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Morphological classification of odontogenic keratocysts using Bouligand–Minkowski fractal descriptors

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ABSTRACT

The Odontogenic keratocyst (OKC) is a cystic lesion of the jaws, which has high growth and recurrence rates compared to other cysts of the jaws (for instance, radicular cyst, which is the most common jaw cyst type). For this reason OKCs are considered by some to be benign neoplasms. There exist two sub-types of OKCs (sporadic and syndromic) and the ability to discriminate between these sub-types, as well as other jaw cysts, is an important task in terms of disease diagnosis and prognosis. With the development of digital pathology, computational algorithms have become central to addressing this type of problem. Considering that only basic feature-based methods have been investigated in this problem before, we propose to use a different approach (the Bouligand–Minkowski descriptors) to assess the success rates achieved on the classification of a database of histological images of the epithelial lining of these cysts. This does not require the level of abstraction necessary to extract histologically-relevant features and therefore has the potential of being more robust than previous approaches. The descriptors were obtained by mapping pixel intensities into a three dimensional cloud of points in discrete space and applying morphological dilations with spheres of increasing radii. The descriptors were computed from the volume of the dilated set and submitted to a machine learning algorithm to classify the samples into diagnostic groups. This approach was capable of discriminating between OKCs and radicular cysts in 98% of images (100% of cases) and between the two sub-types of OKCs in 68% of images (71% of cases). These results improve over previously reported classification rates reported elsewhere and suggest that Bouligand–Minkowski descriptors are useful features to be used in histopathological images of these cysts.

1. Introduction

Cysts are pathological cavities containing fluid or semi-fluid content and lined by epithelial tissue. The diagnosis of jaw cysts is based on histopathological features, firstly to identify to which of the various cyst types a lesion corresponds to (with different origins, behaviour and prognosis), and secondly, to avoid misdiagnosis with other lesions (e.g. as intra osseous squamous cell carcinoma, unicystic ameloblastoma and other tumours) might present with similar radiographical features but require different treatments.

Among the cysts arising in the jaws, two important types are the ‘radicular cyst’ and the ‘odontogenic keratocyst’ (OKC). Radicular cysts are the most common type (55% of odontogenic cysts [10]), they are associated with the roots of teeth with non-vital pulps (e.g. due to advanced dental caries) and have slow growth. OKCs are less frequent (12% of odontogenic cysts [10]), they are not associated with dental disease and have certain characteristics common with neoplasms (e.g. active epithelial growth [21,14,15] and higher recurrence rates). Furthermore, there are two OKC sub-types; they can be solitary cysts (sporadic), or they can be multiple (synchronous or metachronous) as part of a rare autosomal dominant disease, the Gorlin-Goltz or Basal Cell Naevus Syndrome (BCNS) [11,20]. About 85% of syndromic and 30% of sporadic cases have mutations of the PTCH1 gene [13], (similarly to basal cell carcinomas of skin) indicating a potential common pathogenesis across OKC sub-types. However, there seem to be differences in the behaviour of syndromic and sporadic OKCs too, and therefore any morphological evidence that could help differentiating between the two sub-types is of diagnostic and predictive importance.

Woolgar et al. [22,23] summarised some histological differences
between syndromic and sporadic OKCs. While those features are relevant in histopathological terms to understand the organisation of the OKCs, they are difficult to characterise using automated or semi-automated approaches as they require contextual descriptions of cells and tissues that are not directly translated into quantitative features that machine learning algorithms can currently handle. Although an automated technique was proposed to classify other groups of oral cysts in [8], so far in the domain of machine diagnosis the only work addressing the diagnostic differences between syndromic and non-syndromic OKC cysts is [11], where the differences in the morphology of algorithmically segmented cells and the architectural organisation of the epithelial lining of inflammation-free regions of OKCs sub-types were investigated using semi-automated image analysis. Such differences, however, were not sufficient to achieve OKC sub-type classification purposes (60% correct classification rate), although they allowed a good discrimination of OKCs from the most common type of jaw cyst (i.e. radicular cysts) [11]. For expert histopathologists, the latter discrimination should not be a particularly difficult task, but is currently far from becoming automated. Furthermore, the differences between OKC sub-types are more difficult to quantify. If it was possible to rationalise the differences between all these lesions systematically, they could become histological discriminators for cases of BCNS, specially in retrospective analyses, in cases exhibiting low penetrance, and to develop indicative markers of recurrence potential. Advances in digital pathology and virtual slides technology have made attractive the possibility of developing machine learning algorithms to help pathologists gather reproducible morphological evidence for diagnostic purposes, to allow analysis of large datasets with multiple tissue sections and to evaluate the results of treatment modalities.

