Time Series Data Mining for Analyzing Livestock Eamonn Keogh

eamonn@cs.ucr.edu

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Welcome

- Congrats! About 90% of the best minds in data science are implicitly or explicitly working on the problem of getting people to click on ads! You are doing something more interesting and noble.
- If you like this tutorial, you may enjoy my others ;-)

Format

- This tutorial has almost no math or code.
- I want to give you the *intuition* behind the SOTA time series data mining.
- While these ideas are general, I *have* successfully applied them to chickens, insects, seals, cows, penguins, mice etc.
- There is no deep learning in the tutorial!
- Contrary to 100's of paper's claims, there is no evidence that deep learning helps for time series problems.
- I am not going to spend time explaining this, if you are interested, see this talk…

Preview

- What do you want to do with livestock time series?
	- Classification, clustering, summarization, visualization, segmentation, anomaly discovery, repeated pattern discovery, emerging behavior discovery etc.
	- Amazingly, one simple tool, the *Matrix Profile*, is basically all you need to do all the above!
- To analyze time series, there are two different situations, which have almost no overlap:
	- Behaviors are conserved in *shape* (30 min)
	- Behaviors are conserved in *features* (10 min)

Comparing two time series

How similar are these two-time series?

C

Q

 $(Q, C) = \sqrt{\sum_{i} (q_i - c_i)}$ = *n i* $D(Q, C) = \sqrt{\sum_{i} (q_i - c_i)}$ 1 \sum_{i}^{n} \sum_{i}^{n} $\left(q_{i} - c_{i} \right)^{2}$ *Q C D*(*Q*,*C*) Euclidean Distance Metric (ED) Given two time series: $Q = q_1...q_n$ $C = C_1...C_n$ We always z-normalize the data before computing the ED

This is logically equivalent to Pearson's Correlation

Euclidean Distance Metric (ED)

Z-normalization means we are ignoring mean and standard deviation of the data.

At least 99.9% of the time, that is the right thing to do.

So here *D*(*Q*,*C*) = *D*(*Q2*,*C*) = *D*(*Q3*,*C*) =*D*(*Q4*,*C*)

Let us use Apollo to make a running example for this tutorial

The IMU used in the collar tags is InvenSense MPU-92[5](https://www.sciencedirect.com/science/article/pii/S2772375522001277#fn0005)0⁵ at 50 Hz

triaxial accelerometer

To be clear, X , Y and Z do not have any standard meaning for animal work. *Surge*, *Heave* and *Sway* are the biological terms for acceleration directions. (roll, pitch, yaw for rotation, are measures with a gyroscope).

It is rare that X, Y and Z can be made to exactly map to Surge, Heave and Sway, because of the orientation of the sensors against the animal's body.

-0.6 -0.5 -0.4 -0.3 -0.2

-0.1 0

0.1 0.2 0.3

Alternatives to Accelerometers

Markerless tracking of animals has recently become *incredibly* robust and easy.

For this tutorial, I don't care how you got your time series.

http://www.mackenziemathislab.org/deeplabcut

If possible, try to get *labeled* data.

- Could be
	- Apollo vs. Dante
	- Apollo before operation vs. Apollo after operation
	- Apollo on pasture vs. Apollo on grain
	- Apollo during winter vs. Apollo during summer
	- etc.

Try to get *all* labels, *all* metadata at data collection time.

It is very frustrating to find interesting structure in the data but have no way to "go back in time" to discover its cause.

As it happens, for this dataset, we have wonderfully detailed labels, a second-bysecond annotation of the behavior.

Lets zoom-in on 20 seconds of ruminating behavior Call it ApolloYSUB

I have this one-second-long behavior $\int\sqrt{\ }$, I am going to call is ${\tt query}.$

I have reason to think that it is indicative of Bovine spongiform encephalopathy (BSE)

Does this query behavior exist in Apollo? To find out, we will build a *distance profile*.

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This is simply the z-normalized Euclidean distance between the query and every subsequence in the longer time series…

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Lets do a full worked example

I have this dataset, measuring the motion of a dog walking in my lab, first on concrete, then carpet, then concrete…

I also have a query, for the same dog walking, on a different day.

Do you think the query is from when the dog was walking on carpet or concrete or something else…

This task is trivial with MASS code…

>> dist = MASS(dog , carpet_query);
 % compute a distance profile

 \Rightarrow [val loc] = min(dist);

>> disp(['The best matching subsequence starts at ', num2str(loc)])

The best matching subsequence starts at 7479

Below we plot the *16* best matches. Note that they all occur during the carpet walking period. This entire process takes about 1/10,000th of a second.

Note that this example is moving beyond nearest neighbor search, and is really performing *semantic segmentation* into regimes..

Mini-Review: The *distance profile*

- This is a simple idea called: query-by-content/similarity-search/nearest-neighbor search etc.
- You can use the distance profile to find the K-nearest neighbors to any query.
- This is an *incredibly* powerful and useful tool, limited only by your imagination.
- It is by far, the most important subroutine in all of time series data mining.
- Suppose you have multidimensional data? You can just compute the individual distance profiles, sum them, then find the lowest values as the multidimensional nearest neighbor!
- I gave you a fully worked example in the appendix (penguin)
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- The computation of the distance profile is *incredibly* fast.
- We can search the query in 24 hours of our bovine data (4,320,000 datapoints), in well under a second.
- This amazing speed is due to Mueen* (my former PhD student). His algorithm is called MASS*
- You can get MASS in most computer languages.

