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Chemo-inspired Genetic Algorithm for Optimizing the Piecewise Aggregate Approximation

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Time Series.

Abstract: In a previous work we presented DEWPAA: an improved version of the piecewise aggregate approximation

representation method of time series. DEWPAA uses differential evolution to set weights to different segments of the time series according to their information content. In this paper we use a hybrid of bacterial foraging and genetic algorithm (CGA) to set the weights of the different segments in our improved piecewise aggregate approximation. Our experiments show that the new hybrid gives better results in time

series classification.

1 INTRODUCTION

In the last two decades there has been an increasing interest in temporal data, namely, time series. A time series is a chronological collection of observations. This data type is encountered in many scientific and financial applications. The main feature of time series is their high-dimensionality. One of the common approaches to handle the problem of high-dimensionality of time series is to transform them into another domain with a lower-dimensionality followed by an indexing mechanism, called a dimensionality reduction technique or a representation method, applied to this lower-dimensional data.

Several representation methods have been proposed to reduce the dimensionality of time series data. Of those we mention *Discrete Fourier Transformation* (DFT) (Faloutsos *et al.*, 1994), *Discrete Wavelet Transformation* (DWT) (Chan and Fu 1999), *Chebyshev Polynomials* (CHEB) (Cai and Ng, 2004), *Symbolic Aggregate approXimation* (SAX) (Lin *et al.*, 2003), *Piecewise Linear Approximation* (PLA) (Morinaka *et al.*, 2001).

The *Piecewise Aggregate Approximation* (PAA) (Keogh *et al.*, 2000), (Yi and Faloutsos, 2000) is a time series representation method that has been used extensively for its simplicity and its low computational complexity.

In (Muhammad Fuad, M.M., 2012) we applied differential evolution; a popular bio-inspired optimization algorithm, to set weights to different segments of PAA-represented time series, which reflect the information content of each segment. The weights were determined through an optimization process whose output is the weights that maximize the information content of the PAA representation. We showed how this modification improves the performance of PAA.

Differential evolution, however, may suffer from *stagnation*; i.e. the inability of progressing towards global optima. It may also suffer from premature convergence.

In this paper we propose solving the aforementioned optimization problem by applying an alternative optimizer which is a hybrid of two bio-inspired optimization algorithms: genetic algorithm and bacterial foraging. We show how this hybrid yields better results than those obtained when using differential evolution.

The rest of this paper is organized as follows: Section 2 presents related work. The hybrid algorithm is introduced in Section 3, and the comparison with the previous method is conducted in Section 4. We conclude this paper in Section 5.

2 RELATED WORK

PAA is a simple, yet efficient, representation method of time series. PAA reduces the dimensionality of a time series S from n to N dimensions by segmenting the time series into equal-sized frames and mapping each segment to a point that represents the mean of the points that constitute that segment. The similarity measure given in the following equation:

$$d^{N}\left(S,T\right) = \sqrt{\frac{n}{N}} \sqrt{\sum_{i=1}^{N} \left(\overline{s_{i}} - \overline{t_{i}}\right)^{2}} \tag{1}$$

is defined on the lower-dimensional space. This similarity measure is a lower bound of the Euclidean distance defined on the original space. In Figure 1 we show an example of applying PAA to reduce the dimensionality of a time series from n=12 to N=3.

The main drawback of PAA is that it is unable to faithfully represent the original time series due to the information loss that results from the technique PAA uses to reduce the dimensionality, as we showed in (Muhammad Fuad, M.M., 2012). In that paper we proposed weighting different segments of the PAA representation differently according to their information content. We proposed using an alternative similarity measure for PAA which we called the *Weighted Piecewise Aggregate Approximation Distance (WPAAD)*:

$$WPAAD(S,T) = \sqrt{\frac{n}{N}} \sqrt{\sum_{i=1}^{N} w_i (\overline{s_i} - \overline{t_i})^2}$$
 (2)

 $w_i \in [0,1]$

The challenge here is to find an objective measure to determine the values of W_i .

