Using Machine Learning for Classification of Cancer Cells from Raman Spectroscopy

Lerina Aversano¹[®]^a, Mario Luca Bernardi¹[®]^b, Vincenzo Calgano³, Marta Cimitile²[©]^c, Concetta Esposito³, Martina Iammarino¹[®]^d, Marco Pisco³, Sara Spaziani³ and Chiara Verdone¹

> ¹University of Sannio, Department of Engineering, Benevento, Italy ²Unitelma Sapienza University, Rome, Italy ³University of Sannio, Optoelectronic Division - Engineering Department, Benevento, Italy

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Abstract: Since cancer represents one of the leading causes of death worldwide, the development of approaches capable of discerning healthy from diseased cells would be of fundamental importance to support diagnostic and screening techniques. Raman spectroscopy is the most effective molecular analysis technique currently available and provides information on the molecular composition, bonds, chemical environment, phase, and crystalline structure of the samples under examination. This work exploits a combination of Raman spectroscopy and machine learning models to discriminate patients' liver cells between tumor and non-tumor. The research uses real patient data, provided by the Center for Nanophotonics and Optoelectronics for Human Health (CNOS), which analyzed the cells of a patient with liver cancer. Specifically, the dataset has been built through a long data collection process, which first involved the analysis of the cells with Raman spectroscopy and then the training of two classifiers, Decision Tree and Random Forest. The results show good performance for the trained classifiers, especially those relating to the Random Forest, which reaches an accuracy of 90%.

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1 INTRODUCTION

Health institutions' reports clearly highlight that one of the leading causes of death in the world is cancer (Torre et al., 2016; Miller et al., 2016). Just restricting to 2020, we can count about 19.3 million new cases of cancer in the world and about 10 million deaths due to the disease.

Tumors represent a very complex disease and can manifest in distinct phases, (i) the initial or localized disease, in which there is only a single tumor in a single site; (ii) the phase of relapse, whether or not surgery, in which the disease recurs, but always and only in the site where it appeared for the first time and (iii) the disseminated form, in which the malignant cells have exited the organ of origin to colonize other organs even at a distance (metastasis). The main problem for its identification and possible treatment concerns the fact that each type of tumor requires a different approach and, often, also different treatment times.

Research in this regard is always in constant evolution and currently, medicine has several methodologies at its disposal to fight cancer. Active surveillance is reserved for very slow-growing cancers, surgery, radiotherapy that uses X-rays to destroy cancer cells, chemotherapy uses cytotoxic drugs, which are toxic to cells, as they block the division of rapidly replicating cells, without distinguishing between healthy cells and diseased cells, hormone therapy, biological or molecular target drugs are substances capable of "recognizing" the tumor cell and promoting its destruction by the immune system, and finally immunotherapy consists of drugs capable of stimulating the immune system against cancer cells. In general, the earlier a diagnosis is, the more timely and effective the treatment can be. The study (Neal et al., 2015) showed that there is a correlation between diagnosis made in a short time and a favorable outcome for the patient. Therefore, developing approaches capable of distinguishing healthy cells from compromised ones

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^a https://orcid.org/0000-0003-2436-6835

^b https://orcid.org/0000-0002-3223-7032

^c https://orcid.org/0000-0003-2403-8313

^d https://orcid.org/0000-0001-8025-733X

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that can support diagnostic and screening techniques would be fundamental. During the first phase of diagnosis, the most important step is between cancer cells and normal cells. Currently, the methods in use for tumor detection include several approaches including MRI, tomography, endoscopy, different biochemical pathways in combination with mass or optical spectroscopy (Henschke et al., 1999; Sun et al., 2008; Zhu et al., 2008). Unfortunately, however, these require very sensitive times and equipments for the early diagnosis of the tumor.

