RESEARCH ARTICLE

Modeling and Methods of Statistical Processing of a Vector Rhythmocardiosignal

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Abstract:

Aims:
We have developed a new approach to the study of human heart rate, which is based on the use of a vector rhythmocardiosignal, which includes as its component the classical rhythmocardiosignal in the form of a sequence of heart cycle durations in an electrocardiogram.

Background:
Most modern automated heart rate analysis systems are based on a statistical analysis of the rhythmocardiogram, which is an ordered set of R-R interval durations in a recorded electrocardiogram. However, this approach is not very informative, since R-R intervals reflect only the change in the duration of cardiac cycles over time and not the entire set of time intervals between single-phase values of the electrocardiosignal for all its phases.

Objective:
The aim of this paper is to present a mathematical model in the form of a vector of stationary and permanently connected random sequences of a rhythmocardiosignal with an increased resolution for its processing problems. It shows how the vector rhythmocardiosignal is formed and processed in diagnostic systems. The structure of probabilistic characteristics of this model is recorded for statistical analysis of heart rate in modern cardiodiagnostics systems.

Methods:
Based on a new mathematical model of a vector rhythmocardiosignal in the form of a vector of stationary and permanently connected random sequences, new methods for statistical estimation of spectral-correlation characteristics of heart rate with increased resolution have been developed.

Results:
The spectral power densities of the components of the vector rhythmocardiosignal are justified as new diagnostic features when performing rhythm analysis in modern cardiodiagnostics systems, complementing the known signs and increasing the informative value of heart rate analysis in modern cardiodiagnostics systems.

Conclusion:
The structure of probabilistic characteristics of the proposed mathematical model for heart rate analysis in modern cardiodiagnostics systems is studied. It is shown how the vector rhythmocardiosignal is formed, and its statistical processing is carried out on the basis of the proposed mathematical model and developed methods.

Keywords: Vector of stationary and permanently connected random sequences, Methods of statistical estimation, Electrocardiosignal, Vector rhythmocardiosignal, Heart rate, Cardiodiagnostics systems.

1. INTRODUCTION

The heart rate analysis has long been an integral part of not only modern cardiology but also many other areas of biomet-
human. In addition, the heart rate analysis is carried out for early diagnosis of the pathological condition of the fetus, the state of the autonomic system in diabetic patients. Heart rate makes it possible to assess the value of the risk of death in myocardial infarction, the degree of tension of the state of the regulatory process in the human body, etc [1 - 12].

The special efficiency of the heart rate analysis is achieved by using modern computerized diagnostic systems that make it possible to automate the assessment of diagnostic signs and make medical decisions about the human heart rate based on recorded cardiosignals, mainly electrocardiosignals. The accuracy, reliability, information content and speed of functioning of cardiodiagnostic computerized heart rate research systems significantly depend on the adequacy and constructiveness of the mathematical model of heart rate, as well as the accuracy, reliability, information content, speed of methods and algorithms of its analysis in these information systems.

Most methods for processing the classical rhythmocardiosignal in the framework of the stochastic approach are based on three of its probabilistic models, namely, a random variable, a random stationary sequence, and a periodically correlated random sequence are used. These models are based on an approach to describing the heart rate as a sequence of R-R intervals that uses a ritmocardiosignal (classical ritmocardiosignals), which imposes significant limitations on the informative value of heart rate analysis. The essence of this limitation is that the values of R-R intervals, which are the corresponding values of the rhythmocardiosignal, reflect only the change in the duration of cardiac cycles over time, and not the entire set of time intervals between single-phase values of the electrocardiosignal for all its phases, which does not make it possible to describe the heart rate with sufficient information. That is why the approach based on the analysis of the classical rhythmocardiosignal as a sequence of R-R intervals does not allow us to identify more subtle and detailed features of the heart rate in modern computer systems of medical diagnostics.

In the study [13, 14], a new approach to heart rate analysis based on a high-resolution rhythmocardiosignal was developed. As indicated in these works, the classical rhythmocardiosignal is embedded in a rhythmocardiosignal with increased resolution, which is the basis for increasing the level of the information content of heart rate analysis in modern computer systems for functional diagnostics of the human heart state based on a rhythmocardiosignal with increased resolution.

In this approach, the heart rate was represented by a high-resolution rhythmocardiosignal (other names: high-informative rhythmocardiosignal or vector rhythmocardiosignal), the mathematical model of which was a vector of normally distributed random variables. Therefore, the classical rhythmocardiosignal is embedded in a rhythmocardiosignal with increased resolution, which is the basis for the level increase of information content of heart rate analysis in the modern computer systems for functional diagnostics of the human heart state based on a rhythmocardiosignal with increased resolution.

As a mathematical model of a rhythmocardiosignal with increased resolution, in the study [13, 14], is justified the use of a vector of random variables with a normal distribution. This stochastic model can already take into account several phases of the cardiac cycle when analyzing the heart rate. However, this model is a relatively simple mathematical model of a high-resolution rhythmocardiosignal, since it does not allow us to study its temporal dynamics. To take into account the temporal dynamics of a rhythmocardiosignal with increased resolution, it is necessary to use the mathematical apparatus of random sequence theory, namely, to consider it a vector of discrete random sequences.

In this paper, a mathematical model of a rhythmocardiosignal with an increased resolution for its processing problems is described as a vector of stationery and stationary connected random sequences. It shows how the vector rhythmocardiosignal is formed, processed and modeled in diagnostic systems. The structure of probabilistic characteristics of this model is recorded for statistical analysis of heart rate in modern cardiodiagnostics systems.

