of elements match. This technique is extend ed to the m atching o f m ultidimensional sequences in [2 1]. In [2 2], a tim e w arping distance is u sed as a sim ilarity measure with a two-step filtering process: a FastMap in dex filter proceeded by a lower-bound d istance f ilter. T he underlying index structures of [5,21,22] a re b ased o n the tr iangular inequality. The authors of [14] introduce an aligned subsequence matching with a tim e w arping distance. Whereas their approaches are u seful f or lon g data sequences, subsequences not starting or ending at segment boundaries cannot be found.

Similarity m atching bas ed on s hapes of s equences i s proposed in [2,19]. [2] dem onstrates a s hape def inition language (SDL) a nd p rovides a n ind ex str ucture fo r speeding up the ex ecution of SDL qu eries. In [19], th e authors intro duce the no tion of generalized approximate queries that specify the general shapes of data h istories. Whereas both approaches m ay h andle th e v ariations of element values on the time axis, they cannot be used for applications that care about specific element values.

There are also sev eral approach es f or m atching of biological sequences. [4] proposes t o u se a di sk-based suffix tree f or solv ing the sequence alignment problem, and [20] addres ses the problem of discovering patterns in protein d atabases with the sim ilarity measure of a string edit distance. W hile w e f ocus on the sequences of continuous n umeric v alues, th e approach es of [4,20] center on the sequences of ch aracters. Fu rthermore, th e algorithm of [20] u ses a m ain-memory bas ed suffix tree, making it infeasible for a large sequence set.

3. Time warping distance

In general, finding a sim ilarity measure for sequences is not easy because sequences that are qualitatively the same may be quantitatively different. First, the sequences may be of different lengths, making it difficult or impossible to embed the sequences in a metric space and use the Euclidean d istance to d etermine similarity. Second, the sampling rates of sequences may be different: one sequence may be sampled every minute while another sequence is sampled every other minute. Such differences in rates make similarity measures such as cross-correlation unusable.

In this paper, we use a time warping (TW) similarity measure [3,15] that allows sequences to be stretched or compressed along the time axis. TW is a generalization of classical algorithms for comparing discrete sequences to sequences of continuous values, and is used extensively in matching of v oice, au dio an d medical signals (electrocardiograms). T o f ind th e m inimum dif ference between two sequences, T W m aps each element of a sequence to one or more neighboring elements of another sequence. Let us no w give the f ormal d efinition [15] of the time warping distance.

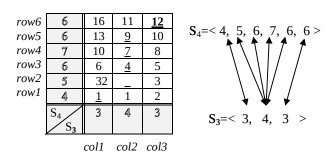
Definition 1. Given any two non-null sequences, S_i and S_j , the time warping distance, $D_{tw}()$, is defined as follows :

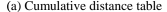
$$\begin{split} D_{tw}(S_i, S_j) &= D_{base}(S_i[1], S_j[1]) + \min \begin{cases} D_{tw}(S_i, S_j[2:-]) \\ D_{tw}(S_i[2:-], S_j) \\ D_{tw}(S_i[2:-], S_j[2:-]) \end{cases} \\ \end{split}$$

In above definition, $S_i[p]$ represents the p^{th} element of S_i and $S_i[p:q]$ denotes the subsequence of S_i including elements in positions p through q. We use the notation $S_i[p:-]$ for the suffix of S_i starting from the p^{th} element. That is, $S_i[p:-]$ is identical to $S_i[p:|S_i|]$ where $|S_i|$ is the length of S_i . $D_{base}()$ on two numeric v alues can be any distance function, but we assume that it is defined as the city-block distance. $D_{tw}(S_i,S_j)$ can be calculated efficiently using a dynamic programming technique [3] bas ed on the recurrence relation $\gamma_{tw}(x, y)$.

Definition 2. Given any two non-null sequences, S_i and S_j , the recurrence relation $\gamma_{tw}(x, y)$ (x =1,2,..., $|S_i|$, y =1,2,..., $|S_j|$) that builds the cumulative time warping distance table for S_i and S_j is defined as follows:

 $\gamma_{tw}(x, y) = D_{base}(S_i[x], S_j[y]) + \min \begin{cases} \gamma_{tw}(x, y-1) \\ \gamma_{tw}(x-1, y) \\ \gamma_{tw}(x-1, y-1) \end{cases}$





(b) Mapping of elements

Figure 1. Time warping distance for $S_3 = \langle 3,4,3 \rangle$ and $S_4 = \langle 4,5,6,7,6,6 \rangle$

The d ynamic p rogramming algo rithm [3] fills in the table of cumulative distances as the computation proceeds. This computation has complexity $O(|S_i||S_j|)$. The final cumulative distance, $\gamma_{tw}(|S_i|,|S_j|)$, is the minimum distance

between S_i and S_j, and the matching of elements can be traced backward in the table by choosing the previous cells with the lowest cumulative distance. Figure 1 shows the cumulative distance table f or two sequences, S₃ = $\langle 3,4,3 \rangle$ and S₄ = $\langle 4,5,6,7,6,6 \rangle$ and the mapping of elements that generates the minimum distance.

