Propentofylline decreases hypothalamic astrogliosis induced by hypercaloric diet in the rat

A propentofilina diminui a astrogliose hipotalâmica induzida pela dieta hipercalórica no rato

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ABSTRACT

Obesity is associated with a chronic and low-grade inflammatory response in the hypothalamus, where astrogliosis occurs with the upregulation of the astrocyte structural protein GFAP. As propentofylline (PPF) has inhibitory effects on astrocyte and microglial activation during inflammation, this study aimed to investigate if this xanthine derivative could decrease the astrocyte reaction induced by a hypercaloric diet (HD). Male Wistar rats were divided into four groups: NDS – rats receiving a normocaloric diet (ND) and daily saline solution; NDP – rats receiving ND and daily PPF (12.5 mg/kg/day, intraperitoneal route); HDS – rats receiving HD and saline solution, HDP – rats receiving HD and PPF. On the 21st day, rats were anesthetized, and perfused, and brains were collected for GFAP immunohistochemical study in the hypothalamus. Results showed that HD induced increased weight gain and hypothalamic astrogliosis. Propentofylline decreased the expression of GFAP in the HDP group, although it did not affect the weight gain induced by this diet.

Keywords: astrocytes; central nervous system; inflammation; obesity; xanthine.

RESUMO

A obesidade está associada com uma resposta inflamatória crônica e de baixo grau no hipotálamo, onde ocorre astrogliose com a superexpressão da proteína astrocitária GFAP. Como a propentofilina (PPF) possui efeitos inibitórios sobre a ativação astrocitária e microglial durante a inflamação, este estudo visou a investigar se esta xantina podia diminuir a reação astrocitária induzida pela dieta hipercalórica (HD). Ratos Wistar machos foram divididos em 4 grupos: NDS – ratos recebendo dieta normocalórica (ND) e solução salina diária; NDP – ratos recebendo ND e PPF diária (12.5 mg/kg/dia, via intraperitoneal); HDS – ratos recebendo HD e solução salina, HDP – ratos recebendo HD e PPF. No 21º dia, os ratos foram perfundidos e os encéfalos, coletados para estudo imuno-histoquímico para a GFAP no hipotálamo. Os resultados mostram que a HD induziu aumento do ganho de peso e astrogliose no hipotálamo. A PPF diminuiu a expressão de GFAP no grupo HD, embora não tenha afetado o ganho de peso induzido por esta dieta.

Palavras-chave: astrócitos; sistema nervoso central; inflamação; obesidade; xantina.

The concept that obesity could affect the central nervous system (CNS), specifically the brain, has recently emerged1. Cognitive abnormalities, specifically learning and memory deficits, have been reported in obesity and typical comorbid disorders12. There are several associations between obesity and neurological disorders, including sleep apnea, anxiety, manic depressive disorders and increased risk of having a cerebrovascular accident1. It also has been reported that obesity is associated with changes in volume or density of gray and white matter, assessed by magnetic resonance imaging13. Obesity-associated structural abnormalities in the brain14 contribute to a higher incidence of diseases, such as dementia15 and Alzheimer's disease16. Although the exact cause remains unknown, it has been assumed that CNS disorders are triggered by multifactorial events including neuroinflammation and oxidative stress17. Neuroinflammation leads to resident glial cells (astrocytes and microglia) activation, which results in invasion of circulating immune cells and production of proinflammatory cytokines, such as tumor necrosis factor-α (TNF-α), interleukin (IL)-1β, and IL-6, and reactive oxygen species18.

Therefore, obesity is characterized by chronic and low-grade inflammation in several tissues, including the CNS, and particularly the hypothalamus, which is known to regulate food intake and energy expenditure19,20. The arcuate and the paraventricular nuclei are two key hypothalamic areas of this central network and play a critical role in maintenance of energy balance21.
Astrocytes are known to be the most numerous and diverse neuroglial cells in the CNS. They are dynamic cells that respond to changes in the CNS by undergoing morphological and functional alterations that affect neuronal activity. In response to CNS insults, astrocytes develop a hypertrophic or reactive phenotype termed astrogliosis, which is characterized by the upregulation of specific structural proteins, such as glial fibrillary acidic protein (GFAP) and vimentin. Hypothalamic astrogliosis has been widely associated with high-fat diet-induced obesity.

