Markov Chains application to the financial-economic time series prediction

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In this research the technology of complex Markov chains is applied to predict financial time series. The main distinction of complex or high-order Markov Chains and simple first-order ones is the existing of aftereffect or memory. The technology proposes prediction with the hierarchy of time discretization intervals and splicing procedure for the prediction results at the different frequency levels to the single prediction output time series. The hierarchy of time discretizations gives a possibility to use fractal properties of the given time series to make prediction on the different frequencies of the series. The prediction results for world’s stock market indices is presented.

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1. INTRODUCTION

Successful modeling and prediction of processes peculiar to complex systems, such as ecological, social, and economical (ESE) ones, remain one of the most relevant problems as applied to the whole complex of natural, human and social sciences ([1–6]).

The diversity of possible approaches to modeling such systems and, usually, more than modest success in the dynamics prediction, compel us to look for the reasons of failure, finding them not only in details, but also in the axiomatics, which relates to problem statement, chosen modeling methods, results interpretation, connections with other scientific directions.

With the appearance of quantum mechanics and relativity theory in early twentieth century new philosophical ideas on physical values, measuring procedures and system state have been established, the ones that are completely different from Newtonian notions [7, 8].

For more than 70 years basic concepts of classical and neoclassical economic theories have been discussed by leading scientists, generating new approaches [9]. The general systems theory has acquired recognition in the middle of the 20th century giving way to development
of the new, systemic, emergent, and quantum in essence approach to investigation of complex objects, which postulates the limited nature of any kind of modeling and is based upon fixed and closed system of axioms [10].

However, the development of this new philosophical basis of ESE systems modeling is still accompanied with numerous difficulties, and new principles are often merely declared.

Current research is devoted to investigation and application of the new modeling and prediction technology, suggested in [11, 12], based on concepts of determined chaos, complex Markov chains and hierarchic (in terms of time scale) organization of calculating procedures.

2. ANALYSIS OF PROMINENT PUBLICATIONS RELEVANT TO THE SUBJECT

Prediction of financial-economic time series is an extremely urgent task. Modern approaches to the problem can be characterized by the following directions: 1) approximation of a time series using an analytical function and extrapolation of the derived function towards future – so-called trend models [13]; 2) investigation of the possible influence various factors might have on the index, which is being predicted, as well as development of econometric or more complicated models using the Group Method of Data Handling (GMDH) [3, 14]; 3) modeling future prices as the decisions-making results using neuronal networks, genetic algorithms, fuzzy sets [14–16].

Unfortunately, these techniques don’t produce stable forecasts, what can be explained by complexity of the investigated systems, constant changes in their structure. Although we are trying to join these directions in one algorithm, it is the latter option that we prefer, with it consisting in creating a model adequate to the process generating a price time series [17]. This very approach gives a chance to approach the complexity of the system, which generates the observed series, develop the model and use its properties as the prognosis.

3. AIMS OF THE PAPER, PROBLEM STATEMENT

Assume the time series is set by a sequence of discrete levels with constant step of time sampling $\Delta t$. We need to generate variants of the time series continuation (prognosis scenarios) according to the relations between the sequences of absolute and relative changes
discovered with the help of complex Markov chains.

4. CLASSICAL MODELING PROBLEMS OF ESE SYSTEMS DYNAMICS

Another peculiar feature of ESE systems, apart from complexity, is a memory, including the long-term one, as well as nonlinear and unstable nature of interactions and components, which makes it harder to predict their future behavior.

Unfortunately, mathematical models based on differential equations have no memory (there is no aftereffect), while for models with memory, where integral interrelations are used, it is not always possible to take into account nonlinearity (the integration procedure is linear by definition).

In reality, in the Cauchy problem future systems behavior is defined by its initial state and doesn’t depend on the way the system reached its current state. However, it is hardly true that future behavior of a real socio-economic or socio-ecological system can be predicted by giving an immediate time “slice” of a variables set that describe its state.

Let us consider possible ways to take into account past events while modeling ESE systems’ dynamics, which goes beyond the boundaries of classical differential and integral equations.

