Recurrent hypoglycemia alters hypothalamic expression of the regulatory proteins FosB and synaptophysin

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Al-Noori S, Sanders NM, Taborsky GJ Jr, Wilkinson CW, Zavosh A, West C, Sanders CM, Figlewicz DP. Recurrent hypoglycemia alters hypothalamic expression of the regulatory proteins FosB and synaptophysin. Am J Physiol Regul Integr Comp Physiol 295: R1446–R1454, 2008. First published August 27, 2008; doi:10.1152/ajpregu.90511.2008.—A limiting factor to the clinical management of diabetes is iatrogenic hypoglycemia. With multiple hypoglycemic episodes, the collective neuroendocrine response that restores euglycemia is impaired. In our animal model of recurrent hypoglycemia (RH), neuroendocrine deficits are accompanied by a decrease in medial hypothalamic activation. Here we tested the hypothesis that the medial hypothalamus may exhibit unique changes in the expression of regulatory proteins in response to RH. We report that expression of the immediate early gene FosB is increased in medial hypothalamic nuclei, anterior hypothalamic nuclei, and posterior paraventricular nucleus of the thalamus (THPVN) of the thalamus following RH. We identified the hypothalamic PVN, a key autonomic output site, among the regions expressing FosB. To identify the subtype(s) of neuronal populations that express FosB, we screened candidate neuropeptides of the PVN for coexpression using dual fluorescence immunohistochemistry. Among the neuropeptides analyzed [including oxytocin, vasopressin, thyrotropin-releasing hormone, and corticotropin-releasing factor (CRF)], FosB was only identified in CRF-positive neurons. Inhibitory γ-aminobutyric acid-positive processes appear to impinge on these FosB-expressing neurons. Finally, we observed a significant decrease in the presynaptic marker synaptophysin within the PVN of RH-treated vs. saline-treated rats, suggesting that rapid alterations of synaptic morphology may occur in association with RH. Collectively, these data suggest that RH stress triggers cellular changes that support synaptic plasticity, in specific neuroanatomical sites, which may contribute to the development of hypoglycemia-associated autonomic failure.

FosB; recurrent hypoglycemia; paraventricular nucleus; corticotropin-releasing factor; synaptophysin

The neuroendocrine counterregulatory response (CRR) that occurs to correct a single hypoglycemic (SH) episode becomes impaired when multiple episodes occur closely in time (recurrent hypoglycemia, RH), a situation often experienced by individuals with diabetes (18, 19). Although iatrogenic hypoglycemia represents a limitation to the clinical management of diabetes (49), the mechanisms underlying the failed neuroendocrine response are not completely understood.

The immediate early gene (IEG) and transcription factor, c-Fos, has served as a marker of neuronal activity in numerous in vivo and in vitro paradigms (36) and has been utilized to document activation of the central nervous system (CNS) as a result of SH in several laboratories (3, 23, 46). In a previous study, we used c-Fos to compare relative activation in the forebrain of rats receiving one or three bouts of hypoglycemia (23). That study revealed that three medial hypothalamic areas [the arcuate nucleus (ARC), the paraventricular nucleus (PVN), and the dorsomedial hypothalamus (DMH)] had decreased c-Fos expression in rats that had experienced three, vs. one, bout(s) of hypoglycemia. Furthermore, we determined that the PVN and DMH play key roles in mediating the hypothalamic-pituitary-adrenal axis (PVN and DMH) and sympathoadrenal (PVN) responses to hypoglycemia (24, 25). Accordingly, impaired activation of these medial hypothalamic areas would be expected to contribute substantially to the impaired CRR to RH. However, whether the decreased medial hypothalamic activation is the result of impaired afferent input [e.g., decreased input from glucensing neurons (40, 53)], impaired excitability of neurons within the PVN and DMH, initiation of intracellular processes that decrease subsequent neuronal excitability, or a combination of these is still unknown.