The purpose of this paper is to investigate a statistical approach to the classification of cyst lining images, based on a machine vision algorithm (Bouligand–Minkowski descriptors) that does not require high levels of segmentation of histologically relevant features.

2. Materials

The material used in this paper consists of an anonymised database of 150 images of haematoxylin and eosin (H & E) sections from formalin fixed and paraffin embedded specimens of developmental (solitary and syndromic OKCs) or inflammatory (radicular cysts) origin. Briefly, the database included 65 images of the lining from 13 cases of sporadic (k) OKCs, 40 images from 8 cases of syndromic (s) OKCs and 45 images from 9 cases of radicular (r) cysts (i.e. 5 non-overlapping images per case and with no sectioning artefacts). The cyst types were previously determined by histopathologists using histological as well as clinical information from the respective patients.

The images were captured using an Olympus BX50 microscope (Olympus Optical Co. Tokyo, Japan) with a x40 objective and a colour camera JVC KY-55B (JVC, Tokyo, Japan) attached to a 24 bit RGB frame grabber (Imaging Technologies IT4PCI, Bedford, MA, USA). With this setup the image size was 768×572 pixels with an inter-pixel distance of 0.31 µm. The images were background corrected for illumination uniformity (by means of the transmittance ratio of the specimen with an empty bright field frame), for camera bias (by subtraction of a non illuminated frame) and for camera shot noise (each image was the average of 32 consecutive frames). The extent of the epithelial lining of the lesions was segmented (from the background and connective capsule) by optical intensity thresholding (described below), and further analysis was done exclusively on the pixel values of the epithelial lining. Fig. 1 shows one sample for each group (sporadic OKC, radicular and syndromic OKC).

Finally, all the stained images were converted to grey values, where the in the intensity G of each pixel is obtained from the R, G and B components using the following weighted sum:

\[ G = 0.2989^*R + 0.5870^*G + 0.1140^*B \]  

(1)

3. Method

We investigated the classification performance of the Bouligand–Minkowski (B-M) descriptors developed in [2], to classify the images of cyst linings into their original diagnostic groups k (sporadic OKCs), s (syndromic OKCs) and r (radicular cysts). To our knowledge, this is the first proposal for using a statistical approach to this task, while previous works used morphological features that require more complicated pipelines to guarantee that the generated histologically-relevant features are segmented accurately. The method consists of three steps applied to each image: 1) segmenting the epithelial lining from the background (i.e. empty space of the cystic lumen and the surrounding cyst fibrous capsule), 2) computing the Bouligand–Minkowski descriptors of the lining image and 3) concatenating and submitting such descriptors to a classifier algorithm to predict the original type of cysts they belonged to. Section 3.1 briefly describes the steps involved in the segmentation, whereas Sections 3.2–4 provide the theoretical background on the B-M descriptors.

3.1. Segmentation

The segmentation of the epithelial lining was performed by a combination of colour deconvolution and morphological operations. First, colour deconvolution separated the main colour components of the hematoxylin and eosin stained regions into two channels. The epithelial lining typically showed more intense staining in the haematoxylin channel than the other parts of the sample. The histogram equalization and thresholding of this channel allowed highlighting the region containing the epithelial cells. Finally, morphological filtering (using binary reconstruction) was used to eliminate small features producing a clean image where the largest region closely corresponded to the epithelial lining of the cysts. Fig. 2 illustrates the procedure and each step involved. More details can be found in [11].