In (Muhammad Fuad, M.M., 2012) we formulated the above problem as an optimization problem where the fitness function is the classification error and the optimization algorithm seeks to set the values of w_i that minimize the classification error.

The optimizer we used in (Muhammad Fuad, M.M., 2012) was the differential evolution, and we called our method the *Differential Evolutionary Weighted Piecewise Aggregate Approximation* (DEWPAA).

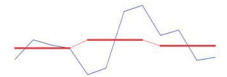


Figure 1: PAA dimensionality reduction.

Differential Evolution (DE): DE is an evolutionary optimization algorithm that has the same computational steps as any other evolutionary algorithm. Its particularity is in the way the population members are perturbed, as DE uses a scaled difference of two other members added to a third to perform this operation.

DE is particularly adapted to solve continuous optimization problems, and it has been successfully used to solve real-life optimization problems.

DE starts by initializing a random population of vectors. After initialization DE, and for each member \vec{T}_i (the *target vector*), creates a *donor vector* \vec{D} as a weighted difference of two other vectors in the population added to a third one. In the next step DE generates a *trial vector* \vec{R} through a crossover operation.

In the next step DE selects which of the trial vector or the target vector will survive in the next generation and which will die out. This selection is based on which of \vec{T}_i and \vec{R} yields a better value of the fitness function.

The above steps repeat for a number of generations or until a stopping criterion terminates the algorithm.

3 CHEMO-INSPIRED GENETIC ALGORITHM

Hybridization of different optimization algorithms has been extensively used to solve different optimization problems. In time series data mining different hybrid methods have been successfully used (Muhammad Fuad, 2014a, 2014b, 2014c). The main advantage that hybridization offers is that the resulting hybrid method benefits from the strengths of the two methods, or it avoids their weaknesses.

In this work we use a hybrid of genetic algorithm and bacterial foraging proposed by (Das and Mishra, 2013) to solve the problem we presented in Section 2.

3.1 The Genetic Algorithm

The Genetic Algorithm (GA) is a famous evolutionary algorithm that has been applied to solve a variety of optimization problems. GA is a population-based global optimization algorithm which mimics the rules of Darwinian selection in that weaker individuals have less chance of surviving the evolution process than stronger ones.

GA captures this concept by adopting a mechanism that preserves the "good" features during the optimization process.

In GA a population of candidate solutions (also called *chromosomes*) explores the search space and exploits this by sharing information. These chromosomes evolve using genetic operations (selection, crossover, mutation, and replacement).

GA starts by randomly initializing a population of chromosomes inside the search space. The fitness function of these chromosomes is evaluated. According to the values of the fitness function new offspring chromosomes are generated through the aforementioned genetic operations. The above steps repeat for a number of generations or until a predefined stopping condition terminates the GA.

3.2 Bacterial Foraging

The foraging bahavior of the *Escherichia coli* (*E. coli*) bacteria has inspired a nature-inspired optimization algorithm called *Bacterial Foraging* (BF). The motivation behind this optimizing algorithm is that in order to perform social foraging, an animal needs communication capabilities. Over a period of time this animal gains advantages which exploit the sensing capabilities of the whole group. This helps the group to predate on a larger prey, or it enables the individuals to get better protection against predators (Kim *et al.*, 2007).

The basis of BF is that animals with poor foraging strategies tend to be eliminated by natural selection and they are either replaced by other individuals with better foraging strategies or they are shaped into ones which have these desirable strategies (Passino, 2002). BF formulates this process as an optimization problem.

The *E. coli* bacterium moves by means of a set of flagella, each driven as a biological motor. The two types of movements the *E. coli* bacteria perform are *swimming* and *tumbling*. The former takes place when the flagella rotate in the counterclockwise direction whereas the latter is achieved by rotating the flagella in the clockwise direction. Figure 1 shows these two movement types. Together they are known as *chemotaxis* (which we will define more formally later in this section). The aim of chemotaxis is to help the bacterium approach or avoid nutrient or noxious substance gradients. This chemotaxis progress can be destroyed by sudden environmental changes which cause the elimination and dispersal of a group of bacteria.