2. MATERIALS AND METHODS

2.1. Electrocardiosignal Mathematical Model in the form of a Conditional Cyclic Random Process

Let’s move on to constructing a mathematical model of a vector rhythmocardiosignal. Since the rhythmocardiosignal is formed from an electrocardiosignal, the mathematical model of the vector rhythmocardiosignal is based on the corresponding model of the electrocardiosignal itself (ECS). According to the study [15], a mathematical model of an electrocardiosignal that is conditional cyclic random process is called a process

\[ \{(\omega, \omega', t), \omega \in \Omega, \omega' \in \Omega', t \in R \} \]

, which is given on the cartesian product of two stochastically independent probabilistic spaces with sets of elementary events \( \Omega \) and \( \Omega' \) on the set of real numbers \( R \), and for which the following conditions are met:

1) There is such a random function \( T(\omega', t, n), \omega' \in \Omega', t \in R, n \in Z \), what for each \( \omega' \), appropriate \( \omega' \)-realisation \( T_{\omega'}(t, n) \) of this function satisfies the conditions of the rhythm function;

2) For each \( \omega' \) is \( \Omega' \) finite-dimensional vectors (\( \xi_{\omega'}(\omega, t_1), \xi_{\omega'}(\omega, t_2), \ldots, \xi_{\omega'}(\omega, t_k) \)) and (\( \xi_{\omega'}(\omega, t_1 + T_{\omega'}(t_2, n)) \)) \( \xi_{\omega'}(\omega, t_2 + T_{\omega'}(t_2, n)), \ldots, \xi_{\omega'}(\omega, t_k + T_{\omega'}(t_k, n)) \), \( n \in Z \), where \( \{t_1, t_2, \ldots, t_k\} \)-multiple separabilities of the process \( \xi_{\omega'}(\omega, t), \omega' \in \Omega', \omega \in \Omega, t \in R \), with all the goals \( k \in N \) is stochastically equivalent in a broad sense;

3) For any different \( \omega'_1 \in \Omega' \) and \( \omega'_2 \in \Omega' \) random processes \( \xi_{\omega'}(\omega, t) \) and \( \xi_{\omega'}(\omega, t) \) are isomorphic to the order and values cyclic random processes.

Realisation of (\( \omega' \)-realisation) a random function \( T(\omega', t, n) \) there is a deterministic function \( T_{\omega'}(t, n) \), which satisfies the conditions of the rhythm function, namely:

- a group of conditions:
  a) \( T_{\omega'}(t, n) > 0 \), if \( n > 0 \) \( T_{\omega'}(t, 1) < \infty \);
b) \( T_{\omega'}(t, n) = 0 \), if \( n = 0 \);

c) \( T_{\omega'}(t, n) < 0 \), if \( n < 0, t \in \mathbb{R} \); for any \( t_1 \in \mathbb{R} \) and \( t_2 \in \mathbb{R} \) for which \( t_1 < t_2 \), for the function \( T_{\omega'}(t, n) \) a strict inequality is fulfilled \( T_{\omega'}(t_1, n) + 1 < T_{\omega'}(t_2, n) + 2, \forall n \in \mathbb{Z} \).

3) The function \( T_{\omega'}(t, n) \) is the smallest in modulus \( \{ |T_{\omega'}(t, n)| \leq |T_{\omega'}(t, n)| \} \) among all such functions \( \{ T_{\omega'}(t, n), \gamma' \in \Gamma \} \), which satisfy the above conditions 1 and 2.

### 2.2. A Generalized Mathematical model of a High-resolution Rhythmocardiosignal

The conditional cyclic random process \( \xi(\omega, \omega', t) \) allows simultaneous consideration of both the stochasticity of the morphological structure of electrocardiosignals (which is important for their statistical morphological analysis), as well as the stochasticity of their rhythmic structure (which is important for the heart rate analysis). Considering that according to such a mathematical model of the electrical signal, information about the heart rate is contained in the rhythm function \( T(\omega', t, n) \) of the conditional cyclic random process \( \xi(\omega, \omega', t) \), and also taking into account the fact that the processing of electrocardiosignals is carried out in a digital system, the analysis of heart rate is reduced to the statistical analysis of the random rhythm function \( T(t_{ml}(\omega'), n), \omega' \in \Omega', t_{ml}(\omega') \in \mathbb{R}, n \in \mathbb{Z} \) of the conditional cyclic random process discrete argument \( \{ \xi(\omega, t \quad (\omega'), \omega' \in \Omega', \omega \in \Omega, t \quad (\omega') \in D(\omega') \} \).

Random rhythm function \( T(t_{n}(\omega'), n) \) is completely defined by the elements of the random domain \( D(\omega') \) according to the formula:

\[
T(t_{ml}(\omega'), n) = t_{m+1,l}(\omega') - t_{ml}(\omega'), \quad m, n \in \mathbb{Z}, \quad l = 1, L, \quad t_{ml}(\omega') \in D(\omega')
\]

When \( n = 1 \), rhythm function \( T(t_{n}(\omega'), 1) \) is calculated in the following way:

\[
T(t_{ml}(\omega'), 1) = t_{m+1,l}(\omega') - t_{ml}(\omega'), \quad m \in \mathbb{Z}, \quad l = 1, L, \quad t_{ml}(\omega') \in D(\omega')
\]

