

# Matrix Profile XX: Finding and Visualizing Time Series Motifs of All Lengths using the Matrix Profile

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**Abstract**— Many time series analytic tasks can be reduced to discovering and then reasoning about conserved structures, or *time series motifs*. Recently, the Matrix Profile has emerged as the state-of-the-art for finding time series motifs, allowing the community to efficiently find time series motifs in large datasets. The matrix profile reduced time series motif discovery to a process requiring a single parameter, the length of time series motifs we expect (or wish) to find. In many cases this is a reasonable limitation as the user may utilize out-of-band information or domain knowledge to set this parameter. However, in truly exploratory data mining, a poor choice of this parameter can result in failing to find unexpected and exploitable regularities in the data. In this work, we introduce the Pan Matrix Profile, a new data structure which contains the nearest neighbor information for *all* subsequences of *all* lengths. This data structure allows the first truly parameter-free motif discovery algorithm in the literature. The sheer volume of information produced by our representation may be overwhelming; thus, we also introduce a novel visualization tool called the motif-heatmap which allows the users to discover and reason about repeated structures at a glance. We demonstrate our ideas on a diverse set of domains including seismology, bioinformatics, transportation and biology.

**Keywords**—Time series, Motif discovery, Anomaly detection

## I. INTRODUCTION

In recent years, the Matrix Profile (MP) has emerged as a promising data structure to support time series data mining. The MP is a simple data structure that contains the nearest neighbor information for all subsequences of a given length in a time series. In the past three years it has been shown that the MP can be used to facilitate the discovery of motifs [19], discords (anomalies) [23], chains (evolving patterns) [22], shapelets [19], snippets [6], regimes [23], and more. However, we argue that the MP has a strong assumption that limits its practicality by requiring the user to specify the subsequence length ahead of time. A data scientist may have a good intuition as to what this subsequence length should be, based on their experience or a first principles model of the system being examined. However, in many cases, particularly for exploratory data mining, the user may have no idea as to the subsequence lengths at which patterns are conserved in the data necessitating the need for *variable-length* motif discovery.

Consider the one-hundred second excerpt of an EOG (Electro-oculogram; the movement of an eye) dataset from a 66-year old healthy male recorded during a sleep study shown in Fig. 1. Here we are tasked with identifying regions corresponding to the “blinking of the eye” in an attempt to remove these regions from a companion EEG dataset (not shown). Because eye blinks are not only unique to the individual but also sensitive to the sensor placement, we cannot use a single “one-size-fits-all”

template. But, given that blinks are typically well-conserved, at least during a single sleep session, we can perform motif discovery to identify an appropriate template. However, the suggested subsequence length for motif discovery is not readily apparent. We may attempt to rely on the current sleep study literature in which case [14] suggests “a duration of 1.5 to 2.5 seconds” as the subsequence length. In Fig 1.*bottom.left* we show that using 2.5 seconds does indeed discover a highly conserved motif that corresponds to an eye blink. Moreover, searching for more examples of this pattern in the full night of sleep data, we find hundreds of additional examples of this shape.

A sleep technician might very well be justified in terminating her search. However, as Fig 1.*bottom.right* shows, this time series has a second type of eye-blink artifact with a subsequence length of five seconds which may not have been considered by the sleep technician due to the high frequency of the 2.5 second motif. The fact that eye blinks can be polymorphic seems underappreciated, but [1] cautions EOG signals can have “*more than one category... classified by shape.*” Missing this second blink artifact would have drastically corrupted the downstream analytics performed on this dataset.

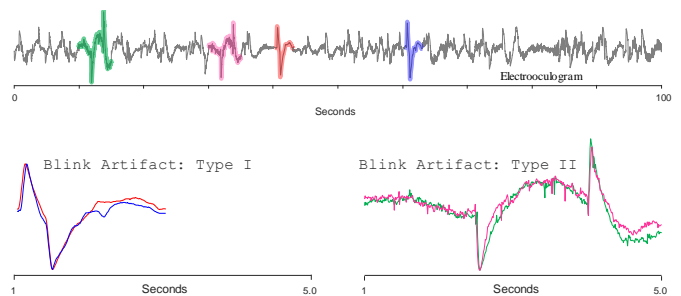


Fig. 1. *top*) One hundred seconds of EOG data. *bottom.left*) A search for the top 2.5-second motif reveals a highly conserved pattern eye-blink-artifact. *bottom.right*) A search for the top 5-second motif reveals another highly conserved pattern, that has no overlap with the first.

This very basic problem exemplified in an EOG dataset is ubiquitous in nearly all domains as the user’s choice limits what regularities can be found in the dataset. In the EOG dataset shown in Fig. 1, the two motif patterns differ by a factor of two; however, as this paper will show, other datasets may contain motifs which can differ by up to two orders of magnitude.

A possible, yet inelegant, solution to this problem is trial and error over different lengths. Beyond being frustratingly time consuming and awkward for the user, there is still a real danger of missing an interesting pattern. Though the definition of a time series motif is fairly robust to minor changes in length [11][12][23], there *will* be some length at which there is a “phase

change”, that is, the location of the motif will “jump” to a different place in the time series. For example, in Section III.A we introduce a dataset with a maximal motif for  $m_1 = 68$ , and a non-related/nonoverlapping maximal motif for  $m_2 = 610$ .

In this work we solve this motif-length sensitivity problem by introducing the Pan Matrix Profile (PMP), a data structure that contains *all* MP information of a time series with length  $n$  for all lengths in a fixed range  $r$ . In addition, we introduce SKIMP (Scalable **K**inetoscopic **M**atrix **P**rofile), an algorithm to compute the PMP with time complexity  $O(n^2r)$  and space complexity  $O(nr)$ . Though untenable for large datasets which require an *exact* solution, SKIMP is computed in an *anytime* fashion allowing for fast *approximate* solutions [24]. In almost all cases, running SKIMP to even one one-hundredth of its full convergence time will produce results that are almost indistinguishable from the final product.