Functional differential lagging equation can serve as a simple example of the dynamic model with memory, where present time is defined by the state variable \( x(t) \) and depends on the past state \( x(t - \tau) \) with constant time lag \( \tau = \text{const} \):

\[
x(t) = f(x(t - \tau)); t \geq t_0,
\]

where \( f(x) \) is the known function, with initial conditions being set for the half-interval \( t_0 - \tau \leq t < t_0 \) by the function \( \phi(t) \):

\[
x(t) = \phi(t); t_0 - \tau \leq t < t_0.
\]

Given the equation 1 has the only solution, defined by recurrent ratios:

\[
x(t) = \begin{cases} 
  f(\phi(t - \tau)); & \text{if } t_0 \leq t < t + \tau; \\
  f(f(\phi(t - \tau))); & \text{if } t_0 + \tau \leq t < t + 2\tau; \\
  f((f(\phi(t - \tau)))); & \text{if } t_0 + 2\tau \leq t < t + 3\tau; \\
  \ldots 
\end{cases}
\]
Using Dirac delta function, as defined by ratios:

\[
\delta(t) = 0, \text{ if } x \neq 0; \quad \int_{-\infty}^{+\infty} \delta(t) dt = 1, \tag{4}
\]

we can formally rewrite equation (1) in the integral form:

\[
x(t) = \int_{-\infty}^{t} dt_1 f(x(t_1)) H(t_1, t); H(t_1, t) \equiv \delta(t_1 - (t - \tau)) ; t \geq t_0. \tag{5}
\]

Delta function is not a function in the conventional interpretation and is related to the class of generalized functions that were mathematically described only in the middle of the last century \[18\] (physics started using this function much earlier). Its classical form is considered to be a limit of the “peak” sequence, with its centre set in the point of origin. The afore-mentioned “peaks” indefinitely converge widthway, indefinitely increase throughout the height and have a unit area.

An approximate classic integral analogue of the equation (5) can be derived by substituting \(\delta(t)\) with an ordinary function - some specific narrow enough “peak” of a unit area, a certain finite width \(\sim \Delta t\) as well as a finite height . The derivative of the Fermi function is one of the possible examples:

\[
\Phi(t) = \frac{1}{1 + e^{x(t)\theta}}; \delta(t) \approx \frac{1}{\theta \left(2 + e^{x(t)\theta} + e^{x(t)\theta}\right)}. \tag{6}
\]

If the system’s state in the moment \(t, x(t)\), is defined not by one, as in (1) but \(k (k = 2, 3, 4, \cdots)\) of her past states \(x(t - \tau_1), x(t - \tau_2), \cdots x(t - \tau_k)\) in the following moments of time \((t - \tau_1), (t - \tau_2), \cdots, (t - \tau_k)\) respectively \((\tau_1 = const, \tau_2 = const, \cdots, \tau_k = const, \tau_1 > \tau_2 > \cdots > \tau_k > 0)\), then instead of (1), (2), (5) we get:

\[
x(t) = f(x(t - \tau_1); x(t - \tau_2); \cdots ; x(t - \tau_k)) ; t \geq t_0; \tag{7}
\]

\[
x(t) = \phi(t); t_0 - \tau_1 \leq t < t_0; \tag{8}
\]

\[
x(t) = \int_{-\infty}^{t} dt_1 \int_{-\infty}^{t} dt_2 \cdots \int_{-\infty}^{t} dt_k f(x(t_1), x(t_2), \cdots, x(t_k)).
\]

\[
\delta ((t_1 - (t - \tau_1)) \delta ((t_2 - (t - \tau_2)) \cdots \delta ((t_k - (t - \tau_k)) ; t \geq t_0. \tag{9}
\]

\[
\delta ((t_1 - (t - \tau_1)) \delta ((t_2 - (t - \tau_2)) \cdots \delta ((t_k - (t - \tau_k)) ; t \geq t_0.
\]
Therefore if the system’s state in the time moment $t$ depends on the infinite sequence of its past states, the integral analogue of the functional differential lagging equation will, generally speaking, contain an integral of the infinite multiplicity. At the same time the infinite amount of past states can relate to both finite $(t - \tau_1; t)$ (short-term memory) and infinite $(-\infty; t)$ (long-term memory) time span.

