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Detection of lobular structures in normal breast tissue

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Abstract

Background: Ongoing research into inflammatory conditions raises an increasing need to evaluate immune cells in histological sections in biologically relevant regions of interest (ROIs). Herein, we compare different approaches to automatically detect lobular structures in human normal breast tissue in digitized whole slide images (WSIs). This automation is required to perform objective and consistent quantitative studies on large data sets.

Methods: In normal breast tissue from nine healthy patients immunohistochemically stained for different markers, we evaluated and compared three different image analysis methods to automatically detect lobular structures in WSIs: (1) a bottom–up approach using the cell–based data for subsequent tissue level classification, (2) a top–down method starting with texture classification at tissue level analysis of cell densities in specific ROIs, and (3) a direct texture classification using deep learning technology.

Results: All three methods result in comparable overall quality allowing automated detection of lobular structures with minor advantage in sensitivity (approach 3), specificity (approach 2), or processing time (approach 1). Combining the outputs of the approaches further improved the precision.

Conclusions: Different approaches of automated ROI detection are feasible and should be selected according to the individual needs of biomarker research. Additionally, detected ROIs could be used as a basis for quantification of immune infiltration in lobular structures.

Keywords: Whole Slide Image, Digital Histopathology, Normal Breast Lobule, Image Analysis, Convolutional Neural Network

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1. Introduction

Lobular structures are the functional units of the resting mammalian breast that further differentiate into milk–producing glands during lactation. The normal anatomical structures are important because breast cancer and premalignant lesions originate in these epithelial structures and there is evidence that the transition between ductal and lobular structures may be particularly susceptible to oncogenic events [1]. In addition to studies on the origins of cancer, the detection of lobules is also relevant for an inflammatory condition referred to as lymphocytic lobulitis (LLO), which has been observed in the adjacent tissue around breast cancer and in prophylactically removed breast tissue without any evidence for cancer in BRCA1/2 mutation carriers [2, 3]. This phenomenon is not yet well understood and deciphering its possible link with hereditary breast cancer may lead to better disease understanding, new prognostic indicators, or novel treatment options. In order to perform an objective, repeatable, and statistically reliable quantitative study of LLO on large data sets, the ability to automatically detect lobules in histological slides is necessary.

Nowadays, such slides can be routinely digitized; the resulting whole slide images (WSIs) can be processed by automated image analysis techniques with the aim to detect lobular structures and to quantify cell numbers [4, 5]. Many works are based on detecting and automatically counting cells for cancer diagnosis, grading, and prognosis [6, 7, 8]. However, to give more insight to the pathologist, it is necessary to evaluate immune cells beyond estimation of their density, for example by object–based recognition of spatial patterns and interactions at high resolution [9]. Thus, our objective is to pave the way for identifying and classifying cells in those areas in image that are most relevant, like lobules in breast cancer and LLO, finally enabling methods to characterize the spatial distribution of different subtypes of immune cells in relation to these to these larger image objects.

In general, lobules are composed of dense areas of epithelial cells in tubular structures, normally with a clear contrast to the lobular stroma. However, lobules can be very different in size, shape, and texture depending on their functional stage (e.g., phase of the menstrual cycle), the degree of immune cell infiltration, and also appear differently according to the used staining method (see Figure 1).

Cancer growth adds to this complexity of tissue structures and makes manual or automated lobule detection even more challenging. As a preparatory work before delving into the structural complexity of cancer–affected tissue, we focus on normal breast tissue of healthy patients in order to evaluate and compare three different image analysis methods to automatically detect lobular structures in immunohistochemically (IHC) stained WSIs. Even without solving the cancer–related challenges, the work addresses an important demand because the evaluation of LLO will lead to new biomarker patterns with diagnostic and prognostic value. From a technological point of view, the task of defining regions of interest (ROI) for further analysis tackles a problem that occurs frequently in medical image analysis: The limited availability of experts...
to perform large-scale manual annotation due to restricted resources of trained pathologists remains an important bottleneck for progress. In the context of LLO, this is evident because the statistical base for experimental evidence has been limited by the available time for pathologists who have to manually select every lobule and therefore could so far only annotate small data sets \[10, 11\]. Thus, overcoming the limitations of manual annotation of WSIs by automation is highly desirable. This work builds on previous work \[12\], where detection of normal lobules in the vicinity of breast cancer was optimized for the purpose of analyzing nuclear expression of estrogen receptor (ER) or progesterone receptor (PR). Grote et al. detected lobules on several segmentation layers using textural, geometric, and relational features, as well as solid tumor using textural features. Other tissue classification techniques have supported the study of pathologies like odontogenic cysts \[13\] and various cancers \[14, 15\]. The identification of general biological structures has received comparatively little attention, although graph-based approaches exist for unsupervised top-down tissue categorization \[16\] and bottom-up biological object identification \[17\].