Using SKIMP, we believe that all algorithms that exploit the MP could be made length-agnostic, that is to say, we can have length-agnostic chains [22], snippets, regimes, etc. However, for clarity and concreteness, in this work we confine our claims to motif and anomaly discovery and leave all other considerations for future work.

The practical application of SKIMP is in facilitating interactive time series analytics on practical problems in bioinformatics, seismology, medicine and industry.

The rest of this paper is organized as follows. In Section II we introduce the relevant notation, background material, and define the PMP data structure. Section III introduces a family of algorithms to compute the PMP and several algorithms to exploit and visualize it. We conduct an extensive empirical evaluation in Section IV. We defer a discussion of related work to V, so the readers intuitions for the issues at hand are more fully developed, before offering conclusions and directions for future work in Section VI.

## II. NOTATION AND BACKGROUND

### A. Time Series Notation

We begin by introducing all the necessary definitions, starting with the data type of interest, *time series*:

**Definition 1:** A time series  $T$  is a sequence of real-valued numbers  $t_i$ :  $T = t_1, t_2, \dots, t_n$  where  $n$  is the length of  $T$ :

We are typically interested not in *global*, but *local* properties of a time series. A local region of a time series is called a *subsequence*:

**Definition 2:** A subsequence  $T_{i,m}$  of a time series  $T$  is a continuous subset of the values from  $T$  of length  $m$  starting from position  $i$ . Formally,  $T_{i,m} = t_i, t_{i+1}, \dots, t_{i+m-1}$ , where  $1 \leq i \leq n-m+1$ .

Given a query subsequence  $T_{i,m}$  and a time series  $T$ , we can compute the distance between  $T_{i,m}$  and *all* the subsequences in  $T$  with length  $m$ . We call this a *distance profile*:

**Definition 3:** A *distance profile*  $D_i$  corresponding to query  $T_{i,m}$  and time series  $T$  is a vector of the Pearson correlation between a given query subsequence  $T_{i,m}$  and each subsequence in time series  $T$  with length  $m$ . Formally,  $D_i = [d_{i,1}, d_{i,2}, \dots, d_{i,n-m+1}]$ , where  $d_{i,j}$  ( $1 \leq j \leq n-m+1$ ) is the distance between  $T_{i,m}$  and  $T_{j,m}$ .

We assume that the distance is measured by Euclidean distance between z-normalized subsequences [23][19]. Once we obtain  $D_i$ , we can extract the nearest neighbor of  $T_{i,m}$  in  $T$ . Note that if the query  $T_{i,m}$  is a subsequence of  $T$ , the  $i^{\text{th}}$  location of distance profile  $D_i$  is zero (i.e.,  $d_{i,i} = 0$ ) and close to zero just to the left and right of  $i$ . This is called a *trivial match* in the literature (See Definition 7). Most of the community follow the suggestion in [3] to avoid such matches by ignoring an “exclusion” zone of length  $m/2$  before and after  $i$ , the location of the query [19].

We wish to find the nearest neighbor of every subsequence in  $T$ . The nearest neighbor information for subsequences with length  $m$  is stored in two meta time series, the *matrix profile*, and the *matrix profile index*:

**Definition 4:** A *matrix profile*  $P$  of time series  $T$  is a vector of the Euclidean distances between every subsequence of  $T_{i,m}$  and its nearest neighbor  $T_{j,m}$  in  $T$ . Formally,  $P_m = [\min(D_1), \min(D_2), \dots, \min(D_{n-m+1})]$ , where  $D_i$  ( $1 \leq i \leq n-m+1$ ) is the distance profile  $D_i$  corresponding to query  $T_{i,m}$  and time series  $T$ .

The  $i$ -th element in the matrix profile  $P$  tells us the Euclidean Distance from subsequence  $T_{i,m}$  to its nearest neighbor in time series  $T$ . However, it does not tell us the *location* of that nearest neighbor; this is stored in the companion *matrix profile index*:

**Definition 5:** A *matrix profile index*  $I$  of time series  $T$  is a vector of integers:  $I = [I_1, I_2, \dots, I_{n-m+1}]$ , where  $I_i = j$  if  $d_{i,j} = \min(D_i)$ .

Fig. 2 illustrates the relationship between distance matrix, distance profile (Definition 3) and matrix profile (Definition 4). Each element of the distance matrix  $d_{i,j}$  is the distance between  $T_{i,m}$  and  $T_{j,m}$  for  $1 \leq i$  and  $j \leq n - m + 1$  of time series  $T$ .

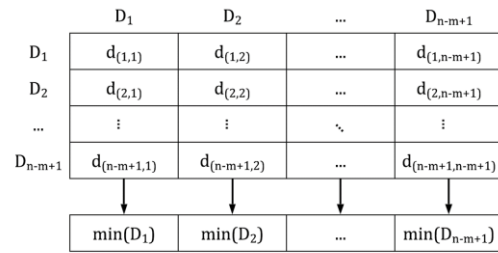


Fig. 2. The relationship between the distance matrix, distance profile, and matrix profile. A distance profile  $D_i$  is a column (also a row) of the distance matrix. The matrix profile stores the minimum (off diagonal) value of each column of the distance matrix; the location of the minimum value within each column is stored in the companion matrix profile index.

Fig. 3 shows a visual example of a distance profile and a matrix profile created from the same time series  $T$ . Note that as we presented it above, the matrix profile uses the z-normalized Euclidean distance [19]. However, this is logically equivalent to the Pearson correlation, and we can convert between them with ease. Some communities prefer to work with Pearson correlation while our work remains agnostic to such considerations.

