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An automated prognosis system for estrogen hormone status assessment in breast cancer tissue samples

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Abstract: Estrogen receptor (ER) status evaluation is a widely applied method in the prognosis of breast cancer. However, testing for the existence of the ER biomarker in a patient’s tumor sample mainly depends on the subjective decisions of the doctors. The aim of this paper is to introduce the usage of a machine learning tool, functional trees (FTs), to attain an ER prognosis of the disease via an objective decision model. For this aim, 27 image files, each of which came from a biopsy sample of an invasive ductal carcinoma patient, were scanned and captured by a light microscope. From these images, 5150 nuclei were segmented with image processing methods. Several attributes, including statistical, wavelet, cooccurrence matrix, and Laws’ texture features, were calculated inside the border area of each nucleus. A FT was trained over the feature dataset using a 10-fold cross-validation and then the obtained model was tested on a separate dataset. The assessment results of the model were compared with those of 2 experts. Consequently, the weighted kappa coefficient indicated a very good agreement (κ = 0.899 and κ = 0.927, P < 0.001) and the Spearman’s rank order correlation showed a high level of correlation (ρ = 0.963 and ρ = 0.943, P < 0.001) between the results of the FT and those of the observers. The Wilcoxon test revealed that there was no significant difference between the results of the experts and the model (P = 0.051 and P = 0.316). Finally, it was concluded from the results that the FT could be used as a tool to support the decision of doctors by indicating consistent outputs and hence contribute to the objectiveness and reproducibility of the assessment results.

Key words: Image processing, nucleus classification, segmentation, functional trees, estrogen receptor status evaluation, breast cancer prognosis, Allred scoring, machine learning

1. Introduction

Breast cancer is the second most frequently diagnosed cancer type among the female population of industrialized countries. For patients who have a cancer diagnosis, the treatment type, duration, cost, effective chemical substance, and survival are decided upon based on the prognosis results. One of the important prognostic factors of this disease is the evaluation of the hormone receptor presence in the tumor section [1,2].

Immunohistochemical (IHC) staining of a sample biopsy section is a common method to assess the presence of an estrogen receptor (ER), since it provides cheap material and is an easy procedure [3]. In this study, a computer-based assessment of the ER status was introduced according to the Allred scoring system, where several scoring alternatives are available in medical practice [4–6]. This scoring system is easy to use and is able to identify low-positive cases [7,8] to avoid having false-negative results.

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Despite its widespread usage for prognosis, the assessment of the ER is done subjectively, relying upon the perception of the observer. Due to physiopsychological variation in the perception of human vision, an observer may give different score values for the same specimen at different times (intraobserver variation) or different observers may give different score values for the same specimen at the same time (interobserver variation). Interpretation subjectivity of observers was reported in [9]. This study included 172 pathologists in Germany, revealing that 24% of the pathologists’ ER interpretations were in fact false negatives. Having false negatives will lead to the consequence that these patients will be labeled as ER-negative and will not receive the benefit of endocrine therapy.

According to a survey about IHC techniques, interpretation variation may come from different laboratory conditions and the diversity of IHC staining procedures, such as the duration of the tissue fixation, type of antigen retrieval, antibody specificity, and dilution or detection systems [5]. Therefore, IHC results lack standardization and reproducibility [10]. Even though different conditions are maintained in the same base, it would not be possible to reach standardization and reproducibility of IHC results unless the interobserver and intraobserver variation of the human factor is overcome or at least minimized. With this motivation in mind, aside from searching for standard procedures in different laboratory conditions, several computer-aided systems and methods have also been presented up to now.

Some of the methods implement global thresholding techniques by the intensity or optical density values of several color spaces [10–22]. Some works focus on texture and morphological features [23–25]. Recently, Krecsák et al. introduced NuclearQuant software, which benefits from color, intensity, and size features for nuclei detection [26]. Tuominen et al. introduced a web-based software for nuclei quantification relying on an image color deconvolution algorithm [27]. In this paper, we use color and gray-level statistical, textural, and spectral features while comparing the classification performance of each feature set separately.

When the literature is searched for ER status evaluation, there are several machine learning methods used in this area of medical implementation. These are k-nearest neighbors with weighted votes [23], radial basis neural networks [24], k-means clustering [25], random forest clustering [28], and probabilistic neural network and support vector machines [29]. These papers refer to different scoring protocols than the one that we used.

There is not a unique scoring protocol that is used as a reference for all of the methods. Calculating the agreement or correlation coefficient over a 3- or 8-range scoring protocol might yield different performance results. For this reason, it is not directly possible to compare the achievements of all of the approaches. However, in a survey in [7], it was affirmed that the Allred scoring protocol is becoming widely accepted in medical laboratories because of its low-level scoring sensitivity with additional scoring ranges and the consideration of intensity variation in staining. Hence, the Allred scoring protocol was implemented as a reference scoring protocol in our work.

We designed a computer-based approach in which several work steps depicted in Figure 1 were automatically realized without user interaction. First, an expert captured a representative region of the ER-stained specimen with a light microscope and recorded the vision as an image file. After that, the work of automated analysis started. In the image file, the stained nuclei were segmented from the other structures of the background tissue. This process was carried out by use of the Otsu thresholding method [30], benefiting from the fact that positive- or negative-stained nuclei have a darker appearance compared to the cytoplasm and other parts of the tissue. However, there were some components such as the stromal cells that looked very similar to the nuclei stain. Those types of components were eliminated with morphological operators, as their sizes were obviously smaller than that of the nuclei. Having obtained a segmented image file that only showed the
detected nuclei, the classification algorithm, i.e. functional trees (FTs) [31], was employed to distinguish the positive and negative nuclei, where the statistical, textural, and spectral features were used as inputs for the classifier. The dimension of the input space for the classifier was reduced before the classification process via a correlation-based feature-elimination method [32]. Once the classifier was identified and labeled for each nucleus according to the type and staining intensity, the prognostic score of the whole image was calculated according to the scoring protocol [3]. The classifier model was trained over the training dataset containing 6 image files. Classification performances over the alternative feature datasets were measured by the average values of a $10 \times 10$-fold cross-validation process leading to a total of 100 runs of the model on each dataset. The prognostic performance of the model was tested on a separate test set composed of 27 image files. The successes of the prognostic scores were measured by comparing the scoring values of the classifier model and those of the 2 experts, where statistical analysis of agreement, correlation, and significance of the difference test was conducted.

In our study, according to the test-run results, the classifier model of the FT revealed a very good agreement ($\kappa = 0.899$ and $\kappa = 0.927$, $P < 0.001$) and showed a high correlation level ($\rho = 0.963$ and $\rho = 0.943$, $P < 0.001$) with the observers. Additionally, the Wilcoxon test indicated that there was no significance difference among these 3 scoring results ($P = 0.051$ and $P = 0.316$). From these statistics, it was concluded that the FT can be used as a tool to help doctors give a reproducible decision according to the Allred scoring protocol.

The other sections of this paper are organized as follows. The materials are explained in Section 2, and the methods, including the segmentation, feature extraction, and classification, are described in Section 3. The results of the feature selection, classification, and prognosis are given in Section 4. The conclusion and discussion are given in Section 5.

