Non-negative matrix factorization for spectral colors using genetic algorithms: Substantially Updated Version

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SUMMARY In this paper we introduce a novel method for non-negative matrix factorization (NMF) using a genetic algorithm. The method finds the optimal basis functions for the spectral colors in both spectral and color spaces. We show that one version of the proposed algorithm works as well as the standard NMF algorithm in spectral space. Further, this algorithm is modified to obtain a functionality to work in color space which the standard NMF is currently not capable of providing. The modification involves optimization in color space reducing the approximation error by a factor of 6 for Macbeth ColorChecker colors. The algorithm can be used in digital camera design. In addition, we propose an algorithm based on multiobjective optimization in both spectral and color space which can be used in digital image archiving.

key words: Genetic algorithm, spectral colors, Macbeth ColorChecker, non-negative matrix factorization

1. Introduction

Non-negative Matrix Factorization (NMF) gives a representation of the latent data structure and reduces the dimensionality of a dataset. There are at least two good physical reasons to use the NMF technique for spectrometry and imaging: First, optical sensors generate non-negative signals, that may benefit from non-negative decomposition, and, second, the optical color filters can be physically implemented only with non-negative spectral characteristics [1]. In multispectral imaging, for example, the technique may be used to calculate digital camera spectral band characteristics and is particularly attractive for spectral reflectance estimation using the Wiener method [2].

NMF solves the problem of non-negative data decomposition [3–6]. NMF has been used in spectral unmixing for non-resolved space object characterization [3]. NMF and a related technique called non-negative tensor factorization (NTF) were also used to define the optimal non-negative representation of spectral colors for different sets including the Macbeth ColorChecker (MCC) and Munsell color set [1, 7]. The present NMF methods represent measured spectra by a few non-negative basis functions. NMF determines non-negative factors $W$ and $Z$ and implements non-negative factorization of a given matrix $X$ as follows: $X \approx WZ$. The size of $X$ is the number of wavelengths times the number of spectra, the columns of $W$ are the basis functions found to minimizing the error of the approximation, and $Z$ is a matrix of weights.

The present work investigates a common form of 3-primary filter systems that are standard in the digital camera industry. Practically speaking, if spectral response profiles of camera color sensors are designed to equal color matching functions of a human Standard Observer (such as $X$, $Y$, $Z$ of the CIE 1931 XYZ color system), then fidelity between a digital image and a scene’s original color should be achieved. In practice, however, such designs do not yield correct digital color capture and reproduction. One reason for such color fidelity failures is that display device controls commonly employ linearly transformed color matching functions $r$, $g$, $b$ of the CIE 1931 RGB color system. Such transformations pose a problem because, while the positive portion of $r$, $g$, $b$ functions yield the full range of colors reproducible by the display, the negative portions of these functions are usually ignored in digital camera designs, which makes the physical reproduction of all colors impossible [8–10]. This mismatch between acquired and displayed image information suggests further study is needed to optimize color matching functions for digital camera color image acquisition. Principal component analysis (PCA) may seem a natural approach to this problem, however, due to orthogonality constraints, PCA basis functions usually contain negative values which complicate physical implementation. Alternatively, the present approach using NMF provides non-negative decomposition which is appropriate and may provide a modeling solution. This seems plausible because the basis functions found by NMF optimized in spectral space provide approximate representations of basis function shapes, wavelength subranges and locations of modes for color matching functions (Fig. 1). Previously published results [1, 7] have provided limited findings towards approximating spectral optimization, however, the present research aims for the accurate approximation of colors, as is needed to fully solve digital capture and color rendering problems.

In this paper we propose a non-negative matrix factorization approach that uses a genetic algorithm (GA) as an optimization technique. Our aim is to find non-negative basis functions that provide accurate color reproduction for a given spectral dataset. Three forms of novel algorithms are developed: non-negative matrix factorization in spectral space; non-negative matrix factorization for color space; and
optimization in both spectral and color spaces. Although the separate algorithms can be thought of as integral to a coherent system, for convenience and ease of presentation, we implement and demonstrate each of these forms here as a separate algorithm. Below we test the utility via a series of demonstrations. We first investigate if the algorithm optimized in spectral space achieves similar performance in accuracy as a standard form of NMF. This provides confidence that our novel algorithm performs similarly when compared to a reference algorithm. Next we modify the algorithm optimized in spectral space to achieve new functionality for color space – an extension that a standard NMF is currently not capable of providing. This straightforward modification achieves optimization in color space, greatly increasing the algorithm’s accuracy, and demonstrates that the modified variant functions properly. Finally, the algorithm is modified to implement both optimization in color and spectral spaces, demonstrating the success and utility of both implementations.