Pay attention that the classic integral lagging equation is one of the Volterra type \[19\]:

$$x(t) = \int_{-\infty}^{t} F(x(\tilde{t}); t; \tilde{t}) \, d\tilde{t},$$ \hspace{1cm} (10)

where $F(x(\tilde{t}); t; \tilde{t})$ - is an arbitrary (generally nonlinear) function of variables $x(\tilde{t}); t; \tilde{t}$, which allows to take into account system’s memory of its past states only in the additive approximation, which becomes evident, if the right section \[10\] is rewritten in the following way:

$$\int_{-\infty}^{t} F(x(\tilde{t}); t; \tilde{t}) \, d\tilde{t} \equiv \int_{t_1}^{t} F(x(\tilde{t}_1); t; \tilde{t}_1) \, d\tilde{t}_1 + \int_{t_2}^{t_1} F(x(\tilde{t}_2); t; \tilde{t}_2) \, d\tilde{t}_2 + \cdots$$ \hspace{1cm} (11)

$$t > t_1 > t_2 > \cdots ; \tilde{t}_1 \in [t_1, t] ; \tilde{t}_2 \in [t_2, t_1] ; \cdots$$

In connection with it note that the equation \[9\] in case of an additive dependency of contemporaneity on the past, i.e. in case:

$$f(x(t - \tau_1); x(t - \tau_2); \cdots) \equiv f_1(x(t - \tau_1)); f_2(x(t - \tau_2)); \cdots$$ \hspace{1cm} (12)

becomes a particular case of the equation \[10\] with the following integrand:

$$F(x(\tilde{t}); t; \tilde{t}) \equiv f_1(x(\tilde{t})) \delta(\tilde{t} - (t - \tau_1)) + f_2(x(\tilde{t})) \delta(\tilde{t} - (t - \tau_2)) + \cdots$$ \hspace{1cm} (13)

Meaningful analysis of nonlinear models dynamics with memory, in which the future is defined by the infinite amount of states in the past is generally possible only in case of a discrete representation. The results of such analysis will be approximated, i.e. will contain uncertainty, which has to be considered endogenous, i.e. internal, and peculiar to this very system.

With a certain level of time sampling, models with memory both \[7\] and \[10\] becomes:

$$x(n + 1) = f(x(n); x(n - 1); x(n - 2) \cdots).$$ \hspace{1cm} (14)
To take into account and quantify the uncertainties, observed in ESE as well as other complex systems probability models are normally used. However their application is based on doubtful hypotheses, while the statistical interpretation of the results is not always informative enough and results might not correspond with the real process occurring within the system. In particular, the well-known problem of $1/f$–noise (look for example [20]), closely connected to the presence of long-term memory in complex systems, implies the absence of the mean temporary value (as a limit of a certain time span converging to infinity, which serves as the basis for averaging) for any process occurring in such kind of system. Therefore such processes can’t have a rigorous statistical substantiation.

5. MODERN CONCEPTS IN ESE SYSTEMS MODELING

New approaches to modeling and prediction of complex nonlinear systems dynamics with memory are based on the use of determined chaos and neural networks technologies (cf. e.g. [15, 21, 22]). Both investigation and realization of such techniques has become possible only with the appearance of quick-operating computers. Use of the recurrent computational process has become the general feature for all these technologies:

$$x_{n+1} = f_n (f_{n-1} (\cdots (f_1 (x_1) \cdots ))), \quad n = 1, 2, \cdots ,$$

(15)

where $f_i(x_i)$ is a certain nonlinear mapping of a multi-dimensional vector $x_i$, $i$ - discrete, real or fictitious, time. Identification of the model 15 is reduced to the determination of functions $f_i(x_i)$, while the differences between the models of determined chaos and neural networks are connected with the function type and methods of its definition (neural network models normally use rather narrow class of $f_i(x_i)$ mappings [16]). Generally speaking, stability or convergence of the process 15 is not required, whereas a single-step set of vector $x_i$ components as well as their time dynamics can be of great interest.

For the particular case of the model 15 introducing corresponding lagged variables, a model 14 can be transformed.

Both determined stable processes, described by integro-differential equations, and random processes, which also include complex Markov chains (CMC), can be formally considered as separate extreme cases of determined chaos models realization 15. Given the sampling scale, which tends to zero, if such a tendency makes sense and corresponding limits exist, we
derive classical differential and integral problem statement. Finite $\Delta t$ allows to get models with discrete time, which in the general case in the corresponding phase space (which also includes lagged variables) can produce both measurable sets (discrete or continuous) that allow probabilistic interpretation and those of the special structure – fractals \[23\], that can’t be always interpreted in that way.

