Age-Dependent Brain Tissue Hydration, Ca Exchange and their Dose-Dependent Ouabain Sensitivity

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Abstract

Tissue hydration, dose-dependent H-ouabain binding, 45Ca\textsuperscript{2+} exchange in rat’s brain cortex, subcortex and cerebellum were studied in three age groups. Age-dependent tissue dehydration in all three zones of brain was due to inhibition of Na\textsuperscript{+}/K\textsuperscript{+} pump. The age-dependence of cell hydration in cortex was more expressed. The curve of dose-dependent ouabain binding consists of three components corresponding to Na\textsuperscript{+}/K\textsuperscript{+} pump isoforms (\(\alpha_1\), \(\alpha_2\), \(\alpha_3\)). Age-dependency of these isoforms was more expressed in cortex than in subcortex and cerebellum. High affinity receptors were depressed in old rats’ brain tissues. Initial 45Ca\textsuperscript{2+} uptake in three brain zones of old rats was depressed as compared to that of young animals. Ouabain at 10\textsuperscript{–9}M has activation effect on 45Ca\textsuperscript{2+} uptake, which was also age dependent. Initial 45Ca\textsuperscript{2+} efflux in cortex and subcortex tissue in old rats was significantly depressed as compared to young ones while in cerebellum the opposite age-dependence was observed. The curves of dose-dependent ouabain effect on 45Ca\textsuperscript{2+} efflux and cell hydration consist of 6 components. However, close correlation between kinetics of 45Ca\textsuperscript{2+} efflux and cell hydration was not observed. It is suggested that brain tissue dehydration in aged animals is a consequence of Na\textsuperscript{+}/K\textsuperscript{+} pump dysfunction induced intracellular calcium elevation. It is suggested that \(\alpha_3\) receptors are functionally connected with intracellular Ca\textsuperscript{2+} buffering systems through intracellular signaling systems and their dysfunction in aged brain is a consequence of [Ca\textsuperscript{2+}] increase. Obtained data allow us to conclude that endogenous nanomolar ouabain-like species circulating in mammals’ blood removing Ca\textsuperscript{2+} from cells could have a beneficial effect on brain of old animals.

Introduction

Although the important physiological role of water in cell functional activity is well known and widely accepted, its messenger role in the generation of various diseases including aging-induced memory loss and the increase of nerve disorders risk has not been paid an adequate attention by researchers. It is known that aging-induced memory loss and the increase of the risk of nerve disorders are accompanied by tissue dehydration. However, whether the age-induced cell dehydration is the primary risk factor for age-related nerve disorders is not clear. The role of cell hydration in the memory processes, as well as the role of cell volume controlling metabolic mechanisms in neuronal dysfunctions has not been investigated in detail.

As the intracellular osmotic pressure in normal cells exceeds the extracellular one, then dysfunction of metabolic energy generating pathways or membrane selective permeability disturbance at first could bring to cell swelling. As the activity of intracellular proteins and gene expressing systems strongly depends on hydration, the latter could be considered as a critical factor for the regulation of intracellular metabolic activity. Our early works showed that membrane proteins having enzymatic, receptor and channel-forming properties can be in functionally active and inactive (reserve) states, the ratio of which is changed depending on cell hydration [1]. Cell swelling (membrane unpacking), leads to the increase of functionally active membrane receptors’ [2,3], membrane enzyme molecules’ [4] and ionic channels’ number in membrane [5], while its dehydration (membrane packing) has an opposite effect. On the basis of these data cell swelling can be considered as primary messenger switching on intracellular signaling system through which the protective reaction of cells is realized.

Among the number of mechanisms involved in cell volume regulation Na\textsuperscript{+}/K\textsuperscript{+} pump has fundamental role, because Na\textsuperscript{+} gradient serves as an energy source for a number of secondary ionic transporters, such as Na\textsuperscript{+}/Ca\textsuperscript{2+}, Na\textsuperscript{+}/H\textsuperscript{+}, sugars, amino acids and osmoletes [6]. It is known that there are two enzyme systems actively involved in metabolic regulation of cell volume associated with cation transport across membrane surface. In this process transport ATPases are indeed the translocating structure and are fueled by the free energy derived from ATP hydrolysis while kinases may regulate translocation via phosphorylation of the transporter molecules through the phosphorylation of associated regulatory structures. The close talking between these two enzyme systems is realized through the intracellular signaling systems, the dysfunction of which leads to generation of cell pathology. The latter is accompanied by corresponding changes of cell hydration. As Na\textsuperscript{+}/K\textsuperscript{+} pump is the most ATP-utilizing machine in cell, it serves as a main regulator of all other ionic pumps’ and kinases’ activity. Therefore factors able to change the balance between ATP hydrolysis and ATP production system (in mitochondria) by changing Na\textsuperscript{+}/K\textsuperscript{+} pump activity could switch on the intracellular signaling systems-induced modulation of cell katabolic and anabolic process. Therefore, the dysfunction of Na\textsuperscript{+}/K\textsuperscript{+} pump controlling cell hydration can be considered as a common gate for cell pathology. However, the nature of mechanism through which Na\textsuperscript{+}/K\textsuperscript{+} pump dysfunction leads to apoptosis and proliferation inhibition in excitable cells stays unclear.

