Characterization and classification of numerical data patterns using Annotated Paraconsistent Logic and the effect of contradiction

Caracterização e classificação de padrões de dados numéricos utilizando a Lógica Paraconsistente Anotada e o conceito da contradição

Caracterización y clasificación de patrones de datos numéricos utilizando Lógica Paraconsistente Anotada y el concepto de contradicción

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Abstract
This work describes the development of a computational mathematical model that uses Annotated Paraconsistent Logic - APL and a concept derived from it, the effect of contradiction, to identify patterns in numerical data for pattern classification purposes. The APL admits paraconsistent and paracomplete logical principles, which allow the manipulation of inconsistent and contradictory data, and its use allowed the identification and quantization of the attribute related to the contradiction. To validate the model, series of Raman spectroscopies obtained after exposure of proteins, lipids and nucleic acids, collected from cutaneous tissue cell samples previously examined for the detection of cancerous lesions, identified as basal carcinoma, melanoma and normal, were used. Initially, the attributes related to contradiction, derivative and median obtained from spectroscopies were identified and quantified. A machine learning process with approximately 31.6% of each type of samples detects a sequence of spectroscopies capable of characterizing and classifying the type of lesion through the chosen attributes. Approximately 68.4% of the samples are used for classification tests. The proposed model identified a segment of spectroscopies where the classification of test samples had a hit rate of 76.92%. As a differential and innovation of this work, the use of APL principles in a complete process of training, learning and classification of patterns for numerical data sets stands out, with flexibility to choose the attributes used for the characterization of patterns, and a quantity of samples of about one third of the total required for characterization.

Keywords: Annotated paraconsistent logic; Machine learning; Pattern classification; Mathematical model; Contradiction effect.

Resumo
Este trabalho descreve o desenvolvimento de um modelo matemático computacional que utiliza a Lógica Paraconsistente Anotada - LPA e um conceito derivado dela, o efeito da contradição, para identificar padrões em dados numéricos para fins de classificação de padrões. A LPA admite princípios lógicos paraconsistentes e paracompletos, que permitem a manipulação de dados inconsistentes e contraditórios, e sua utilização permitiu a identificação e quantização do atributo relacionado à contradição. Para validação do modelo, foram utilizadas séries de espectroscopias Raman obtidas após exposição de proteínas, lipídios e ácidos nucléicos, coletados de amostras de células de tecido cutâneo previamente examinadas para detecção de lesões cancerígenas, identificadas como:
Um processo de aprendizado de máquina com características de padrões, e uma quantidade de amostras de cerca de um terço do total necessário para a caracterização. LPA soporta princípios lógicos paraconsistentes e paracompletos, que permitem a manipulação de dados inconsistentes e contraditórios, e seu uso permitiu a identificação e quantificação do atributo relacionado com a contradição. Para validar o modelo se utilizaram serie de espectroscopia Raman obtidas tras a exposição de proteínas, lípidos e ácidos nucleicos, recolectadas de muestras de células de tejido cutâneo previamente examinadas para la detección de lesiones cancerosas, identificadas como basal, melanoma y carcinoma normal. Se identificaron y quantificaron los atributos relacionados con la contradição, la derivada y la mediana obtenidos en la espectroscopía. Un proceso de aprendizaje automático con el 31,6% de las muestras detecta una secuencia espectroscópica capaz de caracterizar y clasificar el tipo de lesión a través de los atributos elegidos. El 68,4% de las muestras se utilizan para la clasificación. El modelo identificó un segmento de espectroscopia donde la clasificación de las muestras de prueba tenía una tasa de aciertos del 76,92%. Como diferencial e innovación de este trabajo, se destaca el uso de principios LPA en un proceso completo de entrenamiento, aprendizaje y clasificación de patrones para conjuntos de datos numéricos, con flexibilidad para elegir los atributos utilizados para caracterizar los patrones, y una serie de muestras de cerca de un terço do total necessário para a caracterização.

**Palavras-chave:** Lógica paraconsistente anotada; Aprendizado de máquina; Classificação de padrões; Modelo matemático; Efeito da contradição.

**Resumen**

Este artículo describe el desarrollo de un modelo matemático computacional que utiliza lógica paraconsistente anotada - LPA y un concepto derivado de él, el efecto de contradicción, para identificar patrones en datos numéricos con fines de clasificación de patrones. LPA soporta principios lógicos paraconsistentes y paracompletos, que permiten la manipulación de datos inconsistentes y contradictorios, y su uso permitió la identificación y cuantificación del atributo relacionado con la contradição. Para validar el modelo se utilizaron serie de espectroscopia Raman obtenidas tras la exposición de proteínas, lípidos y ácidos nucleicos, recolectadas de muestras de células de tejido cutáneo previamente examinadas para la detección de lesiones cancerosas, identificadas como basal, melanoma y carcinoma normal. Se identificaron y quantificaron los atributos relacionados con la contradição, la derivada y la mediana obtenidos en la espectroscopía. Un proceso de aprendizaje automático con el 31,6% de las muestras detecta una secuencia espectros cópica capaz de caracterizar y clasificar el tipo de lesión a través de los atributos elegidos. El 68,4% de las muestras se utilizan para la clasificación. El modelo identificó un segmento de espectroscopia donde la clasificación de las muestras de prueba tenía una tasa de aciertos del 76,92%. Como diferencial e innovación de este trabajo, se destaca el uso de principios LPA en un proceso completo de entrenamiento, aprendizaje y clasificación de patrones para conjuntos de datos numéricos, con flexibilidad en la elección de atributos utilizados para caracterizar los patrones, y una serie de muestras de un terço do total requerido para caracterizar.