Although c-Fos, because of its rapid activation and deactivation kinetics, functions as a marker for the acute activation induced by a single bout of hypoglycemia, absence of c-Fos expression cannot be used to identify the phenotype of neurons within the PVN and DMH that are impacted by RH. Within the PVN, such neurons might include preautonomic neurons or neuroendocrine cells expressing oxytocin (OT), vasopressin (VP), thyrotropin-releasing hormone (TRH), or corticotropin-releasing factor (CRF) that are implicated in stress responses and regulation of energy homeostasis (5, 6, 26, 35, 38, 42, 51), making them candidates for evaluation of their role in the response to hypoglycemia. In this study, we took advantage of the differential expression profile of another member of the “Fos” family of transcription factors (14, 15, 32) to begin to examine the delayed or longer-acting neuronal alterations that we hypothesize are associated with RH. Although the role of FosB isoforms has been studied most extensively in models of drug addiction, FosB has been implicated in local CNS responses to the pathophysiology of seizure (44) and Parkinson’s disease (2, 13). As studied in the striatum by Nestler and colleagues (14, 15, 32), in the hours subsequent to the transient activation/deactivation time frame for c-Fos, alternative iso-
forms of Fos become expressed, notably FosB and delta FosB, which have longer half-lives, estimated to be in the range of days in vivo. This stability contributes to the potential accumulation of these isoforms following repeated stimulation and may serve to identify neurons that undergo delayed and longer-acting changes. These more stable IEGs can also control transcription of different sets of neuron-regulatory proteins, compared with c-Fos. Thus, with RH, transcription of alternative regulatory proteins that are not affected by SH might occur in the medial hypothalamus. To begin to evaluate this hypothesis, we measured the expression of FosB in the forebrain of rats following one or three bouts of hypoglycemia (or three control infusions). In addition to confirming that the medial hypothalamus is one of the brain regions that has increased FosB expression in association with RH, we used dual immunofluorescence to phenotype affected neurons in the PVN.

Various instances of neural plasticity, both behavioral and functional, are characterized by not only modified responses to repeated exposures of the same stimulus but changes in expression of key functional proteins in the neurons that are targeted by the stimulus. As such, we would predict that, in our model of RH, where repetitive exposure to the stress of hypoglycemia is associated with impaired neuroendocrine responsivity, there are underlying cellular mechanisms of neural adaptation. Thus transcription factors such as FosB could serve not only to simply identify the brain regions and neuronal phenotypes involved in mediating the impaired CRR to RH as discussed above but could represent a molecular step whereby persistent changes in gene expression can be induced and maintained. The functional implication of this for RH would be that neuronal activation is not merely decreased, but that neurons would essentially be reprogrammed.

As one additional index of change of function (as opposed to a simple decrease of activation) within the PVN, we evaluated expression of the synaptic marker synaptophysin (SYN) (41, 52, 61). Changes of SYN expression have been associated with the onset of plasticity in response to a variety of pathophysiological or adaptive changes (39) and challenges in the CNS, including hypoglycemia (52), although in that study the hypothalamus and individual brain regions were not evaluated. There are examples of early onset decreases of SYN and later-onset increases of SYN, which may reflect synaptic remodeling (33). We hypothesized that SYN expression would be altered in our model of acute RH.

**EXPERIMENTAL PROCEDURES**

**Animals**

Male Wistar rats (Simonsen, Gilroy, CA), 350–400 g, were maintained on ad libitum chow and water and a 12:12-h light-dark cycle (6:00 A.M. lights on; 6:00 P.M. lights off), and were studied during the light portion of the cycle. The study protocol was approved by the Institutional Animal Care and Use Committee of the Veterans Affairs Puget Sound Health Care System. After acclimation in the animal facility, rats were implanted with chronic venous catheters (jugular and submaxillary) according to our published methodology (23) to provide venous access for infusion and collection of blood samples (see Experimental protocol below). Rats recovered from the surgery and demonstrated a positive trajectory of weight gain before undergoing the experimental procedure.