3.2. Fractal geometry

A fractal [16] is a set of points embedded in a topological space that exhibits self-similarity (i.e. the characteristic of being exactly or statistically similar independently of the scale of observation). Fractal geometry concerns the study of the properties that are scale-independent and the self-similarity of such objects. The most commonly used property to characterise such objects is the ‘fractal dimension’. This describes the rate of space filling of the set with scale and it can be defined through the Hausdorff-Besicovitch dimension, a concept from Measure Theory, which requires some knowledge of the analytical rules that generated the fractal set.

Fractal scale independence implies infinite amount of morphological detail. Real-world natural objects cannot be infinitely complex, yet some degree of self-similarity and complexity is a common feature. Therefore, it is possible to model such objects by means of fractal geometry concepts (e.g. the fractal dimension) even when there are no well-defined analytical rules (as required by Measure Theory) responsible for generating the sets. An alternative approximation to the Hausdorff-Besicovitch dimension consists of measuring some physical property of the object over a range of scales, e.g. with a ruler with measuring unit length e and counting the number \( N(e) \) of units necessary to measure the length of the object. Such abstraction can be extended to any embedding space endowed with a topological dimension \( D_f \) by enlarging the length \( e \) and recomputing \( N(e) \), to estimate the fractal dimension:

\[ D = D_f - \lim_{\epsilon \to 0} \frac{\log N(e)}{\log 1/e} \]  

(2)

In practice, the limit is estimated from the slope of a straight line fit to
the plot of \( \log N(\epsilon) \) vs. \( \log \frac{1}{\epsilon} \). Quantities other than \( N(\epsilon) \) can be used to estimate the self-similarity of objects using, e.g., methods such as box-counting, B-M dilation and mass-radius [6]. Here we concentrate on the B-M approach because it can be easily implemented and was shown to be useful in characterising a number of biological problems [19,2,7].

3.3. Bouligand–Minkowski dimension

The B-M fractal dimension, originally developed for the analysis of binary sets [6], was adapted here to analyse grey-level images. The procedure consists of two steps. First, the grey level image \( I: [1, M] \times [1, N] \rightarrow \mathbb{R} \) is mapped onto a cloud \( C \) of points within a discretised bounded region \( E \) of the three-dimensional Euclidean space. Each pixel with coordinates \((x, y, z) \in [1, M] \times [1, N]\) and intensity \( z \in \mathbb{R} \) is mapped onto a point \((x, y, z) \in C\). Fig. 3 illustrates such procedure.

Second, the points are gradually dilated by spheres with increasing radii \( r \), starting with \( r=1 \), and the total volume \( V(r) \) of the dilated cloud is measured. By replacing \( N(\epsilon) \) by \( V(r) \) in Eq. (2) and considering that the power-law scales with \( r^2 \) in the B-M dilation, the dimension is estimated by

\[
D_{\text{B-M}} = 3 - \lim_{r \to 0} \frac{\log V(r)}{\log 2r}.
\]

Fig. 1. Images of the three cysts classes investigated. Field width 241 µm. On top, the image as captured with the microscope. At the bottom, the region of interest containing exclusively cyst lining (epithelium), which is where the analysis was actually carried out. The cysts lumen is at the top of the image. The dark elliptic dark structures in the epithelium are the epithelial cell nuclei. Note the similarities between the OKC subtypes.

Fig. 2. Steps involved in the segmentation of the epithelial lining. (a) Original image. (b) Hematoxylin component. (c) Grey-level equalised version of (b). (d) Threshold. (e) Morphological reconstruction to remove "holes". (f) Segmented epithelial lining.
Here, instead of estimating the dimension $DBM$, the method uses the values of $\log V(r)$ within a pre-defined range of radii $r$ as the B-M descriptors. When an image is homogeneous, the procedure results in a regular cloud of points with the plot of $\log V(r)$ vs. $\log r$ approaching a linear function. With heterogeneous images, at some scales the spheres start merging for various values of $r$ and the log-log plot shows a different pattern. This curve therefore is a representation of the image homogeneity at various scales (radii) and this makes it appealing for characterisation of grey-level image complexity [2].

