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Evidence-based Model for Real-time Surveillance of ARDS

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Abstract

Real-time health surveillance becomes important and necessary with the increase of the elderly population to preserve their quality of life. Real-time models aim to provide alerts before the severe illness occurs. Acute respiratory distress syndrome is a crucial disease of the respiratory system that threatens the health of the elderly. This paper proposes a real-time model for the surveillance of ARDS based on belief functions theory. Non-invasive physiological signals are considered such as heart rate, respiratory rate, oxygen saturation and mean airway blood pressure. Different linear and nonlinear parameters are extracted from these signals; then a parameters selection procedure is performed to reduce their dimensionality. Afterwards, classifiers are constructed using parameters distributions defined in the evidence framework. Real-time prediction is then performed by combining all classifiers decisions. As results, high performances are obtained over the testing sets with performances of 77% and 71% for sensitivity and specificity, respectively.

Keywords: Acute respiratory distress syndrome; Evidence-based theory; Linear and nonlinear parameters; Real-time surveillance

1. Introduction

The elderly population is expected to increase from 962 million in 2017 to more than 2 billion by 2050 [1]. Older people are prone to many pathologies that reduce their autonomy and may threaten their lives leading to worldwide economic burden. The most known elderly-related pathologies are cardiovascular diseases, chronic respiratory diseases, neurological and mental disorders [2]. A real-time surveillance for elderly people helps to detect pathologies in advance. Thus it prevents its severity, shortens lengths of hospital stay and reduces mortality [3]. Real-time surveillance consists of

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continuous analysis of health-related data collected at a fixed time interval to reveal
10 upcoming illness. It has been the subject of debate for many researchers [4, 5, 6]. In
this paper, we present a real-time surveillance system for Acute Respiratory Distress
Syndrome (ARDS).

ARDS is a severe disability of the respiratory system characterized by expanded
pulmonary inflammation [7, 8]. It results in insufficiency of gas exchange with the
15 blood, associated with high short-term mortality [9]. Over years, several definitions
had been published until the Berlin Definition was proposed in 2012 [9, 10, 11]. Ac-
cordingly, ARDS is a hypoxemia that develops within one week of new respiratory
inflammation symptoms or a known clinical insult. It is diagnosed using the ratio of
the partial pressure of oxygen in the arterial blood (PaO_2) over the fraction of oxygen
20 in the inspired air (FiO_2). Moreover, ARDS was classified as mild ($200\text{ mmHg} <$
 $PaO_2/FiO_2 \leq 300\text{ mmHg}$), moderate ($100\text{ mmHg} < PaO_2/FiO_2 \leq 200\text{ mmHg}$),
or severe ARDS ($PaO_2/FiO_2 \leq 100\text{ mmHg}$). All having a minimum positive end-
expiratory pressure (PEEP) of 5 cm H₂O.

The risk of ARDS increases for patients who have undergone cardiac surgery or
25 were hospitalized for severe health problems. Depending on the case, some patients
were transferred from intensive care units (ICU) to wards or were completely dis-
charged from hospitals. Therefore, the monitoring and early-identification of those
patients is crucial in order to implement preventative measures. Generally, studies con-
ducted on ARDS have used clinical variables to characterize it or to predict its onset.
30 Some studies have investigated the risk factors associated with ARDS [12, 13], others
have developed predictive algorithms for ARDS using clinical data of ventilated ICU
patients, as in [14]. Other researchers have worked on mortality prediction related to
ARDS [15, 16] or respiratory complications [17, 18].

Furthermore, data mining and machine learning techniques are widely adopted in
35 biomedical applications to analyze, classify and predict medical data [19]. Classical
machine learning techniques such as k-nearest neighbors [20], naive Bayes [21] and
support vector machine [22] are applied on medical data to diagnose or predict a clinical
abnormality in physiological signals. However, there is a lack in the literature of
prediction models and real-time surveillance for ARDS.

40 This paper proposes a novel algorithm for predicting the occurrence of ARDS.
A key point in this surveillance is considering multiple vital signs that are easy to
collect [23, 24]. Identifying ARDS can be done using just one signal [25], but most
of the serious diseases occur with concurrent irregularities in multiple vital signs [6].
Hence, in this study, physiological signals are analyzed continuously to identify the
45 current health state of a patient. Multiple vital signs are used, that are the heart rate,
the respiratory rate, the oxygen saturation and the mean airway blood pressure. Instead
of using the whole signals, linear and non-linear parameters are extracted in real-time
and the most informative ones are selected. After that, a prediction model based on
evidence theory is defined in order to combine information of the extracted parame-
50 ters and generate a decision regarding the health state of the patient in study. This
model is constructed using the multi-parameter intelligent monitoring of intensive care
II database.

The outline of the paper is as follows. In the next section, the patient extraction
procedure is presented, and then the evidence-based method is introduced. Thereafter,

55 the results are illustrated and interpreted. Finally, a conclusion summarizes the paper
and proposes some perspectives.

2. Materials

Subjects are extracted from the multi-parameter intelligent monitoring of intensive
care II (MIMIC II) database. It contains two types of data: monitor waveforms
60 and clinical data, collected over a seven years period. A detailed explanation of the
database can be found in [26, 27]. In this study, the clinical database was considered
first to select subjects, using the values of their PaO_2/FiO_2 ratios. Hence, an
ARDS group is formed using the definition of both moderate and severe ARDS with
 $PaO_2/FiO_2 \leq 200$ mmHg in addition to PEEP levels higher than 5 cmH₂O; while a
65 non-ARDS group included subjects having mild and non ARDS patients with PEEP < 5
cm H₂O. After that, the waveform database is matched to the selected subjects of the
two groups, and their time series are extracted; that are the heart rate (HR), the res-
piratory rate (RR), the oxygen saturation (SpO₂) and the mean airway blood pressure
(MABP). These time series have a frequency of 1 sample/min. This leads to 140 ARDS
70 subjects and 135 non-ARDS subjects.

Among the ARDS subjects, there are 50 subjects that started their records after
ARDS diagnosis. These subjects are irrelevant for this study and thus were eliminated.
Other ARDS subjects started their records before the onset of ARDS but stopped before
ARDS diagnosis; these subjects, of number equal to 18, are also eliminated. Therefore,
75 only the 72 subjects having started records before the ARDS onset and last at least
until ARDS occurrence are considered. After that, a preprocessing step was required
to smooth the signals and remove the noise. Some subjects were removed due to noise
or too short signals. This leads finally to 50 ARDS subjects. The non-ARDS group
is then reduced to have equal number of subjects, leading to 50 non-ARDS subjects
80 as well. Figure 1 illustrates an example of the four extracted time series for an ARDS
subjects. Within the database of 100 subjects, a k -fold cross-validation technique is
considered to set training databases and testing databases [28]. In the following, the
terms *training set* and *testing set* would denote any training database obtained at any
step of the cross-validation process with its corresponding testing database respectively.