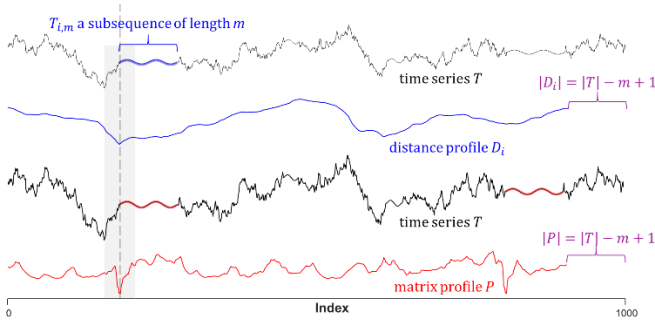


Fig. 3. *top*) A distance profile  $D_i$  created from  $T_{i,m}$  shows the distance between  $T_{i,m}$  and all the subsequences in  $T$ . The values in the dark zone are ignored to avoid trivial matches. *bottom*) The matrix profile  $P$  is the element-wise minimum of all the distance profiles ( $D_i$  is one of them). Note that the two lowest values in  $P$  are at the location of the 1<sup>st</sup> motif in  $T$ .

**Definition 6:** A *Pan Matrix Profile* (PMP) of a time series  $T$  is a matrix whose rows are the matrix profiles  $P_i$  of some time series  $T$ . The PMP is accompanied by a PMP index, recording the location of the nearest neighbor for each MP in the PMP.

To avoid extracting redundant motifs we must understand the issue of *trivial matches*:

**Definition 7:** (Trivial matches): Given a time series  $T$  of length  $n$  containing subsequence  $T_{p,m}$ , if  $T_{p,m}$  scores highly on any scoring function, then  $T_{j,m}$  where  $j \in [\min(1, p - m/2), \max(p + m/2, n)]$  will almost certainly score high on the same function. These spurious high scoring subsequences are *trivial matches*.

To avoid the false positives of trivial matches when finding the top- $K$  matches to a query, we discard some of the patterns using the concept of an *exclusion zone*, a standard practice [3].

### III. COMPUTING THE PAN MATRIX PROFILE

Before introducing algorithms to *compute* the PMP, we introduce motif-heatmaps, a technique to *visualize* the PMP.

#### A. Visualizing the PMP with Motif Heatmaps

While many algorithms treat the classic MP as a “black box” [22], it can be very helpful to visualize the MP for exploratory data analysis. At a quick glance, the MP can be used to visualize the *frequency* and *fidelity* (how well-conserved), and the *location* of motifs in a time series (Fig. 3.*bottom*).

We would like to achieve a similar visualization for the PMP. To achieve this, we propose mapping each MP to a one-dimensional row of a bitmap image, recording  $y$ -axis heights as a color gradient using a heatmap.

We illustrate this, in Fig. 4 using a text string analog. Given the text string:

$T$ : d3icdmy19qicdmnu19a

we compute its “Matrix Profile” at every subsequence length from 1 to 5. Here the colors are discrete because subsequences  $T_{i,m}$  and  $T_{j,m}$  either match or they do not. Corresponding to locations at the apex of each dark triangle, we have maximal motifs of length four beginning at location 3 (icdm) and length two beginning at location 8 (19) which correspond to locations 11 and 17, respectively.

$$PMP_{(i,j)} = \begin{cases} \blacksquare & \text{if } T_{i,m} = T_{j,m} \\ \square & \text{if } T_{i,m} \neq T_{j,m} \end{cases}$$

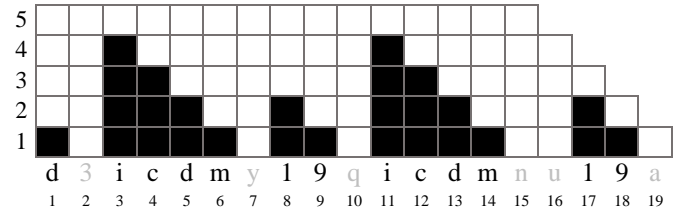


Fig. 4. A binary valued visualization of the PMP where subsequences either match (black) or don't match (white) for subsequence lengths  $m = 1, 2, \dots, 5$ .

In a real-valued time series case, the colors of the PMP can take on subtle gradations of color or grayscale to indicate the degree of similarity.

Though motif-heatmaps perform a similar role as the ubiquitous *dot-plots* used in bioinformatics, they are not directly comparable, as dot-plots are only defined for discrete data, although a handful of papers have suggested discretizing real-valued time series to avail a dot-plot.

#### B. Computing the PMP

We begin with a concrete statement of the problem we wish to address:

**Problem Definition:** Given a time series  $T$  of length  $n$ , and a fixed range of subsequence lengths  $i$  with lower bound  $L$ , upper bound  $U$ , and step size  $S$ , we wish to produce the pan matrix profile PMP whose rows consist of matrix profiles  $P_i$ :

$$PMP = [P_L \ P_{L+S} \ \dots \ P_U]^T$$

In addition, we wish to produce a matrix PMPI whose rows consist of the matrix profile index  $I_i$ :

$$PMPI = [I_L \ I_{L+S} \ \dots \ I_U]^T$$

Before outlining our solution to this problem, we dismiss two apparently *promising* directions. Since the matrix profiles  $P_i$  and  $P_{i+1}$  will be highly related, we may attempt to “cache” some calculations used to compute one in order to reduce the number of computations required to compute the other; however, to produce meaningful results we use z-normalized Euclidean distance (or equivalently, Pearson correlation) [3][12][23][19] which makes such caching impossible. In addition, given a matrix profile  $P_i$ , it is impossible to predict or even produce an upper or lower bound for matrix profile  $P_{i+1}$  since  $\max(D_j)$  for  $P_{i+1}$  may be significantly greater than its value in  $P_i$  as shown in Fig. 5 for a toy dataset with embedded noisy sine waves.