The set of volumes $V(r)$ are obtained by summing the points pertaining to the union of spheres $B(p, r)$ centred at each point $p \in C$ with radius $r$, that is,

$$V(r) = \sum_{p \in E} I_{U_r}(p'),$$

where $I$ is the indicator function and

$$U_r = \bigcup_{p \in C} B(p, r).$$

In discrete spaces (such as digital images), $B(p, r)$ is the (finite) set of points at a distance at most $r$ from $p$. Therefore the number of points in $U$ can be efficiently computed by means of the Euclidean Distance Transform (EDT). In a three-dimensional space, each point $p' \in E$ is transformed into:

$$\text{EDT}(p') = \min_{p \in C} \text{dist}(p, p'),$$

where $\text{dist}(p, p')$ is the Euclidean distance between the vectors. Several methods exist to compute the EDT efficiently [5].

To compute the B-M descriptors $D$ from the EDT, all the possible values of the transform within the region of interest in $E$ are increasingly sorted into a vector $o(k)$ and the descriptors are given by the logarithm of the cumulated sum of points $o'$ such that $\text{EDT}(p') \leq o(k)$:

$$D(k) = \log \# \{ p' \in E \text{ and } \text{EDT}(p') \leq o(k) \}. $$

---

**Fig. 3.** Steps involved in the estimation of the Bouligand-Minkowski fractal dimension of a grey-level image. (a) Greyscale image. (b) Cloud dilation by growing radius. (c) Plot of $\log V(r)$ vs. $\log r$. The dimension is obtained by replacing the limit in Eq. (3) by the slope of a straight line fit to the curve. Here, the cloud points were sub-sampled by 1/10th of the original for display clarity purposes. In the proposed method the entire image is analysed, (a) Original image, (b) Morphological dilation, (c) Bouligand-Minkowski curve.
where \( \# \) expresses for set cardinality. The variable \( k \) indexes all the non-negative values for the EDT within the considered space, so that e.g., if we take \( r \leq 2 \) there are four possible values for \( k \) (1, \( \sqrt{2} \), \( \sqrt{3} \) and 2), and therefore four B-M descriptors are produced. We use \( r \leq 8 \), to provide 64 descriptors, although only the first 50 are used in our experiments, as no significant classification gains were achieved beyond that number.

### 3.5. Assessment

As described in Section 2, the database contains 150 images collected from 30 independent cases (5 images per case).

The B-M descriptors were used for image classification as input parameters of a Linear Discriminant Analysis (LDA) supervised classifier [4]. This method was chosen due to its ability of decorrelating features among the samples. For small values of \( k \) the difference between the growth of \( \Omega(k) \) is subtle, so simpler classifiers (such as k-nearest-neighbours [4]) for instance, might not be capable of identifying patterns useful for the classification.

We used two approaches to define training and testing sets and assess the classification performance, to know, a random 10-fold to evaluate the performance of the classifier when the number of descriptors is varied and a nested 10-10-fold employed to obtain the best classifier setup for the cyst classification. In both cases, the B-M descriptors were subjected to principal component analysis [4] with the aim of decorrelating the image features, as the cumulative way that B-M descriptors are obtained tends to result in some degree of inter-descriptors dependence. If any random fold did not contain at least one sample from each group, that classification round was discarded and the procedure was repeated until satisfying such requirement.

In the first approach, the set of samples is randomly divided into 10 equal sub-sets and in each round of classification one sub-set is used for testing and all the other sub-sets are used for training. The success rate is given by the ratio of the number of images correctly classified and the total number of images in the database. The procedure is repeated 10 times (each time with a different arrangement of the random sub-sets) and the final success rate is given by the total average.

