

# Evolutionary Associative Memories Through Genetic Programming

J. Villegas-Cortez

*Universidad Autónoma Metropolitana - Azcapotzalco, Departamento de Electrónica,  
Av. San Pablo 180 Col. Reynosa, México, D.F., 02200, México,  
e-mail: juanvc@correo.azc.uam.mx*

J.H. Sossa

*Centro de Investigación en Computación del Instituto Politécnico Nacional,  
Av. Juan de Dios Bátiz, esquina con Miguel Othón de Mendizábal,  
Col. Nueva Industrial Vallejo. México D.F. 07738, México,  
e-mail: hsossa@cic.ipn.mx*

C. Avilés-Cruz

*Universidad Autónoma Metropolitana - Azcapotzalco, Departamento de Electrónica,  
Av. San Pablo 180 Col. Reynosa, México D.F., 02200, México.*

G. Olague

*Centro de Investigación Científica y de Educación Superior de Ensenada CICESE,  
Km. 107 Carretera Tijuana-Ensenada, Ensenada, 22860, B.C. México.*

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Associative Memories (AMs) are useful devices designed to recall output patterns from input patterns. Each input-output pair forms an association. Thus, AMs store associations among pairs of patterns. An important feature is that since its origins AMs have been manually designed. This way, during the last 50 years about 26 different models and variations have been reported. In this paper, we illustrate how new models of AMs can be automatically generated through Genetic Programming (GP) based methodology. In particular, GP provides a way to successfully facilitate the search for an AM in the form of a computer program. The efficiency of the proposal was conducted by means of two tests based on binary and real-valued patterns. The experimental results show that it is possible to automatically generate AMs that achieve good results for the selected pattern recognition problems. This opens a new research area that allows, for the first time, synthesizing new AMs to solve specific problems.

*Keywords:* Computer science and technology; neural engineering; image quality; contrast; resolution; noise; image analysis.

Las memorias asociativas (AMs) son estructuras matemáticas específicamente diseñadas para recuperar patrones de entrada con patrones de salida. Cada par asociado (entrada-salida) forma una asociación, es así que la AM almacena las asociaciones entre los pares. Desde sus orígenes las AMs han sido diseñadas manualmente, y durante los últimos 50 años se han reportado un aproximado de 26 modelos de AMs con sus variantes. En este trabajo mostramos un nuevo modelo de AMs que es generado de forma automática por medio de Programación Genética. Este trabajo abre una nueva área de investigación que permite por primera vez sintetizar nuevas AMs para resolver problemas específicos. Para probar la eficiencia de nuestra propuesta la hemos aplicado para los casos de patrones en valores binarios y reales. Los experimentos muestran que es posible la generación automática de AMs para alcanzar buenos resultados para algunos problemas comunes del área de reconocimiento de patrones.

*Descriptores:* Ciencias de la computación y tecnología; ingeniería neuronal; calidad de imagen; contraste; resolución; ruido; análisis de imágenes.

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## 1. Introduction

Associative Memories (AMs) are simple and useful devices designed to recall output patterns in terms of input patterns by means of simple operations. AMs are considered as part of Artificial Neural Networks (ANN), but instead of using complex structures and operators based on transcendental or trigonometric functions, AMs work in two simple phases using two elemental operators, one for association and one for recall. During the association phase the AM is built in terms of the associations  $(X, Y)^k$ , for  $k$  an integer. Every pattern association is carried out by the application of simple operations such as: additions, multiplications, maximums, min-

imums, and others. Associative Memory  $\mathbf{M}$  is represented by a matrix, which is formed from all the pattern associations. The corresponding components  $m_{ij}$  can be seen as the synapses of a simple neural network. Operator  $\mathbf{M}$  is generated from a preset of finite known associations called the fundamental set; this association set is represented as  $\{(X^k; Y^k) | k = 1, \dots, p\}$ , where  $p$  is the number of associations. If  $(X^k = Y^k) \forall k = 1, \dots, p$ , then  $\mathbf{M}$  is considered auto-associative, otherwise it is hetero-associative. If a distorted version of a pattern, denoted as  $\tilde{X}$ , is fed to  $\mathbf{M}$  and the obtained output is exactly  $Y^k$ , then recalling is considered as perfect. The simplicity of AMs mod-