2. Materials

The archives of the Department of Pathology at the Erciyes University Medical Faculty in Turkey were searched in 2007 and 2008, and 40 cases of invasive ductal carcinomas were identified from the files and collected for the study. Hematoxylin and eosin (H&E)-stained sections of formalin-fixed paraffin-embedded tumors (4–5 $\mu$m thick) were used for histological assessments.

The type of the tumor was defined with regard to the World Health Organization Classification of Tumors [33] and the histological grading was performed according to the method of Elston and Ellis [34]. Representative sections with the tumor and the adjacent normal breast tissue (internal control) were processed for ER IHC staining. Tissue sections (4–5 $\mu$m thick) from the paraffin blocks were used for all of the IHC analyses according to the manufacturer’s instructions. Diaminobenzidine tetrahydrochloride (Dako Liquid DAB Plus, K3468, Denmark) was used as a chromogen and the sections were counterstained with Mayer’s hematoxylin.

After having applied the IHC staining of the ER, nuclei that had a positive ER status expression were stained in brownish colors, whereas nuclei with a negative ER expression were stained in bluish colors. Staining
of the ER was evaluated in the nuclei of the malignant cells, where the ER status was scored using the Allred scoring system [3]. With these properties, 40 specimens were collected in total; however, 7 of them were discarded, since they had substantial artifacts, cytoplasmic stain, or scoring disagreements between the expert observers taking part in this study. An example of a discarded image having cytoplasmic staining is shown in Figure 2.

Figure 2. Image part with cytoplasmic stain.

A pathologist (F.A.) analyzed each slide under a light microscope by selecting a representative region of the specimen and capturing the region on each slide with a linear magnification of 40×. As is usually done in this pathology department, she recorded the representative region as a 2048 × 1536 × 24-bit (8 bits in each channel of RGB) JPEG color image (Lecia DMD 108 Microimaging Device).

Taking into account the interobserver variation, 2 experienced pathologists (H.A. and M.K.) manually assessed these collected specimens according to the Allred scoring protocol. In this protocol, there are 2 types of scores determining the total score: the first is the percentage score (PS), calculated according the proportion formula:

\[
\text{proportion} = \frac{N_p}{N_p + N_n},
\]

where \(N_p\) and \(N_n\) are the number of positive- and negative-stained nuclei. The PS score is defined by comparing the proportion value with Table 1.

The second is the intensity score (IS) given by the overall intensity range of the positive-stained nuclei, where negative stain intensity indicates a 0 intensity score when there is no positive-stained nucleus. Weak, moderate, and strong stain intensities in the positive nuclei mean intensity scores of 1, 2, and 3, respectively. By adding up the PS and IS scores, the total score (TS), known as the Allred score, is calculated, as shown in Table 1 [3].

To form a separate training set, 6 images out of 33 were kept aside and were not used in any of the test experiments. Depending on the type and the staining intensity, 2 experts marked some of the nuclei on these images by labeling each nucleus with a dedicated color using Microsoft Paint Brush Software. Among those labeled nuclei, a total of 384 nuclei that were identified by 2 observers with the same type and intensity range were chosen to form the training data. As it was very tedious and difficult work to mark and get enough nuclei for a separate validation set, it was decided to employ a 10-fold cross-validation scheme to validate the
classification performances. The remaining 27 image files were reserved as the test set and the prognostic TSs of the 2 experts for these images were noted.

Table 1. Allred score: total score (TS) = percentage score (PS) + intensity score (IS) (range 0, 2–8).

<table>
<thead>
<tr>
<th>Staining PS</th>
<th>Proportion of positive staining nuclei</th>
<th>IS</th>
<th>Average intensity of positively stained nuclei</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>None</td>
<td>0</td>
<td>None</td>
</tr>
<tr>
<td>1</td>
<td>&lt; 1/100</td>
<td>1</td>
<td>Weak</td>
</tr>
<tr>
<td>2</td>
<td>1/100 to 1/10</td>
<td>2</td>
<td>Moderate/medium</td>
</tr>
<tr>
<td>3</td>
<td>1/10 to 1/3</td>
<td>3</td>
<td>Strong</td>
</tr>
<tr>
<td>4</td>
<td>1/3 to 2/3</td>
<td>3</td>
<td>Strong</td>
</tr>
<tr>
<td>5</td>
<td>&gt; 2/3</td>
<td>3</td>
<td>Strong</td>
</tr>
</tbody>
</table>

3. Methods
3.1. Nuclei segmentation

In the Allred scoring protocol, the positive and negative nuclei numbers were taken into account rather than counting pixels or calculating the stained area measurements. Consequently, the detection and identification of a nucleus in integrity was a basic part of the scoring process.

With that in mind, the primary goal of the segmentation stage was to detect a nucleus from the tissue background. For this aim, a simple yet efficient approach was followed by running the Otsu thresholding method [30] over the image. During the thresholding, we benefited from the intensity variation in the IHC image, which mainly consisted of 2 parts: the stained nuclei, either positive or negative, and the tissue background. The first part was obviously darker than the second part because of the stain effect. Thus, a representative image file was first transformed into an 8-bit gray-scale intensity image and, after that, the Otsu thresholding method was used to separate the darker objects from the lighter background tissue components. As an output, a binary image was produced, where the nuclei were shown in white and the background was shown in black. Afterwards, a morphological opening operator with a disk-shaped structuring element composed of 12 pixels opened the image. Finally, the connected components were counted and labeled in the resulting image, where each connected component showed a nucleus.

While performing segmentation, some exceptional situations must be considered in advance. Therefore, we considered other types of components in the tissue, such as the stromal cells that were similarly darker but were in fact different from the nuclei. Another possibility was having visually joined cell structures, either because of the opening operator or the cytoplasmic stain itself. We also took into account the nuclei that were too small, particularly due to broken parts in the corners of the image frame, or that were deeper inside the thick section of the specimen and hence partly visible.

Considering these possibilities, it was experimentally decided to have a size filter only allowing for the quantification of nuclei at a size of less than 3000 pixels and more than 300 pixels. By having such a morphological filter, it was possible to avoid the quantification of the stromal cells, as they were quite smaller in size than a nucleus. It was also easy to discard the joined cell structures due to overlapping cytoplasmic stain. As a result of this process, there was a loss of information, since some of the nuclei parts, which were either broken in the corners or partly visible on the surface of the focus, were eventually eliminated. However, this intrinsic loss of information did not cause any biased effect over the classifier, since the size filtering was independent of the nucleus positivity or negativity. Hence, if enough nuclei were detected in an image file of a
representative region, the image file could be used as a reference input for all types of the feature set combination experiments.

Finally, the morphological and spatial properties of each segmented nucleus were recorded and stored in a file for further analysis. An example part cropped from a representative region of a specimen is shown in Figure 3.

![Figure 3. Subprocesses of the segmentation stage: a) cropped part of the original image, b) result of Otsu thresholding on the original part, c) result of morphological operations, and d) detected nuclei borders superimposed on the original image part.](image)

### 3.2. Feature extraction

One of the obvious differences between the positive- and negative-stained nuclei is that the positive nuclei have brownish colors, while the negative nuclei have bluish colors. However, this rough definition alone is not enough to separate both types of nuclei when we consider the diversity of the chemical procedure in medical laboratories that do not necessarily have the same color representation of the ER stain. Additionally, it was reported in the literature that the perception of the colors, especially in low-level stain, may lead to a false-negative prognosis [8].