*https://www.cs.unm.edu/~mueen/FastestSimilaritySearch.html

Preview: The *Matrix Profile*

- We are about to learn about the Matrix Profile*
- The Matrix Profile is the best idea in time series data mining in the last decade ;-)
- It is a stunningly simply idea.
- My claim is that once you have the Matrix Profile, almost all time series problems are trivial.
- Longer tutorials are online

*https://www.cs.ucr.edu/~eamonn/MatrixProfile.html

The Matrix Profile has exploded in popularity in the last 3 years

observations of the magnetosphere collected by the Cassini spacecraft in orbit around Saturn.. ..in this case, the best-performing method was the Matrix Profile.. Kiri L. Wagstaf et. al. NASA JPL.2020

(for an industrial IoT problem) *Matrix Profiles perform well with almost no parameterisation needed*. Anton et al ICDM 2018.

While there will never be a mathematical silver bullet, we have discovered that the Matrix Profile, a novel algorithm developed by the Keogh research group at UC-Riverside, is a powerful tool. Andrew Van Benschoten, lead engineer at Target.

If anybody has ever asked you to analyze time series data and to look for new insights then (the Matrix Profile) is definitely the open source tool that you'll want to add to your arsenal Sean Law, Ameritrade. (for) intrusion detection in industrial network traffic, distances as calculated with Matrix Profiles rises significantly during the attacks. ..as a result, time series-based anomaly detection methods are capable of *detecting deviations and anomalies.* Schotten (2019).

The MatrixProfile technique is the state-of-the-art anomaly detection technique for continuous time series. Bart Goethals et. al. (ECML-PKDD 2019).

Based on the concept of Matrix Profile ..without relying on time series synchronization.. the Railway Technologies Laboratory of Virginia Tech has been developing an automated onboard data analysis for the *maintenance track system* Ahmadian et. al. JRC2019

Matrix Profile is the state-of-the-art similarity-based outlier detection method. Christian Jensen et. al. IJCAI-19

we use the exact method based on the Matrix Profile (to assess the effectiveness of therapy) Funkner et al Procedia 2019.

Recently, a research group from UCR have proposed a powerful tool - the Matrix Profile (MP) as a primitive...(we use it for) *fault detection* Jing Zhang et al. ICPHM 2019

Inspecting both graphs one can see that the matrix-profile algorithm was able to identify regions where there is a change on the power level over the observed band. [F Lobao](https://www.researchgate.net/profile/Fabio_Santos_Lobao/publication/336316249_Intelligent_Radio_Spectrum_Monitoring/links/5d9bc29c299bf1c363ff051c/Intelligent-Radio-Spectrum-Monitoring.pdf) 2019.

RAMP builds upon an existing time series data analysis technique called Matrix Profile to detect anomalous distances...collected from scientific workflows in an online manner. Herath et. al. IEEE Big Data 2019

Based on obtained results for the considered data set, matrix profiles turned out to be most suitable for the task of anomaly detection Lohfink et al. VISSEC2019

The computation speed and exactness of the Matrix Profile make it a powerful tool and (our) results back this. Barry & Crane AICS 2019

(examining) *manufacturing batches considering raw amperage* (we found that the) *Matrix Profile highlights anomalies* Hillion & O'Connell of TIBCO Data Science. re:Invent 2019. [

we use the exact method based on the matrix profile to search for motifs can be used to monitor the patient's condition, to assess the effectiveness of therapy or to assess the physician's actions. Funknera et al.

(The Matrix Profile is a) *similarity join to measure the similarity between two given sequences. we opt for the median of the profile array as the representative distance* (3D Dancing Move Synthesis from

Music)" Anh et al. IEEE Robotics and Automation Letters

We were amazed by the power of MP and seek to incorporate it into our framework Ye and Ageno.

..*adopting the concept of (the) Matrix Profile, we conduct the first attempt to*.. J. Zuo et. al. Big Data 20019

The accuracies obtained ...indicate that the Matrix Profile is useful for the task at hand instead of using the CNN features directly Dhruv Batheja

To speed up online bad PMU data detection a fast discovery strategy is introduced based on (the Matrix Profile) Zhu and Hill.

Specifically, ALDI uses the matrix profile method to quantify the similarities of daily subsequences in time series meter data, Zoltan Nagy, Energy & Buildings (2020)

Our two-fold approach first leverages the Matrix Profile technique for time series data mining.. Nichiforov 2020.

*the class of matrix profile algorithms.. ..is a promising approach, as it allows simplified post-processing and analysis steps by examining the resulting matrix profile structure*A. Raoofy et al.

We only require information about the time of several critical incidents to train our methods, as previously. To this end, we employ the Matrix Profle.. Bellas. et al.

a matrix-profile based algorithm applied across all trajectory data against a validation set revealed four significant motifs which we defined as motif A, B, C and D.. Fernandez Alvarez 2020.

The main building block of this (game analytics) algorithm is the matrix profile, Saadat and Sukthankar AAAI2020

We leverage the Matrix Profile (MP),… to create a micro-service-based machinery monitoring solution Naskos et al 2021

SLMAD uses statistical-learning and employs a robust box-plot algorithm and Matrix Profile (MP) to detect anomalies Team from Huawei/UCD.

