Compact color texture descriptor based on rank transform and product ordering in the RGB color space
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Compact color texture descriptor based on rank transform and product ordering in the RGB color space

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Abstract

Color information is generally considered useful for texture analysis. However, an important category of highly effective texture descriptors – namely rank features – has no obvious extension to color spaces, on which no canonical order is defined. In this work, we explore the use of partial orders in conjunction with rank features. We introduce the rank transform based on product ordering, that generalizes the classic rank transform to RGB space by a combined tally of dominated and non-comparable pixels. Experimental results on nine heterogeneous standard databases confirm that our approach outperforms the standard rank transform and its extension to lexicographic and bit mixing total orders, as well as to the preorders based on the Euclidean distance to a reference color. The low computational complexity and compact codebook size of the transform make it suitable for multi-scale approaches.

1. Introduction

Color texture analysis is routinely used in many computer vision applications such as image-based medical diagnosis, land cover classification from satellite imagery, automated quality inspection, grading of agricultural and food products, etc. It has been shown by many authors that taking into account color information generally improves the performance – at least under steady illumination conditions – see for instance [30] and [6]. In the last two decades rank features have emerged as one of the most robust tools for image analysis. Examples of this approach are the well-known local binary patterns and their variants [7, 29], rank transform [38], texture spectrum [18], ranklets [35], directional rank coding [34] and sudoku rank features [17], all of which rely on a “greater than” relation between pairs of gray level values. The extension of such methods to color images has so far been limited by the absence of a natural ordering in the color spaces, where statements like pink is greater than brown don’t make any sense.

In this work we present a novel approach to color texture analysis based on rank features in color space. The key idea
is to replace the order relation over intensity values by a suitable partial order in color space. By doing so we admit that some pairs of colors are not comparable, as will become apparent later. We present an implementation of this idea by extending a classic method, the rank transform (RT), to work with a partial ordering (the product order) in the RGB space. The result is a compact yet highly effective local descriptor for color images.

In the remainder of the paper, after providing some background on partial orders (Sec. 2), we briefly review how rank-based descriptors have been defined in the literature for both grayscale and color texture analysis (Sec. 3). We present the proposed approach in Sec. 4 and validate the method through a set of classification experiments over nine datasets of color texture images (Sec. 5). The results clearly show that our method is more effective at discriminating texture than the original rank transform, either based on gray levels, or on different orders or preorders on the RGB color space. Section 6 concludes the paper and discusses directions for future work.

2. Theoretical background

We recall in this section some basic notions about order relations and briefly review the existing approaches for the ordering of multi-dimensional data.

2.1. Order relations

A total order $\leq$ is a binary relation on a set $X$ for which the following four properties hold for any three elements $a$, $b$ and $c$ of $X$:

- Antisymmetry: $a \leq b \land b \leq a \implies a = c$
- Transitivity: $a \leq b \land b \leq c \implies a \leq c$
- Totality: $a \leq b \lor b \leq a$
- Reflexivity: $a \leq a$

Totality establishes that any two elements of $X$ are comparable; it also implies reflexivity, as can be seen by setting $a = b$. By discarding totality for the weaker requirement of reflexivity we obtain a partial order. This differs from a total order in that there will be elements of $X$ that are not comparable to each other.

Finally, a preorder or quasiorder is a binary relation that is both reflexive and transitive (but not necessarily total or antisymmetric). The lack of antisymmetry means there can be cases where both $a$ weakly dominates $b$ and $b$ weakly dominates $a$ without $a$ and $b$ being equal; see Sec. 3.2 for an example. Total orders and partial orders are special cases of a preorder. Tab. 1 summarizes the definitions just introduced.

<table>
<thead>
<tr>
<th></th>
<th>Preorder</th>
<th>Partial order</th>
<th>Total order</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reflexivity</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Transitivity</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Antisymmetry</td>
<td>–</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Totality</td>
<td>–</td>
<td>–</td>
<td>✓</td>
</tr>
</tbody>
</table>

2.2. Ordering multi-dimensional data

The application of rank features to color or multispectral images is anything but straightforward for multidimensional data lack a natural way of ordering. Research has so far focused on restricted ordering principles or sub-orderings. These can be broadly divided into the four categories summarized below [2].