After presenting our data set and evaluation criterion, we will describe three methods that were developed in the context of lymphocytic lobulitis: (1) a
Bottom–Up Method (MBU) starting with cell detection and ending with tissue classification, (2) a Top–Down Method (MTD) starting with texture classification and ending with cell density characterization, and (3) a direct texture classification Method using Deep Learning technology (MDL). We conclude by comparing their strengths and weaknesses using two characteristically different stainings and by assessing the feasibility of combining the methods.

2. Materials and Methods

We collected tissue samples from a cohort including nine healthy women who underwent reduction mammoplasty due to hypermasty [18]. The paraffin–embedded samples were cut into 3 µm thick sections and stained for the nuclear marker ER and the immune cell marker cluster of differentiation 8 (CD8) that is expressed in the cell cytoplasm and on the outer cell membranes of cytotoxic T lymphocytes, using an automated staining instrument (Ventana Benchmark Ultra). As ER is often expressed in breast cancer, it was chosen as example for a routinely used breast cancer marker important for treatment decisions [19, 20] and CD8 for its relevance in oncoimmunology [21]. Whereas ER+ cells are epithelial cells of lobular structures, CD8+ cytotoxic T lymphocytes may spread close to epithelial compartments, over the full lobular stroma, or even in non–lobular stroma. WSIs were acquired by Aperio AT2 scanner at 40X magnification, scanned images have a resolution of 0.253 µm/pixel. For each WSI, a pathologist (FF) performed annotations of the full epithelial compartment (including lobular and ductal regions) and more detailed annotations to distinguish between lobular and non–lobular regions. This was performed by drawing outlines over the digital images using the software tool Aperio ImageScope. Table 1 summarizes the number of annotated lobules for each case and each staining.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>ER Size</th>
<th>CD8 Size</th>
<th>ER Lobules</th>
<th>CD8 Lobules</th>
</tr>
</thead>
<tbody>
<tr>
<td>NB02</td>
<td>25</td>
<td>20.6 x 20.6</td>
<td>21.6 x 21.2</td>
<td>125</td>
<td>120</td>
</tr>
<tr>
<td>NB05</td>
<td>29</td>
<td>21.1 x 20.1</td>
<td>21.6 x 21.6</td>
<td>32</td>
<td>25</td>
</tr>
<tr>
<td>NB11</td>
<td>19</td>
<td>15.6 x 15.9</td>
<td>14.1 x 15.8</td>
<td>55</td>
<td>35</td>
</tr>
<tr>
<td>NB12</td>
<td>22</td>
<td>17.6 x 15.7</td>
<td>17.1 x 14.2</td>
<td>12</td>
<td>11</td>
</tr>
<tr>
<td>NB16</td>
<td>27</td>
<td>16.1 x 15.5</td>
<td>15.1 x 17.9</td>
<td>54</td>
<td>63</td>
</tr>
<tr>
<td>NB20</td>
<td>27</td>
<td>16.6 x 15.1</td>
<td>17.6 x 15.0</td>
<td>80</td>
<td>76</td>
</tr>
<tr>
<td>NB25</td>
<td>30</td>
<td>20.1 x 16.6</td>
<td>13.6 x 19.9</td>
<td>49</td>
<td>34</td>
</tr>
<tr>
<td>NB28</td>
<td>21</td>
<td>24.6 x 20.9</td>
<td>28.7 x 22.6</td>
<td>19</td>
<td>24</td>
</tr>
<tr>
<td>NB34</td>
<td>28</td>
<td>14.1 x 19.0</td>
<td>16.1 x 19.7</td>
<td>226</td>
<td>238</td>
</tr>
</tbody>
</table>

Table 1: Overview over data set: Size of image (width x height mm) and number of lobules in ground truth. The slight variation in numbers of lobules could be either due to variable composition of the sections at different levels of the paraffin block or due to slightly different annotation of components of lobules, e.g. merging two elements of the glands to one “lobule”.