For each \( \omega' \)-realisation \( \omega' \)-realisation \( D_{\omega'} = \{ t_{ml}(\omega') \in \mathbb{R}, m \in \mathbb{Z}, l = 1, L, L \geq 2 \} \) of the random domain definition \( D_{\omega'} = \{ t_{ml}(\omega') \in \mathbb{R}, m \in \mathbb{Z}, l = 1, L, L \geq 2 \} \) of the conditional cyclic random process of a discrete argument that is given on a probabilistic space \( (\Omega', F', P') \), the following conditions apply: \( t_{ml,1} \) < \( t_{ml,2} \), whether \( m \), or whether \( l_1 < l_2 \), in other cases \( t_{ml,1} < t_{ml,2} \).

\[
\Xi_L(\omega', m) = \{ T_l(\omega', m), \omega' \in \Omega', l = 1, L, m \in \mathbb{Z} \}
\]

where each \( l \)-component of a vector is a random sequence \( T_l(\omega', m) \), the value of which is equal to the value of the random rhythm function \( T(t_{n}(\omega'), 1) \) at moments in time \( t_{n}(\omega') \) from a discrete set \( D_l(\omega') = \{ t_{ml}(\omega') \in \mathbb{R}, m \in \mathbb{Z}, l = const \} \). The set of \( D(\omega') \) is integrated into the \( D(\omega') \) and describes the time distances between the same type \( l \) - phases of the studied electrocardiosignal in its two adjacent cycles, namely:

\[
T_l(\omega', m) = T(t_{ml}(\omega'), 1) = t_{m+1,l}(\omega') - t_{ml}(\omega'), \quad m \in \mathbb{Z}, \quad l = 1, L, \quad t_{ml,l}(\omega') \in D(\omega')
\]

Dimension (the number of components) \( L \) of vector \( \Xi_L(\omega', m) \) determines the resolution of the rhythmocardiosignal and is equal to the number of studied time intervals between pre-selected phases in the electrocardiosignal, which can be identified by segmentation and detection methods when solving the problem of automatic formation of the rhythmocardio signal from the electrocardiosignal [16 - 28]. According to the block diagram shown in Fig. (1), the first block is the determination of the same type of phases corresponding to the boundaries of segments-zones ECS; this stage is implemented on the basis of the use of methods for segmenting cyclic signals. Detection of the same type of phases within certain zones is the next step in the formation of a vector rhythmocardiosignal. At this stage, information is obtained about time points that correspond to the maximum or minimum of characteristic ECS segments, for example, R, P, or T. The final stage in this structure is the formation of a vector rhythmocardiosignal based on the information obtained at the previous stages.
2.3. Updated Mathematical Model of a High-resolution Rhythmocardiosignal and its Probabilistic Characteristics

Let’s move on to substantiating the probabilistic characteristics of the vector $\Xi_l(\omega', m)$ of random sequences. One of the simplest stochastic models that can take into account the dynamics of changes in the rhythmocardiosignal with increased resolution is the vector $\Xi_L(\omega', m) = \{T_l(\omega', m), \omega' \in \Omega', l = 1, L, m \in Z\}$ describing its probabilistic structure, but methods for statistical estimation of the function $F_{\mathcal{P}_{T_l}}(x_1, ..., x_p, m_1, ..., m_p)$ they are too bulky for their practical use in computer diagnostic systems of the functional state of the cardiovascular system of the human body.

Let’s move on to substantiating the probabilistic characteristics of the vector $\Xi_L(\omega', m)$ random sequences. One of the simplest stochastic models that takes into account the dynamics of changes in the rhythmocardiosignal with increased resolution is the vector $\Xi_L(\omega', m) = \{T_l(\omega', m), \omega' \in \Omega', l = 1, L, m \in Z\}$ stationary and stationary-connected random sequences. First of all, note that the vector $\Xi_L(\omega', m)$ of the stationary and stationary-related random sequences, in the partial case, if its components are stationary sequences with independent values, i.e. white noises given on a set of integers, is a well-known model of a rhythmocardiosignal with increased resolution in the form of a random variable vector, which was developed in the studies. However, the hypothesis of independence or uncorrelation of rhythmocardiosignal readings does not correspond to reality, which requires taking into account the stochastic relationship between rhythmocardiosignal readings with increased resolution, and therefore the use of a more complex and more general mathematical model in the form of a vector $\Xi_L(\omega', m)$ stationary and stationary possible random sequences.

Defining property of the vector $\Xi_L(\omega', m)$ stationary and permanently connected random sequences are the invariance of its family of distribution functions to time shifts by an arbitrary integer $k \in Z$. For any distribution function $F_{\mathcal{P}_{T_l}}(x_1, ..., x_p, m_1, ..., m_p)$ of order $p (p \in N)$ of the family of vector distribution functions $\Xi_L(\omega', m)$ for stationary and permanently connected random sequences the following equality holds:

$$x_1, ..., x_p \in R, m_1, ..., m_p \in Z, l_1, ..., l_p \in \left\{1, \frac{1}{L}\right\}, k \in Z \quad (5)$$

Distribution function $F_{\mathcal{P}_{T_l}}(x_1, ..., x_p, m_1, ..., m_p)$ in the case when $l_1 = l_2 = ... = l_p = l$ is a distribution function $F_{\mathcal{P}_{T_l}}(x_1, ..., x_p, m_1, ..., m_p)$ of stationary components $T_l(\omega', m)$ of vector $\Xi_L(\omega', m)$ - that is an automatic order distribution function $\Xi_L(\omega', m)$, which describes the time distances between single-phase readings of an electrocardiosignal for its $l$-phase. If $p = 1$, then we will have a one-dimensional $F_{\mathcal{P}_{T_l}}(x, m)$ automatic distribution function of a stationary random sequence $T_l(\omega', m)$.