Various digital generators of so-called random sequences used in imitational modeling can be an example of determined chaos models that allow probabilistic interpretation.

Let us note that in reality there are no accurate procedures that would give an opportunity to distinguish a “real” random sequence from the pseudorandom one.

6. MARKOV CHAINS PREDICTION TECHNOLOGY

Suppose there is a sequence of a certain system discrete states. From this sequence we can determine transitions probabilities between the two states. Simple Markov chain is a random process, in which the next state probability depends solely on the previous state and is independent from the rest of them. Complex Markov chain, unlike the simple one, stands for the random process, in which the next state probability depends not only on the current, but also on the sequence of several previous states (history). The amount of states in history is the order of the Markov chain.

Theory of simple Markov chains is widely presented in literature, for example \[24\]. As for the high order Markov chains, modern literature \[25\] can offer us a mere definition. Developing complex or high order Markov chain’s properties is not widely presented in modern scientific publications. It’s necessary to mention the papers \[26\] \[27\] where properties of complex Markov Chains are developed, but no prediction algorithm is proposed there. The development of prediction method, based on complex Markov chains, is proposed in this paper.

Markov chain of the higher order can be brought to a simple Markov chain by introducing the notion of a “generalized state” and including a series of consequent system’s states into it. In this case, tools of simple Markov chains can be applied to the complex ones.

Investigated dynamic series is a result of a certain process. It is assumed that this process is determined, which implies the existence of a causal dependence of further states on history. It is impossible to fix and analyze the infinite history, which puts obstacles in the way of an
accurate detection of this influence and making precise predictions.

The problem consists in the maximal use of information, which is contained in the known segment of the time series, and subsequent modeling of the most probable future dynamics scenario.

The observed process is described as a time series of prices $p_t$ with the given sampling time span $\Delta t$

$$p_{ti} = p(t_0 + i\Delta t).$$  \hspace{1cm} (16)

Discrete presentation of the time series is in fact a way of existence of this very system. New prices are formed on the basis of contracts or deals, made on the market in certain discrete moments of time, while the price time series is a series of the averaged price levels during the chosen time intervals. While making a decision each trader, who is an active part of the pricing system, works solely with discrete series of the chosen time interval (e.g. minute, 5-minute, hourly, daily etc.). For $\Delta t \rightarrow 0$ the accuracy of data presentations reaches a certain limit, since for relatively small $\Delta t$ the price leaps in the moment of deal, while staying unchanged and equal to the last deal during the time between the two deals. Hence, the discreteness of time series has to be understood not only as a limited presentation of activity of the complex financial system, but also as one of the principles of its operation [11, 12, 28–30].

The time series of initial conditions has to be turned into a sequence of discrete states. Let us denote the amount of chosen states as $s$, each of them being connected to the change in the quantity of the initial signal (returns). For example, consider the classification with two states, first of which corresponds to positive returns as the price increases, while the second one – to negative as it descends. Generally all possible increments of the initial time series are divided into $s$ groups. Ways of division will be discussed further.

Next we develop predictions for the time series of sampled states. For the given order of the Markov chain and the last generalized state the most probable state is chosen to be the next one. In case if ambiguity occurs while the state of maximum probability is being evaluated, an algorithm is used that allows reducing the amount of possible prediction scenarios. Therefore we get the series of predicted states that can be turned into a sampled sequence of prognostic values.

Evaluation of increments, prediction, and subsequent restoration are conducted for the given hierarchy of time increments $t$. To use the given information as effectively as possible,
The prediction is conducted for time increments \( t = 1, 2, 4, 8, \ldots \), or a more complex hierarchy of increments and subsequent “splicing” of the results derived from different prediction samplings.

The procedure of prediction and splicing is iterative and conducted starting from smaller increments, adding a prediction with the bigger time increment on every step.

As the sampling time step \( t \) increases, the statistics for the investigation of Markov chains decreases, whereas the biggest sampling step, which takes part in the prognostication, limits itself. To supplement the prediction with the low-frequency component the approximation of zero order is being used in the form of a linear trend or a combination of a linear trend and harmonic oscillations \([31, 32]\).

7. PREDICTION CONSTRUCTION ALGORITHM

Let us consider the consequence of operations, required for the prognostic time series construction. To do this we need to set the following parameters:

1) The type of time increments hierarchy (simple – powers of two, complex – product of powers of the first simple numbers).