To evaluate the methods, we calculated a global, pixel–based F1 score (equiv-
alent to Dice similarity \[22\]):

\[
F_1 = \frac{2TP}{2TP + FP + FN}
\]

where \(TP\) is the number of true positives, \(FP\) the number of false positives, and \(FN\) the number of false negatives. We additionally measured the true positive percentages inside the detected and actual lobular objects in order to obtain a more object–based quality estimation.

To the best of our knowledge, there is no equivalent published work that we can use for comparison, but the elementary concepts and tools that we use are well documented \[23\].

2.1. Method 1: Bottom–Up (MBU)

As a starting point, we developed a workflow (see Figure 3) that applies well–established algorithms using Definiens Developer XD 64 2.4 based on the assumption that cell clusters and therefore highest deviations in RGB occur in lobular structures. In the following, we refer to this workflow as MBU. Due to large size of WSIs (up to 100,000 × 100,000 pixels), we used images down–sampled to a resolution of 8 \(\mu\)m/pixel (\(\sim 3.2%\) of the original WSI). Based on a multi–threshold segmentation on a gray layer, tissue regions (remaining: background) are detected, in which afterwards each pixel is classified as lobular or non–lobular area. For this, we generated a color–distribution–range image (CDR) that represents the averaged RGB range in a small part \((s \times s)\) pixels of the down–sampled image cut by a lower and an upper threshold \(t_l\) and \(t_u\); i.e., each image is split into tiles \(s \times s\) pixel large having three layers red, green, blue each in range \([0, 255]\), in which all new pixel values \(x\) are set as defined in equation (2),

\[
x = \begin{cases} 
0 & \text{if } \frac{1}{3}(r_R + r_G + r_B) < t_l \\
255 & \text{if } \frac{1}{3}(r_R + r_G + r_B) > t_u \\
\frac{1}{3}(r_R + r_G + r_B) & \text{otherwise}
\end{cases}
\]

where \(r_X = \max_X - \min_X\) represents the range of layer \(X\) in the corresponding small tile. Thus, a homogeneous area with similar colors in all pixels is set to black, whereas a texture–rich area with large color deviations in these tiny tiles is set to white. Lobular structures in IHC images have a high contrast between epithelial cells in dark blue and very bright lobular stroma. The remaining image is composed of bright stroma with less contrasts and almost white fatty tissue, which is mostly classified as background. Since we do not use a sliding window approach for simplicity, we afterwards split the image into tiles \(l \times l\) pixel large to redefine the pixel values depending on their neighborhood (defined in Figure 2).

In tiles with less than \(t_{wl}\) white pixels or in tiles where more than 50\% (parameter \(t_b\)) of the pixels are black and less than 25\% (parameter \(t_{wu}\)) are white, each pixel value \(x\) is set to \(\frac{5}{2}\) (darker). For the remaining tiles, the
Figure 2: Neighborhood of pixel \((i, j)\) (red: direct neighbors; blue: further neighbors) used in MBU.

decision is separately done for each individual pixel based on its eight direct neighbors and 16 further neighbors, for details see equation (3),

\[
x_{\text{new}} = \begin{cases} 
\frac{x}{2} & \text{if } (|B| > t_b \cdot l^2 \land |W| \geq t_{wu} \cdot l^2 \land \neg N_8) \\
\frac{x + \frac{W}{2}}{2} & \text{if } (|B| \leq t_b \cdot l^2 \land |W| > t_{wu} \cdot l^2 \land (N_8 \lor N_{16}) \lor (|B| > t_b \cdot l^2 \land |W| \geq t_{wu} \cdot l^2 \land N_8)) \\
x & \text{otherwise}
\end{cases} 
\tag{3}
\]