In the case where equality $l_1 = l_2 = ... = l_p = l$ if the distribution function is not executed $F_{\mathcal{P}_{T_l}}(x_1, ..., x_p, m_1, ..., m_p)$ is a compatible distribution function for several (at least two) stationary vector components $\Xi_L(\omega', m)$, which describes the time distances between single-phase readings of an electrocardiosignal as a whole for its different phases.

Family of vector distribution functions $\Xi_L(\omega', m)$ stationary and stationary-connected sequences must fully describe its probabilistic structure, but methods for statistical estimation of the distribution function $F_{\mathcal{P}_{T_l}}(x_1, ..., x_p, m_1, ..., m_p)$ they are too bulky for their practical use in computer diagnostic systems of the functional state of the cardiovascular system of the human body.

Fig. (1). Block diagram of the method of forming a vector rhythmocardiosignal with increased resolution.
Therefore, in addition to the vector distribution functions $\mathcal{Z}_t(\omega', m)$ effective is the use of instantaneous order functions $s = \sum_{j=1}^{p} s_j$, which, if they exist, are also invariant to time shifts (shifts by argument $m$).

$$c_{\tau_1 \tau_2} (m_1 + k, \ldots, m_p + k, m_1, \ldots, m_p \in Z, l_1, \ldots, l_p \in \{1, L\}, k \in Z,$$

where $M$ - operator of mathematical expectation.

If there is a mixed central moment function $r_{\tau_1 \tau_2} (m_1, \ldots, m_p)$ of order $s = \sum_{j=1}^{p} s_j$ of vector $\mathcal{Z}_t(\omega', m)$ of stationary and permanently connected random sequences, then the equality holds for it:

$$r_{\tau_1 \tau_2} (m_1, \ldots, m_p) = M \left\{ \left( T_{\tau_1} (\omega', m_1) - c_{\tau_1} \right)^s \ldots \left( T_{\tau_p} (\omega', m_p) - c_{\tau_p} \right)^s \right\} = r_{\tau_1 \tau_2} (m_1 + k, \ldots, m_p + k, m_1, \ldots, m_p \in Z, l_1, \ldots, l_p \in \{1, L\}, k \in Z.$$

where is the plural $\{c_{\tau_1 \tau_1}, \ldots, c_{\tau_p \tau_p}\}$ is a set of first-order initial moments (mathematical expectations) of stationary random sequences from the set $\{T_{\tau_1} (\omega', m), \ldots, T_{\tau_p} (\omega', m)\}$.

In practice, to analyze a rhythmocardiosignal with increased resolution, it is appropriate to use mixed moment functions of low orders, namely, mixed initial moment functions of the second-order – covariance functions and mixed central moment functions of the second-order – correlation functions. In this case, the initial moment functions of the second order for the vector $\mathcal{Z}_t(\omega', m)$ stationary and permanently connected random sequences are represented as a matrix of covariance functions:

$$C_T = \begin{bmatrix}
    c_{T_1 T_1} (m_1, m_2) & c_{T_1 T_2} (m_1, m_2) & \cdots & c_{T_1 T_p} (m_1, m_2) \\
    c_{T_2 T_1} (m_1, m_2) & c_{T_2 T_2} (m_1, m_2) & \cdots & c_{T_2 T_p} (m_1, m_2) \\
    \vdots & \vdots & \ddots & \vdots \\
    c_{T_2 T_1} (m_1, m_2) & c_{T_2 T_2} (m_1, m_2) & \cdots & c_{T_2 T_p} (m_1, m_2) 
\end{bmatrix},$$

what can be noted more compactly like this:

$$C_T = \begin{bmatrix}
    c_{T_1 T_1} (m_1, m_2), \quad l_1, l_2 = 1, L
\end{bmatrix},$$

where each of its elements is a covariance function $c_{T_1 T_2} (m_1, m_2)$, which is set as:

$$c_{T_1 T_2} (m_1, m_2) = M \{ T_{\tau_1} (\omega', m_1) \cdot T_{\tau_2} (\omega', m_2) \}, m_1, m_2 \in Z, l_1, l_2 \in \{1, L\}.$$

Since the components of the vector $\mathcal{Z}_t(\omega', m)$ random sequences are stationary and permanently connected sequences, then their covariance functions are functions of only one integer argument $u$, which is equal to $u = m_1 - m_2$. Therefore the covariance matrix of this random vector can be represented as follows:

$$C_T = \begin{bmatrix}
    c_{2T_1 T_2} (u), \quad l_1, l_2 = 1, L
\end{bmatrix},$$

where each of its elements is a covariance function $c_{2T_1 T_2} (u)$ which is equal to:

$$c_{2T_1 T_2} (u) = c_{T_1 T_2} (m_1 - m_2), \quad u, m_1, m_2 \in Z, \quad l_1, l_2 \in \{1, L\}.$$

Provided that $l_1 = l_2 = l$, covariance function $c_{\tau_1 \tau_2} (u)$ is an auto-variational function $l$ stationary components $T_l (\omega', m)$ of vector $\mathcal{Z}_t(\omega', m)$ which describes the time distances between single-phase readings of an electrocardiosignal for its phase. If $l_1 \neq l_2$, then the covariance function $c_{2T_1 T_2} (u)$ is a mutual covariance function for two stationary vector components $\tilde{Z}_t(\omega', m)$ which describe the time distances between single-phase electrocardiosignals for $l_1$.
and $l_1$ phase.