In the literature genetic algorithms have been utilized for enhancing NMF initialization [11–13]. For example, Clouch and Boukhetala address situations where an algorithm solves a semi-nonnegative matrix factorization for the case where the data matrix $X$ may have both positive and negative signs [11]. These previous results [11] differ from those of the present investigation in which the data matrix is nonnegative. Other results find that evolutionary algorithms for NMF initialization [12] are more computationally demanding than NMF based on nonnegative double singular–value–decomposition (NNDSVD) [14]. By comparison the present study adopts a new SVD-based initialization method that improves upon approaches that previously used NNDSVD [15]. In addition, one investigation previously combined a genetic algorithm and NMF to assess a Japanese Female Facial Expression image dataset, and showed their algorithm has advantages in terms of relative error, sparsity and orthogonality [13]. However, for the kinds of spectral datasets we use here, while some degree of orthogonality is attractive, strong sparsity is undesirable. This is because spectral datasets typically represent curves that are smooth and broadband. Also, earlier investigations [12, 13] that explored genetic algorithms for initialization of NMF algorithms used NMF algorithms which were strictly based on an updating rule. By comparison, the algorithms employed here use closed-form solutions for initialization and the form of NMF employed uses a genetic algorithm. As a result, the new approach presented here can implement optimization in color space, which is an advance when compared to the previous investigations, which were not designed for and, thus, not capable of such an implementation.

The paper is arranged as follows: Methodology is presented in Section 2, the algorithms are described in Section 3, experimental results are given in Section 4, and conclusions are given in Section 5.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{Spectral curves. a) CIE 1931 color matching functions of the Standard Observer. b) Normalized basis functions calculated using the standard NMF and optimized in a spectral space.}
\end{figure}

### 2. Methodology

Previous work shows the GA is known as a search technique for large spaces and is superior to other optimization methods when requirements for function features like smoothness, continuity and differentiability are violated [16]. In addition, the GA concept is easy to understand and implement, works in noisy environments and supports multiobjective optimization.

The GA contains a population of individuals and assesses the fitness of individuals for each generation. We consider a data matrix $X \in \mathbb{R}^{d \times n}$ and factors $W \in \mathbb{R}^{d \times r}$ and $Z \in \mathbb{R}^{r \times n}$, where $d$ is the number of wavelengths, $n$ is the number of spectral colors, and $r$ is the maximum number of basis functions or the rank of factorization. The number of basis functions can be less or equal to the rank of factorization. We use the GA to define the best subset of variables according to the fitness function:

$$\min_{W, Z} \sum_{i=1}^{n} \|x_i - Wz_i\|^2$$

subject to $W \geq 0; \ z_i \geq 0$.

where $x_i$ and $z_i$ are the columns of the matrices $X$ and $Z$, respectively, and $W \geq 0$ and $z_i \geq 0$ indicate that all entries of $W$ and $z_i$ are non-negative.

On the other hand, GA provides an effective way to select the best variables for color representation minimizing the color difference. The representation of individuals in this case is illustrated in Fig. 2. The individuals or “chromosomes” are represented by real valued vectors that live in a genotype space. Problem–specific solutions are located in the visual phenotype space as colors. The mapping between the spaces is made using encoding and decoding. The encoding is usually not used while decoding the real valued vectors (spectra), but helps to evaluate individuals according to their fitness function. From a spectral color viewpoint the genotype space corresponds to the physical spectral space, while the phenotype space relates to the physiological color space. Decoding is identical to the spectrum-to-color conversion typically used in spectral imaging [17]. Thus, the real–valued vectors/chromosomes are spectra and the problem–specific solutions achieve color
difference minimization. Let \( f() \) be a decoding function, i.e. a spectrum-to-color conversion function, then \( \|\Delta E_{ab}\|^2 = \|f(x_t) - f(W_z_t)\|^2 \), where \( \Delta E_{ab}^t \) is a CIELAB color difference:
\[
\Delta E_{ab}^t = \sqrt{(L_x^t - L_{Wz}^t)^2 + (a_x^t - a_{Wz}^t)^2 + (b_x^t - b_{Wz}^t)^2}.
\]