In the late 1970s, Raman spectroscopy with an optical microscope was introduced and has been used for microanalysis in many fields (Mulvaney and Keating, 2000). Micro-Raman spectroscopy has become an important tool in biology, particularly for singlecell studies (Smith et al., 2016). Raman spectroscopy exploits the interaction of light through a process of diffusion with matter (called scattering) to obtain information on the characteristics of a material and its molecular structure but, infrared spectroscopy is based on the absorption of light, Raman spectroscopy provides information on intra- and intermolecular interactions. Therefore, analyzing Raman spectra, it should be possible to identify differences in molecular compositions and structures between normal and cancer cells and tissues.

At the same time, the technique of machine learning is spreading more and more in different fields of medicine and bioengineering (Aversano et al., 2021b; Ardimento et al., 2021; Aversano et al., 2020). Machine learning algorithms are fast and effective in learning from the data that is provided in input, generating calculation models capable of automatically and rapidly producing evaluation results.

Therefore, the combination of Raman spectroscopy and machine learning could lead to a reduction in the time required for the diagnosis of patient cells, in order to subject the patient to specialist examinations as soon as possible in case of need (Zhang et al., 2021; Zhang et al., 2022).

This research study is placed in this context and has as its objective the classification of cells into the tumor and non-tumor cells analyzed with Raman spectroscopy.

This document is structured as follows. The following section provides some basic information. In Section 2 there is a brief discussion of the related work. The proposed approach is described in Section 4, while the results of the experiment are discussed in Section 5. Finally, in Section 6 and 7, respectively, the threats to validity and conclusions are reported.

2 RELATED WORKS

In recent years, numerous studies have focused on the classification and prediction of human diseases of different types. In particular, machine learning techniques have been largely used for the early diagnosis of many diseases, such as diabetes(G. and K., 2019), heart disease(Karayılan and Kılıç, 2017), Parkinson's disease (Aversano et al., 2020) and thyroid diseases (Aversano et al., 2021a). More recently they have also been used for Covid-19 diagnosis (Rasheed et al., 2021). These works are intended to reduce the time and costs required for the diagnosis and treatment of the patient.

Similarly, the proposed study aims to develop a predictive model of liver cancer in a patient, starting from the Raman spectroscopy analysis of patient cells.

Few other studies have investigated approaches based on the combination of Raman spectroscopy and machine learning with this objective.

The study (Germond et al., 2018) concerns the application of techniques based on machine learning for the classification of cell types. The authors present different approaches to exploit the Raman hyperspectral images: they extract information of the cell from the calculation of the average spectrum (i) and from the wave numbers used to map the distribution of molecular compounds (ii) combining the two previous methods (iii). The adopted classification method is the PCA-DA approach consisting of a principal components analysis (PCA) step followed by discriminant analysis (DA). In addition, authors investigated projection on latent structure (PLS-DA), predictive model K-means and Support Vector Machine (SVM). With spectrum-based classification, the PCA-DA model showed an accuracy of 83.3%, while image-based classification scored an accuracy of 96.3%. The combination of the two approaches archives a 100% accuracy in cell line discrimination in their experimentation.

Like the previous work, also in (Pavillon et al., 2018) the authors used the statistical approach of PCA for the analysis of small cellular changes in response to stimuli, acquiring different parameters with unlabeled microscopy and achieving an accuracy of the model equal to 85

In (Schie et al., 2016) the authors used Raman spectroscopy for the diversification of eukaryotic from prokaryotic cells. Since the former is smaller than the latter, a single Raman spectrum is often enough to generate a dataset sufficient for the training phase. For the latter, more than one spectrum is necessary. Since probing entire cells with Raman spectroscopy using high resolution takes a long time, the authors propose a method that acquires integrated Raman spectra that can cover a large portion of the cell. The approach exploits support vector machines for classification by comparing single spectra with integrated Raman images and spectra of cells. Their results show that the sensitivity of the model can be as high as 90%.