2) Values of \( s \) – the amount of states and \( r \) – the order of the Markov chain. These parameters can be individual for every sampling level; finding of optimal parameters is done experimentally.

3) Threshold values \( \delta \), and minimal number of transitions \( N_{\text{min}} \).

Prediction construction algorithm includes the following steps:

1) Generating hierarchy of time increments - \( t \) sequence. The maximal of them has to correspond to the length of a prognostic interval \( N_{\text{max}} \).

2) For every time increment \( \Delta t \), as the increments increase, a prediction of states and restoration of the time series along the prognostic states is conducted. Current stage includes following actions:

   2.1. Evaluating increments (returns) of the series with \( \Delta t \) sampling.

   2.2. Transforming the time series of increments into the series of state numbers (1..s).

   2.3. Calculating transition probabilities for generalized states.

   2.4. Constructing the series of prognostic states using the procedure of defining the most probable next state.
2.5. Restoring the value series from the state series with $\Delta t$ sampling.

2.6. Splicing the prediction of $\Delta t$ sampling with the time series derived from splicing of the previous layers (with the lesser step $\Delta t$). In case if the current time series is the first one, the unchanged time series will come as a result of splicing.

3) To splice the last spliced time series with the continuation of the linear trend, created along all previously known points.

The time series, spliced with the linear trend, is the result of prediction. Let us consider the stages of the given algorithm in detail.

8. STATES IN COMPLEX MARKOV CHAINS AND APPROACHES FOR DEFINING THEM

In everything that concerns current technology, states are connected to the measuring of a prognostic value. There is a number of ways to classify returns in states, from which the following are suggested. One of them is the classification based on the homogeneity principle as concerning the amount of representatives in classes; based on the homogeneity principle of deviation, as well their combinations for different deviation modules.

Increment or returns of the time series serves as the basis for states classification [32, 33]. Absolute $r_a$ and relative $r_t$ increments of the time series are considered:

$$r_a = p_t - p_{t-\Delta t},$$ (17)

$$r_t = \frac{p_t - p_{t-\Delta t}}{p_t},$$ (18)

where $p_t$ – is the input time series of price dynamics, $\Delta t$ – sampling interval, which is chosen for subsequent analysis. It is known that mathematical expectation of the returns time series equals zero, whereas variation comes as the measure of time series volatility. Based on returns values $r_t$ classification and transformation of values to the time series of discrete states are conducted. One of the classification principles is homogeneity according to the amount of class representatives. This classification divides the set of all increments into $s$ groups equal in number. Calculated with the given sampling, time series increments are then systematized in growth and divided into equal parts. Thus we define limit values $\{r_{lim,i}\}$, which are used afterwards during transformation of the returns into class numbers. Large
number of identical states can cause certain problems, such as identical bounds of several
neighbouring states. It creates a number of states with no representatives, which makes
correction of the division a necessary action. In that way we will reach the largest possible
homogeneity in state division. Classification is conducted along the following algorithm
[34, 35]:

\[ s_t = \{ i \mid r_{lim,i-1} > r_t > r_{lim,i} \} \] (19)

where \( s_t \) is the number of state, which corresponds to the moment of time \( t \), for which the
returns level was computed \( r_t \); \( i \) is the number of state \([1 \ldots s]\), which is characterized by
the interval \([r_{lim,i-1}, r_{lim,i}]\) corresponding to the calculated returns level \( r_t \).

Apart from the returns interval, given by the aforementioned values \([r_{lim,i-1}, r_{lim,i}]\), a
mean returns value is chosen for every state \( r_{avg,i} \), which will be used in time series values
transformation according to the prognostic discrete states.

Another way of dividing the time series into states implies dividing the interval of returns
values into equal parts, from minimal to maximal deviation. In this case homogeneity
according to the amount of representatives in states does not occur. In fact this method
differs from the previous one in terms of defining limit values \( \{r_{lim,i}\} \). Possible combined
ways of division, in case of which the limit value, dependent on standard deviation, is used
instead of maximal and minimal value, and division is conducted homogenously according
to the deviation.

Since the real causal dependence is unknown during the process, to find it adequate
state classification, which would allow to reveal vital dependencies of the time series, is
required. We suggest a couple of ways to divide the time series into states, which in the first
place allow to preserve adequate transition probabilities between states, as well as prevent
averaged deviations inside the states from affecting the accuracy of the derived prediction.

