Evolving Novel Image Features Using Genetic Programming-based Image Transforms

Taras Kowaliw, Wolfgang Banzhaf, Nawwaf Kharma, and Simon Harding

Abstract—In this paper, we use Genetic Programming (GP) to define a set of transforms on the space of greyscale images. The motivation is to allow an evolutionary algorithm means of transforming a set of image patterns into a more classifiable form. To this end, we introduce the notion of a Transform-based Evolvable Feature (TEF), a moment value extracted from a GP-transformed image, used in a classification task. Unlike many previous approaches, the TEF allows the whole image space to be searched and augmented. TEFs are instantiated through Cartesian Genetic Programming, and applied to a medical image classification task, that of detecting OPMD-indicating inclusions in cell images. It is shown that the inclusion of a single TEF allows for significantly superior classification relative to predefined features alone.

I. Introduction

In this paper, we explore means of augmenting the typical use of features in classification problems. Due to increased capacity for computation, we may now, in a training phase, explore means of searching image databases directly for novel database-specific features. Hence, rather than apply only a general set of predefined features, or devoting significant human effort to the careful design of database-specific features, we may use techniques of machine learning in the direct creation of new features well suited to some given problem.

Here, we will attempt to evolve transformations on the space of images, in hopes that particular transforms which emphasize distinguishing characteristics may be found. These transforms will be genetic programs (GPs), optimized through an evolutionary algorithm. A set of moments describing the transformed image will be extracted, and used for classification. The introduction of GP-based transforms to the classification task brings two desireable properties to the pattern recognition process: firstly, if allows for the search of the entire pixel-space of images, not just a collection of predefined features, potentially finding patterns missed by human designers; and secondly, it allows for recombination of this pixel-space view of the database into forms useful to a classifier. This latter point may be viewed as a form of Support Vector Machine, defined on the pixel-space, using a GP kernel.

Our domain of application will be a binary classification problem based on the detection of a form of Muscular Dystrophy in cell nuclei. A database of pre-segmented cell

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images has been procured, and we will attempt to train a classifier to recognize human-defined ground truth. We will show that the addition of a single evolved TEF to our database will improve recognition by 38% relative to a set of predefined features often used for cell classification, obviously allowing evolution to successfully exploit our additional information sources. In so doing, we will provide evidence for the efficacy of an evolutionary design of image features, and the potential for the automation of pattern recognition.

II. REVIEW

A. Predefined Features and Feature Selection for Image Classification

Image classification (recognition) typically uses a set of features to canalize the pattern space. A great deal of effort is spent on the design, selection and weighting of features. Often, practitioners begin with a set of "standard" features, then use some machine learning technique to find the most appropriate choices; these "standard" features often involve statistical moments, or entropy- or histogram-based measures (as in [1]). Domain specific measures are used as well, such as the nucleus-specific measures in the construction of the Wisconsin Breast Cancer Database [2].

Let an image space (some finite, non-trivial square of pixel locations) be denoted I, and an image on that space be denoted f(I). Each pixel $p=(p_x,p_y)\in I$ is associated with an intensity, $f(p)\in[0,1]$. Moments are statistical descriptions of some given data, often used to break down an image into numeric values which can be sent to a machine learning technique. In discrete form, geometric moments are often computed as

$$M_{mn} = \sum_{p \in I} p_x^m p_y^n f(p)$$

Central moments are computed similarly, although relative to the centre of mass:

$$C_{mn} = \sum_{p \in I} (p_x - \bar{x})^m (p_y - \bar{y})^n f(p)$$

where $(\bar{x}, \bar{y}) = (M_{10}, M_{01})$ is the centroid. From these moments it is possible to define a set of scale- and rotation-invariant moments, known as Hu's moments [3].

The *entropy* of an image is typically defined to be the entropy of the image histogram. That is, the histogram (\bar{h}) is interpreted as a probability distribution on the space of pixel values, then the Shannon entropy is computed:

$$E(\bar{h}) = \sum_{i=0}^{255} -h_i \log h_i$$

where we have assumed an 8-bit greyscale image. Entropy can be normalized by dividing by the maximum value, here $\log 256$.

