Automatic Pill Identification from Pillbox Images

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Abstract: There is a vital need for fast and accurate recognition of medicinal tablets and capsules. Efforts to date have centered on automatic segmentation, color and shape identification. Our system combines these with pre-processing before imprint recognition. Using the National Library of Medicine Pillbox database, regression analysis applied to automatic color and shape recognition allows for successful pill identification. Measured errors for the subtasks of segmentation and color recognition for this database are 1.9% and 2.2%, respectively. Imprint recognition with optical character recognition (OCR) is key to exact pill ID, but remains a challenging problem, therefore overall recognition accuracy is not yet known.

1 INTRODUCTION

Adverse reactions to both legally prescribed medications and illicit or abused pills are a present and growing problem (Moore et al., 2007). When patients are brought to medical facilities in a stupor or coma with unidentified pills, rapid pill identification can be lifesaving. Adverse reactions involving anti-hyperglycemic medications, anticoagulants, and narcotics are potentially life-threatening, and all require their own particular care paths. Accordingly, automatic pill identification in emergency rooms and intensive care units could lead to better outcomes for these patients. Additionally, automatic pill identification would give police officers an efficient alternative to the current tedious method of entering each pill’s features into a database search and reduces user input errors on pills with many characters. Furthermore, the manual method, although it has the accuracy of human characterized imprint, color, and shape, fails when healthcare workers and police officers find themselves in locations with no internet access. Thus, there is a widespread need for automatic pill identification.

Recently, large commercial and government pill image databases have become available. These databases allow development and testing of pill identification programs. Among the very few works to appear in the literature, Lee et al. (2012) reported an identification accuracy of approximately 74%. Additionally, Hartl (2010) used the Studierstube ES framework for a mobile phone that focuses on speed. The accuracy and robustness of pill ID systems must be improved before pill identification systems can be utilized in the fields of healthcare and law enforcement.

This report details a pilot system that uses novel segmentation, shape recognition, color, and optical character recognition methods—all applied to pill recognition. In this paper, our model system is the Pillbox database (U.S. National Library of Medicine, 2012). The remainder of this article is organized in the following sections: 2) Automatic segmentation of pills from background, 3) Color identification, 4) Pill shape recognition, 5) Preliminary optical character recognition of imprint, 6) Results, and 7) Conclusions.

2SEGMENTATION OF PILLS

The initial task in pill recognition is segmentation, i.e. separating Pillbox images into distinct pill and background regions (see Figure 1). This involves four steps: 1) Conversion of the captured pill’s image RGB color space into L*a*b* color space, 2) 2D histogram generation along the L*-a* planes, 3)
Clustering via K-means++, and 4) Binary mask generation.

Figure 1: Original pill image (left); Final binary image border mask using our segmentation algorithm (right).

2.1 RGB Image Conversion to L*a*b*

Space

In the RGB color space, the color temperature of light and the demosaicing method affects the perceived color. Therefore, conversion from RGB to L*a*b* color space must be performed to reduce these interferences; this process is achieved through an intermediate color space, XYZ, described below.

The pill color in the database image and that of the captured image are expected to be different in practice, since the camera or the illumination conditions for the two images are different, creating varying colors with no common reference point (Szeliski, 2011). The XYZ color space was created to model the response curve of the human eye, to be used as a common point of reference. By transforming a color represented by an RGB value to an XYZ value, two colors can be compared more easily because of this common reference. L*a*b* is a non-linear re-mapping of the XYZ color space “where differences in luminance or chrominance are more perceptually uniform” (Szeliski, 2011).

Figure 2: Log of the original histogram. This image is the L*-a* histogram that has been modified by taking the log(1+histogramValue) for viewing. This is done to prevent the histogram image from being saturated when any particular histogram bin accumulates above 255.

2.2 2D L*-a* Histogram Generation

After converting to L*a*b* space, a two-dimensional histogram using the L* and a* planes of the image is generated. (Figure 2) To generate the 2D histogram, 256 linearly spaced bins ranging from 0 to 255 for L*, and -127 to 128 for a*, were used along each axis. This results in a histogram that illustrates the number of pixels that have a particular (L*,a*) combination.

2.3 K-Means++ Clustering

Input images are assumed to contain a single pill on a homogenous background; as a result, two clusters in the L*-a* histogram are expected. Therefore, a partitioning technique to minimize the total of Euclidean distances with two defined cluster centroids is applied, known as K-means++ clustering. (Figure 3) A variation of the K-means++ algorithm is defined and then described below.

