Spotting Differences Among Observations

Marko Rak, Tim König and Klaus-Dietz Tönies

Department of Simulation and Graphics, Otto-von-Guericke University, Magdeburg, Germany

{rak, tim.koenig, toennies}@isg.cs.ovgu.de

Keywords: Density Difference, Kernel Density Estimation, Scale Space, Blob Detection, Affine Shape Adaption

Abstract: Identifying differences among the sample distributions of different observations is an important issue in many fields ranging from medicine over biology and chemistry to physics. We address this issue, providing a general framework to detect difference spots of interest in feature space. Such spots occur not only at various locations, they may also come in various shapes and multiple sizes, even at the same location. We deal with these challenges in a scale-space detection framework based on the density function difference of the observations. Our framework is intended for semi-automatic processing, providing human-interpretable interest spots for further investigation of some kind, e.g., for generating hypotheses about the observations. Such interest spots carry valuable information, which we outline at a number of classification scenarios from UCI Machine Learning Repository; namely, classification of benign/malign breast cancer, genuine/forged money and normal/spondylolisthetic/disc-herniated vertebral columns. To this end, we establish a simple decision rule on top of our framework, which bases on the detected spots. Results indicate state-of-the-art classification performance, which underpins the importance of the information that is carried by these interest spots.

1 INTRODUCTION

Sooner or later a large portion of pattern recognition tasks come down to the question What makes X different from Y? Some scenarios of that kind are:

- Detection of forged money based on image-derived features: What makes some sort of forgery different from genuine money?
- Comparison of medical data of healthy and non-healthy subjects for disease detection: What makes data of the healthy different from that of the non-healthy?
- Comparison of document data sets for text retrieval purposes: What makes this set of documents different from another set?

Apart from this, spotting differences in two or more observations is of interest in fields of computational biology, chemistry or physics. Looking at it from a general perspective, such questions generalize to

What makes the samples of group X different from the samples of group Y?

This question usually arises when we deal with grouped samples in some feature space. For humans, answering such questions tends to become more challenging with increasing number of groups, samples and feature space dimensions, up to the point where we miss the forest for the trees. This complexity is not an issue to automatic approaches, which, on the other hand, tend to either overfit or underfit patterns in the data. Therefore, semi-automatic approaches are needed to generate a number of interest spots which are to be looked at in more detail.

We address this issue by a scale-space difference detection framework. Our approach relies on the density difference of group samples in feature space. This enables us to identify spots where one group dominates the other. We draw on kernel density estimators to represent arbitrary density functions. Embedding this into a scale-space representation, we are able to detect spots of different sizes and shapes in feature space in an efficient manner. Our framework:

- applies to d-dimensional feature spaces
- is able to reflect arbitrary density functions
- selects optimal spot locations, sizes and shapes
- is robust to outliers and measurement errors
- produces human-interpretable results

Our presentation is structured as follows. We outline the key idea of our framework in Section 2. The specific parts of our framework are detailed in Section 3, while Section 4 comprises our results on several data sets. In Section 5, we close with a summary of our work, our most important results and an outline of future work.
2 FOUNDATIONS

Searching for differences between the sample distribution of two groups of observations $g$ and $h$, we, quite naturally, seek for spots where the density function $f^g(x)$ of group $g$ dominates the density function $f^h(x)$ of group $h$, or vice versa. Hence, we try to find positive-negative-valued spots of the density difference

$$f^{g-h}(x) = f^g(x) - f^h(x)$$

w.r.t. the underlying feature space $\mathbb{R}^d$ with $x \in \mathbb{R}^d$. Such spots may come in various shapes and sizes. A difference detection framework should be able to deal with these degrees of freedom. Additionally, it must be robust to various sources of error, e.g., from measurement, quantization and outliers.

We propose to superimpose a scale-space representation to the density difference $f^{g-h}(x)$ to achieve the above-mentioned properties. Scale-space frameworks have been shown to robustly handle a wide range of detection tasks for various types of structures, e.g., text strings (Yi and Tian, 2011), persons and animals (Felzenszwalb et al., 2010) in natural scenes, neuron membranes in electron microscopy imaging (Seyedhosseini et al., 2011) or microaneurysms in digital fundus images (Adal et al., 2014). In each of these tasks the function of interest is represented through a grid of values, allowing for an explicit evaluation of the scale-space. However, an explicit grid-based approach becomes intractable for higher-dimensional feature spaces.