Thus, we believe there is no direct way to exploit the redundancy of adjacent matrix profiles to reduce computation. However, as we will show, we do exploit this redundancy to *order* our calculations, and achieve a faster convergence in the early stages of our anytime algorithm [19][24].

Calculating the PMP for a time series  $T$  for a range of subsequences  $r$  reduces to the calculation of  $r$  matrix profiles  $P_1, P_2, \dots, P_r$ . This can easily be calculated using the brute force algorithm outlined in **Error! Reference source not found.**

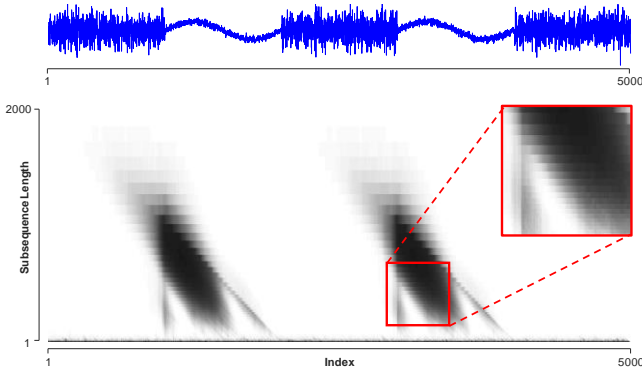


Fig. 5. A real-valued visualization of the Pan Matrix Profile of a time series  $T$  with length 5000 with a magnified “detached motif,”  $P_i < P_{i+1}$  for some index.

TABLE 1: A BRUTE FORCE ALGORITHM TO CREATE THE PMP

<b>Input:</b>	T:	Time series
	L:	Subsequence length lower bound
	U:	Subsequence length upper bound
	S:	Subsequence length range step size
<b>Output:</b>	PMP:	Pan matrix profile
1	R = L : S : U // [L, L+S, L+2S, ..., U]	
2	PMP = [] //  T  x  R  matrix of zeros	
3	for r in R	
4	PMP <sub>r</sub> = BuildMP(T, r) // (Definition 4)	
5	return PMP	

In this algorithm we begin by explicitly specifying the range  $R$  of subsequence lengths we wish to explore (line 1) and then calculate the matrix profile  $P_r$  for consecutive subsequence lengths  $L + iS$  for  $i = 0, \dots, (U - L)/S$  (lines 3-4). In line 4 we use the .MP, currently SCRIMP or STOMP [23][19]. Using this algorithm, we can generate the complete PMP shown in Fig. 6 with maximal motifs at subsequence length  $m_1 = 68$  and  $m_2 = 610$ . An approximation of the PMP is depicted in Fig. 6 after 16 iterations, that is, after calculating  $P_1, P_2, \dots, P_{16}$ , about 2% of the exact PMP has been calculated.

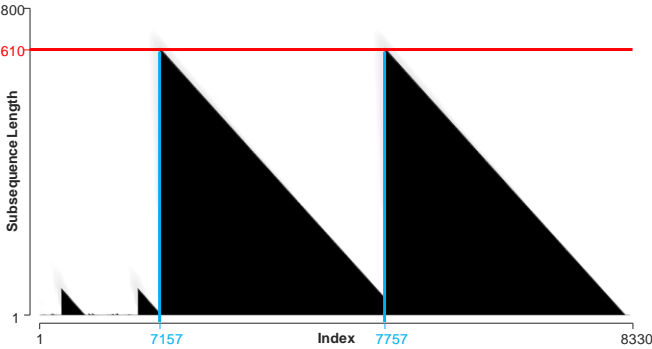


Fig. 6. The PMP of a time series  $T$  with length  $n = 8330$  where we consider subsequence lengths bounded by lower bound  $L = 1$ , upper bound  $U = 800$ , and step size  $S = 1$ . The slice  $m = 610$  corresponds to matrix profile  $P_{610}$  which has minimal values at  $t_1 = 7157$  and  $t_2 = 7757$ . Note that this example is based on real data shown in Section IV.B.

Until relatively recently, computing this would have required  $R$  invocations of an  $O(n^2r)$  algorithm. As we will show in our experimental section,  $R$  could be over 10,000, making this algorithm completely untenable. The STOMP algorithm [23] is able to compute a single MP in just  $O(n^2)$

time, giving us an overall  $O(n^2R)$  algorithm to compute the PMP. This may be tenable for small datasets, especially if we avail the GPU or multicore versions of STOMP that now exist [23]; however, it is clearly limiting given the typical sized datasets that modern data analysts need to deal with.

Though we strongly suspect that  $O(n^2R)$  may be optimal for the task-at-hand, our key insight is while it may take  $O(n^2R)$  time to converge to the exact answer, by carefully ordering the computations, we can typically allow it to converge to 99% of the exact answer, after doing less than 1% of the computations. By computing the PMP with a novel anytime algorithm, we can increase the size of considered datasets by orders of magnitude.

Before continuing, it is important to ward off a possible misunderstanding. There already exist anytime algorithms to compute the MP, in particular STAMP [19] and the more recently introduced SCRIMP [23]. However, these algorithms only compute a *single* MP. To ensure fast convergence of the PMP we must optimize anytime performance at a higher level.

### C. SKIMP: An Anytime Algorithm to Compute PMP

We are finally in a position to introduce SKIMP, which we outline in TABLE 2. Unlike brute-force search, SKIMP recursively subdivides the range into equally spaced regions with increasing granularity (line 1). This has a similar effect to iterating through a balanced binary search tree on the range  $r = [1, 2, \dots, (U - L)/S]$  using breadth-first search.