To check the efficiency of division we conduct the sampling procedure and classify the
increments according to each hierarchy. Having completed that, we restore the time series
using known states for each hierarchy and finish the splicing procedure. Since the state series
correspond to the initial time series, we get the curve, with deviation, caused exclusively
by the state averaging mistake (quantum mistake). Thus, having set a certain value of
state numbers \( s \) and carried out sampling, restoring, and sampling procedures (excluding
prediction), we get absolute sampling (quantum) mistake.
Increasing the number of states, we improve the accuracy of restoration, however one should remember, that the choice of the quantum levels is limited by the fact that the transition probabilities definition with sufficient accuracy is required, which is confirmed by artificial test time series prediction experiments.

9. STEP-BY-STEP PREDICTION PROCEDURE. DEFINING THE MOST PROBABLE STATE ON THE NEXT STEP, PREDICTION SCENARIOS

Predicting procedure uses the most probable state as the next one under current circumstances. Probability matrix of state transitions is used for the afore-mentioned purpose. In this case, you have to take into account that probabilities are calculated with a certain mistake. We cannot precisely compute the probabilities, since it is impossible to derive an infinite time series, and only a part of the time series is known – the known part serves as the basis for probabilities. The second important aspect implies the case of several states with maximal probability.

To prevent the omission of the states, for which the probabilities are computed with a mistake, one should add a state with maximal probability to the states, which are located in the distance of $\delta$ from the maximal one. The value of parameter $\delta$ depends on the probability evaluation mistake and requires experimental refinement.

If $\delta > 0$, the number of states with maximal probability increases in comparison to the value $\delta = 0$. Let us call a couple of neighbouring states with maximal probability a cluster. Cluster states with average deviation values are supposed to have the largest probability. To predict the dynamics, let us confine ourselves to one or two most probable states. To define them a following algorithm is suggested:

1) If levels (discretized increments) create several clusters (cluster is a group of several neighbouring levels – cluster elements, minimal cluster is a single isolated level) with maximal probability, we choose the largest cluster.

2) If the number of cluster elements is odd, as $k_{max}$ we choose a central element.

3) If the number of cluster elements is even, we consider two central cluster elements and choose as $k_{max}$ the one, which is closer to the centre of distribution.

4) If two central cluster elements are equidistant to the centre of distribution, we consider both cases as possible variants of $k_{max}$ values (bifurcation point).
5) If there are several clusters of maximal size, we consider them as new elements, which can also form clusters that will undergo the same steps 1)-4).

This principle is based on the following ideas:

1) If there are two neighbouring states of maximal probability, it is better to take the one, which is closer to the centre of distribution, in order to minimize the risk of occurrence of false linear trends in the prediction.

2) If levels of maximal probability are not the neighbouring ones, at least two variants have to be considered, as it can be connected to the bifurcations that should not be omitted.

3) If the prediction is carried out according to 1) (on all stages of the hierarchy), we receive a certain approximation of the lower limit of the prediction, whereas in case of 2) – we get an approximation of the upper limit.

Hence this algorithm can adequately restore the case of possible bimodal probability distribution, it is proposed to consider 2 prediction scenarios.

In case of the complex Markov chains, probability of the next state depends not only on the previous state, but also on the sequence of \( r \) states, which have occurred before given. In this case, it is necessary to calculate transition probabilities from the sequence of \( r \) states into the \( r + 1 \) state. Formally, these probabilities can be written into the rectangular table of \((r^*, s)\) size.

Having generalized the notion of “present state” and included a sequence of \( r \) preceding states into it, we can reduce Markov chains of \( r \) order to the chain of the first order. Thus transition probabilities can be written into rectangular matrices of \((r^*, r^*)\), that come as transition probability matrices for generalized states.

The process of prediction implies the following: the last state is chosen (in case of Markov chains of an order \( r > 1 \) a sequence of \( r \) latest states is taken). The probability of transition from current state to all possible states is defined. From all possible states a state with maximal probability is chosen. It is possible that several states with maximal probability occur, which can be explained by the bimodal probability distribution. The process of decision-making in this case is described later.

The chosen most probable state is taken as the next prognostic state and the procedure is repeated for the next (last added) state. Thus we receive a time series of prognostic states for the given sampling time \( \Delta t \).