In what follows, we show how a scale-space representation of $f^{g-h}(x)$ can be obtained from kernel density estimates of $f^g(x)$ and $f^h(x)$ in an implicit fashion, expressing the problem by scale-space kernel density estimators. Note that by the usage of kernel density estimates our work is limited to feature spaces with dense filling. We close with a brief discussion on how this can be used to compare observations among more than two groups.

2.1 Scale Space Representation

First, we establish a family $l^{g-h}(x;t)$ of smoothed versions of the density difference $l^{g-h}(x)$. Scale parameter $t \geq 0$ defines the amount of smoothing that is applied to $l^{g-h}(x)$ via convolution with kernel $k_t(x)$ of bandwidth $t$ as stated in

$$l^{g-h}(x;t) = k_t(x) * f^{g-h}(x).$$

For a given scale $t$, spots having a size of about $2\sqrt{t}$ will be highlighted, while smaller ones will be smoothed out. This leads to an efficient spot detection scheme, which will be discussed in Section 3. Let

$$l^g(x;t) = k_t(x) * f^g(x)$$

$$l^h(x;t) = k_t(x) * f^h(x)$$

be the scale-space representations of the group densities $f^g(x)$ and $f^h(x)$. Looking at Equation 2 more closely, we can rewrite $l^{g-h}(x;t)$ equivalently in terms of $l^g(x;t)$ and $l^h(x;t)$ via Equation 3 and 4. This reads

$$l^{g-h}(x;t) = k_t(x) * [f^g(x) - f^h(x)]$$

$$= k_t(x) * [l^g(x;t) - l^h(x;t)].$$

The simple yet powerful relation between the left and the right-hand side of Equation 8 will allow us to evaluate the scale-space representation $l^{g-h}(x)$ implicitly, i.e., using only kernel functions. Of major importance is the choice of the smoothing kernel $k_t(x)$. According to scale-space axioms, $k_t(x)$ should suffice a number of properties, resulting in the uniform Gaussian kernel of Equation 9 as the unique choice, cf. (Babaud et al., 1986; Yuille and Poggio, 1986).

$$\phi_t(x) = \frac{1}{\sqrt{2\pi t}} \exp \left(-\frac{1}{2\pi} x^T x \right)$$

2.2 Kernel Density Estimation

In kernel density estimation, the group density $f^g(x)$ is estimated from its $n^g$ samples by means of a kernel function $K_B^g(x)$. Let $x^g_i \in \mathbb{R}^d$ with $i = 1, \ldots, n^g$ being the group samples. Then, the group density estimate is given by

$$\hat{f}^g(x) = \frac{1}{n^g} \sum_{i=1}^{n^g} K_B^g \left(x - x^g_i\right).$$

Parameter $B^g \in \mathbb{R}^{d \times d}$ is a symmetric positive definite matrix, which controls the influences of samples to the density estimate. Informally speaking, $K_B^g(x)$ applies a smoothing with bandwidth $B^g$ to the "spiky sample relief" in feature space.

Plugging kernel density estimator $\hat{f}^g(x)$ into the scale-space representation $l^g(x;t)$ defines the scale-space kernel density estimator $\hat{l}^g(x;t)$ to be

$$\hat{l}^g(x;t) = k_t(x) * \hat{f}^g(x).$$

Inserting Equation 10 into the above, we can trace down the definition of the scale-space density estima-
tor $\tilde{f}(x;t)$ to the sample level via transformation

$$\tilde{f}(x;t) = k_t(x) * \hat{f}(x) = k_t(x) * \left[ \frac{1}{n^g} \sum_{i=1}^{n^g} K_{B^g}(x - x_i^g) \right] = \frac{1}{n^g} \sum_{i=1}^{n^g} (k_t * K_{B^g})(x - x_i^g).$$

(13)

Though arbitrary kernels can be used, we choose $K_B(x)$ to be a Gaussian kernel $\Phi_B(x)$ due to its convenient algebraic properties. This (potentially non-uniform) kernel is defined as

$$\Phi_B(x) = \frac{1}{\sqrt{\det(2\pi B)}} \exp \left(-\frac{1}{2} x^T B^{-1} x \right).$$