85 3. Evidence-based surveillance approach

The method described in this paper is a real-time method that monitors the ongoing
health state of patients using their vital signs. Let $x_{s,i}(\ell)$, $i = 1, \dots, 4$, denote the four
time series considered in this study for a subject s , recorded at time ℓ , and let t denote
the real-time. Moreover, let $x_{s,i}(a : b) = (x_{s,i}(a), \dots, x_{s,i}(b))$ be a segment of signal i
90 for subject s going from time a to time b .

3.1. Description of the approach

The aim of this work is to set a model Ψ from a training set that distinguishes
in real-time between subjects who are going to develop ARDS labeled “+ 1”, and
subjects who will not, labeled “- 1”. The model Ψ takes as input the collected signals

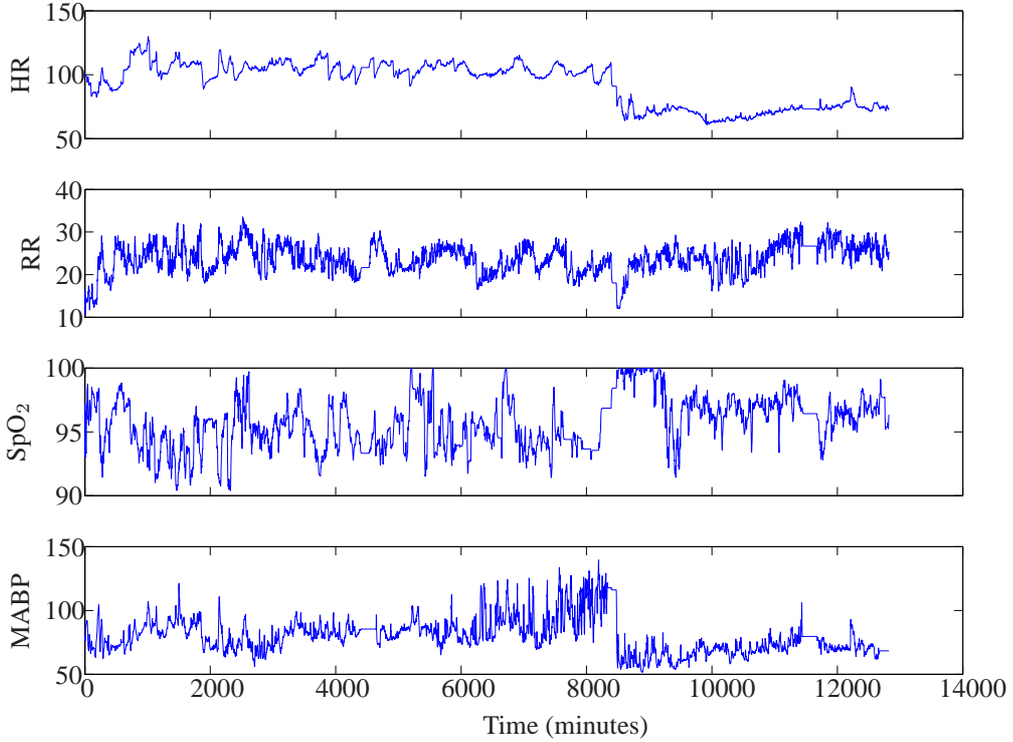


Figure 1: An example of the extracted time series for an ARDS subject

⁹⁵ $x_{s,i}(1 : t)$, going from the beginning of records until the real-time t , and yields as output a decision whether the subject in study is going to develop ARDS. Instead of using the whole signals, the proposed approach considers a fixed window of length τ taken at the end of the signals, leading to four segments $x_{s,i}(t - \tau + 1 : t)$ to be analyzed at each time t . The reason of analyzing the data of a window of length τ

¹⁰⁰ is to have fixed length signals in computations. Moreover, an abnormality is better and faster detected within a short segment, compared to the high length signal. The value of τ could be taken equal to 6, 12 or 24 hours. In this paper, it is obtained by an optimization over the training sets, as shown in the following. Then, a vector of parameters is extracted from these segments at real-time t . The beginning of the signals is yet needed to normalize these parameters. This vector of parameters is then used with the belief functions theory to assign evidence to each health state according to each parameter. The parameters are considered as sources of information, providing an evidence regarding each state of the patient at the given time. Thus, masses are assigned to each state “+ 1” and “- 1” according to each parameter. The masses are obtained by means of mass functions defined using the parameters extracted from the training set at an earlier offline phase. The masses are then combined to assign a final mass to each state. A higher mass assigned to the state “+ 1” at time t means that the subject is predicted to develop ARDS in the future, whereas a higher mass to “- 1” means that the ARDS is not yet predicted at the current time. Therefore, for a new test

¹¹⁰

115 subject given it signals in real-time, a decision will be made on its future health state at the earliest possible time.

3.2. Parameters Extraction

The first step of the prediction model is the extraction of parameters. Having a segment $x_{s,i}(t - \tau + 1 : t)$ from each signal at the real-time t , several linear and non-linear parameters are extracted as follows. In the following, t^- would denote the lower bound of the considered time interval, that is, $t^- = t - \tau + 1$. For linear parameters, we consider:

- The mean defined as

$$\mu_{s,i}(t) = \frac{1}{\tau} \sum_{\ell=t^-}^{\ell=t} x_{s,i}(\ell);$$

- The standard deviation

$$\sigma_{s,i}(t) = \sqrt{\frac{1}{\tau} \sum_{\ell=t^-}^{\ell=t} (x_{s,i}(\ell) - \mu_{s,i}(t))^2};$$

The variance is equal to the total power of spectral analysis. Thus, it reflects all the cyclic components responsible for the variability in the data [29].

- The skewness known as the third moment. It measures the symmetry of the data and it is computed using

$$Sk_{s,i}(t) = \frac{\sum_{\ell=t^-}^{\ell=t} (x_{s,i}(\ell) - \mu_{s,i}(t))^3}{\tau \times \sigma_{s,i}^3(t)};$$

125 It has been proposed that asymmetric tails to the left shown by positive skewness or to the right with negative skewness reflect, respectively, the acceleration or deceleration capacity of the time series data, e.g. for heart rate signal, it is an approximate distinction of vagal and sympathetic effects on the cardiac modulations [30].