To avoid dependence on solely the color luminosity features, we also benefit from alternative attributes such as textural, spectral, and second-order statistical features. On the other hand, after some visual examinations, we decided not to use the morphological features, as there was no significant shape or size difference between the positive- and negative-stained nuclei.
In total, 144 features were extracted from each nucleus, where all of the features were generated from the image area inside the border of the nucleus.

### 3.2.1. First-order statistical features

The statistics on an intensity histogram of an image can be used as descriptive attributes. In this work, we calculated 6 different statistical properties of the intensity histogram, which were the mean/average, contrast, smoothness, uniformity, third moment, and entropy [35]. Consequently, by extracting these features from each RGB channel, and additionally from a gray-level transform, a total of 24 first-order statistical features were computed.

### 3.2.2. Second-order statistical features

Even though it is possible to get information about the intensity distribution of an image, the placement information for some specific intensity values is not known without examining the second-order statistical features. Therefore, structural descriptors like the cooccurrence matrix properties were calculated. To form the intensity cooccurrence matrix from an 8-bit color channel, the intensity values were quantized at 8 different levels; hence, an $8 \times 8$-bit cooccurrence matrix was obtained. During this formation, the pixel of interest and its neighbor on the right-hand side were regarded. Thus, 8 structural features, autocorrelation, contrast, correlation, dissimilarity, energy, entropy, homogeneity, and the sum of squares, were calculated from each cooccurrence matrix [36–38]. Finally, 32 second-order statistical features were generated from the color channels of the RGB and from 1 gray-level transform of the nucleus image.

### 3.2.3. Laws’ texture energy features

The existence of some specific types of intensity variations in an image texture can be determined with small convolution kernels. Stemming from this idea, Laws’ texture energy images indicate the level, edge, spot, wave, and ripple appearance inside an image texture via the convolution kernels. The typical Laws’ kernels used for texture discrimination are generated from 1-dimensional (1-D) kernels [39].

The convolution of a vertical 1-D kernel with a horizontal kernel or by repeating the same operation in reverse order yields a 2-D kernel. Hence, it is possible to generate 25 different 2-D kernels. If the directionality of a pattern is not important when searching the existence of the pattern inside the image texture, then similar 2-D kernels are combined so as to make the search complete with a lower number of 2-D kernels. By referring to “similar kernels”, we mean 2 types of kernels composed of the same 1-D kernels by convolving in a different order.

In our work, we have chosen to use 15 rationally invariant kernels, where 10 kernels came from the mutual convolution rationally variant 1-D kernels and 5 self-convolved kernels were used as they are. Only the self-convolved 2-D kernel was discarded, since it did not have a zero-sum. Thus, 14 rationally invariant 2-D kernels were generated, where 4 kernels were self-convolved and 10 kernels were the output of the mutual convolution.

By applying these kernels to each nucleus image and computing the energy function over them, a total of 56 Laws’ texture energy images were obtained from the channels of the RGB nucleus image and 1 gray-level transform of the nucleus image.

### 3.2.4. Wavelet energy features

The periodicity of any pattern in the intensity variation could be useful to identify an object in an image. For this aim, spectral analysis was used with the idea that every signal could be considered as a linear summation of
some basis functions, where all of the functions were orthogonal to each other and they were frequency-shifted versions of one basic harmonic function.

Fourier transform stems from this approach; however, in this transform, it is not possible to detect the time and localization information of any frequency component inside the signal, while it is possible to check whether or not any frequency component exists. To fill this gap, the wavelet transform is proposed, which includes one additional output dimension that is denoted for different scales of the basis function. By changing the scale of the basis function, the frequency content of the signal, including the information of when or where a specific harmonic occurs, can be obtained without losing the spatial representation of image signals.

By employing the wavelet transform [35] in 2 different scales, 8 different wavelet approximations of the original nucleus image were produced. Later, the energy of each approximated image was calculated. Hence, in total, 32 different wavelet approximation-based energy features were generated from the RGB color channels and gray-level transform of the nucleus image.

3.3. Feature elimination

We produced 144 features from each nucleus at the end of the feature generation stage. Considering the 384 instances in the training set, such a high dimensionality of features as inputs may degrade the classification success or easily cause an overtraining error. Usually, it is suggested that the cardinality of the instances must be 10–30 times bigger than the cardinality of inputs of the classifier to avoid possible overtraining [40]. Moreover, some features do not contribute extra information as they are highly correlated with each other. This redundancy in the inputs increases the computational complexity and time of the operation, where it may also cause a decrease in the classification or prediction accuracy [40]. For these reasons, it was experimentally decided to employ a correlation-based heuristic feature selection algorithm in order to have a smaller input dimensionality for the problem.

This heuristic approach relies on the hypothesis that “good feature subsets contain features highly correlated with the class, yet uncorrelated with each other”. Based on this idea [32], Eq. (2) describes the worth of a subset of features:

\[
Worth_S = \frac{k \cdot r_{cf}}{\sqrt{k + k(k-1) \cdot r_{ff}}},
\]

where \(Worth_S\) is the heuristic worth of a feature subset \(S\) containing \(k\) features, \(r_{cf}\) is the average feature–class correlation, and \(r_{ff}\) is the average feature–feature intercorrelation. The prediction capability of the features in the subset is represented by the numerator, while the existence of intercorrelations among the features of the subset, or in other words the redundancy of the features, is indicated by the denominator. It was stated that “this formula actually is Pearson’s correlation, where all variables have been standardized” [32].

3.4. Classification

3.4.1. Performance measures of classification

Having selected some of the features as inputs for the classifier, it was expected from the model to be able to identify the type and intensity range of a nucleus. There were 4 possibilities as the output, as shown in Table 2. Here, the second possibility was not available in the dataset, and so a classifier was designed to give outputs according to the other 3 possibilities.

To measure the classification performance, we used 4 types of methods: correct classification performance, sensitivity and specificity analysis, confusion matrix, and receiver operating characteristic (ROC) curves. As its
name implies, correct classification performance show us the percentage of instances that are correctly identified by the classifier. Given a predicted class of interest, the sensitivity and specificity values were calculated according to the following expressions:

\[ \text{sensitivity} = \frac{TP}{TP + FN} \times 100 \%, \quad (3) \]
\[ \text{specificity} = \frac{TN}{TN + FP} \times 100 \%, \quad (4) \]

where \( TP, \) \( TN, \) \( FP, \) and \( FN \) are acronyms for true positive, true negative, false positive, and false negative, respectively.