The second ionic transporting mechanism in cell membrane having a crucial role in cell volume regulation is Na\textsuperscript{+}/Ca\textsuperscript{2+} exchange [7-10]. It is known that there is a close correlation between electrogenic Na\textsuperscript{+}/K\textsuperscript{+} pump and cell volume preservation. This mechanism is indeed the translocating structure and is fueled by the free energy derived from ATP hydrolysis while kinases may regulate translocation via phosphorylation of the transporter molecules through the phosphorylation of associated regulatory structures. The close talking between these two enzyme systems is realized through the intracellular signaling systems, the dysfunction of which leads to generation of cell pathology. The latter is accompanied by corresponding changes of cell hydration. As Na\textsuperscript{+}/K\textsuperscript{+} pump is the most ATP-utilizing machine in cell, it serves as a main regulator of all other ionic pumps’ and kinases’ activity. Therefore factors able to change the balance between ATP hydrolysis and ATP production system (in mitochondria) by changing Na\textsuperscript{+}/K\textsuperscript{+} pump activity could switch on the intracellular signaling systems-induced modulation of cell katabolic and anabolic process. Therefore, the dysfunction of Na\textsuperscript{+}/K\textsuperscript{+} pump controlling cell hydration can be considered as a common gate for cell pathology. However, the nature of mechanism through which Na\textsuperscript{+}/K\textsuperscript{+} pump dysfunction leads to apoptosis and proliferation inhibition in excitable cells stays unclear.

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pump and electrogenic Na’/Ca²⁺ exchange which have been described since pioneering work by Baker et al. [11]. At present a close correlation between these two ion transporting mechanisms on the level of different Na’/K’ pump isoforms is known, thanks to great contribution of one of co-authors of this work doctor Blaustein’s group, who identified and characterized different isoforms of Na’/K’ pump [12,13].

As Na’/K’ pump has a crucial role in cell adaptation process, aging can be considered as a consequence of Na’/K’ pump dysfunction [14,10], which leads to the down regulation Ca²⁺ homeostasis [15-18]. Small enhancement of intracellular Ca²⁺ concentration leads to the inactivation of Na’/K’/Mg²⁺-ATPase [19]. It is known that Na’/ K’-ATPase has three isoforms of catalytic (α) subunit having different affinities to cardiac glycoside ouabain: α₁ (low affinity), α₂ (middle affinity) and α₃ (high affinity). They also have a different localization in cells: α₁ is ubiquitously distributed over the surfaces of cells, while high ouabain affinity isoforms are confined to a reticular distribution within the cellular membrane that paralleled underlying endoplasmic or sarcoplasmic reticulum, with Na’/Ca²⁺ exchanger protein. The expression of these isoforms is age-dependent [20-25]. While the role of low-affinity receptors as a working molecule for Na’/K’ pump is well established [25-27], the functional role of high-affinity receptors is still discussable [13]. Detailed investigations [28-30] showed that Na’/Ca²⁺ exchange is regulated through the high-affinity ouabain receptors, while Na’/K’ pump by means of low-affinity ones. The detailed mechanism of correlation between functions of α₁ and α₂ pump subunit and Na’/Ca²⁺ exchange and their functional significance in norm and pathology are not clear.

Therefore, considering all the facts mentioned, the study of age-dependent ouabain sensitivity of cell hydration and Na’/Ca²⁺ exchange could bring us close in understanding the role of different isoforms of Na’/K’ pump in the regulation of cell hydration. For this purpose the effect of rats’ aging (6, 18 weeks and 18 months) on the cell hydration and its ouabain sensitivity, dose-dependent ouabain binding with cell membrane and ⁴⁰Ca²⁺ uptake and efflux in brain cortex, subcortex and cerebellum were studied.

Materials and Methods

All procedures performed on animals were carried out following the protocols approved by Animal Care and Use Committee of Life Sciences International Postgraduate Educational Centre (LSIPEC, Yerevan, Armenia).

Animals

Experiments were performed on three separate groups of male albino rats: young (6 weeks old), adult (18 weeks old) and old (18 months old) weighing 35-40 g, 100-120 g and 250-300 g, respectively. Animals were regularly examined, kept under control of the veterinary in LSIPEC and reserved in a specific pathogen free animal room under optimum conditions of 12 h light/dark cycle, at temperature of 22 ± 2°C, a relative humidity of 50 % and were fed ad libitum on a standard lab chow and water.

For definition of initial water content five animals from each group were chosen and their data were considered as control after intraperitonally injected physiological solution (PS). In subsequent experimental parts of investigation the same number of animals was used. The brain tissue hydration was provoked by simple application of interperitoneal injection of 3 ml DW, isotonic (0.9% NaCl) and hypertonic (2 M Mannitol containing) PS. Study of ⁴⁰Ca²⁺ uptake and the rate constant of ⁴⁰Ca efflux from brain tissues was performed on young and old animals (n=5). All data was received from three independent experiments.

Chemicals

Tyrode’s physiological solution (PS) containing (in mM) 137 NaCl, 5.4 KCl, 1.05 MgCl₂, 5 C₆H₁₂O₆, 11.9 NaHCO₃, 0.42 NaH₂PO₄ and adjusted to pH 7.4 with NaOH was used. For increasing the PS osmolality the non-metabolizing substance- Mannitol [C₆H₁₂O₆ mol. wt. 182.18], 2 M was dissolved in the saline (0.9 % NaCl). All chemicals were obtained from “Medisan” Industrial Chemical Importation Company (Yerevan, Armenia). K-free solution was consisted of 5.4 mM NaCl instead of 5.4 mM KCl. ⁴⁰Ca²⁺ with specific activity: 40mCi/ml was added in K-free solution containing 68.5 mM (50%) NaCl and used for tissues’ samples enriching. Radioactive ⁴⁰Ca-ouabain with specific activity 25.34 Ci/mM and non radioactive one (PerkinElmer, Massachusetts, USA) from 10⁻¹¹ M to 10⁻⁴ M concentrations dissolved in physiological solution was used for intraperitoneal injections and tissues’ incubation. The volume of injected solution was adjusted according to the weight of animals.