**Palabras clave:** Lógica paraconsistente anotada; Aprendizaje automático; Clasificación de patrones; Modelo matemático; Efecto de contradicción.

**1. Introduction**

**1.1 The Annotated Paraconsistent Logic – APL**

Classical Logic is founded in such a way that, given a proposition, it can assume two states, being then considered a binary logic. This condition limits the representation of numerous real-world situations, especially those that imply decision-making based on complex problems. In researches related to Artificial Intelligence, systems projects are studied based on more flexible types of logic and, therefore, adaptable to the complex problems of the real world. Non-Classical Logics admit in their foundations indefiniteness, ambiguities and contradictions, being more adequate to represent complex systems. Paraconsistent Logic belongs to the Non-Classical Logic family and admits contradictory states in its theoretical structure (Da Silva Filho et al., 2008).

Studies on the fundamentals of Annotated Paraconsistent Logic – APL, are described by Da Costa, Subrahmanian and Vago (Da Costa et al., 1991). To represent the admitted states of the APL, a lattice with four vertices can be used, with the so-called constants of annotation in each of the vertices: $V$ - true, $F$ - false, $\perp$ - Inconsistent, $\bot$ - Paracomplete or Indeterminate. Given the set of these constants $\tau = \{ T, V, F, \perp \}$ and the lattice operator $\tau = \leq | \perp $, $\leq , \sim >$ which can be characterized by a Hasse diagram. The operator over $\tau$ is: $\sim : | \perp $ $\rightarrow | \perp $, whose operations are described next to the lattice (Da Silva Filho et al., 2008)- Figure 1.
Figure 1: lattice with 4 vertices (from Hasse) and the states of Paraconsistent Logic.

\[ \sim T = T \] (the negation of an inconsistent proposition is inconsistent)
\[ \sim V = F \] (the negation of a true proposition is false)
\[ \sim F = V \] (the negation of a false proposition is true)
\[ \sim \bot = \bot \] (the negation of a paracomplete proposition is paracomplete)

Source: Authors (2021).

We observe in the lattice the true and false states present also in Classical Logic, and the states that are additionally admitted by the Paraconsistent Logic, inconsistent and paracomplete, whose complements or negation are the values of these states.

1.2 The Paraconsistent Logic Annotated with an Annotation of Two Values – APL2v

APL2v allows the representation of two values \( \mu \) and \( \lambda \) for the states of Paraconsistent Logic related to a given proposition, being \( \mu \) the favorable Evidence Degree and \( \lambda \) the unfavorable Evidence Degree, and thus express with greater amplitude the knowledge about a proposition. The representation in the lattice is done such that \( \tau = \{ (\mu, \lambda) \mid \mu, \lambda \in [0, 1] \subset \mathbb{R} \} \). The operator about \( \tau \) is: \( \sim: |\tau| \to |\tau| \), and:
- \( \bot \) indicates the minimum of \( \tau = (0, 0) \);
- \( T \) indicates the maximum of \( \tau = (1, 1) \).

A four-vertex lattice to represent APL2v is shown in Figure 2, where a proposition is represented by \( P \), the degree of evidence supporting a proposition is represented by \( \mu \), and the degree of evidence unfavorable to a proposition is represented by \( \lambda \) (Da Silva Filho et al., 2008).

Figure 2: four-vertex lattice (from Hasse) and APL2v states.

APL2v then admits two annotations or evidence for each of the APL states, which can vary between the numerical values 0 and 1. When each of the states is effectively characterized, that is, at the vertices, the evidence in favor and against a proposition, represented respectively by \( \mu \) and \( \lambda \), assume the maximum or minimum values.

1.3 Paraconsistent Algorithms

Algorithms that express the principles of Annotated Paraconsistent Logic can be structured through modules or blocks that have inputs that are the evidence for or against a given proposition, and one or more outputs corresponding to the paraconsistent analysis. All numerical values that express the inputs are normalized between 0 and 1. The structuring of APL in blocks allows them to be interconnected, forming Paraconsistent Analysis Networks, so that the output of one block can be the input of another, that is, the resulting paraconsistent analysis becomes the input evidence for a subsequent block. A paraconsistent analysis block is shown in Figure 3.
Figure 3: Representative block of the paraconsistent analysis.

In the paraconsistent analysis block, the degrees of favorable and contrary evidence enter through inputs $\mu$ e $\lambda$, while the output $S$ presents a resultant degree of evidence that depends on the paraconsistent algorithm that is implemented inside the block, and some of these algorithms will be described in this job.

1.4 The representation of certainty and contradiction in APL2v

Through the representative lattice of APL2v shown in Figure 4, it is possible to describe how values related to the evidence of certainty and contradiction of a given proposition are quantified, through the so-called Degree of Certainty – $D_C$ and Degree of Contradiction - $D_{Ct}$ (Da Silva Filho et al., 2008).

Figure 4: The representative lattice of APL2v.

