ENDOCRINE SYSTEM PATHOLOGY
Hypothalamic lesions in mice treated with monosodium glutamate

INTRODUCTION

In an earlier report from this laboratory, neonatal treatment with monosodium glutamate (MSG) was shown to cause a disruption of the hypothalamic-pituitary-gonadal axis which induced Cushing’s obesity, lethargy, hypoagonism and sterility (1,4). The administration of monosodium glutamate (MSG) to mice in the neonatal period induces destructive lesions in the arcuate nuclei of the hypothalamus and results in a syndrome of obesity, stunting and hypothalamic (4). In spite of the arcuate nucleus which plays a major role in neuroendocrine and autonomic integration, treated animals consume less food than controls (1,5,6). Corticosterone level is, however, high in the MSG-obese mouse suggesting that ACTH secretion is not reduced. Whether a regulatory mechanism affecting fat metabolism in the mouse can be localized to the arcuate nucleus, or to other brain areas selectively destroyed by MSG treatment (8), requires further study, especially the topography of acute lesions of hypothalamus. The question if MSG at the same time destroys peptidergic fibers responsible for SRIF synthesis (inhibitor of ACTH secretion) or not, also waits the answer.

Having in mind the contradictory data on hypothalamic lesions in obese mice, the aim of the present work was thus to study the MSG-effect on hypothalamus.

MATERIALS AND METHODS

Twenty newborn C57BL/6J mice were injected daily with MSG (4,4 mg/g body wt) sc, on days 1-10 of life, for the first 5 postnatal days. Another group of 10 mice, which were injected with saline solution, 0.02 ml/g body wt, every 24 h for the first 5 postnatal days, served as controls. As adults, treated animals showed stunted skeletal development, marked obesity and sterility. Pathological changes were also found in several organs associated with endocrine functions. Studies of food consumption failed to demonstrate hyperphagia to explain the obesity. It is postulated that the adult syndrome represents a multifaceted neuroenocrine disturbance arising from the disruption of developing neural centers concerned in the mediation of endocrine function.

RESULTS

Treated animals were quite lethargic as adults, and they lacked the sleekness of body coat, seen in controls. Cushing-type of obesity was evident macroscopically. Basophils of pituitary gland were hyperplastic of large size and with foamy aspect. Cortical cells of the adrenal gland, symmetrically, were also hyperplastic.

Hypothalamic tissue. The arcuate nuclei were completely destroyed along with neuronal constituents in the median eminence. Instead these tissues, gliosis was found, the most important histopathologic indicator of CNS injury. So, astrocytes were hypertrophic and hyperplastic, their nuclei small and dark located in a dense net of processes (glial fibrils). Multifocal small foci of microglial nodules, associated with neuronophagia, were also observed.

DISCUSSION

The hypothalamic lesions induced by neonatal treatment with MSG are primarily in the preoptic and arcuate nuclei and in the median eminence. The arcuate nucleus plays a major role in neuroendocrine and autonomic integration (6), and lesions in this region of the hypothalamus result in disruption of the hypothalamic-pituitary-adenal axis. Corticosteron level is high in the MSG-obese mouse suggesting that ACTH secretion is not reduced. Whether a regulatory mechanism affecting fat metabolism in the mouse can be localized to the arcuate nucleus, or to other brain areas selectively destroyed by MSG treatment (8), requires further study, especially the topography of acute lesions of hypothalamus. The question if MSG at the same time destroy peptidergic fibers responsible for SRIF synthesis (inhibitor of ACTH secretion) or not, also waits the answer.

CONCLUSION

Brain lesions, obesity, endocrine gland dysfunction and hypothalamic multifocal lesions in mice treated with MSG suggest a complex hypothalamic-pituitary-gonadal disturbance.

REFERENCES

Immunophenotypical characteristics of pheochromocytomas

ABSTRACT

Pheochromocytoma is a rare tumor originating from the chromaffin cells in the adrenal medulla and at the extra-adrenal sites in the body (paraganglia). The purpose of this paper is to determine the immunophenotypical characteristics of pheochromocytomas in our biotopic material. Seven cases of pheochromocytomas were analyzed, using standard histological, histochemical and a panel of immunohistochemical stainings. All 7 cases showed marked positivity with synaptophysin and chromogranin. Four cases from 4 stained were positive for NSE and NFP. All epithelial markers were negative in most of the cases, except in two cases where weak signal was noted. Three cases from three stained were positive for S-100. Stainings for LCA and GFAP were negative in most of the cases, except in two cases where weak signal was noted. Three cases from three stained were positive for S-100. Stainings for LCA and GFAP were negative in most of the cases, except in two cases where weak signal was noted. To determine the immunophenotypical characteristics of the analyzed tumors, immunohistochemical analyses with a panel of the above mentioned antibodies were done. All 7 cases showed marked positivity with synaptophysin and chromogranin. Four cases from 4 stained were positive for NSE and NFP. All epithelial markers were negative in most of the cases, except in two cases where weak signal was noted. Three cases from three stained were positive for S-100. Stainings for LCA and GFAP were negative in most of the cases, except in two cases where weak signal was noted.

