Data Reduction Techniques for Sensor Networks

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Abstract

We are inevitably moving into a realm where small and inexpensive wireless devices would be seamlessly embedded in the physical world and form a wireless sensor network in order to perform complex monitoring and computational tasks. Such networks pose new challenges in data processing and dissemination due to the conflict between (i) the abundance of information that can be collected and processed in a distributed fashion among thousands of nodes and (ii) the limited resources (bandwidth, energy) that such devices possess. In this paper we propose a new data reduction technique that exploits the correlation and redundancy among multiple measurements on the same sensor and achieves high degree of data reduction while managing to capture even the smallest details of the recorded measurements. The key to our technique is the *base signal*, a series of values extracted from the real measurements, used for encoding piece-wise linear correlations among the collected data values. We provide efficient algorithms for extracting the base signal features from the data and for encoding the measurements using these features. Our experiments demonstrate that our method by far outperforms standard approximation techniques like Wavelets, Histograms and the Discrete Cosine Transform, on a variety of error metrics and for real datasets from different domains.

1 Introduction

Technological advances in the development of low-power embedded communication devices have made possible scenarios in which thousands of sensor nodes could be seamlessly embedded in the

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physical world and form a wireless sensor network. These sensors would monitor various quantities such as temperature, pressure, humidity, movement, noise levels, chemicals, etc, that would then be periodically transmitted to a *base-station*¹ for further processing and analysis. Applications of such networks span a large variety of domains from collaborative environments to military command and control systems and even home networks.

Large-scale sensor networks require tight data handling and data dissemination techniques. Transmitting a *full-resolution* data feed from each sensor back to the base-station, is often prohibitive due to (i) limited bandwidth that may not be sufficient to sustain a continuous feed from all sensors and (ii) increased power consumption due to wireless multi-hop communication.

In order to minimize the volume of data transmitted, we can apply two well known ideas: aggregation and approximation. Aggregation works by summarizing the measured information in the form of simple statistics like average, maximum, minimum etc that are then transmitted to the base-station over regular intervals. Aggregation is an effective mean to reduce the volume of data, but can be rather crude for applications that need detailed historical information, e.g. military surveillance. When data feeds exhibit a large degree of redundancy, approximation is a less intrusive form of data reduction in which the underlying data feed is replaced by an approximate signal tailored to the application needs. The tradeoff is then between the size of the approximate signal and its precision compared to the real-time information monitored by the sensor.

In this paper we present a new data reduction algorithm for the dissemination of approximate measurements over sensor networks. Our techniques build on the observation that the values of the collected measurements exhibit similar patterns over time, or that different measurements are naturally correlated, as is the case between pressure and humidity. At the core of our approximation lies the notion of a *base signal*, a set of values from the collected measurements that capture prominent features of the data. Following the construction of the *base signal*, the collected data is partitioned into intervals that can be efficiently approximated as linear projections of some part of the *base signal*. As we will show in this paper, our techniques provide:

• Increased accuracy when compared to other approximation techniques for the same reduction factor.

¹A base-station may represent any node of the network with increased storage, battery and processing capabilities.

• Adaptability to different error metrics: Our algorithms can be adapted with only minor modifications, which do not alter their time complexity, to minimize different error metrics, such as the sum squared error, sum squared relative error, and maximum error of the approximation.

Our contributions are summarized as follows:

1. We introduce a new approximation scheme that encodes piece-wise correlations among the data values. Such correlations are often linear in nature and can be easily captured by standard techniques like linear regression. We exploit correlations both within the values of a single measurement (ex: periodicity, self-similarity) as well as among values of different quantities (ex: pressure and humidity).

2. We introduce the concept of the *base signal* that is analogous to a *carrier-wave* in radio-frequency transmissions and is used for encoding the measurements. We explore the technical challenges of (i) constructing the base signal, (ii) approximating the recorded measurements by exploring piece-wise correlations amongst them and the base signal, and (iii) dynamically updating the base signal to capture new data trends in subsequent transmissions.

3. We provide an efficient algorithm (Self-Based Regression or SBR) that answers all questions above, while balancing the cost of transmitting new (or updated) base signal values with the gains of using them for approximating the data values. For a dataset containing n measurements to approximate, the SBR algorithm takes $O(n^{1.5})$ time and requires linear space, while its running time scales linearly to the size of both the transmitted data and the base signal.

4. We provide an extensive experimental study of our framework using real datasets from different application domains and make direct comparisons against previously studied approximation techniques like the Wavelet and Discrete Cosine transforms and Histograms. In all datasets our method achieves substantially lower approximation errors for the same data reduction factor.

5. We have adapted ideas from the Singular Value Decomposition and the Discrete Cosine transform for constructing alternative base signals. Our experiments demonstrate that the base signal features selected by SBR outperform these techniques. Furthermore, we show that SBR makes near-optimal choices when selecting the number of features to include in the base signal.

The rest of the paper is organized as follows. Section 2 presents related work. In Section 3

we state our problem and sketch the basics of our techniques, while in Section 4 we describe our framework in more details. Section 5 contains our experiments, while Section 6 contains concluding remarks and future directions.

2 Related Work

In recent years there has been a flurry of research in the area of sensor networks. Some of the most important issues addressed include network self-configuration [4], data discovery [9, 12] and in-network query processing [14, 10, 30, 17].

The benefits of in-network data aggregation are investigated in [14, 17, 30]. The main idea is to build an aggregation tree which the results will follow. Non-leaf nodes of the tree aggregate the values of their children before transmitting the aggregate result to their parents. In [17] additional issues are also addressed, such as determining when a node becomes $active^2$ and designing query processing techniques for aggregates with different characteristics.

Sensor nodes are small devices that "measure" their environment and communicate streams of low-level values to a base station for further processing and archiving. These streams are then used to construct a higher-level model of the environment. This process makes historical data equally important to current values [8]. In this paper we propose approximation as a less intrusive data reduction method that is more suited for applications in which a long-term historical record of measurements from each sensor is required.

Recently, there has been increasing interest in studying the general principles over continuous queries in data streams [6, 13, 20, 28, 31]. Olston et. al in [21, 2] study the tradeoff between precision and performance when querying replicated, cached data. In [3] the users register continuous queries with strict precision constraints at a central *stream processor*, which, in turn installs filters at the remote data sources. These filters adapt to changes in the streams to minimize update rates. An online algorithm for minimizing the update cost while the query can be answered within an error bound is presented in [26]. The authors of [25] study a probabilistic query evaluation method that places appropriate confidence in the query answer to quantify the *uncertainty* of the recorded data

values.

