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A Non-negative Matrix Factorization Framework for Combining Multiple Image Segmentations

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Abstract

Segmentation, or partitioning images into internally homogeneous regions, is an important first step in many Computer Vision tasks. In this paper, we attack the segmentation problem using an ensemble of low cost image segmentations. These segmentations are reconciled by applying recent techniques from the consensus clustering literature which exploit a Non-negative Matrix Factorization (NMF) framework. We describe extensions to these methods that scale them for large images and incorporate smoothness constraints. This framework allows us to uniformly combine segmentations from different algorithms or feature modalities, while avoiding significant parameter tuning for the specific image being segmented. We demonstrate that combining multiple “naive” image segmentations derived from k-means clustering compares favorably with more advanced Mean Shift and Efficient Graph Based Segmentation algorithms. The approach is evaluated on the Berkeley image segmentation dataset.

1. Introduction

Image segmentation, or partitioning an image into regions with internal segment coherence, has a long history in the Computer Vision literature and yet still has no generally accepted solution. The goal is to represent the image with fewer, more meaningful parts, which makes processing more tractable and relevant to subsequent vision tasks such as object localization and identification or content-based image retrieval.

In general, we want to assign pixels to segments based on their similarity to other members of that segment and dissimilarity to those of other segments. This implies a distance or homogeneity metric based on some set of cues which may include image distance, various region properties such as color and texture, or boundary/gradient information. Often these different feature modalities give us different results for the pixel and region similarity required to define segments, and considerable work has been devoted to combining them [1, 3, 12, 13].

Clustering techniques are a natural approach to computing image segmentations and a great variety of methods have been applied to the problem. Generic clustering techniques such as K-means often have difficulty integrating the spatial continuity or smoothness implied by image segmentation. These limitations are overcome by more expensive graph-based techniques such as spectral clustering [6] or normalized cuts [14, 15], which explicitly represent neighborhood linkages. Amongst the many segmentation algorithms proposed in the literature, few work well on natural images, and all are finely tuned to work on certain sets of images. For example, algorithms which work well segmenting biological images rarely work well on outdoor scenes.

Our work is motivated by the observation that even though a single segmentation algorithm or similarity metric by itself might produce some poor segments, there often exist sub-parts of the image which it explains well. So if diverse segmentations explain different parts of the image well, an ensemble of these could produce a superior consensus segmentation than any of the original segmentations. Our contribution in this paper is to provide a general framework for combining segmentations from heterogeneous sources of information, while guiding the fusion process using additional a-priori domain knowledge.

The problem of combining multiple segmentations can
be posed as a cluster ensemble problem. While classifier ensembles have been widely used in the Machine Learning and Data Mining communities, researchers have only recently started exploring cluster ensemble problems [16]. This is primarily because the cluster ensemble problem is inherently more difficult, since we no longer have well defined classes. From a linear algebra point of view clustering has been studied as a matrix factorization problem. Traditionally, SVD based methods have been used for this purpose. However, for image data (which is non-negative), the bases produced by these methods are not easily interpretable since they do not enforce non-negativity constraints. Non-Negative Matrix Factorizations (NMF) [8] produces a matrix factorization which respects non negative constraints, thereby producing directly interpretable and more representative bases. On a parallel front, recently Li et al. [9] have shown that consensus clustering (an algorithm for solving cluster ensembles) may be posed as a NMF problem. In this paper we propose the use of NMFs for finding the consensus segmentation. We also explore incorporating domain constraints in the consensus process to produce higher quality segmentation maps.

Cho and Meer [2] also proposed a consensus segmentation approach. The system is based on a bottom up Region Adjacency Graph (RAG) pyramid method which merges regions until a threshold on similarity is reached. The base segmentations are generated using a grayscale consistency metric, these are then used to compute a co-occurrence probability field for pixels grouped together in the segmentations. The probability field is in turn used as the metric to compute the final consensus segmentation again via the RAG pyramid. Our approach is related in that we search for an assignment of pixels to segments which best matches the mean co-occurrence \( \bar{M}_{ij} \) (see 2) for each pixel or object pair. We determine the final segmentation using NMF rather than a multiscale approach, which allows us to avoid setting thresholds on region similarity.

Zhang et al. [20] propose combining an ensemble of Spectral Clustering results computed using randomly generated scale parameters to construct a consensus segmentation of SAR images. The authors propose several approaches for combining segmentation maps including a majority voting scheme and a hypergraph-based metaclustering algorithm. They conclude that, of the voting and hypergraph techniques, the segmentation which maximizes sum of the normalized mutual information between the base segmentations and the consensus is the best solution. However, it has been shown in [9] that the NMF approach outperforms both the naive voting scheme and the more advanced hypergraph approach.