From the Degree of Certainty and the Degree of Contradiction, it is possible to define the so-called Interval of Certainty - $\phi$, which is an indicator of how much certainty or contradiction, originated by the favorable Degree of Evidence - $\mu$, and the Degree prevail in the paraconsistent analysis of Unfavorable Evidence - $\lambda$, of a given proposition. The Certainty Interval is also accompanied by a sign (+) indicating that the $D_{Ct}$ tends to the True logic state or (-) indicating that the $D_C$ tends to the False logical state. The Certainty Interval is defined as:

$$\phi_{(a)} = 1 - |D_{Ct}|$$

(3)
1.5 Paraconsistent Analysis Algorithm

In a paraconsistent analysis, the Real Degree of Certainty - RDC, is the result of the analysis where, at the end of it, it is desired to present a pure certainty value, that is, without influence of the effects of uncertainties. In calculating the Degree of Real Certainty, the result of the paraconsistent analysis must be subtracted from the value attributed to the effect of the influence of inconsistencies arising from conflicting information, free from the effect of contradiction (Da Silva Filho et al., 2008). Figure 5 shows the APL2v lattice with the intersection point between DC and DCT. For an analysis with null DCT, the DC value would be maximum and equal to 1. Since the distance between the maximum DC value and the intersection point is called d, the projection of d on the Degrees of Certainty axis is the value of Degree of Real Certainty - DRCC. The Resulting Degree of Evidence - μER is the normalized DRCC value. The Resulting Evidence Interval - φE represents, in the paraconsistent analysis, the same concept demonstrated by the Certainty Interval defined in equation (3), but it is calculated from the Degree of Contradiction normalized between values 0 and 1, called in the algorithm of μCTR.

Figure 5: APL2v lattice with the projection of the Degree of Real Certainty - DRCC.

The lattice shows the Degree of Certainty and the Degree of Contradiction, the distance d between the maximum value of DC and the intersection point of the degrees of certainty and contradiction, and the projection of the distance d on the DC axis, which corresponds to the Degree of Real Certainty. Next, Algorithm 1 of paraconsistent analysis.

Algorithm 1: paraconsistent analysis - start:
1. Insert favorable evidence μ and contrary λ of the paraconsistent analysis;
   \[ 0 \leq \mu \leq 1; \quad 0 \leq \lambda \leq 1; \]
2. Calculate the Degree of Certainty - DC:
   \[ DC = \mu - \lambda; \]
3. Calculate the Degree of Contradiction - DCT:
   \[ DCT = (\mu + \lambda) - 1; \]
4. Calculate the Degree of Evidence - μE (for \( \lambda = 1 - \lambda \)):
   \[ \muE = \frac{(\mu - (1 - \lambda) + \lambda)}{2} = \mu + \frac{\lambda}{2}. \]
5. Calculate the distance - d:
   \[ d = \sqrt{1 - |GC|^2} + GCT^2; \]
6. Calculate the Normalized Degree of Contradiction - μCTR (for \( \lambda = 1 - \lambda \)):
   \[ \muCTR = \frac{GC + \lambda}{2}. \]
7. Calculate the Resulting Evidence Interval - φE:
   \[ \phiE = 1 - |2\cdot\muCTR - 1|; \]
8. Determine outputs S1 and S2:
if \( q_{E} \leq 0.25 \) or \( d > 1 \), so:
\[
\{ S_1 = 0.5; S_2 = q_{E(i+1)} \}; \quad \text{go to step 11;}
\]
otherwise go to the next step;

9. Determining the Degree of Real Certainty - \( D_{RC} \):
\[
\begin{align*}
& \text{if } G_{C} > 0, \text{ so: } G_{CR} = 1 - d; \\
& \text{if } G_{C} < 0, \text{ so: } G_{CR} = d - 1;
\end{align*}
\]

10. Calculate the Resulting Degree of Evidence - \( \mu_{ER} \):
\[
\mu_{ER} = \frac{G_{CR} + 1}{2}
\]

**Algorithm 1 - paraconsistent analysis - end.**

1.6 Algorithm Extractor of the Effects of Contra
diction

The Contradiction Effects Extractor algorithm is used when there is a set of evidence for a given proposition subject to contradictions, and through the algorithm a paraconsistent analysis of this set of evidence is performed, subtracting the effects caused by the contradictions and presenting a single level of evidence resulting representative of the whole. The resulting numerical value is a Degree of Certainty from which the effects of the contradiction are subtracted. Algorithm 2 used in the contradiction effects extraction process is described below:

**Algorithm 2: Contradiction Effects Extractor – beginning:**

1. Present the Degrees of Evidence of the study group - 0,0 \( \leq \mu \leq 1,0 \):
\[
G_{\text{est}} = \{ \mu_A, \mu_B, \mu_C, \ldots, \mu_n \};
\]

2. Select the highest Degree of Evidence value of the study group:
\[
M_{\text{max}} = \max \{ \mu_A, \mu_B, \mu_C, \ldots, \mu_n \};
\]

3. Select the lowest Degree of Evidence value of the study group:
\[
M_{\text{min}} = \min \{ \mu_A, \mu_B, \mu_C, \ldots, \mu_n \};
\]

4. Perform steps 1 to 10 of the paraconsistent analysis and return the Resulting Degree of Evidence between the selected maximum and minimum values, \( \mu_{\text{max}} \) and \( \mu_{\text{min}} \):
\[
\mu_{R1} = M_{\text{max}} \bigcirc M_{\text{min}};
\]

5. Add the obtained value \( \mu_{R1} \) for the evidence group under study excluding the maximum values - \( \mu_{\text{max}} \) and minimum - \( \mu_{\text{min}} \) previously selected:
\[
G_{\text{est}} = \{ \mu_A, \mu_B, \mu_C, \ldots, \mu_n, \mu_{R1} \};
\]

6. Return to item 2 until the study group has only one element, then considering the resulting value of the analysis.
\[
G_{\text{est}} = \mu_{\text{ER}};
\]

**Algorithm Extractor of the Effects of Contradiction - end.**

1.7 Algorithm for quantizing the attributes of the spectroscopy series

The training and learning process values attributes of the spectroscopy series, and one of the attributes, the absence of contradiction, is obtained from the principles of APL2v.