Tissue preparation

It is well known that anesthetics with different chemical and pharmacological profiles [31-33] significantly affect metabolic processes which play an important role in regulation of cell volume [12,34]. Therefore, in present experiments the animals were sharply immobilized by freezing method (dipping their noses into liquid nitrogen for 3-4 sec) and decapitated [35]. After such procedure the full absence of somatic reflexes on extra stimuli was recorded. Investigated tissues were isolated and dissected into the samples weighing from 50 to 70 mg. Both for cortex, subcortex and cerebellum tissues 25 samples were chosen for each experimental group.

Definition of water content of brain tissues

Water content of brain tissues was performed by traditional “tissue drying” method. After measuring the tissue wet weight (w.w.) it was dried in thermostat (Factory of Medical Equipment, Odessa, Ukraine) during 24 h at 105°C for determination of dry weight (d.w.). The quantity of water in 1 g of d.w. of tissue was counted by the following equation: (w.w. – d.w.)/d.w. ± 0.1 g.

Counting of ⁴⁰Ca-ouabain receptors in membrane

²⁰⁶H-ouabain solution is usually used to estimate the number of membrane ouabain receptors. It is assumed that each binding site (receptor) in membrane binds one ouabain molecule. Dried tissues’ samples were homogenized in 50 μl of 68% HNO₃ solution, then 2 ml of Bray’s scintillation fluid was added and chemoluminescence of samples was quantified with 1450 MicroBeta liquid scintillation counter (Wallac Oy, Turku, Finland). Number of ouabain receptors was defined per mg of dry weight.

Measurement of ⁶⁰⁰Ca²⁺ uptake and ⁶⁰⁰Ca²⁺ efflux rate constant

Measuring of ⁶⁰⁰Ca²⁺ uptake was made in vivo conditions. For this purpose in Tyrode’s physiological solution 0.0115 mM CaCl₂ from 1.8 mM was substituted by ⁶⁰⁰Ca²⁺ the radioactivity of which was 11.2 mCi/l. Then young and old animals were intraperitoneally injected by 0.187 mCi/g of body weight. After 30 min they were decapitated and brain samples were incubated for 30 min in PS (as a control) and 10⁻⁴ M ⁶⁰⁰Ca²⁺ ouabain solutions. After that all samples were dried in thermostat
during 24 h at 105°C. After definition of samples' dry weight they were homogenized in 50 µl of 68% HNO₃ solution, then 2 ml of Bray’s scintillation fluid was added and chemiluminescence’s of samples was quantified with 1450 MicroBeta liquid scintillation counter (Wallac Oy, Turku, Finland). “Ca²⁺” content was defined per mg of dry mass.

For measurement of “Ca²⁺” efflux all samples of brain tissues were incubated during 60 min in K-free (containing 50% NaCl) solution where “Ca²⁺” was added (10.77 µl “Ca²⁺” for 100ml K-free solution) for tissue enriching by “Ca²⁺”. Then “Ca²⁺”-enriched samples were washed three times (in K-free solution) for 10.5 and 5 min, respectively, for removing the extracellular “Ca²⁺”. Brain tissues samples were divided on two parts. First 25 samples were placed in thermostat during 24 h at 105°C. Second 25 samples were incubated in ouabain solution during 30 min. Then all samples were dried in thermostat during 24 h at 105°C. After definition of samples dry weight they were homogenized in 50 µl of 68% HNO₃ solution, then 2 ml of Bray’s scintillation fluid was added and chemiluminescence’s of samples was quantified with 1450 MicroBeta liquid scintillation counter (Wallac Oy, Turku, Finland). “Ca²⁺” content was defined per mg of dry mass and the rate constant of “Ca²⁺” efflux was calculated as an exit part of absorbed “Ca²⁺” for control and experimental data.

Statistick analysis

The Microsoft Excel and Sigma-Plot (Version 8.02A, NY, USA) were used for data analysis. Significance in comparison with the control group was calculated with Student’s paired t-test with the following symbols (*P < 0.05; **P < 0.01; ***P < 0.001).

Results

Before the main study of the ouabain effect on brain tissue hydration four independent experiments were performed to determine the dynamics of hydration changes in brain tissue in three age groups. The results showed variability between the mean values of initial hydration in each age group. However, the dynamics of change for the mean hydration value was the same for each age group in all experiments. The overall mean hydration value in all three age groups is very close to the data presented for the fourth group of experiments. On Figure 1 the initial water content in tissue of three brain zones in three age groups is presented. As can be seen the initial water content in tissues of young animals was higher than in adult and old ones. It worth to note that initial water content in cortex tissue was higher than in subcortex and cerebellum tissues in all age groups.

