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Methemoglobinemia as biomarker and precursor of brain capillary oxidative damage link to ferric iron accumulation and originator of neurodegenerative diseases

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Abstract
Our main aim is to make a contribution to the establishment of sources of oxidants as the key factors in understanding the role that oxidants and oxidative stress play in the pathogenesis of neurodegeneration, i.e. in the progression of Alzheimer’s disease (AD). Our original observation pointed out the main difference between the hemoglobin and methemoglobin degradation, did from the heme oxygenation when a hemoglobin results in ferrous (Fe2+) iron, but methemoglobin catabolism produces ferric (Fe3+) iron. The methemoglobin plays a role of the carrier, the donor of cytotoxic and redox-active ferric (Fe3+) iron, and it also acts as the originator of neurodegenerative diseases. The abundant and permanent source of redox-active ferric (Fe3+) iron which, without Ferrous-Ferric inversions, has “in situ” direct impact on endothelial small vessels in the brain accumulates and increases the rate of capillary endothelial cell apoptosis and possibly crosses into brain parenchyma to the astrocytes, glia, neurons, and other neuronal cells. Our understanding of the transport and neuronal accumulation of ferric (Fe3+) iron points to how microvessels are organized into a well-structured neurovascular unit with harmful consequences to the brain. Our previously conducted research found that the neonatal jaundice incidence (p=0.034), heart murmur at a later age (p=0.011) and mild disorders in children and adults such as dyslalia and learning/memory impairments (p=0.002) were significantly higher than in children and adults of control mothers without pregnancy methemoglobinemia. The consequence are performed as initial brain iron harmful effects from the mother-fetal pregnancy methemoglobinemia complication, and according to our hypothesis in humans could be followed with the neuronal death, the disease aging process, and leading finally to the severe disorders as AD, PD and other neurodegenerative diseases.

Introduction
It is still not known why iron levels are abnormally high in some regions of the brain in neurodegenerative disorders and it is also not clear whether iron accumulation in the brain is the initial event that causes neuronal death or is a consequence of the disease process. According to the previous research, we consider that, at least in some neurodegenerative disorders, brain iron misregulation is the initial cause of neuronal death and that this misregulation might be the linked to either genetic or non-genetic factors.

Ferric-iron brain accumulation as a cause of neurodegenerative brain disease: A new insight to understanding the mechanism of iron transport
Our research focused on people who continuously inhale strong oxidants such as NOx (NO and NO2) and it showed that these oxidants can reversibly oxidize oxyhemoglobin (Fe2+) to methemoglobin (Fe3+). Irreversible methemoglobinemia can occur because of the disruption of the oxidant/antioxidant balance, supported by the synergic SO2 metabolites, leading to the degradation of antioxidant thiols [1]. Methemoglobin by itself and heme have prooxidant properties and induce structural and functional changes in the vascular endothelium [2,3]. These changes include cell growth arrest, senescence, morphological alterations and cell apoptosis, and they lead to both vessel trombosis and endothelial cell denudation under the influence of redox-active ferric iron (Fe3+), which is a product of heme-oxygenase, responsible for methemoglobin-heme degradation.

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The toxicity of $\text{H}_2\text{O}_2$ is also dependent upon its reaction with ferrous iron to form hydroxyl radicals by the Fenton reaction. The ferrous iron needed for this reaction is formed by the reduction of cellular ferric iron by superoxide ions [6]. We would like to emphasize the difference between physiological hemoglobin catabolism and pathological methemoglobin catabolism, because their different final products, ferrous and ferric iron, have distinct characteristics. Ferrous iron has the potential for catalyzing and generating highly cytotoxic hydroxyl radicals as the Fenton reactions (ferrous iron + $\text{H}_2\text{O}_2$ $\rightarrow$ ferric iron + OH + OH). Ferric iron is then reduced back to ferrous iron, peroxide radical and proton by the same hydrogen peroxide.

