Endocrinological Differences between Hatano High- and Low-Avoidance Rats during Early Two-Way Avoidance Acquisition

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Abstract: Hatano high (HAA)- and low (LAA)-avoidance rats were selected from Sprague-Dawley rats genetically on the basis of their active avoidance behavior in a shuttle-box test. The purpose of this study was to investigate stress-related alterations of hormones corticotropin-releasing hormone (CRH), arginine-vasopressin (AVP), prolactin, and adrenocorticotropin (ACTH) in the brain and blood during early avoidance acquisition using two lines of Hatano rats. In paraventricular nucleus (PVN) of the hypothalamus, the CRH levels in HAA rats were significantly increased after shuttle-box tasks compared with before the tasks, whereas the CRH levels in LAA rats significantly decreased after shuttle-box tasks compared with before the tasks. In the HAA rats, the CRH and AVP levels in the median eminence decreased after shuttle-box tasks, whereas there were no significant differences in the levels between before and after shuttle-box tasks in LAA rats. The plasma concentrations of ACTH were significantly higher in HAA rats than in LAA rats after shuttle-box tasks. These results show that the response of CRH-ACTH was higher in HAA rats than in LAA rats. This phenotype may be an important reason for the high avoidance rates of shuttle-box tasks in HAA rats. These endocrine differences in early avoidance acquisition may be involved in regulation of their avoidance responses in the shuttle-box task.

Key words: ACTH, AVP, CRH, Hatano rat strains, prolactin

(Received 23 March 2011 / Accepted 12 June 2011)

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Introduction

Two lines of Hatano rats have been genetically selected and bred from Sprague-Dawley rats on the basis of their performance in shuttle-box tasks [31]. High- and low-avoidance animals (HAA and LAA) were selected on the basis of high and low rates of avoidance response, respectively, because the shuttle-box test produced individual differences. The Hatano lines also show differences in maze performance [31], behavioral development [32], and stress-induced gastric ulceration [4]. We previously investigated the effect of cross-fostering on shuttle-box avoidance and the open-field and Biel water maze learning tests [32]. These tests were unaffected by cross-fostering. Therefore, the genetic regulations rather than the epigenetic mechanisms are more important in the two Hatano lines.

Furthermore, these rats have endocrinological strain differences. The responses to restraint stress of the adrenocorticotropin (ACTH) secretions were higher in HAA rats than in LAA rats, and those of the corticosterone and prolactin secretions to restraint stress were lower in HAA rats than in LAA rats [2]. Regulation of endocrine systems in the brain is a complicated process involving stress and memory. In particular, the amygdala and hypothalamus are brain structures that are deeply involved in stress response and regulation of the hypothalamus-pituitary-adrenal (HPA) axis [23, 30]. Corticotropin releasing hormone (CRH) from the paraventricular nucleus (PVN) of the hypothalamus and/or circulating plasma corticosterone change as a consequence of restraint stress [24]. CRH has arousal properties, which are also part of learning and memory [22, 27]. Therefore, we hypothesized that there are strain differences in the hypothalamic and amygdaloid contents and regulation of CRH in the two lines of Hatano rats during early two-way avoidance acquisition.

CRH increases coproduction, corelease and cosecretion of arginine vasopressin (AVP) [38], and CRH induces ACTH release from the pituitary gland, which results in the adrenocortical release of corticosterone [21, 35]. AVP is involved in learning and memory processes, and enhances memory consolidation and retrieval processes [8, 13]. Passive avoidance behavior is affected by AVP [40]. In the present study, we also focused on the hypothalamic contents of AVP during early two-way avoidance acquisition.

The present study was conducted to clarify strain differences in endocrinological characteristics in the brain of two lines of Hatano rat in relation to the stress and learning of avoidance behavior.

Materials and Methods

Animals and experimental procedures

The animal experiment in this study was conducted in accordance with “Guideline for Animal Experiment in Hatano Research Institute, Food and Drug Safety Center.”