The second approach differs from the first one in that instead of running over all possible numbers of features, an ideal number is chosen by a second 10-fold procedure performed only over the testing set. In this way, the classification involves two nested 10-folds, an external one responsible for the classification and an internal procedure where the optimum number of descriptors is determined.

### 4. Results

The results are reported below by groupings (k: sporadic OKCs, r: radicular cysts and s: syndromic OKCs) having diagnostic relevance (k×r×s, ks×r and k×s).

All the results are expressed in terms of classification accuracy \( \text{Acc} \), which is defined by the following expression:
Acc = \frac{TP}{TP + TN}

where \( TP \) is the number of true positives, i.e., the number of samples/cases assigned to a particular group and actually pertaining to that group and \( TN \) is the number of true negatives, i.e., the number of samples/cases not assigned to a particular group and that actually do not come from that group. In this way, \( TP + TN \) is the total number of samples and therefore \( Acc \) can also be interpreted as the ratio of images correctly classified with respect to the total number of samples in the database.

The first test evaluates the success rate of the classification (percentage of images correctly classified) assessed for a number of descriptors ranging between 1 and 50. In addition to the original unprocessed images, we also applied three different pre-processing operations: histogram equalization, Gaussian filter blurring (kernel of radius=5 pixels and standard deviation=1) and sub-sampling size reduction by half, to test the robustness of the classification subject to possible variations in image quality. Histogram equalization enhances the contrast of images, balancing excessively dark or clear images. Blurring and subsampling simulate practical situations where the images are collected with sub-optimal microscope focusing and at lower resolution, respectively. The objective of verifying the performance in such conditions is only to assess how much and in which case such operations affect the final result. For the purpose of discussion and comparison only the results for the original images were considered. Fig. 4 shows the plot of success rates for the discrimination \( k \times r \times s \), Fig. 5 for \( k \times s \times r \) and Fig. 6 for \( k \times s \).

At this point, we also tested the possibility of using only the \( H \) component of the image (Hematoxylin) as that dye is mostly uptaken by cell nuclei. The \( H \) component was extracted from the \( H&E \) image using a procedure called colour deconvolution. This involves an orthonormal transformation of the Red-Green-Blue pixel components, once the individual dye vectors (in this case \( H \) and \( E \) separately) have been determined. The procedure enables determining the contribution of each individual dye used in the staining. Further details can be found in [18]. Tables 5–8 list the results obtained for the same types of pre-processed images. In general the average performance is similar to that achieved using the original image.

Table 9 shows the success rates analysed case-wise (the database...
contains 30 individual cases, with 5 images from each). For each case, the percentage of samples correctly classified as k, r or s is taken into account. The classifier is said to identify the lesion type if more than 50% of a case are correctly classified. In general, all the success rates are now larger than in the sample-wise classification, even because the classification problem is easier in fact, given the reduced number of elements to be discriminated (30 against 150 in the sample-wise test).

Tables 10–12 show the confusion matrices for the case-wise classification of k×r×s, ks×r and k×s, respectively.

Finally, Table 13 shows the contingency tables comparing the classification performance of the approach proposed in this work to that previously achieved in [11]. The same table shows the respective p-values of a McNemar test [3] for rejecting the null hypothesis that both classifiers are equivalent. In all cases it was demonstrated that the difference in the classification performance is statistically significant with an error within the confidence interval.