We denote \( [L_x^t, a_x^t, b_x^t]^T = f(x) \) and \( [L_{Wz}^t, a_{Wz}^t, b_{Wz}^t]^T = f(W_z) \). In this case the fitness function in color space is as follows:
\[
\text{minimize } W, Z \sum_{i=1}^n \|f(x_t) - f(W_z_t)\|^2 \tag{2}
\]
subject to \( W \geq 0; z_t \geq 0 \).

3. Algorithms

Three algorithms based on fitness functions are developed. The first algorithm is for basis functions optimized in a spectral space (GA-NMF-S). The second algorithm is for basis functions optimized in a color space (GA-NMF-C). The third algorithm uses multiobjective optimization for both the spectral and color spaces (GA-NMF-M). The pseudo-code for all these algorithms is as follows:

\[
t := 0;
\]
\[
\text{initialize } P(t); \quad \text{evaluate } P(t);
\]
\[
\text{while not terminate do}
\]
\[
P'(t) := \text{select-mates}(P(t));
\]
\[
P''(t) := \text{crossover}(P'(t), pc);
\]
\[
P'''(t) := \text{mutate}(P''(t), pm);
\]
\[
P''''(t) := \text{smooth}(P'''(t), ps);
\]
\[
\text{evaluate}(P''''(t));
\]
\[
P(t) := P''''(t);
\]
\[
t := t + 1;
\]
\[
\text{end}
\]

Algorithm: GA-NMF-\(i\) algorithm, \(i = S, C, M\).

Let \( P(t) \) denote a population consisting of individuals, and \( pc, pm \) and \( ps \) the probabilities for operations: “crossover”, “mutate”, and “smooth”, respectively. We use two kinds of populations to define the columns of \( W \), i.e. basis functions, and the rows of \( Z \). For both populations the sequence of procedures presented in the algorithm is the same and, apart from the operation “mutate”, all procedures are identical.

3.1 Initialization

The presented Algorithm is similar to general GA except for the initialization and the operation “smooth”. The initialization is important because it can affect algorithm performance. In our earlier study we used a random initialization of \( W \) with Gaussian functions [18]. These functions were utilized based on a priori knowledge that the resultant basis functions should be bell-shaped and to provide a diversity of population that is needed for evolution. For this purpose Gaussian distributions are more desirable than, for example, piecewise linear functions. However, random initialization of the GA, as well as the standard NMF, leads to variations in observed results. To avoid random initialization influences and to improve performance of the standard NMF a non-negative singular value decomposition (SVD) for initialization has been proposed [15]. We adopted this approach for GA-NMF and modified our operation “initialization”, and used singular value decomposition:
\[
X = U S V^T,
\]
where \( S \) is the diagonal matrix of singular values, \( S = \text{diag}(\sigma_1, \sigma_2, \ldots, \sigma_n) \), \( \sigma_1 > \sigma_2 > \ldots, \sigma_n \), the columns of matrices \( U \) and \( V \) are the left and right singular vectors of \( S \), respectively, and \( T \) is transpose.

Let us assume that the rank \( r \) of factorization is less than \( n, r < n \), and compute a fraction (\(< 90\%) of the total data variance to be retained. The retained variance in percents is as follows:
\[
\frac{\sum_{i=1}^r \sigma_i}{\sum_{i=1}^n \sigma_i} \times 100\% < 90\%.
\]

We then define matrices \( U_r = (u_1, u_2, \ldots, u_r) \) and \( S_r = \text{diag}(\sigma_1, \sigma_2, \ldots, \sigma_r) \) consisting of the first \( r \) vectors and the first \( r \) singular values of matrices \( U \) and \( S \), respectively.

Finally, we initialize matrices
\[
W_{\text{init}} = |U_r|, \quad Z_{\text{init}} = |S_r V^T|,
\]
where \(|A|\) denotes a matrix consisting of the absolute values of the entries of \( A \).