The *fractal dimension* of an image, as computed through a box-counting algorithm, is a rough measure of self-similarity of an image. First, the image is thresholded, to produce a black and white version. Next, a sequence of boxes are computed, those boxes being used as discretizations of the image. For each box size, the number of boxes required to cover the image is computed. Finally, the log over the box sizes is plotted against the log of the number of boxes: the coefficient of the line of best fit (by linear regression) is considered the fractal dimension.

Following previous work with cell classification [2], we also may define several features specifically for cell images. Given a pre-segmented cell image, we may define the *perimeter*, *compactness*, *mean radius*, *radius variance*, and *compactness* of the cell boundary. Further, having divided the image into nuclear interior and exterior, we can compute the *area* and greyscale *variance* of the interior.

B. Feature Creation

By feature creation, we refer to the creation of numerical methods drawn from the raw images in a database. This is a contrast with some usage, which refers to the pre-processing of values of predefined features prior to classification (as in [4]).

The most significant research dealing with the issue of feature creation may be divided into four categories.

Historically speaking, the first category contains methods that did not use artificial evolution in any form. This category include the work of [5], [6], [7], [8]. All of them used mask-or pixel-based features; evaluated them using a reference pattern; had no or limited higher-level feature generation; and were not ready for real-world application.

The second category comprises methods that employed polynomial functions of features extracted from the target pattern. These include [9], [10], [11]. All of these techniques employ primitive features that are statistical functions of discrete (or discretizable) data signals; create complex features, in the form of polynomial functions of the primitive features; and most have applications in the area of machine fault diagnosis.

A third category covers those methods that involved true Genetic Programming techniques. The most notable early efforts in this area includes that of [12], [13]. Both methods evolve a Lisp-style program which controls a roving agent that uses any or all of five evolvable Boolean functions or masks to correctly identify a character. What distinguishes these works is that they evolve both the features and the classifier that uses them.

The final and most modern category is that of Evolvable Pattern Recognizers. These are perhaps the most ambitious research projects. Two efforts are worthy of special mention: HELPR [14] and CellNet [1], [15]. HELPR evolves feature detectors, but the classification module is completely separate from the feature extraction module. CellNet blurs the line

between feature selection and the construction of binary classifiers out of these features. Other differences exist, but, both attempts are the only systems, that we know of, that aim at using artificial evolution to synthesize complete recognition systems with minimum human intervention.

C. Genetic Programming and Image Processing

Cartesian Genetic Programming was originally developed by Miller and Thomson [16] for the purpose of evolving digital circuits. It represents a program as a directed graph, or, in our specific case, provides a graph mapping inputs to an output. One of the benefits of this type of representation is the implicit re-use of nodes in the directed graph. The technique is also similar to Parallel Distributed GP, which was independently developed by Poli [17], and also to Linear GP developed by Banzhaf [18]. Harding and Banzhaf have used Cartesian Genetic Programming to reverse-engineer common image filters [19]. CGP has also been used to configure the logic blocks in specialized, parallel hardware developed, applied to the re-evolution of known image filters [20], [21]. Evolved image filters have been used in real-world problems, such as the detection of mud slides [22].

III. THE MODEL

Here we describe the use of CGP as a means of defining transformations on the space of images, and the use of these transformations as evolvable features. Theoretically, nearly any form of Genetic Programming could have been used, CGP was chosen due to previous image processing success. Further, although we apply to medical images, we expect this approach to be applicable to a wider range of image classification tasks.

We use CGP as described by Harding and Banzhaf [19], utilizing a one-dimensional topology, and infinite levels-back. Each node in the graph has two connections to previous nodes or inputs (x and y), and uses the function set $\{max\{x,y\}, min\{x,y\}, x\cdot y, x+y, x/y, x-y, const., |x|, x^2, x^y\}$, where division is "safe" (i.e. returns 1 when the denominator is very close to 0). The number of inputs is a system parameter, and drawn from a square neighbourhood surrounding a given pixel. The inputs are drawn from this square neighbourhood from the top left corner, left to right, top to bottom. The output is the value of the final node in the graph.

An example image of a CGP with 16 inputs is shown in Figure 1. Note several forms of "neutral" code: the green, blue, and pink nodes use one or less of their two inputs; the second-last node (grey) does not connect to the output node (pink). Removing redundancy, this CGP reduces to the function:

$$\phi(i_5, i_6, i_{12}) = (i_5 i_6)^{(34.72(i_{12})^2)}$$

CGP graphs are easily evolvable. Crossover is defined classically (i.e. single-point swap), treating the graph as a list of nodes. Mutation operates on the node types, connectivity, and neighbourhood size, treating each element in turn with equal probability.