During initialization, Arthur’s “D 2 weighting” method is employed (Step A) as a more intelligent starting point of the two cluster centroids, instead of random center initializations (Arthur and Vassilvitskii, 2007). The algorithm then iterates through each point on the histogram and assigns it to the nearest cluster centroid (Step B) (Xu and Wunsch, 2005). Once each point is assigned, the centroids are recomputed based on newly assigned points (Step C). The point allocation and centroid recalculation of Steps B and C repeat until some termination condition is met (Step D). The clustering terminates when either 20 centroid recalculations have occurred or the centroids move less than 0.01 spatial units.

Steps A-D:
A. Initialize clusters C₁ and C₂, with centroids κ₁ and κ₂ respectively, based on K-means++ algorithm
i. Choose an initial centroid, κ₁, uniformly at random from data set, \( \chi \in \mathbb{R}^2 \)
ii. Let \( D(x) \) indicate the smallest distance from data point \( x \) to the closest chosen centroid, \( \kappa_1 \).
Choose the second centroid, \( \kappa_2 \), by selecting \( c_2 = x' \in \chi \) with probability \( \frac{D(x')^2}{\sum_{x \in \chi} D(x)^2} \)
B. Assign each point in the data set to the nearest cluster centroid, i.e.
for \( j = 1, ..., N \):
\( x_i \in \kappa_1, \text{if } ||x_j - \kappa_1|| < ||x_j - \kappa_2|| \)
\( x_i \in \kappa_2, \text{else} \)
C. Recalculate the cluster centroids, \( \kappa_1 \) and \( \kappa_2 \), based on the current point assignments
D. Repeat Steps B, C until one of two termination conditions is reached:
Because the K-means++ algorithm is sensitive to initialization, the entire clustering process is typically run multiple times. The “best” clustering result can then be chosen based on a compactness score (Equation 1), which is the total sum of squared error (SSE) for every point to its centroid. Real-time application is of importance here, so the K-means++ clustering is limited to three attempts. The clustering attempt with the lowest corresponding compactness is selected to generate the binary mask.

The compactness score is shown by Equation 1 (Itseez, 2012).

\[
\sum_{i=1}^{N} \| x_i - \mu_i \|^2 \quad \text{where } i \in \{1, 2\}
\]  

(1)

Figure 3: Segmented histogram. This image shows how the histogram has been segmented into two clusters using the K-means++ algorithm. The red points are the centroids of each cluster.

2.4 Binary Mask Generation

Once the best clustering result is chosen, then the binary pill mask is generated. Previously, each pixel was assigned a label corresponding to its cluster. A blank binary image of identical size as the original is first created. The “background” cluster is then determined by finding the cluster that has the most member pixels contacting the image edge. This cluster’s pixels are assigned a value of 0 on the binary mask, while members of the second cluster, which theoretically correspond to the pill, are assigned a value of 1. The result is shown in Figure 1. Note that this assumes that the image fully captures the pill. The binary pill mask is then used in further pill characterization steps.

3 COLOR RECOGNITION

Seven hundred forty-four images were gathered from the National Library of Medicine’s Pillbox website (http://pillbox.nlm.nih.gov) with both front and back views. These high-quality images were used as the basis to develop a method to recognize color. The idea of histogram vector multiplication as a method for object recognition (Gonzalez and Woods, 2008) led to investigation of a similar approach for color histograms. Initially, histograms previously used for pill segmentation based on XYZ and L*a*b* color spaces resulted in an accuracy of 86.3% using logistic regression. Using the HSV color space which, like L*a*b*, represents luma and chroma separately, along with the captured pill’s image RGB values, resulted in an increase in color recognition to 98.1%.

For each channel of the HSV color space, along with red, yellow, and blue chromaticities (See Equations 2-4, Table 1, and Figures 4 and 5), histograms were calculated for every pill. All histograms consisted of 80 bins ranging from 0-360 for Hue, 0-1 for Saturation, and 0-255 for Value and scaled chromaticity.

\[
C_R = \frac{R}{R + G + B}
\]

(2)

\[
C_B = \frac{B}{R + G + B}
\]

(3)

\[
C_V = \frac{R + G}{2(R + G + B)}
\]

(4)

Table 1: Histogram Channels.

<table>
<thead>
<tr>
<th>Orig.</th>
<th>C_R</th>
<th>C_B</th>
<th>C_V</th>
<th>H</th>
<th>S</th>
<th>V</th>
</tr>
</thead>
</table>

Figure 4: Blue, red, yellow chromaticity histograms.

To reduce the effects of pill imprints, a histogram filter was developed to remove small peaks.