To estimate the age-dependent dynamics of Na⁺/K⁺ pump-dependent cell hydration changes in three zones of brain tissue in all age groups under the effect of 10⁻⁴ M ouabain concentration were studied. For these experiments new animal groups (young, adult and aged) were taken. In each group five animals were decapitated after 30 min of 10⁻⁴ M ouabain intraperitoneal injection and water content was determined in brain zones of all investigated animals. Data presented on Figure 2 demonstrate that ouabain at concentration of 10⁻⁴ M has total hydration effect on all brain zones of three age groups (Figure 2A-2C) as compared to sham data. However, 10⁻⁴ M ouabain induced tissue hydration was less pronounced in young animals than in adult and old animals. These data are in harmony with literature data that in early postnatal period of development the Na⁺/K⁺ pump expression is rather weak and it increases during periods of maturation [12]. Therefore, a more pronounced ouabain-induced hydration of brain tissues in adult and old animals as compared to young ones can be explained by more expressed Na⁺/K⁺ pump in maturated animals [36,37]. Observed

Figure 1: Age dependent initial level of water content in young, adult and aged rats’ brain tissues (cortex, subcortex and cerebellum). Ordinate indicates the mean value of tissues’ water content. As a control the data of young group animals was taken. Error bars indicate the standard error of the mean (SEM) for three independent experiments. The symbols (*) and (**) indicate P<0.01 and P<0.001, respectively.

Figure 2: Age dependent change of brain tissues water content (A, B, C) in young, adult and aged animals after 30 min ouabain 10⁻⁴ M intraperitoneal injection. Ordinates indicate the mean value of tissues’ water content. On figure right side (D, E, F) delta (Δ) determines the difference between water content received from sham and ouabain poisoned data. As a sham data the mean values showed on Figure 2 was taken. Error bars indicate the standard error of the mean (SEM) for three independent experiments. The symbols (*) and (**) indicate P< 0.05, P<0.01 and P<0.001, respectively. A difference which is not statistically significant means that P>0.05.
differences of brain tissues hydration in three groups of animals can be more clearly seen on right colon of Figure 2D-2F where the differences (Δ) between water content of sham and ouabain-poisoned animals are presented. It is worth to note that in adult animals cortex tissue hydration is higher than that of subcortex and cerebellum.

In the next series of experiments the correlation between dose-dependent 'H-ouabain binding and brain tissue hydration in three ages' animal groups were studied, which would let us to clarify the role of different isoforms of catalytic subunit in determination of observed age-dependent changes of tissue hydration.

As shown in Figure 3A-3C on dose-dependent 'H-ouabain binding curves of all investigated brain tissues in young, adult and old animals three types of ouabain receptors having different kinetics of binding (10^{-10}-10^{-4} M, 10^{-8}-10^{-4} M and 10^{-6}-10^{-4} M) can be distinguished. These data are confirmed by literature data on the existence of three types (α1, α2, and α3) of Na+/K-ATPase catalytic subunit in excitable cell membrane [11]. As shown on the Figure 3, the kinetics of investigated brain tissues is similar in young (Figure 3A) and old (Figure 3C) age groups. However, in adult group the kinetics of ouabain binding of cortex tissue where higher hydration was observed than in subcortex and cerebellum (Figure 2) the number of ouabain molecules is much higher in cortex tissue, than in subcortex and cerebellum (Figure 3B). It should be noted that in cortex tissue of adult animals 'H-ouabain binding kinetics is distinctly higher expressed in range of α3 isofrom (10^{-10}-10^{-4} M). The reversed dose-dependent decrease of ouabain binding in all three zones of brain tissues in young and less pronounced in tissues of subcortex and cerebellum of adults seems strange and difficult for interpretation. In old rats the dose dependency of ouabain binding is not visible in all three zones of brain tissues (Figure 3C), which indicates on their dysfunction.

It is known that ouabain binding depends both on the number and the affinity of receptors. Previous experiments performed on snail nerve ganglia showed that the number of ouabain receptors increases with cell swelling and decreases with cell shrinkage [4]. To find out the existence of such correlation between cell hydration and number of ouabain receptors in cell membrane of rats' brain tissue a series of experiments on the influence of different osmolarity solutions was performed on adult rats. As can be seen on Table 1, the intraperitontial injection of distilled water leads to the increase of brain tissue hydration in all three investigated zones of brain; while the same volume of hypertonic solution (mannitol) injection leads to cell hydration. A close correlation between cell hydration and number of 'H-ouabain binding molecules was observed. Distilled water-induced cell hydration was accompanied by increase, while hypertonic solution-induced cell dehydration-by decrease of the number of ouabain molecules binding with membrane. It is interesting to note that cortex tissue hydration and the number of ouabain receptors are more sensitive to both hypo- and hypertonic solution-induced stress than that of subcortex and cerebellum. From these data can be concluded that cell dehydration induced decrease of ouabain receptors in membrane can serve as one of mechanisms determining the weakening of Na+/K+ pump activity (Figure 2) in aged animals (Figure 1).

In order to find out the individual role of observed three populations of ouabain receptors with different affinity (α1, α2, and α3) in determination of age-dependent cell dehydration the dose-dependent effect of ouabain on cell hydration in brain tissues of young and old animals was studied.