Our NMF framework provides a flexible general method for combining a set of maps as well as additional constraints such as smoothness or potentially boundary information. The base maps themselves can arise from any combination of segmentation algorithm and feature modality. Techniques such as ours, which use connectivity (co-occurrence) matrices between pixels, present a problem due to their size. We describe a method to scale the problem by essentially computing regions which have a preconsensus, spatially linked pixels which belong to one segment in all segmentation maps. These superpixels or objects allow us to compute consensus segmentations even for large images.

Finally, Sections 2 and 3 present details of the proposed approach and Section 4 presents the evaluation of our system on the Berkeley image segmentation database and comparison of our results to those for Mean Shift [4] and Efficient Graph-based Segmentation [5].

### 2. Consensus Segmentation Framework

The consensus segmentation problem seeks to reconcile \( T \) different segmentations (base segmentations) of a \( p \times q \) image. Equivalently, the consensus segmentation is a segmentation closest to all \( T \) segmentations. Let \( B = \{S^1, S^2, \ldots, S^T\} \) be the set of base segmentations. For each segmentation \( S^t \), we have \( K \) segments \( \{S^t_1, S^t_2, \ldots, S^t_K\} \), where \( K \) is not necessarily the same for each segmentation \( S^t \), and every pixel must belong to some segment \( S^t_k \) for each segmentation \( S^t \). By representing each segmentation \( S^t \) as a \( p \times q \) connectivity matrix, \( M \):

\[
M^t_{ij} = \begin{cases} 
1 & (i, j) \in S^t_k \\
0 & \text{Otherwise}
\end{cases}
\]

we can compute the distance (\( \Delta \)) between any two segmentations \( S^1 \) and \( S^2 \) as:

\[
\Delta(S^1, S^2) = \sum_{i=1}^{pq} \sum_{j=1}^{pq} \delta_{ij}(S^1, S^2)
\]

where \( \delta_{ij} \) is the pairwise pixel distance:

\[
\delta_{ij}(S^1, S^2) = \begin{cases} 
1 & (i,j) \in S^1_k \text{and}(i,j) \notin S^2_k \\
1 & (i,j) \in S^2_k \text{and}(i,j) \notin S^1_k \\
0 & \text{Otherwise}
\end{cases}
\]

or equivalently

\[
\delta_{ij}(S^1, S^2) = (M^t_{ij} - M^{S^2}_{ij})^2
\]

Now, the problem of finding the consensus segmentation can be formulated as the following optimization [7]

\[
\min_{S^*} \frac{1}{T} \sum_{t=1}^{T} \Delta(S^t, S^*) = \min_{S^*} \frac{1}{T} \sum_{t=1}^{T} \sum_{i,j=1}^{pq} [M^t_{ij} - M^{S^t}_{ij}]^2
\]
equivalently,
\[
\min_U \sum_{i,j=1}^{pq} (\tilde{M}_{ij} - U_{ij})^2
\]
(6)
where, \( \tilde{M} = \frac{1}{T} \sum_{t=1}^{T} M_{ij}(S^t) \) and we adopt \( U_{ij} = MS_{ij} \) as the solution of the optimization problem for notational simplicity.

Unfortunately, we also have constraints on \( U \) that need to be dealt with. Consider any three pixels \( i, j, \) and \( l \). Suppose \( U_{ij} = 1 \); that \( i \) and \( j \) belong to the same segment. If \( j \) and \( l \) belong to the same segment, then \( i \) and \( l \) must also belong to the same segment. However, if \( j \) and \( l \) do not belong to the same segment, then \( i \) and \( l \) cannot belong to the same segment. Now, consider the case where \( i \) and \( j \) belong to separate segments. We can now have \( i \) in the same segment as \( l, j \) in the same segment as \( k \), or none of them in the same segment. These constraints can be expressed as [9]:
\[
\begin{align*}
U_{ij} + U_{jl} - U_{il} & \leq 1 \quad (7) \\
U_{ij} - U_{jl} + U_{il} & \leq 1 \quad (8) \\
- U_{ij} + U_{jl} + U_{il} & \leq 1 \quad (9)
\end{align*}
\]
Note that the above constraints are indexed by individual pixels. Thus there are 3 constraints per pixel of the image. The constrained optimization problem turns out to be np-hard [10].