Marginal ordering (M-ordering) Multivariate ordering consists of sorting the data separately by one or more components. Lexicographical ordering is an example of M-ordering: for instance in the RGB space one could sort the color data first by R, then by G and finally by B (or following any other sequence obtained by a permutation of the channels). The major shortcoming of this approach, however, is that colors close to each other in the color space can end up very far apart after the ordering.

Reduced (aggregate) ordering (R-ordering) In this case multivariate data are preliminarily converted to univariate through some ad hoc functions. One could for instance establish a reference point (pivot) in the multivariate space and sort the data by their distance to it.

Conditional (sequential) ordering (C-ordering) In C-ordering the members of a random multivariate sample are sorted according to the corresponding values of another (usually univariate) random sample. An example of C-ordering is provided by statistically equivalent blocks, which consists of splitting a multidimensional space into subspaces or blocks through the use of one-dimensional cutting functions [32].

Partial ordering (P-ordering) Partial ordering differs from total ordering in that the totality condition is not required: in practice we accept that there will be pairs of elements incomparable to each other. We have already introduced partial ordering in Sec. 2.1; we shall discuss the partial ordering used in this work (i.e. product order) in Sec. 4.2.

Interestingly, most approaches to deal with order in color space belong to the first two categories. For instance, rank-
ing based on the gray level can be either seen as a marginal ordering in the HSV space (along the V axis) or as a reduced or aggregate ordering in the RGB space, where the aggregating function is the standard color to grayscale conversion.

3. Rank-based texture features

Rank features are defined in terms of the relative order (rank) among pixels, with the actual pixel values being discarded. By definition rank features are invariant to any monotonic change in the illumination intensity, a feature that makes them particularly suited for dealing with noise and sensor non-linearities [20].

Though most of the rank-based descriptors proposed in the literature are intended for grayscale image analysis, a number of methods for color images have also been described. We briefly review the two classes in the following subsections.

3.1. Grayscale analysis

Local image descriptors based on pixel intensity can either rely on a full ranking or thresholding. In the first case neighboring pixels are ranked according to their grayscale value and features are extracted from the rank signature. Examples of this approach are: rank transform [38], complete rank transform [15, 10], ranklets [35], n-tuples [19], directional rank coding [34] and local intensity order pattern [37]. In the second case features are computed from pairwise comparisons of pixel values. Most descriptors belonging to this group can actually be considered as LBP variants [7, 29].

3.2. Color analysis

The extension of grayscale rank-based methods to color images usually involves computing the ranking on each color channel separately and/or on pairs of color channels [6]. A classic example are the opponent-color local binary patterns proposed by Mäenpää and Pietikäinen [31]. A similar approach was introduced by Bianconi et al. [5] to compute rotation-invariant color texture features from ranklets [35]. More involved procedures to compute intra- and inter-channel features have been proposed by Lee et al. [27] (local color vector binary patterns), and, more recently, by Cusano et al. [12] (local angular patterns).

An alternative strategy is that of inducing an arbitrary ordering (typically M- or R-ordering) on the multivariate color data. Ledoux et al. [26] is an up-to-date, thorough investigation on this class of methods. In the remainder of this section we describe the three approaches that we considered for the comparative study, i.e.: lexicographic order, bit mixing and reference color.

Lexicographic order This well-known color comparison relation is a total order based on priorities between the components of a vector [11]. Using the priorities $R \succ G \succ B$ where $\succ$ means “has priority over”, a color $c_0 = (r_0, g_0, b_0)$ in the RGB space is less than or equal to a color $c_1 = (r_1, g_1, b_1)$ according to the following rule:

$$c_0 \leq c_1 \iff (r_0 < r_1) \lor [(r_0 = r_1) \land (g_0 < g_1)]$$

$$\lor [(r_0 = r_1) \land (g_0 = g_1) \land (b_0 \leq b_1)] \quad (1)$$

In the experiments we considered the following priority sequences: $R \succ G \succ B$, $B \succ G \succ R$, and $G \succ B \succ R$. Obviously, the main shortcoming of this approach is that the color order induced strongly depends on how the priorities are defined.