This study has shown that even extremely low ouabain concentrations (10^{-4}-10^{-3} M) have a strong modulation effect on cell hydration in all three age animal groups: the activation of high affinity ouabain receptors causes dehydration of brain tissue in young and older rats, hydration - in adults. The dose-dependent ouabain effect on tissue hydration in young and old animals is clearly seen on Figure 4A-4C, on which the dependence of ouabain induced effect is compared with control one. As can be seen on this figure the curve of the dose-dependent ouabain effect on brain tissue hydration in range of 10^{-8}-10^{-4} consists of 6 components, namely 10^{-8}-10^{-6}, 10^{-6}-10^{-4}, 10^{-4}-10^{-2}, 10^{-2}-10^{-1}, 10^{-1}-10^{-0}, whereas the dose-dependent ouabain binding curve consists of 3 components. It is interesting to note that in old animals these components are more pronounced than in young ones. From these data can be concluded that there may be minimum 6 mechanisms involved in cell volume regulation having different ouabain sensitivity. Such heterogeneity within the family of Na+/K+ α1, α2, α3 isoforms indicates that ouabain receptors having the same affinity are functionally connected with different metabolic mechanisms involved in cell volume regulation.

As brain aging leads to the increase of intracellular Ca ions concentration [38] which has a crucial role in regulation of membrane receptors affinity to agonists, the study of the correlation between cell hydration and Ca\(^{2+}\) uptake and efflux as well as their ouabain sensitivity in young and old rats was the subject for the next series of experiments.

In Figure 5A-5C data on Ca\(^{2+}\) uptake by brain tissue in normal and 10\(^{-4}\) M ouabain poisoned young and old rats are presented. As can be seen from these data, in all investigated tissues of young animals the intensity of Ca\(^{2+}\) uptake is higher than that of old ones. In 10\(^{-4}\) M ouabain poisoned young and old animals Ca\(^{2+}\) uptake is higher as compared to control ones. The 10\(^{-4}\) M ouabain concentration influence is much higher than that of 10\(^{-4}\) M in cortex and cerebellum.
in both ages' animals (Figure 5A and 5C) while in subcortex of young rats it was depressed (Figure 5B). The differences of $^{45}\text{Ca}^{2+}$ uptake age-dependence in different zones of normal and poisoned animals' brain tissues can be seen more clearly on Figure 5D-5F.

It is known that $^{45}\text{Ca}^{2+}$ uptake could take place by two pathways: by potential activated $^{45}\text{Ca}^{2+}$ channels and by Na'/Ca$^{2+}$ exchange in reverse mode [30]. As low concentration ouabain is unable to inactivate Na'/K$^+$ pump and depolarize the membrane, the potential-dependent $^{45}\text{Ca}^{2+}$ channels induced uptake of ions is excluded. Therefore Na'/Ca$^{2+}$ exchange in reverse mode can be the only pathway through which low ouabain could stimulate $^{45}\text{Ca}^{2+}$ uptake. These data are in harmony with our previous data performed on snail neuronal ganglia on the uptake activation effect of low concentration ouabain ($<10^{-4}$ M) on $^{22}\text{Na}^{+}$ efflux and $^{45}\text{Ca}^{2+}$ uptake [4,39].

As energy source for driving Na'/Ca$^{2+}$ exchange in reverse mode is electrochemical potential for Ca$^{2+}$ ($E_{Ca}$) which exceeds the electrochemical potential for Na$^+$ ($E_{Na}$), low ouabain-induced activation of $^{45}\text{Ca}^{2+}$ uptake can be explained by increasing $E_{Ca}$ because of Ca$^{2+}$ adsorption by intracellular structure or by activation Ca$^{2+}$ pump.

The study of dose-dependent effect of ouabain on $^{45}\text{Ca}^{2+}$ efflux from young and older rats' brain tissues indicates that the initial rate constant of $^{45}\text{Ca}^{2+}$ efflux from cortex and subcortex tissues in young animals is much higher than in old ones, while in cerebellum tissue it is higher in old animals (Figure 6A). Figure 6B shows the difference between values of $^{45}\text{Ca}^{2+}$ efflux from the investigated brain zones in old and young animals. It is interesting to note that in cerebellum age dependence of $^{45}\text{Ca}^{2+}$ efflux has reverse character in contrast to cortex and subcortex tissues (Figure 6B). As in case of dose-dependent ouabain effect on brain tissue hydration (Figure 4) the curve of dose-dependent ouabain effect on $^{45}\text{Ca}^{2+}$ efflux also consists of 6 components (Figure 7). However, there is no direct correlation between kinetics of dose-dependent ouabain effects on $^{45}\text{Ca}^{2+}$ efflux and cell hydration. It is interesting to note that in range $10^{-11}$-$10^{-9}$ M ouabain i.e. in zones of high affinity receptors ($a_i$ and $a_o$) of $^{45}\text{Ca}^{2+}$ uptake.
the dose-dependent dynamics of $^{45}\text{Ca}^2+$ efflux in young and older rats has opposite character while at higher concentrations its dynamics has similar character (Figure 7A).

In subcortex tissue (Figure 7B) the dynamics of dose dependent ouabain effect on $^{45}\text{Ca}^2+$ efflux in young and old animals has similar character, but the rate constant of $^{45}\text{Ca}^2+$ efflux was much higher in older animals than in young ones. In contrast to cortex and subcortex tissues the dynamics of $^{45}\text{Ca}^2+$ efflux from cerebellum tissue has highly variable character (Figure 7C). Ouabain at all concentrations has inhibitory effect on $^{45}\text{Ca}^2+$ efflux in old animals while in young animals this effect is observed only at $10^{-11}$; $10^{-8}$ and $10^{-4}$ M. Such reverse age dependency of $^{45}\text{Ca}^2+$ efflux in cerebellum tissue of normal (Figure 6) and ouabain poisoned animals (Figure 7) seems very interesting and could serve as a special subject for a more detailed investigation.