els is a result of a great development, which has been carried out during the last 50 years; for some examples, refer to [6,22,24,14,15,25,23,31]. Most of these models have several limitations: limited storage capacity, difficulty to deal with more than one type of pattern (binary, integer or real-valued), lack of robustness to different kinds of noises (additive, subtractive, mixed, Gaussian, etc.); and above all, most of these models work only for the purpose for which they were specially designed and not for any other [32]. All these models have been manually developed. The process of designing one model could take one or two years.

A first attempt to automatically generate AMs has been reported in [16]. This method was conceived for the case of bidirectional associative memories (BAM). On the other hand, the closest work to the approach presented in this paper is concerned with the automatic design of Artificial Neural Networks [13], which uses some kind of genetic programming (GP) for this purpose. Another reported work applies Particle Swarm Optimization [4]. Bearing in mind the relevant results obtained with GP techniques versus Genetic Algorithms [21], in this work we describe a GP-based methodology to automatically generate AMs. GP has been successfully applied in a huge range of areas. In particular, we can mention some related to computer vision [1,29,10,12,27,28,35,36,20,26].

In this paper, we deal with the problem of automatically generating AMs through GP; previous results can be found in [33]. We improve our initial technique and provide a better description of the problems involved in the recall of binary and real-valued patterns.

The rest of the paper is organized as follows. In Sec. 2 we provide a brief description of our first approach fully described in Ref 33. In Sec. 3 we present our new co-evolutionary based-GP methodology for the automatic generation of AM through GP. In Sec. 4 we show the experimental results for some representative databases. Finally, in Sec. 5 we present the conclusions and suggestions for further research.

## 2. Automatic design of AMs through GP

The proposed model for generating AMs using evolutionary algorithms rests on the following two-stage rule. (1) Evolution starts as a simple ANN with a pre-established topology that is evolved from more than one connection [37]. (2) Later, evolution of architecture is developed with the aim of generating different topological structures. Common AM models fix the connection structure and consider the full aspects of every design such as the capacity of memory recall, as well as noise suppression. Our proposal is based on the linear associator concept introduced in [7], in order to consider the general association aspects between pattern sets, but bearing in mind the local association parts focused on the connectivity of every synapse that is carried out between the vector pattern components. Taking into account our first work, we present a very important modification using the co-evolutionary paradigm. As we will see, this allows modeling the two processes in AM designing: association and recall, with the idea of preserving the local pattern aspects.