TABLE 2: THE SKIMP ALGORITHM

<b>Input:</b>	T:	Time series
	L:	Subsequence length lower bound
	U:	Subsequence length upper bound
	S:	Subsequence length range step size
<b>Output:</b>	PMP:	Pan matrix profile
1	R = BFS(BuildBalancedBST(L, U, S))	
2	PMP = [] //  T  x  R  matrix of zeros	
3	for r in R	
4	PMP <sub>r</sub> = BuildMP(T, r) // (Definition 4)	
5	return PMP	

As the initialized PMP (line 2) is approximated for each subsequence  $r$  in  $R$  (line 3-4), we see a “blocky” approximation of triangles being progressively refined exemplified in Fig. 7. The example shown is slightly cleaned and contrived for clear display in this limited format of presentation, but [25] contains videos of this process created on several real-world datasets. This process is reminiscent of the classic idea of progressive refinement of raster images [15]; however, in that case, the limiting factor was bandwidth, while for us it is CPU time.

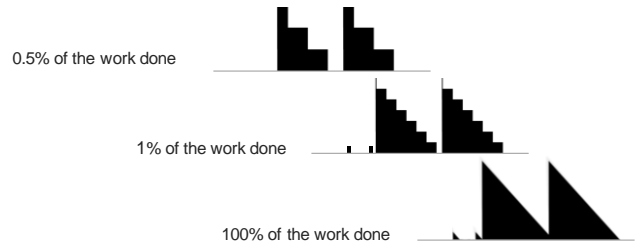


Fig. 7. A visual trace of the PMP shown in Fig. 6 approximated using known PMP values. *top*) After four iterations a single motif has been located. *middle*) A new motif with a significantly smaller subsequence length is located while the initially discovered motif is refined. *bottom*) The complete PMP as shown in Fig. 6.



As we will show in Section IV, SKIMP’s approximation of the PMP depicted in Fig. 6, and many other real-world datasets converge significantly faster than brute-force search, typically achieving 99% accuracy in less than 1% of the work.

#### D. Ranking Motifs of Different Lengths

Given that SKIMP will allow us the possibility to find motifs of any length in the range  $r$ , it is natural to ask how we can rank motifs of different lengths. In many cases, we envision that a higher-level algorithm will make requests for motifs of different lengths, based on its own criteria (which could possibly include out-of-band information), thus absolving us of the responsibility to address this question. Nevertheless, it is an interesting question to answer. For a handful of user cases, especially on relatively small datasets, we envision this being an interactive process. Thus, we have built an interactive and visual tool to allow a user to explore and discover multiple length motifs. Nevertheless, we clearly need an algorithm that allows us to meaningfully rank motifs of different lengths.

#### E. Computing PMP with Unbounded $U$

The PMP allows for the first truly parameter-free algorithm for finding time series motifs (we could envision several algorithms to find motifs from the PMP, in Section III.C we gave one such example). While  $L$  is bounded by the shortest logical subsequence length, and  $S$  simply affects the desired level of granularity, the reader may argue that the value of  $U$  is a parameter and could be as long as  $n/2$ . However,  $U$  is only a parameter in a very weak sense, so long as it is larger than the length of the longest motif in the data, its value is inconsequential. For example, in our termite DNA example in Section IV.B the longest motif has length 610. As we did not know this in advance, we set  $U$  to a very conservative 2,400. This clearly worked, but one could argue that about 75% of the computations (from 618 to 2,400) were wasted. Can we prevent such wasted computations?

If we assume that we have a test to detect when the first row (from  $L$  upwards) of the PMP is devoid of meaningful motifs, then a simple algorithm suggests itself. We can compute this test on the  $MP_i$  with  $i = L$ , then iteratively double  $i$ , computing then testing  $MP_i$  until the test fails. We can then use  $i$  as the value of  $U$ , and simply call the SKIMP algorithm.

Note that since we have already computed  $\log_2(U)$  of the  $|R| = (U - L)/S$  MPs that SKIMP will compute, we can slightly modify SKIMP to ingest these MP and avoid recomputing them.

This idea is predicated on the assumption that we have a test to detect when the first row of the PMP is devoid of meaningful motifs. One way to achieve this, is to set a threshold for the correlation. We can calculate the maximum correlation for each subsequence and if the correlation falls below the threshold then we stop calculating the PMP for the larger subsequence length as outlined in TABLE 3.

Using this algorithm Fig. 8 indicates a suitable upper bound at subsequence length 660 when using a maximum correlation threshold of  $t = 0.988$  on a mtDNA sequence of *Coptotermes suzhouensis* (a termite) depicted in Fig. 13. Gratifyingly, we see dip just after 612, the objectively correct location [10].

TABLE 3: FINDING A SUITABLE UPPERBOUND FOR  $U$

<b>Input:</b>		T: Time Series
		t: Threshold
<b>Output:</b>		S: Maximum subsequence length
1	k = 8	// 8 is the shortest sensible motif
2	c = inf	
3	while (c >= t)	// t is set with domain knowledge
4	c = maximum(PCmatrixProfile(T, k))	
5	k = k * 2	// iteratively double
6	return k	

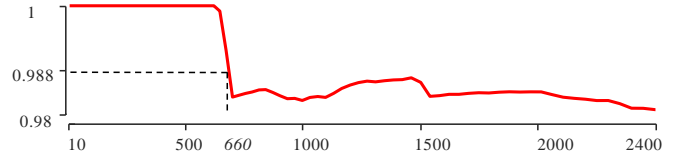


Fig. 8. The maximum correlation for different subsequence length from a mtDNA sequence of *C. suzhouensis* depicted in Fig. 13. Setting the threshold to 0.988 stops calculation of PMP for lengths greater than 660.

This opens the question of how we can set  $t$ . While this is a domain dependent value, it seems to be robust *within* a single domain. For example,  $t = 0.988$  worked well on the termite DNA, the soybean DNA (Fig. 17), and all other DNA data we considered.