 $^{^{2}}$ An active node can receive, process and transmit data. At this mode the sensor drains significantly more energy than when it is *idle*.

Approximate processing techniques have been widely studied. Histograms (e.g. [22, 24]) have been extensively used by query optimizers to estimate the selectivity of queries, and recently in tools for providing fast approximate answers to queries. Wavelets are a mathematical tool for the hierarchical decomposition of functions, with applications in image and signal processing. More recently, Wavelets have been applied successfully in answering range-sum aggregate queries over data cubes [29], in selectivity estimation [19] and in approximate query processing [5]. The Discrete Cosine Transform (DCT) [1] constitutes the basis of the *mpeg* encoding algorithm and has also been used to construct compressed multi-dimensional histograms [16]. Linear regression has been recently used in [7] for on-line multidimensional analysis of data streams.

3 Preliminaries

3.1 Characteristics of Sensor Networks

Recent technological advances have made possible the development of low-cost sensor nodes with heavily integrated sensing, processing and communication capabilities. Networked together in an ad-hoc fashion, hundreds of such nodes can be used for a variety of monitoring applications such as military surveillance, equipment monitoring or medical sensing.

Information about the environment is gathered using a series of sensing elements connected to an analog-to-digital converter. Examples include microphones for acoustic sensing, accelerometers, temperature sensors etc. Once enough data is collected, it is processed locally and periodically forwarded to a base station, using a multi-hop routing protocol [27].

The processing subsystem on the nodes depends on the nature of the application. Applications such as military reconnaissance that require significant processing to be performed at the nodes use sensor nodes with significant processing power. As an example, an improved model of the commonly used StrongARM 1100 processor (μ AMPS [27] and HiDRA nodes) reaches a frequency of 400 MHz and can support up to 64 MB of memory.

As the processing and storage capabilities of sensor nodes tend to follow Moore's Law their communication and power subsystems become the major bottleneck of their design. For example, over the last years, the energy capacity of the batteries used in such nodes has exhibited a mere 2-3% annual growth.³ The main source of energy consumption in a node is the data transmission process. There are several reasons for this:

The energy drain during transmission is much larger than the consumption during processing [9].
 As an example, on a Berkeley MICA Mote sending one bit of data costs as much energy as 1,000
 CPU instructions [18].

2. Transmission ranges between nodes are fairly short. The transmitted data may thus require to traverse multiple hops to reach the base station. This retransmission process at each intermediate node is very costly.

3. Nodes often use broadcast protocols over radio frequencies [17]. Due to the high density of nodes, transmitted messages are not only received by the intended node, but by all nodes in the vicinity of the sender, thus increasing the overall power consumption.

Even on applications where battery lifetime is not a concern (ex: military surveillance sensing nodes attached to moving vehicles with practically infinite power supply) available bandwidth may not sustain a continuous feed of measurements for all sensors deployed in the terrain. The design of data reduction protocols that effectively reduce the amount of data transmitted in the network is thus essential when the goal is to meet the application's bandwidth constraints or to increase the network's lifetime.

3.2 Data Model and Processing

In order not to deplete their power supply (and to conserve bandwidth), the sensors do not continuously transmit every new measurement they take but rather wait till enough data is collected and then forward it to the base station [27]. This form of batch processing allows them to power-down their radio transmitter and prolong their lifetime in a way analogous to [17].

Within a sensor, the recorded data is depicted in a two dimensional array where each row i stores sampled values of a distinct quantity. Informally, each row i is a time series $\vec{Y_i}$ of samples from quantity i collected by the sensor. The array has N rows, N being the number of recorded quantities and M columns, where M depends on the available memory.⁴

³http://nesl.ee.ucla.edu/courses/ee202a/2002f/lectures/L07.ppt

⁴We here assume that all quantities are sampled with the same frequency. This simplifies notation, however, our framework also applies when each quantity is recorded on a different schedule.



Figure 1: Transfer of approximate data values and of the base signal from each sensor to the base station

As more measurements are obtained, the sensor's memory buffers become full. At this point the latest $N \ge M$ values are processed and each row i (of length M) is approximated by a much smaller set of B_i values, i.e. $B_i \ll M$. The resulting "compressed" representation, of total size equal to $B = \sum_{i=1}^{N} B_i$, is then transmitted to the base station. The base station maintains the data in this compact representation by appending the latest "chunk" to a log file. A separate file exists for each sensor that is in contact with the base station. The entire process is illustrated in Figure 1.

Each sensor allocates a small amount of memory of size M_{base} for what we call the base signal. This is a compact ordered collection of values of prominent features that we extract from the recorded values and are used as a base reference in the approximate representation that is transmitted to the base station (details will be given in the next section). The data values that the sensor transmits to the base station are encoded using the in-memory values of the base signal at the time of the transmission. The base signal may be updated at each transmission to ensure that it will be able to capture newly observed data features and that the obtained approximation will be of good quality. When such updates occur they are transmitted along with the data values and appended in a special log file that is unique for each sensor. This allows the base station to reconstruct (approximately) the series \vec{Y}_i at any given point in the past.

3.3 Our Optimization Problem

We can think of the base signal as a dictionary of features used to describe the data values. The richer the pool of features we store in the base signal the better the approximation. On the other

Configuration Parameters						
Ν	Number of input signals					
М	Measurements per input signal					
Input Parameters						
TotalBand	Total bandwidth per transmission					
M_{base}	Buffer size for base signal values					
Derived/Calo	culated Parameters					
$n = N \times M$	Size of in-memory data					
$W = \sqrt{n}$	Size of each base interval					
В	Compressed Data Size					
maxIns	Maximum number of base intervals					
	inserted in current transmission					
Ins	Number of base intervals actually					
	inserted in the current transmission					

Table 1: Configuration, input and derived parameters of our algorithms

hand, these features have to be (i) kept in the memory of the sensor to be used as a reference by the reduction algorithm and (ii) sent to the base station in order for it to be able to reconstruct the values. Thus, for a target bandwidth constraint (number of values that can be transmitted) the more insert and update operations on the base signal that we perform, the less bandwidth that is left available for approximating the data values. Moreover, the time to perform the data approximation increases, in our algorithms, linearly with the size of the base signal.