The substantial difference between the intracellular ferric iron originates from Fenton reaction, and ferric iron originates from methemoglobin catabolism influence a level of methemoglobin cellular uptake which increases the methemoglobinemia leading to ferricIron-induced oxidative stress injury.

We consider that during methemoglobin catabolism, the final ferric iron product is a significantly added source of ferric iron derived from the Fenton reaction and its continuous formation has an impact upon the brain’s neurovascular unit.

According to the suggestions made in the previous research, we consider that, at least in some disorders, brain iron misregulation is the initial cause of neuronal death and that this misregulation might be the result of either genetic or non-genetic factors [7], and is considered to have autoimmune determinants [8]. Our previous research results suggest that methemoglobin plays a particularly important role as the carrier and donor of the cytotoxic and redox-active ferric (Fe$^{3+}$) iron, and determines how iron is transported intracellularly (Figures 1-4).
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Results

Our results point out the consequence of the toxic environmental oxidants, caused by brain damage with a view to the role of methemoglobin catabolism in pregnancy as the source of ferric (Fe³⁺) iron form concentrated in various brain regions. Methemoglobin and hemolysis both occur as a result of oxidative stress, but the prevalent difference between them is that methemoglobin is a reversible phenomenon (oxidant–antioxidant balance) whereas hemolysis, which occurs as a result of oxidative stress on the erythrocyte membrane, is an irreversible event. Methemoglobinemia can additionally exacerbate an existing anemia, stimulating hypoxia that may be additionally dangerous. Our prospective study of methemoglobin in pregnancy revealed a significant rise the level of methemoglobin >1.5 g/L (r=0.72, p < 0.01) in the air-polluted exposure period, which can be explained on the basis of an oxidant–antioxidant imbalance, resulting in methemoglobinemia [15]. Methemoglobinemia and stillbirths recorded throughout exposure period are significantly higher than those recorded in the control period (p = 0.0205) and the frequencies of reproductive loss were significantly lower in the control than in the exposure period (p < 0.05) [16].

As we have found no evidence of the consequences of mother methemoglobinemia on the fetus, the second objective was to direct attention to methemoglobin as an early biomarker of the environmental toxic of oxidative stress effects, which puts pregnancy at risk and may later impair the health of newborns, children and adolescents. The conducted proper research found that neonatal jaundice incidence (p=0.034), heart murmur at a later age (p=0.011) and mild disorders in children and adults such as dyslalia and learning/memory impairments (p=0.002) were significantly higher than in children and adults of control mothers without pregnancy methemoglobinemia [17]. Lavezzi AM, Mohorovic L, Alfonsi G, Coma MF, Matturri L, Brain iron accumulation in unexplained fetal and infant death victims with smoker mothers. The possible involvement of maternal methemoglobinemia. Lino Rossi Research Center for the study and prevention of unexpected perinatal death and SIDS, Department of Surgical, Reconstructive and Diagnostic Sciences, University of Milan, Italy.

Discussion

On the basis of our observation, we point out the specific cellular methemoglobin and heme catabolism when the last product is the ferric iron which, together with cytotoxic and paramagnetic property, has a notable role “in-situ”. We suggest that ferric iron and ferric iron-induced oxidative stress constitute a common mechanism involved in the development of neurodegeneration, and we also suggest the initial cause of neuronal death as a result of environmental toxic factors. The experiments showed that ferric and ferrous iron can enter cells via different pathways, however they do not indicate which pathway is dominant in humans [19].