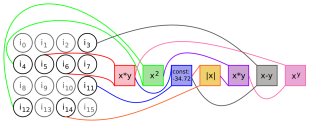


Fig.

ILLUSTRATION OF A CGP GRAPH WITH 16 INPUTS (CIRCLES) AND 7 NODES (SQUARES). THE OUTPUT IS THE VALUE IN THE FINAL NODE.

Fig. 2

EXAMPLES OF HEALTHY (top row) AND SICK (bottom row) CELL

IMAGES.

A. CGP Image Transformations

We consider the application of a CGP graph to a pixel and its neighbourhood. As a neighbourhood, we will consider squares surrounding the sources pixel of integer size.

We shall write the output of an arbitrary CGP graph G applied to a list of k values as $G(x_1,...,x_k)$. Given some location $p \in I$, we apply CGP graph G to pixel p and neighbourhood of size n, denoted $\bar{p} = \{p_0, p_1, ..., p_{n^2}\}$, as follows:

$$G(f(\bar{p})) = G(f(p_0), f(p_1), ..., f(p_{n^2}))$$
(1)

i.e. we retrieve the pixel values for the k^2 pixel neighbours of pixel p and feed them to the CGP graph in order. Note that if f(q) does not exist for some pixel q (beyond the edges of the image, say), then we return the value f(q) = -1. Further, if $G(f(\bar{p})) \not\in [0,1]$, we replace by the closest boundary value, either 0 or 1.

Given some image f(I), we can define a new image, G(f(I)) as follows: Let I' be an image space of same dimensions as I. For each pixel $p' \in I'$, let $f(p') = G(f(\bar{p}))$. Hence, every CGP graph G can be viewed as a function on the space of images.

This is but one of many different sorts of filter functions, and hence is a limit on the space of what can be (easily) represented. Generalization is a future endeavour of ours, but beyond the scope of this research.

IV. IMAGE PATTERNS AND EVALUATION

Features are evaluated on the basis of their ability to separate images in a target database. The database chosen, CellsDB¹, was collected by the Centre hospitalier de l'Université de Montréal (CHUM), where the causes and associated symptoms for OPMD at the genetic and cellular level have been studied extensively [23]. Intranuclear inclusions (INIs) have been detected via both pathological studies and electron microscopy; These INIs were tubular, about 8.5 nm in external and 3 nm in inner diameters, up to 0.25 μ m in length, and converged to form tangles or palisades [24]. Detection of these inclusions is expected to

¹Since undertaking this work, it was shown that some of the images in CellsDB were misclassified: In proceeding discussions, several images from the training set, and two images from the validation set were mislabeled. This will not effect reported validation accuracies by more that 1%.

lead to the detection of OPMD. Hence, we seek to be able to find features which separate images of cells on the basis of whether or not they contain INIs. Detecting INIs, as opposed to other intranuclear patterns, is a difficult task, requiring training for human classification.

CellsDB was collected and prepared by Tarundeep Dhot at the Centre for the Study of Brain Diseases at CHUM, and is pre-segmented so that each image contains a single cell; It is a collection of images of cells taken at 10x, 20x, and 40x zoom, divided into two categories associated with the presence or absence of inclusions indicating OPMD: "healthy" and "sick". An example of some cell images may be seen in Figure 2. Data from the CellsDB has previously been used for image processing [25], [26].

We wish to award fitness to any particular feature based on its ability to distinguish between healthy and sick cells. To do so, we convert the database to a set of features, then attempt to classify the cells using some given classifier and evaluation technique. The classifier will return a false positive rate, FPR, for both classes, which we combine to a measure of *sensitivity-specificity*.

$$SS = (1 - FPR("healthy"))(1 - FPR("sick"))$$
 (2)

Hence, $SS \in [0,1]$ is maximized for perfect recognition. It is trivial to include weights for the various classes into such a function, as may be necessary for medical application, and it is unlikely that evolvability will be effected by such weighting.

The features and evaluation method changed between training and validation runs. The pattern database was broken into two sets: 186 healthy and 200 sick cell images for training, and 200 healthy and 200 sick cell images for validation. All classifiers and evaluation techniques were implemented via the Weka machine learning system, version 3.5.7 [27].