In the next section we present an efficient algorithm that decides (i) how large the base signal needs to be at each transmission (ii) what new features to be included in it (iii) which older features are not relevant any more and (iv) how to best approximate the data measurements using these features. The only user input needed by the algorithm is the target bandwidth constraint and the maximum buffer size of the base signal values.

4 The SBR Data Reduction Framework

We now describe our framework in more detail. We start with a motivational example that demonstrates the intuition behind our techniques. Subsection 4.2 presents the primitive operations required by our framework while the *SBR* algorithm is presented in subsection 4.3. Table 1 contains a brief description of the parameters used in our algorithms.



Figure 2: Example of two correlated signals (Stock Market)



Figure 3: XY scatter plot of Industrial (X axis) vs Insurance (Y axis)

4.1 Motivational Example

Many real signals are correlated. We expect this to be particularly true for measurements taken by a sensor, especially if they are physical quantities like temperature, dew-point, pressure etc. The same is often true in other domains. For example, in Figure 2 we plot the average Industrial and Insurance indexes from the New York stock market for 128 consecutive days.⁵ Both signals show similar trends, i.e. they go up and down together. Figure 3 depicts a XY scatter plot of the same values. This is created by pairing values of the Industrial (X-coordinate) and Insurance (Y-coordinate) indexes, of the same day, and plotting these points in a two-dimensional plane. The strong correlation among these values makes most points lie on a straight line. This observation motivates our work. Assuming that the Industrial index (call it \vec{X}) is given to us in a time-series of 128 values, we can approximate the other time-series (Insurance: \vec{Y}) as:

$$\vec{Y'} = a * \vec{X} + b$$

The coefficients a and b are determined by the condition that the sum of the square residuals, or equivalently the L_2 error norm $||\vec{Y'} - \vec{Y}||_2$, is minimized. This is nothing more than standard linear regression. However, unlike previous methods, we will not attempt to approximate each time-series independently using regression. In Figure 2 we see that the series themselves are not linear, i.e. they would be poorly approximated with a linear model. Instead, we will use regression

⁵Data at http://www.marketdata.nasdaq.com/mr4b.html

to approximate piece-wise correlations of each series to a base signal that we will choose accordingly. In the example of Figure 3 the base signal can be the Industrial index (\vec{X}) and the approximation of the Insurance index will be just two values (a, b). In practice the base signal may be much smaller than the complete time series, since it only needs to contain the "important" trends of the target signal \vec{Y} . For instance, in case \vec{Y} is periodic, a sample of the period would suffice. Our algorithm breaks the latest measurements obtained by the sensor into small intervals (of varying sizes) and looks for intervals of the same length in the base signal that are linearly correlated. At the same time, the base signal values are evaluated and may get updated with features from the newly collected measurements when necessary.

4.2 **Primitives of our Implementation**

Piece-wise Approximation of Measurements

We here assume that the base signal \vec{X} is given to us. We will approximate the latest $N \times M$ measurements in $\vec{Y}_1, \ldots, \vec{Y}_N$ using $B \ge 4 \times N$ values. We later describe how to construct the base signal.

To simplify notation, we model the collected data as a single series \vec{Y} that is simply the concatenation of the N series $\vec{Y_i}$. Our technique relies on breaking \vec{Y} into B/4 intervals and "mapping" each one to an interval of the base signal of equal length.⁶ The algorithm works recursively. It starts with a single interval for each row of the collected data. In each iteration, the interval with the largest error in the approximation is selected and divided in two halves, until the "budget" of B/4 intervals is exhausted. An interval I is a data structure with six entries:

- start, length: these define the scope of the interval; i.e. I represents values of Y[i], with i in [start, start + length).
- *shift*: it defines the part of the base signal that is used to approximate the values of *I*; the interval *I* is mapped to segment [*shift*, *shift* + *length*) in \vec{X} .
- *a*, *b*, *err*: the first two are the regression parameters, while *err* is the sum squared error (*sse*) of the approximation.

Subroutine Regression() shown in Algorithm 1 lies in the core of our method. This function

⁶This mapping requires 4 values per interval, thus the division by 4.

Algorithm 1 Regression Subroutine

Algorithm 2 BestMap Subroutine

Require: \vec{X} , \vec{Y} , Interval I, W 1: I.shift = -12: Perform standard linear regression on I and set the values of I.a, I.b and I.err 3: if $I.length \leq 2 \times W$ then {Shift I over \vec{X} and find segment for which} 4: {regression error is minimized} 5:for shift in $0..length(\vec{X}) - I.length - 1$ do 6: $(a,b,err) = \text{Regression}(\vec{X}, \vec{Y}, shift, I.start, I.length)$ 7: if err is minimum error so far then 8: Update values of I.a, I.b, I.err and I.shift 9: end if 10:end for 11: 12: end if

pairs a segment of the base signal between values $[start_x, start_x + length)$ with values of Y between $[start_y, start_y + length)$, as in Figure 3, and computes the regression parameters a, b as well as the (sse) error of the approximation $\vec{Y'} = a\vec{X} + b$ in this range. Each value Y[i] with index i in $[start_y, start_y + length)$ is approximated as $aX[start_x + i - start_y] + b$.

It should be noted that the Regression() subroutine calculates the optimal a,b values that minimize the sum squared error of the approximation. If the desired error metric is different, then the formulas need to be appropriately modified. In the Appendix we present the necessary modifications for two interesting optimization problems: minimizing the sum squared relative error, and minimizing the maximum absolute error of the approximation. The modified algorithms run in O(length) time and require O(1) and O(length) space, respectively.

Subroutine BestMap() of Algorithm 2 looks for the best way to approximate an interval I.

It shifts I over \vec{X} and calculates the regression parameters and the approximation error for the *shift* parameter that produces the smallest error. This algorithm contains two deviations from our previous discussion. First, it also considers approximating each interval I using standard linear regression, and uses a negative value for the I.shift parameter to denote this. Second, it performs the shifting process over the base signal only for intervals with a maximum length of $2 \times W$, where W is a parameter that denotes the length of the intervals that constitute the base signal.⁷ The last modification is performed both to reduce the time complexity of the algorithm to $O(I.length + W \times M_{base})$, and because of the reduced likelihood that large intervals will be accurately mapped to multiple consecutive intervals of the base signal.

The core approximation algorithm GetIntervals() is given in Algorithm 3. The approximation obtained is returned as a list of B/4 intervals in *iJist*. This list is maintained sorted (priority queue) based on the sse of each interval. \vec{X} is the current base signal. The complete algorithm runs in $O(NMlog(\frac{B}{N}) + B \times M_{base} \times W)$ time. The logarithmic factor in the above formula is produced because the size of the intervals in the algorithm is repeatedly halved.