- The kurtosis that describes the shape of the probability distribution and is calculated using the fourth moment of the data,

$$Kt_{s,i}(t) = \frac{\sum_{\ell=t^-}^{\ell=t} (x_{s,i}(\ell) - \mu_{s,i}(t))^4}{\tau \times \sigma_{s,i}^4(t)}.$$

130 The kurtosis measures the concentration of the data around the mean and reflects the rigidity of the signal [31].

On the other hand, non linear parameters are also considered, such as:

- The sample entropy (*SampEn*), that illustrates the amount of complexity in data. It is easily applied to any type of time series, including physiological data such as heart rate variability and blood pressure variability [32, 33]. It is computed using the negative natural logarithm of the conditional probability that a segment of length N having repeated itself for u samples within a tolerance r will also repeat itself for $u + 1$ samples [34]. Hence, for the segment $x_{s,i}$ of length $N = \tau$, *SampEn* is computed as follows,

$$SampEn_{s,i}(u, r, \tau, t) = -\ln \left(\frac{h_{s,i}(u+1, r)}{h_{s,i}(u, r)} \right),$$

where u is the length of sequences to be compared, r the tolerance for accepting and $h(u, r)$ is the number of pairs of patterns sequences of $x_{s,i}$ of length u whose distance is less than r .

135

- The detrended fluctuation analysis (DFA), that measures the scaling behavior of a segment [35]. It is defined by the short and long range scaling exponents DFA_1 and DFA_2 . To perform detrended fluctuation analysis for a segment $x_{s,i}(t - \tau + 1 : t)$, the integrated series $y(u)$, $u = 1, \dots, \tau$, is first computed

$$y(u) = \sum_{\ell=1}^{\ell=u} [x_{s,i}(\ell + t - \tau) - \mu_{s,i}(t)].$$

Then, $y(u)$ is divided into time segments of size n , and a local trend $y_n(u)$ is obtained by a least-squares line fit and subtracted from $y(u)$. The fluctuation $f(n)$ for segments of length n is calculated as follows:

$$f(n) = \sqrt{\frac{1}{\tau} \sum_{u=1}^{u=\tau} (y(u) - y_n(u))^2}.$$

The fluctuation $f(n)$ is computed for different segment sizes n , then a graph of $\log f(n)$ against $\log n$ is constructed. Finally, the scaling exponents DFA_1 and DFA_2 define, as their names denote, the slope of the relation between $\log f(n)$ and $\log n$ for ranges of low and high values of n .

140

DFA has been used to assess the self-similar correlations of the time series [36]. It has also been described as a measure of “roughness” in the signals, with higher scaling exponents representing a smoother time series [37].

Having the four measured signals, these computations lead to seven parameters per signal and thus twenty-eight parameters, for each subject s at each time t . Let $p_{s,j}(t)$, $j = 1, \dots, 28$, denote these parameters. For many health conditions, the trends in vital signs differ among patients. Thus, normal is a relative state for each patient, that differs with age, weight, medical history, etc. Hence, instead of using the raw values of the parameters, a normalization phase is performed according to each patient. Therefore, a first segment of length τ is considered from each signal, $x_{s,i}(1 : \tau)$. It corresponds to the initial state of the patient that is assumed to be its stable one.

150

Then, the different parameters are computed for the initial segments. Let $p_{s,j}^{(0)}$ denote their initial values. Then, for each real-time measurement, the obtained parameters are normalized, by computing their ratio to the parameters of the first segment $p_{s,j}(t)/p_{s,j}^{(0)}$. In the following, $p_{s,j}$ would denote the normalized parameter, for simplicity.

155 Note that, a parameter selection would be performed in the offline phase using the training set to keep only informative one. More details are given in Section 3.4. In the following, let J be the set of indices of the selected parameters, and thus $p_{s,j}(t)$, $j \in J$, are the normalized parameters extracted at real-time t from a subject s signals.

3.3. ARDS prediction with Belief functions

The theory of belief functions, also called Dempster-Shafer theory, is introduced for the analysis of imperfect information[38, 39]. This theory makes it possible to illustrate the uncertainty and imprecision of information. It also takes into account ambiguities and conflicts between sources. The belief functions theory operates on a frame of discernment Ω which consists of a finite set containing mutually exclusive propositions. In this study, there are two groups of subjects, ARDS labeled “+1” and non-ARDS labeled “-1”. Then $\Omega = \{+1, -1\}$. Let 2^Ω be the set of all possible subsets of Ω , it is then defined by:

$$2^\Omega = \{\emptyset, \{+1\}, \{-1\}, \{+1, -1\}\}.$$

160 While \emptyset denotes that the state of the subject is neither ARDS, nor non-ARDS, which means impossibility, $\{+1, -1\}$ denotes ignorance and thus ambiguity.

3.3.1. Basic concept

The main concept relies on the modeling of the evidence provided by the extracted parameters. Information given by a source, that is a parameter, can be represented by a *basic belief assignment* (BBA), also named mass function [40]. Let $m_j(\cdot)$ be the mass function associated to the parameter p_j . Then, $m_j(\omega, s, t)$ reflects the strength of evidence supporting a subset $\omega \in 2^\Omega$ according to the value of the parameter p_j at time t . In other words, $m_j(\omega, s, t)$ represents the part of evidence saying that the state of the subject s falls in ω at time t . The mass function should have the following properties

170 $\forall s, t$:

$$\begin{cases} m_j(\emptyset, s, t) = 0, \\ m_j(\omega, s, t) \rightarrow [0, 1], \text{ for } \omega \in 2^\Omega, \\ \sum_{\omega \in 2^\Omega} m_j(\omega, s, t) = 1. \end{cases} \quad (1)$$

In order to construct the mass functions related to each parameter, the training sets are considered. For ARDS subjects of a given training set, the last segments of their signals are used to compute their parameters $p_{s,j} = p_{s,j}(T_s)$, T_s being the length of the signals of an ARDS subject s . The reason of this choice is that the end of the signals precedes directly the occurrence of ARDS and thus contains mostly its instability. For non-ARDS subjects, the more the segments approaches the beginning of the signals, the more we are sure of the stability of the subject, since it is possible that a subject labeled non-ARDS develops the ARDS after recordings. Then, the parameters of non-ARDS are computed using segments selected from the wholly beginning of the signals,