We found that all these similarity or randomness measures can be estimated with variants of the highly efficient Matrix Profile (MP) algorithm. `

- Let us return to our small bovine example.
- Let us pick a random subsequence of length one second, I happened to pick location 215
- Let us find its nearest-neighbor distance (excluding *itself*) to anywhere else in the time series.
- (we could use the distance profile to do that)
- The distance was 6.1

- Let do that again
- Let us pick a random subsequence of length one second, I happened to pick location 547
- Let us find its nearest-neighbor distance (excluding *itself*)
- (we could use the distance profile to do that)
- The distance was 2.2

- Let us do this for every location!
- The resulting curve is called the *Matrix Profile*

- There are some parts of the Matrix Profile that have special names
- The highest location is called the *Time Series Discord*

- There are some parts of the Matrix Profile that have special names
- The highest location is called the *Time Series Discord*
- The lowest locations (there will always be a tie) is called the *Time Series Motif* pair

Reading a Matrix Profile

Where you see relatively low values, you know that the subsequence in the original time series must have (at least one) relatively similar subsequence elsewhere in the data (such regions are "motifs" or reoccurring patterns)

Where you see relatively high values, you know that the subsequence in the original time series must be unique in its shape (such areas are "discords" or anomalies).

Zebra Finch

(Zebra Finch Vocalizations in MFCC, 100 day old male)

Can you see any conserved behavior here?

Motif discovery can often surprise you.

While it is clear that this time series is not random, we did not expect the motifs to be so well conserved or repeated so many times. There is evidence of a *vocabulary*, and maybe even a *grammar*…

Taxi Example: Part I

Below is the hourly average of the number of NYC taxi passengers over 75 days in Fall of 2014.

Lets compute the Matrix Profile for it, we choose a subsequence length corresponding to two days…. (next slide)

[a] http://futuredata.stanford.edu/ASAP/extended.pdf

Taxi Example: Part II

- The highest value corresponds to Thanksgiving
- We find a secondary peak around Nov $6th$, what could it be? Daylight Saving Time! The clock going backwards one hour, gives an *apparent* doubling of taxi load.
- We find a tertiary peak around Oct 13th, what could it be? Columbus Day! Columbus Day is largely ignored in much of America, but still a big deal in NY, with its large Italian American community.

A quick example of the amazing utility of motif discovery (next 5 slides)

Australian Fur Seal (*Arctocephalus pusillus*)

This is an interesting dataset; can we find motifs in it?

We can find *many* interesting motifs in this data.

Suppose we label them A, B, C, D etc.

We can then ask the question, is there any patterns in the occurrence of these motifs?

In fact, there is, when we see D, we almost always see C within a few seconds…

Then

What does this mean?

Thirty - six seconds

Thirty - six seconds

Thirty - six seconds

Let us return to Apollo

Let's compute the Matrix Profile for this dataset

To do so, we have to compute the Euclidean Distance between the approximately 24 billion possible pairwise combinations of subsequences.

Because of an amazing algorithm called SCRIMP++, we can do this in seconds!

If you see a subsequence (X) such that $D(X, \sqrt{v}) < (3 * 1.29)$ **Then** Print('Bovine is **ruminating!**')

Mini Review I

- What I have shown you is *amazing* if you think about it.
	- Knowing *nothing* about bovine behavior…
	- Using less than eight lines of code in total.
	- Using one minute of brain power, and a few seconds of CPU power.
	- I built a tool to correctly annotate complex behavior in a bovine.
- Was this a fluke?
- Let quickly do it again.
- This time with chickens.
- This time I have four years of chicken data!

Mini Review II

-
- Here is the template the motif discovery found \sim
- It was associated with *Dustbathing*
- I used it to search a 12,679,054,727 datapoint (four full years) archive of chicken behavior for the one thousand best matches.
- It gets better.
- My data is annotated with two types of chicken has-mites|no-mites.
- I can see that Dustbathing is more common in the has-mites class, so have learned that chickens with mites will dustbath more.

Mini Review III

- With just the *Distance Profile* and the *Matrix Profile*, you can solve many (most/all) problems in livestock data analysis.
- Once installed on your machine, these are both one line of code!
- There is a large and growing community of Matrix Profile users, so these tools exist in most languages/platforms.

The *Matrix Profile* generalizes to multiple time series (i.e. joins) so you can ask questions that *compare* and contrast behaviors:

- What patterns occur in *Males*, **but not** *Females* (join *discord*)
- What patterns occur in *Hereford*, **and** *Holstein* (join *motif*)
- What patterns occur before *Denuding*, **but not** after *Denuding* (join *discord*)
- The *Matrix Profile* generalizes to other useful primitives, *Chains*, *Novelets*, *Shapelets*, *Platos, Snippets, FLOSS*….

Switching Gears a Little

Bad news ;-(Sometimes there are behaviors that are not captured well in *shape*.

Think of human behaviors: walking/running/swimming/cycling all have characteristic *shapes*. *Dynamic Behaviors*

But there is no *shape* for reading/watchingTV/resting/sleeping etc.

Instead, we have to consider *features*.

Instead of pulling out subsequence *shapes*, lets pull out subsequences, and measure *features*…

We can then use the many algorithms that ingest *feature vectors*, nearest neighbor, decision trees, naïve Bayes,…

This begs the question, which features to use?

There are more than 9,000 suggested features for time series! [1]

Many of these features are redundant with each other, or just useless…

Jones and Fulcher searched through all these features to find a small subset that:

- Are mostly non-redundant
- Provide discrimination between semantically different $\frac{1}{3}$ and $\frac{1}{9.72}$ and $\frac{1}{9.74}$ and $\frac{1}{9.76}$ classes in most real-world domains

The small set is called Catch-22 [2]

I think Catch-22 is one of the best ideas in time series data mining in the last decade..