From the creation of six histograms for each pill, the pills were grouped according to each of the defined pill colors (white, yellow, orange, pink,
Determination of the color of a pill starts with using the pill segmentation mask to calculate the normalized histogram for the six channels. Each of the pill’s six histograms is then vector multiplied with its respective template histogram for each of the eleven predefined pill colors. For each color model, six scalar values resulting from the corresponding vector calculation are used as inputs into a logistic regression model.

Logistic regression, or the logit model, is a statistical analysis method by which the probability of an event occurrence is calculated based on predictor variables fitted to a logistic function. The logistic function is defined as:

$$f(z) = \frac{1}{1 + e^{-z}}$$

$$f_i(z_i) \in (0, 1)$$

Each of the $\beta_{n,i}$ coefficients is determined using maximum likelihood estimation and represents the weight of the predictor variable, $x_n$. $f(z)$ represents the probability of the outcome of any item and $z$ represents the measure of the total contribution of all independent variables in the model (Menard, 2001).

Here, $\beta_{0,i}$ is defined as the intercept for the $i^{th}$ defined color model, and $\beta_{n,i}$ as the regression coefficients of $i^{th}$ color model. The $\beta$’s were previously determined using a training set to create the logistic regression model for each of the defined colors. The variables $x_{n,i}$ are the histogram scalar values, which were previously described, for the $i^{th}$ color. Once the $f_i$ are calculated for each of the 11 defined colors, recognition of a pill color corresponded to the color yielding the maximum $f_i$.

In the case of a capsule with two colors, the same technique is used, with each half of the pill processed individually. First, the minimum bounding rectangle of the pill’s segmentation mask is calculated. Next, the pill mask is cut in half along the major axis. The two masks are used to process the pill as previously discussed.

4 SHAPE RECOGNITION

Classifying shapes was found to be most promising when done using Hu invariant moments (Hu, 1962). There are seven Hu moments, each independent of rotation and scale. Using binomial regression methods similar to those in Section 3, shapes were matched using Hu moments.

Difficulties in shape recognition were similar to those encountered in color recognition, namely that shapes are not always clearly defined, as is the case for numerous capsules and tablets. Our solution was to create an addition shape label that grouped together those similarly shaped pill to train our model to use this new label as a parameter to distinguish shapes.

5 IMPRINT RECOGNITION

The process of extracting imprint information from a pill is one which requires the consideration of several factors, including the luminance relationship between inscription and the pill. Once that information is known, two of four morphological operators are applied to the image before using an Optical Character Recognition (OCR) engine, Tesseract (Smith, 2012). Inaccuracies such as misplaced characters may sometimes occur. To compensate for this, a basic string matching algorithm is applied to the OCR output.

5.1 Imprint Extraction

Imprint extraction begins by determining pill color and luminance characteristics. First, if the capsule has two halves of different colors, the halves are processed separately. The half-capsule shape is automatically identified and separated into two pictures. Each picture includes each half capsule and therefore consists of two colors: one for the pill and one for the homogenous background, allowing the pictures to be processed in parallel.

Once the area of interest is acquired, the color image is converted into a gray-scale image. Luminance information allows determination of whether the capsule is darker than the text or vice versa. The appropriate gray-scale morphological operators are then applied.
Depending on the relative luminance of the pill and text, two of four morphological operators are applied to the image (Equations 6-11). The Black Hat operator locates areas of an image that are darker than their surroundings (Figure 6a), whereas the Top Hat operator locates areas that are lighter than their surroundings (Figure 6b) (Gonzalez and Woods, 2008). Applying either dilation or erosion to the image before the Black Hat/Top Hat operator often improves OCR results. For dark text on a light pill, dilation is used; for light text on a dark pill, erosion is used. Previous work has shown that the number of dilation or erosion iterations affects the results on a pill-by-pill basis. After the appropriate operators are applied to a given image, it is passed to the OCR, where text extraction is attempted.

Top Hat
\[ t_p = I - (I \ast h) \]  
(6)

Black Hat
\[ b_h = (I \ast b) - I \]  
(7)

Closing
\[ A \cdot B = (A \oplus B) \ominus B \]  
(8)

Opening
\[ A \ast B = (A \ominus B) \oplus B \]  
(9)

Dilation
\[ A \oplus B = \{z \in (B) \cap A \neq \emptyset\} \]  
(10)

Erosion
\[ A \ominus B = \{z \in (B) \cap A \neq \emptyset\} \]  
(11)

Figure 6a: Pill requiring Black Hat operation.  
Figure 6b: Results of Black Hat operation.