180 following the first segments. Let S_{+1} and S_{-1} be the sets of indices of ARDS and non-ARDS subjects of the training set respectively. Then, normalized parameters databases $p_{s,j}$, $s \in S_{+1}$, and $p_{s',j}$, $s' \in S_{-1}$, are obtained. Afterwards, for a parameter type j , the pdfs are estimated using (1) only ARDS parameters values $p_{s,j}$, $s \in S_{+1}$, (2) only non-ARDS values $p_{s',j}$, $s' \in S_{-1}$ and (3) parameters of both classes $p_{s,j}$, $s \in S_{+1} \cup S_{-1}$. Let $Q_{j,\{+1\}}(\cdot)$, $Q_{j,\{-1\}}(\cdot)$ and $Q_{j,\{+1,-1\}}(\cdot)$ denote the estimated pdfs. 185 These distributions could be obtained by fitting the histograms of the parameters to all the classical distributions. Then, in the online phase, having a computed parameter $p_{s,j}(t)$ at the real-time t , the mass assigned to each subset of 2^Ω is given by

$$m_j(\omega, s, t) = \frac{Q_{j,\omega}(p_{s,j}(t))}{\sum_{\omega' \in 2^\Omega, \omega' \neq \emptyset} Q_{j,\omega'}(p_{s,j}(t))}, \omega \in 2^\Omega, \omega \neq \emptyset, \quad (2)$$

whereas $m_j(\emptyset, s, t) = 0$. Figure 2 illustrates an example of such a computation. The 190 plot shows the distributions of the normalized standard deviation of respiratory rate signals parameter for ARDS subjects of a training database in straight thick line, the one for non-ARDS subjects in dashed line and the one of all subjects in straight thin line. Moreover, the plot shows the mass assignment for two new parameter values $p_2^{(1)}$ and $p_2^{(2)}$. The first parameter value falls in the distribution of ARDS subjects and the second is in the middle of ARDS and non-ARDS distributions. When assigning the 195 masses of each value from Eq. (2), the assigned mass of $p_2^{(1)}$ from $Q_{j,\{+1\}}$ is higher than those of $Q_{j,\{-1\}}$ and $Q_{j,\{+1,-1\}}$. Thus, the decision obtained from the parameter $p_2^{(1)}$ is the state of $\{+1\}$. On the other hand, for parameter $p_2^{(2)}$, the distributions of $\{+1\}$ and $\{-1\}$ are too close, leading to similar masses from both. However, higher 200 mass is associated in this case to the distribution of $\{+1, -1\}$, thus introducing the ambiguity of such case. The mass assigned to $\{+1, -1\}$ is higher than those of $\{+1\}$ and $\{-1\}$ in all the cases where these distributions are too close. This figure illustrates the effectiveness of the evidence-theory concept by inserting ambiguities when dealing with close data and thus avoiding erroneous decisions.

205 3.3.2. Discounting

Since we are working on parameters extracted from human bodies, the information cannot be completely reliable. Hence, a discounting operation is needed to take into consideration the reliability of the information provided by each parameter [39, 41]. This operation transforms each mass function to a less informative one, based on the 210 degree of reliability of the source. Moreover, in this approach, it is possible to estimate its reliability according to each state, leading to a contextual discounting [42, 43]. Let $\beta_{j,\{+1\}} = 1 - \alpha_{j,\{+1\}}$ and $\beta_{j,\{-1\}} = 1 - \alpha_{j,\{-1\}}$ be the degrees of reliability of a parameter p_j knowing that the true state is “+ 1” and “- 1” respectively. Then the

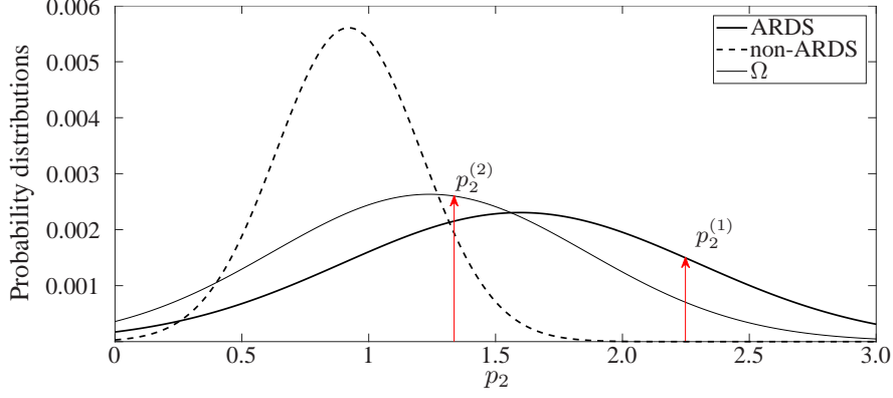


Figure 2: Mass assignment of new computations in the case of the normalized standard deviation of respiratory rate signals.

discounted mass function ${}^\alpha m_j$ assigned to p_j is given by

$$\begin{cases} {}^\alpha m_j(\{+1\}, s, t) = \beta_{j, \{-1\}} m_j(\{+1\}, s, t), \\ {}^\alpha m_j(\{-1\}, s, t) = \beta_{j, \{+1\}} m_j(\{-1\}, s, t), \\ {}^\alpha m_j(\Omega, s, t) = m_j(\Omega, s, t) + \alpha_{j, \{-1\}} m_j(\{+1\}, s, t) + \alpha_{j, \{+1\}} m_j(\{-1\}, s, t), \\ {}^\alpha m_j(\emptyset, s, t) = 0. \end{cases} \quad (3)$$

215 The error rates $\alpha_{j, \{+1\}}$ and $\alpha_{j, \{-1\}}$ could be estimated using the distribution functions $Q_{j, \omega}$. Knowing that a subject s state is equal to “+1”, an error is made in evidence assignment if a higher mass is given to any subset of Ω other than $\{+1\}$. It is also the same for $\{-1\}$. Then, $\alpha_{j, \omega}$, $\omega = \{+1\}$ or $\{-1\}$, is computed for all parameter values, where $Q_{j, \omega}(p)$ is less than any $Q_{j, \omega'}(p)$, for $\omega' \neq \omega \in 2^\Omega$. Therefore, it is
220 defined as follows:

$$\alpha_{j, \omega} = \epsilon_{j, \omega} = \int_{D_{j, \omega}} Q_{j, \omega}(p) dp, \quad (4)$$

where $D_{j, \omega} = \{p | Q_{j, \omega}(p) < Q_{j, \omega'}(p), \forall \omega' \in 2^\Omega, \omega' \neq \omega\}$.