Following [9], we use an alternate specification of the above optimization problem using row stochastic(rows sum to 1) indicator matrices \( H = \{0, 1\}^{n \times k} \). It is easy to see that \( U = HH^T \). Our consensus segmentation problem now becomes:
\[
\min_H \| \tilde{M} - HH^T \|^2 \quad (10)
\]
with \( H \) restricted to the space of indicator matrices. However, since restricting \( H \) to be an indicator matrix is hard, we could reformulate the above problem as follows:
\[
\min_{H^T H = D, H \geq 0} \| \tilde{M} - H H^T \|^2 \quad (11)
\]
where \( D = \text{diag}(H^T H) \). By restricting \( H^T H \) to be diagonal we indirectly enforce the constraint that each row of \( H \) can have only one non zero element. However, this formulation involves a priori knowledge of \( D \) (cluster sizes) which is usually unavailable. As a result Eqn 11 is further reformulated as:
\[
\min_{H^T H = D, H \geq 0} \| \tilde{M} - \tilde{H} D \tilde{H}^T \|^2 \quad \text{s.t. } D \text{ Diagonal} \quad (12)
\]
where \( H H^T = \tilde{H} D \tilde{H}^T \). \( D \) is now obtained as a solution to the optimization problem. In practice, the constraint on \( D \) being diagonal is relaxed to \( D \) being any symmetric non-negative matrix, recasting the above problem as the familiar orthogonal nonnegative matrix tri-factorization problem, which is solved using iterative solution techniques.

**Scaling.**The algorithm as described above, however does not lend itself to be used practically for image segmentation. The major problem in adopting the above algorithm is the \( M \) matrix. For a \( p \times q \) image, the corresponding \( M \) matrix has \( pq \times pq \) entries. This quadratic growth in storage severely limits the size of images which can be processed by the algorithm. For instance, we found that, at best, we could work on 70 \( \times \) 70 images on a computer with 2GB of RAM. To alleviate the scaling problem we observe that the NMF problem may be interpreted as finding the closest connectivity matrix to the given arbitrary (not necessarily a connectivity matrix) \( \tilde{M} \) matrix. Now, if a certain set of pixels always occur in the same segment, establishing connectivity to any one of them is equivalent to explicitly establishing connectivity to all of them. Thus such sets can be collapsed into singleton entries in the \( \tilde{M} \) matrix. We refer to the entries in the \( \tilde{M} \) matrix as “objects”. Note that these objects cover a wide gamut of sizes from large sets to singleton pixels. The dimension of the matrix to be processed now is \( n \times n \), where \( n < pq \) is the number of objects\(^1\).

We find that employing such a scheme results in considerable savings both in terms of memory and computational cost, allowing us to comfortably processes \( 321 \times 481 \) images. Figure In displays the various objects (\( n = 1409 \)) of a \( 321 \times 481 \) image.

**Smoothness Constraints.**Smoothness is incorporated in the consensus framework through additional constraints using the Penalized Matrix Factorization [19] formulation. We now minimize the following augmented objective function:
\[
J = \min_{\tilde{H}^T H = D, H \geq 0} \| \tilde{M} - \tilde{H} D \tilde{H}^T \|^2_F + \text{tr}(\tilde{H}^T \Theta \tilde{H}) \quad (13)
\]
where \( \Theta \) is a \( n \times n \) matrix which encodes the smoothness constraints.

We compute the pairwise overlap \( d_{ij} \) between the entries of \( \tilde{M} \) (objects). We define overlap as the length(in pixels) of the shared boundary between two objects. The values \( \Theta_{ij} \) are computed as follows:
\[
\Theta_{ij} = \frac{1}{1 + e^{-\frac{d_{ij}}{1500}}} \quad (14)
\]
Thus, objects having a smaller overlap would have a larger \( \Theta_{ij} \) values\(^2\). The logistic function has the effect of normalizing the values in \( \Theta \) to lie in the \( \{0, 1\} \) range. Minimizing the augmented objective function has the effect of encouraging neighboring objects to have the same cluster label, thereby preferring smoother solutions to noisier ones.

\(^1\)In our experiments we found that for images with \( 321 \times 481 \) pixels, \( n \) never exceeded 2000 and was frequently \( \leq 1500 \)

\(^2\)A small value is added to \( d_{ij} \) when \( i \) and \( j \) do not overlap. In our experiments we found a value of 0.1 to be effective.
The optimization problem of equation 13 does not result in closed form multiplicative updates needed for solving NMF problems. Instead, following [19] we solve a relaxed version of equation 13 which only enforces the non-negativity of \( \hat{H} \). The necessary update equations are:

\[
D = (\hat{\hat{H}}^T \hat{H})^{-1} \hat{H}^T \hat{H} (\hat{\hat{H}}^T \hat{H})^{-1}
\]