Adult male rats (13–15 weeks) from each strain, HAA rats (n=53) and LAA rats (n=49), were used. The animals were kept under a 12-h light-dark cycle (lights on from 7:00 AM to 19:00 PM) at a temperature of 23–25°C and relative humidity of 55 ± 5%. Food (CE-2, Clea Japan, Inc., Tokyo, Japan) and water were available ad libitum. Animals were killed by decapitation before and after shuttle-box test. Trunk blood was collected in heparinized tubes containing aprotinin and centrifuged immediately, and plasma was separated and stored at −20°C until assayed for ACTH and corticosterone. Brains were frozen on dry ice, and punched hypothalamic amygdala and PVN were obtained from 50-µm cryostat sections [36]. The stalk-median eminence was dissected with a pair of fine scissors under a stereoscopic microscope. These tissues and the anterior and posterior pituitary glands were homogenized in 0.85% (N/V) NaCl solution and centrifuged at 25,000 × g for 30 min at 4°C. The supernatants were stored at −20°C until assayed for CRH, ACTH, and AVP. The pellet was solubilized in 0.5 M PBS and analyzed for protein content by the method of Bradford [7].

Shuttle-box avoidance test

The rats were tested in a shuttle-box (TX-401L, Unicom Inc., Chiba, Japan) consisting of two 50 × 20 × 25 cm (l × w × h) compartments separated by a central hurdle (height 1.8 cm). For each trial, a 3-s conditioning stimulus (CS) consisting of a buzzer and light was followed by 3-s of the unconditioned stimulus consisting of the CS plus a 10-mA electric shock delivered through
the grid floor. The number of responses in which animals moved to the other side during the CS was recorded. After being initially exposed to a 5-min session of free exploration in the shuttle-box, the rats were submitted to 60 conditioning trials of avoidance conditioning on 3 consecutive days. The avoidance rate was estimated as the ratio of the number of avoidance responses to the number of all trials on Days 1 and 3.

Radioimmunoassay (RIA)

Concentrations of ACTH [25], corticosterone [39] and CRH [37] were measured by double-antibody RIAs using ^125^I-labeled radioligands as described previously. The intra- and inter-assay coefficients of variation were 11.3 and 11.9% for ACTH, and 9.8 and 17.5% for corticosterone, and 9.5 and 16.4% for CRH, respectively. Concentrations of AVP were measured with a commercial RIA kit purchased from Mitsubishi Chemical Corporation (Tokyo, Japan). The intra- and inter-assay coefficients of variation were 6.9 and 11.2% for AVP.

Statistical analysis

All data are expressed as standard errors of means (SEM). Statistical comparison between two strains was performed by two-way analysis of variance (strain × time) for repeated measures followed by Tukey-Kramer tests when appropriate. *P*<0.05 was considered to be statistically significant.

Results

Avoidance responses (Fig. 1)

A significant interaction of strain × time (strain × time, *F*(1, 61)=4.82, *P*=0.03; strain, *F*(1, 61)=12.70, *P*=0.0007; time, *F*(1, 61)=1.71, *P*=0.20) was found. Tukey-Kramer post hoc tests revealed no differences between HAA rats and LAA rats in avoidance rates on Day 1. The avoidance rates in HAA rats increased significantly on Day 3 compared with Day 1. On the other hand, the avoidance rates in LAA rats did not change between Days 1 and 3.

Hormone levels in plasma (Fig. 2)

Plasma concentrations of ACTH (a), prolactin (b), and corticosterone (c) were measured after shuttle-box tests both on Days 1 and 3 compared with before the tests in both strains. Main effects of strain and time were observed for ACTH, prolactin, and corticosterone, but no interaction effect of strain × time was found (strain × time, *F*(1, 54)=1.21, *P*=0.31; strain, *F*(1, 54)=4.63, *P*=0.04; time, *F*(1, 54)=21.05, *P*<0.0001; for ACTH; strain × time, *F*(1, 57)=0.56, *P*=0.58; strain, *F*(1, 57)=11.07, *P*=0.002; time, *F*(1, 57)=77.58, *P*<0.0001; for corticosterone; strain × time, *F*(1, 46)=0.05, *P*=0.96; strain, *F*(1, 46)=4.12, *P*=0.04; time, *F*(1, 46)=7.94, *P*=0.001 for prolactin). The concent-
trations of ACTH were higher in HAA rats than in LAA rats after shuttle-box tasks on Days 1 and 3. The concentrations of corticosterone were significantly higher in LAA rats than in HAA rats after shuttle-box tasks on Days 1 and 3. The concentrations of prolactin were significantly higher in LAA rats than in HAA rats after shuttle-box tasks on Day 1.