#### F. Extracting the Top- $K$ motifs

An almost trivial application of the PMP is our algorithm for extracting the length agnostic top- $K$  motifs as outlined in TABLE 4.

TABLE 4: DISCOVERING TOP-K MOTIFS (ANOMALIES)

<b>Input:</b>		PMP: Pan Matrix Profile
		T: Time series
		k: Number of motifs
<b>Output:</b>		TM: top- $K$ motifs
1	TM ← {}	
2	while  TM  < k	
3	[idx, s] = maximum(PMP)	
4	if TM is not covering T[idx:idx + s]	
5	TM ← T[idx: idx + s]	
6	Apply exclusion zone // Definition 7	
7	return TM	

Given a time series  $T$ , the corresponding  $PMP$  for some range, and a user-defined value for the number of motifs  $k$ , our algorithm returns the subsequences  $T_{i,m}$  which correspond to the top- $k$  motifs of the PMP. To extract the top- $k$  motifs, we repeatedly search the  $PMP$  for its minimum value (line 3) and then add the corresponding subsequence to our top- $k$  motifs  $TM$  only if  $TM$  does not span the subsequence (lines 4-5). Afterwards, we apply an exclusion zone using the recovered subsequences to ensure we do not find a trivial match (line 6). Using this algorithm, we discovered the top-2 motifs for the mitochondrial DNA sequence shown in Fig. 13  $T_{7713,615}$  and  $T_{7092,70}$  and their respective nearest neighbors  $T_{7148,615}$  and  $T_{6900,70}$  which is in near perfect agreement with the ground truth noted in [10] and will be illustrated in Fig. 13.

#### G. Anomaly Detection

The previous algorithms for detecting subsequence length agnostic top- $k$  motifs can be easily modified to detect anomalies

with variable lengths. By “inverting” the PMP,  $PMP' = 1 - PMP$  the top- $k$  motifs produced by the algorithm would correspond to the top- $k$  anomalies. We exemplify this algorithm’s ability to perform anomaly detection on an automated pedestrian counting system developed in Taipei to better understand pedestrian activity within the municipality. This information examines how people use different city locations at different times of day to better inform decision-making and infrastructure planning. We extract data from the Xindian District Office as shown in Fig. 9.



Fig. 9. Pedestrian count data from Taipei Xindian District Office metro station starting on December 2015 and ending at March 2017.

Fig. 10 shows one fairly typical week of this behavior.

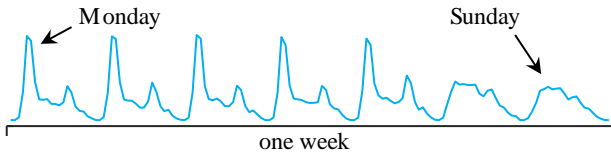


Fig. 10. One fairly typical week data of the pedestrian counting data of Taipei.

After computing the PMP for this data from  $L = 20$  points (~one day) to  $U = 200$  points (~10 days), we can then extract the top- $k$  anomalies using the modified version of the top- $k$  motifs algorithm described in Section F.

Fig. 11 shows the top-4 anomalies that exist in this dataset with the anomaly shown in red.

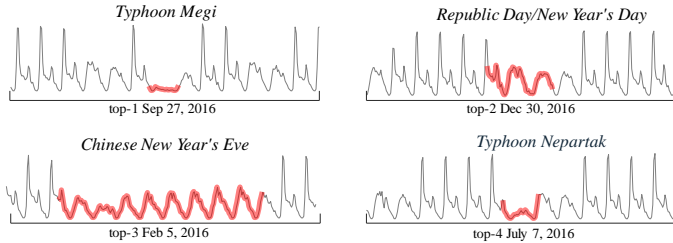


Fig. 11. The top-4 anomalies (red) detected for the data shown in Fig. 9.

Note that these four anomalies represent four different lengths. The first anomalies length is 27 points (~one day). We found the ground truth for this event which is described by [24] as “On September 26/27, 2016 ... Typhoon Megi ... (made) landfall on Taiwan’s southeast coast” The second and third anomalies belong to Republic day/New year’s day and Chinese New Year’s Eve with a length of 59 (~2.5 days) and 185 points (~one week). The last anomalies’ length is 32 points (~1.5 days) which is reported by [4] as “The storm is predicted to make landfall on the island nation on July 7.”

If we set the length of classic anomaly detection algorithm to one day, it fails to find Chinese New Year’s Eve or Republic Day/New year’s day. In contrast, if we set the length of classic anomaly detection to one week, we cannot detect the other anomalies are that are present in Fig. 11. This strongly motivates the need for computing similarity search for variable length.

## IV. EXPERIMENTAL EVALUATION

To ensure that our experiments are reproducible, we have built a website [25] which contains all data/code/raw spreadsheets for the results, in addition to many experiments that are omitted here for brevity. Unless otherwise stated, all experiments were run on a Dell XPS 8920, with Intel Core i7-7700 CPU @ 3.6GHz and 64GB RAM.

### A. A Benchmark for the All-Length Motif Problem

To concretely ground our ideas throughout this paper, let us consider a motivating problem introduced to us by (blinded). They are interested in finding motifs in time series from a large industrial distillation column. The apparatus is massive with great thermal and mechanical inertia, so that it suffices to sample it once per minute (1/60 Hertz). When doing analytics, it is common for them to consider data from the previous year; thus, we have a time series with  $n = 525,600$  points. Let us call this dataset DisCol. Occasionally, DisCol is searched for motifs which are used in downstream analytics to perform root-cause analysis. Though most patterns last for about a day, the fast cooling process of the apparatus by a summer rain shown can induce patterns lasting for only a few hours; thus, there is great uncertainty in the potential length of motifs motivating the desire to identify motifs between the length of one hour ( $L = 60$  minutes) and one day ( $U = 1,440$  minutes), a range of  $|R| = 1,380$  values.