<table>
<thead>
<tr>
<th>Table 2. Classes to which a nucleus may belong.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative nuclei in a negative, nonstained specimen (Class 1)</td>
</tr>
<tr>
<td>Positive nuclei in a weakly stained specimen (not available in the data)</td>
</tr>
<tr>
<td>Positive nuclei in a moderately stained specimen (Class 2)</td>
</tr>
<tr>
<td>Positive nuclei in a strongly stained specimen (Class 3)</td>
</tr>
</tbody>
</table>

The performance measures mentioned above are not enough to judge the achievement of the classifier. It is also necessary to know how well the classifier is able to fit to different datasets and how good of a generalization capability the classifier has. For this aim, validation techniques are used, usually by employing an independent dataset as a validation set, which is different from that of the training. However, instead of a separate validation dataset, the \( k \)-fold cross-validation technique on the training dataset can be used in some applications like this one, where it is very tedious and labor-intensive work to get enough data for an additional separate dataset. Therefore, we employed a 10-fold \( (k = 10) \) cross-validation in our work. During the execution of the \( k \)-fold cross-validation, the training dataset is decomposed into \( k \) different parts and one of those parts is separated as a test set, while the remaining \( (k - 1) \) parts are used in the training. This work step is repeated \( k \) times until all parts are used exactly once as a separate test set. Thus, one validation cycle is completed. There may be many possible subsets of the same dataset and the formation of these subsets may affect the performance outputs of the classifier while following this approach. Having considered this condition, in our work, the \( k \)-fold cross-validation cycle was repeated 10 times as a precaution to compensate for any possible bias effect that may come from the formation of the combinations of the subsets. After that, the average results of all of the runs were indicated.

In short, we employed a 10-fold cross-validation on the training data and ran this cycle 10 times to get the average results. Furthermore, during the training and validation processes, stratification was maintained to ensure that the instances were evenly distributed for each class in the subset.

3.5. Classifier model

A decision tree is a tree-like graph-based search model to make decisions. This model recursively partitions the data space into subregions, where each region is represented by the leaf or branch nodes of a tree model. Decision trees are nonparametric learning models, as they are not initially structured or they do not need any parametric form of class densities. On the other hand, to take the final form, a decision tree needs data instances of inputs and outputs together. In that sense, decision trees are supervised learning models [40].
In a decision tree model, rather than using the whole feature set jointly at the same time, different subsets of the features are used in different nodes or branches of the tree model. This search strategy, employing step-by-step reasoning according to a decision hierarchy, reveals a compact and easily understandable decision model for humans. Therefore, decision tree models are known as white-box models [41]; hence, they are the preferred adapting decision support tool for medical applications. Moreover, the interpretation of tree models by humans is useful, as it is easily possible to convert a decision tree model into a base of if-then-else rules [40].

However, tree-based models have intrinsic shortcomings as well. As an example, a binary decision tree may be addressed, which partitions the data space into 2 subregions at each decision node. A binary decision tree represents the data space as the sum of multidimensional rectangles. In the case of having a problem that involves linear and nonlinear combinations of inputs, this tree model can be quite complex, whereas it is possible to form an easier decision model with the use of linear or nonlinear functions. Therefore, to benefit from the advantages of tree models, while being able to handle linear and nonlinear relations in an easy way, some variants of decision trees are proposed.

One of the variants is the FT. For classification problems, multivariate trees can be used by means of a combination of attributes at the decision nodes. For regression problems, model tree algorithms employ linear models at the leaf nodes. In classification problems, multivariate decisions are usually taken at the inner nodes. In regression problems, multivariate predictions are performed at the leaf nodes. The main idea of FTs is to allow for combinations of attributes in both the inner nodes and leaf nodes. This approach is a joint version of multivariate trees and model trees, where multivariate tests are done in decision nodes and class predictions are made using linear functions at the leaf nodes. Multivariate tests in decision nodes are used to grow the tree, whereas functional leaves are built while pruning the tree.

The FT combines a univariate decision tree with linear functions by means of constructive induction, which discovers new attributes from the training data to form a higher-dimensional input space [42]. An attribute constructor is in reality another type of classifier model over the existing data [43].

The use of a constructive attribute operator in classification problems is described in the literature with the implementation of a cascade generalization scheme for multivariate trees [44]. FTs are closely an extension of this approach, where additionally functional leaf nodes are included, and the regression domain and the classification are also covered [31]. Therefore, here the basic classifier framework is first decided in the chosen notation and after that the FT is introduced relying on the cascade generalization concept.

Given a dataset \( D = (\tilde{x}_n, y_n) \), \( \tilde{x}_n \) is an input vector \( (n = 1...N) \) and \( y_n \) is an output value, meaning a membership to one of the several class values \( y_n \in \{C_1, C_2, ..., C_c\} \). A classifier function \( Cf \) can be defined to form a model \( Cf(D) \) using training data \( D \). This model takes inputs from the input space and estimates the outputs in the output space. \( Cf(\tilde{x}, D) \) represents a predictor classifier trained over data \( D \), which takes an input vector \( \tilde{x} \) and is supposed to give an output value \( y \).

The main difference of the cascade generalization framework from a basic classifier is that the predictor \( Cf(\tilde{x}, D) \) outputs another vector \( \tilde{p} \) instead of a single output value \( y \). Here, \( \tilde{p} \) presents a conditional probability distribution \( [p_1, p_2, ..., p_c] \), where \( \tilde{p} = P(y = C_i|\tilde{x}) \). In this case, the output class predicted for the input is to be the one maximizing output probability distribution.

Another difference of the cascade generalization framework is the constructor operator \( \Phi(\tilde{x}, A(Cf(D), D)) \). This operator takes an example vector \( \tilde{x} \) as an input and produces output probabilities for each class with \( A(Cf(D), D) \), where \( A(Cf(D), D) \) represents the application of the predictor model \( Cf(D) \) over the training data \( D \) and \( \tilde{x} \) is a single input vector. The probability distribution for each class label is evaluated as a new
attribute; hence, new attributes are generated and added to existing actual ones, making the input set larger. When the \( \Phi \) operator is applied to the whole instance of the dataset \( D \), then a larger training set \( D' \) is obtained. The number of the instances in \( D \) and \( D' \) are the same, yet \( D' \) has more attributes than \( D \), where the cardinality existing classes and new attributes are the same.

Having more than one classifier, a sequential composition is performed, in which the \( \Phi \) operator is applied in each composition step using different classifiers. Given a training set \( L \) and a test \( T \), and classifier 1 \( C_{f1} \) and classifier 2 \( C_{f2} \), the sequential composition generates a training set \( L_{new}^{1} \) and a test set \( T_{new}^{1} \) by employing \( C_{f1} \):

\[
L_{new}^{1} = \Phi(L, A(C_{f1}(L), L)),
\]

\[
T_{new}^{1} = \Phi(T, A(C_{f1}(L), T)).
\]

Here, the prediction model \( C_{f1}(L) \) is applied to the training and test datasets, modifying them to generate new datasets at composition level 1, which is indicated by the superscript of the dataset name. Another classifier, \( C_{f2} \), learns from dataset \( L_{new}^{1} \) and is applied to the test dataset \( T_{new}^{1} \), i.e. \( A(C_{f2}(L_{new}^{1}), T_{new}^{1}) \). When the operation sequences of several classifiers are shown with dedicated operator symbol \( \nabla \), then the sequence of the composition can be formally expressed as:

\[
C_{f2} \nabla C_{f1} = A(C_{f2}(L_{new}^{1}), T_{new}^{1}).
\]

In the case of having more than 2 classifiers, a composition of \( n \) classifiers is represented by:

\[
C_{fn} \nabla C_{fn-1} \nabla C_{fn-2} \ldots \nabla C_{f1} = A(C_{fn}(L_{new}^{n-1}), T_{new}^{n-1}).
\]

The final model is given by the classifier \( C_{fn} \) after having done \((n - 1)\) levels of compositions. Using several classifiers and functions combined in such a scheme, it is expected to benefit from the knowledge of the representation of different approaches in the input space and the diverse searching capabilities of several methods together.