For each interval in i_list a record with four values (*I.start*, *I.shift*, *I.a*, *I.b*) is transmitted to the base station. The base station will sort the intervals based on *I.start* and, thus, there is no need to transmit their length. It is interesting to note that the **GetIntervals**() algorithm decides dynamically how many intervals it will use to approximate each of the N rows of the collected data, allocating more intervals to signals that are harder to approximate accurately.

Selecting Data Features for Inclusion in the Base Signal

We focus on the time when the sensor's memory is filled with $N \ge M$ values, as depicted in Figure 1. We assume that the buffer allocated to the base signal is of size M_{base} . This buffer is organized as a list of intervals (called *base intervals*) of the same length W. For simplicity, we assume that both M and M_{base} are multiples of W. We note here that in Algorithm 3 the base signal is presented as a series of M_{base} values, which is simply the concatenation of the base intervals in the buffers.

The GetBase() algorithm (Algorithm 4) lies in the core of the initialization and update procedure of the base signal. The algorithm receives as inputs the N signals, each of size M, the size W of each base interval, and the maximum number of intervals maxIns that can be inserted

⁷This will become more clear later in our discussion.

Algorithm 3 GetIntervals Algorithm

Require: $\vec{X}, \vec{Y_1}, \ldots, \vec{Y_N}, B, W$ 1: $i_list = ()$ 2: $\vec{Y} = concat(\vec{Y_1}, \dots, \vec{Y_N})$ {Virtual assignment} 3: {Create an interval for each row \vec{Y}_i (*M* values each)} 4: for i in 1..N do $(I.start, I.length) = ((i-1) \times M, M)$ 5: $\operatorname{BestMap}(\vec{X}, \vec{Y}, I, W)$ 6: $i_list.push(I);$ 7: 8: end for 9: $num_intervals = N$ 10: while $num_intervals++ < B / 4$ do 11: $\{i_list \text{ is sorted on decreasing order of } I.err\}$ $I = i_list.pop()$ 12:{Break I in 2 pieces} 13: $(I_{left}.start, I_{left}.length) = (I.start, I.length/2)$ 14:BestMap $(\vec{X}, \vec{Y}, I_{left}, W)$ 15: $(I_{right}.start, I_{right}.length) =$ 16:(I.start+I.length/2, I.length/2) $BestMap(X, Y, I_{right}, W)$ 17: $i_list.push(I_{left})$ 18:19: $i_list.push(I_{right})$ 20: end while 21: return *i_list*

in our base signal, where $maxIns = \frac{\min\{M_{base}, TotalBand\}}{W}$. Each input signal \vec{Y}_i is broken into $\frac{M}{W}$ non-overlapping intervals of size W. This provides a "dictionary" of $\frac{N*M}{W}$ candidate base intervals (CBIs). The algorithm will choose maxIns CBIs out of this dictionary to be inserted into a candidate update base signal. We will describe in subsection 4.3 how to determine how many of these CBIs will ultimately be inserted into the base signal.

Each CBI $Cand_i$ can be used to approximate any other CBI $Cand_j$, which is in-fact part of some \vec{Y}_k , using regression. We consider such an approximation to be beneficial, only if the error of the approximation is smaller than the error of approximating $Cand_j$ using standard linear regression. In Algorithm 4 we denote the latter error as $LinearErr(Cand_j)$. The benefit of using $Cand_i$ to approximate $Cand_j$ is the reduction in error that we get compared to $LinearErr(Cand_j)$.

The CBIs are stored in an unordered list Q. At each step of the algorithm, the CBI in Q with the largest benefit is selected for inclusion in the candidate update base signal stored in base_list. After each selection, the benefits of the remaining CBIs in Q have to be properly updated. As we mentioned, the benefit of using $Cand_i$ to approximate $Cand_j$ is originally equal to the reduction

Algorithm 4 GetBase() Algorithm

Require: $\vec{Y_1}, \ldots, \vec{Y_N}, W, M, maxIns$ 1: Create $K = \frac{N \times M}{W}$ CBIs of width W 2: For each CBI $Cand_i$, set its benefit to 0 3: Maintain unsorted list Q with CBIs 4: Maintain list base_list with selected stored intervals 5: $LinearErr(Cand_i)$ is the error of approximating $Cand_i$ using standard linear regression 6: for i in 1..K do for j in 1..K do 7: {Calculate error of approximating the j-th} 8: {CBI by using as base the i-th CBI} 9: $error = Regression(Cand_i, Cand_i, 0, 0, W)$ 10:if $err \leq Linear Err(Cand_i)$ then 11: $Cand_i.benefit + = LinearErr(Cand_i)$ -error 12:end if 13:14:end for 15: $Q.insert(Cand_i)$ 16: end for for i in 1..maxIns do 17:C = Q.popBestInterval()18:19: $base_list.insert(C)$ 20:for j in 1..|Q| do adjust(Q[j], benefit, C)21:22:end for 23: end for 24: return base_list

in error that we get compared to $LinearErr(Cand_j)$. However, at an intermediate step of the algorithm, some CBIs have already been selected for inclusion in the candidate update base signal. By using these stored CBIs, many of the remaining CBIs can now be better approximated than by using standard linear regression. Thus, the benefit of using $Cand_i$ to approximate $Cand_j$ has to be adjusted, to depict the reduction in error that we get when compared to the best approximation for $Cand_i$ that we have so far, by using the current candidate update base signal.

An example is presented in Figure 4. In this small example we consider just 3 CBIs, out of which we need to pick which two to select. In the left part of the figure, we present the benefits of each of the 3 CBIs. The first CBI has the largest total benefit, and is thus selected. In the right part of the figure, the adjusted benefits of the remaining CBIs are presented. Notice that now, the third CBI will be selected, even though initially it had a lower benefit than the second CBI.