For training runs, a set of 16 features were computed for the TEF-transformed images: $\{M_{00}, M_{10}, M_{01}, C_{11}, C_{20}, C_{02}, C_{12}, C_{21}, C_{22}, H_1, ..., H_7\}$, where H_i is Hu's *i*th invariant moment. These 16 features of transformed images were trained and evaluated on the training set of images using 5-fold cross validation.

For validation runs, a set of 40 features were computed: Firstly, the 16 + 8 predefined moments applied to

untransformed images, consisting of the 16 aforementioned moments, along with the set {threshholded area, variance, mean radius, radius variance, perimeter, compactness, entropy, fractal dimension}. Secondly, the 16 aforementioned moments applied to the transformed images. These 40 features were trained and evaluated on the validation set of images using 10-fold cross validation.

The reason that a smaller set of features was used in the definition of the training evaluation is as follows: randomization of the database prior to classification is a useful feature to help prevent overfitting, but leads to a stochastic fitness function. Beginning with a set of features which is already adept at classification, as the predefined features are, makes the variance due to stochasticity greater than fitness gains in early evolution, hence hindering the selection operator.

V. EXPERIMENTS

A. Initial Classification Experiments

Initial classification attempts were undertaken using the 24 predefined features and moments on CellsDB. The classifiers used included Decision Trees, Ridor rules, and k-NN. These were run 40 times and averaged to include the variance associated with randomization of the database patterns. Next, we evaluated the performance of randomly-generated TEFs under classification. For each of the above classifiers, a randomly generated TEF was instantiated, and the 16 additional moments generated by the transform were added to the database. This was repeated 40 times, and averaged. The results are shown in Table I.

As can be seen, there is little difference between the SS value for simply the pre-defined features and moments, and the inclusion of a randomly defined TEF. However, the variance of values increases under the addition of a TEF, as the new feature values allow for helpful or misleading / confusing information to the classifier. Note the additional time requirements for additional moment calculations, which clearly dominate the process.

Several feature evaluation routines were used in an attempt to improve classification accuracy (Information gain attribute evaluation, χ^2 attribute evaluation, and principal component evaluation), but none significantly improved performance (indeed, most significantly lessened classifier accuracy).

B. TEF Evolution

We used a standard evolutionary algorithm, as described by Eiben and Smith [28]. Following an informal parameter search, the following parameters were chosen for the evolutionary algorithm:

initial pop. size	400	pop. size	200
prob. mutation	0.02	prob. crossover	0.6
prop. elite	0.01	tournament size	3
mask size	6×6	CGP graph size	100

Each EA was run for 50 generations, using a 1-NN classifier. 40 runs were undertaken in total.

Evolution was quite successful at increasing fitness (training SS), which increased from a mean best fitness of 0.523

in the first generation (averaged over 40 runs, s.d. 0.026), to a mean best fitness of 0.620 (s.d. 0.046). In a few cases, evolution optimized training SS at the expense of validation SS, but a good general increase in the latter was seen in most runs. Mean best validation SS for the final generation was 0.676 (s.d. 0.052), and maximized at a value of 0.766. This is a 32% improvement over the expected performance of the best classifier found for predefined features alone.

C. Best Discovered Transform

In this section, we have extracted the best evolved transform and will explore it in more detail. The best individual of the final (50th) generation of the highest validation fitness run was selected. This best individual had a sensitivity-specificity of 0.766, encompassing true positive rates of 0.878 for healthy cells and 0.873 for sick cells.

The same evolved attribute values, when evaluated using a J48 Decision Tree instead of 1-NN, gives a sensitivity-specificity of 0.801, encompassing true positive rates of 0.912 for healthy cells and 0.878 for sick cells, or a 38% improvement over the expected performance of the best classifier found for predefined features alone.

The best transform, once neutral code is removed, may be written as the following neighbourhood function:

$$o(i_0, ..., i_{35}) = \min\{i_{11} - i_{15}, (i_9 - \max\{i_6, i_{25}\})^{i_8}\}\$$

The graph view is shown in Figure 3. This function returns black (0) nearly always, except when both $i_{11}-i_{15}$ and $(i_9-\max\{i_6,i_{25}\})^{i_8}$ are (relatively) high. The first subsection ensures variance on the right of the neighbourhood is high (excluding inclusions that are too large), and the second subsection ensures that the left and bottom-left are mostly white, excluding inclusions too small. Hence, we detect the right half of inclusions of the proper size and variance. Note that this function works for several different microscope zoom levels simultaneously.