By reading the last column of each row of the cumulative distance table, we get the distance between S_i and any prefix of S_j . That is, the distance between S_i and $S_j[1:q]$ (q=1,2,...,| S_j |) is obtained from the last column of the qth row. In the above example, $D_{tw}(S_3, S_4[1:4])$ is 8, as seen in the last column of the row 4. Thus, determination as to whether the time warping distance of two sequences is greater th an a distan ce-threshold ε does not require building the entire cumulative distance table, as proven by the following theorem.

Theorem 1. If all columns of the last row of the cumulative distance table h ave v alues g reater than a distance-threshold ε , adding more rows on this table does not yield the new values less than or equal to ε . **Proof.** The proof is given in [13].

Let us look at the table shown in Figure 1. If ε is 3, after inspecting the row 3, we can d etermine that the distance between S₃ and S₄ is greater than ε because all columns of the row 3 h ave v alues g reater th an 3. Therefore, we do not have to fill the remaining three rows. In subsequent sections, we use Theorem 1 to reduce the search space of an index structure.

4. Similarity search using a suffix tree

In this section, we propose to use a suffix tree (ST) as an in dex stru cture f or sim ilarity search es w ith a time warping di stance. Bef ore des cribing t he m ethods f or constructing and searching a su ffix tree, w e present th e definition and the internal structure of a suffix tree.

A trie is an indexing structure used for indexing sets of keywords o f vary ing sizes. A suf fix trie [17] is a trie whose set of keywords comprises the suffixes o f a single sequence. Nodes w ith a sin gle outgoing edge can be collapsed, yielding the structure known as the suffix tree [17]. A suffix tree is generalized [4,20] to allow multiple sequences to be stored in the same tree. Each suffix of a sequence is represented by a leaf node. Precisely, the suffix St[p:-] is expressed by a leaf node labeled with (t,p). The edges are labeled with subsequences such that the concatenation of the edge labels on the path from the root to the leaf (t,p) becomes St[p:-]. The concatenation of the edge labels on the path from the root to the internal node, N_i, represents the longest common prefix of the suffixes represented by the leaf nodes u nder N_i. We use the

notation lab el(N_i,N_j) for the con catenated labels on the path f rom N_i to N_j. Figure 2 shows the suffix tree constructed from two sequences, $S_5 = \langle 4,5,6,7,6,6 \rangle$ and $S_6 = \langle 4,6,7,8 \rangle$, where '\$' is used as an end marker of a suffix.

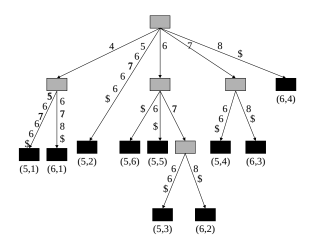


Figure 2. Suffix tree from $S_s = <4,5,6,7,6,6>$ and $S_s = <4,6,7,8>$

4.1. Index construction

A suffix tree for multiple sequences can be constructed by addin g a special sequence separator symbol to the alphabet. The sequences to be in cluded in the tree are concatenated, separated from each other by this separator symbol. Then, the ordinary suffix tree algorithm is applied to the con catenated sequence. The suffix tree created using this process has to be kept in main memory during construction. Therefore, this ap proach is not realistic to a large sequence set.

To remedy the problem, we use an incremental diskbased s uffix t ree construction m ethod proposed in [4]. Two suffix trees, representing two disjoint sets of sequences, are merged to produce a single suffix tree by pre-order traversal of both suffix trees and combining the paths corres ponding t o common subsequences. A suffix tree for a large set of sequences can be constructed by performing a series of binary m erges of s uffix t rees of increasing size. The merge operation of two suffix trees can s upport di sk-based representations in l imited main memory.