Mixed second-order central moment functions for a vector

$$ R_T = \begin{bmatrix} r_{2\tau_1 \tau_1} (m_1, m_2) & r_{2\tau_1 \tau_2} (m_1, m_2) & \cdots & r_{2\tau_1 \tau_p} (m_1, m_2) \\ r_{2\tau_1 \tau_1} (m_1, m_2) & r_{2\tau_2 \tau_2} (m_1, m_2) & \cdots & r_{2\tau_2 \tau_p} (m_1, m_2) \\ \vdots & \vdots & \ddots & \vdots \\ r_{2\tau_p \tau_1} (m_1, m_2) & r_{2\tau_p \tau_2} (m_1, m_2) & \cdots & r_{2\tau_p \tau_p} (m_1, m_2) \end{bmatrix}, $$

which can be written more compactly like this:

$$ R_T = \begin{bmatrix} r_{2\tau_1 \tau_1} (m_1, m_2), & l_1, l_2 = 1, L \end{bmatrix}, $$

where each of its elements is a correlation function $r_{2\tau_1 \tau_l} (m_2, m_2)$, which is set as follows:

$$ r_{2\tau_1 \tau_l} (m_1, m_2) = M \left\{ \left( T_{l_1} (\omega', m_1) - c_{1\tau_l} \right) \cdot \left( T_{l_2} (\omega', m_2) - c_{1\tau_l} \right) \right\}, \ m_1, m_2 \in Z, \ l_1, l_2 \in \{1, L\} $$

Since the components of the vector $\mathbf{z}_l (\omega', m)$ random sequences are stationary and stationary connected sequences, then their correlation functions are functions of only one integer argument $u$, which is equal to $u = m_1 - m_2$. Therefore the correlation matrix of this random vector can be represented as:

$$ R_T = \begin{bmatrix} r_{2\tau_1 \tau_2} (u), & l_1, l_2 = 1, L \end{bmatrix}, $$

where each of its elements is a correlation function $r_{2\tau_1 \tau_2} (u)$, which is equal to:

$$ r_{2\tau_1 \tau_2} (u) = r_{2\tau_1 \tau_2} (m_1 - m_2), \ u, m_1, m_2 \in Z, \ l_1, l_2 \in \{1, L\}. $$

If $l_1 = l_2 = l$, are correlation function $r_{2\tau_l \tau_l} (u)$ is an autocorrelation function $l$-stationary components $T_l (\omega', m)$ of vector $\mathbf{z}_l (\omega', m)$, which describes the time distances between single-phase readings of an electrocardiosignal for its $l$-phase.

If $l_1 \neq l_2$, then the correlation function $r_{2\tau_1 \tau_2} (u)$ is a mutual correlation function for two stationary components of the vector $\mathbf{z}_l (\omega', m)$ which describe the time distances between single-phase electrocardiosignals for $l_1$ and $l_2$ phase.

**2.4. Statistical Estimates of Probabilistic Characteristics of a High-resolution Rhythmocardiosignal**

Let’s write down the formula expressions for calculating the implementations of statistical estimates of probabilistic characteristics of a rhythmocardiosignal with increased resolution. The formula expression for calculating the implementation of a statistical estimate $\mathbf{z}_{\tau_1 \tau_2} (m_1, m_2)$ of covariance function $\mathbf{c}_{\tau_1 \tau_2} (m_1, m_2)$ of two stationary and stationary-related random sequences $T_{\tau_1} (\omega', m)$ and $T_{\tau_2} (\omega', m)$, which describe the time distances between single-phase electrocardiosignals for $l_1$-th and $l_2$-th phases, namely:

$$ M \cdot 1 \sum_{k=0}^{M-1} T_{\tau_1} (m_1 + k) \cdot T_{\tau_2} (m_2 + k), \ m_1, m_2 \in \{1, M-1\}, \ l_1, l_2 \in \{1, L\} $$

If in the formula (18) $p = 1$, then $l_1 = l_2 = \cdots = l_1 = l$, then we get an expression for calculating the implementation of a statistical estimate $\mathbf{c}_{\tau_1 \tau_1}$ of the initial moment ($l$-order $T_{\tau_1} (\omega', m)$) and $T_{\tau_2} (\omega', m)$ of the stationary random sequence
\( T_1(\omega', m) \) namely:

\[ \hat{c}_{T_1} = \frac{1}{M-M_1+1} \sum_{k=0}^{M-M_1} T_1^{s}(k), \ 1 \in \{1, L\} \]  

(19)

If in the formula (19) \( s = 1 \), then we get an expression for calculating the implementation of a statistical estimate \( c_{T_1} \) of the initial moment of the first order \( c_{T_1} \) (mathematical expectation) of a stationary random sequence \( T_{(\omega', m)} \), namely:

\[ \hat{c}_{T_1} = \frac{1}{M-M_1+1} \sum_{k=0}^{M-M_1} T_1^{s}(k), \ 1 \in \{1, L\} \]  

(20)

The formula expression for calculating the implementation of a statistical estimate of the correlation function \( r_{T_1} \) of two stationary and stationary-related random sequences \( T_{(\omega', m)} \) and \( T_{(\omega', m)}' \) which describe the time distances between single-phase electrocardiosignals for \( l_1 \) and \( l_2 \) phases, namely:

\[ r_{T_1}(m_1, m_2) = \frac{1}{M-M_1+1} \sum_{k=0}^{M-M_1} \left( T_{1}(m_1+k) - \hat{c}_{T_1} \right) \left( T_{1}(m_2+k) - \hat{c}_{T_1} \right), \ m_1, m_2 \in \{1, M\}, \ 1_1, 1_2 \in \{1, L\} \]  

(21)