Sequential spectra are expressed by \( x_i \) and \( x_{(i+1)} \), where \( i \) is the position of each spectrum in the dataset, the following attributes will be calculated: increasing slew rate, decreasing slew rate and zero slew rate; the median that represents an approximation to the mean value of the series, while the resulting value of the contradiction extractor asserts a Degree of Certainty to the dataset free from the effects of the contradiction. These attributes are calculated as described in Algorithm 3, presented below:

**Algorithm 3: Quantization of spectroscopy series attributes - start:**

given the variables: \( a = \) increasing rate of change; \( b = \) decreasing rate of change; \( c = \) zero slew rate; \( d = \) Degree of Certainty free from the effects of contradiction; \( e = \) average;

\( a = 0; b = 0; c = 0; \)

1. Calculating the increasing rate of change:
for \( i = 0; i < 10; i = i + 1 \)
describe how the potential uncertainty and inconsistency, and with Paraconsistent Logic, the analysis of data obtained by Raman spectroscopy, data segments where their attributes can be quantized and, in this way, patterns for these series can be characterized. Demonstrate the development of a computational model based on Paraconsistent Logic and that uses a machine learning process; to present a characteristic of non-classical logic, the contradiction, to be quantized in the segments identified in the spectroscopy series. To present a

\[ \begin{align*}
\text{1.8 The use of APL2v in intelligent systems} & \\
\text{Some works that use some of the APL2v principles proposed in this research are described below. Theoretical concepts of Paraconsistent Logic originated in Philosophy and that derive states such as inconsistency and contradiction have been approached in different works so that its principles are used in different areas of Science and Engineering (Priest, 2002; Norihiro & Heinrich, 2012; Carnielli et al., 2018). Studies involving Artificial Neural Networks based on Paraconsistent Logic suggest that they can be applied in decision systems that use pattern recognition (Martínez et al., 2021). An autonomous mobile module using ultrasound distance sensing, and whose trajectory is defined from inconsistencies and contradictions valuations, demonstrated that the APL2v principles could be used beyond theoretical limits (Da Silva Filho et al., 2001). A model to analyze cephalometric variables in orthodontics (Mario et al., 2010), was developed from Paraconsistent Artificial Neural Networks (Mario, 2003), and describes how the potential use of APL2v can be expanded to the medical field. An architecture that uses Paraconsistent Artificial Neural Cells for Learning – PANCL (Mario, 2003) and the algorithm that extracts the effects of contradictions, which, given a data set representative of evidence, returns a single value that is equivalent to the absence of contradiction in a data set (Da Silva Filho et al., 2011), are two principles used to demonstrate the concept of a filter that analyzes the signal-to-noise ratio in telecommunications (Carvalho Jr. et al., 2018). The treatment of signals equivalent to a proportional and integral controller used in an automation system, is structured through the concepts of APL2v, and promising results were obtained compared to a conventional feedback control (Coelho et al., 2019).}
\\
\text{More recent theoretical studies evaluate the use of Artificial Neural Networks in conjunctio} & \\
\text{with Paraconsistent Logic for use in decision-making systems (Martínez et al., 2021).}
\\
\text{The attributes of Paraconsistent Logic were used in the analysis of data obtained by Raman spectroscopies to discriminate cutaneous tissue samples, and in the first work in this area a paraconsistent analysis network was described and implemented for this purpose (Da Silva Filho et al., 2016), and a second work compared the result obtained from the use of LPA2v with a method that uses multivariate statistics based on principal component analysis – PCA (Garcia et al., 2019). Both works showed promising results.}
\\
\text{Demonstrating the scope of Paraconsistent Logic, its principles have also been used to represent Quantum Mechanics phenomena (Da Costa & De Ronde, 2013; Da Silva Filho, 2016).}
\\
\text{1.9 Objective} & \\
\text{The model must identify, in numerical series resulting from Raman spectroscopy, data segments where their attributes can be quantized and, in this way, patterns for these series can be characterized. Demonstrate the development of a computational model based on Paraconsistent Logic and that uses a machine learning process; to present a characteristic of non-classical logic, the contradiction, to be quantized in the segments identified in the spectroscopy series. To present a}
\end{align*} \]
decision-making system that selects the segments with the best assertiveness for pattern classification and through which it is possible to correlate the patterns characterized in the machine learning process to the series of spectroscopy that did not participate in the learning process, through tests of generalization.

2. Methodology

The development of this work is related to an experimental and quantitative research, whose numerical data analyzed were made available from the methodology used in the experiment developed by Silveira et al. (2012). For the construction of the computational mathematical model, which will be referred to as computational model, a Java programming language was used. Spectroscopy data were stored in an EXCEL® spreadsheet. The methodology used in the work can be described through the following sequence of procedures:

a) With the availability of spectroscopy for NORMAL, CBC and MEL classifications, the computational model searches for these data according to the classification and then samples are separated for characterization and samples for tests. The sampling method used is based on holdout (Faceli et al., 2019), as the training and test data do not mix, but because of the availability of samples, approximately one third of the total pre-classified samples is used for training, and the rest for testing. The consequences for this type of procedure are discussed in Baranauskas and Monard (2000);

b) The computational application uses the separate samples for the characterization to quantize the chosen attributes, through sequential segments of 20 wavenumbers. A specific algorithm in the computational model performs a scan so that all segments have quantized spectroscopies. The same process is performed for test samples;

c) Considering that the data analyzed in this work are multivariate, that is, they have more than one attribute, in general, covariance and correlation measures would be used to analyze the relationship between more than two attributes (Faceli et al, 2019). As a differential, this work uses a Paraconsistent Analysis Network to calculate the evidence of similarity between the attributes of the characterization and test data, after their quantization in the previous step. The evidence of similarity is calculated for sequences of 20 wavenumbers, until the calculation is carried out for all;

d) Then, all calculated values of evidence of similarity are exported to an EXCEL® spreadsheet, indexed with the respective segments, and a decision system implemented in the spreadsheet identifies the segments where the assertiveness of the correct classification between the training and test segments is greater than 75%.