where \(|B|\) and \(|W|\) are the number of black and white pixels and \(N_i\) is true when at least one of \(i\) neighbors is 255 (obviously, we set values larger than 255 to 255). All used parameter values are listed in Table 2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Used value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(s)</td>
<td>size of small tiles</td>
<td>5 \text{ i.e., } 1600 \mu m^2</td>
</tr>
<tr>
<td>(l)</td>
<td>size of large tiles</td>
<td>50 \text{ i.e., } 160,000 \mu m^2</td>
</tr>
<tr>
<td>(t_l)</td>
<td>lower threshold to cut CDR</td>
<td>25</td>
</tr>
<tr>
<td>(t_u)</td>
<td>upper threshold to cut CDR</td>
<td>150</td>
</tr>
<tr>
<td>(t_{wl})</td>
<td>lower threshold for white pixels</td>
<td>0.0212</td>
</tr>
<tr>
<td>(t_{wm})</td>
<td>middle threshold of white pixels</td>
<td>0.1248</td>
</tr>
<tr>
<td>(t_{wu})</td>
<td>upper threshold of white pixels</td>
<td>0.2500</td>
</tr>
<tr>
<td>(t_b)</td>
<td>threshold of black pixels</td>
<td>0.5000</td>
</tr>
</tbody>
</table>

Table 2: Parameters used in MBU.

The cut CDR was included as additional layer into our Definiens rule set and
averaged with a layer which represents clusters of well–shaped nuclei. For this combination, the two layers are normalized to the same range to allow an equal weighting of both. If a pixel in this combined layer is non–dark, it is classified as lobular. The nuclei layer is generated by a k–means clustering to find areas with high epithelial cell densities using a robust nucleus detection by Definiens\cite{24} and excluding stromal and immune cells based on size and shape. Due to small size of cells, the clustering is done on a higher resolution of 1.14 \( \mu m \)/pixel (~ 22.1\% of the original WSI). Here, we assume that the nuclei of epithelial cells in lobular structures are usually round and 20–50 pixels large (i.e., a diameter of about 5–10 \( \mu m \)).

![Diagram](image_url)

Figure 3: Overview of MBU processing a single WSI. The classification into lobular and non–lobular structures is based on a combination of nuclei clusters in a medium resolution (1.14 \( \mu m \)/pixel ~ 22.1\% of original WSI) and a cut color–distribution–range (CDR) image, which represents the range of RGB in tiles of \( s \times s \) pixels, in a low resolution (8 \( \mu m \)/pixel ~ 3.2\% of original WSI).

2.2. Method 2: Top–Down (MTD)

The second approach (Figure 4), in the following called MTD, has two basic steps where first candidate regions are identified in low resolution and then refined in medium resolution. In the first step, lobule candidate regions are detected from a texture image in low resolution (5.06 \( \mu m \)/pixel or 5\% of the original image). The texture image is produced by a texture–based classification using local binary patterns and local variance as features, as described in \cite{15}. A
support vector machine (SVM) is used for the classification. For training of the SVM model, small image subsets (from images not contained in our test set but belonging to the same series) containing only lobular tissue or only other tissue are used. The output of the classification is a gray value image showing the probability of lobular tissue. This image is thresholded to obtain the candidate regions with a threshold (60) chosen so that most of the lobules in the images from which the training subsets were taken are inside the candidate regions.

The second step, the refinement of lobule candidates, is done using algorithms from Definiens Developer XD 64 2.4; it is based on nuclear density, which is higher in lobular areas than in the surrounding tissue. In medium resolution (0.84 μm/pixel or 30%), the image in the candidate regions is filtered using a Laplacian of Gaussian (LoG) filter whose parameters (9 × 9 kernel, σ = 1.768) are set to give maximum response for blobs of the average size of epithelial nuclei. The LoG image is thresholded using a fixed threshold to select an initial set of nuclei. The detected nuclei are filtered according to elliptic fit (≥ 0.8) so that only round nuclei remain. This nuclei detection does not attempt to detect all nuclei; the goal is rather to reliably detect a sufficient subset of epithelial nuclei for nuclear density estimation. The detected nuclei are used to produce a nuclear density channel using a sliding window approach (window size 41 × 41 pixel). Using the density channel, the candidate regions are shrunk iteratively until the remaining objects meet a minimum density criterion (average density ≥ 0.05). For further refinement, stromal and epithelial areas inside the lobules are separated by thresholding a smoothed gray value image. As threshold, the average of gray values in all refined lobule regions is used. Lobules that contain very little epithelial tissue (≤ 10%) after this step are eliminated. In a post-processing step, the borders of regions are smoothed by growing and shrinking, small regions are eliminated and small holes are filled.