In the GetBase() algorithm, for each of the $K = \frac{N \times M}{W}$ CBIs, we first estimate its benefit for

	App	Total		
CBI	1	2	3	Benefit
1	1	0.95	0.50	2.45
2	0.8	1	0.55	2.35
3	0.6	0.65	1	2.25

	Appr	Total	
CBI	2	3	Benefit
2	0.05	0.05	0.10
3	0	0.5	0.50

Initial Benefits of CBIs

Adjusted Benefits of Non-Stored CBIs

Figure 4: Example of the GetBase() Algorithm

approximating all the other CBIs. Each such approximation requires O(W) time, thus resulting in a total complexity of $O(\frac{N^2M^2}{W})$. Then, for each of the maxIns selected CBIs, detecting the one with the largest benefit requires O(K) time (we do not sort the CBIs). After each selection, adjusting the benefits of the remaining CBIs requires time $O(K^2)$. Thus, the overall running time complexity of the algorithm is $O(\frac{N^2M^2}{W} + maxIns \times \frac{N^2M^2}{W^2})$, while its space requirements is $O(\frac{N^2M^2}{W^2})$.

For $n = N \times M$ being the size of the data, a value of $W = \sqrt{n}$ used by the SBR algorithm (described in the next subsection) results in a running time of $O(n^{1.5})$ for GetBase() and space of O(n), since $maxIns \times W \leq TotalBand \leq n$. In case of severe memory constraints, we can easily modify the GetBase() algorithm to only store for each CBI the smallest error of approximating it using at each step the current base signal. The only modification will be to replace Lines 20-22 of the GetBase() algorithm with a double for-loop similar to the one of Lines 6-16, and alter the calculation of each CBI's benefit to take into account the error of the best approximation that we have for each CBI so far. This modified algorithm requires $O(\sqrt{n})$ space and has a running time of $O(maxIns \times n^{1.5})$.

4.3 The SBR Algorithm

We now present the Self-Based Regression (SBR) algorithm, which performs the approximation of the data values. The algorithm receives as input the latest $n = N \times M$ data values, a bandwidth constraint TotalBand (number of values to transmit, including any base signal values), the maximum size of the base signal M_{base} and the current base signal \vec{X} of size $|\vec{X}| \leq M_{base}$.⁸ From these parameters the user/application has to provide only TotalBand and M_{base} . The SBR algorithm must then make the following decisions:

^{1.} Decide how many, and which base intervals to insert into the base signal. Recall that any such

⁸At the first transmission the current base signal will be empty.

Algorithm 5 SBR Algorithm

Require: $\vec{X}, \vec{Y_1}, \ldots, \vec{Y_N}, M, Total Band, M_{base}$

1: $maxIns = \frac{\min\{M_{base}, TotalBand\}}{W}$

2: $W = \sqrt{N \times M}$

3: $base_list = GetBase(\vec{Y_1}, \dots, \vec{Y_N}, W, M, maxIns)$

- 4: {Errors[i] is the approximation error after inserting}
- 5: {the first i CBIs of base_list in the base signal}
- 6: Initialize Errors[i] = UNDEFINED $\forall i \in [0..maxIns]$ 7: $Ins = \text{Search}(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, W, M, TotalBand, base_list,$ Errors, 0, maxIns)
- 8: Form \vec{X}_{new} by appending the *Ins* first intervals of the base_list to \vec{X}
- 9: $B = TotalBand Ins \times (W + 1)$
- 10: $GetIntervals(\vec{X}_{new}, \vec{Y}_1, \dots, \vec{Y}_N, B, W)$
- 11: if $|\vec{X}_{new}| > M_{base}$ then
- Evict $Repl = \frac{|\vec{X}_{new}| M_{base}}{W}$ intervals of \vec{X}_{new} that also belonged to \vec{X} using a LFU replacement policy 12:
- Replace evicted intervals with the last *Repl* intervals of \vec{X}_{new} 13:
- 14: **end if**
- 15: $\vec{X} = \vec{X}_{new}$

Transmit the inserted base intervals, their offsets in the base signal and the regression intervals 16:

Algorithm 6 CalculateError SubRoutine

Require: $\vec{X}, \vec{Y_1}, \ldots, \vec{Y_N}, B, W, Errors, pos$ 1: if Errors[pos] == UNDEFINED then list' = $GetIntervals(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, B - pos \times W, W)$ 2: $\operatorname{Errors}[\operatorname{pos}] = \operatorname{sum} \operatorname{of} \operatorname{errors} \operatorname{in} \operatorname{list}'$ 3: 4: **end if**

base intervals need to be transmitted to the base station.

2. If the above procedure causes the size of the base signal to exceed M_{base} , then some base intervals need to be evicted from the base signal, in order to keep its maximum size at M_{base} .

3. Decide how to best approximate the data values given the updated base signal.

We here have to emphasize that it is not always desirable to insert a large number of base intervals into the base signal. Since any inserted base interval needs to be communicated to the base station, the larger the number of such intervals, the smaller the number of intervals that can be used to approximate the N signals by the GetIntervals() algorithm, since the overall bandwidth consumption is upper-bounded by the *TotalBand* parameter.

The SBR algorithm is presented in Algorithm 5. It initially calls the GetBase() subroutine to select a set of $maxIns = \frac{\min\{M_{base}, TotalBand\}}{W}$ CBIs. It then performs a binary search on this list, to determine the number of CBIs that will ultimately be inserted into the base signal. This search terminates when the algorithm determines a number of intervals Ins, such that the error of the

Algorithm 7 Search SubRoutine

Require: $\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, W, B, base \ Jist, Errors, start, end$ 1: if end == start then return start 2: 3: end if 4: middle = (start + end) / 2 5: $CalculateError(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, B, W, middle)$ 6: $CalculateError(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, B, W, start)$ if Errors[middle] > Errors[start] then 7: $CalculateError(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, B, W, end)$ 8: if Errors[end] > Errors[start] then 9: return Search $(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, W, M, B, base list, Errors, start, middle)$ 10:else 11:return Search $(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, W, M, B, base \ list, Errors, middle, end)$ 12:13:end if 14:else $CalculateError(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, B, W, middle + 1)$ if Errors[middle + 1] < Errors[middle] then 15:16:return Search $(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, W, M, B, base \ list, Errors, middle + 1, end)$ 17:18:else return Search $(\vec{X}, \vec{Y_1}, \dots, \vec{Y_N}, W, M, B, base \ list, Errors, start, middle)$ 19:20:end if 21: end if

approximation when inserting the first Ins intervals of the aforementioned list in the base signal is lower than inserting either the first Ins - 1 intervals, or the first Ins + 1 intervals into the base signal. This is achieved through the call to function **Search()** at Line 7, which is presented in Algorithm 7. The approximation of the N signals is then performed by using the concatenation of the previous base signal with these Ins intervals. After this step, if the size of the base signal now exceeds M_{base} , then enough base intervals of the old base signal are evicted from the base signal using a Least Frequently Used (LFU) replacement policy. Any newly inserted base interval will thus either occupy an empty position of the base signal, or replace another base interval. Each transmission includes exactly TotalBand values:

1. The *Ins* newly inserted base intervals, and their position in the base signal in which they were ultimately inserted ($Ins \times (W + 1)$ values in total).