(15)

\[
\hat{H}_{ij} = \hat{H}_{ij} \left[ \frac{(\hat{H} D \hat{H})^+_{ij} + (\hat{H} (D \hat{H}^T \hat{H})^{-1})_{ij}}{(\hat{H} D \hat{H})^{-1}_{ij} + (\hat{H} (D \hat{H}^T \hat{H})_{ij}^+) + (\Theta \hat{H})_{ij}} \right]
\]

(16)

where

\[
(\hat{H} D \hat{H}) = (\hat{H} D \hat{H})^+ - (\hat{H} D \hat{H})^{-1}
\]

(17)

\[
(\hat{H} D \hat{H}) = (\hat{H} D \hat{H})^+ - (\hat{H} D \hat{H})^{-1}
\]

(18)

The effect of the smoothness constraints on the resulting consensus segmentation is shown in Figures 1o and 1p.

3. Algorithmic Details

**Base segmentation diversity.** We employ k-means to generate the base segmentations of the ensemble. Diversity in the base segmentations is an important prerequisite for avoiding degenerate solutions. We incorporate diversity in our base segmentations in three ways:

Firstly, we inject diversity in the feature space by using four different color spaces:

1. Hue-Saturation: We use Hue and Saturation from the Hue-Saturation-Intensity color space. We leave out intensity in the hope of buying robustness to variation in lighting condition.
2. YIQ color space: This is the NTSC luminance (Y) and chrominance (I and Q) color space.
3. YCbCr color space: This is yet another luminance (Y) and chrominance (C_b and C_r) color space.
4. H1213 color space: The transformation is given by:

\[
I_1 = \frac{R + G + B}{3}; I_2 = \frac{R - B}{2}; I_3 = \frac{2G - R - B}{4}
\]

(19)

There is no particular reason for selecting the above four color spaces over any other set of color spaces. Any set of features which produce diverse segmentations may be used in their place. Secondly, the base segmentations are computed at three different scales. This is achieved by using different k values in the k-means clustering. In this work we used values of \{4, 6, 8\}. We, thus form an ensemble of 12 segmentations. Finally, we randomly initialize k-means and run it only for a modest number(100) of iterations. Terminating k-means when the algorithm might not have converged not only adds instability but is also computationally more efficient. Figure 1 shows the 12 base segmentation and the resulting final segmentation for an image from the Berkeley image segmentation database.

**Iterative Optimization.** Akin to other NMF optimization algorithms an iterative scheme is used in this paper. The \( \hat{H} \) matrix is initialized by clustering the matrix using \( k \)-means. Next the initial estimate of \( D \) is computed using 15 while fixing \( \hat{H} \) to it’s initial value. This is followed by \( \hat{H} \) computation using 16 while keeping \( D \) fixed at it’s previously computed value. This process of alternating minimization is repeated till the change in \( J \) falls below a certain threshold \( \epsilon \). We set \( \epsilon = 10^{-3} \) in our experiments.

4. Experiments and Results

We compare consensus segmentation against two other widely used segmentation algorithms, the efficient graph based segmentation (GBIS) algorithm [5] and the mean shift algorithm [4]. GBIS treats the image as a graph. Segmentation is achieved by splitting the graph into a collection of connected components. Two connected components are merged when the weight of the edge connecting the two components is less than the maximum weight in either components’ minimum spanning tree, plus some constant user controlled parameter \( M \). The publicly available implementation [5] of this algorithm has two other tunable parameters, \( \sigma \) a smoothing parameter and \( \min \) the minimum number of pixels in segment. In this paper we fix \( \sigma = 0.8 \) and vary \( M \) through \{100,200,300,400,500\} and \( \min \) through \{20,50,100,300,500\}.

The Mean Shift (MS) algorithm involves a mean shift filtering of the image data followed by a clustering of the filtered data. The mean shift filtering is a search for modes of the underlying pdf of the image data. In this paper we have used the open source EDISON [4] implementation of the mean shift segmentation. The EDISON system converts the original RGB image into the LUV space. The mean shift filtering is carried out in a 5 dimensional feature space, containing the \((x,y)\) image coordinates and the LUV values. The algorithm has three tunable parameters spatial bandwidth \( (h_s) \), color bandwidth \( (h_c) \) and \( \min \). We, following popular trend [18] set \( h_s = 7 \) and vary \( h_c \) through \{3, 5, 7, 9, 11, 13, 15\} and the range of \( \min \) is chosen to be the same as GBIS.