Propentofylline (PPF) is a xanthine derivative that has demonstrated profound neuroprotective, antiproliferative and anti-inflammatory effects in several animal models. Clinically, it has shown efficacy in degenerative vascular dementia and as a potential adjuvant treatment in Alzheimer’s disease, schizophrenia and multiple sclerosis. It probably depresses activation of microglial cells and astrocytes, which is associated with neuronal damage during inflammation and hypoxia, and consequently decreases glial production and release of damaging proinflammatory factors. It has previously been seen that PPF decreased the peripheral astrogliosis observed following gliotoxic injury in the rat brainstem.

Thus, the objective of the present study was to investigate if PPF could attenuate the hypothalamic astrogliosis induced by a hypercaloric diet (HD) as seen in previous studies.

**METHODS**

The animal procedures were performed in accordance with the guidelines of the Committee on Care and Use of Laboratory Animal Resources and Brazilian Institutional Ethics Committee guidelines (University Paulista, protocol number 376/15, CEUA/ICS/UNIP). All efforts were made to minimize animal suffering.

Twenty-four adult male Wistar rats, 12 weeks of age and weighing 300-350 g, were divided into four groups (n = 6 in each): NDS group – rats receiving a normocaloric diet (ND) and treated with 0.9% saline solution; NDP group – rats receiving the ND and treated with PPF; HDS group – rats receiving a hypercaloric diet (HD) and treated with 0.9% saline solution, HDP group – rats receiving the HD and treated with PPF.

Upon arrival at our laboratory, rats were housed in microisolator cages (four rats per cage) under controlled temperatures (22–25°C) and humidity (50–65%), in artificially-lit rooms on a 12-hour light-dark cycle (lights on 07:00) and food and water ad libitum.

Rats from the HDS and HDP groups received free access to the HD, which was the highly palatable liquid diet supplied by Ensure (Abbott Laboratorios do Brasil, São Paulo, Brazil; total of 1 kcal mL⁻¹). Each 231-kcal bottle contained 1.7 g polyunsaturated fat, 3.59 g monounsaturated fat, and 2.2 g saturated fat, with no trans fat. It was presented in a graduated cylinder with a stopper, with 600 mL per bottle. The ND (nutritionally balanced food) was irradiated laboratory chow (Nuvilab; Sogorb Indústria e Comércio, São Paulo, Brazil, with the following values per 100 g of solid food item: 4.2 kcal g⁻¹, 56% carbohydrate, 19% protein, 4.5% cellulose, 5% vitamins and 3.5% g total fat). This was given to all experimental groups, including those receiving the HD. Rats from the HD and ND diet groups were housed four per cage and Ensure and laboratory chow were made available to each cage as a whole. Both diets were replenished daily.

The animals were weighed weekly for 21 days and weight gain was defined as the ratio between the final weight and initial weight.

Rats from the NDP and HDP groups were treated daily with a PPF solution (20 mg/ml, Agener União Química, São Paulo - SP) by intraperitoneal route using 12.5 mg/kg/day during the experimental period. Rats from the NDS and HDS groups received 0.9% saline solution intraperitoneal injections daily.

On the 21st day, the rats were anesthetized (thiopental, 60 mg/kg intraperitoneally, Thiopentax, Cristália, Itapira, Brazil) and submitted to intracardiac perfusion with 4% glutaraldehyde in 0.1 M Sorensen phosphate buffer (pH 7.4). The brains were collected and fixed in 10% buffered formalin for 72 hours. Coronal sections of each brain were made to reach the periventricular area of the hypothalamus (using the following coordinates - 5.64 mm from interaural line and -3.36 mm from the bregma). The tissue was embedded in paraffin for processing for conventional histological procedures. The GFAP immunohistochemistry was performed using the chain polymer-conjugated staining method (Dako EnVision System). We used polyclonal rabbit anti-GFAP immunoglobulin (1:50; Z033401, Dako, Glostrup, Denmark) as the primary antibody followed by the EnVision+ Kit (HRP, Rabbit, DAB+, K4011, Dako, Glostrup, Denmark). Three sections (5 µm thick) per rat were made and, from each individual section and using a 40x objective, ten photomicrographs of the periventricular hypothalamicus were taken. The area of GFAP+ cells and their processes, marked in brown, was automatically calculated, in pixels, using Image-ProPlus 6.0 software (Media Cybernetics, Silver Spring, USA) calibrated with digital color filters such that only positive cells were included and background staining was excluded from the measurement.