3.3.3. Combination

Once the discounted masses of the different parameters are computed, they can be combined using the normalized conjunctive rule of combination from the Dempster-Shafer theory. By doing this computation, one obtains a more informative mass function, leading to more efficient estimation. The combination of these masses is denoted by \oplus and defined by the following equation:
225

$$m_{\oplus}(\omega, s, t) = \frac{\sum_{\cap \omega^{(j)} = \omega} \prod_{j \in J} ({}^\alpha m_j(\omega^{(j)}, s, t))}{1 - \sum_{\cap \omega^{(j)} = \emptyset} \prod_{j \in J} ({}^\alpha m_j(\omega^{(j)}, s, t))}. \quad (5)$$

3.3.4. Decision making

Finally, to make a decision on the health state of a subject s , the mass function m_{\oplus} defined on Ω is transformed to a probability measure over the singletons with the pignistic transformation defined by:

$$BetP(\omega, s, t) = \sum_{\omega' \in 2^{\Omega}, \omega \subseteq \omega'} \frac{m_{\oplus}(\omega', s, t)}{|\omega'|}, \forall \omega \in \Omega, \quad (6)$$

for $\omega = \{+1\}$ or $\{-1\}$. In this way, the final mass of $\{+1, -1\}$ subset is divided between $\{+1\}$ and $\{-1\}$. The state having the highest pignistic level is selected for each subject at every time t . Thus, a decision is made over the health state of subject s whether he will develop ARDS based on its recordings, and an alert is generated for a positive decision.

To make a final decision from the real-time analysis about the health state of a patient, it is also possible to wait for successive positive decisions. Here, a threshold must be defined as being the number of successive ARDS decisions needed to decide if the patient is going to develop ARDS. This is done by applying the algorithm iteratively on the total length of the signals s of the training set, then different numbers of successive positive decisions are considered and the performance indexes are computed for each number, that are the sensitivity (Se) and specificity (Sp) indexes, also called true positive rate and true negative rate, respectively. The threshold of successive ARDS decisions is obtained thereafter by maximizing the Youden index, $Youden = Se + Sp - 100$, over all the considered numbers.

Waiting for successive positive estimations overcomes detection of isolated abnormalities and leads to a higher specificity of the algorithm.

3.4. Selection of parameters

Instead of considering all the extracted parameters in the model, it is relevant to select the most informative ones with less error rates. This part of the algorithm consists of ranking the parameters according to their error rates and then performing a sequential forward selection (SFS) of parameters [44]. Hence, an error rate ϵ_j is computed for each parameter p_j , which is the average of conditional errors, $\epsilon_j = (\epsilon_{j, \{+1\}} + \epsilon_{j, \{-1\}})/2$. The conditional errors $\epsilon_{j, \omega}$ are computed using (4). A global ranking of parameters is performed. Parameters from all signals are ordered from the lowest error obtained to the highest one.

Once the parameters are ranked, a sequential forward selection procedure is performed. One parameter is added at a time and performance indexes of the model are computed. For each added parameter, the belief model is applied in real-time for all the subjects of the training set and a decision is generated for each of them. The sensitivity and specificity indexes are computed. Thereafter, A curve is plotted and updated at each iteration illustrating the sum of the performance indexes obtained versus the number of parameters. Whenever a new value added to the curve shows a decrease in the performance higher than a value ϵ , the SFS procedure ends and the parameters obtained are maintained.

In addition, in order to take into consideration the dependence of the parameters, the correlation between the added parameter and the existing parameters is tested and

Table 1: The most significant parameters for each signal, parameters having a p-value < 0.02 are shown. Mean \pm standard deviation are shown for each parameter.

Signal	Parameters	ARDS group	non ARDS group
HR	<i>mean</i>	1.0297 ± 0.1439	1.0218 ± 0.0935
	<i>Sk</i>	-1.875 ± 6	-3.3629 ± 31.2769
	<i>Kt</i>	1.0628 ± 0.4082	1.2803 ± 0.6892
	<i>SampEn</i>	1.2277 ± 0.6079	1.0231 ± 0.1371
RR	σ	1.6226 ± 1.4729	0.9209 ± 0.2851
	<i>Sk</i>	-0.8592 ± 10.6885	1.0231 ± 3.3186
	<i>Kt</i>	1.0172 ± 0.5052	1.3381 ± 1.2061
	<i>SampEn</i>	1.4088 ± 0.5191	1.0281 ± 0.2
SpO ₂	<i>Sk</i>	0.6881 ± 4.0655	3.1707 ± 14.2335
	<i>Kt</i>	1.0871 ± 0.5252	1.0907 ± 0.8359
	<i>SampEn</i>	1.2836 ± 0.6092	1.5598 ± 0.982
	<i>DFA</i> ₁	0.8678 ± 0.1777	1.0172 ± 0.2767
	<i>DFA</i> ₂	1.0289 ± 0.1349	1.0194 ± 0.0667
MABP	μ	1.0262 ± 0.1246	0.9868 ± 0.072
	<i>Sk</i>	0.6396 ± 1.6238	0.5796 ± 8.2055
	<i>DFA</i> ₁	0.9432 ± 0.1459	1.0743 ± 0.3192
	<i>DFA</i> ₂	0.9985 ± 0.0178	1.0022 ± 0.0397

270 compared to the selection procedure alone. If any correlation coefficient between the parameters presents a p-value ≤ 0.05 then the parameter is considered correlated with an existing one, thus it is eliminated from the selection procedure.

4. Results

275 As mentioned above, two types of parameters were extracted from each time series. Linear parameters are the mean μ , the standard deviation σ , the skewness *Sk* and the kurtosis *Kt*. Non linear parameters are the sample entropy *SampEn*, and both factors from the detrended fluctuation analysis *DFA*₁ and *DFA*₂. For sample entropy, the value of *u* is taken equal to 5, and *r* equals 0.2, as mentioned in [45]. For the detrended fluctuation analysis, *DFA*₁ is calculated between the range $n = 4$ and $n = 16$ and *DFA*₂ is calculated between $n = 16$ and $n = 64$, as shown in [35]. The results are
280 obtained by a 5 fold cross validation repeated 10 times. For the illustration of the approach, presented in Sections 4.1, 4.2 and 4.4, only one training and one test sets are considered, then all the sets are considered for the performance evaluation in Section 4.5.