Prior to the introduction of the Matrix Profile, the only exact algorithm to find all such high-dimensional motifs was brute force search, which would take  $O(n^2r^2)$  time. The factor  $r$  appears both as the subsequence length (more conventionally denoted  $m$ ), and the number of times we must run the motif search. Concretely, on our desktop, this would require about 48 years. Using the recently introduced STOMP algorithm, which can find motifs of a fixed length in time independent of that length, this can be reduced to  $O(n^2r)$  time, or about 23 hours. However, since STOMP is a batch algorithm, it is natural to ask how quickly we can converge to an acceptable approximation of the final PMP.

We have created a proxy for the data in question by editing together some publicly available industrial benchmarks from a similar process. In Fig. 12 we show how fast SKIMP converges on this dataset.

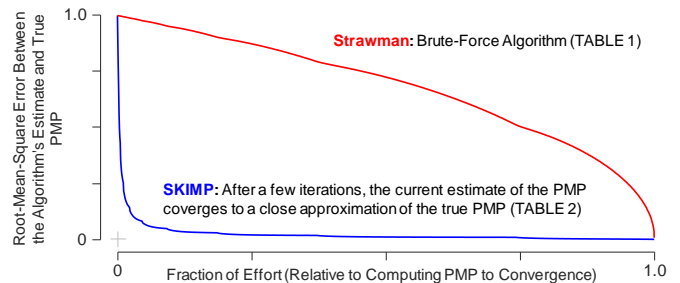


Fig. 12. The root-mean-square error of our approximation of the fully converged PMP when using SKIMP and the brute-force algorithms. When compared to the ten hours required to fully compute the PMP using the brute-force algorithm (Error! Reference source not found.), SKIMP required less than 41 min, or 3% of the required effort to achieve an approximation with less than 10% error. A video showing the convergence of the PMP approximation using SKIMP is available at [25].

In an ideal case, there should exist an approximation of the PMP which converges to the exact PMP using a small fraction of the effort required to compute the complete PMP. In less than 3% of the required effort, SKIMP was able to approximate the full PMP with less than a 10% root-mean-square error. Though it may seem as if the root-mean-square error plot depicted in Fig. 18 converges slower than the plot depicted in Fig. 12 this plot is significantly shorter (by an order of magnitude) but still only requires a handful of iterations before converging to the complete PMP.

### B. DNA-Based Benchmarks for the All-Length Motif Problem

To demonstrate the utility and correctness of time series motif length discovery, we exploit a technique long used by the time series data mining community. By converting discrete DNA sequences to real-valued time series, we can explore the time series space of the DNA sequence and then attempt to confirm our findings with molecular biologists. In Fig. 13 we show the complete mitochondrial DNA sequence of *Coptotermes suzhouensis*, a subterranean termite pest of wooden structures as a real-valued time series and its corresponding PMP as a motif-heatmap in Fig. 14.

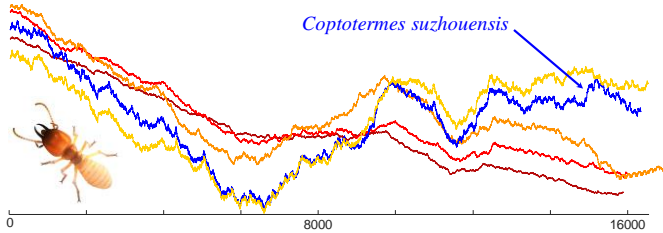


Fig. 13. The mitochondrial DNA sequence of five randomly chosen insects, including the 16,326 bp mitochondrial DNA sequence of *Coptotermes suzhouensis* (blue).

These results are suggestive of a strongly conserved motif of a length of about 610 in the center of the sequence. In a recent paper announcing the complete mitochondrial genome of this insect, the authors noted that the mitogenome had two repeat units, **A** and **B**. Unit **A** is just 66 bp long, however “The **B** repeats consisted of one complete unit **B1** (562 bp) and a partial unit **B2** (38 bp)” [10]. The reader will appreciate that  $562 + 38$  sums to 610, which is just 0.03% less than our suggested motif length of 612 for this dataset.

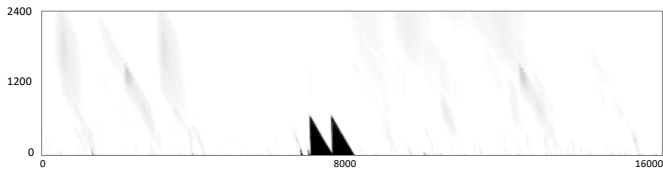


Fig. 14. A motif-heatmap of (the time series representation of) the mitochondrial DNA sequence of *Coptotermes suzhouensis* in Fig. 13 from  $L=10$  to  $U=2,400$ .

Moreover, if we zoom in as shown in Fig. 15, the **A** motif is also clearly visible.

The reader may wonder if DNA time series is “too easy” given the level of conservation observed. To address this, we can revisit our termite example. This time, before converting the DNA string to time series, we randomly changed every base with a one in sixty-four probability, simulating a high mutation

rate. As the motif heatmap in Fig. 16 reveals, this level of noise makes no appreciable difference in our ability to find the motifs.

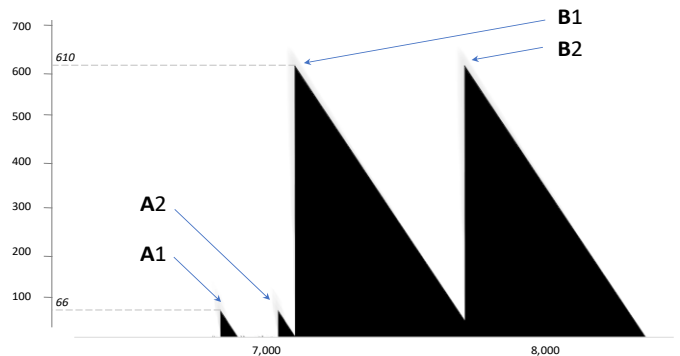


Fig. 15. A zoom-in of the center bottom of Fig. 14 shows that the motif heatmap also discovered the much shorter **A** motif.