2.3. Method 3: Deep Learning (MDL)

A machine learning algorithm describes how to identify patterns in existing data (learning) and uses this acquired knowledge to make predictions on new data[25]. A deep learning algorithm is a machine learning algorithm that can learn a hierarchical description of the data with multiple sublevels of nonlinear features[26].

In recent years, a type of deep learning architecture optimized for 2D data called convolutional neural networks (CNNs) have provided state-of-the-art results in various applications of machine learning–based image analysis, from general scene labeling[28] to cancer classification[29] and mitosis detection[30]. A CNN has the ability to learn a hierarchical description of visual patterns from a set of annotated examples, and then make accurate predictions for new visual inputs. By combining patch–based image analysis with CNNs, we are able to automatically detect lobular structures in normal breast histological images. The process is summarized in Figure 5.

For patch–based image analysis, we assume that it is possible to predict the class of a pixel by observing its neighboring pixels. In order to keep computation time low, a pathologist visually estimated the lowest level of detail at which he
could reliably distinguish epithelial tissue from other elements to be 4 μm/pixel. Likewise, we settled on a square neighborhood of size 128 μm which is enough to cover a full cross-section of a duct with its surroundings. It is then possible to generate, for each pixel, a “raw” or “featureless” description of the patch centered on this pixel. In other words, to each pixel we associate a square RGB patch described by $32 \times 32 \times 3 = 3072$ values. In images annotated by a pathologist, each patch can be considered positive or negative according to its central pixel: if a pixel is in an annotated lobular region, then its patch is positive, otherwise it is negative.

Our CNN is a function that can predict the class of a patch (positive or negative) based on its featureless description. This function has a fixed form defined in the “CIFAR10 Quick” example provided with the software Caffe[31]. The associated deep network architecture is designed to perform multiclass classification of small RGB images[32], which is conveniently similar to the task of patch classification. We can use it as a black box although its parameters (weights) need to be learned during supervised training. In order to estimate the prediction error of the method on unseen data for a given staining, we perform a 3-fold cross-validation using nine annotated images arbitrarily divided in three groups: $A=$\{NB02, NB05, NB11\}, $B=$\{NB12, NB16, NB20\}, and $C=$\{NB25, NB28, NB34\} (see Table[1] for general information on ground truth contents). In fold 0, the method is trained on $B \cup C$ and evaluated on $A$; in fold 1, the
method is trained on $A \cup C$ and evaluated on $B$; and in fold 2, the method is trained on $A \cup B$ and evaluated on $C$. For any fold, training consists of 20 iterations using error backpropagation with adaptive learning rate (ADAGRAD) on a dataset composed of 600,000 patches (50,000 for each class and each image). Given as input an image at 4 μm/pixel, this method will output a binary image where positive and negative pixels have different values. An additional post-processing step is used to denoise the result by removing small elements below a size threshold. In each fold, this threshold is learned by maximizing the macro-F1 score (average of F1 scores for each class) on the 6 training images.

2.4. Combination

We notice that the missclassified objects are not the same for each method. This suggests that they potentially offer complementary information about the patterns to be classified. As stated in [34, 35], if we have many different classifiers it is sensible to consider using them in a combination in the hope of increasing the overall accuracy.
In order to evaluate whether a combination of our methods will improve the results, we use the detection results of every single method in the form of binary masks to produce a combined result. In a pixel–based combination of masks (in low resolution, 5.06 µm/pixel or 5%), all pixels which belong to detected areas for at least two of the three methods are labeled as detected. This majority voting procedure is useful to eliminate false detections that are made by only one method.

3. Results and discussion

The overall performance of all methods for the stainings ER and CD8 is given in Table 3 as F1 score (measured pixel–based) for each of the 18 test images as well as the average for each staining. The highest F1 scores averaged over both stainings were obtained by MTD (0.59) and MDL (0.60).