- 1. Fulcher B.D., Little M.A., Jones N.S . Highly comparative timeseries analysis: the empirical structure of time series and their methods. J. Roy. Soc. Interface 10, 20130048 (2013)
- 2. Lubba C, Sethi S, Knaute P, Schultz S, Fulcher B, Jones NS (2019) catch22: CAnonical Time-series CHaracteristics. Data Min Knowl Disc 33(6):1821–1852

Unfortunately, the names of the Catch-22 features are poor, with no real mnemonic value.

- 1. Fulcher B.D., Little M.A., Jones N.S . Highly comparative timeseries analysis: the empirical structure of time series and their methods. J. Roy. Soc. Interface 10, 20130048 (2013)
- 2. Lubba C, Sethi S, Knaute P, Schultz S, Fulcher B, Jones NS (2019) catch22: CAnonical Time-series CHaracteristics. Data Min Knowl Disc 33(6):1821–1852

Lets see how this works.

Lets take a time series subsequence from Class A

Lets measure its 22 features, and summarize them with a color-coded bar chart…

Now let us compare the feature vector to two unknown instances, '1' and '2'

Which one is also in Class A?

Which one is also in Class A?

It is of course, Unknown 2

Which one is also in Class A?

It is of course Unknown 2

An important note

In this example the class happens to be preserved in *both* shape and feature.

More generally, there are time series that are conserved *only* in features.

Think of human behaviors:

Conserved in **shape**: walking/running/swimming/cycling

Conserved in **feature**: reading/watchingTV/resting/sleeping Once you have your feature vectors, you can (among *many* other things) do the analogue of Distance Profile and Matrix Profile.

Lets see a quick example:

We have a mouse walking on a circular treadmill..

Let's build a catch-22 Matrix Profile. The high values should be when there are anomalies in the data…

This works, beautifully

This works, beautifully

Summary I

- For data that has conserved *shape*, the Distance Profile and the Matrix Profile will solve 90 to 100% of your task at hand.
- For data that has conserved *features*, using catch-22 with an appropriate algorithm will get you a long way. You can do a little better sometimes:
	- Reducing the 22 features to an even smaller subset
	- Augmenting the features with one or two custom features.
- We only looked at univariate data, but you normally have many dimensions, say X, Y, Z, roll, pitch, yaw, temperature, pressure..
	- The Distance Profile and the Matrix Profile trivially generalize to multidimensions
	- However, in most cases, you can solve your problem using only one dimension.
	- Using catch22 with multidimension is possible, but less well understood.

Summary II

- I edit and review for the top data mining conferences and journals.
- I am disappointed at how rarely we see papers on livestock problems.
- However, you *can* publish such papers in top venues, I have done so.
- Please consider sending your best work to the best venues.
- If you are going to publish, share your data!
- If you have detailed questions, feel free to email me anytime.
- If you want an editable version of these slides for teaching a class, just ask.

Summary III

- Thanks to all my *great* students
- Thanks to VistaMilk SFI Research Centre for inviting me
- Thanks to Apollo

- Thank you for listening to me after your long day
- If you are interested in *insects* in AG (crop and livestock nuisance pests), I have both an academic interest, and a small company that make the best insects sensors in the world www.farmsense.io

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eamonn@cs.ucr.edu

Questions?

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Backup Slides Below

•Matrix Profile I: All Pairs Similarity Joins for Time Series: A Unifying View that Includes Motifs, Discords and Shapelets. Chin-Chia Michael Yeh, Yan Zhu, Liudmila Ulanova, Nurjahan Begum, Yifei Ding, Hoang Anh Dau, Diego Furtado Silva, Abdullah Mueen, Eamonn Keogh (2016). IEEE ICDM 2016. [\[pdf](https://www.cs.ucr.edu/~eamonn/PID4481997_extend_Matrix%20Profile_I.pdf)] [\[slides\]](https://www.cs.ucr.edu/~eamonn/matrix_profile_i.pptx)

•Matrix Profile II: Exploiting a Novel Algorithm and GPUs to break the one Hundred Million Barrier for Time Series Motifs and Joins. Yan Zhu, Zachary Zimmerman, Nader Shakibay Senobari, Chin-Chia Michael Yeh, Gareth Funning, Abdullah Mueen, Philip Berisk and Eamonn Keogh (2016). EEE ICDM 2016. [\[pdf](https://www.cs.ucr.edu/~eamonn/STOMP_GPU_final_submission_camera_ready.pdf)] [\[slides\]](https://www.cs.ucr.edu/~eamonn/icdm_stomp.pptx) **Shortlisted for best paper award.**

•Matrix Profile III: The Matrix Profile allows Visualization of Salient Subsequences in Massive Time Series. Chin-Chia Michael Yeh, Helga Van Herle, Eamonn Keogh (2016). IEEE ICDM 2016. [[pdf\]](https://www.cs.ucr.edu/~eamonn/PID4481999_Matrix%20Profile_III.pdf) [\[slides\]](https://www.cs.ucr.edu/~eamonn/matrix_profile_iii.pptx) Supporting [Page](https://sites.google.com/site/salientsubs/).

•Matrix Profile IIII: Using Weakly Labeled Time Series to Predict Outcomes. Chin-Chia Michael Yeh, Nickolas Kavantzas and; Eamonn Keogh. VLDB 2017 [\[pdf](https://www.cs.ucr.edu/~eamonn/WeaklyLabeledTimeSeries.pdf)] Munich Germany.

•Matrix Profile V: A Generic Technique to Incorporate Domain Knowledge into Motif Discovery. Hoang Anh Dau and Eamonn Keogh. [\[pdf](https://www.cs.ucr.edu/~eamonn/guided-motif-KDD17-new-format-10-pages-v005.pdf)] KDD'17, Halifax, Canada.