RESULTS

The analyzed operative material had dimensions varying from 22 to 200 mm, and the largest tumor mass weighed 1200 grams. The standard histological analyses showed tumor processes composed of chromaffin cells arranged around stromal blood vessels with abundant granular cytoplasm and large nuclei containing one or more nucleoli. The cellular and nuclear pleomorphism varied from case to case, as well as the presence and number of mitotic figures. In order to determine the immunophenotypical characteristics of the analyzed tumors, immunohistochemical analyses with a panel of the above mentioned antibodies were done. All 7 cases showed marked positivity with synaptophysin and chromogranin. Four cases from 4 stained were positive for NSE and NFP. All epithelial markers were negative in most of the cases, except in two cases where weak signal was noted. Three cases from three stained were positive for S-100. Stainings for LCA and GFAP were negative, and for bcl2, as a valuable prognostic antibody, were also negative. Most valuable were the immunohistochemical stainings with synaptophysin, chromogranin, neuron-specific enolase (NSE), neurofilament protein (NFP), S-100 protein, and bcl2 protein, which showed marked positivity in all stained tissue specimens.

DISCUSSION

According to the available literature (1), the incidence of pheochromocytomas is 1 case in year on 1-2 million population, but other authors (2) consider that the frequency of this entity is higher. Our series contains 4 cases analyzed during 12 months on 2 million population. We can not explain this higher frequency and it could be a subject of a further more extensive study. In the retrieved literature, only a few articles treating the immunohistochemical properties of the pheochromocytomas were found. The following antibodies were used for routine diagnosis of these tumors: NSE, S-100, cytokeratins, HMB 45. Most of the cases were positive for NSE and S-100 protein (3,4). In a study of 102 patients, in 29 cases treated as malignant, expression of bcl2 protein was found, and the rest were negative (5). In our analyzed 7 cases, most of the cases were positive for synaptophysin and chromogranin, as well as NSE, NFP and S-100 protein, which means that the panel of antibodies useful for diagnosis of pheochromocytoma could be extended with the above mentioned. The positivity for chromogranin in tissue specimens corresponds to the found increased levels of plasma chromogranin detected by RIA (6,7).

CONCLUSION

Immunohistochemical analyses are most valuable in the diagnosis of the tumors of chromaffin cells originating from the adrenal medulla and extra-adrenal sites, and could be applied especially when their histological appearance is unclear, cells are highly anaplastic, and the tumor is necrotic.

REFERENCE

We previously showed an increased level of progesterone in adult female rats following acute ethanol treatment. To determine whether this effect is the result of activation of adrenal glands we morphologically analysed the adrenal cortex in ovariectomized rats. In addition, a possible involvement of the opioid system in observed phenomenon was tested. Adult female Wistar rats were ovariectomized and three weeks after surgery intraperitoneally with: a) ethanol (4 g/kg), b) naltrexone (5 mg/kg) followed 45 min later by ethanol (4 g/kg) and c) naltrexone (5 mg/kg) followed 45 min later by saline. Untreated and saline-injected rats were used as controls. The animals were sacrificed 1/2 h after ethanol administration. Paraffin sections were stained with hematoxylin-eosin and analysed using stereological measurements. Morphometric analysis revealed that a single dose of ethanol significantly increased the volume density of zona glomerulosa and zona fasciculata and decreased the volume density of zona reticularis. Pretreatment with naltrexone prevented these ethanol-induced changes. The results indicate that the previously described increased level of progesterone following acute ethanol treatment, is a result of the activation of the adrenal glands and suggest the involvement of opioid receptors in this phenomenon. (This work was supported by the Ministry of Science and Technology of the Republic of Serbia.)
The calcitonin content in medullary thyroid carcinoma (MTC) appears to be the best marker for tumor behavior. Carcinoembryonic antigen (CEA) is also a reliable marker of follow-up in patients with MTC. In this study 6 cases of MTC, belonging to MEN II syndrome, were analyzed. MTC were associated with pheochromocytomas. The patients were between 32 and 46 years of age, their average age being 38.6 and they were three males and three females. The expression of CT and CEA was investigated using the APAAP method, with appropriate antibodies (DAKO) being applied. The diameter of tumors varied from 1 to 4 cm. Histological appearance of MTC was classical, with plentiful stromal amyloid in 3 cases and trabecular, glandular, and carcinoid-like, respectively. Analysis of CT and CEA expression is done. All MTC showed CT immunopositivity, among them 4 MTC were moderately to intensely positive and in 2 MTC CT positivity was mild. Intensive CT positivity correlated with classical histologic pattern and infrequent mitotic figures. MTC with numerous mitoses were mildly CT immunopositive. There was no correlation between CT positivity and stromal amyloid content. All MTC showed intense CEA immunopositivity but without any correlation between CEA positivity and histologic pattern, stromal amyloid content, calcifications of mitotic figures, respectively. In this study the correlation between CT immunopositivity and histologic pattern and mitotic figures among MTC inside MEN II syndrome was found. There was no correlation between CEA positivity and histologic pattern, stromal amyloid content, calcifications or mitotic figures, respectively.