Discussion
At present it is a proven fact that metabolic control of cell volume is a dynamic parameter regulating cell’s various functions (1,6,40-42). It is widely recognized that although neural cell swelling can be observed under physiological conditions, for example as a result of neurotransmission or intense neuronal discharge (4,43-45), larger changes are encountered during pathological conditions, including aging. It is known also that neuronal dehydration in older adults is a reliable predictor of increasing frailty, progressive deterioration in cognitive function and an overall reduction in quality of life [46-49].

There are a number of hypotheses on regard of detailed role of age-dependent cell dehydration in deterioration in cognitive function of brain. However, because of our weak knowledge on the detailed metabolic cell volume controlling mechanism(s) dysfunction of which brings to neuronal dehydration in aging animal is the main barrier in understanding the real functional role of neuronal dehydration in dysfunction of brain cognitive function in aging. As Na$^+$/K$^+$ pump dysfunction can be considered as one of the reasons for aging in present work we have made an attempt to find out the individual role of different pump isoforms having different affinity to ouabain (α$\text{1,2,3}$) [12] in generation of age-dependent dehydration in rats’ brain tissues.

Data obtained in present work indicate that water content of brain cortex tissues is higher than that of subcortex and cerebellum in all age groups (Figure 1). Although initial water content of investigated tissues of young rats is higher than in adult ones (Figure 1A-1C), $10^{-6}$ M ouabain poisoning-induced cell hydration which is due to Na$^+$/K$^+$ pump inhibition is significantly less than in maturated animals (Figure 2D-2F). The data that age-dependent changes of ouabain-induced brain tissues’ hydration have a bell-shaped dynamics (young<adults>aged) are in harmony with literature data on the reciprocal relationship between the expression of Na$^+$/K$^+$ pump and Na$^+$/Ca$^{2+}$ exchanger proteins in neuronal development. In prenatal and early postnatal periods the expression proteins of Na$^+$/Ca$^{2+}$ exchanger prevails, while by maturating the expression Na$^+$/K$^+$ pump protein increases, reaching its maximum in adults and then intervenes its dysfunction in aging [12]. From this point of view the high water content in brain tissues of young animals can be explained by higher activity of electrogenic Na$^+$/Ca$^{2+}$ exchange, working in stochiometry of 3 Na$^+$/1Ca$^{2+}$ [10] while the brain tissue dehydration in adult animals can be explained by activation of electrogenic Na$^+$/K$^+$ pump [36,37]. Therefore, it is predictable that $10^{-6}$ M ouabain induced inhibition of Na$^+$/K$^+$ pump leads to lower hydration in brain tissues of young than maturated animals (Figure 2).
It is known that brain aging causes increase of [Ca++] ions [38] leading to reverse functioning of Na+/Ca++ exchange [10], having a dehydration effect on cells [7]. The reverse functioning of Na+/Ca++ exchange from one side could have cell dehydration and from the other side cell hydration effect through [Ca++], induced Na+/K+ pump inhibition [21]. Therefore, in aging animals Na+/Ca++ exchange would have a prevailing role in the regulation of neuronal volume because of dysfunction of Na+/K+ pump.

Obtained data on age-dependent kinetics of dose-dependent ouabain binding indicate that in young animals there is a strict difference between kinetics of ouabain receptors having different affinity to ouabain, which correspond to three Na+/K+ pump isoforms (α1,α2,α3) described in literature [11]. It is worth to note that data on age-dependent changes of ouabain molecules' number could not be considered as marker for changes of expressed quantity of ouabain receptors in membrane, because of age-dependent membrane lipid composition changes, having strong modulation effect on membrane receptors affinity to agonist. However the dose-dependent kinetics of ouabain binding with α1,α2,α3 receptors in brain tissue of three animal ages, could give information on age-dependent changes of their functional activity.

It seems extremely interesting that the process of maturation leads to significant changes in ouabain binding in brain cortex while the changes in subcortex and cerebellum of adult animals was negligible (Figure 3B), as in case of hydration (Figure 2D). It can be explained by aging-induced increase of brain cognition function in maturation, in which cortex has a crucial role. The disappearance of dose-dependent ouabain binding with α1 receptors (10^-11-10^-8 M) in the investigated tissues of aged animals could be interpreted as a marker for the dysfunction of these receptors. Such dysfunction can be explained by the depression of their affinity in result of age-dependent increase of intracellular Ca ions [38], having inhibitory effect on Na+/K+ ATPase activity [21]. These data correspond to literature data that α1 receptors have a higher affinity to intracellular Ca ions than α2 and α3 receptors [12].

Data obtained on the close correlation between cell hydration and number of ouabain receptors in membrane (Table 1) allow us to suggest that aging alongside with decreasing of ouabain receptors affinity could bring also to decrease of ouabain receptors number as a result of brain tissue dehydration. The fact that aging-induced dehydration could have a more pronounced depressing effect on α1 receptors than on others indicates their deeper localization in membrane as compared to α2 and α3 [21,42]. The fact that osmo sensitivity of α1 receptors in cortex is higher than in subcortex and cerebellum allows us to speculate on the possible role of these receptors in determination of single neuronal memory.

