Evolution of stress hormones and proteins in relation to developmental origins of health and disease

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Abstract
The mini-review is presented about the evolution of stress hormones (the components of hypothalamo-pituitary-adrenal axis) and stress proteins (heat shock proteins and others), as related to the concept of developmental origins of health and disease. The data on somatic growth retardation by glucocorticoids as mediators and targets of programming / imprinting phenomena are discussed in evolutionary aspect. Finally, two theoretical constructs are briefly introduced, the onto- and phylopatogenic model, with the last one having some perspectives of elaboration, as referred to biological evolution.

Introduction
At present two different types of stress are discussed: physiologic and cellular one [1]. Physiologic stress is known to occur on the level of the whole organism of animals and humans, and its mechanisms involve the following hormones: corticolderin (corticotropin-releasing factor, CRF) and arginine-vasopressin from the hypothalamus, adrenocorticotropic hormone (ACTH) from hypophysis and glucocorticoids (GC: cortisol, corticosterone) from adrenal cortex, composing the so called hypothalamo-pituitary-adrenal (HPA) axis. In addition, mechanisms of physiologic stress involve also catecholamines (adrenaline, noradrenaline), as well as oxytocin and many other bioregulators, but nevertheless, the fundamental role is attributed to HPA hormones [1,2].

On the other hand, cell reactions to stress are realized by means of so-called stress proteins, mainly by heat shock proteins (HSP), as well as metallothioneins (MT), annexins and some others. Today the interactions between stress hormones and proteins continue to be poorly understood [1,2].

Beginning from the end of eighties of the last century, the concept emerged on developmental origins of health and disease (DOHaD), in which the central position is occupied by programming / imprinting phenomena. These phenomena take place in those situations when the action of certain factors including stress, in perinatal and some other periods of development provokes long-term consequences expressed partially in higher predisposition to a number of disorders already in adult state and in senescence. There exist important data showing principal role of GC in programming / imprinting phenomena [3,4].

However, quite reasonable question can emerge: when these phenomena appeared in the evolution? Obviously, this question is related to evaluation of the phylogeny of GC and other HPA hormones and perhaps, of stress proteins also. Therefore, in the present work we aimed at description of the evolution of stress hormones and proteins, focusing the main attention on the role of GC and other hormones in programming / imprinting phenomena in the light of DOHaD concept.

Evolution of hypothalamo-pituitary-adrenal axis hormones
During the last decades significant progress occurred in our understanding of the phylogeny of stress hormones. This progress was related principally to determination of primary structures of hormonal polypeptides in various species of animals and to sequencing of DNA in these species, as well as to using special computer softwares for comparisons of the sequences of aminoacid residues in polypeptides and of nucleotides in DNA, in order to reveal the degree of their similarity. Although GC are steroid hormones and therefore, such methods are not applicable to them, nevertheless, protein receptors of GC and their genes can be compared by these technologies.

And here quite interesting surprise is waiting for us: receptors of GC and enzymes for their biosynthesis were not found in the invertebrates [5], hence these hormones are unlikely to participate in programming / imprinting phenomena, even if such phenomena really exist in these species. Moreover, possible participation of GC in mechanisms of aging [6] also becomes doubtful for such species.

What for another stress hormones? Phylogeny of corticolderin (CRF) and related peptides: urocortin / urotensin I and urocortins type 2 and 3, as well as of their receptors CRF1 and CRF2 may be associated with two genome duplications during appearance of vertebrates in evolution [7,8]. Phylogeny of ACTH precursor proopiomelanocortin (POMC) is also associated with two genome duplications and emergence of related polypeptides: proenkephalin, prodynorphin and proprorphain [9], whereas the phylogeny of POMC is related probably to so called tandem duplication of a gene, in this case of melanocyte-stimulating hormone-MSH [10].

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The evolution of 4 out of 5 types of melanocortin receptors (MC1R, MC2R, MC3R, MC4R) also appears to involve two genome duplications, whereas the emergence of MC5R involves, perhaps, tandem duplication of a gene. It should be noted, nevertheless, that only the type MC2R participates in binding and signaling of ACTH, whereas other types are receptors of various MSH forms. For such aim a special factor, melanocortin receptor-associated protein (MRAP) type 1 is employed, in order to enhance the affinity of MC2R to ACTH. Probably, coevolution of POMC and related protein precursors may occur, in parallel with melanocortin receptors [10].

What for receptors of GC and other steroid hormones, in this case, at the very beginning nuclear receptors emerge for binding lipophilic compounds with low affinity [5], thereafter receptors for estrogens and progestins appear first, being followed by receptors for androgens and corticoids [11], and corticoid receptors are involved at first in mineralocorticoid regulation and only later on separate receptors for mineralocorticoids and glucocorticoids emerge. The appearance in evolution of specialized mineralocorticoid hormone – aldosterone is rather late evolutionary event, and for blocking the action of GC on mineralocorticoid receptors a special mechanism is used on the basis of enzymatic inactivation of GC by means of 11beta-hydroxysteroid dehydrogenase on pre-receptor stage.

Let's discuss briefly now the evolution of principal stress proteins.