<table>
<thead>
<tr>
<th>Staining</th>
<th>Method</th>
<th>NB02</th>
<th>NB05</th>
<th>NB11</th>
<th>NB12</th>
<th>NB16</th>
</tr>
</thead>
<tbody>
<tr>
<td>ER</td>
<td>MBU</td>
<td>.78</td>
<td>.17</td>
<td>.28</td>
<td>.42</td>
<td>.54</td>
</tr>
<tr>
<td></td>
<td>MTD</td>
<td>.73</td>
<td>.51</td>
<td>.54</td>
<td>.73</td>
<td>.76</td>
</tr>
<tr>
<td></td>
<td>MDL</td>
<td>.69</td>
<td>.67</td>
<td>.56</td>
<td>.24</td>
<td>.80</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>.81</td>
<td>.67</td>
<td>.62</td>
<td>.71</td>
<td>.80</td>
</tr>
<tr>
<td>CD8</td>
<td>MBU</td>
<td>.49</td>
<td>.39</td>
<td>.50</td>
<td>.59</td>
<td>.63</td>
</tr>
<tr>
<td></td>
<td>MTD</td>
<td>.65</td>
<td>.42</td>
<td>.58</td>
<td>.68</td>
<td>.57</td>
</tr>
<tr>
<td></td>
<td>MDL</td>
<td>.59</td>
<td>.51</td>
<td>.60</td>
<td>.50</td>
<td>.77</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>.74</td>
<td>.58</td>
<td>.66</td>
<td>.68</td>
<td>.74</td>
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</table>

<table>
<thead>
<tr>
<th>Staining</th>
<th>Method</th>
<th>NB20</th>
<th>NB25</th>
<th>NB28</th>
<th>NB34</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>ER</td>
<td>MBU</td>
<td>.38</td>
<td>.37</td>
<td>.66</td>
<td>.77</td>
<td>.40±.23</td>
</tr>
<tr>
<td></td>
<td>MTD</td>
<td>.71</td>
<td>.57</td>
<td>.61</td>
<td>.84</td>
<td>.67±.11</td>
</tr>
<tr>
<td></td>
<td>MDL</td>
<td>.62</td>
<td>.76</td>
<td>.59</td>
<td>.80</td>
<td>.64±.16</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>.73</td>
<td>.62</td>
<td>.70</td>
<td>.86</td>
<td>.72±.08</td>
</tr>
<tr>
<td>CD8</td>
<td>MBU</td>
<td>.20</td>
<td>.36</td>
<td>.33</td>
<td>.81</td>
<td>.42±.18</td>
</tr>
<tr>
<td></td>
<td>MTD</td>
<td>.45</td>
<td>.37</td>
<td>.20</td>
<td>.70</td>
<td>.51±.17</td>
</tr>
<tr>
<td></td>
<td>MDL</td>
<td>.66</td>
<td>.61</td>
<td>.04</td>
<td>.75</td>
<td>.56±.20</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>.68</td>
<td>.47</td>
<td>.41</td>
<td>.83</td>
<td>.64±.13</td>
</tr>
</tbody>
</table>

Table 3: Comparison of F1 scores for the different methods applied to ER and CD8 images (NBx).

3.1. Visual comparison

Figures 6 to 8 show lobule detection results and comparisons with ground truth for the image with the best F1 score for each method. For all three methods, the best results were achieved in one of the NB34 images (CD8 for MBU, ER for both other methods). This case contains many compact lobules with dense epithelial tissue, which makes them stand out against the surrounding tissue quite clearly. In addition, most of its lobules are positive for both ER and CD8
(i.e., many epithelial cells stained for ER are colored in brown and/or several brown colored CD8+ T lymphocytes are in close contact to epithelial cells) further increasing their differences to the background. Some characteristics of the different methods can be observed in these images. Correctly detected lobules from both MBU (Figure 6) and MTD (Figure 7) tend to have small stripes of false negative area around them, i.e., the detected area is usually smaller than the area that was manually outlined as ground truth. In the case of MBU, false negative areas within ground truth lobular objects occur due to the pixel–based classification instead of segmentation approaches using e.g., watershed or region growing. Intra–lobular stroma is usually not classified as lobular area due to missing contrasts by epithelial cells. This could be solved by additionally including some growing (and shrinking) procedures to reach smoother regions. The main advantage of MBU is that it is very fast (less than 2 min per image on a computer with a 3.6 GHz CPU, 32 GB RAM, using preprocessed nuclei detection) provided that the time–consuming nuclei detection has already been done, which is necessary anyway for cell population analysis. As mentioned before, we applied a robust method performing well for segmenting nuclei and classifying into different cell types by Definiens, which requires approximately 11 hours per image (single core usage in 3.4 GHz cluster environment with 384 GB RAM). In applications that do not quantify cells, it is also possible to use a rough and fast nuclei segmentation in MBU. For MTD, on average 38 min per image are needed on a computer with a 3.6 GHz CPU, 32 GB RAM. In contrast to MBU and MTD, there are practically no false negative regions in the result from MDL (Figure 8). The largest region labeled as false negative is actually a clearly stromal region inside a lobule, which is part of our ground truth.
Figure 6: Image with best result for MBU. For the MBU, the best result was in NB34 CD8, with an F1 score of 0.81. Left outlines of detected lobules in green, right evaluation result (green: true positive areas, blue: true negative areas, red: false positive areas, orange: false negative areas). Scale bar is 2 mm.
Figure 7: Image with best result for MTD. For the MTD, the best result was in NB34 ER, with an F1 score of 0.84. Left outlines of detected lobules in green, right evaluation result (green: true positive areas, blue: true negative areas, red: false positive areas, orange: false negative areas). Scale bar is 2 mm.
Figure 8: Image with best result for MDL. For MDL, the best result was in NB34 ER, with an F1 score of 0.80. Green lines show outlines of detected lobules. Left outlines of detected lobules in green, right evaluation result (green: true positive areas, blue: true negative areas, red: false positive areas, orange: false negative areas). Scale bar is 2 mm.