Two suffix t rees for S_i and S_j are merged with complexity O($|S_i|+|S_j|$), hence the suffix tree f or M sequences is constructed with complexity O(M \overline{L}) where \overline{L} is the average length of M sequences. The total number of nodes in a suffix tree is constrained due to two factors: 1) there are O(M \overline{L}) leaf nodes and 2) the degree of any internal node is at least 2. T herefore, the m aximum number of n odes and ov erall space requ irement of the suffix tree is linear in $M\overline{L}$ [17].

4.2. Search algorithm: SimSearch-ST

A suffix tree (ST) is a u seful index structure to locate subsequences th at are ex actly m atched to a query sequence Q. T o find exactly matched subsequences, Q is traversed from the root of the tree and trav ersal is terminated when the end of Q is reached or a node is reached bey ond which f urther trav ersal is n ot possible. Exact searches are performed in O(|Q|). Even though the exact matching algorithm of a su ffix tree is sim ple and fast, it cannot be directly applied to the p roblem we are going to solve in this paper.

Problem Defin ition: Given M sequences $S_1, S_2, ..., S_M$ of arbitrary lengths, a qu ery sequence Q and a u ser-given distance-threshold ε , find the subsequences $S_i[p:q]$ (i = 1,2,...,M) whose time warping distances from Q are less than or equal to ε .

Our proposed similarity search algorithm SimSearch-ST is given in Algorithm 1. The search starts from the root of a suffix tree and continues the depth-first traversal until all subsequences whose time warping distances are less than or equal to ε are found.

Input : Root Node R, Q, ε Output : answerSet
cumDistTable ← NULL; answerSet ← Filter-ST (R, Q, ɛ , cumDistTable);
return answerSet;

Algorithm 1. SimSearch-ST

The actual f iltering p rocess is executed in Filter-ST shown in Algorithm 2. When Filter-ST visits a node N, it inspects each child node CN_i to find a new answer and to determine whether further depth traversal is needed or not. For simpler explanation, we assume that the edge between two nodes, N and CN_i , is labeled with a single value.

To find a new answer, Filter-ST builds a cum ulative distance table for Q and label(N, CN_i). If N is a root node, the table is built from the bottom. Otherwise, the table is constructed by augmenting a new row on the table T that has been accumulated from the root to N. The function AddRow(T, Q, 1 abel(N, CN_i), $D_{tw}()$) builds a n ew cumulative distance table, u sing the distance function $D_{tw}()$, by augmenting a new row corresponding to label(N, CN_i) on T. S uppose that the rth row is the newly added row. If the last co lumn of the rth row has the value less

than or equal to $\boldsymbol{\epsilon}$, label(GetRoot(CN_i), CN_i)) is inserted into the answer set.

To determine if further depth traversal is needed, Filter-ST uses Theorem 1. If at least one column of the r^{th} row has a v alue n ot greater th an ε , the search continues down the tree to find more answers. Otherwise, the search moves to the next child of N.

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\begin{array}{ll} \textbf{Input} & : \text{Node N, Q, } \boldsymbol{\epsilon}, \text{Cumulative Distance Table T} \\ \textbf{Output} : answerSet \\ \textbf{answerSet} \leftarrow \{\}; \\ \text{CN} \leftarrow \text{GetChildren(N)} \\ \textbf{for } i \leftarrow 1 \text{ to } |\text{CN}| \text{ do } \{ \\ & \text{CT}_i \leftarrow \text{AddRow(T, Q, label(N, \text{CN}_i), D_{tw}())}; \\ & \text{Let } dist \text{ be the last column value of the new row;} \\ & \text{Let } mDist \text{ be the minimum column value of the new row;} \\ & \textbf{if } dist \leq \boldsymbol{\epsilon}, \text{ insert label(GetRoot(CN_i), N_i) into answerSet;} \\ & \textbf{if } mDist \leq \boldsymbol{\epsilon}, \\ & \text{answerSet} \leftarrow \text{answerSet} \cup \text{Filter-ST(CN}_i, \text{Q, } \boldsymbol{\epsilon}, \text{CT}_i); \\ \} \\ \textbf{return answerSet;} \end{array}
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Algorithm 2. Filter-ST

4.3. Algorithm analysis

Before analyzing the complexity of SimSearch-ST, let us examine the complexity of sequential scanning. Sequential scan ning reads each sequence and builds as many cumulative distance tables as the number of suffixes contained in the sequence. The complexity of building a cumulative distance table f or the query sequence Q and the suffix of length L is O(L|Q|). For M sequences whose average length is \overline{L} , there are M \overline{L} suffixes and their average length is $(\overline{L}+1)/2$. Therefore, the com plexity of sequential scanning is $O(M\overline{L}^2|Q|)$.