The study (Hsu et al., 2020) deals with stem cells, which can self-renew and differentiate into multiple cell types, allowing the evaluation of pharmaceutical effects and allowing the treatment of various neural diseases. The authors propose a platform exploiting Raman-labeled spectroscopy to classify cells into the different classes of neural cells (from induced cell stem). For this reason, the authors used several classification models (i.e., Support Vector Machine, Random Forest, K-Nearest Neighbor, and the Stochastic Gradient (SGB) enhancement model) that achieve an average accuracy of 97.5%.

In (Ren, 2020), the authors used normal breast cells and prostate cancer cells and an NGK machine learning approach to obtain a prediction value between 87% and 89% without considering the outliers.

The study (Lussier et al., 2019) reports an approach based on the combination of Raman spectroscopy and Deep Learning for the analysis at the same time of at least eight in vitro metabolites close to different cell lines. Analyzing these components is fundamental because it allows research on living cells for responses to inflammation and wound healing. The authors propose a supervised ANN neural network that assigns multiple spectra to the same metabolite. The network consists of two convolutional layers and two pooling layers and uses the softmax function at the output. The results show good model performance, which achieves an accuracy of 86.8%.

Our study aims to classify tumor cells from nontumor cells through Machine Learning methods for the first time. Moreover, the proposed model was tested on true patients' records as provided by the Center for Nanophotonics and Optoelectronics for Human Health (CNOS).

3 RAMAN SPECTROSCOPY

Raman spectroscopy is a widely used spectroscopic method. It is an analysis technique providing information on the chemical structure and molecular interactions of a target sample (e.g., a tissue segment or even a single cell). It works by pointing a laser beam at a sample. Like is shown in Figure 1, a scattered

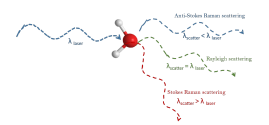


Figure 1: Scattering processes occurring when light interacts with a molecule.

light excites molecules in the sample and the scattering effect is observed. Then, the scattered light is collected by an optical system including a microscope objective, and decomposed by the spectrograph.

A small amount of the scattered light shifts in energy from the laser frequency because of interactions between the incident electromagnetic waves and the vibrational energy levels of the molecules in the sample. Plotting the intensity of the shifted light against the frequency produces a Raman spectrum of the sample.

The measured scattered light showed a broader spectrum with additional wavelengths. A second filter (emission filter) behind the probe allowed blocking the incident wavelength. The observed residual scattered light could now be clearly distinguished from the incident light. Raman spectra are usually plotted according to the laser frequency, meaning the Rayleigh band falls between 0 cm and 1 cm. On this scale, the band positions sit at the frequencies corresponding to the energy levels of varying functional group vibrations.

Therefore, in summary, the light exits the system hit by the laser in three components. A part of the radiation spreads elastically in all directions without loss of energy, i.e. at the same frequency as the incident radiation (elastic scattering or Rayleigh). A large portion passes through the sample and a very small portion is diffused inelastically. This diffusion can be of three types: anelastic ceding diffusion (Raman Stokes diffusion); Acquisition of anelastic diffusion (Raman anti-Stokes diffusion); and Energy in the interaction with the molecule.

As for the excitation source, an intense monochromatic beam is used. Monochromatic because the frequency shifts of the radiation diffused by the incident radiation are very small and therefore the source must be monochromatic to facilitate observation. Intense because the intensity of the diffused radiation is very low and therefore the incident radiation must have a much greater intensity.

4 PROPOSED APPROACH

In this section, we report the proposed approach, which aims to classify a cell as a tumor or not a tumor. First, we describe the process used for data collection, then the machine learning algorithms used and the parameters we have set for their operation, and finally, the metrics used for the validation of the model.

4.1 Data Collection

The data for the research was provided by the Center for Nanophotonics and Optoelectronics for Human Health (CNOS), which have analyzed the cells of a patient with liver cancer (hepatocarcinoma) using RAMAN spectroscopy. In this regard, two tissue samples from the liver have been taken from the patient, one in the area affected by the tumor, and another located in a part distant from the tumor but belonging to the same liver tissue as the patient.