Homoscedasticity was verified using the Bartlett’s test. Normality was verified using the Kolmogorov-Smirnov test. Two-way ANOVA followed by Tukey’s test (for body weight gain) or the Sidak test (for GFAP expression) were used to analyze data with two factors (diets and treatments). Results were expressed as means ± standard errors of the means. In all cases, results were considered significant at p < 0.05.
RESULTS

Increased body weight gain by the 21st day was seen in the HDS and HDP groups relative to the NDS and NDP groups (p < 0.0001) (Figure 1). As for the hypothalamic GFAP expression, differences between diets (p < 0.0001) and treatments (p < 0.0001) were observed, with interaction between factors (p < 0.0001). The Sidak post-test indicated an increased GFAP expression in the HDS group relative to the other experimental groups (NDS, NDP and HDP) (Figure 2). Propentofylline was able to decrease the astrogliosis induced by the HD in the HDP group, although the expression of GFAP in this group was still greater compared to the NDS and NDP groups (p < 0.01).

The expression of GFAP in hypothalamic astrocytes from the different groups are seen in Figure 3.

DISCUSSION

Hypothalamic inflammation may exert a paradoxical effect on energy homeostasis depending on the time course, the involved proinflammatory signals, and the degree of inflammation. An intense acute inflammatory response, as in an infection, induces a state of negative energy balance, although chronic and mild inflammation, as seen in obesity, has the opposite effect9. In our study, an inflammatory response was achieved in the hypothalamus, as suggested by the strong increase in the astrocyte expression of GFAP, indicative of intense astrogliosis in response to the HD. The differences in the weight gain seen in rats fed with the ND or HD from the beginning of the experiment until its conclusion 21 days later, confirmed the effect of the HD we used in creating overweight or obese rats.

Figure 1. Weight gain (in grams) in 21 days. Groups comprise the NDS (normocaloric diet + saline), NDP (normocaloric diet + propentofylline), HDS (hypercaloric diet + saline) and HDP (hypercaloric diet + propentofylline). Data are expressed as means ± standard errors of the means. ****p < 0.0001.

Figure 2. GFAP immunostaining (represented as total count of pixels) in groups comprising the NDS (normocaloric diet + saline), NDP (normocaloric diet + propentofylline), HDS (hypercaloric diet + saline) and HDP (hypercaloric diet + propentofylline). Data are expressed as means ± standard errors of the means. ** p < 0.01; **** p < 0.0001.

Figure 3. Astrocyte GFAP expression in the periventricular zone of the hypothalamus. Groups comprise the NDS (normocaloric diet + saline), NDP (normocaloric diet + propentofylline), HDS (hypercaloric diet + saline) and HDP (hypercaloric diet + propentofylline). 3v - third ventricle; e - ependymal cells; arrowheads - astrocyte processes. GFAP immunohistochemistry.
Both leptin (an anorexigenic hormone primarily produced by adipose tissue) and ghrelin (the most important orexigenic hormone identified to date) are capable of inducing rapid modifications in synaptic inputs to neurons that secrete the anorexigenic pro-opiomelanocortin and the orexigenic neuropeptide Y in the arcuate nucleus of the hypothalamus. However, food intake and body weight maintenance depend on the proper functioning of several hypothalamic areas, such as the ventromedial (controlling appetite, body weight and insulin regulation), the lateral complex (appetite and body weight control), the arcuate (feedback and control of anterior pituitary, including GH and TSH secretion), the posterior (thermoregulation), the supraoptic (fluid balance), the suprachiasmatic (biological rhythms), the preoptic and the lateral (lateral anterior thermoregulation) and the paraventricular (fluid balance, anterior pituitary and autonomic control, including regulation of sympathetic induced lipolysis and thermogenesis). Given the complexity of its circuitry we have chosen to consider the whole hypothalamus as a unique structure of special interest in the observation of astrocytic GFAP expression. It would be simplistic to consider only the arcuate nucleus, the ventromedial hypothalamus or the paraventricular nucleus as important centers for controlling feeding and metabolism.

Hypothalamic astrocytes express various types of leptin receptors, indicating direct effects of the hormones on these cells. High-fat intake and obesity are associated with activation of inflammatory signaling pathways in the hypothalamus involving astrocytes and microglia and resulting in leptin and insulin resistance and disturbed control of homeostasis. In obesity, increased expression of leptin receptors in hypothalamic astrocytes may suggest the sequestration of leptin by astrocytes thus leading to decreased leptin signaling to neurons.