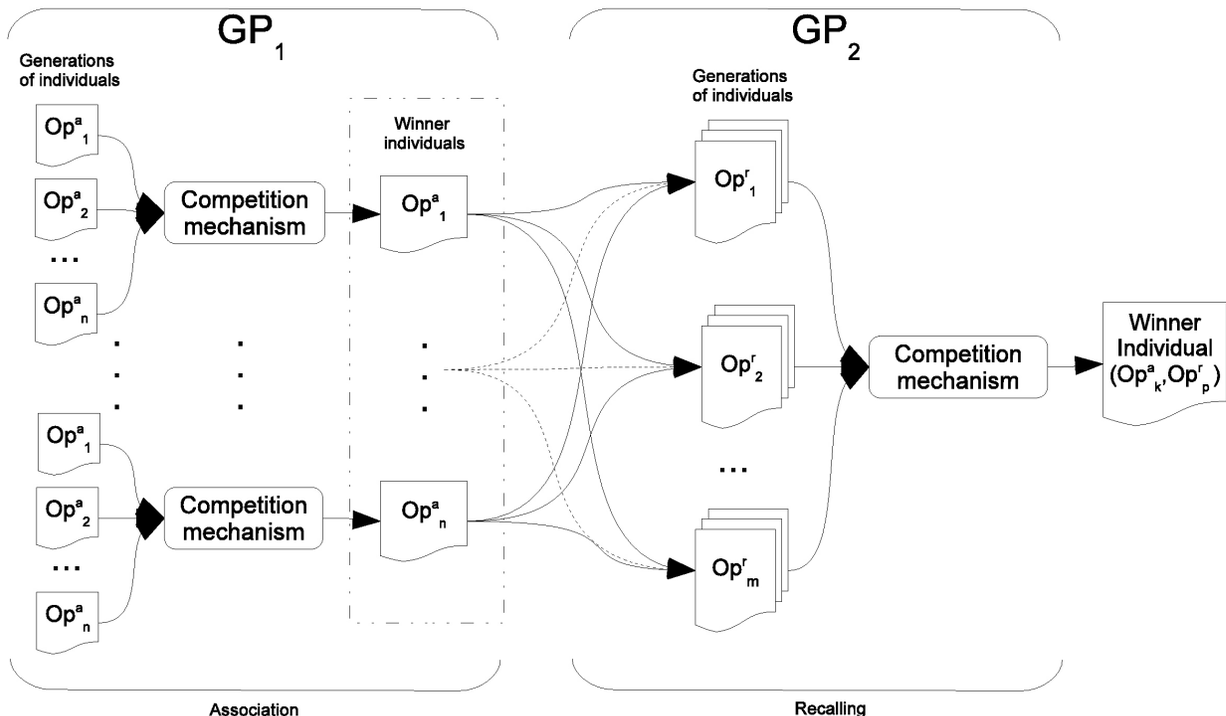


FIGURE 1. Framework of our cooperative co-evolutionary process.

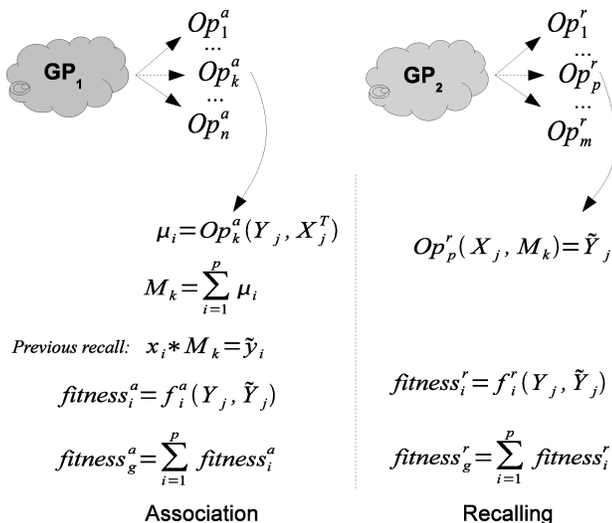


FIGURE 2. Proposed co-evolutionary model for the GP-development of AMs.

Our proposal outlines the evolutionary process in three steps and two phases. First, we define the function operators used during association; second, we define the function operators and terminal elements for recalling, considering the fundamental aspects of the association matrix previously built; and thirdly, we implement the co-evolutionary approach [2] by joining two evolved individuals to create a new one.

### 3. Design of AMs

Our proposed GP based co-evolutionary model comprises two populations, one for association, and one for recalling. This is a first contribution with respect to previous works that considered only one evolutionary process [33]. In this paper, we consider two sequential processes instead of only one bearing in mind the cooperative co-evolutionary paradigm, we consider as more adequate to reach our goal of developing a simulated correlation between species. Thus, population represents a separate part of the problem, in such a way that every species cooperates to have a global solution in two considered phases. Hence, every individual is a possible solution for the problem in the pre-set common environment. The basic idea of co-evolution consists in using the concept of dividing and conquering the problem (system) into many sub-problems (sub-systems), and evolving them separately, then later combining them for implementing the entire solution [11].

Figure 1 shows the framework for implementing the cooperative co-evolutionary process. Here, the solutions (appropriate AMs fitting one pattern set with a perfect recall) come from two search spaces. The first process ( $GP_1$ ) is a clustered process developed to produce individuals for the association stage of the AM, where every individual identified by each cluster is the winner of one evolutionary process. Then, all winner individuals participate in the solution of the second process ( $GP_2$ , the recalling one), by recombining

ing their learned features. After the whole process is completed we get an individual global winner.