Series of spectroscopies obtained through the methodology described in the work developed by Silveira et al. (2012) were used, which describes the use of Raman spectroscopy in cutaneous tissue biopsy fragments in the infrared range with a wavelength of 830 nm, power of 50mW to 200mW and exposure of 20 seconds. Of the total samples, 96 have basal carcinoma classification - CBC, 19 samples have melanoma classification - MEL and 30 samples have normal cell classification - NORMAL. The numerical threshold values of the spectroscopy are within an approximate range of the minimum value -0.6 and the maximum value +0.6. The average values of the spectroscopy sets for each of the three classifications used in the learning process are shown in Graph 1.
**Graph 1:** Mean values of Raman spectroscopies used to build the reference standards for the classification of skin lesions of the types NORMAL, MEL and CBC.

The graph shows that the Raman spectroscopy wave numbers for this experiment are in the range between 378.4 cm\(^{-1}\) and 1779.6 cm\(^{-1}\). For some wavenumber intervals the Raman spectroscopy values are very close, and in other intervals they are quite different for the NORMAL, CBC and MEL classifications.

The machine learning process performed by the computational model uses reference samples to characterize the CBC, MEL and NORMAL patterns, and simultaneously uses test samples to identify the segments of the wavenumber series with greater precision in identifying the patterns. The total number of samples of each type of classification was equal to the amount of MEL spectroscopy available - 19. Of this total, the averages of 6 samples, approximately 31.6%, were used to characterize the reference samples, while 13 samples or approximately 68.4% of the total were used for testing. Figure 6 is a representation of how the available samples were used to build the computational model.
The figure shows a representation of the Raman spectroscopy values stored in an EXCEL® spreadsheet. The first three columns on the left show the values of the samples used for training for the NORMAL, CBC and MEL classifications. Six samples were used for each type of classification, corresponding to wave numbers between 378.4 cm\(^{-1}\) and 1779.6 cm\(^{-1}\). The columns on the right are a representation of the 13 samples submitted for testing for each of the ratings and same wavenumbers.

### 2.1 Paraconsistent Analysis Network

For the characterization of the standards of the reference and test samples, a Paraconsistent Analysis Network is used. The network architecture presents two sets of symmetric blocks, each one to receive reference and test pairs of 10 wavenumber, so that the analysis carried out by the network totals a total of 20 reference and test pairs of wavenumber. Through the inputs \(\mu_1\) and \(\lambda_1\) of the Paraconsistent Analysis Network, the increasing rate of change parameter is quantified. Via the \(\mu_2\) and \(\lambda_2\) inputs, the parameter equivalent to the decreasing rate of change is quantified. Inputs \(\mu_3\) and \(\lambda_3\) quantify the null rate of change. Inputs \(\mu_4\) and \(\lambda_4\) quantify the value equivalent to the extraction of the contradiction. Through the \(\mu_8\) and \(\lambda_8\) inputs, they quantify the median (Larson & Farber, 2010) of the wavenumber sequence. Output S5 is the resulting value of the increasing and decreasing rates of slew, while output S6 adds the null rate of slew to this analysis. Output S7 adds to the analysis the extraction of contradiction, which through output S9 adds the median in the paraconsistent analysis. Through the symmetry of the Paraconsistent Analysis Network, the analyzes referring to 10 spectra are inserted in the \(\mu\) and \(\lambda\) inputs of the last block of the network. For the two analyzed segments to have equivalent weights in the last block, the input \(\lambda\) is inserted so that its value is complemented, equivalent to a true evidence, that is, \(\lambda = 1 - \lambda\). The Paraconsistent Analysis Algorithm is then executed from step 1 to step 4. The output of the Paraconsistent Analysis Network presents a value that represents the evidence of how much a segment of 20 test wavenumber has similarity with one of the reference standards, CBC, MEL and NORMAL. The Paraconsistent Analysis Network is shown in Figure 7.
Then there is the Paraconsistent Analysis Network, formed by blocks of paraconsistent analysis, whose algorithms were previously presented, and which allow the interconnection of the values of the numerical data attributes related to Raman spectroscopy, to present in the output a resulting evidence equivalent to similarity between spectroscopy segments, which allows to identify for the pairs of training and test segments if they belong to the same classifications.

2.2 Model functionality to quantify similarities between Raman spectroscopies

The computational model starts scanning the wavenumber segments from the first and increments to the twentieth, with $n_1$ being the identifier of the first spectrum and $n_{20}$ being the identifier of the twentieth spectrum of the first segment. The second segment starts from the second spectrum $n_2$ and has the twenty-first spectrum as its last spectrum, identified as $n_{21}$. Each available spectroscopic series consists of 1178 wavenumbers. As the algorithm scans every 20 wavenumbers, this corresponds to 1158 segments of 20 wavenumbers. Each segment of the wavenumber inserted in the Paraconsistent Analysis Network, presents in the output an evidence of similarity between the segment in relation to the reference standards and the analyzed test segment. This evidence is then directed to a decision-making system, which checks whether the value of evidence between the test and reference spectra segment had a value greater than 0.75 for the standard to which the test segment belongs, CBC, MEL or NORMAL. The decision system nominally identifies all wavenumber segments and has an assertiveness greater than 75%. An overview of the computational model is shown in Figure 8.
**Figure 8:** Overview of the computational model to identify series of wavenumber segments with better performance in pattern characterization.