Astrocytes control both glucose and lipid transport and metabolism in the CNS. Apolipoprotein E, the most abundant lipid transporter in the CNS, is produced mainly in astrocytes, acting as a satiety factor in the hypothalamus, possibly mediating some of the inhibitory effects of leptin. It is widely known that astrocytes oxidize fatty acids to produce ketone bodies, which constitute an energy source for neurons. On the other hand, ketones mediate leptin and insulin signaling in the hypothalamus thus affecting energy homeostasis.

It remains to be determined if morphological changes in astrocytes and in the number of synaptic inputs to neurons of the arcuate nucleus are a cause or a consequence of the increased weight gain and its complications. Exposure to a high-fat diet was associated with reactive gliosis and this affected the structure of the blood-brain barrier, such that the pro-opiomelanocortin and neuropeptide Y cell bodies and dendrites became less accessible to blood vessels. It is unclear whether synaptic inputs are first removed and astrocytes respond by taking up the vacant space and/or if they actively remove synapses and occupy new spaces.

The astrogliosis observed in our investigation, due to administration of the HD, is known to be accompanied by increased cytokine expression and attenuation of leptin signaling in the hypothalamic neurons. Activated astrocytes may lose their homeostatic functions upon exposure to stressors, decreasing glutamate uptake and increasing the expression of deleterious proinflammatory molecules such as cytokines, nitric oxide, prostaglandins, among others, as an injury response.

Many different types of signaling molecules, including cytokines, are able to trigger and/or regulate astroglial reactions and can be released by all cell types of the CNS tissue, including invasive inflammatory/immune cells, neurons, microglia, oligodendrocyte lineage cells, pericytes, endothelia and other astrocytes.

In the CNS, PPF may serve as a glial modulator, with direct actions on microglia, and dose dependently decreases microglial proliferation and expression of inflammatory cytokines, such as TNF-α and IL-1β, as seen in response to lipopolysaccharide stimulation in vitro.

Known mechanisms of PPF include inhibition of cyclic AMP (cAMP) and cyclic GMP phosphodiesterases, and action as a reuptake inhibitor for the purine nucleoside and neurotransmitter adenosine by blocking the action of membrane nucleoside transporters. This may lead to increased intracellular cAMP levels and greater extracellular concentrations of adenosine, which stimulates adenosinergic neurotransmission and adenosine 2 receptor-mediated cAMP synthesis.

Regulation of cytokine production includes the adenylate cyclase-cAMP-protein kinase pathway. Propentofylline, a type III-IV specific phosphodiesterase inhibitor, although it decreases, in a dose-dependent manner, the production of the inflammatory cytokines TNF-α, IL-1 and IL-6 by mouse microglia stimulated by lipopolysaccharide in vitro, increases up to two or three times the production of the inhibitory cytokine IL-10, which suppresses cytokine release by microglia and macrophages and attenuates astroglial reactivity in vivo.

A Ca ++-dependent and excessive activation of glial cells is usually found in neuroinflammation and, in this context, increased levels of adenosine induced by PPF administration may perform a regulatory role on these Ca ++- and cAMP-dependent molecular signaling pathways that determine many cell-related functions, such as cellular proliferation rate, differentiation state, cytokine production, among others. A strengthening of the cAMP signaling, which can be achieved by adenosine agonists and by PPF, stimulates the production of trophic factors in astrocytes, apparently preventing a deleterious and secondary astrocytic activation caused by previous microglial upregulation. Although not entirely understood, it has been accepted that drugs that elevate extracellular adenosine and/or block the degradation of cyclic nucleotides, like PPF, may be used to counteract glia-related damage in CNS pathological processes.

Thus, morphometric analysis of GFAP expression in the present study unequivocally demonstrated that PPF...
decreased astrocytic activation 21 days after the onset of administration of the HD, probably by simultaneously suppressing the release of proinflammatory molecules, such as the above-mentioned TNF-α, IL-1β and IL-6, which may trigger and promote astrogliosis during inflammation, and by increasing secretion of the anti-inflammatory cytokine IL-10. In conclusion, our results clearly indicate that PPF may have a role in reducing astrocytic overactivation following hypercaloric or high-fat diets. In turn, PPF did not show any effect on reducing the weight gain induced by the HD.

References

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