**Experimental protocol.** We utilized our published methodology (23) for inducing one or three bouts of hypoglycemia in a 2-day procedure. Rats were studied in the conscious, non-food-deprived, unstressed condition, with blood samples being collected remotely and volume replaced remotely with whole blood obtained from donor rats. Before the experiment, rats were brought into the experiment room and placed in acrylic testing chambers with clean bedding, for habituation to the testing conditions. On experimental days, rats were placed in acrylic chambers, and intravenous lines were hooked up. Rats were settled for ~90 min before drawing a *time 0* baseline sample, with subsequent infusion of insulin (0.25 U·100 g−1·h−1), or saline vehicle, at a rate of 1.146 ml/h using a microinfusion syringe pump (KD Scientific, Holliston, MA). On day 1, rats received a 2-h insulin (or saline) infusion in the late A.M. This was followed by a 60-min interval in which they were given chow and allowed to feed; we have determined that this feeding interval is sufficient to rescue plasma glucose levels back to within normoglycemia (80–85 mg/dl) by the start of the second infusion. A second 2-h insulin or saline infusion was then carried out, and rats were returned to home cages where they had ad libitum access to chow overnight. On day 2, at ~9:00 A.M., rats were returned to their experimental chambers and, after baseline blood (*time 0*) collection, were subjected to either a third infusion of insulin or saline. Blood samples were then collected at 90 and/or 120 min thereafter for the measurement of glucose and the neuroendocrine CRR. Plasma samples obtained were stored at ~80°C until assayed. Blood for the catecholamine assays was collected on EGTA-glutathione (2.3:1.5 mg/ml; Sigma). Tubes for glucagon assays contained 50 μl of 1 M benzamidine (Sigma) and 1 unit heparin. Blood for glucose, adrenocorticotropic hormone (ACTH), and corticosterone assays was collected on EDTA and aprotinin (1.7 tissue inhibitor unit; Sigma). The assays have been described previously. Briefly, a radioenzymatic method as described in Evans et al. (23) was used for determination of plasma epinephrine and norepinephrine. Plasma glucose was measured using the Beckman Glucose Analyzer. Glucagon was assayed by the Linco glucagon RIA kit (Linco Research, St. Charles, MO). Measurements of ACTH were made using the Nichols Institute Diagnostics immunoradiometric assay kit (Nichols Institute Diagnostics, San Juan Capistrano, CA). A radioimmunoassay procedure was used for plasma corticosterone measurement as described in van Dijk et al. (60).

Following the last infusion on day 2, rats were deeply anesthetized with either intraperitoneal pentobarbital sodium injection (50 mg·ml−1·kg−1) or isoflurane inhalation and perfused with 0.9% NaCl followed with 0.9% buffered cold 4% paraformaldehyde solution. One group of rats was killed 90 min following a single infusion of saline or insulin (SH). This corresponds to the timing of our previous determination of c-fos in response to a single bout of hypoglycemia (23). Note that essentially no FosB expression was observed following a single infusion of saline and, therefore, no quantitation was carried out for this group. The remaining groups were killed 90 min after the third infusion (saline or RH) on day 2. This time point should reflect an accumulation of FosB isoforms, as well as the c-fos response to a third bout of hypoglycemia. Brains were removed and postfixated overnight in the same fixative and subsequently placed in 30% sucrose-PBS solution. Brains were sectioned on a cryostat (Leica CM 3050S cryostat) for immunohistochemistry.

**Immunohistochemistry.** Atlas-matched 40-μm sections were evaluated for comparison between insulin- and saline-treated rats, based upon Paxinos and Watson (47). Initial screen of forebrain areas revealed virtually no detectable FosB in most brain regions; therefore, we focused upon a limited number of areas for semiquantitation and for neuronal phenotyping (atlas figures 24–31, which include hypothalamus [ventromedial hypothalamic nucleus (VMH)], −2.12 to −3.14 mm; DMH, −2.56 to −3.14; PVN, −1.8 to −2.8; anterior...
hypothalamic area, −2.12 to −2.3]; posterior PVN of the thalamus (THPV; −3.14 to −3.30), a region that is implicated in the habituation of the neuroendocrine response to other stressors (9), and is also activated in response to hypoglycemia (58); and anterior PVN of the thalamus (THPVA; −1.4 to −2.12) and central amygdala (−2.12 to −2.3) as control regions implicated in stress circuitry but not significantly responsive to hypoglycemia in our model.

FosB immunocytochemistry and quantitation. The laboratory’s established methodology for c-Fos quantitative immunocytochemistry (23) was adapted for quantitation of FosB. Sections were washed in PBS for 45 min; goat serum-Triton X-100 buffer (3% goat serum-0.3% Triton X-100 in PBS) for 30 min; gelatin blocking buffer (0.5% gelatin in goat serum-Triton X-100 buffer) for 60 min; and goat serum-Triton X-100 buffer for 30 min. Free-floating sections were then incubated with primary antibody (1:2,000) in goat serum-Triton X-100 buffer for 40–48 h at 4°C. Following incubation with primary antibody, tissue was washed at room temperature in PBS for 90 min, transferred to secondary antibody solution (5 μl secondary antibody/ml goat serum-Triton X-100 buffer) for 60 min, and then washed for 60 min in PBS. Sections were incubated with avidin-biotin solution at room temperature for 60 min, transferred to dianaminobenzidine solution for hand mixing up to 2 min maximum, and washed in PBS for 30 min. Sections were mounted onto glass slides and coverslipped using Permount mounting medium. For quantitation (at ×40 magnification), atlas-matched sections from control or hypoglycemic brains were selected. ImagePro Plus software (Media Cybernetic) was utilized to capture an image of the desired area. An area was delineated for counting, and threshold for positive cell counts was established. The identical area and background (threshold) were used for sections from control and hypoglycemic brains to prevent between-session changes in background setting.