The figure represents the entire sequence of steps initially described in the methodology, that is, from reading the data in the spreadsheets, quantizing attributes and calculating the similarity between training and test spectroscopy segments, passing through the decision system implemented again in the environment from the spreadsheet, to the identification of the wavenumber segments whose assertiveness in the classification exceeds the 75% rate.

The decision system consists of EXCEL® spreadsheets that receive the resulting evidence values calculated through the Paraconsistent Analysis Network. Through the use of formulas and conditionals applied to the cells that receive the evidence values, the wavenumber segments are identified with greater precision. Figure 9 shows the spreadsheet used to identify the NORMAL classification segments. Similar spreadsheets were used for the CBC and MEL ratings.
Figure 9: decision-making system implemented through EXCEL® spreadsheets.

Source: Authors (2021).

The decision system represented in the figure where it is implemented in the spreadsheet identifies the segments where the assertiveness of the correct classification between the training and testing segments is greater than 75%. The complete functionality of the computational model for identifying the wavenumber segments is described using Algorithm 4:

Algorithm 4: description of the general functionality of the model - top:
1. Reads reference spectroscopies and test spectroscopies;
2. Characterizes and quantifies the values of reference attributes through evidence;
3. Quantify the similarity between reference and test standards for spectroscopy segments;
4. Identify spectroscopy segments with similarity values between reference and test greater than 75%.

description of the general functionality of the model – end.

3. Results and Discussion

An evaluation of the model's performance can be done by analyzing the responses of the Resulting Degrees of Evidence values, in the output of the Paraconsistent Analysis Network, for a segment of spectroscopy. In this section of the network, spectroscopy segments with different performance in terms of output value were selected for each type of classification, for a more comprehensive view of the model's behavior. Then, for each classification related to diagnoses, 6 samples were used to characterize the reference standards and 13 samples for testing. These quantities in the proportion of 33% and 66% respectively, are based on available samples, and in the case of MEL diagnosis, there were a total of 19 samples.

The results are presented in order to demonstrate the general behavior of the model as a function of the spectroscopy segments inserted in it, that is, high and low performance spectroscopy segments, for each type of classification.

Afterwards, after the model had calculated the accuracy for each spectroscopy segment, a specific spectroscopy segment that presented the best performance for the three classifications was identified. In this case, additional samples that did not participate in the machine learning process were used, 11 for the NORMAL classification and 77 samples for the CBC classification.

The model functionality compares the Degree of Evidence resulting from the tested classification with the other two classifications, and the answer is the highest resulting Degree of Evidence among the three classifications. When the highest resulting degree of evidence corresponds to the type of classification tested, the model's answer is correct. Otherwise, the answer is considered wrong.
3.1 Model results for CBC classification

The representative values of the precision of the Degrees of Evidence Resulting from the CBC classification test samples as a function of the Raman wave numbers are shown in Graph 2. For example, for the 20 wavenumbers spectroscopy from 827.8 to 854.4 the precision is 100%, that is, 13 CBC test samples were classified correctly. For the segment of 20 wavenumbers from 947.4 to 972.7 the precision is 84.62%, that is, for 11 of the 13 samples tested the classification was done correctly. It is also possible to observe that there are segments with low precision, for example for the segment that starts in the 1680.1 wavenumber, with an accuracy of 7.7%, that is, only one test sample was classified correctly. In the chart, these mentioned points are indicated with arrows. We can then have a complete view of the model's performance for the CBC classification, for the entire range of wavenumbers.

Graph 2: accuracy of the model for the CBC classification as a function of the Raman wave numbers segments.

![Graph 2](image)

Source: Authors (2021).

Then there is the resulting graph of the decision system for the CBC classification brings the assertiveness rates as a function of the Raman wave numbers, pointing to the beginning of the segments with the best and worst performance.

Graph 3 shows the Resulting Degree of Evidence result for each of the 13 samples classified as CBC for the 20 wavenumber segment, 827.8 to 854.4, where the accuracy is 100%. The Resulting Degree of Evidence is always higher when the standard similarity of the test samples is compared to the CBC reference standard.
Graph 3: Degrees of Resulting Evidence for the 13 CBC test samples, compared with the NORMAL, CBC and MEL reference standards, wavenumber segments range from 827.8 to 854.4.

![Graph 3](image)

Source: Authors (2021).

According to the graph, it can be seen that the result obtained by the Paraconsistent Analysis Network for the 13 test samples previously classified as CBC was correct, as they present a higher resulting value of evidence for the classification to which they belong.

Graph 4 shows the Resulting Degrees of Evidence for the segment of 20 wavenumber segments, 947.4 to 972.7, with an accuracy of 84.62%, that is, for 11 of the 13 CBC samples tested, the classification was done correctly.

Graph 4: Degrees of Resulting Evidence for the 13 CBC test samples, compared with the NORMAL, CBC and MEL reference standards, wavenumber segments from 947.4 to 972.7.

![Graph 4](image)

Source: Authors (2021).