The cells have been supplied to the laboratory by the National Cancer Institute IRCCS G. Pascale Foundation¹. These are real cells that have not undergone any preliminary process and that compared to cell lines, which correspond to a cell taken from the patient and replicated thousands of times, have not been cultured or immortalized. Therefore, the lack of replication ensures that there is no loss of information related to their fundamental properties.

For the analysis using Raman spectroscopy, LabRAM HR Evolution has been used, a system that, thanks to the Raman effect, can obtain high spatial and spectral resolution spectra using ultra-fast confocal images. This instrument offers a wide range of wavelengths, from 200 to 2200 nm, and can reach frequencies of the order of 10 cm^{-1} using an ultra-low frequency module. For the processing and setup of the measurements, the instrument has been supplied with the LabSpec software.

During the measurement, the first phase consists of the self-calibration of the instrument to set the right parameters to obtain the Raman effect. The instrument also has a Rayleigh filter which takes care of filtering the non-informative vibrations, thus showing only the Raman Stokes vibrations. In the second phase, instead, the cell suitable for measurement is searched for using a resolution of 10x. In the figure 2 it's possible to see the candidate cells for possible measurement.

The selection of the suitable cell for measurement is made based on the experience of the operator, who



Figure 2: Candidate cells for measurement with 10x resolution.



Figure 3: Visualization of a cell with resolution 100x and laser positioning.

chooses the one that has the best visual characteristics, excluding cells with irregular shapes or cells that are involved in the cell division process. Once the cell is selected, 100x resolution is used for a more detailed view of the cell At this point, the operator moves the laser towards a point within its nucleus, making 5 measurements for each cell at different points within the nucleus of the same cell. The figure 3 shows the cell with 100x resolution, where the green dot represents the laser that is manually guided by the operator.

The measurements carried out concern the Finger Print Region which captures the Raman spectrum in the region ranging from 600 cm^{-1} up to 1800 cm^{-1} . There is also another region for acquisitions, the High Wawe Length Region which instead ranges from 1800 cm^{-1} up to 3100^{-1} . In other cases, it is possible to make measurements on the union of the two regions, therefore from 600 cm^{-1} up to 3100 cm^{-1} .

Once the data was collected, these have been subjected to preprocessing operations: vector normalization, and windowing.

Having available a vector space that has an internal product and a norm, the vector normalization has been performed because it allows obtaining a unitary norm to the vector space In addition to normalization, the removal of outliers, background, and sub-

¹https://www.alleanzacontroilcancro.it/en/istituto/istitutonazionale-tumori-fondazione-pascale/

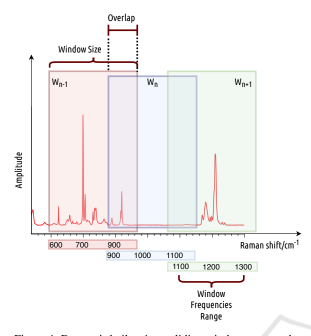


Figure 4: Dataset is built using a sliding windows approach.

strate was also performed.

The windowing technique, exemplified in Figure t 4, has been used to divide the sample into windows of varying widths on which the ensemble is trained. In particular, the window size ranges from 2 samples to 90 samples. Data is divided into multiple windows, characterized by different sizes and overlaps. In this regard we have used four levels of overlap: (*i*) no overlap; (*ii*) overlap of 0.5, meaning that windows overlap by half of their size; (*iii*) overlap of 0.75, meaning that windows overlap by $\frac{3}{4}$ of their size and (*iv*) leave-one-out overlap, meaning that the overlap is almost total advancing the window by just one sample.