•Matrix Profile VI: Meaningful Multidimensional Motif Discovery. Chin-Chia Michael Yeh, Nickolas Kavantzas, Eamonn Keogh. [\[pdf\]](http://www.cs.ucr.edu/~eamonn/Motif_Discovery_ICDM.pdf) ICDM 2017.

•Matrix Profile VII: Time Series Chains: A New Primitive for Time Series Data Mining. Yan Zhu, Makoto Imamura, Daniel Nikovski, and Eamonn Keogh. [\[pdf\]](http://www.cs.ucr.edu/~eamonn/chains_ICDM.pdf) ICDM 2017. **Winner best paper award.** [\[slides\]](https://www.cs.ucr.edu/~eamonn/Slides_for_chains_website.pptx)

•Matrix Profile VIII: Domain Agnostic Online Semantic Segmentation at Superhuman Performance Levels. Shaghayegh Gharghabi, Yifei Ding, Chin-Chia Michael Yeh, Kaveh Kamgar, Liudmila Ulanova, and Eamonn Keogh. [\[pdf\]](http://www.cs.ucr.edu/~eamonn/Segmentation_ICDM.pdf) ICDM 2017.

•Matrix Profile IX: Admissible Time Series Motif Discovery with Missing Data [temp [link](https://arxiv.org/abs/1802.05472)]. Yan Zhu, Abdullah Mueen and Eamonn Keogh.TKDE 2020

Matrix Profile X: VALMOD - Scalable Discovery of Variable-Length Motifs in Data Series. Michele Linardi ,Yan Zhu ,Themis Palpanas and Eamonn Keogh. SIGMOD 2018.

•Matrix Profile XI: SCRIMP++: Time Series Motif Discovery at Interactive Speed. Yan Zhu, Chin-Chia Michael Yeh, Zachary Zimmerman, Kaveh Kamgar and Eamonn Keogh, ICDM 2018. **[**[PDF](https://www.cs.ucr.edu/~eamonn/SCRIMP_ICDM_camera_ready_updated.pdf)**]**

•Matrix Profile XII: MPdist: A Novel Time Series Distance Measure to Allow Data Mining in More Challenging Scenarios. Shaghayegh Gharghabi, Shima Imani, Anthony Bagnall, Amirali Darvishzadeh, Eamonn Keogh. ICDM 2018. [expanded version [PDF\]](https://www.cs.ucr.edu/~eamonn/MPdist_Expanded.pdf)

•Matrix Profile XIII: Time Series Snippets: A New Primitive for Time Series Data Mining. Shima Imani, Frank Madrid, Wei Ding, Scott Crouter, Eamonn Keogh. IEEE Big Knowledge 2018. [expanded version [PDF](https://www.cs.ucr.edu/~eamonn/Time_Series_Snippets_10pages.pdf)]. Matrix Profile XIV: Scaling Time Series Motif Discovery with GPUs: Breaking the Quintillion Pairwise Comparisons a Day Barrier. SoCC 2018. [[pdf\]](https://www.cs.ucr.edu/~eamonn/SCAMP-camera-ready-final1.pdf) [This paper had an interesting [history\]](https://www.reddit.com/r/datamining/comments/9egby9/i_was_denied_a_review_at_vldb/)

•Matrix Profile XV: Time Series Consensus Motifs: A New Primitive for Finding Repeated Structure in Time Series Sets. Kaveh Kamgar, Shaghayegh Gharghabi, and Eamonn Keogh (2019). IEEE ICDM 2019. [\[pdf\]](https://www.cs.ucr.edu/~eamonn/consensus_Motif_ICDM_Long_version.pdf) •Matrix Profile XVI: Time Series Semantic Motifs: A New Primitive for Finding Higher-Level Structure in Time Series. Shima Imani and Eamonn Keogh (2019). IEEE ICDM 2019. [[pdf\]](https://www.cs.ucr.edu/~eamonn/semantic_motif_icdm.pdf)

•Matrix Profile XVII:Indexing the Matrix Profile to Allow Arbitrary Range Queries. Yan Zhu, Chin-Chia Michael Yeh, Zachary Zimmerman, Eamonn J. Keogh ICDE 2020.

Matrix Profile XVIII: Time Series Mining in the Face of Fast Moving Streams using a Learned Approximate Matrix Profile. Zachary Zimmerman, Nader Shakibay Senobari, Gareth Funning, Evangelos Papalexakis, Samet Oymak, Philip Brisk, and Eamonn Keogh (2019). IEEE ICDM 2019. [[pdf](https://www.cs.ucr.edu/~eamonn/LAMP_Camera_Ready2.pdf)]

•Matrix Profile XIX: Efficient and Effective Labeling of Massive Time Series Archives. Frank Madrid, Shailendra Singh, Quentin Chesnais, Kerry Mauck and Eamonn Keogh. DSAA 2019: International Conference on Data Science and Advanced Analytics [[pdf\]](https://www.cs.ucr.edu/~eamonn/Efficient%20and%20Effective%20Labeling%20of%20Massive%20Entomological%20Datasets.pdf).