(15)

Using the above, the right-hand side of Equation 14 simplifies further because of the Gaussian’s cascade convolution property. Eventually, the scale-space kernel density estimator $\hat{f}(x;t)$ is given by Equation 16, where $I \in \mathbb{R}^{d \times d}$ is the identity.

$$\hat{f}(x;t) = \frac{1}{n^g} \sum_{i=1}^{n^g} \Phi_{I+B^g}(x - x_i^g).$$

(16)

Using this estimator, the scale-space representation $F^g(x;t)$ of group density $f^g(x)$ and analogously that of group $h$ can be estimated for any $(x;t)$ in an implicit fashion. Consequently, this allows us to estimate the scale-space representation $F^{g-h}(x;t)$ of the density difference $f^{g-h}(x)$ via Equation 7 by means of kernel functions only.

### 2.3 Bandwidth Selection

When regarding bandwidth selection in such a scale-space representation, we see that the impact of different choices for bandwidth matrix $B$ vanishes as scale $t$ increases. This can be seen when comparing matrices $I + 0$ and $I + B$ where $0$ represents the zero matrix, i.e., no bandwidth selection at all. We observe that relative differences between them become negligible once $\|I\| \gg \|B\|$. This is especially true for large sample sizes, because the bandwidth will then tend towards zero for any reasonable bandwidth selector anyway. Hence, we may actually consider setting $B$ to $0$ for certain problems, as we typically search for differences that fall above some lower bound for $t$.

Literature bares extensive work on bandwidth matrix selection, for example, based on plug-in estimators (Duong and Hazelton, 2003; Wand and Jones, 1994) or biased, unbiased and smoothed cross-validation estimators (Duong and Hazelton, 2005; Sain et al., 1992). All of these integrate well with our scale-space density difference framework. However, in view of the argument above, we propose to compromise between a full bandwidth optimization and having no bandwidth at all. We define $B^g = b^g I$ and use an unbiased least-squares cross-validation to set up the bandwidth estimate for group $g$. For Gaussian kernels, this leads to the optimization of 17, cf. (Duong and Hazelton, 2005), which we achieved by golden section search over $b^g$.

$$\arg \min_{b^g} \frac{1}{n^g \sqrt{\det(4\pi B^g)}} + \frac{1}{n^g(n^g - 1)} \sum_{i=1}^{n^g} \sum_{j \neq i}^{n^g} (\Phi_{2B^g} - 2\Phi_{B^g})(x_i^g - x_j^g).$$

(17)

### 2.4 Multiple Groups

If differences among more than two groups shall be detected, we can reduce the comparison to a number of two-group problems. We can consider two typical use cases, namely one group vs. another and one group vs. rest. Which of the two is more suitable depends on the specific task at hand. Let us illustrate this using two medical scenarios. Assume we have a number of groups which represent patients having different diseases that are hard to discriminate in differential diagnosis. Then we may consider the second use case, to generate clues on markers that make one disease different from the others. In contrast, if these groups represent stages of a disease, potentially including a healthy control group, then we may consider the first use case, comparing only subsequent stages to give clues on markers of the disease’s progress.

### 3 METHOD

To identify the positive-negative-valued spots of a density difference, we apply the concept of blob detection, which is well-known in computer vision, to the scale-space representation derived in Section 2. In scale-space blob detection, some blobness criterion is applied to the scale-space representation, seeking for local optima of the function of interest w.r.t. space and scale. This directly leads to an efficient detection scheme that identifies a spot’s location and size. The latter corresponds to the detection scale.

In a grid-representable problem we can evaluate blobness densely over the scale-space grid and identify interesting spots directly using the grid neighborhood. This is intractable here, which is why we rely on a more refined three-stage approach. First, we trace the local spatial optima of the density difference.
through scales of the scale-space representation. Second, we identify the interesting spots by evaluating their blobnness along the dendrogram of optima that was obtained during the first stage. Having selected spots and therefore knowing their locations and sizes, we finally calculate an elliptical shape estimate for each spot in a third stage.

Spots obtained in this fashion characterize elliptical regions in feature space as outlined in Figure 1. The representation of such regions, i.e., location, size and shape, as well as its strength, i.e., its scale-space density difference value, are easily interpretable by humans, which allows to look at them in more detail using some other method. This also renders a limitation of our work, because non-elliptical regions may only be approximated by elliptical ones. We now give a detailed description of the three stages.