Because of high sensitivity of cell hydration to ouabain in present work was possible to indicate that the family of α1,α2,α3 ouabain receptors is not homogenic, i.e. there can be distinguished minimum 6 components, namely sensitive to 10^-11-10^-6 M, 10^-6-10^-4 M, 10^-4-10^-2 M, 10^-2-10^-1 M, 10^-1-10^0 M (Figure 4). These data are in harmony with literature data on the existence of caveolae in cell membrane having different metabolic functions [50].

The fact that direct correlation between cell hydration and ouabain binding was absent in the described experiments (Figure 4) indicates that the age-dependent depression of ouabain binding cannot be a result of cell dehydration-induced decrease of ouabain receptors' number in membrane only, there are other age-dependent factors leading to decrease of ouabain receptor's affinity to agonist. Obtained data indicate also the involvement of intracellular mechanisms of cell volume regulation having different sensitivity to ouabain. The literature data on higher ouabain sensitivity of intracellular Ca buffering structure [51,52] allow us to consider this buffering system responsible for age-dependent changes of dose-dependent ouabain effect on cell hydration. This suggestion is confirmed by the study of dose-dependent ouabain effect on 46Ca+ uptake (Figure 5) and efflux (Figure 6 and 7).

Pioneering work by Baker et al. [10] performed on perfused axon from Loligo forbesi, described in detail the correlation between Na+K+ pump and Na+/Ca++ exchange. There were distinguished two components of 22Na efflux: ouabain sensitive (Na+K+ pump) and ouabain non sensitive (Na+/Ca++ exchange) components. Our work performed on whole (non-perfused) and preliminary 22Na enriched snail neurones has shown that ouabain high concentration (>10^-4 M) has inhibitory effect on 22Na efflux as in axon, but its lower concentrations (<10^-4 M) have activation effect on 22Na efflux [4], which is due to the activation of Na+/Ca++ exchange in reverse mode [39]. It is obvious that such differences of ouabain sensitivity of Na+/Ca++ exchange in internally perfused squid axon and whole snail neuron can be explained by different membrane properties of squid axon and snail neuronal membrane or by ouabain-induced changes of Ca ions' gradient on

<table>
<thead>
<tr>
<th>Brain Zones</th>
<th>Control 1 (intact animals)</th>
<th>DW injection</th>
<th>%</th>
<th>P value</th>
<th>Control 2 (PS injection)</th>
<th>%</th>
<th>P value</th>
<th>Mannitol injection</th>
<th>%</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydration [water cont. g/g d.w.] (mg.)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Cortex</td>
<td>2.82 ± 0.05</td>
<td>4.4 ± 0.074</td>
<td>142</td>
<td>P = 0.4</td>
<td>3.1 ± 0.0374</td>
<td>110</td>
<td>P = 0.0000078</td>
<td>2.28 ± 0.022</td>
<td>26.5</td>
<td>P = 0.00039</td>
</tr>
<tr>
<td>Stem</td>
<td>3.09 ± 0.075</td>
<td>3.5 ± 0.072</td>
<td>134</td>
<td>P = 0.3</td>
<td>2.6 ± 0.055</td>
<td>15.8</td>
<td>P = 0.000025</td>
<td>2.28 ± 0.018</td>
<td>12.3</td>
<td>P = 0.0006</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>2.75 ± 0.041</td>
<td>3.2 ± 0.08</td>
<td>123</td>
<td>P = 0.8</td>
<td>2.6 ± 0.06</td>
<td>5.5</td>
<td>P = 0.0001</td>
<td>2.145 ± 0.019</td>
<td>17.5</td>
<td>P = 0.053</td>
</tr>
</tbody>
</table>

Table 1: Correlation between brain tissues hydration and the number of ouabain molecules in experiments with intraperitoneally injected distilled water (DW), physiological solution (PS) and mannitol.
membrane by modulation Ca-buffering intracellular systems which were removed in axonal experiments. Data obtained in presented experiments on clear age-dependent depressing of \( ^{45}\)Ca\(^{+}\)/Ca\(^{2+}\) uptake (Figure 5) and \( ^{45}\)Ca\(^{+}\) efflux (Figure 6 and 7) in brain tissues of rats conform the second explanation. As initial level of intracellular Ca ions is higher in brain tissues of old animals than in that of young ones [38] data obtained in present work on the lower intensity of \( ^{45}\)Ca\(^{2+}\) uptake in brain tissues of older animals than that of young ones is predictable (Figure 5).

Data that low ouabain concentration (10\(^{-6}\) M) which is unable to inactivate Na\(^{+}/K\(^{+}\) pump, but activates \( ^{45}\)Ca\(^{2+}\) uptake in brain cortex and cerebellum tissues, exclude the explanation of this effect by increase of intracellular Na ions [53]. It is known that \( ^{45}\)Ca\(^{2+}\) uptake by neuron can be realized also by potential-dependent Ca ionic channels [7]. The fact that 10\(^{-6}\) M ouabain concentration has more pronounced activation effect on \( ^{45}\)Ca\(^{2+}\) uptake than 10\(^{-4}\) M ouabain, when membrane potential is depolarized because of inactivation of electrogenic Na\(^{+}/K\(^{+}\) pump [54], excludes involvement of potential activated Ca channels in the realization of ouabain-induced \( ^{45}\)Ca\(^{2+}\) uptake stimulation. Therefore, as the difference between electrochemical gradients of these two ions (E\(_{\text{Na},2}\) and E\(_{\text{Ca},2}\)) serves energy for Na\(^{+}\)/Ca\(^{2+}\) exchange, the activation of Na\(^{+}/Ca\(^{2+}\) exchange in reverse mode could be explained only by the increase of E\(_{\text{Na},2}\) as a result of cell intracellular Ca adsorption. The fact that low-ouabain induced activation effect on \( ^{45}\)Ca\(^{2+}\) uptake in old animals weakens can be considered as a result of aging induced depression of Ca adsorption properties of intracellular structure because of their being Ca-saturated by high [Ca\(^{2+}\)], [38]. It is interesting that aging has a more pronounced inhibitory effect on \( ^{45}\)Ca efflux in subcortical than cortex tissue, while in cerebellum activation of \( ^{45}\)Ca\(^{2+}\) efflux was observed (Figure 6B).