The final dataset that we have used consists of 364 frequencies, and each of them corresponds to a sample of amplitude in the various records of the dataset. Specifically, within the dataset used, each record has:

- a progressive number representing the cell number and the measurement number, separated by a dot. These two numbers, taken together, identify one of the five points measured in the cell nucleus (for instance, "1.1" identifies the first measurement of the first cell);
- the measurement samples that represent the amplitudes corresponding to the various frequencies.
- the class label to predict, which identifies the type of cell, and can take two values: *Tumor* if it is a measurement made on a tumor cell, *Non Tumor* if the cell is healthy.

4.2 Data Augmentation

To mitigate the problem of the reduced size of the dataset, a data augmentation step has been performed. It is a set of techniques that extend the available dataset without actually collecting new elements: data augmentation applies random controlled changes to the already existing data, making modified copies. Augmented data can be either slightly modified copies of already existing data or synthetic data created starting from the initial dataset (van Dyk and Meng, 2001). Therefore, with this technique, we have increased the set of spectral data. In summary, for spectral data, data augmentation is mainly carried out with changes in the slope of the spectrum, random multiplication of the amplitudes, addition of random offsets, or, as achieved in this study, with the shift of the wavelengths of the spectra. Another advantage of this technique is that it reduces the phenomenon of overfitting. Therefore, in this work an augmentation was carried out based on the random generation of new spectra shifted in wavelength, with the help of the *aug_xshift()* function.

More specifically, to use this function we have set the following parameters:

- the spectrum, randomly selected from the set of spectra available;
- the shift range specifies the range of possible random shift values for the spectrum. In this case, 6 indicates the left limit at -6 and the right limit at +6, corresponding to the interval [-6, + 6];
- the quantity, the number of new spectra generated starting from the original spectrum. In this case, we have opted for a value of 1 corresponding to the default value of the function, so only a new spectrum will be generated;
- the classes to predict, 1 in the case of tumor cell spectra, 0 for non-tumor cells.

Therefore, an equal number of new spectra were generated for the two prediction classes, where each spectrum was randomly selected from the available set. With this technique, we conducted an additional analysis, using a dataset made up of 632 samples as the input of the classifiers, doubling the initial dataset.

4.3 Classification Methods and Setup

For the classification of cells into the tumor and nontumor cells, different classifiers have been used, varying the algorithm, the window size, the window overlap, and the adopted features. The goal is to train a model that can support the diagnosis of liver cancer

Metrics	Formula	Description							
		The accuracy of the model on the test set, it defines true							
Accuracy	$A = \frac{TP + TN}{TP + FP + TN + FN} (1)$	positives (TP) and true negatives (TN) as correctly clas-							
neeuracy	$A = \frac{1}{TP + FP + TN + FN} $ (1)	sified instances while false positives (FP) and false nega-							
		tives (FN) as classified instances incorrectly							
		The ability of a classifier not to label as positive an in-							
Precision	$\mathbf{P} = \frac{TP}{TP + FP} \ (2)$	stance that is in fact negative, it represents the ratio be-							
Frecision		tween correct forecasts (TP) and the total forecasts (TP +							
		FP).							
		The sensitivity of the model, it represents the ratio be-							
Recall	$R = \frac{TP}{TP + FN} (3)$	tween the correct predictions for a given class on the total							
		of cases in which the class is verified							
		The harmonic and weighted average of the Precision and							
F1 Score	$F1 = 2 * \frac{Precision*Recall}{Precision+Recall}$ (4)	Recall. A classifier gets a high F1-Score only for high							
		Precision and Recall values.							

Table 1: Validation Metrics.

effectively. More specifically, we have used learning algorithms based on decision trees.

The *Decision Tree Classifier* (DTC) (Rokach and Maimon, 2014) is an algorithm for which classification functions are learned in the form of a tree. Within the tree, the variables are represented by the nodes, the possible value for that property is represented by an arc at a child node and the expected value for a given class based on the values of the other properties is represented by a leaf. For each iteration, each attribute is evaluated and the information gain is calculated, the attribute that gets the most information gain is set as the new tree node. Once at the leaf node, the algorithm assigns the class to which they belong to the remaining instances. If there are multiple classes, a probability distribution is assigned.