Heme, the major functional form of iron, is synthesized in the mitochondria. Smith MA, et al. suggests that iron is able to participate oxidation in in-situ and it readily catalyzes an H₂O₂-dependent oxidation, and they indicate that iron accumulation could be an important contributor towards the oxidative damage of Alzheimer’s disease [20]. According to our point of view, our work supports the above statement about the importance of disturbed heme metabolism when the heme oxygenase-1, which is an enzyme that catalyzes the
conversion of methemoglobin and heme to ferric iron, is increased in Alzheimer’s disease, suggesting the increased heme turnover as a source of redox-active iron. Perry G, et al. have found that while mitochondrial DNA and cytchrome C oxidase activity are increased in Alzheimer’s disease, the number of mitochondria is decreased, indicating the accelerated mitochondria turnover and suggesting mitochondrial dysfunction as a potentially inseparable component of the initiation and progression of Alzheimer’s disease [21]. It was also discovered that oxidative damage occurs primarily within the cytoplasm rather than in mitochondria. This leads to the hypothesis that mitochondria, as a source, provide hydrogen peroxide, which, as an intermediate, once in the cytoplasm, will be converted into highly reactive hydroxyl radicals through Fenton reaction in the presence of metal ion and cause damage to the cytoplasm [22]. Cell apoptosis is initiated by extracellular and intracellular signaling pathways, the death receptor- and the mitochondria-mediated pathway.

Various pathologies can result from oxidative stress-induced apoptotic signaling, consequently leading to ROS increases and/or antioxidant decreases, the disruption of intracellular redox homeostasis, and irreversible oxidative modifications of lipid, protein, or DNA [23]. Furthermore, iron participates in diverse pathologic processes by way of the Fenton reaction, which catalyzes the formation of reactive oxygen species (ROS). To test the hypothesis that this reaction accelerates apoptosis, Jacob AK, et al. used human umbilical vein endothelial cells (HUVECs) as surrogates for the microvasculature in vivo. HUVECs were loaded with Fe [III] (ferric chloride and ferric ammonium citrate) and executed apoptosis after a heat shock stimulus [24]. Brain iron is a major contributor to magnetic resonance imaging (MRI) contrast in the normal gray matter. Non-heme brain iron is present mainly in the form of ferritin. The quantitation of non-heme brain ferric iron indicated by MRI helps in the diagnosis and monitoring of different neurological diseases [25]. Most of the brain’s non-heme iron is believed to present as a storage pool consisting of ferritin or hemosiderin and also as a product of methemoglobin catabolism [26]. However, the concentration of transferrin–bound iron is always far too small to affect MRI. This fact suggests that the role of methemoglobin catabolism as the source of ferric iron (FeIII) form concentrated in various brain regions should be considered. Nowadays, non-heme-bound Fe3+ is quantified by using Magnetic Resonance Imaging (MRI), thanks to its paramagnetic properties. It is believed that most non-heme-bound iron is deposited in the form of ferritin, hemosiderin, or methemoglobin catabolic products, whereas transferrin-bound iron concentration is always low and cannot be detected by MRI [27]. Recent research results indicate a ferricytride-magnetite core-shell ferritin structure. It was also found that the magnetite in the ferritin iron core is not a source of free toxic ferrous iron, as it had been previously believed. Therefore, the presence of magnetite in the ferritin cores of patients with Alzheimer’s disease is not the cause of their increased brain ferrous iron (II) concentration [28].

Conclusion

Abundant in the source of cytotoxic and redox-active ferric (Fe3+) iron which without Ferrous- Ferric inversions, “in situ” as a cause of iron–induced cellular oxidative stress, have a direct and specific impact on the brain endothelial small vessels, and increase the rate of endothelial cell apoptosis and so make possible the accumulation of methemoglobin, heme and ferric iron, in brain parenchyma. In conclusion, we point out the importance of methemoglobinemia not only as the biomarker and precursor of environmental oxidants’ noticeable effects, but also as a carrier and donor of the redox-active ferric iron form. We identify ferric iron as the originator of the accumulation of brain parenchyma that has an important role in crossing the brain’s microvessels to the neurons (neurovascular unit damage), causing neuronal death, continuous ageing process, and finally leading to severe neurodegenerative disorders such as AD, PD, MS and other diseases. Nevertheless, as to the relation between environmental oxidants and the pathogenesis of neurodegenerative diseases, need further research.

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