False positive areas also occur for all methods; these are often located in the small space between adjacent lobules. Correctly detected lobules from MDL almost always have a small stripe of false positive area around them, i.e., the detected area is larger than the ground truth area. Although false positive or false negative borders around correctly detected lobules are quite narrow, they contribute to the measurements shown in Table 3, so that the detection quality needed for applications, as judged by visual inspection, may be better than these numbers suggest.

Some typical examples of detection problems for the different methods can be seen in Figures 9 to 11. In Figure 9, a subset of NB02, CD8 contains a false positive area from MBU, where the tissue has an altered structure due to the IHC processing. Such regions often cause false positive detections. Due to calculation of CDR, the method strongly relies on textures in which the color range is quite high, which could be caused for example by tissue processing artifacts due to antigen retrieval (bright pixels directly neighbored to dark pixels).
Figure 9: Example for false positive detection in MBU. Left outlines of detected lobules in green, right evaluation result (green: true positive areas, blue: true negative areas, red: false positive areas, orange: false negative areas). Example subset from NB02 CD8. False positive detections are caused by changes in tissue structure due to IHC processing. Scale bar is 1 mm.

Figure 10: Example for false negative detection in MTD. Left outlines of detected lobules in green, right evaluation result (green: true positive areas, blue: true negative areas, red: false positive areas, orange: false negative areas). Example subset from NB16 CD8. Lobules are missed from detection because measured nuclear density was too low: epithelial tissue is sparse and nuclei have low contrast in image. Scale bar is 1 mm.
Figure 11: Example for false positive detection in MDL. Left outlines of detected lobules in green, right evaluation result (green: true positive areas, blue: true negative areas, red: false positive areas, orange: false negative areas). Example from NB28 CD8. Scale bar is 2 mm.

Figure 10 shows an example of false negative detection from MTD. This method relies on the measurement of nuclear density from detected nuclei. In the example, the lobule structure is relatively loose, meaning that epithelial tissue is sparse. In addition, the epithelial nuclei do not have large contrast to their surroundings, so that some are missed in the nuclei detection step. As a result, the nuclear density in the given area is below the required threshold, and the region was eliminated from the detection result. False positive detections (not shown) are often caused by vessels containing blood aggregations. These regions, though hardly stained, have a very similar texture to that of lobules, which leads to their detection.

Figure 11 shows an image with a large false positive area from MDL. This method depends heavily on the quality of the training set: classes should be balanced, examples should be correctly labeled and they should also be representative of the variety that can be found in unseen data. This last criterion is the hardest to achieve and image NB28 CD8 exhibits a stromal texture that is not found in the others. This could be an artifact resulting from the physical process used to prepare the slide. While it is an infrequent event, it does happen more than once in the rest of our images.

Another disadvantage of MDL is that it requires a specific hardware, but the resulting speed is satisfying. Using a Nvidia Quadro K4200 GPU with 4 GB of memory, training for one fold takes less than two hours and evaluation for an
image takes four minutes on average.