### Table 1
Classification accuracy in the discrimination between all the types of cysts here considered (k×r×s), between radicular (r) and OKC (ks×r) and between both types of OKCs (k×s) (k: sporadic and s: syndromic OKs).

|            | k×r×s | ks×r | k×s |
|------------|-------|------|-----|
| Original   | 72 ± 2% | 98 ± 1% | 68 ± 3% |
| Equalised  | 74 ± 2% | 95 ± 1% | 73 ± 2% |
| Blurred    | 70 ± 1% | 94 ± 2% | 72 ± 2% |
| Sub-sampled| 69 ± 2% | 92 ± 2% | 62 ± 4% |

### Table 2
Confusion matrices obtained from the classification of cysts k×r×s.

|            | k  | r  | s  |
|------------|----|----|----|
| k          | 0.66 | 0.00 | 0.34 |
| r          | 0.04 | 0.88 | 0.08 |
| s          | 0.36 | 0.00 | 0.64 |
| Normal     |    |    |    |
| k          | 0.73 | 0.03 | 0.34 |
| r          | 0.10 | 0.86 | 0.04 |
| s          | 0.33 | 0.05 | 0.62 |
| Equalised  |    |    |    |
| k          | 0.63 | 0.06 | 0.31 |
| r          | 0.06 | 0.82 | 0.11 |
| s          | 0.29 | 0.02 | 0.69 |
| Blurred    |    |    |    |
| k          | 0.64 | 0.06 | 0.30 |
| r          | 0.05 | 0.84 | 0.11 |
| s          | 0.36 | 0.04 | 0.60 |
| Sub-sampled|    |    |    |
5. Discussion

Odontogenic cysts arise from proliferation of epithelial cell rests commonly found in the jaw bones. These cells were originally part of the tooth-forming apparatus and subsequently remain inactive through life, but can occasionally proliferate spontaneously or in response to inflammatory processes occurring nearby.

### Table 3
Confusion matrices obtained from the classification of cysts $k \times r$

|     | $k$ | $r$ |
|-----|-----|-----|
| $k$ | 1.00| 0.00|
| $r$ | 0.07| 0.93|
| Normal | | |

### Table 4
Confusion matrices obtained from the classification of cysts $k \times c \times s$

|     | $k$ | $s$ |
|-----|-----|-----|
| $k$ | 0.69| 0.31|
| $s$ | 0.34| 0.66|
| Normal | | |

### Table 5
Classification accuracy in the discrimination between all the types of cysts using only the H (hematoxylin) component of the microscopy image.

|     | $k \times r \times s$ | $k \times r$ | $k \times s$ |
|-----|-----------------------|--------------|--------------|
| Original | 73 ± 3% | 97 ± 1% | 60 ± 3% |
| Equalised | 78 ± 2% | 95 ± 1% | 75 ± 2% |
| Blurred | 65 ± 2% | 92 ± 1% | 64 ± 4% |
| Sub-sampled | 71 ± 2% | 93 ± 1% | 60 ± 4% |

### Table 6
Confusion matrices obtained from the classification of cysts $k \times r \times s$ using the H component.

|     | $k$ | $r$ | $s$ |
|-----|-----|-----|-----|
| $k$ | 0.72| 0.02| 0.26|
| $r$ | 0.06| 0.91| 0.03|
| $s$ | 0.43| 0.01| 0.56|
| Normal | | | |

### Table 7
Confusion matrices obtained from the classification of cysts $k \times r$ using the H component.

|     | $k$ | $r$ |
|-----|-----|-----|
| $k$ | 0.98| 0.02|
| $r$ | 0.07| 0.93|
| Normal | | |

Traditional histopathology is based on the interpretation of the morphological appearance of cells and tissues, and while this is the current diagnostic gold standard, the assessment of morphological features through visual observation has unavoidable elements of subjectivity. For instance, cell and tissues shapes and sizes and staining patterns are difficult to assess consistently across observers, making the reproducibility of diagnostic procedures less robust than desired. The arrival of digital imaging and digital pathology, however, has opened the opportunity to treat histopathological digital images as numerical data sets which can be processed by algorithms for image reconstruction, enhancement and classification. The rationale for the development of automated diagnosis therefore aims to more accurate and reproducible diagnosis and to analyse more tissue samples than is
Currently possible.