According to the graph, it can be seen that the result obtained by the Paraconsistent Analysis Network for 11 test samples previously classified as CBC is correct, as they present a greater value of resulting evidence for the classification to
which they belong. For sample 6 the result found was the NORMAL classification and for sample 10 the result found was MEL.

Graph 5 shows the Resulting Degrees of Evidence for the wavenumber segments from 1680.1 to 1696.3 with only one test sample 4 correctly classified, accuracy of 7.7%.

**Graph 5:** Degrees of Resulting Evidence for the 13 CBC test samples, compared with the NORMAL, CBC and MEL reference standards, wavenumber segments from 1680.1 to 1696.3.

Therefore, this graph represents the result of a wavenumber segment with low performance: for samples 1 to 3, 5 to 7, 9, and 11 to 13 the classification result was NORMAL. For samples 8 and 10 the classification result was MEL. Only samples 4 have the correct classification, which is CBC.

### 3.2 Model results for NORMAL classification

Samples of test results with NORMAL classification are shown in Figure 6. Initially, the accuracy of the model is shown for the Degrees of Resulting Evidence as a function of the wavenumber segments. Accuracy ranged from 100% to 7.7%. Next, the Resulting Degrees of Evidence results are shown for the segments of 20 wavenumbers from 580.5 to 609.8 with 100% accuracy, from 474.7 to 505 with 76.92% accuracy, and for the segment of wavenumbers that from 1470.1 to 1488.9, with an accuracy of 7.7%.
Figure 6: NORMAL Classification: accuracy and Degrees of Resulting Evidence according to wavenumber segments starting at 580.5, 474.7 and 1470.1.

Figure 6a shows the performance of test samples for NORMAL classification as a function of wavenumber segments. Many segments achieved 100% accuracy. In 6b are the results of the 13 test samples for the wavenumber range starting at 580.5 and all classifications are correct. In 6c we have the results for the wave segment starting at 474.7 and 10 of the 13 samples are correctly classified. Sample 3 is classified as MEL and samples 1 and 6 are classified as CBC. In 6d are the results of the wavenumber segment that starts at 1470.1, and which has a low performance. Only sample 11 is classified correctly. Sample 6 is classified as MEL and the other 11 samples were classified as CBC.

3.3 Model results for MEL classification

Samples of test results with MEL classification are shown in Figure 7. Initially, the accuracy of the model for the Degrees of Resulting Evidence as a function of the wavenumber segments is shown. The accuracy ranged between 92.31% and 7.7%. In the sequence, the results of the Resulting Degrees of Evidence are shown for the segments of 20 wavenumbers from 691.4 to 719.6 with an accuracy of 92.31%, from 1060.9 to 1084.8 with an accuracy of 76.92 %, and for the wavenumber segments starting at 1337.6 to 1358.1, with a precision of 23.07%.
Figure 7: MEL Classification: accuracy Degrees of Resulting Evidence for the spectroscopy segments starting at 691.4, 1060.9 and 1337.6.

Figure 7a shows the performance of test samples for MEL classification as a function of wavenumber segments. Only one segment surpassed the 90% accuracy. In 6b are the results of the 13 test samples for the wavenumber range that starts at 1337.6 and is precisely the one that performs best in the MEL classification. In this segment 12 samples were correctly classified as MEL and only sample 5 was classified as CBC. In 6c we have the results for the wave segment starting at 1060.9 and 10 of the 13 samples are correctly classified. Sample 8 is rated CBC and samples 6 and 9 are rated NORMAL. In 7d are the results of the wavenumber segment starting at 1337.6. Three samples 2, 11 and 12 are classified correctly. Samples 5 to 10 are classified as CBC and samples 1 to 4 as well as 13 are classified as NORMAL.

3.4 Wavenumber segments with the best and worst accuracy for the three classifications

When performing the experiments to evaluate the model's performance for the NORMAL, CBC and MEL classifications, it can be seen that there is a segment, which starts at the wave numbers 1232 to 1253.8, where the model has the best accuracy considering the three ratings, with 84.61% for the NORMAL rating, 92.3% for the CBC rating and 76.92% for the MEL rating. It is also possible to verify a wavenumber segment of spectroscopy, from 1748.4 to 1763.7, with worse accuracy if the three classifications are considered, being 38.46% for the NORMAL classification, 15.38% for the CBC classification and 7.69% for the MEL classification. Figure 8 highlights these spectroscopic segments for the NORMAL, CBC and MEL reference samples. In the sequence of Figure 8, the Resulting Evidence Degrees are presented for the wavenumber segment from 1232 to 1253.8 respectively for the NORMAL, CBC and MEL test samples.
In 8a we have the mean values of samples from NORMAL, CBC and MEL spectroscopy, as a function of wave number. The segment from wave number 1232 stands out, with the best performance for the three types of classification, and the segment starting from wave number 1748.4, this one with the worst performance for the three types of classification. For the segment with the best performance, starting in 1232, we have in 8b the results for the NORMAL classification, with 11 samples classified correctly, in 8c we have the results of the CBC classification, with 12 samples classified correctly, and in 8d the results of the MEL classification, with 10 samples correctly classified.

### 3.5 Tests with samples not used in the machine learning process for the most accurate wavenumber segment

Of the total available samples, 11 samples of the NORMAL classification were not used in the learning process. These samples were tested by the model to verify the response in what was identified as the best performing segment, 1232 to 1253.8. The performance achieved was 81.82%, that is, 9 samples were correctly classified. This result is shown in Graph 6.
Graph 6: Degrees of Resulting Evidence and Degrees of Contradiction Standardized for the 11 NORMAL samples not previously used in the experiment, spectroscopies segment from 1232 to 1253.4.