The rank \( r \) defines the maximum number of basis functions. In experiments we use the number \( r_b \) of basis functions, \( r_b \leq r \).

The initialization made for the standard NMF can be used to generate only one individual. Therefore, we modify this approach to initialize the whole population. We employ SVD initialization followed by NMF with an update rule called local non-negative matrix factorization (LNMF) [15]:
\[
Z \leftarrow \sqrt{Z} \ast (W^T(X \ast (WZ))).
\]
\[
W \leftarrow W \ast ((XZ^T) / (WZZ^T)),
\]
where \( \ast \) and \( ./ \) indicate element-wise multiplication and division, respectively.
We take each even solution of the first 100 iterations of the SVD-NMF algorithm to obtain the population size 50. To determine the population size we follow previous work suggesting that the optimal population size for problems coded as bitstrings is approximately the length of the string [19]. Though we use real valued strings we follow this rule. Hence, the population size, i.e. 50, is approximately defined by the number of wavelengths, i.e. 61.

The solution obtained at iteration 100 does not dominate earlier iterations. The cross-correlation coefficients for the solutions of the MCC dataset at iterations 20 and 100 are as follows: 0.96, 0.98 and 0.97 for short, middle and long wavelengths, respectively. This means that recombination (crossover and mutation) easily achieves a better solution than the best one obtained at initialization.

3.2 Selection, crossover and mutation

The operation “select–mates” makes the tournament selection of parents. In single objective optimization we randomly select the group of 15 individuals from the population. The group size is experimentally defined. Then, within this group the individual with the best fitness value is selected as a parent. For the selection of parents in multiobjective optimization two individuals are used. The selection procedure is given in Section 3.4.

The operation “crossover” is based on uniform crossover with probability \(pc = 0.9\).

The operation “mutate” is different for \(W\) and \(Z\). For individuals of \(W\), the gene value \(w_{ij}\) is randomly selected and changed using a random value \(a \sim \mathcal{N}(0, 1)\) as follows:

\[
w_{ij} = w_{ij} + 0.0005a.
\]

Similarly for \(z_{ij} = z_{ij} + 0.2a\). When the value \(w_{ij}\) or \(z_{ij}\) become less than zero it is set to zero. The selected value for probability of mutation is \(pm = 1/61\).

The designed algorithms have a deterministic initialization, but selection is still random. Therefore the results have small variations at each algorithm run which can be neglected. For example for three datasets used in this study and spectral optimization described in Section 4, the confidence interval of the average MSE is \([0.1386 \times 10^{-2}, 0.1414\times 10^{-2}]\) (MCC dataset), \([0.8377\times 10^{-2}, 0.8623\times 10^{-2}]\) (paint dataset) and \([0.1829, 0.1869]\) (Munsell dataset) at confidence level equal to 95%.

3.3 Smoothing

Crossover and mutation can produce abrupt changes in the values of a chromosome’s neighboring genes, leading to multi–peaked basis functions. To solve this problem we introduce a smoothing operation “smooth” for columns of \(W\) and for rows of \(Z\) with a simple averaging window comprising three elements. Testing shows that a good probability of smoothing is about 10 times less than the probability of crossover, i.e. \(ps = 0.1\).

3.4 Evaluation and fitness functions

For algorithms GA-NMF-S and GA-NMF-C, fitness functions are identical to the objective functions Eqs. (1) and (2), respectively.

In multiobjective optimization (GA-NMF-M) a Pareto ranking approach is utilized [20]. The objective value of each solution is not assigned in this case. The population is ranked according to a dominance rule in the objective space. The operation “evaluate” is replaced by three operations: “evaluateS”, “evaluateC” and “rank”:

\[
E_s(t) := \text{evaluateS}\left(P(t)\right);
E_c(t) := \text{evaluateC}\left(P(t)\right);
\text{rank}\left(E_s(t), E_c(t)\right);
\]

The operations “evaluateS” and “evaluateC” calculate the objective functions \(E_s(t)\) (Eq. (1)) and \(E_c(t)\) (Eq. (2)), respectively. The solutions are evaluated using the operation “rank” which calculates the fitness value based on the solution rank in population [20, 21].