Within this context, automated diagnosis can be approached from two different perspectives. One is the interpretation of digital images in terms of histologically-relevant concepts. This type of intelligent imaging requires algorithms that associate pixel values into discrete regions with special histological meaning (such as cells and nuclei, see for example [17,1] to then compare and measure those against known spatial and temporal models of normality and disease [12]. Such model-interpreting computations aim to approach algorithmically the tasks pathologists perform. An alternative is a data-driven, ‘machine vision’ approach based on the analysis of the image data itself, without need of identifying histologically-relevant structures.

Both approaches have their benefits. Ideally, intelligent imaging might provide the means to enable histopathological generalisations to be made. The data-analysis approach, on the other hand, can be simpler to implement by avoiding intermediate, higher level interpretation of the image data. However the solutions encountered might not be specific to the biological problem and prevent any generalisation (in pathology terms) of the solutions achieved.

Here we concentrated on an ‘image data’ analysis approach to the problem of classifying sporadic, syndromic OKC and radicular cyst images solely based on the staining patterns of the epithelial lining of

| Table 8 |
| --- |
| Confusion matrices obtained from the classification of cysts k×s using the H component. |
| | k | s |
| k | 0.66 | 0.34 |
| s | 0.49 | 0.51 |
| Normal |
| k | 0.77 | 0.23 |
| s | 0.28 | 0.72 |
| Equalised |
| k | 0.68 | 0.32 |
| s | 0.48 | 0.52 |
| Blurred |
| k | 0.65 | 0.35 |
| s | 0.48 | 0.52 |
| Sub-sampled |

| Table 9 |
| --- |
| Classification accuracy in the case-wise discrimination between all the types of cysts. |
| k×r×s | k×r | k×s |
| Original | 76 ± 6% | 100 ± 1% | 71 ± 4% |
| Equalised | 83 ± 4% | 97 ± 1% | 75 ± 5% |
| Blurred | 79 ± 5% | 99 ± 2% | 80 ± 4% |
| Sub-sampled | 79 ± 5% | 98 ± 3% | 67 ± 8% |

| Table 10 |
| --- |
| Confusion matrices obtained from the case-wise classification of cysts k×r×s. |
| k | r | s |
| k | 0.68 | 0.00 | 0.32 |
| r | 0.00 | 0.98 | 0.02 |
| s | 0.34 | 0.00 | 0.66 |
| Normal |
| k | 0.81 | 0.01 | 0.18 |
| r | 0.01 | 0.98 | 0.01 |
| s | 0.31 | 0.00 | 0.69 |
| Equalised |
| k | 0.74 | 0.03 | 0.23 |
| r | 0.00 | 0.99 | 0.01 |
| s | 0.35 | 0.00 | 0.65 |
| Blurred |
| k | 0.72 | 0.02 | 0.25 |
| r | 0.00 | 0.97 | 0.03 |
| s | 0.30 | 0.01 | 0.69 |
| Sub-sampled |

| Table 11 |
| --- |
| Confusion matrices obtained from the case-wise classification of cysts k×r. |
| k | r |
| ks | k | s |
| Normal |
| k | 0.99 | 0.01 |
| r | 0.00 | 1.00 |
| Equalised |
| k | 0.99 | 0.01 |
| r | 0.07 | 0.93 |
| Blurred |
| k | 0.99 | 0.01 |
| r | 0.00 | 1.00 |
| Sub-sampled |

| Table 12 |
| --- |
| Confusion matrices obtained from the case-wise classification of cysts k×s. |
| k | s |
| Normal |
| k | 0.75 | 0.25 |
| s | 0.33 | 0.65 |
| Equalised |
| k | 0.78 | 0.22 |
| s | 0.30 | 0.70 |
| Blurred |
| k | 0.81 | 0.19 |
| s | 0.21 | 0.79 |
| Sub-sampled |

|   |   |   |
|---|---|---|
| k | 0.69 | 0.31 |
| s | 0.36 | 0.64 |

regions with special histological meaning (such as cells and nuclei, see for example [17,1] to then compare and measure those against known spatial and temporal models of normality and disease [12]. Such model-interpreting computations aim to approach algorithmically the tasks pathologists perform. An alternative is a data-driven, ‘machine vision’ approach based on the analysis of the image data itself, without need of identifying histologically-relevant structures.