The *Random Forest Classifier* (RFC) (Breiman, 2001) is a classifier obtained by aggregating multiple decision trees, which are trained with a random subset of the training dataset. Each tree produces a prediction of the class. Once all the predictions have been produced, the result will be the one that appears most frequently.

To train the classifiers we have used these parameters:

- DecisionTreeClassifier(criterion = 'entropy', max_dept = 5000) where the criterion parameter sets the way the quality of the split function is evaluated. We have used entropy which sets the criterion based on informational gain. The max_dept parameter indicates the maximum depth of the decision tree. In our case, a depth of 5000 was set.
- RandomForestClassifier($max_dept = 1000$ $n_estimators = 1000$), where the max_dept parameter has the same meaning as in the previous case and has been set to a depth of 1000, while $n_estimators$ represents the number of trees belonging to the forest, in our case 1000.

The dataset construction and the classifiers have

been built in Python, using the open-source scikitlearn library for machine learning algorithms and the Seglearn library to perform windows generation with different sizes and overlaps.

4.4 Validation

To validate the performance of the different classifiers, we have used the metrics reported in Table 1. More specifically, the first column contains the name of the metrics, the second the formula to calculate it, and the third a brief description.

5 DISCUSSION OF RESULTS

In this section, we report the results obtained. Specifically, we have conducted, for each classification algorithm, five different experiments, with a different set of features provided in input to the classifiers:

- original dataset, as provided to us by the CNOS center;
- addition of the base frequency (i.e., the lowest frequency of each window) to the windows samples;
- addition of the frequency range (i.e., the lowest and the highest frequencies of each window) to the windows samples;
- addition of all the frequencies to the windows samples (hence doubling the features);
- original dataset, as provided to us by the CNOS center, extended with data augmentation (hence doubling the samples).

In the second, third, and fourth experiments, we have added more information to allow the classifier to learn relationships among the peaks in the spectrum to the frequencies where they happen.

For each experiment conducted, the results are shown in Tables 2, 3, 4, 5, 6. Each table shows the

results obtained with both classifiers used, the Decision Tree and the Random Forest. Respectively, the first six columns refer to the first classifier, and those following to the second. For each experiment, we report the four cases in which we found the best results.

More specifically, in the first two columns, colored in yellow, we report the parameters we have set for the windows size and overlap, and in the following columns the metrics we used for model validation. In particular, accuracy, precision, recall, and F measure.

Therefore, Table 2 reports the results obtained by training and evaluating the model on the starting dataset, containing the progressive number representing the cell number and the measurement number, the amplitudes corresponding to the various wavenumbers, and the type of cell (tumor or non-tumor). As you can see in the table, with the Decision Tree the best performances have been obtained by setting the window size to 85 and an overlap equal to 50%. In this case, the model has an accuracy of 70%, a precision of 75%, recall of 77%, and F measure of 76%.

With the Random Forest, the results are better, the F-measure oscillates between 85% and 86% for all four of the best cases reported. In particular, with this classifier, the best results were obtained by setting the window size to 60 and considering a null overlap. In this case, the validation metrics are respectively 82%, 83%, 88%, and 85%. These results attest that the classifier, specifically Random Forest, albeit with a modestly sized dataset, can classify the instances with some success.

In Table 3 we show the results obtained for the second experiment. In particular, in addition to the information previously described, we have added the lowest frequency of each window as an additional feature. With these input data, the Decision Tree classifier has got the best results with window size 80 and null overlap, it reaches an F-score equal to 74%. Therefore, compared to the previous experiment, in this case, the additional information did not help, but led to a lowering, albeit minimal, of the model validation metrics.

This did not happen in the case of Random Forest, where the F-score reached 86% in the best case. In particular, the best case is the one with a window size of 60 and null overlap.