4.1. Statistical analysis

285 First, a statistical analysis is performed to identify parameters that show a significant difference between ARDS and non-ARDS groups. For each parameter, values were compared between both groups using the two-sample F-test. A p-value < 0.02 was considered as significant. Results are presented in Table 1. Different linear and

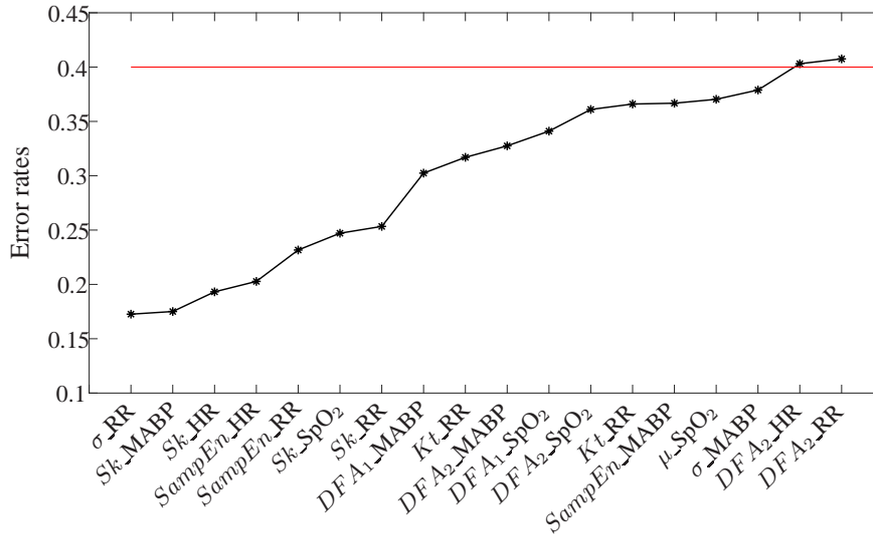


Figure 3: Ranking of parameters according to error rates. Only parameters having an error lower than 0.4 are included in the figure.

non-linear parameters for each signal has shown significant difference between groups. It can be noted from the table that distributions of non ARDS group are more asymmetric than ARDS group for all the signals from the values of skewness. In addition to being asymmetric, kurtosis shows that distributions of non ARDS group for HR, RR and SpO₂ have heavy tails. Moreover, RR's distribution is widely spread in ARDS group according to σ . For ARDS group, sample entropy is higher in HR and RR signals; while it is lower in SpO₂. Detrended fluctuation analysis factors are generally higher in non ARDS group for SpO₂ and MABP signal.

4.2. Parameters Selection

Since there is a high number of parameters, a ranking of these parameters based on the error rates is performed. The ranking is illustrated in Figure 3. Parameters having error rates higher than 0.4 are not presented. The results obtained from the error rate ranking in the figure are confirmed from the statistical analysis shown in Table 1. As shown, most of the parameters that have low errors in Figure 3 also showed a significant difference in the table. This figure shows that the standard deviation extracted from RR signals presented the lowest error rate among all parameters extracted from the four signals, thus ranking at the top.

From these ranked parameters, a sequential feature selection is considered to reduce the input vector of the belief computations. Figure 4 illustrates the selection procedure on the training set for both cases with and without correlations. The plot shows the accuracy as a function of the indicators of the ranked parameters. The accuracy is defined by the proportion of correctly identified subjects among the total number of subjects. A marker is placed on the curves when a parameter is selected. Starting with σ from RR, the parameters presented in Figure 3 are added sequentially until

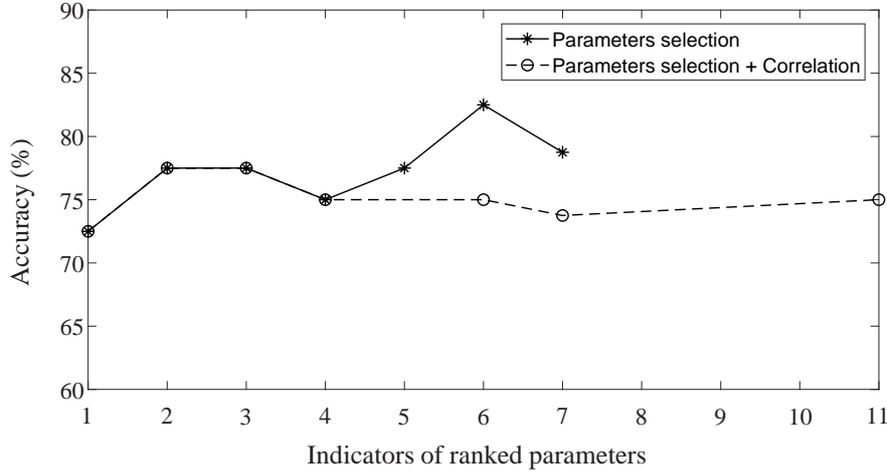


Figure 4: The parameters selection procedure using the simple procedure and the correlation extension procedure.

the accuracy decreases more than $\epsilon > 5\%$. For the correlation study, if a parameter shows a correlation with any existing parameter with a p-value lower than 0.05 it is excluded from the selection. The simple selection procedure finished on the insertion of the seventh parameter that is Sk from RR. Thus, six parameters are selected in the simple SFS procedure which are Sk and $SampEn$ from HR, σ and $SampEn$ from RR, Sk from both SpO_2 and MABP. However, there is no decrease in the accuracy of the model in the correlation study but the accuracy is lower than that obtained from the simple selection.

4.3. Selection of the alert generation threshold

In order to make a decision on the health state of a new test subject, a threshold must be defined from the training set. Therefore, we sweep the value of successive decisions on the interval $[1, \dots, 30]$ hours and the performances Se and Sp are computed. The threshold that maximizes the Youden index is then considered. Figure 5 illustrates the ROC curve of the proposed algorithm by taking the fusion of the first six parameters obtained in the previous section. The plot shows that the optimal cut-off value is obtained for threshold = 20 hours with $Se = 81.4\%$ and $Sp = 83.78\%$.