Since for stationary and stationary-related random sequences, the correlation functions are the functions of only one integer argument \( u \), which is equal to \( u = m_1 - m_2 \), then

\[ r_{T_1}(u) = \frac{1}{M-M_1+1} \sum_{k=0}^{M-M_1} \left( T_{1}(k+u) - \hat{c}_{T_1} \right) \left( T_{1}(k+u) - \hat{c}_{T_1} \right), \ u = 0, M_1 - 1, \ m_1, m_2 \in \{1, M\}, \ 1_1, 1_2 \in \{1, L\} \]  

(22)

If in the formula (22) \( u = 0 \), then \( l_1 = l_1 = l \), then we have the expression for calculating the implementation of the variance estimate of the stationary random sequence \( T_{(\omega', m)} \), namely:

\[ \hat{p}_{T_1} = \frac{1}{M-1} \sum_{k=1}^{M} \left( T_{1}(\omega', k) - c_{T_1} \right)^2, \ 1 \in \{1, L\}. \]  

(23)

In order to reduce the number of diagnostic signs for a high-resolution rhythmocardiogram it is necessary to take into account the fact of symmetry of the matrix of correlation functions

\[ \sigma_{2T_1}(u) = \sigma_{2T_1}(u), \ l_1, l_2 = 1, L \]  

of the estimated matrix of correlation functions

\[ \hat{R}_T = \left[ \hat{p}_{2T_1}(u), \ l_1, l_2 = 1, L \right], \]  

which indicates the adequacy evaluation of only those elements of the matrix \( \hat{R}_T \) which lie on its diagonal and above the diagonal, namely, such an ordered totality

\[ \hat{R}_T = \left[ \hat{p}_{2T_1}(u), \ l_1 = 1, L, l_2 = 1, L \right]. \]  

On the diagonal of this matrix, when \( l_1 = l_2 \), estimates of autocorrelation functions are placed, and the elements of matrix \( \hat{R}_T \), which are placed above its diagonal, namely, when \( l_1 = l_2 \), are estimates of inter-correlation functions. Therefore, the matrix

\[ \hat{R}_T = \left[ \hat{p}_{2T_1}(u), \ l_1, l_2 = 1, L \right], \]  

without losing its information content, can be replaced with a triangular matrix

\[ \hat{R}_T = \left[ \hat{p}_{2T_1}(u), \ l_1 = 1, L, l_2 = 1, L \right]. \]  

Another way to reduce the number of diagnostic features in information systems for the analysis of heart rate for the main rhythmocardiograms with increased resolution is to use spectral decompositions of the triangular matrix elements of themselves

\[ \hat{R}_T = \left[ \hat{p}_{2T_1}(u), \ l_1 = 1, L, l_2 = 1, L \right] \]  

in particular, by using the discrete Fourier transform of autocorrelation estimates and inter-correlation functions from this matrix. Namely, instead of a triangular matrix

\[ \hat{R}_T = \left[ \hat{p}_{2T_1}(u), \ l_1 = 1, L, l_2 = 1, L \right] \]  

of the correlation functions a triangular matrix can be used the elements of which are Fourier images of the corresponding estimates of the correlation functions from the matrix \( \hat{R}_T \). Namely, Fourier images from The Matrix \( \hat{S}_T \) are calculated like this:

\[ \hat{S}_{2T_1}(u) = \sum_{u=0}^{M_1-1} \hat{p}_{2T_1}(u) \cdot e^{\frac{-j\omega u}{M_1}}, \ v = 0, M_1 - 1, \ l_1 = 1, L, \ l_2 = 1, L, \ j = \sqrt{-1} \]  

(24)
Based on Bessel's inequality, we will not choose the entire set as diagnostic signs $\{s_{2\nu_1\nu_2}(\nu), \nu = 0, M_1 - 1\}$ function counts $s_{2\nu_1\nu_2}(\nu)$, but only a certain subset of their first ones $M_2 (M_2 << M_1)$ of counts $s_{2\nu_1\nu_2}(\nu), \nu = 0, M_2 - 1$ which contribute to the full energy of evaluation $\hat{f}_{2\nu_1\nu_2}(u)$ of the correlation function not less than 95%.

3. RESULTS AND DISCUSSION

Based on the above mathematical model and methods of processing a high-resolution rhythmocardiosignal, a multifunctional software package for modeling and automated analysis of a wide class of cyclic heart signals for the needs of functional medical diagnostics has been upgraded. Namely, as a component of this software package, a system of computer programs has been developed for the automated formation and statistical analysis of heart rate based on a vector rhythmocardiosignal (rhythmocardiosignal with increased resolution), which expanded the functionality of the existing software package and made it possible to automatically analyze the heart rate with increased information content. A typical structural and functional diagram of the software for processing ECS is shown in Fig. (2). A dashed line in this block diagram highlights the blocks that are emphasized in this article. The software package is implemented in the programming language Object Pascal.

According to the blocks presented in the block diagram, ECS processing includes evaluating the segmental structure using segmentation methods, for example [29]. Evaluation of the rhythm function by interpolating the rhythmic structure (discrete rhythm function), based on the method [30].

Further, the development of the ECS branches out into two stages (two problems are solved). The first stage performs morphological analysis, which, according to this structure, provides for statistical processing of ECS, normalization of statistical estimates and their decomposition in the Chebyshev basis, and decision-making based on the obtained morphological features. This stage is described in [31]. The second stage performs rhythm analysis and consists in forming a vector rhythmocardiosignal, statistical processing of the vector and spectral analysis of the obtained statistical estimates.