### 3.1 Scale Tracing

Assume we are given an equidistant scale sampling, containing non-negative scales $t_1, \ldots, t_n$ in increasing order and we search for spots where group $g$ dominates $h$. More precisely, we search for the non-negatively valued maxima of $l^{g-h}(x; t_i)$. The opposite case, i.e., group $h$ dominates $g$, is equivalent.

Let us further assume that we know the spatial local maxima of the density difference $l^{g-h}(x; t_{i-1})$ for a certain scale $t_{i-1}$ and we want to estimate those of the current scale $t_i$. This can be done taking the previous local maxima as initial points and optimizing each w.r.t. $l^{g-h}(x; t_i)$. In the first scale, we take the samples of group $g$ themselves. As some maxima may converge to the same location, we merge them together, feeding unique locations as initials into the next scale $t_{i+1}$ only. We also drop any negatively-valued locations as these are not of interest to our task. They will not become of interest for any higher scale either, because local extrema will not enhance as scale increases, cf. (Lindeberg, 1998). Since derivatives are simple to evaluate for Gaussian kernels, we can use Newton’s method for spatial optimization.

We can assemble gradient \( \frac{\partial^2 l^{g-h}}{\partial x \partial x^T} \) and Hessian \( \frac{\partial^2 \Phi_B}{\partial x^2} \) sample-wise using

\[
\frac{\partial}{\partial x} \Phi_B(x) = -\Phi_B(x) B^{-1} x \quad \text{and} \quad (18)
\]

\[
\frac{\partial^2 \Phi_B}{\partial x \partial x^T} = \Phi_B(x) (B^{-1} x x^T B^{-1} - B^{-1}) \quad \text{and} \quad (19)
\]

Iterating this process through all scales, we form a discreet dendrogram of the maxima over scales. A dendrogram branching means that a maxima formed from two (or more) maxima from the preceding scale.

### 3.2 Spot Detection

The maxima of interest are derived from a scale-normalized blobnness criterion $c_t(x; t)$. Two main criteria, namely the determinant of the Hessian (Bretzner and Lindeberg, 1998) and the trace of the Hessian (Lindeberg, 1998) have been discussed in literature. We focus on the former, which is given in Equation $20^1$, as it has been shown to provide better scale selection properties under affine transformation of the feature space, cf. (Lindeberg, 1998).

\[
c_t(x; t) = t^{nd} \left\{ (-1)^d \det \left( \frac{\partial^2 \Phi_B}{\partial x \partial x^T} l^{g-h}(x; t) \right) \right\} \quad (20)
\]

\[
c_t(x; t) = t^{nd} c(x; t) \quad (21)
\]

\(^1(-1)^d\) handles even and odd dimensions consistently.
Because the maxima are already spatially optimal, we can search for spots that maximize $c_f(x;t)$ w.r.t. the dendrogram neighborhood only. Parameter $\gamma \geq 0$ can be used to introduce a size bias, shifting the detected spot towards smaller or larger scales. The definition of $\gamma$ highly depends on the type of spot that we are looking for, cf. (Lindeberg, 1996). This is impractical when we seek for spots of, for example, small and large skewness or extreme kurtosis at the same time.

Addressing the parameter issue, we search for all spots that maximize $c_f(x;t)$ locally w.r.t. some $\gamma \in [0, \infty)$. Some dendrogram spot $s$ with scale-space coordinates $(x_s; t_s)$ is locally maximal if there exists a $\gamma$-interval such that its blobness $c_f(x_s; t_s)$ is larger than that of every spot in its dendrogram neighborhood $\mathcal{N}(s)$. This leads to a number of inequalities, which can be written as

$$
t_f^\gamma c(x_s; t_s) > \max_{s \in \mathcal{N}(s)} t_f^\gamma c(x_s; t_s), \quad \text{or} \quad (22)
$$

$$
\gamma d \log \frac{t_s}{t_n} > \max_{s \in \mathcal{N}(s)} \frac{c(x_s; t_s)}{c(x_n; t_n)}, \quad \text{or} \quad (23)
$$