Although the dysfunction of \( ^{45}\)Ca\(^{2+}\) efflux in aging animals can be considered as a proven fact, its reason is not clear yet. It is known that Ca ions can be removed from cells by two mechanisms: Ca-pump and Na\(^{+}/Ca\(^{2+}\) exchange in forward mode. Notwithstanding the affinity of Ca-pump is higher than Na\(^{+}/Ca\(^{2+}\) exchange because the latter has incomparably higher rate than Ca-pump, it is suggested that Na\(^{+}/Ca\(^{2+}\) exchange has a more essential role in regulation of [Ca\(^{2+}\)] homeostasis [12,34]. There is a great number of literature data about the crucial role of \( \alpha_1 \), \( \alpha_2 \) isoforms of Na\(^{+}/K\(^{+}\) in regulation of Na\(^{+}/Ca\(^{2+}\) exchange. However, the detailed mechanism of their interaction stays unclear.

The existence of 6 components on curve of dose-dependent ouabain effect on \( ^{45}\)Ca\(^{2+}\) efflux (Figure 7), as in case of cell hydration (Figure 4), indicates that in the background of dose-dependent ouabain-induced modulation of cell hydration and \( ^{45}\)Ca\(^{2+}\) efflux are common intracellular mechanism(s) having the same age-dependent character. Data on heterogeneity within the family of previously described three \( \alpha_1 \), \( \alpha_2 \), \( \alpha_3 \) isoforms and clear age-dependency are new and could serve as a target for detailed investigation of metabolic mechanisms, dysfunction of which leads to cell pathology, including aging.

The fact that ouabain has a reverse dose-dependent activation on \( ^{45}\)Ca\(^{2+}\) uptake (Figure 5) and inhibition on \( ^{45}\)Ca\(^{2+}\) efflux (Figure 7) in young animals' brain tissues indicates that ouabain-induced increase of [Ca\(^{2+}\)], which has inhibitory effect on receptors affinity to agonist, is responsible for reverse dose dependent decrease of ouabain binding in range of 10\(^{-7}\)-10\(^{-8}\) M concentration (Figure 3A). Data on dose-dependent weakening of ouabain-induced cell dehydration in range of 10\(^{-11}\)-10\(^{-10}\) M (Figure 4), leading to increase of the number of receptors in membrane (Table 1) could serve as an evidence for this suggestion. Obtained data on age-dependent dysfunction of dose-dependent ouabain binding accompanied by inactivation of \( ^{45}\)Ca\(^{2+}\) efflux (Figure 6) and literature data on the high affinity of \( \alpha_1 \) receptors to Ca ions [53] confirm also this suggestion. However, a question rises which of these two phenomena is primary. On the basis of data presented in this work we suggest that the dysfunction of \( \alpha_1 \) Na\(^{+}/K\(^{+}\) pump isoform in aged animals is a consequence of [Ca\(^{2+}\)] increase.

10\(^{-4}\) M ouabain, having fully inhibitory effect on Na\(^{+}/K\(^{+}\) pump, activates \( ^{45}\)Ca\(^{2+}\) efflux in aging animals (Figure 7A). Therefore, ouabain-induced activation of \( ^{45}\)Ca\(^{2+}\) efflux in brain cortex of old animals can be explained by release of Ca\(^{2+}\) by intracellular storage. It is clear that data on activation effect of ouabain on \( ^{45}\)Ca\(^{2+}\) efflux in cortex of aged rats (Figure 6) cannot be explained by ouabain-induced inactivation of Na\(^{+}/K\(^{+}\) pump.

Since intracellular Ca\(^{2+}\) buffer system is extra sensitive to external and internal signals it can be suggested as mechanism responsible for ouabain-induced elevation of \( ^{45}\)Ca\(^{2+}\) efflux in old and inactivation in young rats' brain cortex and subcortex tissues. The reason of reverse age dependency of \( ^{45}\)Ca\(^{2+}\) efflux and its ouabain sensitivity in cerebellum observed in present work stays unclear and needs a more detailed investigation.

Taking together data obtained in present work on reversal ouabain effects, especially low ouabain concentration, on intracellular Ca buffer system and brain tissue hydration in young and old animals and literature data on constant circulation of endogenic ouabain like compound in blood [55], it can be concluded on possible beneficial effect of this compound on aged organisms and hazardous effect on young organisms, considering Ca-releasing from neurons in old and promotion of Ca enrichment process in young animal's brain. It is obvious that this suggestion can be considered as a working hypothesis which is the subject of our current study.

References