Both approaches have their benefits. Ideally, intelligent imaging might provide the means to enable histopathological generalisations to be made. The data-analysis approach, on the other hand, can be simpler to implement by avoiding intermediate, higher level interpretation of the image data. However the solutions encountered might not be specific to the biological problem and prevent any generalisation (in pathology terms) of the solutions achieved.

Here we concentrated on an ‘image data’ analysis approach to the problem of classifying sporadic, syndromic OKC and radicular cyst images solely based on the staining patterns of the epithelial lining of
these lesions. This is a difficult task that, to date, has not been resolved satisfactorily by neither expert observers nor algorithmic approaches. Our analysis based on the B-M descriptors achieved better results classifying OKCs vs. radicular cysts than those previously reported in [11]/(98% (Table 1) against around 95%), or in [9]. When discriminating between the three types of cysts, our approach achieved a better success rate (72%, Table 1), compared to 66% in [11]. The previous reported success rate in the discrimination of solitary from syndrome OKCs was also outperformed by the approach here described (68% (Table 1) vs. 60%).

We have used a nested cross-validation procedure which is a more challenging than the standard cross-validation, ensuring a fair feature selection stage with no a priori knowledge about images in the training set and using a similar number of images for training and testing sets. The patient-wise analysis, had an average success rate of 76% for k×r×s (average between the two majority success rates in Table 9), 100% for the patient-wise analysis, had an average success rate of 76% for k×r×s set and using a similar number of images for training and testing sets.

We applied the state-of-the-art Bouligand-Minkowski descriptors to classify the epithelial lining of the sub-types of OKCs and discriminate between OKCs and radicular cysts. Unlike previous attempts to resolve this problem, we used a statistical approach to analyse and quantify the complex patterns of pixel intensities in the image. This outperformed the previous sample classification results and demonstrates that in addition to histologically-relevant structures (such as those in [11]) the distribution pixel intensities also carry useful diagnostic importance for image characterisation. This highlights the value of texture-based image analysis to describe complex histological images. The results achieved in the classification of cyst types are of high relevance to histopathology as they can form the basis for automatic diagnosis with known levels of accuracy, making the diagnostic process more reproducible and less time-consuming. This also opens the possibility to automatic pre-screening of larger numbers of histological samples from the same case than it is currently possible, therefore making diagnosis more robust.

6. Conclusion

We applied the state-of-the-art Bouligand-Minkowski descriptors to classify the epithelial lining of the sub-types of OKCs and discriminate between OKCs and radicular cysts. Unlike previous attempts to resolve this problem, we used a statistical approach to analyse and quantify the complex patterns of pixel intensities in the image. This outperformed the previous sample classification results and demonstrates that in addition to histologically-relevant structures (such as those in [11]) the distribution pixel intensities also carry useful diagnostic importance for image characterisation. This highlights the value of texture-based image analysis to describe complex histological images.

The results achieved in the classification of cyst types are of high relevance to histopathology as they can form the basis for automatic diagnosis with known levels of accuracy, making the diagnostic process more reproducible and less time-consuming. This also opens the possibility to automatic pre-screening of larger numbers of histological samples from the same case than it is currently possible, therefore making diagnosis more robust.

### Table 13

|        | BM incorrect | BM correct |
|--------|--------------|------------|
| Ref [11] incorrect | 21           | 30         |
| Ref [11] correct    | 16           | 83         |
| p-value=0.0390      |              |            |

|        | BM incorrect | BM correct |
|--------|--------------|------------|
| Ref [11] incorrect | 1            | 7          |
| Ref [11] correct    | 1             | 141        |
| p-value=0.0339      |              |            |

|        | BM incorrect | BM correct |
|--------|--------------|------------|
| Ref [11] incorrect | 16           | 30         |
| Ref [11] correct    | 15           | 44         |
| p-value=0.0253      |              |            |

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