The latter can be solved easily for the $\gamma$-interval, if any. We can now identify our interest spots by looking for the maxima along the dendrogram that locally maximize the width of the $\gamma$-interval. More precisely, let $w_f(x_s; t_s)$ be the width of the $\gamma$-interval for dendrogram spot $s$, then $s$ is of interest if the dendrogram Laplacian of $w_f(x; t)$ is negative at $(x_s, t_s)$, or equivalently, if

$$
w_f(x_s; t_s) > \frac{1}{|\mathcal{N}(s)|} \sum_{n \in \mathcal{N}(s)} w_f(x_n; t_n). \quad (24)$$

Intuitively, a spot is of interest if its $\gamma$-interval width is above neighborhood average. This is the only assumption we can make without imposing limitations on the results. Interest spots identified in this way will be dendrogram segments, each ranging over a number of consecutive scales.

### 3.3 Shape Adaptation

Shape estimation can be done in an iterative manner for each interest spot. The iteration alternatingly updates the current shape estimate based on a measure of anisotropy around the spot and then corrects the bandwidth of the scale-space smoothing kernel according to this estimate, eventually reaching a fixed point. The second moment matrix of the function of interest is typically used as an anisotropy measure, e.g., in (Lindeberg and Garding, 1994) and (Mikolajczyk and Schmid, 2004). Since it requires spatial integration of the scale-space representation around the interest spot, this measure is not feasible here.

We adapted the Hessian-based approach of (Lake-mond et al., 2012) to $d$-dimensional problems. The aim is to make the scale-space representation isotropic around the interest spot, iteratively moving any anisotropy into the symmetric positive definite shape matrix $S \in \mathbb{R}^{d \times d}$ of the smoothing kernel’s bandwidth $t$. Thus, we lift the problem into a generalized scale-space representation $t^{\gamma-h}(x; tS)$ of non-uniform scale-space kernels, which requires us to replace the definition of $\Phi_t(x)$ by that of $\Phi_t^\gamma(x)$.

Starting with the isotropic $S_1 = I$, we decompose the current Hessian via

$$
\frac{\partial^2}{\partial x \partial x^T} t^{\gamma-h}(\cdot; tS) = \Phi_t^\gamma(x) = \Phi_t^\gamma(x) = \Phi_t^\gamma(x)
$$

and start all over again. Iteration terminates when isotropy is reached. More precisely: when the ratio of minimal and maximal eigenvalue of the Hessian approaches one, which usually happens within a few iterations.

### 4 EXPERIMENTS

We next demonstrate that interest spots carry valuable information about a data set. Due to the lack of data sets that match our particular detection task a ground truth comparison is impossible. Certainly, artificially constructed problems are an exception. However, the generalizability of results is at least questionable for such problems. Therefore, we chose to benchmark our approach indirectly via a number of classification tasks. The rational is that results that are comparable to those of well-established classifiers should underpin the importance of the identified interest spots.

We next show how to use these interest spots for classification using a simple decision rule and detail the data sets that were used. We then investigate parameters of our approach and discuss the results of the classification tasks in comparison to decision trees, Fisher’s linear discriminant analysis, $k$-nearest neighbors with optimized $k$ and support vector machines with linear and cubic kernels. All experiments were performed via leave-one-out cross-validation.
Figure 2: Feature space decision boundaries (black plane curves) obtained from group likelihood criterion for the two-dimensional two-group problem of Figure 1; Red squares and blue circles visualize the samples of each group; Red/blue paths outline the dendrogram of scale-space density difference optima for the red/blue group dominating the other group; Interesting spots of each dendrogram are printed thick; Red/blue ellipses characterize the shape for each of the interest spots.

Figure 3: Sample group likelihoods and decision boundary (black diagonal line) for the two-group problem of Figure 1.

4.1 Decision Rule

To perform classification we establish a simple decision rule based on interest spots that were detected using the one group vs. rest use case. Therefore, we define a group likelihood criterion as follows. For each group $g$, having the set of interest spots $I_g$, we define

$$p_g(x) = \max_{s \in I_g} I^{g-h}(x_s, t_s S_s) \cdot \exp \left( -\frac{1}{2} (x - x_s)^T (t_s S_s)^{-1} (x - x_s) \right). \quad (28)$$

This is a quite natural trade-off, where the first factor favors spots $s$ with high density difference, while the latter factor favors spots with small Mahalanobis distance to the location $x$ that is investigated. We may also think of $p_g(x)$ as an exponential approximation of the scale-space density difference using interesting spots only. Given this, our decision rule simply takes the group that maximizes the group likelihood for the location of interest $x$. Figure 2 and Figure 3 illustrate the decision boundary obtained from this rule.