Fitness function of the whole process is implemented as the minimum of all the individual pairs after the second phase. The components of our new model are defined in terms of both functions and terminal sets per every population for each GP evolutionary process, see Fig. 2. Thus, the co-evolutionary model is composed of the following aspects:

- $Op_k^a$  represents the evolutionary operator used in each pattern association. Its representation is generated as an individual in the form of a tree. This correspond to the encoded genotype. The local association matrix  $\mu_i$  is integrated by the  $Op_k^a$  operator acting on the components of each vector of the local association  $\{Y_j, X_j^T\}$ .
- $M_k$  describes the general association matrix. It is taken as the sum of all local associations and provides the cumulative knowledge inspired by the perceptron principle.
- $T_a$  is the *Terminal Set* for the association stage. It consists of a set of nodes taking the vector entries, such as  $T_a = \{x_j, y_j\}$ . These nodes belong to each pattern-vector  $\{X\}$  and  $\{Y\}$ ; this is in order to have a local encoded correlation input.
- $F_a$  is the *Function Set* for the association stage. These functions have been defined by considering the solutions reported in literature in order to target the space of possible structures with the aim of producing individuals similar in performance to the reviewed AM models:  $F_a = \{+, -, \wedge, \vee, \times\}$  where  $\wedge, \vee$ , and  $\times$  are the *minimum, maximum* and *multiplication* operators
- $Op_p^r$  is the evolutionary operator for pattern recall. It comprises the input vector  $X_j$ , as well as the matrix association  $M_k$  generated by the previous evolutionary association operator.
- $T_r$  is the *Terminal Set* for the recalling stage,  $T_r = \{v, R_1, R_2, \dots, R_m, M_k\}$ ; with  $v \in X$ , as the input vector, and  $R_i$  as the  $i$ th-row-vector belonging to  $M_k$ , which is the corresponding  $k$ th-AM.
- $F_r$  represents the *Function Set* for the recalling stage.  $F_r = \{+, -, \wedge, \vee, \otimes\}$ ;  $\otimes$  is defined as the multiplication operator between vector components,  $\otimes(X, Y) = [x_1 * y_1, x_2 * y_2, \dots, x_n * y_n]$ , and when the association matrix is considered is defined as follows  $\otimes(X, M_k) = X * M_k$ ; thus it satisfies the dimensionality of the multiplication operator between matrices.
- $\tilde{Y}_j$  is the approximated  $Y_j$  pattern resulting from the application obtained with the operator rule  $Op_p^r$  in the recalling stage.

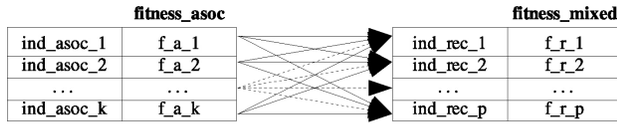


FIGURE 3. Fitness model for the GP-development of AMs.

TABLE I. Winner individuals (pairs) of the association rules (first column in competition), with its corresponding recalling rule (second column), and the global fitness attained in cooperation. The indexed operators in the first column are related to each association rule. Operator  $Op_1^a$  is related to Fig. 7(a); numbers two and three are related to Figs. 7(b) and 8. Operator  $Op_1^r$  in the second column is related to the operator depicted in Fig. 9(a), and so on.

Association rule	Recalling rule	Global Fitness
<i>Competition</i>		
$(Op_1^a, Op_2^a, Op_3^a) := Op_1^a$	$Op_1^r$	1
<i>Competition</i>		
$(Op_1^a, Op_2^a, Op_3^a) := Op_1^a$	$Op_2^r$	1
<i>Competition</i>		
$(Op_1^a, Op_2^a, Op_3^a) := Op_1^a$	$Op_3^r$	1