In addition to the FT model, we also examined the classification of the performance of some other models: the J48 decision tree of the Waikato Environment for Knowledge Analysis (WEKA), which is an implementation of the C4.5 algorithm [45], and several support vector machine (SVM) models that are based on linear, quadratic, and radial base functions [46].

As stated before, a FT is able to use linear combinations of the attributes in the nodes and leaves, while a univariate decision tree makes a simple probabilistic value test over a single attribute in a node. Thus, while a FT has the interpretation easiness of decision trees, it is also possible to exploit the generalization capability of linear regression models without having complex tree structures. This benefit can be exemplified over a dataset of this prognosis application. When the J48 decision tree is constructed over the same dataset, such a decision tree is formed as is given in Figure 4. The tree was built with a confidence factor of 0.25 and pruning was enabled.

On the other hand, the FT method yields just a single node over the same dataset after the growing and pruning processes are completed. This single node contains regression functions, of which the bias and weight values for the features are given in Table 3. Having nominal class labels in the data is not an obstacle when using FTs, whereas it is necessary to double the number of classes by dichotomizing the existing nominal labels so as to use linear regression models.
Both of the tree models are intuitive and easy to understand. However, if compared, the latter is intrinsically more immune to overfitting by means of its smaller structure. This is because the more complex model has too many components to be able to efficiently describe a relation; hence, the overfitting problem tends to occur more, and eventually it ends up memorizing rather than learning.

In addition to these tree models, another method known as SVM was employed over the same data. We chose a sequential minimal optimization algorithm [46] for training a support vector classifier, which has been provided by WEKA. When we examined the performance of several SVM models (linear, quadratic, and radial base function), we observed a slight increase in the performance with low-degree polynomial kernel functions, but much bigger models need more features than a FT. The smallest SVM model was the linear model, which is shown in Table 4 with the weights.

The attributes in Figure 4 and Tables 3 and 4 were actually some selected features explained in the section on feature elimination results. It was clear even without knowing the meanings of these attributes that the FT gave the simplest model, needing fewer attributes than the alternatives. Therefore, the FT model was chosen to get the Allred scores of the test images. However, the correct classification percentages of all of the models are given in the results section.
4. Results

After having a trained model, there was no need of supervising the interaction for any of the images. All of the test images were assessed automatically by the computer model. During the computer-based assessment, 5150 nuclei were detected from 27 test image files, where an average of 191 nuclei were segmented and classified in each image file. All of the codes implementing segmentation, feature extraction, nuclei detection, and finally prognosis were written and executed in a MATLAB software environment, The MathWorks increment implementation of the FT classifier and dimension reduction processes on the data were realized by means of Java routines from the WEKA library [47]. Statistical tests and graphics were performed in the R statistical programming language. All of the mentioned software applications were conducted on a PC that had an Intel Pentium M Processor with 1.73 GHz and 1.25 GB DDR2 RAM. To allow other researchers to replicate the same methods and experiments in their datasets, we addressed the WEKA routines for the dimension reduction and the classifiers at the appendix with initial settings for the experimental methods.

4.1. Feature generation and selection results

As was detailed in the section on feature generation, there were 144 features computed from each nucleus. To sum up, the features can be grouped by these titles: 1st-order gray-level, 2nd-order gray-level, Laws’ gray-level, wavelet gray-level, 1st-order color, 2nd-order color, Laws’ color, and wavelet color features. Furthermore, 3 new groups were evaluated in the experiments for their classification performance, which were features selected from the gray level, color, and complete set as a result of the dimension reduction process.

The experiments were carried out using the WEKA Experiment Utility [47], and the accuracy of the FT method over the different feature groups is indicated in Table 5. In the first column of Table 5, a variety of different feature groups are addressed, while in the second column, the correct classification performance of the FT algorithm is shown. The indicated results were the averages of 10 × 10-fold cross-validated runs of the FT method.

It is clear from Table 5 that the color features were significantly more successful in terms of the correct classification rates than the gray-level features. In addition, among the color features, the wavelet features were better than the others, with a correct classification rate of 97.34%.

<table>
<thead>
<tr>
<th>Classifier for Class 1 and Class 2 weights and variables</th>
<th>Classifier for Class 1 and Class 3 weights and variables</th>
<th>Classifier for Class 2 and Class 3 weights and variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.6555 R-AvInt</td>
<td>0.9832 R-AvInt</td>
<td>0.2458 R-AvInt</td>
</tr>
<tr>
<td>-1.7299 B-AvInt</td>
<td>-1.6367 B-AvInt</td>
<td>-3.1246 B-AvInt</td>
</tr>
<tr>
<td>1.5779 R-Hom</td>
<td>0.8466 R-Hom</td>
<td>1.863 R-Hom</td>
</tr>
<tr>
<td>-0.0356 B-Autoc</td>
<td>-0.1566 B-Autoc</td>
<td>-1.2469 B-Autoc</td>
</tr>
<tr>
<td>2.6781 R-EnAp1</td>
<td>0.9358 R-EnAp1</td>
<td>0.8085 R-EnAp1</td>
</tr>
<tr>
<td>2.3948 R-EnAp2</td>
<td>0.861 R-EnAp2</td>
<td>1.131 R-EnAp2</td>
</tr>
<tr>
<td>-0.4596 B-EnDg2</td>
<td>-1.1557 B-EnDg2</td>
<td>-2.8398 B-EnDg2</td>
</tr>
<tr>
<td>1.4834 R-R5L5</td>
<td>0.7278 R-R5L5</td>
<td>0.6264 R-R5L5</td>
</tr>
<tr>
<td>-0.445 B-E5L5</td>
<td>-0.8425 B-E5L5</td>
<td>-0.4056 B-E5L5</td>
</tr>
<tr>
<td>-1.027 bias</td>
<td>0.7128 bias</td>
<td>0.9939 bias</td>
</tr>
</tbody>
</table>
Having applied a correlation-based feature elimination method, the number of features was reduced from 144 to 11. The correct classification rate of the classifier on this subset of features was 96.67%. Despite a significant dimension reduction, the result was very close to that of the wavelet color features, which had the maximum rate for the classification experiments. The selected features for the prognosis stage of the study are given below in Table 6.