Further according to the received state sequence and known initial value the time series is
being restored for the given time sampling $\Delta t$. In this case every state implies $\Delta t$ points of the time series. On the stage of state classification every state was connected to the average increment $r_{\text{avg},i}$, which is added to the value of the last point in the time series, and the next discrete point is computed. Intermediate points are filled as linear interpolation of two known neighbouring points. Algorithm of $y_t$ time series values restoration according to the initial price $p_t$ and a series of average increments $r_{\text{avg},ik}$, corresponding to the prognostic states $s_k$, can be given by a sequence of calculations:

$$
y_t = p_t,$$
$$y_{t+1} = y_t + r_{\text{avg},i}/\Delta t = p_t + r_{\text{avg},i}/\Delta t,$$
$$y_{t+2} = y_{t+1} + r_{\text{avg},i}/\Delta t = p_t + 2r_{\text{avg},i}/\Delta t,$$
$$\ldots$$
$$y_{t+n\Delta t-1} = y_{t+n\Delta t-2} + r_{\text{avg},i}/\Delta t = p_t + (\Delta t - 1)r_{\text{avg},i}/t,$$
$$y_{t+n\Delta t} = y_{t+n\Delta t-1} + r_{\text{avg},i}/\Delta t = p_t + tr_{\text{avg},i}/\Delta t = p_t + r_{\text{avg},i},$$

$$\ldots$$

(20)

10. TIME INCREMENTS HIERARCHY AND SPLICING PROCEDURE

Time series increments will be computed with different steps. For example, analogous to the discrete Fourier transform, time increments are equal the powers of 2 are considered. First, we calculate increments as a remainder of two nearest neighbouring time series values, then next nearest values are considered with the step of 2, 4, 8, 16 etc. Let us mark this difference in time as $\Delta t$.

For every $\Delta t$ we conduct an increment time series transformation leading to a time series of states. Further we predict the future sequence of states and restore the time series with the given sampling rate according to the prognostic series of states.

Time series, received as a result of restoring for different $\Delta t$, undergo the splicing procedure, which gives out an actual prognostic time series.

Thus an increment hierarchy is chosen, where each one is responsible for its own sampling rate, which serves as a basis for predicting, restoring and splicing.
The splicing process implies the following. The procedure is iterative. With every next (along with the increasing step) sampling time the series corrects itself, driving the prediction, formed under lower $\Delta t$, to its actual point. Transformations that are conducted during splicing can be written down in the form of the following calculations.

Suppose the splicing procedure has been finished for all time increments $\Delta t < \Delta t_i$, the prediction has been done under the $\Delta t_i$ sampling according to formulae and as a result a time series $y_i$ has been derived. Let us consider the iterative splicing procedure of the received series $y_i$ with the series, acquired during all preceding splicing procedures $g_i$.

Since the series $y_i$ contains system points only in moments aliquot to $\Delta t_i$, and other points of the series are interpolated, the process of splicing implies the substitution of these interpolated points with the values of system points from previous $\Delta t < \Delta t_i$, which are contained in the series of results of previous splicing procedures $g_i$. Splicing algorithm can be written in the sequence of computations:

\[
\begin{align*}
  z_t &= g_t = p_t, \\
  z_{t+1} &= g_{t+1} + \frac{(y_{t+\Delta t_i} - g_{t+\Delta t_i})}{\Delta t_i}, \\
  z_{t+2} &= g_{t+2} + \frac{2(y_{t+\Delta t_i} - g_{t+\Delta t_i})}{\Delta t_i}, \\
  \vdots \\
  z_{t+\Delta t - 1} &= g_{t+\Delta t - 1} + \frac{(\Delta t - 1)(y_{t+\Delta t_i} - g_{t+\Delta t_i})}{\Delta t_i}, \\
  z_{t+\Delta t} &= g_{t+\Delta t} + \frac{(\Delta t)(y_{t+\Delta t_i} - g_{t+\Delta t_i})}{\Delta t_i} = y_{t+\Delta t_i}, \\
  z_{t+\Delta t+1} &= g_{t+\Delta t+1} + \frac{((y_{t+2\Delta t_i} - g_{t+2\Delta t_i}) - (y_{t+\Delta t_i} - g_{t+\Delta t_i}))}{\Delta t_i}, \\
  z_{t+\Delta t+2} &= g_{t+\Delta t+2} + \frac{2((y_{t+2\Delta t_i} - g_{t+2\Delta t_i}) - (y_{t+\Delta t_i} - g_{t+\Delta t_i}))}{\Delta t_i}, \\
  \vdots \\
  z_{t+n\Delta t-1} &= g_{t+n\Delta t-1} + \frac{(\Delta t-1)}{\Delta t} \left( (y_{t-n\Delta t} - g_{t-n\Delta t}) - (y_{t-(n-1)\Delta t} - g_{t-(n-1)\Delta t}) \right), \\
  z_{t+n\Delta t} &= g_{t+n\Delta t} + \frac{((y_{t-n\Delta t} - g_{t+n\Delta t}) - (y_{t-(n-1)\Delta t} - g_{t+(n-1)\Delta t}))}{\Delta t_i} = g_{t+(n-1)\Delta t} - y_{t+(n-1)\Delta t} - y_{t-n\Delta t}. 
\end{align*}
\]
11. RESULTS OF STOCK INDICES PREDICTION