In the third experiment, the frequency range (i.e., lowest and highest frequencies of the windows) have been added as additional features. Table 4 shows that in the case of the Decision Tree there is still no increase in the F-score, this being almost 75% in the best case, the one with a window size of 50 and zero overlap. In the original dataset, however, the highest F-score with this model was 76%. With the Ran-

dom Forest instead there continues to be a slight increase in the best case (i.e., window size 75 and zero overlap). Specifically, it shows a slight improvement in the F-score equal to 86% and a strong increase in the Recall that reaches 91%. Therefore, these results show that with the addition of this information Random Forest is able to learn relationships among peaks and the frequencies better than other classifiers, managing to further minimize the classification errors of the input instances.

Table 5 instead shows the results related to the fourth experiment where all the frequencies corresponding to each window have been added as additional features (doubling the features). As in the two previous cases, for the Decision Tree, there are no substantial variations in the scores obtained for the validation metrics, which remain stable. On the contrary, with Random Forest we continue to have a small increase in the F-score. In fact, in the best case, with a window size equal to 75 and without overlap, we obtain an increase of 0.3% compared to the experiment conducted including the frequency intervals. These improvements, albeit small, say that the classifier is improving its performance during the training phase.

Finally, the last experiment has been conducted using the data augmentation technique, the results of which are shown in Table 6. The results show a decisive improvement in the case of the Decision Tree, which in the best case with a window size of 70 and without overlap obtains a substantial increase in accuracy, going from 70% on the initial dataset to 82% in this case. The other metrics, on the other hand, remain stable. This is also true for the Random Forest, which in the best case has a window size of 75 and an overlap of 50%. With these parameters we obtain an accuracy of almost 90%, precision almost 81%, recall 84%, and F-score 82%.

Therefore, it is possible to note that among the classifiers chosen, the one that best fits the data is the Random Forest, which in all five experiments conducted obtained the best results with an overlap equal to 0 and which is the best performance ever. were obtained in the fourth experiment, where the classifier received the starting dataset as input with the addition of all the frequencies of the window under examination.

6 THREATS TO VALIDITY

The proposed study suffers from three types of threats to validity: internal, external, and constructive.

Threat to internal validity could be classification errors due to incorrect data labeling. This risk is

	Decision Tree						Random Forest						
Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score	Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score		
85	0	68,95%	72,84%	77,29%	75,00%	90	0.5	81,17%	82,71%	87,41%	84,99%		
90	0.5	70,03%	76,14%	74,07%	75,09%	75	0	81,32%	80,86%	90,39%	85,36%		
80	0	69,47%	73,84%	76,42%	75,11%	55	0	81,90%	82,48%	88,95%	85,59%		
85	0.5	70,33%	75,00%	77,04%	76,00%	60	0	82,25%	83,11%	88,66%	85,79%		

Table 2: Results on the original Dataset.

Table 3: Results with the base Frequency.

	Decision Tree							Random Forest							
Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score	Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score				
80	0.75	68,42%	71,89%	74,94%	73,38%	75	0.75	80,89%	78,17%	92,41%	84,69%				
65	0.75	68,87%	72,20%	75,31%	73,72%	75	0	81,32%	80,86%	90,39%	85,36%				
75	0.75	69,91%	72,83%	75,60%	74,19%	55	0	82,43%	82,97%	89,24%	85,99%				
80	0	68,95%	74,03%	74,67%	74,35%	60	0	82,60%	83,75%	88,37%	86,00%				

Table 4:	Results	with	the	interval	Frec	uency.

Decision Tree						Random Forest						
Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score	Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score	
75	0.75	69,42%	72,79%	74,30%	73,54%	75	0.75	80,58%	77,91%	92,19%	84,45%	
75	0	68,95%	74,67%	73,36%	74,01%	60	0	82,25%	83,47%	88,08%	85,71%	
80	0.75	69,48%	72,95%	75,42%	74,17%	55	0	82,43%	82,80%	89,53%	86,03%	
50	0	70,33%	77,37%	72,59%	74,90%	75	0	82,37%	81,64%	91,27%	86,19%	

Table 5: Results with all Frequencies.