SimSearch-ST is computationally less expensive than sequential scan ning du e to bran ch-pruning (based on Theorem 1) and sharing cumulative distance tables for all suffixes that have common prefixes. Thus, the complexity of SimSearch-ST is $O(\frac{M \overline{L}^2 |Q|}{R_d R_p})$, where R_d (≥ 1) is the reduction f actor du e to sh aring th e cumulative distance tables, and R_p (≥ 1) is the reduction factor gained from the branch-pruning. R_d grows as the length and the number of common prefixes of the suffixes contained in a suffix tree increase. G iven k suffixe s, $\alpha_1, \alpha_2, ..., \alpha_k$, whose f irst t elements are the same, the construction of k cumulative distance tables requires the computation of $|Q||\alpha_1| + |Q||\alpha_2|$ $+ ... + |Q||\alpha_k|$ cells. Ho wever, it is red uced to t|Q| + $|Q|(|\alpha_1|-t) + |Q|(|\alpha_2|-t) + ... + |Q|(|\alpha_k|-t)$ if the cum ulative distance table for Q and the common prefix of length t is shared by k suffixes. In this case, R_d can be expressed as $R_d = (|\alpha_1|+|\alpha_2|+...+|\alpha_k|) / ((|\alpha_1|+|\alpha_2|+...+|\alpha_k|) - t(k-1)).$

While R_d is determined by the distribution of element values, R_p is decided by the number of answers required by a user. That is, R_p increases as the distance-threshold ε decreases. If ε is so small that just one or two subsequences may be an swers, only the topmost part of a suffix tree may be v isited during the query processing. In another ex treme case w here ε is large enough for all subsequences to be answers, all nodes of a suffix tree need to be visited, thus making $R_p = 1$. In the worst case where there is no common subsequence and the branch-pruning cannot help, both values of R_d and R_p are 1, and therefore the complexity of SimSearch-ST becomes the same as that of sequential scanning.

5. Similarity search using categorization

In this section we introduce the co ncept o f categorization to decrease the n umber of v alues that elements can take and thus increase the length and the number of common subsequences. A s explained in the previous section, if the length and the number of common subsequences increase, the in dex size an d th e qu ery processing time are reduced. T o g et th e categ orized representations of element values, we first generate the set of categories and determine their ranges. Then, we convert every element v alue to the symbol of the corresponding category. For example, given two categories $C_1 = [0.1, 3.9]$ and C₂ = [4.0,10.0], S₇ = $\langle 5.27, 2.56, 3.85 \rangle$ is transformed to $CS_7 = \langle C_2, C_1, C_1 \rangle$ where CS_7 denotes the converted sequence of S_7 . Thus, sequences of continuous values are converted to sequences of discrete symbols.

5.1. Categorization method

In this work, the following two categorization methods have been chosen and experimented for their simple implementations, albeit oth er categ orization approaches like the type abstraction hierarchy (TAH) [6] and the kmeans algorithm may also be used to categorize element values.

The first method is an equal-length categorization. As the name im plies, all the categories have equal interval length (MAX-MIN) / c. Here, MIN is the smallest element value found in the set of sequences, MAX is the largest element value found in the set of sequences, and c is the number of categories given as the input parameter to the categorization algorithm. This categorization approach is simple and f ast, b ut lo ses information about value and frequency distributions of the sequences. The secon d m ethod is a maximum-entropy categorization. T he en tropy [18] of a categorization is defined as $H(C) = -\sum_{i=1}^{c} P(C_i) \log P(C_i)$ where $P(C_i)$ is the probability that an elem ent is included in the i th category. T o minimize the loss of information about the sequences, this categorization m ethod decides th e category boundaries that g enerate the maximum entropy value. The boundaries can be determined easily by making all categories include the same number of elements $(P(C_1) = P(C_2) = ... = P(C_c))$.

It is not easy to determine the number of categories: too many categ ories do n ot h elp m uch to increase the number of common subsequences, but likewise, too few categories do n ot h elp m uch to redu ce the query processing time because of the decreased filtering rate of the index. A simple strategy is to do many experiments on the s et of s equences and det ermine t he best number of categories using the cost function $W_tC_t + W_sC_s$. Here, C_t and C_s are costs for processing the query and storing the index, respectiv ely, an d W t and W s are their relative weights. T he d etermination o f these w eights is application-dependent.

5.2. Index construction

After converting element values into discrete symbols, we build a suffix tree from the set of converted sequences. We denote the resultant tree ST_C . ST_C is constructed using the same construction a lgorithm use d fo r a n o rdinary suffix tree.