2. $\frac{TotalBand-Ins \times (W+1)}{4}$ intervals of four values each (start, shift plus the two regression parameters).

The running time complexity of the SBR algorithm is $O(n^{1.5} + (nlog(\frac{TotalBand}{N}) + TotalBand \times \sqrt{n} \times M_{base}) \times \log(maxIns))$, where $maxIns = \frac{\min\{M_{base}, TotalBand\}}{\sqrt{n}}$. Thus the entire algorithm has a modest $O(n^{1.5})$ dependency on the data size, while its running time scales linearly with the size

of the transmitted data TotalBand and the (maximum) size of the base signal M_{base} .

5 Experiments

In this section, we provide a thorough analysis of our techniques. In subsection 5.1 we describe the datasets we used. In subsection 5.2 we compare the SBR algorithm against standard approximation techniques (Wavelets, DCT, Histograms). Finally, in subsection 5.3 we compare the GetBase() algorithm against alternative base-signal constructions, while in subsection 5.4 we present an analysis of the SBR algorithm.

5.1 Dataset Description

For the experiments we used the following real datasets:

- Phone Call Data: Includes the number of long distance calls originating from 15 states (AZ, CA, CO, CT, FL, GA, IL, IN, MD, MN, MO, NJ, NY, TX, WA). For each state we provide the number of calls per minute for a period of 19 days (data from AT&T's network).
- Weather Data: Includes the air temperature, dewpoint temperature, wind speed, wind peak, solar irradiance and relative humidity weather measurements for the station in the university of Washington, and for year 2002 (http://www-kl2.atmos.washington.edu/kl2/grayskies).
- Stock Data: Includes information on all trades performed in a minute basis over April 3 and April 4 of year 2000. The approximated measure in our experiments is the trade value of the stock.

5.2 Comparison to Alternative Techniques

5.2.1 Experimental Setup

For this experiment we used all three datasets described in Section 5.1. From the *Stock* data, we extracted the trade values of the following ten (N=10) stocks: Microsoft, Oracle, Intel, Dell, Yahoo, Nokia, Cisco, WorldCom, Ariba and Legato Systems. For each stock we created a random sample of 20480 of its trade values, and then split each sample in ten files of 2048 values each. The first of these ten files of each stock was used for the initial creation of our base signal, while the

Compression		Weat	her Data			Stoc	k Data					
Ratio	SBR Wavelets DCT Histograms			SBR Wavelets DCT Histograms SB				SBR Wavelets DCT			Histograms	
5%	1.160	2.187	35.835	27.692	0.089	0.123	0.232	0.283				
10%	0.403	0.824	20.169	11.294	0.033	0.056	0.208	0.233				
15%	0.209	0.514	14.328	5.432	0.017	0.034	0.192	0.214				
20%	0.118	0.356	10.774	3.009	0.009	0.022	0.179	0.199				
25%	0.069	0.258	8.975	1.507	0.006	0.015	0.166	0.182				
30%	0.043	0.191	6.526	0.995	0.003	0.011	0.153	0.169				

Table 2: Average SSE Error Varying the Compression for Weather and Stock Datasets

remaining files were used to simulate nine update operations. For the *Weather* dataset, we selected the first 40960 records and then split the data measurements of each signal into ten files of 4096 values each. For the *Phone Call* dataset, the aggregates for each state (N=15) were broken into ten files of 2560 values each.

In our experiments we compared the accuracy of SBR against the approximations obtained by using the *Wavelet* decomposition [5, 29], equi-depth Histograms [23] and the DCT. The Fourier transform was also considered, but produced consistently larger errors than DCT and is thus omitted. For a fair comparison we set the space used by all methods to the exact same amount.

For all methods we considered both treating each bunch of updates as a group of N series \vec{Y}_i , each of length M, and, alternatively, concatenating the signals into a single series Y of length $N \times M$. For Wavelets, we found out that this produced in most cases significantly more accurate results than by dividing the space equally among the N signals (by a factor of 5 in many cases) because some signals needed more wavelet coefficients than others to be approximated well. For Wavelets, we also considered a 2-dimensional decomposition of the $N \times M$ values, which produced worse results than the 1-dimensional decomposition. In the tables we present the best results achieved by each method.

5.2.2 Comparison Varying the Compression Ratio

We varied the compression ratio (size of the transmitted data TotalBand over the data size n) from 5% to 30%. In this experiment we set M_{base} to 2048 values for the *Phone Call* and the *Stocks* datasets and to 3456 values for the *Weather* dataset. In Tables 2 and 3 we present the results.

In all datasets SBR produces significantly more accurate results than the other approximations. The difference is larger for the *Phone Call* dataset which contained the largest values. As the size

Compression	Average SSE Error					al Sum Squa	ared Rel	ative Error				
Ratio	SBR Wavelets DCT Histograms		SBR Wavelets DCT Histograms		SBR Wavelets DCT Histogram		SBR Wavelets DCT H		SBR	Wavelets	DCT	$\operatorname{Histograms}$
5%	9,631	29,938	15,714	$165,\!241$	922	38,477	9,019	$139{,}528$				
10%	5,071	12,349	$10,\!173$	$45,\!610$	503	$19,\!186$	3,002	62,337				
15%	3,192	$7,\!998$	6,767	23,311	325	12,885	1,400	36,812				
20%	2,170	5,821	$5,\!661$	$15,\!581$	222	10,954	$1,\!192$	34,820				
25%	1,527	4,468	4,791	$11,\!340$	158	6,915	823	$33,\!237$				
$\overline{30\%}$	1,091	$3,\!537$	$4,\!157$	8,689	116	3,865	721	30,010				

Table 3: Errors Varying the Compression Ratio for Phone Dataset

of transmitted data increases, the error in our method decreases more sharply, and is up to 4.4 times smaller than the error of Wavelets. The DCT and the Histogram approximations produced much larger errors is most cases.