	Decisio	n Tree	Random Forest								
Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score	Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score
50	0	69,28%	76,52%	71,60%	73,98%	80	1	80,77%	78,56%	91,59%	84,57%
60	0	68,89%	74,78%	73,26%	74,01%	55	0	82,07%	82,70%	88,95%	85,71%
75	0.75	70,22%	73,46%	75,05%	74,25%	60	0	82,25%	83,47%	88,08%	85,71%
80	0	68,95%	74,03%	74,67%	74,35%	75	0	82,63%	81,71%	91,70%	86,42%

Table 6: Results with Data Augmentation.

	Decision Tree							Random Forest						
Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score	Windows Size	Overlap	Accuracy	Precision	Recall	F1-Score			
75	0	83,38%	69,19%	70,19%	69,69%	85	0.75	89,97%	83,36%	81,18%	82,25%			
85	0.5	82,56%	68,59%	71,65%	70,09%	85	1	89,76%	80,65%	84,13%	82,35%			
75	0.5	82,25%	66,60%	74,19%	70,19%	90	1	89,82%	80,51%	84,43%	82,42%			
70	0	82,39%	69,18%	72,14%	70,63%	75	0.5	89,98%	80,98%	84,19%	82,55%			

strongly mitigated because the data set used was provided by a specialized center (i.e., the Center for Nanophotonics and Optoelectronics for Human Health (CNOS)) which analyzed the cells of a patient under treatment at a known institute of national prestige (National Cancer Institute IRCCS G. Pascale Foundation).

On the other hand, the generalization of the results is about the threat to external validity. A limitation of the study is represented by the classification carried out on the cells of a single patient, for which the dataset does not contain a very high number of instances. To avoid this threat, we have used the data augmentation technique.

Finally, threats to construct validity could be rep-

resented by inaccuracies or omissions made during the construction phase of the dataset. To mitigate this problem, the cells have been analyzed using Raman spectroscopy.

7 CONCLUSIONS AND FUTURE WORK

This paper addressed an important issue because nowadays oncological diseases represent the leading cause of death in the world. Developing a system that can help the oncologist in the evaluation of tumor markers, can lead soon to tools that greatly speed up the time in diagnosing the pathology. The study carried out on real cells opens the door to "diagnosis in real-time", through detection using the Raman spectrum, a non-invasive and non-destructive technique for the patient.

The proposed approach, based on the combination of Raman spectroscopy and the use of machine learning models, allows obtaining data on the patient's cells to be identified as "malignant" or not in a matter of minutes. This methodology does not aim to replace the work of the doctor who remains at the center of the diagnosis and treatment process but is a tool made available to him.

The main contribution of this work consists in the use of a dataset containing real information about a patient under treatment at the National CancerInstitute IRCCS G. Pascale Foundation, whose cells have been analyzed by the Center for Nanophotonics and Optoelectronics for Human Health (CNOS).

Therefore, the proposed approach has been tested on an overall dataset containing 364 wavenumbers where each corresponds to a sample of amplitude across the various records of the dataset. The results show good performance of the Random Forest Classifier which in the case of data augmentation reached an accuracy of 89.98%.

The limitation of the study concerns the fact that the classification was carried out on cells relating to a single patient, because some were collected in the tumor area, and others in adjacent but healthy areas.

So, in the future it could be interesting to investigate in three different directions: classification of cells from different patients but with the same pathology to assess whether the pathology has similar traits in different patients (i); classification of cells from different patients but with different tumor pathologies to evaluate if there is an indicator, that is a set of biological components, common for all oncological pathologies (ii); classification of cells from healthy patients and patients suffering from oncological diseases to understand if some tumor traits are also present in healthy patients, avoiding the disease with effective prevention therapy (iii).

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