5.3. A modified distance function: **D**_{tw-lb}()

Whereas the edges of a su ffix tree are labeled w ith numeric values, the edges of ST_C are labeled with discrete symbols. As a resu lt, th e ex act tim e w arping distan ce between a query sequ ence of n umeric v alues and an y subsequence con tained in ST_C cannot be computed. Therefore, we introduce the new distance function $D_{tw-lb}()$.

Definition 3. Given two non-null sequences, S_i and S_j , the distance f unction $D_{tw-lb}(S_i, CS_j)$ that returns the lower-bound distance of $D_{tw}(S_i, S_j)$ is defined as follows:

$$\begin{split} D_{tw-lb}(S_{is}CS_{j}) &= D_{base-lb}(S_{i}[1],CS_{j}[1]) + \min \begin{cases} D_{tw}(S_{i},CS_{j}[2:-]) \\ D_{tw}(S_{i}[2:-],CS_{j}) \\ D_{tw}(S_{i}[2:-],CS_{j}[2:-]) \end{cases} \\ \\ D_{base-lb}(a,B) &= \begin{cases} 0 & (\text{if } B.lb \leq a \leq B.ub) \\ a - B.ub & (\text{if } a > B.ub) \\ B.lb - a & (\text{if } a < B.lb) \end{cases} \end{split}$$

where 'a' is the numeric value corresponding to $S_i[1]$ and

'B' is the category symbol corresponding to $CS_i[1]$.

In the definition of D $_{base-lb}$ (), B.lb and B .ub are the minimum and the maximum element values, respectively, found in the category B. As shown in Figure 3, D $_{base-lb}$ (a, B) returns the possible minimum distance between a and B.

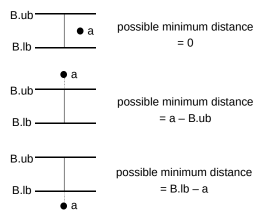


Figure 3. Minimum distance between a and B

To prevent false dismissals, the distance returned from $D_{tw-lb}(S_i, CS_j)$ should always be less than o r equal to the distance computed by $D_{tw}(S_i, S_j)$. Theorem 2 states this fact.

Theorem 2. For any two non-null sequences, S_i and S_j , the following inequality holds.

 $D_{tw-lb}(S_i, CS_j) \le D_{tw}(S_i, S_j)$ **Proof.** The proof is given in [13]. ■

5.4. Search algorithm: SimSearch-ST_c

 $\label{eq:linear_state} \begin{array}{l} \mbox{Input} : \mbox{Root Node R, Q, ϵ} \\ \mbox{Output} : \mbox{answerSet} \\ \mbox{cumDistTable} \leftarrow \mbox{NULL}; \\ \mbox{candidateSet} \leftarrow \mbox{null} \mbox{NULL}; \\ \mbox{candidateSet} \leftarrow \mbox{Filter-ST}_{C} (R, Q, ϵ, \mbox{cumDistTable}); \\ \mbox{answerSet} \leftarrow \mbox{PostProcess(candidateSet, Q, ϵ)} \\ \mbox{return} \mbox{answerSet}; \end{array}$

Algorithm 3. SimSearch-ST $_{c}$

The algorithm SimSearch-ST needs to be modified to reflect the categorized representation of element v alues. Our proposed search algorithm SimSearch-ST_c is shown in Algorithm 3. Note that element v alues of a query sequence are not converted to discrete symbols.

To f ind th e can didate su bsequences w hose lower-

bound distances to the query sequence Q are w ithin ε , Filter-ST_c is called recursively. Filter-ST_c is the same as Filter-ST except that the former uses D_{tw-lb} () to build the cumulative distance table while the latter uses D_{tw} (). Since the lower-bound d istance is used f or f iltering, the subsequences whose exact tim e w arping distan ces are larger than ε may be included in the candidate answer set. These subsequences are called *false a larms*. For each answer con tained in the can didate an swer set, the algorithm PostProcess retrieves the actual subsequences and computes their exact time warping distances. Finally, the subsequences whose actual time warping distances are not larger than $\boldsymbol{\varepsilon}$ are returned as answers. Algorithms Filter-ST_c and PostProcess are omitted due to space limitations.