**Table 5.** Different feature datasets and classification performances.

<table>
<thead>
<tr>
<th>Data</th>
<th>Number of features</th>
<th>FT correct classification rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>First-order gray features</td>
<td>6</td>
<td>59.33</td>
</tr>
<tr>
<td>Second-order gray features</td>
<td>8</td>
<td>54.17</td>
</tr>
<tr>
<td>Wavelet gray features</td>
<td>8</td>
<td>61.45</td>
</tr>
<tr>
<td>Laws’ gray features</td>
<td>14</td>
<td>51.77</td>
</tr>
<tr>
<td>First-order color features</td>
<td>18</td>
<td>94.93</td>
</tr>
<tr>
<td>Second-order color features</td>
<td>24</td>
<td>90.45</td>
</tr>
<tr>
<td>Wavelet color features</td>
<td>24</td>
<td>97.34</td>
</tr>
<tr>
<td>Laws’ color features</td>
<td>42</td>
<td>92.81</td>
</tr>
</tbody>
</table>

**Table 6.** Selected color-based features.

- Average intensity in the red channel (R-AvInt)
- Average intensity in the blue channel (B-AvInt)
- Homogeneity of cooccurrence matrix in the red channel (R-Hom)
- Autocorrelation of cooccurrence matrix in the blue channel (B-Auto)
- Energy of approximated wavelet transform of the image at scale 1 in the red channel (R-EnAp1)
- Energy of approximated wavelet transform of the image at scale 2 in the red channel (R-EnAp2)
- Energy of approximated wavelet transform of the image at scale 1 in the blue channel (B-EnAp1)
- Energy of approximated wavelet transform of the image at scale 2 in the blue channel (B-EnAp2)
- Energy of wavelet transform for diagonal detail of the image at scale 2 in the blue channel (B-EnDg2)
- Energy of R5L5 Laws’ kernel convolution of the image in the red channel (R-R5L5)
- Energy of E5L5 Laws’ kernel convolution of the image in the blue channel (B-E5L5)

**4.2. Classification results**

As a result of the feature elimination stage, the reduced feature set, composed of 11 features, was taken into account in the experiments. In this feature set, the performance results of each classifier were compared, having 10 × 10-fold cross-validated runs. It was seen that the correct classification rate for the J48 tree was 93.73%, and for the SVM with radial basis functions it was 84.05%. These results were statistically worse than the other results: FT, 96.67%; SVM linear, 97.50%; and SVM quadratic, 96.80%. There was no statistically significant difference among the FT, SVM linear, and SVM quadratic according to the 2-tailed paired t-test. For comparisons of the paired t-test, the significance threshold level was accepted as $P < 0.05$. Even though the resulting models were not simpler than the FT, the classification accuracy benefits were also negligibly small or worse.

While generating the model of the FT classifier with the 10-fold cross-validation scheme, the sensitivity and specificity outputs of every interim FT model in each test fold were recorded. When all of the validation steps were completed, the average values of these outputs were calculated. The output classifier model was the one that was built over the full training dataset and the other interim models, formed during each validation step, were used for statistical aims and were later discarded. Applying this scheme, the average sensitivity value of the FT classifier over the reduced feature set was 0.969 and the average specificity value was 0.984. These
values indicated for the classifier that it was highly probable that it would correctly recognize each nucleus and staining type. Moreover, having examined the confusion matrix of the FT classifier in Table 7, it was seen that there were few misclassified instances out of the 384 training set nuclei.

Table 7. FT classifier confusion matrix.

<table>
<thead>
<tr>
<th>Actual\predicted</th>
<th>Class 1</th>
<th>Class 2</th>
<th>Class 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class 1 (none)</td>
<td>127</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Class 2 (moderate)</td>
<td>2</td>
<td>123</td>
<td>3</td>
</tr>
<tr>
<td>Class 3 (strong)</td>
<td>0</td>
<td>6</td>
<td>122</td>
</tr>
</tbody>
</table>

The ROC curve is measured to show the performance of a classifier with regard to several threshold values defining the decision. As the threshold value was changed, the separation capability of the classifier between the target class and the remaining classes is depicted in Figure 5, where it can be observed that all of the classes (nonstained nuclei, moderately stained, and strongly stained positive nuclei) were robust against the changes of the threshold.

Finally, it was concluded from the classification performance measures that the FT method was able to give good separation results among the 3 classes.

4.3. Assessment results

We calculated the weighted kappa statistics over the assessment scores to measure the level of agreement between the observers and the computer-based approach. As explained in the Section 2, ordinal scales were used as a reference for scoring. Therefore, in this work, a quadratic scale of weights contributed to the calculation of the kappa statistics by considering different levels of agreements.

Kappa statistics can take values in the scale of [−1 1], where negative values mean disagreement, 0 indicates agreement by chance, and the positive values imply agreement. In this range, −1 indicates perfect disagreement and 1 shows perfect agreement. Possible results according to kappa statistics and their interpretation are shown in Table 8 [48].

In addition to measuring the agreement, we also calculated a pair-wise correlation among the assessors by means of the Spearman’s rank order correlation coefficient. This is a well-known interrater agreement measure.
Furthermore, the Wilcoxon pair-wise signed rank test was employed to see if there was a statistically significant difference among the 3 assessors. Even though having a high correlation means a relation between the scores, it is also necessary to see whether or not the mean values of scores are close enough to each other, because it is possible to have good correlation between 2 series, each of which has very different mean values. In this sense, a significance test is supplemental to the agreement and correlation measures. For the significance test, we tested a null hypothesis, assuming that the average values of the assessments did not differ from each other. The decision threshold was defined as 0.05. If the P-value was bigger than 0.05, then the null hypotheses was accepted as correct; otherwise, it was not.

<table>
<thead>
<tr>
<th>Value of K</th>
<th>Strength of agreement</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 0.20</td>
<td>Poor</td>
</tr>
<tr>
<td>0.21–0.40</td>
<td>Fair</td>
</tr>
<tr>
<td>0.41–0.60</td>
<td>Moderate</td>
</tr>
<tr>
<td>0.61–0.80</td>
<td>Good</td>
</tr>
<tr>
<td>0.81–1.00</td>
<td>Very good</td>
</tr>
</tbody>
</table>

Table 8. Interpretation of Kappa values.

Referring to the Allred scoring protocol, the assessments of the computer-based approach and the observers are given in Table 9. All of the measures for the assessments, i.e. the agreement, correlation, and significance test results, are demonstrated in Table 10.

When the kappa statistic results in Table 10 were compared with the reference values in Table 8, it was observed that there exists a very good agreement between the assessments of the pathologist observers and the computer-based method. In addition, it was noted that highly correlated scores were obtained among the 3 assessors. Finally, considering the significance test outputs, it was concluded that there was no statistically significant difference among the assessors.

5. Discussion and conclusions

In our work, we found the kappa results ($\kappa = 0.899$ and $\kappa = 0.927, P < 0.001$), Spearman’s rank order correlation coefficients ($\rho = 0.963$ and $\rho = 0.943, P < 0.001$) and Wilcoxon significance test results ($P = 0.051$ and $P = 0.316$) between the Allred scores of experts and the proposed automated model. Among the test images, 25 of them (25/27) were correctly dichotomized as positive or negative compared to the average of the assessment scores of the 2 experts. Unfortunately, it is not directly possible to make a comparison between what we achieved and what was reported in the literature, because there are a variety of performance measures reported for ER prognosis and a unified approach has not been adopted for this aim. Additionally, different scoring procedures and decision cutoff values have been applied in the literature. Nonetheless, we will mention similar works and their results so as to provide an overview of the dedicated area of literature. Hence, we think that computer-based prognostic systems and the emerging need of objective benchmarking for the results of different laboratories will contribute to the standardization of performance measures. This also may enforce standardization in the substages of clinical works from the fixation of tissues to the analysis of IHC results.