**Phylogeny of heat shock proteins, metallothioneins and annexins**

In contrast to hormones of physiologic stress, many proteins of cell stress are found in the most primitive early life forms on the planet Earth. In fact, HSP were identified practically in all species studied, from bacteria to higher metazoans [12]. Probably, this situation is related to extremely important their functions in proteostasis, i.e. providing stability and renaturation of various proteins, as well as in proteolysis with ubiquitin participation in those cases when renaturation cannot be performed. Therefore it is not surprising that the degree of similarity of HSP70 and HSP90 in E.coli and vertebrates is close to 50%, whereas yeast cells already possess practically all necessary HSP [13].

In addition, MT are also quite ancient proteins, although the authors that analyzed their phylogeny, admit the complexity of such task, because of heterogeneity of MT, expressed in different affinity to Zn²⁺ and Cu⁺ ions [14]. Nevertheless, the emergence of 4 types of MT in higher vertebrates appears to involve also two genome duplications. However, as in all other variants of such explanation, the next question would be: how did the most ancient, early precursor of MT emerge?

To this aim, an attempt was made for analysis of life evolution on the planet Earth, beginning from the very onset of its emergence [14] and separating the whole time scale to three epochs, really giant in their extension:

- with high Fe content and low concentrations of Zn and Cu in water, when bacteria species with anaerobic metabolism dominated, due to atmosphere without oxygen (from 4 to 2,4 billion years ago);
- with increasing Zn and Cu concentration and decreasing Fe content in separate aqueous compartments with more oxygenated atmosphere, when aerobic bacteria and unicellular eukaryotes emerged (from 2,4 to 0,5 billion years ago);
- with low Fe content and rather high Zn and Cu concentrations in water, when metazoans emerged in fully oxygenated atmosphere (from 0,5 billion years ago to the moment close to present).

Therefore, the emergence and phylogeny of MT may be related to transition from the first out of these 3 epochs to the second one and partially, from the second to third one, in order to providing the transport of Zn and Cu, in association with the necessity of regulation by their ions of gene expression and various enzymatic activities.

What for annexins / lipocortins, they appear to exist only in metazoans, and their phylogeny involved probably tandem duplication of a gene [15].

Let's discuss now, how the phylogeny of stress hormones and proteins may be related to programming / imprinting phenomena, as well as to aging and strategies of life history.

**DOHaD concept, aging and the evolution of stress hormones and proteins**

If GC are involved in programming / imprinting phenomena as principal factors, then it becomes clear that such phenomena are possible only in vertebrates. As a matter of fact, one of main details of such GC involvement is growth inhibition in organs and whole body, with concomitant induction of mature, more differentiated tissue phenotype.

It is interesting that one of the authors who demonstrated growth-inhibitory GC action was a pharmacologist from St. Petersburg, M.V. Nezhentsev, using hydrocortisone administered to rats [16]. We were able to evaluate growth-inhibitory action of more active GC dexamethasone in rats only in the nineties of the last century and at the beginning of the current one [17,18].

However, in two small studies we could show also growth-inhibitory influence of dexamethasone in chicken [19] and canine pups [20], and in the first case we observed short-term growth retardation by means of mininknemometry, whereas in the second case the disruption of growth proportionality in the members was established under the influence of GC.

Moreover, we were able to collect together rather scarce literature data about growth-inhibitory GC action in various species [19]. All these data indicate the conservation in evolution of the mechanism of GC involvement in programming / imprinting phenomena on the basis of growth retardation.

What for aging, the role of GC and stress in its occurrence is not totally clear even in higher mammals and humans [6]. However, the suggestion is possible that species with greater longevity possess finely regulated reactions to stress, without inducing the negative allostatic consequences in the long term. Nevertheless, significant efforts still should be made for revealing the role of stress and its hormonal and proteinaceous mediators in phylogenetic aspects of aging. This topic becomes even more important, if to take into account that in modern gerontology the data obtained on species with short lifespan, like nematodes and Drosophila are considered as significant for other species, in spite of the fact that these invertebrates have principal differences with vertebrates and humans in the mechanisms of stress, for example, because of the absence of GC in the former.

In addition, many invertebrates, as well as a number of vertebrates have principally different (as compared to humans and some other vertebrates) strategies of redistribution of energetic and material resources between the maintenance of the body, growth and reproduction (the so-called r- and K-species) [21]. It is clear that not
only the occurrence of aging is different in them, but the contribution of stress and its hormonal and proteinaceous mediators may be quite different also.

Finally, the strategies of life history may be quite different, what should be considered in the usage even of mammals like rats and mice as the models of hormonal regulation of ontogeny in humans. Indeed, taking into account that rats are born in less mature state, as compared to human, the administration of GC to neonatal rats, as a matter of fact, is modeling their use in the 3rd trimester of gestation in humans or in premature newborns [22].

Conclusion

Therefore, even if the programming / imprinting phenomena are manifest by invertebrates, they should occur without participation of GC. What for vertebrates, the role of GC in such phenomena demonstrates evolutionary conservative mode, but it may represent some differences, as related to the strategies of life history.

Currently we are elaborating the ontopathogenic model in the framework of DOHaD concept. This model considers the pathology of various disorders during the whole ontogeny, with the possibility of programming / imprinting and embedding phenomena in critical periods of development, provoking the consequences till adult state and even senescence [23,24]. However, another theoretical model is possible, the phylopathogenic one [25], with multi- and intergenerational transfer of the risk of different pathologies. However, for its elaboration we should estimate at first the evolutionary aspects of various diseases.

References