5.5. Algorithm analysis

The complexity of $SimSearch-ST_c$ is represented as $O(\frac{M \overline{L}^2 |Q|}{B d B n} + n \overline{L} |Q|)$ where n is the number of

subsequences requiring the post-processing. Hen ce, th e left expression represents the time f or f iltering and the right expression represents the time for post-processing. SimSearch-ST, SimSearch-ST_c has Compared to performance improvements due to a larger value of R_d, despite the extra time for post-processing

6. Similarity search using a sparse suffix tree

A suffix tree th at stores on ly a su bset of suffixes is called a sparse su ffix tree [12]. Sin ce the size of a su ffix tree is linear in the number of leaves, a sparse su ffix tree is smaller than an o riginal suffix tree. Suffixes inserted into a tree are called stored suffixes, and suffixes no t inserted into a tree are called non-stored suffixes. The reduction of the index size by storing on ly a subset of suffixes is measured by the compaction ratio $r (0 \le r < 1)$ that is defined as r = (the number of non-stored suffixes) /(the number of suffixes). In this section, we propose to use a sparse suffix tree to f urther reduce the index size and accelerate the query processing.

6.1. Index construction

Similar to ST_{C} , a sparse suffix tree is built from the set of categ orized sequ ences. How ever, unlike ST_c, only suffixes w hose fi rst va lues a re different from their immediately preceding values are stored in a sparse suffix tree. That is, the suffix $CS_i[p:-]$ is stored only if $CS_i[p] \neq$ $CS_{i}[p-1]$. For example, for $CS_{8} = \langle C_{1}, C_{1}, C_{1}, C_{3}, C_{2}, C_{2} \rangle$, only the three suf fixes (CS₈[1:-], CS₈[4:-], and CS₈[5:-])

are stored in a sparse su ffix tree. We denote the resultant tree SST_C.

6.2. A modified distance function: D_{tw-lb2}()

Suppose that we have the cumulative distance table for S_i and CS_j where S_i and CS_j are located along the x-axis and the y-axis, respectively. While we can get the distance between S_i and any *prefix* of CS_i by reading the last column of each row, there is no direct way to compute the distance betw een S_i and any *suffix* of CS_i except by building a new table. However, if the first N elements of CS_i have the same value, we can obtain the lower-bound distance of D_{tw-lb}(S_i, CS_j[p:-]) (p=2,3,...,N) using a new distance function, D_{tw-lb2}(S_i, CS_i[p:-]).

Definition 4. For any two non-null sequences, S_i and CS_i , if the first N elements of CS_j have the same value, then the distance f unction D $_{tw-lb2}(S_i, CS_j[p:-])$ (p=2,3,...,N) that returns the lower-bound distance of $D_{tw-lb}(S_i, CS_i[p:-])$ is defined as follows:

$$\mathbf{D}_{\text{tw-lb2}}(\mathbf{S}_i, \mathbf{C}\mathbf{S}_j[p:-]) = \mathbf{D}_{\text{tw-lb}}(\mathbf{S}_i, \mathbf{C}\mathbf{S}_j) - (p-1) * \mathbf{D}_{\text{base-lb}}(\mathbf{S}_i[1], \mathbf{C}\mathbf{S}_j[1])$$

If we know the value of $D_{tw-lb}(S_i, CS_j)$, then $D_{tw-lb2}(S_i, CS_j)$ $CS_i[p:-]$) can be computed with complexity O(1). The distance returned from $D_{tw-lb2}(S_i, CS_i[p:-])$ is always less than or equal to D_{tw-lb}(S_i, CS_i[p:-]). The following theorem states this fact.

Theorem 3. For any two non-null sequences, S_i and S_j, if the first N elements of CS_i have the same value, then the following inequality holds for p = 2,3,...N:

$$D_{tw-lb2}(S_i, CS_j[p:-]) \le D_{tw-lb}(S_i, CS_j[p:-]) \le D_{tw}(S_i, S_j[p:-])$$

Proof. The proof is given in [13].

6.3. Search algorithm: SimSearch-SST_c

The algorithm SimSearch-ST_c needs to be modified to reflect the fact that there are some suffixes not stored in the index. If SimSearch-ST_c is applied to a SST_C without modification, the sub sequences contained in the non-stored suffixes may not be included in the answer set even if similar to a targ et query sequence. Therefore, the steps of f inding an d processin g th e su bsequences contained in the non-stored suffixes need to be added to the SimSearch-ST_c.

The proposed algorithm SimSearch-SST_c includes the filtering step and the post-processing step. During the filtering step, D tw-lb() is used to calcu late distances between Q and the subsequences contained in the stored suffixes, and $D_{tw-lb2}()$ is used to compute distances between Q and the subsequences contained in the non-stored suffixes. During the post-processing, $D_{tw}()$ is applied to the subsequences in cluded in the candidate answer set. A detail description of the SimSearch-SST_c algorithm is in [13].