Bit mixing This is a total order widely used in multivalued morphological filtering [11]. This paradigm associates an integer value to each color from the binary representation of their chromatic components. For convenience let us denote a color in the RGB space as $c = (c_0, c_1, c_2)$ and the $n$-th bit of the $k$-th component as $c_{k,n}$, with $0 \leq k \leq 2$ and $0 \leq n \leq 7$ (we have implicitly assumed that intensity is discretized to 8-bits). We can build up a 24-bit integer by mixing the bits of the three chromatic components like this:

$$N(c) = \sum_{n=0}^{7} 2^{3n} \sum_{k=0}^{2} 2^{k} c_{\pi(k),n} \quad (2)$$

where $\pi$ represents a permutation of the chromatic channels (represented by their indices $\{0, 1, 2\}$). The bit mixing order is then defined as:

$$c_0 \leq c_1 \iff N(c_0) \leq N(c_1) \quad (3)$$

In the experiments we took into account the following bit-mixing permutations: RGB, BGR, and GBR. Similarly to the lexicographical order, the bit mixing order strongly depends on how the color components are permuted.

Reference color This preorder relies on the Euclidean distance between a color and a reference color, referred to as $c_{\text{ref}}$. The relation that compares two colors $c_0$ and $c_1$ with respect to the reference color is defined as [25]:

$$c_0 \preceq c_1 \iff \|c_0 - c_{\text{ref}}\| \leq \|c_1 - c_{\text{ref}}\| \quad (4)$$

Herein we used two reference colors: black $(0, 0, 0)$ and a shade of gray $(127, 127, 127)$. Needless to say that the reference color should be judiciously chosen since this choice may have a great impact on the resulting order.
4. Proposed approach

In the following subsections we provide a description and mathematical formulation of the image descriptor proposed in this paper. But before getting into the technicalities of the method we recall the basics of the rank transform (Sec. 4.1) and product order (Sec. 4.2).

4.1. Rank transform

The rank transform was first proposed by Zabih and Woodfill [38]. In this model the image is scanned through a sliding window of $N \times N$ pixels and the feature value at each position is the number of pixels in the periphery of the local neighborhood. The intensity of the central pixel, $I_c$, is the RGB triplet corresponding to the central pixel: $I_c = (r_c, g_c, b_c)$.

The neighborhood, $X$, where the RGB coordinates of two pixels, the product order relation between the central pixel and a peripheral pixel is defined as $\leq_P$, which results in a set of possible patterns. Image features are the dense, orderless distribution over the set of all possible patterns. Note that the features so calculated are intrinsically invariant to rotation and to any permutation of the peripheral pixels.

4.2. Product order

Given an ordered set $X$, one can induce a partial ordering on the Cartesian product $X \times X \times X$ [14]. This is referred to as the product order (also called component-wise order). If $X$ is the set of intensity levels of one channel of an RGB image (for an 8-bit intensity scale such set would be formed by the integers from 0 to 255) this can be extended to order colors in a straightforward manner. Given two triplets $c_0 = (r_0, g_0, b_0)$ and $c_1 = (r_1, g_1, b_1)$ representing the RGB coordinates of two pixels, the product order relation establishes that:

$$ c_0 \preceq_P c_1 \iff r_0 \leq r_1 \land g_0 \leq g_1 \land b_0 \leq b_1 \quad (7) $$

Conversely, when two colors are not comparable we shall write:

$$ c_0 \preceq_P c_1 \iff c_0 \not\preceq_P c_1 \land c_1 \not\preceq_P c_0 \quad (8) $$

Note that in the above equations we use symbol $\preceq_P$ rather than $\preceq$ to emphasize the fact that this binary relation is a partial order, and subscript $\times$ to signal that this particular partial order is the product order.