First, the Pareto ranking technique calculates a set of nondominant fronts \(F_i\), where \(i = 1, 2, \ldots, v\) for the whole population, where lower ranks correspond to better solutions. Hence, \(F_1\) is a Pareto front. Second, we use the ranking technique for solutions belonging to the particular front. We adopt the crowding distance approach [20] and modify it introducing a more efficient hypervolume measure [22]. For three sequential solutions/points located in the front, the hypervolume measure for the central point is \(v = d_c d_s\), where \(d_c\) and \(d_s\) are distances between the two extreme points related to \(E_c\) and \(E_s\), respectively.

For the parent selection we randomly take two solutions. If solutions are from different nondominant fronts, the solution with the lowest rank is selected. Otherwise, the solution with the higher hypervolume measure wins.

3.5 Elitism

Here, “Elitism” relates to the problem of keeping good solutions during optimization. In single objective optimization we visually monitor the objective function value over time and select algorithm parameters achieving a good convergence rate. Therefore, we do not maintain elitist solutions in the single objective optimization case.

For multiobjective optimization we use the elitism strategy combining the old population and the offspring. Then, we apply deterministic selection keeping the best solutions to obtain the new population with the size of 50.

4. Experiments

The experiments were conducted with stimulus measures using optimization in two spaces: spectral and color spaces. The measured number of wavelength of spectra was taken at 5nm intervals in the humanly visible range (400-700nm). The number of basis functions tested was three.
Four light sources (D65, Halogen, LED and Xenon) were utilized in tests. Three datasets of measured reflectance spectra were used in experiments (Tab. 1). For the paint dataset we increased the number of wavelengths originally taken at 10nm from 31 to 61 using linear interpolation.

NMF can produce ambiguous solutions with respect to factors $W$ and $Z$. To avoid this problem, two normalizations of calculated basis functions were completed. That is, we first divided all elements in each column of $W$ by their sum [5], and subsequently normalized all basis functions to make their total maximum equal unity.

Table 1: Spectral datasets. The first dimension of “Size” equals number of wavelengths (#w) and the second one equals the number of samples (#s).

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Size=#w×#s</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCC</td>
<td>61×24</td>
</tr>
<tr>
<td>Munsell color dataset [23]</td>
<td>61×1269</td>
</tr>
<tr>
<td>Paint dataset [24]</td>
<td>61×91</td>
</tr>
</tbody>
</table>

4.1 Spectral space

Non-negative matrix factorization was implemented in the spectral space. The Algorithm used fitness function according to Eq. (1) and the initialization described in Section 3.

The parameters for GA were as follows. The individual length for columns of $W$ was 61 (number of wavelengths #w). The individual length for rows of $Z$ corresponded to the number of samples (#s) (Tab.1). The number of iterations was 200,000, although for the Munsell set it was 1,000,000. The population size was 50 for all tests.

We initially compare GA-NMF-S with a standard NMF algorithm which can be divided into several classes: Multiplicative update algorithms, gradient descent algorithms, and alternating least squares (ALS) algorithms. For these algorithms the statements about convergence (global or local) have not been proven, although it has been suggested that the ALS algorithm will converge to a local minimum in certain special cases [3]. The multiplicative update algorithms are the first well–known NMF algorithms which often converge in practice. They are, generally speaking, known to be a standard baseline against which any new algorithm is compared. Therefore, a multiplicative update algorithm was selected for comparison with GA. Note that convergence of GA has not been proven, and it is known that GA can be found to converge to a local minimum. However, we have confidence in the present solutions due to their wellformedness compared to known chromatic response functions of human observers, and because GA algorithms in general are widely used to obtain high-quality solutions for difficult and complex optimization problems.

For comparison we give the result for MCC using the standard NMF algorithm, 200,000 iterations (Fig. 1b) [4]. The result for GA-NMF-S is shown in Fig. 3a. The results are similar. In addition, the MSE values for each spectral color are given (Tab. 2). The GA-NMF results are better than those of the standard NMF.