### 5.2 Integration of Tesseract

Once a given image has been processed to emphasize the text on the pill, it is passed to an open-source character recognition engine known as Tesseract (Smith, 2012). Tesseract analyzes the image and returns the identified characters. Typically, the OCR output will contain some mistakes, as 100% accuracy is uncommon. A solution to this problem is approximate string matching with limited vocabulary. Since only pills are considered in this project, the possibilities of text are limited to the imprints that are found on pills. As a result, it is possible to construct a dictionary that includes only the possible text outputs. By using dynamic programming to implement the edit distance match method (Apostolico & Galil, 1997), each OCR result can be compared to the dictionary entries. This method indicates which word in the dictionary is most similar to a given OCR output; essentially, it takes what may only be a partial match and finds the dictionary word that it most closely resembles. The next step in the previous example involves passing the image in Figure 6b to Tesseract. Due to the curvature of the pill, part of the text is missing and, as expected, the OCR returns only the partial match, “yVATSOi 3159.” However, by performing string matching with limited vocabulary, the correct imprint, “WATSON 3159,” is obtained.

### 6 RESULTS

The segmentation section of this algorithm was evaluated by comparing it to a set of ideal segmentation masks for 50 Pillbox images. By applying a threshold across the RGB planes and mathematically intersecting the results, an ideal pill mask was obtained for each pill image. Note that the “ideal” mask can easily be found in the Pillbox images because the background is uniformly black. A percentage error for the K-Means++ segmented images was found by locating all pixel locations where the two masks differed and dividing that value by the total non-zero pixels in the ideal mask. The algorithm’s average error was 4.05%, with a median error of 2.2% in the image set.

In reference to color, our methods and images from the Pillbox database achieved a high level of accuracy based on using multiple color spaces and classification using logistic regression. In the situation that a color was asserted only when z was greater than zero, probit regression accurately identified 95.8% of pills and logistic regression’s accuracy was as high as 96.8%. When z was less than zero, probit regression identified 96.6% correct while logistic regression showed 98% accuracy.

With regard to pill shape, when z was greater than zero, probit and logistic regression showed accuracies of 64.9% and 88.5%, respectively; when less than zero, probit and logistic regression showed accuracies of 65.5% and 90.9%, respectively.

In terms of type of pill, capsule was matched with 98.9% accuracy and tablet at 99.6%. A negative factor contributing to this yield is that the “tablet” and “oval” shapes sometimes overlap.

Considering imprint, the raw OCR output often contains words with several inaccuracies; ideally, these mistakes would simply be fixed by finding the dictionary word with the minimum number of edits between an OCR word and a dictionary word. If there are multiple dictionary words that yield the same number of minimum edits, the final output string will be incorrect. To quantify the results of the imprint extraction, the edit distance match method
was used to count the number of edits necessary for the final output string to be transformed into a 100% match. An average of 2.48 edits per sample was found to be necessary to have optimal results.

7 CONCLUSIONS

This report outlines a pill identification system that achieves a higher degree of automatic identification than previously reported. Further improvement is needed prior to practical application.

7.1 More Image Testing

Images in other databases, especially those taken in the field, have variable lighting and focus. It is likely that our successful segmentation accuracy, with a median error of 2.2%, will fail when algorithms are applied to other images. Other algorithms and additional color space dimensions such as the b* dimension in L*a*b* color space will be attempted.

7.2 Color Recognition Improvements

Color recognition accuracy measured by logistic regression, with a current error of 1.9%, is expected to fail with other images. Future steps to improve color recognition are more image blurring, RGB histogram normalization before processing, and adding L*a*b* to the current list of channels. We will explore other adaptive methods to ensure that data is not lost in the averaging method.

7.3 Shape Recognition Improvements

A secondary problem concerns shape recognition. Of twelve shape types, the three most common are prevalent enough such that uncommon shapes, e.g. teardrop or pentagonal shapes, are under-selected. A special function for these uncommonly shaped pills is needed.

7.4 Imprint Recognition Improvements

The algorithm for imprint extraction that has been outlined suggests a two-part system. First, the image should be processed in order to improve raw OCR results. Secondly, the OCR output string should be analyzed to limit the final output to a finite vocabulary. Preliminary efforts have been inconclusive. The optimal number of mathematical morphology operations, such as repeated dilation or erosion to produce the best results for a given image, is not known. This currently relies largely on human input. The techniques in string matching could also be improved in returning only the relevant information and excluding words of little value.

7.5 Improvements for a Practical System

Multiple challenges must be met to complete a working system. Fusion of the information from shape, color, and character determination will be needed. The images in the Pillbox database are of higher quality than can be obtained with a smartphone under real-life conditions. Overcoming non-ideal lighting, irregular positioning, and limited resolution are additional challenges that must be met before a practical system is available for health and law enforcement.

REFERENCES


algorithms. *IEEE Transactions on Neural Networks*, 16(3), 645-78.