The fitness function which provides the qualification to every process as an essential evolutionary step in the whole process is requested. For our purposes the fitness value is applied at the end of both stages (association and recall). The fitness function covers the sum of all the local fitness (related to local associations), and all the recalling set (one recalling for every pattern relation), as shown in Fig. 2. For this evolutionary approach we considered the normalized correlation coefficient between the goal ( $Y$ ) and the source processed pattern ( $\tilde{Y}$ ) [18]. The fitness function  $f$ , known as *similarity* ( $0 \leq f \leq 1$ ), is defined as follows:

$$f = \frac{Y \cdot \tilde{Y}}{\sqrt{Y \cdot Y} \sqrt{\tilde{Y} \cdot \tilde{Y}}} \quad (1)$$

where  $Y$  and  $\tilde{Y}$  are vectors of size  $1 \times N$ , and  $Y \cdot \tilde{Y}$  is given by the following equation:

$$Y \cdot \tilde{Y} = \frac{1}{N} \sum_{j=1}^N Y(1, j) \cdot \tilde{Y}(1, j) \quad (2)$$

Function  $f$  tries to maximize the number of matching component vectors  $Y$  and  $\tilde{Y}$ . This seems a reasonable choice for the fitness function. The optimum is found when  $f = 1$  corresponds to the matching of all pixels. The worst case takes place for  $f = 0$ , implying that not a single pixel matches.

Thus, we used Eq. (1) as our fitness function, and was applied in the evaluation of every generated individual  $\mu_i$ . For the association stage, the training one, between the source set  $X$ , and the goal set  $Y$ , both conformed our *fundamental sets*. The fitness evaluation is carried out in three steps: First, the *association* between pattern sets  $X$  and  $Y$  using the first stage

operator  $Op_k^a$ ; second, we fix the *multiplication* operator for the recalling task just to have a first estimation of *multiplication* operator. After this we have one recalled pattern set,  $\hat{Y}$  and this local fitness is considered to have the winner individual for the first evolutionary process  $GP_1$ , which is taken into count for the  $n$  batch processes (see Fig. 1, the association block). Second, the  $n$ -winner individuals from the first process are considered for the second  $GP$  process, the recalling-one. At the end of this block, the fitness function is applied again to get the winner individual for every  $m$ -population as a local estimation to each operator  $Op_a^r$ . Here we wave a pair of operators as one individual ( $Op_k^a, Op_a^r$ ), thirdly, we consider the global fitness for each pair of operators as the high-fitness grade for the solution pair. To follow the explanation, refer to Figs. 1, 2 and 3.

### 3.1. GP experiments and parameters

All the experiments were implemented on workstations with 64-bit architectures using Matlab with GPLab toolbox ver. 3.0 [17]. The co-evolutionary process was composed of two phases, association and recalling. They were performed in two batch processes  $n$  and  $m$ : GP1 and GP2, respectively. After 50 generations with 70 individuals per every evolutionary process, we got one evolutionary solution for our problem.

For the experiments we considered the GP parameters similar to those suggested in [8], rate values of 0.7 and 0.3 for the crossover and mutation operators respectively were set up. In order to initialize populations, a ramped-half-and-half initialization method for the mutation was used.

## 4. Results and analysis

In this section we present the set of experiments we have performed to test the efficiency of the proposed methodology. For this we have used two datasets. In the first case we show how the proposal works. In the second we show the application of the proposal to solve one classic pattern recognition problem. For the first experiment we used simple vector sets consisting of binary sets of digit characters from 0 to 9, within a matrix size of  $7 \times 5$ , as shown in Fig. 4.

For these simple images we considered the line vector per matrix where  $X$  and  $Y$  are the source and the goal vector sets, respectively, using an auto-associative relationship.

The application of our methodology generates a set of rules for association, and another set of rules for recall. Each of these rules is co-evolved with the aim of maximizing the global fitness. The winner is one pair set as shown in Fig. 5. The AM shown in Fig. 5 is only one example of pairs selected from a vast production of individuals, which fits to this kind of pattern perfectly.

The solution depicted in Fig. 5 shows that an important describing feature for this particular pattern set, is row number 7, which belongs to the association matrix  $M_k$ . It is the only significant one during recalling regarding the auto-





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