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KEYWORDS: Carcinoma Medullary; Thyroid Neoplasms; Multiple Endocrine Neoplasia; Immunohistochemistry; Calcitonin

The differential diagnosis between adrenocortical adenoma and well-differentiated adrenocortical carcinoma may be the challenge even for the experienced pathologist. The aim of the study is to make histomorphological and immunohistochemical analysis of 27 nonfunctional adrenocortical carcinomas (ACC) obtained from Institute of Endocrinology, Clinical Center of Serbia, Belgrade in the 1986. to 1997. period. The patients were between 31 and 74 years of age, their average age being 54, and they were 16 males and 11 females. The expression of cytokeratin and vimentin was investigated using the PAP method, with appropriate antibodies (DAKO) being applied. The diameter of ACC was 5 cm and above. On gross examination the tumors proved to be ill-defined and showed numerous foci of necrosis, hemorrhage and cystic softening. The tumor histology (HE) revealed predominantly large, polygonal cells arranged in trabecular or alveolar patterns. Pleomorphism, atypia and numerous mitotic figures were present, as well as the invasion of the capsule. Cytokeratin reactivity was present in 12 cases (5 cases - weak immunopositivity, and 7 cases - moderate). 18 ACC were vimentin immunopositive (7-intense, 3-moderate, 8-weak). Immunohistochemistry is not of great value in confirming the diagnosis of ACC.

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KEYWORDS: Carcinoma, Adrenal Cortical; Immunohistochemistry; Keratin; Vimentin
Pheochromocytomas are rare neoplasms, solitarily or as a part of Multiple Endocrine Neoplasms syndrome, von Hippel-Lindau disease and neurofibromatosis. It consists of chromatofile cells which secrete catecholamines and manifested as a paroxysmal hypertensive crisis. Most frequently it is unilateral, situated in the adrenal gland, rarely intraabdominal beneath the bifurcation of the aorta or in the wall of the urinary bladder. The authors reported one case of a woman who had been operated at the Clinic of Urology of Serbian Clinical Center. A 38 years old woman have paroxysmal hypertensive crisis, after clinical and biochemical tests was found a tumor in the hilus of the left kidney. She had been operated in February in 1999, at Clinic of Urology of Serbian Clinical Center in Belgrade when a total nephrectomy and adrenalectomy were performed. On gross inspection, the kidney was sized 12x6x5cm, with a tumor mass in the size of 13x10x7 cm in the hilus of the kidney, nodular and lobular, gray-brown on cut section, with focal necrosis, haemorrhages and small cysts. The material was pathohistologically analysed by histochemistry and immunohistochemistry methods. Microscopically was found pheochromocytoma which consist of chromaffine tumor cells with granular cytoplasm. The tumor cells were arranged in trabecules and alveolar in form, with cellular and nuclear pleomorphism, lining the wall of capillaries and sinusoids. Rarely there were giant and bizarre cells and atypical mitotic figures. Deposits of tumor cells were found in the branches of the renal vein. The postoperative course was without complications, recidives and metastases. We conclude that pheochromocytoma may be a cause of hypertensive crisis, and its treatment is always surgical.

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Keywords: Pheochromocytoma; Hypertension; Adrenal glands