Flagella rotating counterclockwise: swimming



Flagella rotating clockwise: tumbling

Figure 2: The swimming and tumbling movements.

BF finds the minimum of a function $f(\theta)$; $\theta \in \mathbb{R}^{nbp}$ (nbp is the number of parameters) by applying four mechanisms; chemotaxis, swarming, reproduction, and elimination-dispersal.

The position of each member of the population of N_b bacteria at the j^{th} chemotactic step, k^{th} reproduction step, and l^{th} elimination-dispersal event is denoted by $P(i, j, k) = \left\{ \theta^i(j, k, l) / i = 1, 2, ..., N_b \right\}$

We now describe the four mechanisms we mentioned earlier in this section:

• **Chemotaxis:** Let $\theta^i(j,k,l)$ be the i^{th} bacterium at the j^{th} chemotactic step, k^{th} reproduction step, and l^{th} elimination-dispersal event, then the movement of the bacterium can be represented by:

$$\theta^{i}(j+1,k,l) = \theta^{i}(j,k,l) + C(i) \frac{\Delta(i)}{\sqrt{\Delta^{T}(i)\Delta(i)}}$$
(3)

where Δ is a vector in the random direction whose elements lie in the interval [-1, 1].

• **Swarming:** *E. coli* bacteria demonstrate a swarming behavior as they travel in rings which move up the nutrient medium when they are placed in the center of a semisolid matrix with a single nutrient chemo-effecter. When simulated

Table 1: The symbols used in the description of bacterial foraging.

N_b	The number of bacteria in the population
N_c	The number of chemotactic steps
N_s	The swimming length
N_{re}	The number of reproduction steps
N_{ed}	The number of elimination-dispersal events
P_{ed}	The probability of elimination-dispersal
C(i)	The size of the step taken in the random direction determined by the tumble

by a high level of succinate the bacteria release an attractant aspartate which helps them aggregate into groups and thus move as a swarm. The cell-to-cell signal in the swam can be represented by the following function:

$$f_{cc}(\theta, P(j, k, l)) = \sum_{i=1}^{N_b} f_{cc}(\theta, \theta^i(j, k, l))) =$$

$$\sum_{i=1}^{N_b} \left[-d_{\text{attractant}} \cdot \exp\left(-\omega_{\text{attractant}} \sum_{m=1}^{nbp} (\theta_m - \theta_m^i)^2\right) \right] +$$

$$\sum_{i=1}^{N_b} \left[-h_{\text{repellant}} \cdot \exp\left(-\omega_{\text{repellant}} \sum_{m=1}^{nbp} (\theta_m - \theta_m^i)^2\right) \right]$$

$$(4)$$

where the coefficients $d_{attractant}$, $\omega_{attractant}$, $h_{repellant}$, $\omega_{repellant}$ are control parameters.

The objective function $f_{cc}(\theta, P(j,k,l))$ is added to the original objective function to represent a *time varying* objective function in that if many cells come close together there will be a high amount of attractant and hence an increasing likelihood that other cells will move towards the group. This produces the swarming effect (Passino, 2002).

- **Reproduction:** Through this process the least healthy bacteria die out and the healthier ones will replicate themselves. This guarantees that the size of the bacterial swam will remain constant.
- Elimination and dispersal: There might be a gradual or sudden change in the environment where the bacteria live. As a result, a small percentage of the bacteria in a certain region will be liquidated or a group might be dispersed into another location. This has two effects on chemotaxis: the first is destroying the chemotactic progress, the second is that the new bacteria might be placed at locations with a better food source, thus assisting chemotaxis.

3.3 CGA

GA has the advantage of quickly locating high performance regions of vast and complex search spaces, but they are not well suited for fine-tuning solutions (Gendreau and Potvin, 2005), (Kazarlis *et al.*, 2001).