4.4. Selection of window size

As described in the real-time model we have used a fixed-length window for extracting the parameters vector for all subjects. Thus, the selection of the size of the window is important in this study. A short window may not contain enough information and a long window may eliminate an abnormal change in the signal. Different window sizes are evaluated in this study, values are considered in the range $[4, 6, \dots, 24]$ hours. Figure 6 illustrates the accuracy for each window size (in hours) as well as the number of selected parameters and the threshold shown as a couple (number of parameters,

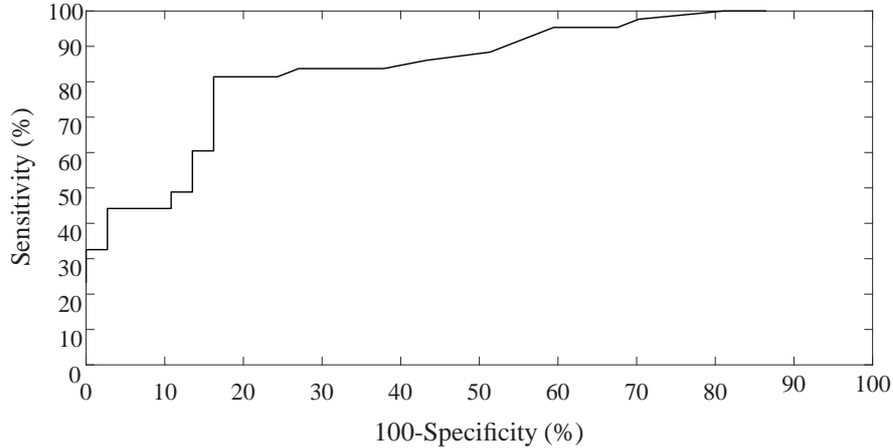


Figure 5: ROC curve for the selection of the threshold for alert generating.

threshold). As shown in the figure, window size = 24 hours presents the highest accuracy of 82.5% on the considered training set. These values are then considered for the following.

340 4.5. Performance Evaluation

After the illustration of the extraction and selection of the parameters, the model is tested on the test sets. In this section, results are obtained by a 5-fold cross validation repeated 10 times. Table 2 presents the performance of each added phase to obtain at the end the performance of the complete proposed model. The modeling of
 345 the data by associating and combining masses from all the parameters leads to a sensitivity of 70.23% and a specificity of 71.74% over the training sets and $Se = 70.16\%$ and $Sp = 67.89\%$ over the test sets. However, when the parameter selection phase was performed, an enhancement in the sensitivity was noted over the training sets ($Se = 78.10\%$, $Sp = 66.91\%$) and the test sets ($Se = 72.59\%$, $Sp = 61.16\%$). In addition,
 350 tion, discounting the mass functions using the error rates improved the performance to 81.65% of sensitivity and 74.72% of specificity over the training sets and 77.24% and 71.25% for sensitivity and specificity respectively over the test sets. However, when the correlation was considered in the selection phase the performances decreased with
 355 $Se = 80.18\%$ and $Sp = 69.29\%$ over the training sets and 74.10%, 61.12% over the test sets.

4.6. Comparison to the state-of-the-art methods

The proposed model is compared to some classification techniques such as Bayesian hypothesis test, k-nearest neighbors (kNN) and SVM. The Bayesian hypothesis test is based on Bayes theorem. It is a probability model that consists of factorizing a joint
 360 probability distribution into a set of conditional distributions for each variable. The k-nearest neighbors is one of the basic algorithms in machine learning. It classifies a new observation to a class by choosing the majority class among the k nearest neighbors in

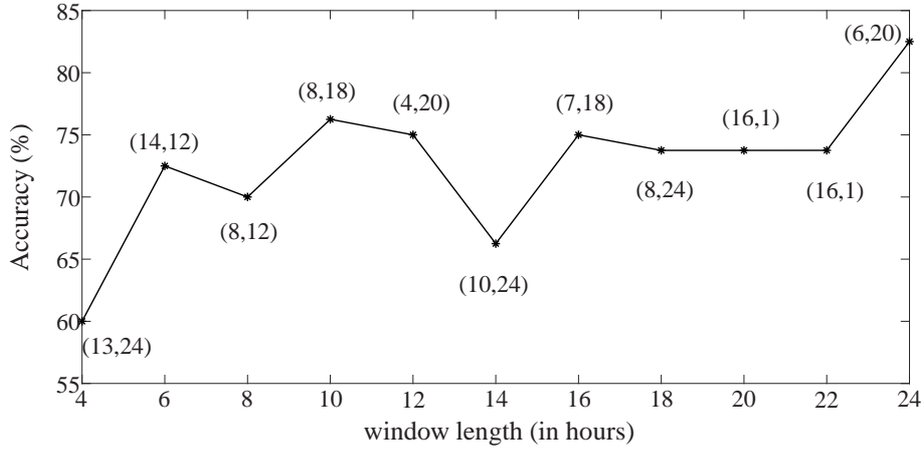


Figure 6: Changes in the accuracy over the training set with windows length. Values in the figure denote (number of parameters, threshold).

Table 2: Performance of the proposed model and the influence of each phase.

Method	training set		test set	
	Se(%)	Sp(%)	Se(%)	Sp(%)
BF without discounting, nor selection	70.23	71.74	70.16	67.89
BF without discounting	78.10	66.91	72.59	61.16
complete BF model	81.65	74.72	77.24	71.25
complete BF model+corr	80.18	69.29	74.10	61.12

the training set. A ten-fold cross validation was performed to obtain k that minimizes the test loss. The optimal k was found to be 8. As for SVM, it has proven to be a good model to classify data. It can be used to differentiate the data using a decision boundary that produces the optimal separation of classes. Table 3 shows the performance indexes of the proposed model compared to these described techniques. As shown, the proposed model outperforms the kNN, the SVM as well as the bayesian tests, that are a simplest form of the evidence theory where the ambiguities between the sets are not considered. These results show the importance of the proposed model.

Table 3: Comparison to state-of-the-art classification methods

Method	Sensitivity (%)	Specificity (%)
KNN	59.20	79.75
Bayesian test	67.74	65
SVM	58.67	74.06
The proposed Model	77.24	71.25

5. Discussion

This study proposed a real-time model, based on the belief functions theory, for predicting ARDS prior to its onset. ARDS results from inflammatory alveolar injury occurring progressively [46]. The process begins with the destruction of the barriers of the alveolar membranes resulting by increasing the alveolar permeability. Then, injuries progress causing decreased pulmonary compliance which leads to pulmonary hypertension and refractory hypoxia [47, 48]. Moreover, ARDS has been linked to clinical features such as severe dyspnea, tachypnea and hypoxemia according to [49]. Other studies on the clinical profile of ARDS had shown that it is associated with breathlessness, hypotension, tachypnea [50] and tachycardia [51]. Although, there are no clear clinical features for ARDS, it is evident that ARDS is linked to cardiorespiratory and cardiovascular mechanisms. Therefore, the four vital signs that are considered are the heart rate, the respiratory rate, the oxygen saturation and the mean airway blood pressure. However, there is a lack in the literature for the characterization of ARDS using feature extraction.