6.4. Algorithm analysis

The complexity of SimSearch-SST_c is represented as $O(\frac{(1-r)M\bar{L}^{2}|Q|}{R_{d}R_{o}} + rM\bar{L} + n\bar{L}|Q|)$ where n is the number

of subsequences requiring the post-processing, and r is the compaction ratio of a SST_c. Thus, $(1-r)M\overline{L}$ is the number of the stored suffixes, and $rM\overline{L}$ is the number of the non-stored suffixes. Compared w ith SimSearch-ST_c, SimSearch-SST_c reduces the query processing time by decreasing the number of cumulative distance tables generated during the tree traversal, at the cost of larger n.

7. Experimental results

To study the perf ormance of our sim ilarity search algorithms, we conducted se veral experiments on stock data sequences ex tracted f rom S&P 500 (http://biz.swcp.com/stocks/) and o n the artif icial d ata sequences. The s tock dat a w ere bas ed on t heir dai ly closing prices. A total of 545 s tock s equences was used with an av erage l ength of 232. T he ex pression for generating the artificial sequences was defined as $S_i[p] =$ $S_i[p-1] + Z_p$ where Z_p (p=1,2,...) are independent, identically distributed random variables. Twenty percent of the query sequences were extracted from the stocks whose av erage pri ces w ere bel ow \$30, 50% f rom t he stocks whose average prices were between \$30 and \$60, and 30% from the remaining stocks. The query sequences for the artificial sequences were obtained in a similar manner. The average length of the query sequences was set to 20. All experiments except for the scalability test in Section 7.3 w ere performed on both the stock and the artificial sequences.

7.1. Index siz e and query processing time w ith increasing number of categories

Table 1 shows the sizes of the proposed indices built from the stock sequences, where EL is the equal-length categorization an d ME is the maximum-entropy categorization. While the size of ST is not affected by the number of categories, ST_C and SST_C become larger as the number of categories increases. ST_C and SST_C are smaller than ST due to the in creased n umber of com mon subsequences, and SST_C is smaller than ST_C due to the decreased number of suffixes stored in the index.

Table 1. Index sizes with selected number of categories

# c	Index Size (Kbytes)				
categories	ST	ST _C		SST _C	
ies		EL	ME	EL	ME
10		5,360	10,534	262	<u>914</u>
20		7,982	15,879	8 50	2,355
40		12,362	26,069	2,685	7,108
80		18,817	41,288	3,985	18,317
120	158,512	26,888	51,942	7,657	28,842
160		32,860	59,927	11,620	37,922
200		37,837	66,357	15,416	45,449
250		43,413	72,937	20,326	5 3,535
300		48,087	78,297	24,905	60,345

Table 2 shows the average query processing times of the three proposed s imilarity s earch al gorithms with the average distan ce-tolerance of 30. On the w hole, as th e number of categories increases, the executions of SimSearch-ST_c and SimSearch-SST_c become faster. However, their executions slow down when the number of categories ex ceeds a certain th reshold. T his th reshold value m ay be u sed as the optimal number of categories. For example, 200 is the optimal number of categories for SimSearch-SST_c with EL and 120 is for SimSearch-SST_c with ME. Using similar-sized indices, SimSearch-SST_c based on ME yields better performance than SimSearch-SST_c based on EL. We obtained similar conclusions from experiments on the artificial sequences.

Table 2. Average query processing times with selected number of categories

# c	Average Query Processing Time (sec)				
categories	Sim Search- ST	$SimSearch-ST_c$		$SimSearch-SST_c$	
ies		EL	ME	EL	ME
10		241.94	84.09	215.73	75.53
20		122.63	35.57	122.75	30.90
40		54.89	25.88	49.61	20.65
80		30.57	21.05	25.90	18.40
120	5 5.30	26.03	20.93	21.30	20.80
160		23.08	21.60	19.13	23.49
200		21.42	22.41	<u>18.63</u>	26.53
250		21.19	23.67	19.08	30.49
300		20.65	25.04	19.55	34.15

7.2. Comparison with sequential scanning

Based on the results from Section 7.1, we chose MEbased S ST_c as our index structure and compared its similarity search algorithm with sequential scanning. Table 3 shows their average query processing times with increasing distance-threshold (ε) from 5 t o 50. A bout 50 answers were returned when $\varepsilon = 5$ and about 350,000 answers were reported when $\varepsilon = 50$. Here, SeqScan is sequential scanning and SimSearch-SST_c (k) represents the proposed algorithm with k categories. From Table 1, we know that SST_c with 10, 20, and 80 categories require about 50%, 100%, and 1,000% s paces of databas e size (1,896 Kbytes), respectively. Our proposed technique is up to 4.2 times faster with 10 categories, 11.1 times faster with 20 cat egories, an d 34.7 t imes f aster w ith 80 categories than the sequential scanning.