Western blotting. Brains were removed and snap-frozen in methy- lbutane. Thick (280 μm) cryosections of the tissue were prepared, and punches of the specific brain regions (see above for atlas coordinates) were taken and used to make homogenates using RIPA lysis buffer (Santa Cruz Kit) according to standard techniques (manufacturer’s specifications). Protein estimates were performed using a BCA assay kit (Pierce, Rockford, IL). Samples were run on 10% Tris–HCl precast Ready Gels (Bio-Rad, Hercules, CA) and transferred to polyvinylidene difluoride membranes (Bio-Rad) for 2 h at 100 volts. Blots were blocked in Superblock Blocking Buffer (Pierce) made up in PBS, washed three times in Tris-buffered saline (TBS) containing 0.1% Tween 20 (Sigma, St. Louis, MO) (TBS-T), and incubated in primary antibody solution overnight at 4°C. Blots were washed again three times in TBS-T and incubated for 1–2 h at room temperature in secondary antibody solution. After multiple washes in TBS-T, blots were subjected to detection using the Supersignal West Femto ECL kit (Pierce) and developed on film (x-omat blueXB-1; Kodak). Digital images of blots were obtained using an Epson 836XL high-resolution document scanner, and the bands were quantitated using Bio-Rad’s Quantity One analysis software.

Materials

Humulin R regular insulin, human USP rDNA origin ( Elli Lilly, Indianapolis, IN), 0.25 U/100 g made up in saline, was used to induce hypoglycemia. Primary antibodies used included rabbit anti-FosB (1:500) (sc-48) (Santa Cruz Biotechnology, Santa Cruz, CA), mouse anti-FosB (1:500) (Abcam, Cambridge, MA), mouse anti-GAD (1:1,000) (Chemicon, Temecula, CA), mouse anti-OT (1:1,000) and mouse anti-VP (1:500) (generous gifts from Dr. Ann-Judith Silverman, Columbia University), rabbit anti-TRH (1:500) (#PBL rCRF; generous gift from Dr. Wylie Vale, Salk Institute), mouse anti-SYN (1:1,000) (Chemicon), and rabbit anti-pre-pro-TRH (1:10,000) (generous gift from Dr. Eva Redei, Northwestern University). Secondary antibodies used included Cy3-conjugated goat anti-rabbit or anti-mouse (Jackson Immunoresearch, West Grove, PA), Alexa fluor 488 goat anti-mouse or anti-rabbit IgG ( Molecular Probes, Eugene, OR), and biotinylated anti-rabbit IgG (Vector Laboratories). All secondary antibodies were diluted at 1:500.

RESULTS

Neuroendocrine Responses to SH and RH

Table 1 shows the metabolic and neuroendocrine response to three bouts of saline (control), SH, or RH in rats for which

<table>
<thead>
<tr>
<th>Glucose, mg/dl</th>
<th>Glucagon, pg/ml</th>
<th>Epi, pg/ml</th>
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<tr>
<td><strong>Saline (n = 9)</strong></td>
<td></td>
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<tr>
<td>Time 0</td>
<td>107±2</td>
<td>99±15</td>
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<tr>
<td>Time 120 min</td>
<td>101±1</td>
<td>101±3</td>
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<tr>
<td><strong>SH (n = 9)</strong></td>
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<tr>
<td>Time 0</td>
<td>105±2</td>
<td>131±7</td>
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<tr>
<td>Time 120 min</td>
<td>31±2</td>
<td>574±94</td>
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<tr>
<td><strong>RH (n = 9)</strong></td>
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<tr>
<td>Time 0</td>
<td>106±3</td>
<td>78±10</td>
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<tr>
<td>Time 120 min</td>
<td>31±2</td>
<td>211±24</td>
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Values are means ± SE; n, no. of rats. Epi, epinephrine; SH, single hypoglycemic; RH, recurrent hypoglycemia. Data were obtained from rats studied for FosB quantitation (Fig. 1).
FosB was quantitated (Fig. 1). As expected, epinephrine and glucagon levels were elevated in hypoglycemic rats compared with the values of saline-infused control rats. The RH responses are comparable to our published data (23) and lower than those after SH. Table 2 shows plasma glucose and neuroendocrine data, in response to RH or three infusions of saline, in rats studied for the neural phenotype of Fos-B-expressing cells within the PVN (Figs. 2–5). Glucagon and epinephrine data are comparable to those reported in Table 1, consistent with a blunted response to RH. ACTH responses to RH are substantially lower than those we have reported previously as a response to SH (211±1100622 pg/ml) and perhaps reflect the impaired CRR to RH, along with impaired glucagon and epinephrine. Corticosterone responses, as we have previously reported (23), are essentially normal after RH.