As an example, Fig. (3) shows a general view of the program interface for evaluating the autocorrelation function and the cross-correlation function of the components of a vector rhythmocardiosignal.

**Fig. (2).** Structural and functional diagram of software for heart rate analysis with increased information content.
Fig. (3). Example of a program interface for evaluating the autocorrelation function and the cross-correlation function of a vector rhythmocardiosignal component.

Figs. (4 and 5) show the stages of formation from the ECS vector rhythmocardiosignal, and Figs. (6 and 7) show the results of statistical processing of the rhythmocardiosignal with increased information content, by statistical evaluation of its corresponding statistical characteristics corresponding to the blocks of the structural and functional scheme 2.

![Graphs showing statistical processing results](image)

Fig. (4). The results of processing: a) several cycles of the studied electrocardiosignal; b) the rhythmic structure of the electrocardiosignal.
Fig (5). results of processing A) graphs of relative errors in the formation of high-resolution rhythmocardiogram samples corresponding to the R-R intervals of the electrocardiogram; B) graphs of relative errors in the formation of high-resolution rhythmocardiogram samples corresponding to the T-T intervals of the electrocardiogram.

Fig. (6). Schedule of implementation $T_{1\omega'}(m)$, $T_{2\omega'}(m)$, component of the vector rhythmocardiosignal of the first component $T_1(\omega', m)$ and the second component $T_2(\omega', m)$, that describing the duration accordingly: a) P - intervals of electrocardiosignal; b) R - intervals of electrocardiosignal.

Fig. (7). Histograms of implementation $T_{1\omega'}(m)$, $T_{2\omega'}(m)$ component of the vector rhythmocardiosignal of the first component $T_1(\omega', m)$ and the second component $T_2(\omega', m)$, describing the duration accordingly: a) P - intervals of electrocardiosignal; b) R - intervals of electrocardiosignal.

Fig. (8) shows the results of the spectral decomposition of statistical estimates of the power spectral densities of the components of the vector rhythmocardiosignal.
Fig. (8). Schedule of implementations \( \hat{r}_{2T_1}^{T_1}(u) \) and \( \hat{r}_{2T_2}^{T_2}(u) \) statistical estimates of autocorrelation functions \( r_{2T_1}^{T_1}(u) \) and \( r_{2T_2}^{T_2}(u) \) of the first component \( T_1(\omega', m) \) and the second component \( T_2(\omega', m) \), what describing the duration accordingly: a) \( P \) - intervals of electrocardiosignal; b) \( R \) - intervals of electrocardiosignal.

Fig. (5A) shows graphs of relative errors in the formation of high-resolution rhythmocardiogram samples corresponding to R–R intervals and obtained on the basis of the method of segmentation and detection of extreme values of electrocardiogram zones, based on Brodsky-Darkhovsky statistics (indicated on the graph with bold dots) and on the basis of the method based on the use of a first-order difference function (indicated on the graph with triangles). Fig. (5B) shows graphs of relative errors in the formation of high-resolution rhythmocardiogram samples corresponding to T–T intervals, and obtained on the basis of the method of segmentation and detection of extreme values of electrocardiogram zones, based on Brodsky-Darkhovsky statistics (indicated on the graph with bold dots) and on the basis of the method based on the use of a first-order difference function (indicated on the graph with triangles).

Analyzing the graphs of relative errors in the formation of a high-resolution rhythmocardiogram, which are presented in Fig. (5), it can be argued that the method of automatic formation of a high-resolution rhythmocardiogram, which is based on Brodsky-Darkhovsky Statistics, has higher accuracy compared to a similar method based on the use of a first-order difference function.

From the registered electrocardiogram according to the method of automatic formation of a rhythmocardiogram with increased accuracy, the implementation of \( \mathbf{\Sigma}_4(\omega', m) = \{ T_1(\omega', m), \omega' \in \Omega', l = 1,4, m = 1,245 \} \) of four components of the vector \( \mathbf{\Sigma}_4(\omega', m) = \{ T_1(\omega', m), \omega' \in \Omega', l = 1,4, m = 1,245 \} \) stationary and stationary related random sequences. The first component of \( T_1(\omega', m) \) this vector is a random stationary sequence describing durations \( P \) in the electrocardiosignal for all its 245 recorded cycles. Second component \( T_2(\omega', m) \) of this vector is a random stationary sequence describing durations \( R \) in the electrocardiosignal. Third component \( T_3(\omega', m) \) this vector is a random stationary sequence describing durations \( T \) in the electrocardiosignal. Fourth component \( T_4(\omega', m) \) of this vector is a random stationary sequence describing durations R–R - intervals in the electrocardiosignal. As an example, the implementation schedule \( T_{1\omega}(m) \) of the first component is shown in Fig. (6) A, and the implementation graph \( T_{2\omega}(m) \) of the second component is shown in Fig. (6B).

The justification of statistical hypotheses about the stationarity of the mathematical expectation and variance of the components of the vector rhythmocardiosignal have been tested. Namely, the statistical hypotheses about the invariance of the mathematical expectation and variance of the components of the vector rhythmocardiosignal have been tested by applying well-known statistical criteria for checking the equality of mathematical expectations and variances of two random variables represented by their samples (as a sample, two sections of each component of the vector rhythmocardiosignal were taken. The Student's criterion (for mathematical expectation of the vector rhythmocardiosignal component) and Fischer's criterion (for variance of the vector rhythmocardiosignal component) have been used as a statistical criterion for testing hypotheses about stationarity. The results of 13 of the 15 tests performed with a confidence level of 0.95 indicate the consistency of the hypothesis about the stationarity of the components of the vector rhythmocardiosignal, which can be considered verification of a new mathematical model of the rhythmocardiosignal with increased resolution in the form of a vector of stationary and stationary-related random sequences.