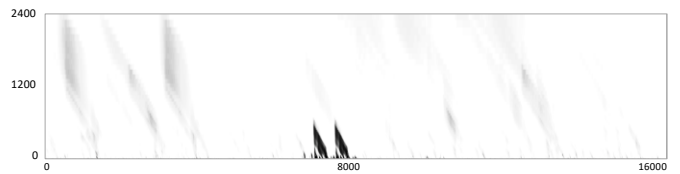


Fig. 16. (contrast with Fig. 14) A motif-heatmap of (the time series representation of) the mtDNA of sequence *Coptotermes suzhouensis* after 1-in-64 bases were randomly changed.

Above we have demonstrated our technique on an insect which was chosen to be visually clear in two-column format, but it is arguably too simple to really challenge our algorithms. In order to stress test SKIMP, we can turn to plant mtDNA. It has long been noted that “Unlike the relatively simple mitochondrial genomes of animals, the genomes of nonparasitic flowering plant mitochondria are large and complex.” [1]. Thus, we consider the mitochondrial genome of Soybean (*Glycine max*) [2]. Because it is 402,540 bp long and has repeats that differ over three orders of magnitude in length, it is difficult to do it full justice in this paper. In the accompanying web materials [25], we show a video of our methods applied to it, and here we content ourselves with a figure that allows us to see only the longer motifs.

From the literature we know that repeats in plants may be as long as 10,000, thus we consider  $L = 1,000$ ,  $U = 10,000$ , with  $S = 1$ . To run this dataset to convergence requires about 42 days, however, as the video at [25] shows, in about half a day, the basic shape of the final motif heatmap has already emerged.

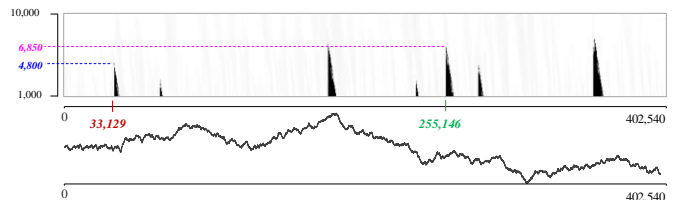


Fig. 17. *bottom*) The mtDNA of Soybean in a time series representation. *top*) The motif-heatmap of Soybean. The location and length of two motifs are highlighted. As [2] discovered, the (location/length) of the first occurrence of **R2** is (33,155/4,692), which is very close to our result of (33,129/4,800). Similarly, [2] notes the (location/length) of **R1** is (255,146/6,502), we discovered a motif the exactly the right location 255,146, with a slightly different length 6,850.



This experiment offers strong evidence of the utility of our anytime approach. In addition, SKIMP was able to approximate the PMP with less than 10% root-mean-square error when performing less than 4% of the required effort as shown in Fig. 18.

Note that such DNA repeats could also be visualized using dot-plots. However, recall that dot-plots require  $O(n^2)$  space, whereas motif-heat maps require only  $O(nr)$  space, and their long aspect ratio is amenable to panning interactions when dealing with long sequences. More importantly, dot-plots are only well defined for discrete strings while motif-heat maps facilitate the visualization of *real-valued* data.

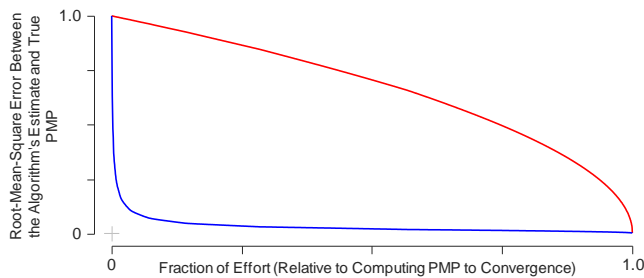


Fig. 18. The change in the root-mean-square error when approximating the PMP after performing a fraction of the work required to compute the PMP to convergence on the mtDBA Soybean time series representation. Using less than 4% of the time required to compute the entire PMP, SKIMP found an approximation to the PMP which had a root-mean-square error of less than 10%. A video showing the converging of SKIMP is available at [25].

## V. RELATED WORK

The literature on time series motif discovery is large and growing, see [19] and the references therein. However, to the best of our knowledge, there are no other algorithms that can approximately or exactly discover *all* motifs of arbitrary lengths.

In a sequence of papers, Lin and colleagues introduce a series of tools to allow interactive discovery of variable-length time series patterns [20]. However, this work is not directly comparable to PMP. First, because they use a discretized representation of the data (for efficiency), they are always condemned to finding approximate answers. Second, the system only returns information about a small *subset* of the patterns, whereas PMP contains exact distances for *all* subsequences of *all* lengths. Finally, there are many parameters to be set and choices to be made in the grammar inference algorithm. However, like us, the authors see great value in attempting to visualize the results of the motif search.

## VI. CONCLUSIONS

We have introduced the first practical technique to find motifs and discords [19] for all lengths. Given the glut of information that this provides, we have also introduced a novel visualization that allows a practitioner to understand the location, length, and fidelity of all motifs in her dataset. We have shown that these new tools allow us to find useful conserved structures and anomalies in domains as diverse as bioinformatics, transportation, and seismology.

In future work plan to investigate the implications of our ideas for other algorithms that exploit the matrix profile, including chain discovery [22] and segmentation.

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