Finally, our consensus algorithm has two tunable parameters. We apply a Gaussian filter on our images, and the standard deviation of the filter \( \sigma \) is the first parameter, while the number of color clusters \( C \) present in an image is the second parameter. \( \sigma \) is varied through \{0.5, 0.75, 1.0, 1.25\}, while \( k \) varies from image to image. In our experiments \( C \) took an integer value between 3 and 9 depending on the image.

For quantifying the performance of the segmentation algorithms, we use the Probabilistic Rand Index (PRI) proposed in [17]. We compare an image segmentation
Figure 1: An image from the Berkeley database, base segmentations and the resulting Consensus Segmentations (CS)

$S_{test}$ with a set of “ground truth” human segmentations \( \{H^1, H^2, \ldots, H^M\} \). The human segmentations are obtained from the Berkeley Image Segmentation Dataset [11], which contains a test set of 100 images.

For completeness we briefly describe the computation of PRI. A segmentation is considered “good” if it agrees with the human segmentations provided. The PRI score increases if the labels \( l_i \) and \( l_j \) of two pixels \( i \) and \( j \) are the same, i.e. if they are classified in the same segment of \( S_{test} \), and they are also classified in the same segment for a human segmentation \( H^h \). The score is hurt if this is not the case.

Formally, PRI is computed as:

\[
\text{PR}(S_{test}, \{H^1, \ldots, H^M\}) = \frac{1}{(N^2)} \sum_{i<j} [\mathcal{I}(l_i^{S_{test}} = l_j^{S_{test}}) p_{ij}
+ \mathcal{I}(l_i^{S_{test}} \neq l_j^{S_{test}})(1 - p_{ij})]
\]

(20)

Where, \( \mathcal{I} \) is the identity function and

\[
p_{ij} = \frac{1}{(N^2)} \sum_{i<j} [\mathcal{I}(l_i^h = l_j^h)]
\]

(21)

PRI takes values in the range \([0, 1]\), with a value of 0 resulting when \( S_{test} \) and \( \{H^1, \ldots, H^M\} \) having no similarities, and a value of 1 when \( S_{test} \) matches \( \{H^1, \ldots, H^M\} \) exactly.
<table>
<thead>
<tr>
<th>Segmentation</th>
<th>Mean PRI</th>
<th>Median PRI</th>
<th>St. Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kmeans</td>
<td>0.7070</td>
<td>0.7246</td>
<td>0.11590</td>
</tr>
<tr>
<td>MS</td>
<td>0.7627</td>
<td>0.8044</td>
<td>0.14135</td>
</tr>
<tr>
<td>GBIS</td>
<td>0.7759</td>
<td>0.8024</td>
<td>0.13246</td>
</tr>
<tr>
<td>Consensus</td>
<td>0.7602</td>
<td>0.8016</td>
<td>0.14242</td>
</tr>
</tbody>
</table>

Table 1: Comparison on the Berkeley test images.

The PRI values obtained for the three segmentation algorithms as well as the best $k$-means base segmentation are presented in Table 1. Figure 2 displays segmentations of a representative subset of the Berkeley test images, for visual comparison. The consensus algorithm significantly outperforms the best base segmentation, thus providing a quantitative measure of utility of the consensus process. Consensus also performs comparably with both MS and GBIS. A t-test at 95% confidence interval reveals that there is no statistical difference between the performance of the consensus algorithm and that of MS, while GBIS performs marginally better. GBIS scores significantly higher on images containing a high amount of detail. While consensus is prone to placing spurious boundaries in regions of high color variability, GBIS is able to identify such regions and incorporates adaptive thresholding to avoid placing spurious boundaries. Figures 3a, 3b and 3c illustrate this point. Even though we achieve a more semantic segmentation of the objects of interest our score suffers due to the additional details captured as compared to the human “ground truth” segmentations. Incorporating textural constraints in $\Theta$ or incorporating more expensive textural segmentations in the ensemble would help alleviate this problem.

5. Conclusion and Future Work

This paper presents a penalized non-negative matrix factorization framework for combining multiple image segmentations. A method to scale the framework for large images is also presented. We find that the performance of an ensemble of rudimentary $k$-means segmentations is able to perform competitively against popular state of the art segmentation algorithms.

In the future, we plan to explore alternate means of scaling the matrix factorization problem, such as blocking large connectivity matrices. We also plan to further explore the space of possible constraints which can be successfully incorporated in our framework to provide superior quality segmentations. We are especially interested in textural constraints, which would let us better deal with the problem of spurious boundaries.

6. Acknowledgements

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References


Figure 2: Images from the Berkeley test set and their segmentations
Figure 3: Segmentations and their PRI scores. The PRI scores are on the lower right corner of the segmentations.