As seen in Table 1, the Allred scoring protocol is very sensitive to low-level staining variations because of the PS score ranges (0%–1%, 1%–10%, 10%–33%, 33%–66%, and 66%–100%). Given a positive nuclei detection percentage of even less than 2%, there may be 3 different prognostic scores according to the Allred scoring protocol. Considering the sensitiveness to such small intervals in terms of nucleus detection, having 1 or 2 wrongly detected nuclei could easily alter the whole prognostic score, while some other protocols cited in the
Table 9. Allred scores on test images.

<table>
<thead>
<tr>
<th>Images</th>
<th>FT</th>
<th>Obs. 1</th>
<th>Obs. 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>6</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>5</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>5</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>4</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>7</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>8</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>10</td>
<td>6</td>
<td>6</td>
<td>7</td>
</tr>
<tr>
<td>11</td>
<td>5</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>12</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>13</td>
<td>7</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>14</td>
<td>7</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>15</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>16</td>
<td>6</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>17</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>18</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>19</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>20</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>21</td>
<td>6</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>22</td>
<td>7</td>
<td>8</td>
<td>7</td>
</tr>
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<td>23</td>
<td>7</td>
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</tr>
<tr>
<td>25</td>
<td>6</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>26</td>
<td>5</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>27</td>
<td>7</td>
<td>8</td>
<td>8</td>
</tr>
</tbody>
</table>

Table 10. Assessment results of observers and classifiers (asterisk signifies P < 0.001).

<table>
<thead>
<tr>
<th>Assessments</th>
<th>Weighted kappa agreement</th>
<th>Spearman’s correlation</th>
<th>Significance of difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>obser1–obser2</td>
<td>0.944*</td>
<td>0.948*</td>
<td>P = 0.148</td>
</tr>
<tr>
<td>FT–obser1</td>
<td>0.899*</td>
<td>0.963*</td>
<td>P = 0.051</td>
</tr>
<tr>
<td>FT–obser2</td>
<td>0.927*</td>
<td>0.943*</td>
<td>P = 0.316</td>
</tr>
</tbody>
</table>

The Allred scoring protocol is more sensitive to variations in low percentage cases and it is harder to get an agreement between the observers according to this protocol. Despite its difficulty in implementation by a computer-based approach, the Allred protocol is becoming widely practiced in medical laboratories, as it is clinically validated [3,8]. Large-scale clinical trials suggest the use of this protocol [6]. The importance of having sensitive techniques detecting low levels of ER and the focus of the Allred score on the low levels have been reported in the literature [7]. Considering these facts, in our work, we proposed a computer-based prognosis method relying on the Allred Protocol. This protocol has
not been used in these previous studies, which also realized nuclei detection-based ER prognosis [23–25,27–29].

To the best of our knowledge, there are a few studies [25,26] detecting and counting nuclei while applying Allred scoring for ER assessment. In a similar work, Krečsák et al. presented a computer-based automated approach for ER status evaluation [26] according to the Allred scoring protocol. Even though a higher agreement score ($\kappa = 0.981$) was reported in that paper, unlike our study, the reported nilpotent quotient algorithm needs user interaction and recalibration for the image to be assessed. Additionally, the color, intensity, and size features of the nuclei are considered in this recent work, whereas our study also computes additional features like wavelet transform energy features, giving higher correct classification rates according to Table 5 compared to the first-order statistical features. In another work that conducted nucleus detection and quantification with the same protocol, Sharangpani et al. [25] found a strong correlation between the algorithm-based values and the subjective measurements (intraclass correlation: 0.77; 95% CI: 0.59–0.95) for the ER and the progesterone receptor percentage nuclear positivity. In our work, the automated model achieved a higher level of correlation between the scores of the experts and the model.

Some of the studies used nuclei detection and counting approaches, as we did. These studies, given below, referred to different scoring protocols. Kostopoulos et al. employed an unsupervised segmentation method, maintaining an adequate level of agreement (Kendall’s $W = 0.79$) between an automated computer-based system and physician evaluations [25]. The same authors introduced a color texture-based image analysis method resulting in an agreement level of Kendall’s $W = 0.875$, $P < 0.001$ [23]. In another work by the same authors, a high correlation value ($\rho = 0.89$, $P < 0.001$) was reported using Spearman’s rank order correlation between the assessments of a histopathologist and an image analysis system that was based on texture energy features [29]. Schnorrenberg et al. proposed a computer-aided detection system, the biopsy analysis support system (BASS), that achieved Spearman’s rank order correlation levels of $0.78 < \rho < 0.86$ ($P < 0.001$) for strongly and very strongly stained nuclei. For the weakly stained nuclei, the correlation between the BASS system and the 2 experts was lower ($\rho = 0.51$ and $\rho = 0.38$, $P < 0.001$) [24].

Some previous studies in the literature relied on global threshold techniques, pixel- or area-based measures, for assessment [10,12–22]. Among these studies, we focus on those giving agreement or correlation results between automated systems and the observers, as we have these types of results. We excluded the results of Charpin et al., Furukawa et al., Hatanaka et al., and Lehr et al., as they compared automated systems with biochemical procedures [10,17,13] or cytometric analysis [15]. In these works, the usage of some commercial systems, i.e. SAMBA 200 by Charpin et al., WinROOF by Hatanaka et al., Adobe Photoshop by Lehr et al., and CAS 200 by Furukawa et al., were proposed. For the ER evaluation, the agreement rate between the automated quantitative coronary analysis image analysis system and the manual scoring was reported as $\kappa = 0.84$ by Diaz and Sneige [10]. Comparing the dichotomized scores between the automated Ariol machines and the visual scores of 2 pathologists, the highest agreement rate was reported as $\kappa = 0.9021$ (95% CI: 0.8854–0.9180) by Turbin et al. [12]. Gokhale et al. examined 2 commercially available systems: the ChromaVision Automated Cellular Imaging System (ACIS) and the Applied Imaging Ariol SL-50. The highest correlation level according to the gamma statistics for the ER scores between the observers and the automated systems was obtained by the ACIS as $\gamma = 0.91$ in [19]. Mofidi et al. observed a close correlation between the median optical density-based Adobe Photoshop mask implementation and the percentage positivity assessed manually ($r^2 = 0.844$, $P < 0.0001$) [14]. McClelland et al. reported a correlation level for all of the tumors as $r = 0.919$, $P < 0.01$, $n = 94$ between the percentage measurements of the positively stained nuclear area by the CAS 100 system and the manually determined percentage of the stained cell nuclei, where a lower value of $r = 0.821$, 1216
P < 0.01, n = 68 was reported for the ER-positive cases only [16]. Chung et al. recorded a mean linear regression of R = 0.8903 by comparing AQUA software results in a logarithmic scale with gold standard percentage scores in a study of 29 slides from 11 classic cases [18]. Camp et al. found a high degree of correlation (R = 0.884) between the AQUA-based automated system and the pathologist’s evaluation. They also compared the variability of a pathologist’s evaluation and the automated analysis and reported that the automated analysis had a slightly better reproducibility (R = 0.824 versus R = 0.732) [20]. Lloyd et al. investigated 2 imaging systems, Definiens (Munich, Germany) and Aperio Technologies (Vista, CA, USA), and concluded, without giving statistical details, that both algorithms scored 10/10 cases within the range of the pathologist’s visual labeling index [21]. Bolton et al. found that the agreement between the results of the pathologist and the automated negative/positive and categorical scores were excellent for ER-α (κ range = 0.86–0.91), and lower levels of agreement were seen for ER-β categorical scores (κ = 0.80–0.86). In this work, the performances of 3 different automated systems for IHC scoring were assessed: TMAx (Beecher Instruments, Sun Prairie, WI, USA), Ariol (Applied Imaging, Grand Rapids, MI, USA), and TMALab II (Aperio) [22]. By utilizing the color deconvolution algorithm, Tuominen et al. introduced the ImmunoRatio, which is a publicly available web-based IHC analysis tool. It is reported that the calibrated ImmunoRatio has a strong linear relation (r = 0.98) with visual assessments in terms of the percentage positive nuclei of a test set, including several types of biomarkers: estrogen, progesterone, and Ki-67 [27]. According to the results of Rexhepaj et al., an excellent correlation was observed between the percentages of positive tumor nuclei by image analysis and manual analysis (Spearman’s ρ = 0.9, P < 0.001). In the same paper, a strong correlation between image analysis and manual assessment was indicated for the ER scores (Spearman’s ρ = 0.74, P < 0.001) [28]. Sharangpani et al. published a summary table for the properties of commercially available tools in addition to their custom-made algorithm [25]. A dedicated review of commercial devices can be found in [49]. Prasad et al. introduced a pixel color quantification-based in-house application, TissueQuant, that provides the Spearman’s correlation (r = 0.53) between the automated system and the visual scores for ER expression [50]. The common point of all of these tools is that they are pixel- or area-based measures or global thresholding techniques for the luminosity features.