**CRH and AVP levels in amygdala (Fig. 3)**

A main effect of time for CRH levels \( (F_{(1, 22)}=4.87, P=0.02) \) was found, but no main effects were observed for strain \( (F_{(1, 22)}=0.30, P=0.59) \) or strain \( \times \) time \( (F_{(1, 22)}=1.67, P=0.21) \) in the amygdala. The concentrations of CRH in LAA rats significantly decreased after shuttle-box tasks on Days 1 and 3 as compared with before the tasks, whereas a reduction in the CRH concentrations in HAA rats was observed only on Day 1 after the shuttle-box test (a). No main effects of strain \( \times \) time, strain or time were found for the AVP levels (strain \( \times \) time, \( F_{(1, 23)}=0.25, P=0.78 \); strain, \( F_{(1, 23)}=0.001, P=0.98 \); time, \( F_{(1, 23)}=2.75, P=0.09 \)) in the amygdala.

**CRH levels in the PVN (Fig. 4)**

A main effect of strain \( \times \) time for CRH in the PVN \( (F_{(1, 22)}=3.67, P=0.04) \) was observed. The concentrations of CRH were significantly higher in LAA rats than in HAA rats before the shuttle-box tasks. The CRH levels in HAA rats were significantly increased after the shuttle-box tasks on Day 3 as compared with before the tasks, whereas the CRH levels in LAA rats were significantly decreased after shuttle-box tasks on Day 3 compared with before the tasks.

**CRH and AVP levels in the median eminence (Fig. 5)**

Main effects of strain and time were found for CRH in the median eminence, but no interaction effect of strain \( \times \) time was observed (strain \( \times \) time, \( F_{(1, 54)}=2.18, P=0.12 \); strain, \( F_{(1, 54)}=5.21, P=0.03 \); time, \( F_{(1, 54)}=7.64, P=0.001 \)). Furthermore, significant interaction of strain \( \times \) time was observed for AVP in the median eminence (strain \( \times \) time, \( F_{(1, 22)}=3.84, P=0.04 \)). The concentrations of CRH and AVP were significantly lower in LAA rats than in HAA rats before the shuttle-box tasks.
Hormone levels in pituitary glands (Fig. 6)

Main effects of strain and time were found for ACTH, but no interaction effect of strain × time was observed (strain × time, $F_{(1,54)}=2.50$, $P=0.09$; strain, $F_{(1,54)}=10.92$, $P=0.002$; time, $F_{(1,54)}=4.62$, $P=0.01$) in the pituitary gland. The concentrations of ACTH were significantly increased after shuttle-box tasks on Day 3 as compared with before the tasks in HAA rats, whereas there were no significant differences in ACTH levels between before and after shuttle-box tasks in LAA rats (a). The concentrations of ACTH were significantly lower in LAA rats than in HAA rats after shuttle-box tasks on the Day 3. Main effects of strain were observed for prolactin, but no interaction effect of time or strain × time was found (strain × time, $F_{(1,54)}=2.43$, $P=0.10$; strain, $F_{(1,54)}=8.87$, $P=0.004$; time, $F_{(1,54)}=1.51$, $P=0.23$) in the pituitary gland. The pituitary concentrations of prolactin were significantly higher in LAA rats than in HAA rats before shuttle-box tasks (b). The prolactin levels in LAA rats tended to be lower after shuttle-box tasks than before the tasks.