4.2 Data Sets

We carried out our experiments on three classification data sets taken from UCI Machine Learning Repository. A brief summary of them is given in Table 1. In the first task, we distinguish between benign and malign breast cancer based on manually graded cytological characteristics, cf. (Wolberg and Mangasarian, 1990). In the second task, we distinguish between genuine and forged money based on wavelet-transform-derived features from photographs of banknote-like specimen, cf. (Glock et al., 2009). In the third task, we differentiate among normal, spondylolisthetic and disc-herniated vertebral columns based on biomechanical attributes derived from shape and orientation of the pelvis and the lumbar vertebral column, cf. (Berthonnaud et al., 2005).

4.3 Parameter Investigation

Before detailing classification results, we investigate two aspects of our approach. Firstly, we inspect the importance of bandwidth selection, benchmarking no kernel density bandwidth against the least-squares cross-validation technique that we use. Secondly, we determine the influence of the scale sampling rate. For the latter we space $n+1$ scales for various $n$ equidistantly from zero to

$$t_n = F_{\chi^2}^{-1}(1-\epsilon|d) \max_g \left( \sqrt{\det(\Sigma_g)} \right), \quad (29)$$

where $F_{\chi^2}^{-1}(\cdot|d)$ is the cumulative inverse-$\chi^2$ distribution with $d$ degrees of freedom and $\Sigma_g$ is the covariance matrix of group $g$. Intuitively, $t_n$ captures the extent of the group with largest variance up to a small $\epsilon$, i.e., here $1.5 \cdot 10^{-8}$. 
To investigate the two aspects, we compare classification accuracies with and without bandwidth selection as well as sampling rates ranging from $n = 100$ to $n = 300$ in steps of 25. From the results, which are given in Table 2, we observe that bandwidth selection is almost negligible for the Breast Cancer (BC) and the Banknote Authentication (BA) data set. However, the impact is substantial throughout all scale sampling rates for the Vertebral Column (VC) data set. This may be due to the comparatively small number of samples per group for this data set. Regarding the second aspect, we observe that for the BA and VC data set the classification accuracy slightly increases when the scale sampling rate rises. Regarding the BC data set, accuracy remains relatively stable, except for the lower rates. From the results we conclude that bandwidth selection is a necessary part for interest spot detection. We further recommend $n \geq 200$, because accuracy starts to saturate at this point for all data sets. For the remaining experiments we used bandwidth selection and a sampling rate of $n = 200$.

### 4.4 Classification Results

A comparison of classification accuracies of our decision rule against the aforementioned classifiers is given in Table 3. For the BC data set we observe that except for the support vector machine (SVM) with cubic kernel all approaches were highly accurate, scoring between 94% and 97% with our decision rule being topmost. Even more similar to each other are results for the BA data set, where all approaches score between 97% and 99%, with ours lying in the middle of this range. Results are most diverse for the VC data sets. Here, the SVM with cubic kernel again performs significantly worse than the rest, which all score between 80% and 85%, while our decision rule peaks at 88%. Other research showed similar scores on the given data sets. For example the artificial neural networks based on pareto-differential evolution in (Abbass, 2002) obtained 98% accuracy for the BC data set, while (Rocha Neto et al., 2011) achieved 83% to 85% accuracy on the VC data set with SVMs with different kernels. These results suggest that our interest points carry information about a data set that are similarly important than the information carried by the well-established classifiers.

Confusion tables for our decision rule are given in Table 4 for all data sets. As can be seen, our approach gave balanced inter-group results for the BC and the BA data set. We obtained only small inaccuracies for the recall of the benign (96%) and genuine (97%) groups as well as for the precision of the malign (94%) and forged (96%) groups. Results for the VC data set were more diverse. Here, a number of samples with disc herniation were mistaken for being normal, lowering the recall of the herniated group (86%) noticeably. However, more severe inter-group imbalances were caused by the normal samples, which were relatively often mistaken for being spondylolisthetic or herniated discs. Thus, recall for the normal group (76%) and precision for the herniated group (74%) decreased significantly. The latter is to some degree caused by a handful of strong outliers from the normal group that fall into either of the other groups, which can already be seen from the group likelihood plot in Figure 4. This finding was made by others as well, cf. (Rocha Neto and Barreto, 2009).