It is instructive to show that the product order in the RGB space has an intuitive geometric interpretation. Let us consider a generic color $c_0 = (r_0, g_0, b_0)$ in the RGB space as shown in Fig. 2, where points $(0, 0, 0)$ and $(1, 1, 1)$ respectively indicate colors ‘black’ and ‘white’. We have that:

- Any color in the (closed) green region dominates $c_0$
- Any color in the (closed) brown region is dominated by $c_0$
- Any other color outside the two above regions is incomparable to $c_0$

Though the product order (as well as other types of partial orders) can be applied to any color space, in this work we restrict the study to partial ordering in the RGB space.

4.3. Rank transform based on product order (RTPO)

Let $\phi$ be a function that takes in two RGB triplets and returns 1 when the first one is dominated by the second as per the product order and 0 otherwise, and $\psi$ another function that flags (with a 1) the situation in which the arguments are not comparable:

$$ \phi(c_j, c_c) = \begin{cases} 1, & \text{if } c_j \preceq_P c_c \\ 0, & \text{otherwise} \end{cases} \quad (9) $$

$$ \psi(c_j, c_c) = \begin{cases} 1, & \text{if } c_j \not\preceq_P c_c \\ 0, & \text{otherwise} \end{cases} \quad (10) $$

where $c_c$ is the RGB triplet corresponding to the central pixel and $c_j$, $0 \leq j \leq P - 1$, is the RGB triplet of the $j$-th peripheral pixel. Notice that in Eq. 10 we replaced the symbol $\preceq$ introduced in Eq. 7 by $\preceq_P$ to be consistent with the strict inequality employed in the original implementation of the rank transform [38]. Those functions are handy to compute the number of peripheral pixels that are dominated by $(\lambda)$ and not comparable to $(\nu)$ the central pixel:

$$ \lambda = \sum_{j=0}^{P-1} \phi(c_j, c_c) \quad (11) $$

$$ \nu = \sum_{j=0}^{P-1} \psi(c_j, c_c) \quad (12) $$

1035
Both are non negative integer numbers, and are constrained to $0 \leq \lambda + \nu \leq P$. The kernel function that assigns a label to a local pattern can be expressed as a function of those numbers:

$$f_{\text{RTPO}}(x) = \nu + \lambda + \frac{\lambda(2P + 3 - \lambda)}{2} \quad (13)$$

where $x$ in this case represents an array containing the RGB triplets of the pixels in the local neighborhood.

To further illustrate the kernel above, in Tab. 2 we display the pattern labels for a $3 \times 3$ neighborhood as a function of $\lambda$ and $\nu$. Clearly the number of possible features $F$ can be computed as the number of ways to distribute $n$ unlabelled balls into $k$ labelled urns $[3, \text{Th. 4.2.6}]$; the balls being the peripheral pixels and the urns the possible outcomes of the comparison with the central pixel, i.e.: dominates, is dominated or non comparable. Since $n = P$ and $k = 3$ we have:

$$F = \frac{(n + k - 1)}{k - 1} = \frac{(P + 2)}{2} = \frac{(P + 2)(P + 1)}{2} \quad (14)$$

Thus the dimensions of the histograms that result from the neighborhoods of size $3 \times 3$, $5 \times 5$ and $7 \times 7$ are 45, 153 and 325, respectively.

5. Experimental results

To assess the effectiveness of the proposed approach we ran a set of image classification experiments on nine datasets of color texture images from different sources. Tab. 3 summarizes the main properties of each dataset along with references/URLs.

<table>
<thead>
<tr>
<th>$\nu$</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
<td>7</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>9</td>
<td>10</td>
<td>11</td>
<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
<td>16</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>17</td>
<td>18</td>
<td>19</td>
<td>20</td>
<td>21</td>
<td>22</td>
<td>23</td>
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<td></td>
</tr>
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<td>3</td>
<td>24</td>
<td>25</td>
<td>26</td>
<td>27</td>
<td>28</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>4</td>
<td>30</td>
<td>31</td>
<td>32</td>
<td>33</td>
<td>34</td>
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<td>35</td>
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<td>40</td>
<td>41</td>
<td></td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>42</td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>8</td>
<td>44</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

Our primary goal was to comparatively assess, using a set-up as simple as possible, the performance of RTPO against that of the RT based on gray levels (original RT) and on the orders and preorders described in Sec. 3.2. To this end we used a simple $3 \times 3$ square neighborhood of pixels and the nearest neighbor classifier (1-NN) with Euclidean distance. The use of this classifier is standard practice in comparative studies – see for instance $[22, 28, 13]$ –
Table 3. Datasets used in the experiments: round-up table