We repeated the experiment for the *Phone Call* dataset, computing this time the sum-squared relative error. The results are also shown in Table 3. The modified **Regression()** algorithm is presented in the Appendix. Depending on the compression ratio, our method was up to 49 times better than Wavelets, 9.8 times better than DCT and 258 times better than Histograms. We notice here that for this comparison we used straight-forward Wavelets that are optimal only under the sum-squared-error. Garofalakis and Gibbons in [11] describe novel algorithms for minimizing, among other metrics, the relative error of a Wavelet-based approximation. Except for cases of very skewed datasets, they observe a reduction of the mean relative error up to 3 times over regular Wavelets. These improvements were seen for very coarse approximations (i.e. for a compression ratio of 5% or less) where our method already has an advantage of 42-1 over regular Wavelets. For more space, their techniques are a lot closer to regular Wavelets.

5.2.3 Mixing The Datasets

At this experiment we tried mixing data from different datasets, to reduce the amount of correlation among the approximated signals. We thus created a dataset that contains phone call data from three states (AZ, CA and FL), three types of meteorological measurements (air temperature, pressure and solar irradiance), and data from three stocks (Microsoft, Intel and Oracle). For each of these data series we created ten files of 2048 values each. We then varied the compression ratio of all algorithms from 5% to 30% and set M_{base} to 2048 values. In Table 4 we present the average sum squared and total sum squared relative errors for all methods. The improvements of the SBR

Compression	Average SSE Error					Total Sum Squared Relative Erre				
Ratio	SBR Wavelets DCT Histogra		BR Wavelets DCT Histograms		SBR Wavelets DCT Histogra		SBR	Wavelets	DCT	$\operatorname{Histograms}$
5%	2,900	8,094	$12,\!677$	$199,\!150$	113	20,974	$29,\!625$	$182,\!027$		
10%	918	3,020	7,146	46,805	37	$11,\!054$	8,653	43,701		
15%	364	1,582	4,757	23,711	17	5,481	4,825	26,068		
20%	139	894	3,814	$14,\!157$	9	5,310	3,339	14,780		
25%	46	516	3,120	10,486	5	$5,\!172$	$6,\!115$	11,118		
30%	11	297	2,680	6,894	3	5,109	1,579	9,591		

Table 4: Errors for Varying Compression Ratios for the Mixed Dataset

	$\mathbf{Error} \ \mathbf{Oet} \mathbf{Base}()$							
Dataset	GetBaseSVD()	Linear Regression	GetBaseDCT()					
Weather	10.55	4.47	6.44					
Phone	1.13	1.32	1.19					
Stock	2.08	2.77	2.99					

Table 5: Comparison to Alternative Base Signals

algorithm were even larger in this case. The SBR algorithm produced up to 27 times smaller average sum squared errors than the closest competitor, while the improvement reached up to 1034 times for the total sum squared relative error.

5.3 Alternative Base Signal Constructions

In the Appendix we present two alternative algorithms to GetBase(). The first, denoted as GetBaseSVD(), is based on the Singular Value Decomposition. The second algorithm, GetBaseDCT(), uses the basis of the Discrete Cosine Transform (DCT), which is a collection of cosine functions. Finally, a third alternative for SBR, is to do standard linear regression without using a specially constructed base signal. For the later case, no bandwidth is lost for sending base signal values and we do not need the *I.shift* pointer. Thus we can send exactly *TotalBand*/3 intervals for a bandwidth limit *TotalBand*. Similarly, the DCT-base consists of cosine functions and its values are constructed on the fly and are thus neither stored in memory, nor are they transmitted to the base station.

In Table 5 we compare the approximations obtained by using the base signals computed in algorithm GetBase() with the base signal from the alternative constructions. We need to emphasize here that for this experiment we modified the BestMap() function not to use linear regression as an alternative to using the base signal (so that the differences among GetBase(), GetBaseSVD(),

		Transmission								
Dataset	1	2	3	4	5	6	7	8	9	10
Weather	7	6	0	3	1	4	0	1	1	1
Phone	8	6	0	1	0	0	2	0	0	0
Stock	6	0	0	2	1	2	0	1	0	0

Table 6: Number of Inserted Base Intervals per Transmission



Figure 5: Average Running Time vs TotalBand

GetBaseDCT() and linear regression are not diffused). Using the BestMap() function as presented in Section 4.2 would thus further improve the results of our method. The compression ratio was set to 10%. We notice that GetBase() performs a lot better in the *Weather* dataset, up to 10 times better than the alternative algorithms. For the *Phone Call* and the *Stock* data the differences are smaller but still significant.

5.4 Analysis of SBR

We now analyze several characteristics of the SBR algorithm, including its running time, the number of base intervals it selects for inclusion in the base signal and the quality of its decisions.

In Figure 5 we plot the average time of each transmission operation for the Stock dataset, when the size of the transmitted data is varied from 5% to 30% of the data size, and for an experimental setup similar to the one of Section 5.2.2. Since we have not yet ported our code to the StrongARM platform, we executed this experiment on a Irix machine using a 300MHz processor. As expected (see Section 4.3) the running time scales linearly with the size of the transmitted data. Notice that SBR is significantly faster when greater reduction is obtained. For many practical applications, we expect to use a compression ratio of 10% or less.

The SBR algorithm dynamically decides the number of base signal values to use for an upper bound M_{base} . We now compare SBR against a straight-forward implementation that populates all



Figure 6: SSE error vs base signal size

the available space for the base signal. In Figure 6 we plot the error of only the initial transmission as the size of the base signal is varied, manually, from 1 to 30 intervals for the *Phone, Stock* and *Weather* datasets. For this initial transmission we populated the entire space of the base signal using the **GetBase()** algorithm. For each dataset we also show the selection that the SBR algorithm made, when deciding how many base intervals to populate. For presentation purposes the errors for each dataset have been divided by the error of the approximation when using just one interval. We set the size of each stock, phone and weather data file to 3072, 2048 and 5120 values, correspondingly, in order for all datasets to have exactly the same size, and the TotalBand value to 5012, which results to a compression ratio of about 16%.

The fixed value of the compression ratio implies that an increase in the size of the base signal results in a decrease in the number of intervals used to approximate the data values in order to keep the total space constant. After some point, the benefit of storing more intervals for the base signal is outweighted by the increase in the error that we get due to the reduced number of intervals used for the approximation. It is interesting to see that the optimal case occurs for a base size of between 7 (for the *Weather* dataset) and 9 base intervals (for the *Stock* dataset), which correspond to just 2.9% to 3.75% of the data size at the first transmission. The SBR algorithm made the optimal choice for the *Phone* and *Weather* datasets and produced a near-optimal solution for the *Stock* dataset (it selected to insert 6 base intervals, instead of 9). We remind that the M_{base} base signal values need to be kept in the memory of the sensor in order to perform the approximation. Our results suggest that a very small fraction of memory needs to be sacrificed for these values.