•Matrix Profile XX: Finding and Visualizing Time Series Motifs of All Lengths using the Matrix Profile. Frank Madrid, Shima Imani, Ryan Mercer, Zacharay Zimmerman, Nader Shakibay, Eamonn Keogh. IEEE Big Knowledge 2019 [\[pdf](https://www.cs.ucr.edu/~eamonn/PAN_SKIMP%20%28Matrix%20Profile%20XX%29.pdf)]

•Matrix Profile XXI: MERLIN: Parameter-Free Discovery of Arbitrary Length Anomalies in Massive Time Series Archives. Takaaki Nakamura, Makoto Imamura, Ryan Mercer and Eamonn Keogh. ICDM 2020 [[pdf\[](https://www.cs.ucr.edu/~eamonn/MERLIN_Long_version_for_website.pdf) Matrix Profile XXII: Exact Discovery of Time Series Motifs under DTW. Sara Alaee, Ryan Mercer, Kaveh Kamgar, Eamonn Keogh. ICDM 2020 [[pdf](https://www.cs.ucr.edu/~eamonn/DTWMotifsSWAMP_Expanded.pdf)]

•Matrix Profile XXIII: Contrast Profile: A Novel Time Series Primitive that Allows Real World Classification. Ryan Mercer, Sara Alaee, Alireza Abdoli, Shailendra Singh, Amy Murillo, Eamonn Keogh. ICDM 2021 [[pdf\]](https://www.cs.ucr.edu/~eamonn/ContrastProfile_V048_names.pdf) •Matrix Profile XXIV:Scaling Time Series Anomaly Detection to Trillions of Datapoints and Ultra-fast Arriving Data Streams. Yue Lu, Renjie Wu, Abdullah Mueen, Maria A. Zuluaga and Eamonn Keogh. ACM SIGKDD 2022 [[pdf\]](https://www.cs.ucr.edu/~eamonn/DAMP_long_version.pdf) •Matrix Profile XXV: Introducing Novelets: A Primitive that Allows Online Detection of Emerging Behaviors in Time Series. Ryan Mercer and Eamonn Keogh. ICDM 2022. [[pdf\]](https://www.dropbox.com/sh/qtmb8uxqntpt1e4/AAABdxjushcLJRw4_ppDTwuYa/Novelets_V0029_CameraReady.pdf?dl=0)

•Matrix Profile XXVI: Mplots: Scaling Time Series Similarity Matrices to Massive Data. Maryam Shahcheraghi, Ryan Mercer, João Manuel de Almeida Rodrigues, Audrey Der, Hugo Filipe Silveira Gamboa, Zachary Zimmerman and Eamonn Keogh. ICDM 2022.[[pdf\]](https://www.dropbox.com/sh/qtmb8uxqntpt1e4/AAD6YfOpVLf0C4PuGxMNXvJ-a/MPLOT_V0044.pdf?dl=0)

•Matrix Profile XXVII: A Novel Distance Measure for Comparing Long Time Series. Audrey Der, Chin-Chia Michael Yeh , Renjie Wu , Junpeng Wang , Yan Zheng , Zhongfang Zhuang, Liang Wang , Wei Zhang, Eamonn Keogh. ICKG 2022.

Best match in 2D space

Pressure

Orientation (Magn)

I have 262,144 data points that record a penguin's orientation (MagX) and the water/air pressure as he hunts for fish. **Question**: Does he ever change his bearing leftwards as he reaches the apex of his dive. This is easy to describe as a multidimensional search. The apex of a dive is just an approximately parabolic shape. I can create this with $_{query_pressure}$ = zscore([[-500:500].^2]*-1)'; it looks like this I can create *bearing leftwards* with a straight rising line, like this query_MagX = zscore([[-500:500]])'; It looks like this.

> bumps? They are wingstokes as the bird "flies" underwater

We have seen elsewhere in this document how to search for a 1D pattern. For this 2D case, all we have to do is add the two distance profiles together, before we find the minimum value. Note that the best match location in 2D is different to either of the 1D queries.

We can do this for 3D or 4D…

However, there are some caveats. In brief, it almost never makes sense to do multidimensional time series search in more than 3 or 4D. See Matrix Profile VI: M. Yeh ICDM 2017 and "Weighting" B. Hu, ICDM 2013. In addition, in some cases we may want to weight the dimensions differently, even though they are both z-normalized Euclidean Distance. What are the periodic

load penguintest.mat figure;, hold on;

```
query pressure = zscore([[-500:500].<sup>^2]*-1</sup>)';
dist p = MASS V2(penguintest(:,1), query pressure);
query MagX = zscore([[-500:500]])';
dist \overline{m} = MASS V2(penguintest(:,2), query MagX);
[val,loc] = min([dist m + dist p]); % find best match location in 2D
plot(zscore(penguintest(loc:loc+length(query_MagX),2)),'color',[0.85 0.32 0.09])
plot(zscore(query_MagX),'m') 
plot(zscore(penguintest(loc:loc+length(query_pressure),1)),'b')
plot(zscore(query_pressure),'g') 
title(['Best matching sequence, pressure/MagX, is at ', num2str(loc)])
```


Let us run the Matrix Profile, looking for four-second long motifs…

- >> load eog_sample.mat
- >> [matrixProfile profileIndex, motifIndex, discordIndex] = interactiveMatrixProfileVer3 website(eog_sample, 400);

The code takes a while to fully converge, but in just a few seconds, we see some stunningly well conserved motifs…

Having found the motifs, we can ask, what are they? A quick glance at a paper by Noureddin et. al. locates a very similar pattern (with some time warping) called eye-blinkartifact.

Note that there may be more examples of each motif. We should take one of the above, and use MASS to find the top 100 neighbors… See *Have we ever seen a pattern that looks just like this?*. We can also adjust the range parameter *r* inside the motif extraction code.

What are the three most unusual days in this three-month long dataset?

The datasets is Taxi demand, in New York City, in the last three months of the year.

We choose 100 datapoints, which is about *two* days long (the exact values do not matter much here).