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Classes</th>
<th>Samples per class</th>
<th>Sample size (px)</th>
<th>Variations in imaging conditions</th>
<th>Sample images</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Kather</td>
<td>8</td>
<td>625</td>
<td>150 × 150</td>
<td>None</td>
<td></td>
<td>[23]</td>
</tr>
<tr>
<td>2</td>
<td>KTH-TIPS2b</td>
<td>11</td>
<td>432</td>
<td>200 × 200</td>
<td>4 ☀, 9 ☀, 3 ☐</td>
<td></td>
<td>[8]</td>
</tr>
<tr>
<td>3</td>
<td>Kylberg-Sintorn</td>
<td>25</td>
<td>216</td>
<td>1728 × 1728</td>
<td>9 ☐</td>
<td></td>
<td>[24]</td>
</tr>
<tr>
<td>4</td>
<td>MondialMarmi</td>
<td>25</td>
<td>40</td>
<td>1500 × 1500</td>
<td>10 ☐</td>
<td></td>
<td>[4]</td>
</tr>
<tr>
<td>5</td>
<td>Outex-13</td>
<td>68</td>
<td>20</td>
<td>128 × 128</td>
<td>None</td>
<td></td>
<td>[33]</td>
</tr>
<tr>
<td>6</td>
<td>PapSmear</td>
<td>2</td>
<td>204</td>
<td>67 × 92 – 300 × 768</td>
<td>Unspecified</td>
<td></td>
<td>[21]</td>
</tr>
<tr>
<td>7</td>
<td>PlantLeaves</td>
<td>20</td>
<td>60</td>
<td>128 × 128</td>
<td>None</td>
<td></td>
<td>[9]</td>
</tr>
<tr>
<td>8</td>
<td>RawFooT</td>
<td>68</td>
<td>184</td>
<td>400 × 400</td>
<td>46 ☀</td>
<td></td>
<td>[13]</td>
</tr>
<tr>
<td>9</td>
<td>STex</td>
<td>476</td>
<td>16</td>
<td>128 × 128</td>
<td>Unspecified</td>
<td></td>
<td>[36]</td>
</tr>
</tbody>
</table>

KEY TO SYMBOLS: ☀ = illumination, ☐ = rotation, ☐ = scale, ☐ = viewpoint

due to the absence of tuning parameters, ease of implementation and other desirable asymptotic properties. Accuracy estimation was based on split-half validation with stratified sampling over 100 random splits. For each subdivision into train and test set we computed the fraction of samples of the test set correctly classified and averaged the results over the 100 splits.

The results obtained are summarized in the box plot of Fig. 3. As can be seen, in most datasets RTPO outperformed the other methods by a large margin. We also notice that the overall accuracy varied a lot from one dataset to another: this is due to the fact that the image datasets differ from each other in a number of significant factors such as the overall number of classes, the imaging conditions and the stationariness of the textures. To test whether there was significant difference between the accuracy of RTPO and that of the other descriptors we performed a two-sided Wilcoxon signed-rank test at significance level $\alpha = 1\%$. The results (Tab. 4) show that RTPO significantly outperformed the other descriptors in eight datasets out of nine, whereas in the remaining one the difference was very slight ($<0.1\%$).

We also performed a second group of experiments to determine how much the accuracy of RTPO features could be improved by using a multi-resolution approach and more sophisticated classifiers such as decision trees and support vector machines. This experiment included datasets ‘Kather’, ‘KTH-TIPS2b’, ‘Outex-13’ and ‘RawFooT’. The multi-scale approach consisted of concatenating the feature vectors resulting from employing the three different neighborhoods shown in Fig. 1. Optimal parameters $C$ and $\gamma$ for the rbf-SVM classifier were estimated, for each descriptors and image dataset, through exhaustive grid search. The parameters of the decision tree classifier (i.e.: maximum depth of the tree, minimum number of samples required to split an internal node and minimum number of samples required to be at a leaf node) were determined in a similar fashion. In the case of decision trees the dimension of the feature vectors was preliminary reduced through PCA before classification.