Algorithms were implemented as Matlab routines installed on a 2.5GHz laptop computer running Windows operating system. The computational time for standard NMF obtained 101s versus 220s for the GA-NMF-S. Below we modify the GA-NMF algorithm to obtain a new functionality to work in color space which the standard NMF does not provide. The standard NMF is based on an update rule working in the physical spectral space and is not capable of working in color space. It will be shown that the optimal basis functions derived in color space approximate color appearance similarity significantly better.

Though our main purpose is to study three-band (primary or filter) systems we additionally test the potential of our method for a four-band system. We note that SVD initialization defines the rank of matrix factorization [15]. According to this approach the rank cannot be higher than four,
Moreover, the curves are less smooth. Similar results decayed more rapidly (Fig. 5a) compared to those of curves in the basis functions tails for the Munsell dataset derived for the same light source (D65 illuminant) for both datasets, we observed the basis functions tails for the Munsell dataset using the GA-NMF-C algorithm given 200,000 iterations and 1,000,000 iterations, respectively. Assuming the same light source (D65 illuminant) for both datasets, we observed the basis functions tails for the Munsell dataset decayed more rapidly (Fig. 5a) compared to those of curves in Fig. 3d. Moreover, the curves are less smooth. Similar results were obtained for the paint dataset. For GA-NMF-S (paint dataset) ΔE_{avg} = 15.05 and for GA-NMF-C ΔE_{avg} = 7.56. For GA-NMF-S (Munsell dataset) ΔE_{avg} = 12.53 and for GA-NMF-C ΔE_{avg} = 8.52.

4.3 Multiobjective optimization

We also investigated the advantage of GA in multiobjective optimization (GA-NMF-M). In this experiment we search for optimal basis functions for both spectra and colors using criterion based on the Pareto front described in Section 3. The results are presented in Fig. 5b, Tabs. 2, and 3. The results represent a trade-off between two marginal cases: spectral and color optima. The algorithm is less accurate than corresponding single objective algorithms in color space.

We suppose that the solution given by the GA-NMF-S algorithm is close to optimal. The approximation obtained from the multiobjective algorithm can be considered good because the ratio indicating fit, defined as the average MSE of the GA-NMF-S algorithm over the average MSE of multiobjective optimization, is 0.82, i.e. close to unity. In addition, the average MSE of multiobjective optimization is better than the average MSE of the standard NMF (Tab. 2). The best rank solution of the GA-NMF-M algorithm gives \( E_r = 71.85 \) and \( E_s = 0.042 \) at iteration 200,000 while \( E_r = 1.34 \times 10^4 \) and \( E_s = 17.7 \) at iteration 300 which are far from the optimum. Based on these findings we consider the multiobjective optimization to be efficient.

The above results give us confidence that the multiobjective algorithm can be effectively used in digital image archiving for the purposes of employing one set of basis functions, simplifying the task of deriving two sets representing spectral and color space solutions.

5. Conclusions

We developed new algorithms for NMF using a GA approach. One algorithm presented, GA-NMF-S, finds optimal basis functions for a spectral dataset in the spectral do-
main. That algorithm’s performance was shown to be similar to the standard NMF algorithm. A second algorithm was introduced, GA-NMF-C, that was shown to provide optimal basis functions for a color space. To the best of our knowledge this is a new result. The basis functions from GA-NMF-C were shown to be bell shaped and more accurately capture the mode locations of the spectral curves. In addition, the second algorithm accurately approximates colors. Finally, multiobjective optimization in both spectral and color spaces was implemented using the third algorithm modification GA-NMF-M. Investigations of the latter implementation found that the multiobjective algorithm can be used in digital image archiving for the purposes of employing one set of basis functions, and thereby simplifying the task of deriving two sets representing spectral and color space solutions.

An important feature of the present approach is that the procedures are both general and scalable, in the sense that they are not bound to the use of CIE 1931 color space, per se, or to the choice of particular R.G.B display primaries, or constrained by the 5nm measurement intervals present in the wavelength measurements in our spectral datasets. Thus, the methods described here can be generalized and make use of any color appearance spaces that has some form of color difference metric, and across a reasonable set of naturalistic illuminants. Moreover, as seen by the demonstration of 4 basis functions approximated using these techniques, the present approach can be used to evaluate and approximate color processing dimensions that differ from that commonly used when developing three-parameter device profiles as a model of the color perception of a standard normal human observer.

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References

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