The other classifiers performed similarly balanced on the BA and BC data set. Major differences occurred on the VC data set only. A precision/recall compar-
Table 4: Confusion table for predicted/actual groups of our decision rule for data sets of Table 1

(a) Breast Cancer

<table>
<thead>
<tr>
<th></th>
<th>ben.</th>
<th>mal.</th>
<th>ben.</th>
<th>mal.</th>
</tr>
</thead>
<tbody>
<tr>
<td>P A</td>
<td>429</td>
<td>4</td>
<td>15</td>
<td>235</td>
</tr>
<tr>
<td></td>
<td>99</td>
<td></td>
<td>94</td>
<td></td>
</tr>
<tr>
<td></td>
<td>96</td>
<td>98</td>
<td>[%]</td>
<td></td>
</tr>
</tbody>
</table>

(b) Banknote Authentication

<table>
<thead>
<tr>
<th></th>
<th>gen.</th>
<th>for.</th>
<th>100</th>
<th>96</th>
</tr>
</thead>
<tbody>
<tr>
<td>P A</td>
<td>742</td>
<td>0</td>
<td>20</td>
<td>610</td>
</tr>
<tr>
<td></td>
<td>97</td>
<td>100</td>
<td>[%]</td>
<td></td>
</tr>
</tbody>
</table>

(c) Vertebral Column

<table>
<thead>
<tr>
<th></th>
<th>norm.</th>
<th>spon.</th>
<th>hern.</th>
<th>76</th>
<th>1</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>P A</td>
<td></td>
<td></td>
<td></td>
<td>100</td>
<td>96</td>
<td>94</td>
</tr>
<tr>
<td></td>
<td>14</td>
<td>4</td>
<td>52</td>
<td>74</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>97</td>
<td>96</td>
<td>86</td>
<td>[%]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 4: Sample group likelihoods and decision boundary (black diagonal line) for the Vertebral Column data set of Table 1; Normal, spondylolisthetic and herniated discs in blue, magenta and red, respectively.

We observe that the precision of the normal and the herniated group are significantly lower (gap > 12\%) than that of the spondylolisthetic group for all classifiers except for our decision rule, for which at least the normal group is predicted with a similar precision. Regarding the recall we note an even more unbalanced behavior. Here, a strict ordering from spondylolisthetic over normal to herniated disks occurs. The differences of the recall of spondylolisthetic and normal are significant (gap > 16\%) and those between normal and herniated are even larger (gap > 18\%) among all classifiers that we compared against. The recalls for our decision rule are distributed differently, ordering the herniated before the normal group. Also the magnitude of differences is less significant (gaps \approx 10\%) for our decision rule. Results of this comparison indicate that the information that is carried by our interest points tends to be more balanced among groups than the information carried by the well-established classifiers that we compared against.

5 CONCLUSION

We proposed a detection framework that is able to identify differences among the sample distributions of different observations. Potential applications are manifold, touching fields such as medicine, biology, chemistry and physics. Our approach bases on the density function difference of the observations in feature space, seeking to identify spots where one observation dominates the other. Superimposing a scale-space framework to the density difference, we are able to detect interest spots of various locations, size and shapes in an efficient manner.

Our framework is intended for semi-automatic processing, providing human-interpretable interest spots for potential further investigation of some kind. We outlined that these interest spots carry valuable information about a data set at a number of classification tasks from the UCI Machine Learning Repository. To this end, we established a simple decision rule on top of our framework. Results indicate state-of-the-art performance of our approach, which underpins the importance of the information that is carried by these interest spots.

In the future, we plan to extend our work to support repetitive features such as angles, which currently is a limitation of our approach. Modifying our notion of distance, we would then be able to cope with problems defined on, e.g., a sphere or torus. Future work may also include the migration of other types of scale-space detectors to density difference problems. This includes the notion of ridges, valleys and zero-crossings, leading to richer sources of information.

ACKNOWLEDGEMENTS

This research was partially funded by the project “Visual Analytics in Public Health” (TO 166/13-2) of the German Research Foundation.
REFERENCES