- >> load taxi 3 months.txt
- >> [matrixProfile, profileIndex, motifIndex, discordIndex] = interactiveMatrixProfileVer3 website(taxi 3 months ,100);

The code pops up the matrix profile tool, and one second later, we are done! The three most unusual days correspond to the three highest values of the matrix profile (i.e. the *discords*), but what are they?

- The highest value corresponds to Thanksgiving
- We find a secondary peak around Nov 6th, what could it be? Daylight Saving Time! The clock going backwards one hour, gives an *apparent* doubling of taxi load.
- We find a tertiary peak around Oct 13th, what could it be? Columbus Day! Columbus Day is largely ignored in much of America, but still a big deal in NY, with its large Italian American community.

Is there any pattern that is common to these two time series?

Lets assume that the common pattern is 3 seconds, or 300 datapoints long. Let us concatenate the two time series, and smooth them (just for visualization purposes, we don't really need to) Now let us find the top motif, but insist that one motif comes before 24289, and one after…

The top join motif shows a highly conserved pattern.

It is the famous bass line $\frac{1}{2}$. If $\frac{1}{2}$ from *Under Pressure* by Queen which was plagiarized by Vanilla Ice.

The concept for this example comes from Dr. Diego Furtado Silva.

Find the most conserved pattern that happens at least once every two days in this dataset

The question is a little underspecified, as the length for the conserved patterns was not given. Let us try two hours, which is about 800 data points.

The full 20,000 datapoints represents about 14 days of electrical demand data for a house in the U.K. Thus we first need to divide it into approximate 2 day chunks.

>> load TwoWeekElectrical

```
>> seven two day chunks = divide data(T);
```
Now we just need to call the consensus motif code.

```
>> consensus motifs = consensusMotifs(seven two day chunks,800); % 800 is the length of subsequence
```
The code returns the seven time series below. Note that the basic pattern is highly conserved, given how noisy the data is. The similarity between the items can be better seen if we cluster the time series with a single linkage dendrogram.

If you had to summarize this long time series with just two shorter examples, what would they be?

The dataset is 3 years of Italian power demand data which represents the hourly electrical power demand of a small Italian city for 3 years beginning on Jan $1st$ 1995.

an aik ma dalam dia aktifa 1940 (1940) (1940) (1940) (1940) (1940) (1940) (1940) (1940) (1940) (1940) (1940) (1 i ait eanaananan an - - - - ana**un an ta'a am an ta'a am am am am a** an tan an am an - - - an an am am am am am a 6 MF Br. Tha ON the market off Officer entrance the annual

Jan/1/1995 May/31/1998

We just need to call Time Series Snippets algorithm…

```
>> load('ItalianPowerDemand.mat')
```

```
>> [fraction, snippet, snippetidx]= snippetfinder(data(:, 4), 2, 200, 30);
```
It will pop open three windows, which are snippet 1, snippet 2 and the regime bar.

We searched for the top-2 snippets of length 200. This was our quick "eyeballing" guess as to the length of a week, but it is actually about 8.3 days. Note that the snippets are not align to start at the same day of the week (this is a trivial constraint to add if desired).

We obtain the "regime bar," which tells us which snippet "explains" which region of data. As it happens, Snippets seem to represent *summer* and *winter* regimes respectively.

Are there any patterns that appear as time reversed versions of themselves in my data?

Lets us load the data, and concatenate it to itself, after flipping left to right. We can then search for a join motif, that spans 5046, the length of the original time series. If we find a good join motif, it means that the conserved pattern is time reversed!

```
>> load('mfcc.mat')
\gg length (mfcc1(1,:))
   = 5046>> interactiveMatrixProfileAB(([mfcc1(1,:)'; flipud(mfcc1(1,:)')]), 150, 5046); % This will spawn this plot ->
```
The top join motif shows a highly conserved pattern. Why would a pattern occur time reversed?

play the music 'backward' the second time around". MFC8"Symphony "*The most extraordinary of all canonic movements from this* time is of course from Symphony No. 47. Here Haydn writes out **14:53** The data is the 1st MFCC of this piece of music. ¹ ¹⁵⁰*only one reprise of a two-reprise form, and the performer must*

0 1000 2000 3000 4000 5000

In this dataset, at time stamp 7,500, bleeding was induced in an otherwise healthy pig. This changes the pig's APB measurement, but only *very* slightly. Could we find the location of the change, if we were not told it? Moreover, can we do this with no domain knowledge? In other words, can we detect regime changes in time series?

```
>> TS = load('PigInternalBleedingDatasetArtPressureFluidFilled 100 7501.txt');
>> CAC = RunSegmentation(TS, SL); %SL is the length of subsequence
>> plot(CAC,'c')
\Rightarrow [\sim, loc] = min(CAC) %value of loc is 7460 which is the approximation of exact value 7500
```
Here, we choose SL to be 100, approximately the length of one period of arterial pressure (or the period of whatever repeated patterns you have in your data), however, up to half or twice that value would work just as well. The output curve, the CAC, minimizes at just the right place. How does it do it? In brief, if we examine the pointers in the Matrix Profile Index, we will find that very few will cross *over* the location of a regime change (most healthy beats have a nearest neighbor that is another healthy beat, most "bleeding" beats have a nearest neighbor that is another "bleeding" beat), it is this lack of pointers that cross over the regime change that is what the CAC is measuring.

Are there any patterns that repeat in my data, but at two distinct lengths?

See also "*Is there any pattern that is common to these two time series?*"

We can solve this with a quick and dirty trick. The code interactiveMatrixProfileAB(T,m, crossover) searches time series T for a motif of length m, such that one of the motif pair occurs before \csc{v} and one occurs after \csc{v} We can take a time series and append it to a *rescaled* copy itself, setting the to the length of the original time series. Now when we find motifs, we are finding one at the original scale, *and* one at the rescaled size.