In feature extraction techniques, parameters are extracted from raw data signals and pertinent ones are selected to enhance the fusion performance. The characterization of signals using the extraction of parameters has been widely applied in biomedical applications. For instance, an analysis of heart rate variability during syncope was conducted by extracting parameters such as the mean, the standard deviation of the RR interval and non linear parameters such as the sample entropy and the detrended fluctuation analysis [52]. Moreover, linear and non-linear parameters were extracted from heart rate variability signals for ARDS subjects to analyze their response to alveolar recruitment manoeuver [53]. Only few studies have considered other cardiovascular signals than heart rate, like blood pressure and oxygen saturation signals [31, 54]. A characterization of heart beat and blood pressure for diabetes patients using time domain parameters such as the mean, the standard deviation, the skewness and the kurtosis was studied in [31]. In addition, detrended fluctuation analysis was extracted from blood pressure signal to study its dynamics through surgical procedure in rats [55]. In [33], a characterization of periventricular leukomalacia, a brain injury affecting infants, is proposed by extracting minimum, maximum, mean, variance, skewness, kurtosis, energy of wavelet coefficients and sample entropy from heart rate, mean arterial blood pressure and oxygen saturation. From what preceded, we can conclude that the extraction of parameters from different physiological signals might help in the characterization of physiological aspects related to several pathologies, such as ARDS.

Hence, in this paper, linear and non-linear parameters were extracted from the time series as mean, standard deviation, skewness, kurtosis, sample entropy and detrended fluctuation analysis. Linear parameters provide information regarding the distributions of the data. Distributions were shown to be asymmetric, having wide tails and being widely spread. This verifies that biomedical data are not gaussian [56]. Conversely, it was found that heart rate and respiratory rate had higher sample entropy in ARDS group, thus increased complexity [57]. However, oxygen saturation presented lower complexity in ARDS subjects. In other terms, SpO_2 presents more similarity in the data. From DFA_1 and DFA_2 values, we can notice that SpO_2 and MABP had lower correlations in ARDS subjects [36].

Moreover, feature fusion is performed using the belief functions framework. The proposed belief functions framework defines ARDS and non-ARDS groups based on the probability distributions of the data according to each parameter. In addition, these models consider supper subsets instead of considering only singletons subsets, thus they introduce the notion of ambiguity between choosing an ARDS or non-ARDS states. The belief theory also takes into account the reliability of each source of information presented by the discounting operation. The reliability of the sources is a very important characteristic, especially when working with parameters extracted from physiological signals. On top of these comes the importance of belief functions framework, that is the modeling of the data even with missing recordings. It often happens that some signals are missing in some part of the recordings because of disconnected electrodes or a sudden movement of the patient. Unlike classical data mining techniques, the belief functions guarantee that missing parameters due to missing signals will not affect the overall performance of features fusion.

From the results, the reduction of the number of parameters, by performing a selection procedure, improved the sensitivity of the model and reduced the complexity of the computations as well. This way only parameters having low error rates are included in the model and the combination that presents the best local accuracy is considered. Thus, the modeling of the data is done using the minimum amount of parameters instead of using all the extracted parameters. Then, the inclusion of the conditional reliability of each source has led to an enhancement in the accuracy of the model over the training and the test sets. Each parameter gives information regarding the patient and the reliability of the parameter itself in distinguishing between states. Finally, 77% of ARDS patients are correctly identified in real-time before the occurrence of ARDS and only 29% of non-ARDS patients are misclassified and considered as possible developers of ARDS. The obtained results demonstrate, beside the already existing studies on ARDS, that ARDS is linked to cardiorespiratory and cardiovascular mechanisms. More importantly, ARDS can be predicted using non-invasive physiological signals instead of using clinical exams or variables that necessitate the presence of subjects in hospitals for long period [15, 16, 14].

It is shown from the results that this model outperformed the classical classifiers tested similarly on the data. It is worth noting that the bayesian tests perform only on singletons and the state membership is determined by computing the likelihood ratio. However, the added ambiguity set in the proposed evidence-based model may affect the final decision since it is considered in both discounting and combination phases. Thus, the advantage of such models over the classical classifiers, especially the bayesian tests, is the inclusion of higher order sets that illustrate the ambiguity intervals.

Previous studies were conducted on the prediction of ARDS or the mortality related to ARDS using computed tomographic images [16] or clinical variables [15, 14], that necessitate clinical exams or the presence of subjects in the hospitals for long periods. However, the approach proposed in this paper uses only vital signs that can be collected using non-invasive techniques, thus without disturbing the subjects. Moreover, these studies present general statistical models to detect ARDS assuming that all subjects will present similar changes; while this approach is a subject-based one that considers an initial stable state for each subject and computes the ongoing parameters with respect to the stable state. In addition, an information fusion was handled using the evidence

theory to consider the ambiguity and unreliability of the sources. Finally, this approach achieved good performances for both sensitivity and specificity indexes for the real-time surveillance of ARDS.

465 6. Conclusion

This paper proposes an evidence-based model for the prediction of ARDS in intensive care unit patients. This work presents several contributions. Non invasive vital signs are included in the study for many reasons, the facility of their acquisition, the possible integration in home surveillance systems and the link between these signals and the risk factors associated with ARDS. Linear and non-linear parameters were computed and considered the source of information, since they can give information about the properties of a signal more than the signal itself. Imprecision and unreliability of information sources were considered by the belief functions model. It assigns masses to each subset computed from a predefined training set. Then, these masses were discounted and combined according to the reliability of each source of information. This model is then extended to reduce the dimension of the input by performing a parameter selection procedure. This model has achieved high performances in both ARDS and non-ARDS groups. In addition, it outperforms state of the art techniques. Further work must be done to extract more parameters that provide evidence on different characteristics in the signals. Moreover, it would be interesting to establish the relationships between the changes in time series and the cardiovascular mechanism. A further study on the parameters selection will also be done, to take into consideration the dependance of parameters from the beginning of the selection process.

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