While non-OPMD indicating inclusions are still highlighted using this function, the OPMD-indicating inclusions are highlighted with more intensity, hence allowing for a greater recognition of the cells than original image features alone. Examples of output are shown in Figure 4.

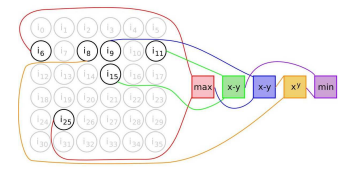


Fig. 3
ILLUSTRATION OF BEST DISCOVERED CGP.

TABLE I Comparison of classification on standard database (\emptyset) to database augmented by a randomly-generated TEF. All figures averaged over 40 runs, with standard deviation in brackets.

classifier	transform	SS	TPR(H)	TPR(S)	time ms
decision	Ø	0.4945 (0.0201)	0.7223 (0.0349)	0.6858 (0.0335)	369.3 (101.1)
tree	random	0.5213 (0.0800)	0.7236 (0.0633)	0.7177 (0.0534)	77883.4 (9135.9)
ridor	Ø	0.4606 (0.0236)	0.7025 (0.0841)	0.6646 (0.0829)	473.3 (109.6)
	random	0.5034 (0.0586)	0.7061 (0.0735)	0.7172 (0.0825)	76374.3 (12943.5)
1-NN	Ø	0.5166 (0.0117)	0.7323 (0.0137)	0.7056 (0.0126)	601.9 (73.2)
	random	0.5527 (0.0698)	0.7585 (0.0472)	0.7262 (0.0464)	99860.2 (16031.5)
5-NN	Ø	0.5803 (0.0157)	0.7735 (0.0182)	0.7503 (0.0122)	174.4 (99.6)
	random	0.6108 (0.0670)	0.7993 (0.0582)	0.7623 (0.0302)	77368.5 (13933.3)

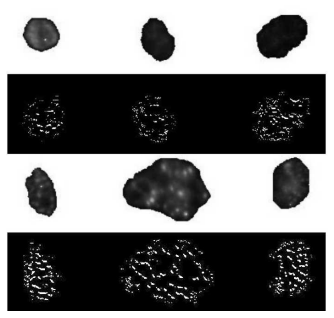


Fig. 4

EXAMPLES OF OUTPUT OF BEST DISCOVERED TEF: VIEW OF ORIGINAL CELL IMAGES (ON WHITE) AND IMAGES SUBJECTED TO ACTION BY CGP (ON BLACK). TOP ROW ARE HEALTHY, BOTTOM ROW ARE SICK.

TRANSFORMED IMAGES HAVE BEEN SUBJECTED TO CONTRAST STRETCHING TO MAKE DISTINCTION BETWEEN BLACK AND DARK GREY MORE VISIBLE.

Feature selection was run on the database of the original moments and the evolved moments, using an Information Gain Attribute Evaluator. The evaluator selected 30 of the features as significant, discarding the other 10. The top ten ranked attributes, by information gain, were all moments of the evolved transformed image.

VI. CONCLUSIONS AND FUTURE DIRECTIONS

In this paper, we have defined a mechanism for the automated discovery of novel database-specific features via evolutionary computation. This system has been applied to a new and significant recognition problem in medical imaging, and shown to significantly improve overall performance. In the short term, we have discovered a set of features that will achieve a 91% recognition for healthy cell images, and a 88% recognition for sick cell images, a 38% improvement over predefined features alone. More generally, we have achieved a significant advance in the capacity of automated systems to automatically adapt to a given pattern database without human expertise, or, perhaps, with greater efficacy than human designs.

To properly evaluate the power of this new approach, several future directions immediately suggest themselves: (a) to apply the same framework to several different databases, showing both generality and the capacity for automated adaptation; and (b) to extend the system to use a collection of features rather than a single graph, or to evolve the choice of moments simultaneously. These steps bear potential to advance the goal of a fully-automated pattern recognition system. Both steps are currently being explored at the Memorial University of Newfoundland.

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