The results of this second group of experiments are graphically displayed in Fig. 4. Here the labels ‘1’, ‘2’ and ‘3’ respectively denote: the features obtained with the $3 \times 3$ neighborhood, the concatenation of the features obtained with the $3 \times 3$ and $5 \times 5$ neighborhoods and the concatenation of the features obtained with the $3 \times 3$, $5 \times 5$ and $7 \times 7$ neighborhoods. The total number of features by the three configurations respectively is 45, 198 and 523.

The plots in Fig. 4 clearly show that the overall accuracy significantly improved by adopting the multi-resolution approach and SVM classification. Decision trees, however, did not provide any benefit compared with nearest-neighbor classification.
Figure 3. Box plots of the success rates obtained on the different datasets. RT: rank transform on gray levels; Lex: RT on lexicographic orders; Mix: RT on bit mixing orders; Ref: RT on reference color preorders; RTPO: the proposed enhanced RT over the product order.

Table 4. Results of the Wilcoxon signed-rank test at significance level 1%. For all the datasets the rank transform based on the product order is compared to the best performing benchmark descriptor.

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Best preorder</th>
<th>Mean accuracy</th>
<th>p-value</th>
<th>Significant?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Best preorder</td>
<td>Partial order</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Kather</td>
<td>Ref 127</td>
<td>64.67%</td>
<td>77.40%</td>
<td>3.88e-18</td>
</tr>
<tr>
<td>2</td>
<td>KTH-TIPS2b</td>
<td>Lex GBR</td>
<td>73.57%</td>
<td>87.71%</td>
<td>3.88e-18</td>
</tr>
<tr>
<td>3</td>
<td>Kylberg-Sintorn</td>
<td>Lex RGB</td>
<td>99.07%</td>
<td>99.80%</td>
<td>3.84e-18</td>
</tr>
<tr>
<td>4</td>
<td>MondialMarmi</td>
<td>Lex BGR</td>
<td><strong>100.00%</strong></td>
<td>99.94%</td>
<td>1.83e-05</td>
</tr>
<tr>
<td>5</td>
<td>Outex-13</td>
<td>Lex RGB</td>
<td>64.26%</td>
<td>64.84%</td>
<td>3.74e-03</td>
</tr>
<tr>
<td>6</td>
<td>PapSmear</td>
<td>Ref 0</td>
<td>66.24%</td>
<td><strong>83.48%</strong></td>
<td>3.85e-18</td>
</tr>
<tr>
<td>7</td>
<td>PlantLeaves</td>
<td>Ref 0</td>
<td>50.74%</td>
<td><strong>63.41%</strong></td>
<td>3.88e-18</td>
</tr>
<tr>
<td>8</td>
<td>RawFooT</td>
<td>Ref 0</td>
<td>80.51%</td>
<td><strong>92.29%</strong></td>
<td>3.89e-18</td>
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<tr>
<td>9</td>
<td>STex</td>
<td>Ref 0</td>
<td>33.14%</td>
<td><strong>69.15%</strong></td>
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</table>
6. Conclusions and future work

In this paper we have introduced RTPO (rank transform based on product order), a compact yet effective local image descriptor for color textures. Our approach generalizes a classic descriptor (rank transform) to RGB color images via a particular type of partial order, i.e. the product order.

In a set of color texture classification experiments RTPO proved more effective than the original rank transform – either based on grayscale on other arbitrary orders and pre-orders on the RGB space. We found that the improvement was statistically significant. The highest success rates were obtained with a multi-scale approach and SVM classification. Remarkably, the proposed method provides a compact representation of color texture, the number of features for square neighborhoods of size $3 \times 3$, $5 \times 5$ and $7 \times 7$ respectively being 45, 153 and 325.

As far as we know the use of partial orders is entirely novel in this context. The results established so far seem very promising and encourage further investigation. In particular, we plan to utilize the product order with other rank features to make them suitable for color texture analysis. Another direction for future research is the investigation of other types of partial orders (different than the product order) that may better integrate color information into rank features.

References