 Table 3. Comparison of sequential scanning and our algorithm with selected distance-threshold

dis threa	Query Processing Time (sec)				
distance- threshold(ε)	Seq Scan	Sim Search- SST(10)	Sim Search- SST(20)	Sim Search- SST(80)	
5	206.04	48.96	18.61	5.94	
10	210.48	54.63	21.24	9 .01	
20	217.14	71.31	27.18	14.15	
30	217.45	75.62	30.98	18.49	
40	218.13	79.85	34.89	22.71	
50	218.96	81.94	38.29	27.08	

7.3. Scalability study

To study the scalability of our approach, we compared the execution times of ME-based $SimSearch-SST_c$ with that of sequ ential scan ning by increasing the average length and the number of the artificial sequences. First, we increased the average length of the sequences from 200 to 1,000 while keeping the number of the sequences equal to 200. And we changed the number of sequences from 1000 to 10,000 while maintaining t he av erage l ength of sequences equal to 200. F or bot h ex periments, t he numbers of categories were chosen to make the index size smaller than the database size. A s shown in Figure 4 and Figure 5, the performance improvements of SimSearch-**SST**_c is maintained for both long sequences and large number of sequ ences. Note that the qu ery processin g times for both sequential scanning and SimSearch-SST_c increase qu adratically w ith respect to the average sequence length and linearly with respect to the number of sequences.

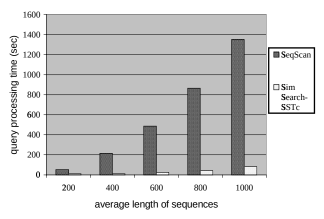


Figure 4. Comparison of sequential scanning and our algorithm with selected length of sequences

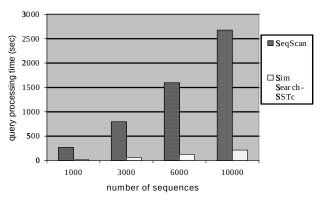


Figure 5. Comparison of sequential scanning and our algorithm with selected number of sequences

8. Conclusion

In this paper we have proposed an indexing method based on a di sk-based s uffix t ree, f or f ast ret rieval of similar su bsequences w ithout f alse dism issals. Because the sampling rates and the lengths of sequences may be different, t he propos ed m ethod u ses a t ime warping distance as a sim ilarity measure that allows stretching or compressing of sequ ences alon g th e time axis. Experiments on the stock and the artificial sequences have shown that our proposed technique can be a few orders of magnitude f aster than seq uential scanning. The contributions of our work are: 1) ex tending the exact matching algorithm of a suf fix tree to similarity searches with a tim e w arping similarity m easure, 2) ap plying the ideas of categorization and sparse suffix tree to make an index stru cture m ore com pact, an d 3) in troducing two lower-bound time warping distance functions, D_{tw-lb}() and $D_{tw-lb2}()$, to filter out dissimilar subsequences without false dismissals.

The index space can be reduced further if we know the minimum and maximum lengths of t he queries. U sing a warping w indow con straint [3], we can calcu late the minimum and maximum lengths of t he a nswers. The suffixes whose lengths are shorter than the minimum answer length need not be inserted into the index. For the suffixes whose lengths are longer than the maximum, only the prefixes whose lengths are equal to the maximum length need to be stored in the index.

The subsequences found by similarity searches can be used for p redictions, hy pothesis te sting, c lustering a nd rule discov ery. For ex ample, in the m edical domain, retrieved subsequences can be used for predicting the disease evolution patterns of a p atient; in the w eb environment, they can be used to discov er user web-site visiting patterns.

Our approach can be ex tended to the sequences of multivariate num eric values. Multivariate values are converted into m ulti-dimensional cells using m ulti-dimensional categorization methods such as m ultiple-attribute ty pe abstraction h ierarchy (MT AH) [6]. Then, the sam e in dex con struction and query processing techniques are applied to the set of converted sequences. We are currently working in this d irection for retrieving similar medical image subsequences [7,8].

9. References

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