Discussion

The present study clearly demonstrated that HAA rats and LAA rats have differences in the pattern of hypothalamic hormonal secretion and circulating hormone levels during early avoidance acquisition (Fig. 7). Synthesized CRH and AVP in parvocellular neurons of the PVN coordinate HPA system activity and project to the external layer of the median eminence, where CRH and AVP are released into the portal blood [1]. The CRH and AVP are the primary hypothalamic and hypophysiotropic factors regulating basal and stress-induced release of pituitary ACTH and mediating the stress response by activation of the HPA axis [29, 34]. Before shuttle-box tasks, the concentrations of CRH in the PVN were significantly higher in LAA rats than in HAA rats (Fig. 4). The CRH levels in HAA rats were significantly increased after shuttle-box tasks on Day 3 before the tasks, whereas the CRH levels in LAA rats significantly decreased after shuttle-box tasks on Day 3 compared with before...
the tasks. The concentrations of CRH and AVP in the median eminence were significantly lower in LAA rats than in HAA rats before shuttle-box tasks (Fig. 5). In HAA rats, the CRH and AVP levels in the median eminence decreased after shuttle-box tasks, whereas there were no significant differences in the levels between before and after shuttle-box tasks in LAA rats (Fig. 5). Cell bodies of CRH neurons are primarily localized in the PVN. CRH neurons project to the median eminence and release CRH to the portal vessel. These results suggest that CRH in LAA rats is the sufficiently synthesized in the PVN; however, the secretions of CRH from the median eminence were low.

The plasma concentrations of ACTH were significantly higher in HAA rats than in LAA rats after shuttle-box tasks (Fig. 2). ACTH improves the deficient acquisition of shuttle-box avoidance behavior of hypophysectomized rats, delay extinction of shuttle-box avoidance behavior, and pole-jumping avoidance behavior, and facilitates passive avoidance behavior of intact rats [28]. Therefore, the high avoidance rate of shuttle-box tasks in HAA rats may be due to the high response of CRH-ACTH.

On the other hand, the adrenal response of corticosterone release to ACTH secretion after shuttle-box tasks was higher in LAA rats than in HAA rats on Day 3 (Fig. 2), as found in the previous study of restraint stress [3, 4]. These results indicate that the response of CRH-ACTH is higher in HAA rats than in LAA rats, whereas the response of corticosterone is lower in HAA rats than in LAA rats. Stress-evoked increases in the plasma corticosterone are correlated with impaired memory in a hippocampal-dependent task [42], and mineralocorticoid receptors are indispensable for nongenomic modulation of hippocampal glutamate transmission by corticosterone [26]. The high response of corticosterone to stress or the differential expression levels of central mineralocorticoid receptors may be a reason the low avoidance rates of shuttle-box tasks in LAA rats.

Prolactin is an antistress hormone [14–16]. Formation of gastric ulcers is observed more frequently in HAA rats than in LAA rats 7 h after restraint stress in water, whereas the plasma prolactin levels and numbers of long-form prolactin receptors in the PVN are significantly increased in LAA rats than in HAA rats [3, 4]. In agreement with the previous studies, the secretion of prolactin was higher in LAA rats than in HAA rats in the present study (Figs. 2 and 6). Thus, LAA rats may be resistant to stress compared with HAA rat.

It is also known that Roman and Syracuse rats, which were selectively bred for differential shuttle-box acquisition, originated from the Wistar and Long-Evans strains, respectively [6, 9]. In Roman and Syracuse rats, the response of the hypothalamo-pituitary-adrenal axis is lower in high-avoidance rats than in low-avoidance rats during stress [5, 10, 12, 20, 41]. The Roman and Syracuse rat strains differ in terms of anxiety, and this is one of the reasons for different results for avoidance behavior [11, 43]. The low avoidance rats of Roman lines display mostly freezing responses in the shuttle-box. However, Hatano LAA rats do not display freezing, anxiety behavior and depression behavior [32, 33]. Therefore, Hatano rats have characteristics that differ from those of Roman and Syracuse rats.

In summary, the present study clearly demonstrated strain differences in the characteristics of brain hormone responses between the two Hatano lines during early avoidance acquisition. The response of CRH-ACTH was higher in HAA rats than in LAA rats. This phenotype may be one of the important reasons for high avoidance rates of shuttle-box tasks in HAA rats. These two Hatano rats may be useful animals for study of the effects of stress-induced hormones on memory and retention regardless of anxiety and depression.

Acknowledgments

We thank Dr. Atsushi Tohei for advice on CRH measurement. This work was supported by a Grant-in-Aid for Scientific Research (B-1831044; Japan and Thailand Joint Research) from the Japan Society for the Promotion of Science.
References


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