For the same data setup, we report in Table 6 the number of inserted base intervals during the 10

transmissions. As we can see, most base intervals are inserted during the first two transmissions. We notice that there are many transmissions on which no new base intervals are inserted, and that the different datasets seem to contain a widely different number of features, with the Weather dataset containing the most features, and the Stock dataset containing the fewest.

6 Conclusions

We presented a new data reduction technique designed for data disseminated over sensor networks. Our method splits the recorded series into intervals of variable length and then encodes each of them using an artificially constructed *base signal*. The values of the base signal are extracted from the real measurements and maintained dynamically as data changes. In our experiments we used real datasets from a variety of fields (weather, stock and phone call data). Using the sum-squared error and the sum-squared relative error of the approximation, our method significantly outperformed in accuracy approximations obtained by using Wavelets, DCT and Histograms.

A key to our method is the use of the base signal for encoding piece-wise linear correlations among the data values. We emphasize here that our method does not only apply to linear datasets; in fact none of the data we used are linear in nature. Linearity is exploited when encoding the correlations of the data values and the base signal. An interesting question is to what extent nonlinear encodings over the base signal values would benefit the approximations obtained without sacrificing complexity. We plan to investigate this path in the future.

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Appendix

Alternative Base Signal Constructions

We here present the two alternative algorithms for obtaining a base-signal from the data in more details.

Construction Using SVD

SVD involves computing the eigenvectors and eigenvalues of a given $N \times n$ matrix R. It can be proven that any real matrix can be written as:

$$R = U \times \Lambda \times V^t$$

where U is a column-orthonormal $N \times r$ matrix, r is the rank of matrix R, Λ is a diagonal $r \times r$ matrix of the eigenvalues λ_i of R and V is a column-orthonormal $n \times r$ matrix. By definition $U^t \times U = V^t \times V = I$, where I is the identity matrix. It can be shown that the columns of V are the eigenvectors of matrix $R^t \times R$. Similarly, the eigenvalues of $R^t \times R$ are the squares of λ_i s i.e.

$$R^t \times R = V \times \Lambda^2 \times V^t$$

For R=A (our collected measurements), $R^t \times R$ captures the similarities among the columns of A (each collected sample). SVD can be used for approximating $R^t \times R$ by keeping the first few eigenvectors (columns of matrix V). Informally, each eigenvector captures linear trends among the rows of A (the \vec{Y}_i s), see [15] for an application of this observation in a different context.

We here propose the use of SVD as a competitor to the GetBase() algorithm for generating a base signal from the data. We sketch the new algorithm (GetBaseSVD()) bellow.

- 1. For each row of A, list all non-overlapping intervals of length W. This gives us $\frac{M}{W}$ intervals per row and $n = \frac{N \times M}{W}$ intervals overall.
- 2. Build an $n \times W$ matrix R whose rows are the intervals of the previous step.
- 3. Compute the SVD of $R = U \times \Lambda \times V^t$. Return the first *Store* columns of V.

By definition, V is an $r \times W$ matrix (r=rank(R)) of the eigenvectors of $R^t \times R$. The eigenvectors are ordered from left to right in V. The first column of V contains the eigenvector (of length W) that corresponds to the largest eigenvalue of $R^t \times R$. The algorithm returns the top-*Store* eigenvectors of total size *Store* × W. These constitute the base signal from GetBaseSVD().

Construction Using DCT

The base signal can be constructed from the basis-vectors of standard mathematical transforms. As an example we present a base signal construction, motivated by the Discrete Cosine Transform (DCT). Assuming we are to use base intervals, each of length W, we enumerate all frequencies f such that $0 \le f \le W$. For each frequency f, we define a base interval with values $cos(\frac{(2i+1)\pi}{2W}f)$, where $0 \le i < W$. We call this algorithm GetBaseDCT(). We notice we do not need to store these intervals implicitly as they can be computed on the fly.

Handling Other Error Metrics

We now present the necessary modifications to the Regression algorithm of Section 4.2 when the desired error metric involves minimizing the sum squared relative errors, or the maximum absolute error of the approximation.

The Regression algorithm approximates the value $Y[i + start_y]$ as $a \times X[i + start_x] + b$. The

relative error induced by this approximation is:

$$\frac{|Y[i+start_y] - a \times X[i+start_x] - b|}{max\{c, |Y[i+start_y]|\}}$$

The c value serves as a *sanity* bound, and helps avoid very large relative error values when the $Y[i + start_y]$ value is either zero, or close to zero. The Regression algorithm that minimizes the sum squared relative error of the approximation is presented in Algorithm 8.

Algorithm 8 Regression Subroutine that Minimizes the Sum of the Squared Relative Errors
Require: \vec{X} , \vec{Y} , start_x, start_y, length, sanity
1: {Compute Regression Parameters}
2: $sum_x = \sum_{0 \le i < length} \frac{X[i+start_x]}{max\{sanity, Y[i+start_y] \}}$
3: $sum_y = \sum_{0 \le i < length} \frac{Y[i+start_y]}{max\{sanity, Y[i+start_y] \}}$
4: $sum_xy = \sum_{0 \le i < length} \frac{X[i+start_x]Y[i+start_y]}{max\{sanity, Y[i+start_y] \}}$
5: $sum_x2 = \sum_{0 \le i < length} \frac{X[i+start_x]^2}{max\{sanity, Y[i+start_y] \}}$
6: $sum_z = \sum_{0 \le i < length} \frac{1}{max\{sanity, Y[i+start_y] \}}$
7: $a = \frac{sum_z \times sum_x y - sum_x \times sum_y}{sum_z \times sum_x x \times sum_y}$
8: $b = \frac{sum_z - sum_x}{sum_z}$
{Compute sum squared relative error of signal}
$\{\vec{Y'} = a\vec{X} + b \text{ in range } [start_y \dots start_y + length)\}$
9: $err = \sum_{i=0}^{length-1} \left(\frac{Y[i+start_y] - (a \times X[i+start_x]+b)}{max\{sanity, Y[i+start_y] \}} \right)^2$
10: return (a,b,err)

Calculating the a,b parameters that minimize the maximum absolute error of the approximation is somewhat harder to accomplish. The solution is based on the well known Chebyshev approximation problem, which can be solved with a randomized linear programming algorithm in O(length)randomized expected time and O(length) space.