There have been several attempts to hybridize GA with other optimization algorithms. In (Mahfoud and Goldberg, 1995) the authors present the *Parallel Recombinative Simulated Annealing* (PRSA) which combines elements from the *simulated annealing* algorithm with others from GA. Another hybrid was presented in (Lee and Lee, 2005). This method hybridizes GA with *Ant Colony Optimization* (ACO).

On the other hand, BF possesses a poor convergence behavior over multi-modal and rough fitness landscapes. Its performance is also heavily affected with the growth of problem dimensionality (Biswas *et al.*, 2007).

To take advantage of the two optimizers, (Kim *et al.*, 2007) proposed a hybrid of GA and BF (called *GA-BF*). They validated their method on several test functions.

In another paper (Das and Mishra, 2013) proposed another hybrid of GA and BF which they called the *Chemo-inspired Genetic Algorithm* (CGA). Their motivation is that chemotaxis actually contributes the most in the search process, so instead of taking the whole BF to hybridize with GA, they only integrate the chemotaxis step in the hybrid with GA. CGA has five major steps: selection, crossover, mutation, elitism and chemotaxis. In addition, CGA employs three mechanisms: a) adaptive step size b) squeezed search space c) fitness function criterion.

4 EXPERIMENTS

We conducted intensive experiments to compare CGA with DEWPAA. The experiments were the same as those conducted in (Muhammad Fuad, M.M., 2012) and (Muhammad Fuad, M.M., 2013); i.e. classification task experiments of time series. The aim of the experiments is to show that using CGA in the optimization process of determining the weights of the segments in equation (2) will result in a lower classification error than that of DEWPAA.

As in (Muhammad Fuad, M.M., 2012), we computed the classification error using WPAAD for different compression ratios (1:8,1:12,1:16). We conducted our experiments using the UCR archive (Keogh *et al.*, 2011), which is the same archive used to test DEWPAA.

The two methods were tested on a classification task based on the first nearest-neighbor (1-NN) rule using leaving-one-out cross validation.

Table 2 shows some of the results of our experiments. As we can see from the table, CGA outperforms DEWPAA on almost all the datasets tested and for the different compression ratios.

5 CONCLUSIONS

In this paper we used a hybrid of genetic algorithm and bacterial foraging (CGA) to calculate the weights given to different segments of the time

Dataset	Method	Compression Ratio		
Dataset		1:8	1:12	1:16
Lighting7	CGA	0.329	0.342	0.356
Lighting /	DEWPAA	0.370	0.397	0.427
MedicalImages	CGA	0.308	0.308	0.321
Wiedicaiiiiages	DEWPAA	0.337	0.337	0.378
Gun_Point	CGA	0.06	0.06	0.06
Guii_Foiiit	DEWPAA	0.053	0.067	0.087
Coffee	CGA	0.179	0.179	0.179
Confee	DEWPAA	0.179	0.179	0.25
Lighting2	CGA	0.164	0.180	0.197
Lighting2	DEWPAA	0.213	0.230	0.246
MALLAT	CGA	0.077	0.082	0.094
WALLAT	DEWPAA	0.094	0.094	0.095
FacesUCR	CGA	0.238	0.238	0.238
racesock	DEWPAA	0.238	0.316	0.366
FISH	CGA	0.194	0.194	0.194
11311	DEWPAA	0.194	0.229	0.240
synthetic_control	CGA	0.067	0.067	0.100
synthetic_control	DEWPAA	0.110	0.113	0.113
CBF	CGA	0.017	0.021	0.021
СБГ	DEWPAA	0.017	0.021	0.037
ECG	CGA	0.090	0.100	0.100
LCG	DEWPAA	0.110	0.120	0.130
Trace	CGA	0.100	0.120	0.120
Trace	DEWPAA	0.140	0.180	0.190

Table 2: Comparison of the classification error between CGA and DEWPAA for different compression ratios.

series represented by the piecewise aggregate approximation representation method. The weights were obtained through an optimization process using chemo-inspired genetic algorithm (CGA) as optimizer and the fitness function is the classification error. Compared with differential evolution, another optimizer that we used in a previous work to solve the same optimization problem, CGA gives better results.

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