In this section we offer the results of stock indices prediction. The stock's indices databases are available from [36]. Point 2000 indicates the starting moment of the prognosis: March 24, 2011. The green line on the next figures indicates real indice’s or price’s values. Our software for time series forecasting by the proposed methods is available from our website: [http://kafek.at.ua/MarkovChains1_2_20100505.rar](http://kafek.at.ua/MarkovChains1_2_20100505.rar).

![Dow Jones Industrial Average - DJI (USA) and FTSE 100 (Great Britain)](image1)

Figure 1: Stock indices prediction. a) Dow Jones Industrial Average - DJI (USA). b) FTSE 100 (Great Britain)

![Morgan Stanley (USA) and BNP Paribas (France)](image2)

Figure 2: Financial company’s share prices forecasting. a) Morgan Stanley (USA). b) BNP Paribas (France)
Prediction time series with different input learning set’s length are shown at the fig. 3 and 4. Prediction series for DJI at the figure 3 are more correlated, than FTSE index at the figure 4. At the subplot b) of the above mentioned plots the mean value and standard deviations of the prediction’s series are presented. The time of prediction series beginning on the next figures is the point 1000 and correspond to October 14, 2011.

Figure 3: Dow Jones Industrial Average - DJI (USA). a) Prediction series, calculated with different learning set’s length. b) Mean value and standard deviation for prediction series.

Figure 4: FTSE 100 index prediction. a) Prediction series, calculated with different learning set’s length. b) Mean value and standard deviation for prediction series.
The normalization procedure is proposed in order to compare indices and its prediction series with different absolute values. The normalized values are calculated with the following formula:

$$y_n(t) = \frac{y(t) - \min (y(t))}{\max (y(t)) - \min (y(t))}.$$  \hspace{1cm} (22)

Normalized prediction time series are shown in the fig.5 (America), fig.6 (Europe, developed countries), fig.7 (Europe, PIIGS), fig.8 (Asian markets). All the figures contain mean time series, which are weighted average of countries' stock indices predictions, weighted with GDP values [37] for the corresponding countries.

Figure 5: Normalized mean values for the prediction series of America’s stock indices. Brazil (BVSP), Mexico (MXX), Canada (GSPTSE), Argentina (MERV), USA (S&P 500)
Figure 6: Normalized mean values for the prediction series of European stock indices. Developed countries: FTSE (Great Britain), DAX (Germany) FCHI (France), Netherlands (AEX).

Figure 7: Normalized mean values for the prediction series of European stock indices. Portugal (PSI20), Italy (FTSEMIB), Ireland (ISEQ), Greece (GD) and Spain (IBEX).
Figure 8: Normalized mean values for the prediction series of Asian stock indices. China (SSEC, HSI), Korea (KS11), Japan (NIKKEI), India (BSESN), New Zealand (NZ50).

Figure 9: Mean values of normalized World’s powerful economies indices prediction series.
12. CONCLUSIONS AND FURTHER WORK

Current paper suggests an algorithm of time series prediction based on complex Markov chains. Hierarchy of time increments principle allows to use the information, which is contained in the time series during the prognosis construction, to its fullest. Experimental work on stock market indices time series prediction shows the efficiency of the algorithm and confirms the relevance of further research of the offered method.

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