On the contrary, these studies based on global threshold techniques, pixel- or area-based percentage measures, nuclei detection, and counting-based approaches are used by experts in real medical practice. In that sense, our approach is more closely related to real medical applications by giving outputs directly fitting the interpretations of the human experts. Moreover, by utilizing not only the luminosity thresholding but also pattern recognition methods, the discrimination efficiency of the positive versus the negative nuclei was increased, as was seen in the results of the feature elimination stage, where 10 out of 11 selected features were texture-based and only 1 remaining feature was an intensity feature.

In addition, using such a computer-based tool, it was easier to have a nuclei quantification analysis report for each specimen. Since there is diversity in medical laboratories regarding scoring systems and cutoff values for status evaluation, there is an emerging trend to ask the laboratories to give their quantification values, such as the number of detected nuclei or positivity percentage, rather than just expressing the qualitative assessment results as negative or positive. For this reason, we think that this computer-based approach can make it easier for experts to record and report the documentation of quantitative data, justifying their decision, which is also a benefit providing the reproducibility of a prognosis.

Custom-made software and algorithms add value to the research of standardized automated IHC analysis. However, the application of these approaches by other researchers may involve time and extra effort. Considering this fact, we adopted a compact approach by utilizing one of the existing machine learning libraries [47]. We
also addressed open-source Java codes to allow other researchers to replicate the method that we proposed here. In such an integrated and shared application environment allowing for ready-to-use libraries, it would be easier to explore the achievement of alternative machine learning methods. Hence, we expect that the research in this area might increase by facilitating the usage of different machine learning methods in available tools. Moreover, the need for benchmarking of the results from several machine learning algorithms on the same basis may lead to standard assessment performance measures.

Another contribution of this paper was the use of the FT method as a classifier for this medical application. Even though there might be many different alternative classifiers for this study, it was taken into account that tree-based approaches give compactly stored robust models. Unlike other classifiers described as a black box because of their complicated structure, tree models are easy to understand for experts and could also provide insight into the existing data structure [40,41]. On the other hand, it is known that decision trees have some drawbacks, such as the difficulty in designing an optimal tree with fewer nodes and the need to define the search space with parallel orthogonal rectangles in the case of using binary trees. To circumvent these weaknesses [31], the FT method was intentionally chosen from among the other tree-based methods.

Having implemented the FT classifier, it was observed in the assessment results that there was a very good agreement between the scores of the observers and the computer-based approach. In addition, the assessments of all of the raters were highly correlated and they did not have a statistically significant difference. However, it was also noteworthy that the agreement between the FT classifier and observer 1 was slightly lower and the significance of difference was nearly at the border of the threshold. Still, all of these findings showed the practical potential of the work. It could be possible to increase the accuracy further, with a wider range of data and optimized substages such as an investigation performance of several of the dimension reduction techniques or several types of classifiers over the problem. Furthermore, although the preprocessing of the images was not carried out in this work, the effects of the preprocessing, performance comparison of different segmentation procedures, and morphological operators can be investigated as a future study.

As to the limitations of this application, 4 main points must be addressed. First, the region of interest on a slide must be chosen by an expert in advance. Second, the same fixation procedures must be applied for all of the tissue samples. Third, the images that have a cytoplasmic stain, artifacts, or blurred patterns must be discarded, because some parts of the stained cytoplasm are detected as positive nuclei by the model. Hence, it erroneously tends to give positive scores when an image with cytoplasmic stain is involved. As the last limitation, some additional time for the algorithmic process must be given at least once, as supervised classifier algorithms need a training process before their implementation.

All in all, after obtaining the experimental results, it was seen that image analysis with FT classifiers could be useful to pathologists as a support tool. This may reduce intralaboratory variation and contribute to the increase of the interobserver and intraobserver reproducibility for ER status evaluation according to the Allred scoring protocol.

6. Appendix

WEKA routines used in this work for the classifiers and feature elimination technique were given with the initial settings in Table 11, and these codes can be freely experimented with [47].
Table 11. Classifiers and Java routines of WEKA with initial settings.

<table>
<thead>
<tr>
<th>Classifiers or algorithms</th>
<th>WEKA routines and initial settings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimension reduction</td>
<td>weka.attributeSelection.CfsSubsetEval</td>
</tr>
<tr>
<td>Functional trees</td>
<td>trees.FT -I 15 -F 0 -M 15 -W 0.0</td>
</tr>
<tr>
<td>C4.5 decision tree</td>
<td>trees.J48 -C 0.25 -M 2</td>
</tr>
<tr>
<td>SVM linear</td>
<td>functions.SMO -C 1.0 -L 0.0010 -P 1.0E-12 -N 0 -V -1 -W 1</td>
</tr>
<tr>
<td></td>
<td>-K &quot;functions.supportVector.PolyKernel -C 250007 -E 1.0&quot;</td>
</tr>
<tr>
<td>SVM quadratic</td>
<td>functions.SMO -C 1.0 -L 0.0010 -P 1.0E-12 -N 0 -V -1 -W 1</td>
</tr>
<tr>
<td></td>
<td>-K &quot;functions.supportVector.PolyKernel -C 250007 -E 2.0&quot;</td>
</tr>
<tr>
<td>SVM radial base function</td>
<td>functions.SMO -C 1.0 -L 0.0010 -P 1.0E-12 -N 0 -V -1 -W 1</td>
</tr>
<tr>
<td></td>
<td>-K &quot;functions.supportVector.RBFKernel -C 250007 -G 0.01&quot;</td>
</tr>
</tbody>
</table>

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References