Voltage reading

In this case, I want to know if any of my insect behaviors happens at length 5,000 and at 10,000, so I type…

```
>> load insectvolts.mat % load some insect epg data
```
>> interactiveMatrixProfileAB([insectvolts ; insectvolts(1:2:end)], 5000, length(insectvolts)); % search the appended data

No need to let it converge, after a few seconds we have our answer…

How can I optimize similarity search in a long time series?

see also "*How do I quickly search this long dataset for this pattern, if an approximate search is acceptable?*"

Suppose you want to find a query inside a long time series, say of length 67,000,000.

First trick: MASS (and several other FFT and DWT ideas) have their best case when the data length is a power of two, so pad the data to make it a power of two (padding with zeros works fine).

Second trick: MASS V3 is a piecewise version of MASS that performs better when the size of the pieces are well aligned with the hardware. You need to tune a single parameter, but the parameter can only be a power of two, so you can search over say 2^{10} to 2^{20} . Once you find a good value, you can hardcode it for your machine.

What is the right length for motifs in this dataset?

This is a very interesting question, which more than most, deserves a long explanation. However, to be brief and pragmatic. Let us revisit the EOG dataset. Recall that we choose 4 seconds as the motif length, which I happen to know (from reading papers on the topic) is a good choice.

```
>> load eog_sample.mat
```

```
>> [MP profileIndex, motifIndex, discordIndex] = interactiveMatrixProfileVer3 website(eog_sample, 400);
Let us look at the Matrix Profile, and the top motif we find (bottom left). The results seem to make sense.
```
However, suppose in contrast that we knew nothing about the domain, and had chosen a motif length that was much too long, say length 3,000 (bottom right). How could we know that we had picked a length that was too long? There are two clues:

- Obviously, the motifs themselves will be less well conserved visually.
- The Matrix Profile itself offers useful clues. It tells us how "specially well conserved" the motif is ($min(MP)$) relative to the *average* subsequence ($mean(MP)$). As the ratio of these two numbers is approaches zero, it suggests a stunningly well conserved motif in the midst of others unconserved data. However, as the ratio of these two numbers is approaches one, it suggest that the "motif" is no better conserved than we would expect by random chance. In practice, we rarely compute these ratios, as is visually obvious that the MP looks "flat".

I need to find motifs faster! Part I

Part of the solution might be to use GPUs, see [a][b].

Moreover, it is important to understand, we almost *never* need to compute the Matrix Profile to completion, the anytime SCRIMP++ converges so fast, that in general we just run it 1% (or less) of convergence.

Nevertheless, sometimes you might want to compute the converged Matrix Profile. There is a faster algorithm for this. It exploits some of the ideas in [b], and it exploits the fact that it does not need to waste the overhead needed to make anytime updates, to achieve about an order of magnitude speedup.

For consistency with our other tools, when the fast code finish, it pops open the same plot.

Have we ever seen a pattern that looks just like this, but possibly at a different length?

www.physionality.com/www.com/www.com/www.com/www.com/www.com/www.com/www.com/www.com/www.com/www.com/

In our insect data, a basic feeding primitive looks like this: // we can model it with something like: [1:600].^0.2

We have a theory that a certain higher level behavior will result in "A long primitive, followed by a shorter and smaller primitive, followed by another long primitive", like this.. ye can model it with: [[[1:600].^0.2] [[1:300].^0.2] [[1:600].^0.2]]

However, we don't know how long the whole pattern could be...

The function to the right can solve this problem. It simply brute forces a MASS test for all lengths within a range (here 100 to 300%) at a given step size (here 5%). There may be faster techniques, but MASS is so fast, they may not be worth bothering with. A critical trick is to *normalize* the comparisons at

different lengths (see Appendix).

>> load insectvolts.mat % load some insect epq data >> query = $([[[1:600].^0.2] [[1:300].^0.2] [[1:600].^0.2]])$; >> uniform scaling search (smooth (insectvolts, 10), query);

Even with this unoptimized approach. We can search two hours of data (at 100hz), with a long query (upto to 4,500 datapoints), in well under 10 seconds

This looks like a lot of code, but most of it is for plotting

plot(zscore(TAG(best_match_loc:best_match_loc+length(QUERY(1:100/best_match_scale:end))-1)),'g'); set(gca, 'Xlim', [1 length(QUERY(1:100/best match scale:end))]); title(['The best match is found when we rescale to ', num2str(best match scale), '%']);

2.017 2.018 2.019 2.02 2.021

Khale

Normal **Breathin**

This is a dataset of respiration from a sleep study. Each breath appears to be about 360 data points long. So lets search *for time series chains* of length 360…

```
>> load respiration.mat
>> TSC1 demo(respiration, 360);
```
The algorithm finds the highlighted chains below.

Note the increasing "gulp" artifact that happens between cycles. Also note that it begins to happen earlier and earlier in the cycle. What does this mean?

Here is the (lightly edited) annotation of Dr. Gregory Mason (LA BioMed/UCLA) an expert on cardiopulmonary interactions. *"The gulps are attempts to inspire against an obstruction coming the back of the tongue. The large signals are from the machine which do not necessarily reach the patient, the small gulps are pathologic attempts to breathe. Why does it increase? With each successive breath the patient tries harder to inspire. It finally is 'synchronized' and you don't see the small patient signal, and this event cycles over and over. The cycling is best seen without treatment if one looks up "crescendo snoring," a hallmark of obstructive sleep apnea."*