The graph shows the result of the segment with the best performance, 1232, for 11 additional samples from the NORMAL classification that did not participate in the training events and tests carried out previously, with 9 samples classified correctly. This event corresponds to a generalization test.

A number of 77 samples of the CBC classification not previously used in the learning process were submitted to the model to verify the response in the spectroscopy segment from 1232 to 1253.8. The performance achieved was 84.42%, that is, 65 samples were correctly classified. This result is shown in Graph 7.

Graph 7: Degrees of Resulting Evidence for the 77 CBC samples not previously used in the learning process, spectroscopies segment from 1232 to 1253.4.
The graph shows the result of the segment with the best performance, 1232, for 77 additional samples from the CBC classification that did not participate in the training events and tests carried out previously, with 65 samples classified correctly, that is, all points where the corresponding line of the CBC classification has higher numeric values. This event also corresponds to a generalization test.

3.6 Discussion

The computational model proposed in this work achieved the objectives of identifying spectroscopic segments that could characterize patterns from previously defined attributes, some common to numerical series, such as the derivative and median, and another parameter that was quantized due to the use of LPA, or that is, the extraction of the effects of the contradiction. The possibility of choosing attributes that will characterize patterns can be an advantage over machine learning processes where this type of functionality is not explicit. In the specific case of the data sets used for the experiments in this work, it can be seen that in the segment with the best performance for classification, from frequency 1232 onwards, variation rates with variation rates with very evident differences, which allows us to better characterize this type of attribute to characterize a pattern. On the other hand, in the segment of spectroscopy with the worst performance, from frequency 1748.4, it can be seen that the rates of change are similar, making it difficult to characterize patterns. Figure 9 shows these details.

The use of Annotated Paraconsistent Logic allowed aggregating and quantifying evidence related to the attributes, through the use of a Paraconsistent Analysis Network, as well as quantifying the similarity between the attributes used to characterize the patterns of reference samples and tests of the types of classifications used in the process through the Resulting Degree of Evidence.

The developed model treated the Raman data without any correlation between their spectroscopy values at a given frequency and the respective classification of skin lesions, that is, the spectroscopies were treated only as numerical series, and from these the model quantified the similarity between the reference samples and tests for segments of 20 sequential spectroscopies.

Within the universe of samples available for the characterization of reference and test standards, the model was able to identify Raman spectroscopy segments with minimum accuracy results of 76.92% for the spectroscopy segment with the best performance for sample classification of tests.
Figure 9: Details of the wavenumber segments 1232 to 1253.8 and 1748.4 to 1763.7.

The graph represents a sampling of spectroscopy as a function of wave numbers, highlighting the segments where the wave numbers with the best performance for correct classification, 1232, and with the worst performance, 1748.4, begin. In the segment started in 1232, significant differences can be seen, such as the attribute of the derivative, favoring the classifications. In the segment that begins in 1748.4, the variations are similar for the three types of classification, making it difficult to quantify the differences and, consequently, carry out the classifications.

4. Conclusion

Through this work, it was possible to demonstrate the development of a computational model to identify parts of a data set that can characterize patterns with better performance based on predefined attributes. This is a differentiated approach within existing machine learning processes, firstly because it is possible to define attributes based on the characteristics of the data and identify specific parts of the set where the use of these attributes allows a characterization and subsequent classification of patterns where the accuracy is higher, anticipating performance and allowing the assessment of whether it is satisfactory for the application and also enabling the redefinition of attributes in search of the possibility of effective classification of patterns.

The use of Annotated Paraconsistent Logic brings the innovation of adding to the model the manipulation of concepts related to Non-Classical Logics, as was the case shown here of manipulating the contradiction present in a dataset. Another innovative aspect related to Paraconsistent Logic is that the Paraconsistent Analysis Network allows variations by aggregating evidence related to attributes, as well as their increment or even substitution. The network architecture can be adapted according to the characteristics of the analyzed data.

As demonstrated in the work, when parts of a dataset with the best performance for classification were identified, the proportion of samples to characterize a pattern, approximately 33% of the total, is significantly lower than in traditional machine learning processes. This is a promising feature of the model, as the amount used in learning can be a variable that can be adjusted in the process for characterizing patterns, as well as the definition of attributes itself.
Another product derived from the model's functionality is the possibility of identifying, from the tests between samples of the data sets, if there will be segments where the precision is significant for the classification.

It is concluded then that the proposed model brings innovations in relation to the already existing machine learning processes used for characterization and classification of patterns, both in the possibility of defining attributes of the data sets, as well as in the flexibility of aggregating these attributes through an architecture of network, in addition to the possibility of adding to the process the quantization of properties that are part of the real world but are difficult to abstract, as was the case of the contradiction, manipulated by the model described here.

Future work complementary to this can be done using other types of datasets. Another important aspect of this study, which could be improved for better conclusions regarding its validation, is in relation to the number of samples. The results presented here were marked by the small amount of pre-classified samples, especially the MEL classification samples, making a generalization test for this type of classification impossible. In this work, a holdout adaptation was used for the sampling of classification and test data, but if more samples are available, cross-validation and bootstrap techniques can be used, and then a more comprehensive evaluation of the model can be effected.

The results obtained, with identification of segments of wave numbers that perform better than 75% of correct answers for data classifications related to diagnoses, particularly involving skin lesions from Raman spectroscopy, allow an optimistic perspective in relation to the study carried out, and the amplitude of its application to aid diagnosis of this and other nature, as well as the classification of patterns of other types of numerical data with the adaptation of the model to the respective attributes.

References