3.2. True positives in lobular structures

Histograms from per–lobule analysis of true positive percentages (Figure 12) show the characteristics of the different methods in a more quantified manner. In Figure 12 rows 1 and 3, histograms of percentage of ground truth area in detected lobules are shown, indicating how correct the detections are and how many false detections occurred. The first bin (0–5%) in the histograms of row 1 and 3 shows how many false positive objects appear in the results of the different methods. In all three methods, more false positive objects were detected in CD8 than in ER (2–3 times as many). Most false detections are observed in MBU, but on the other hand, the correctly found lobules are more cleanly detected (contain less non–lobular tissue) than in the other methods. For MTD, this picture is more heterogeneous, and in the results of MDL, most lobules contain some amount of non–lobular tissue (see also Figure 5).

Figure 12 rows 2 and 4 show histograms of percentage of detected area in ground truth lobules, indicating how complete the detection is for existing ground truth lobules and how many ground truth lobules are missed from the detection. The first bin in the histograms of row 2 and 4 shows how many lobules are missed. It can be seen that MDL has the least total losses. Most ground truth lobules are completely or almost completely detected, and there are very few where parts are missing. In the other two methods, the total losses are higher and the detection is more heterogeneous: there are many lobules where only parts of the ground truth lobules are detected. As described above, missing parts are often located along the borders of lobules. While MDL shows no substantial differences between ER and CD8 images, MBU and MTD show some differences: for MBU, ER has more missed detections than CD8, as well as relatively few complete detections. For MTD, CD8 has more missed detections and fewer complete detections.

In summary, MDL shows a high completeness in detection — hardly any lobule is missed — but most of the detected lobule objects also contain non–lobular tissue. MBU has a number of missed as well as false detections, but correctly detected lobules are relatively clean. MTD has more missed detections than MDL, but also more objects that contain mostly lobular tissue.
3.3. Combination

The top part of Figure 13 visualizes the effect of the combination of results. The F1 score results can be found in Table 3 (row “Combination” for both stainings). The results show that in two thirds of the cases the F1 score of the combination is higher than that of every single method. In the others, it is second best, often very close to the best single result. An exception is NB25 (both stainings), where the combination result is still second best but with a larger distance to the best single method.

This case has a large amount of ductal tissue, which is incorporated in the ground truth but for the most part only detected by MDL (Figure 13, bottom). If the target of an analysis are primarily the lobules but not the ducts, the combination seems to be the better choice also in this case. Further, using both the
combination results and the results of MDL might be helpful in distinguishing lobular and ductal tissue.

Figure 13: Combination of methods. White: labeled as lobular by all three methods; yellow: labeled as lobular by MBU and MTD; magenta: labeled as lobular by MBU and MDL; cyan: labeled as lobular by MTD and MDL; red: labeled as lobular by MBU only; green: labeled as lobular by MTD only; blue: labeled as lobular by MDL only. Gray lines: border lines of ground truth. Top: A subset of NB02 CD8 shows the effect of the combination. Detections by one method only (red, green, blue) often lie outside of the ground truth. Ground truth regions are almost completely covered, in this example often by all three methods or by combinations of MBU and MDL (magenta), sometimes by combinations of MTD and MDL (cyan) or combinations of MBU and MTD (yellow). Bottom: In NB25 ER, large ductal areas are only detected by MDL (blue). Scale bars are 2 mm.

4. Conclusion

In order to automate the detection of lobular structures in digital WSIs of normal breast tissue, we developed and compared several image analysis methods: two methods combining manually designed features and cell detection and a featureless machine learning–based image analysis.

We showed that despite a moderate variance for any given staining, each gives good results based on visual check of accuracy, and that the output could be used as basis for further image analysis (e.g., cell populations identification and quantification). We also showed that combining all the methods by pixel–
level majority voting improves precision and might help with subclassification of lobular tissue into lobules and ducts. We will leverage the knowledge gained in this study to tackle the issue of lobular structure detection in cancerous breast tissue and expand the concept of ROI–targeted immune cell detection in oncoimmunology.

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6. Author contributions

GA developed MDL, AG MTD, and NSS MBU. GA, AG, and NSS designed the evaluation and combination and prepared the manuscript. RS applied the nuclei detection used in MBU and established exchange of image analysis data. FF provided the expert annotations and pathological input. GF, CW, BN, and FF helped in study design and edited the manuscript. All authors read and approved the final manuscript.

7. Conflict of interest statement

RS is a full-time employee of Definiens AG. GA, NSS, BN, GF, FF, CW and AG did not report any conflict of interest.
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