To check the stationary components of the vector for their normality, Fig. (7) shows histograms for implementations \( T_{\omega}(m), T_{\omega}(m) \) corresponding stationary vector components \( \mathbf{\Sigma}_4(\omega', m) \).
Table 1. Shows the results of statistical evaluation of mathematical expectations of stationary vector components $\mathbf{\Xi}_x(\omega', m)$.

| Stationary Component Number | Significance of Implementing a Statistical Estimate of Mathematical Expectation |
|-----------------------------|--------------------------------------------------------------------------------|
| 1                           | $C_1 = 14.88$                                                                     |
| 2                           | $C_2 = 25.02$                                                                     |
| 3                           | $C_3 = 73.82$                                                                     |
| 4                           | $C_4 = 799.51$                                                                    |

Fig. (9). Schedule of implementations $\hat{S}_{2T_1T_1}(v)$ and $\hat{S}_{2T_2T_2}(v)$ statistical estimates of power spectral densities $S_{2T_1T_1}(v)$ and $S_{2T_2T_2}(v)$ of the first component $T_1(\omega', m)$ and the second component $T_2(\omega', m)$, what describing the duration accordingly: a) $P$- intervals of electrocardiosignal; b) $R$- intervals of electrocardiosignal.

Fig. (8). Shows graphs of implementations $\hat{r}_{2T_1T_1}(u)$ and $\hat{r}_{2T_2T_2}(u)$ statistical estimates of autocorrelation functions $r_{2T_1T_1}(u)$ and $r_{2T_2T_2}(u)$ of the first component $T_1(\omega', m)$ and the second component $T_2(\omega', m)$, what describing the duration accordingly: a) $P$- intervals of electrocardiosignal; b) $R$- intervals of electrocardiosignal.

To test the hypothesis of the normality of the distribution of stationary components of a random vector $\mathbf{\Xi}_x(\omega', m)$ according to the Pearson consent criterion, it has been found that these results do not contradict the hypothesis of the normality of its distribution. Normality of the vector distribution $\mathbf{\Xi}_x(\omega', m)$ is the basis for substantiating diagnostic features in systems for the analysis of heart rate using a high-resolution rhythmocardiogram within the framework of spectral correlation theory, which significantly reduces the computational complexity of such an analysis. In this case, to estimate the probabilistic structure of the vector $\mathbf{\Xi}_x(\omega', m)$ stationary and stationary-related random sequences, it is sufficient to perform a statistical estimation of its mathematical expectations according to Formula (20) and the matrix of correlation functions $R_T = \begin{bmatrix} r_{2T_1T_1}(u), & l_1, l_2 = 1, L \end{bmatrix}$ according to the Formula (22).

Fig. (9) shows graphs of implementations $\hat{S}_{2\gamma_1\gamma_1}(v)$ and $\hat{S}_{2\gamma_2\gamma_2}(v)$ statistical estimates of power spectral densities $S_{2\gamma_1\gamma_1}(v)$ and $S_{2\gamma_2\gamma_2}(v)$ of the first component $\gamma_1(\omega', m)$ and the second component $\gamma_2(\omega', m)$, what describing the duration accordingly: a) $P$- intervals of electrocardiosignal; b) $R$- intervals of electrocardiosignal.

CONCLUSION

The paper presents a new mathematical model of a rhythmocardiogram with increased resolution in the form of a vector of stationary and stationary-related random sequences, which, in comparison with the known mathematical models of heart rate, allows to increase the level of information content of automated heart rate analysis and is logically consistent with...
the stochastic mathematical model of an electrocardiosignal in the form of a conditional cyclic random process. The mathematical model of a high-resolution rhythmocardiosignal has been verified by testing the statistical hypotheses about stationarity for the normal distribution of components of a high-resolution rhythmocardiosignal, which has been the basis for reducing the computational complexity of statistical methods for the analysis of heart rate in computer systems of medical diagnostics.

The statistical methods for the analysis of high-resolution rhythmocardiosignals have been developed, which are based on their new mathematical model in the form of a vector of stationary and permanently connected random sequences, namely, expressions are recorded for calculating implementations of statistical estimates of the vector of mathematical expectations and a matrix of correlation functions of components of the vector of the rhythmocardiosignal. A number of new diagnostic features in computer systems of medical diagnostics based on vector rhythmocardiosignals for assessing the state of the cardiovascular system and adaptive-regulatory mechanisms of the human body as a whole have been justified. Namely, the known diagnostic features of the vector rhythmocardiosignal are supplemented with such new diagnostic features as the matrix of correlation functions and the matrix of spectral power densities of stationary components of the high-resolution rhythmocardiosignal, which by reflecting the stochastic temporal dynamics of the heart rate make it possible to increase the level of information content of heart rate analysis in modern cardiodiagnostic systems.

Based on the new mathematical model and methods developed in the dissertation for processing high-resolution rhythmocardiosignals, a multifunctional software package for modelling and automated analysis of a wide class of cyclic heart signals for the needs of functional medical diagnostics has been upgraded. Namely, as a component of this software package, a system of computer programs has been developed for automated formation and statistical analysis of heart rate based on a vector rhythmocardiosignal, which expanded the functionality of the existing software package and made it possible to perform heart rate analysis with increased information content year analysis systems.

**CONFLICT OF INTEREST**

The authors declare no conflict of interest, financial or otherwise.

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