**C-Fos and FosB Responses to SH and RH**

Figure 1A demonstrates c-Fos increases with SH compared with RH (Fig. 1A, axis on left). As we have reported previously, c-Fos is decreased in the PVN (P<0.015) and DMH (P<0.04) in response to RH vs. SH. There is no detectable c-Fos in the VMH in response to either SH or RH. In contrast, FosB (Fig. 1A, axis on right) levels are increased following RH.

**Table 2. Plasma glucose, glucagon, Epi, ACTH, and CORT responses to RH or saline (control) infusions**

<table>
<thead>
<tr>
<th></th>
<th>Glucose, mg/dl</th>
<th>Glucagon, pg/ml</th>
<th>Epi, pg/ml</th>
<th>ACTH, pg/ml</th>
<th>CORT, μg/dl</th>
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<tr>
<td><strong>Saline (n = 7–8)</strong></td>
<td></td>
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<td></td>
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<tr>
<td>Time 0</td>
<td>107±7±1100557</td>
<td>87±1100519</td>
<td>19±110053</td>
<td>4.9±1.5</td>
<td></td>
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<tr>
<td>Time 90 min</td>
<td>119±21100574</td>
<td>67±1100525</td>
<td>17±3</td>
<td>5.5±2.0</td>
<td></td>
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<tr>
<td>Time 120 min</td>
<td>118±31100578</td>
<td>76±1100521</td>
<td>19±3</td>
<td>9.6±1.6</td>
<td></td>
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<tr>
<td><strong>Insulin (n = 7–8)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Time 0</td>
<td>114±51100574</td>
<td>90±1100518</td>
<td>17±2</td>
<td>5.4±1.1</td>
<td></td>
</tr>
<tr>
<td>Time 90 min</td>
<td>46±51100574</td>
<td>1,658±1100555</td>
<td>93±13</td>
<td>26.9±2.1</td>
<td></td>
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<tr>
<td>Time 120 min</td>
<td>34±41100574</td>
<td>2,333±11005479</td>
<td>88±10</td>
<td>32.4±2.5</td>
<td></td>
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Values are means ± SE; n, no. of rats. ACTH, adrenocorticotropic hormone; CORT, corticosterone. Data were obtained from rats studied for immunocytochemical phenotyping (Figs. 2–5).

Fig. 1. Recurrent hypoglycemia (RH) induces FosB expression. A: quantitative comparison of c-Fos (axes on left) and FosB (axes on right) cell counts following single hypoglycemic (SH) and RH in the paraventricular nucleus (PVN), ventromedial hypothalamic nucleus (VMH), and dorsomedial hypothalamic nucleus (DMH). B: FosB immunopositive neuron counts following three infusions of saline or insulin. Statistically significant increases in the number of FosB-positive cells in the RH-treated group compared with the saline control group were observed in all of the hypothalamic areas indicated and the posterior PVN of the thalamus (PVP). FosB expression was not increased in the other brain areas, as evaluated by either one- or two-tailed t-tests (hypothesis: increased FosB expression in RH- vs. saline-treated rats, one-tailed t-test; change in either direction of FosB expression in RH- vs. saline-treated rats, two-tailed t-test).

C: immunofluorescence images taken at mid-PVN levels illustrating the expression of FosB in the PVN of a RH-treated rat (right) compared with a saline-treated rat (left). The dotted white line demarcates the ventral perimeter of the PVN, whereas “V” is placed over the region of the ventricle. D: immunofluorescence image of the caudal PVN; the dotted white line demarcates the lateral parvocellular subdivision.

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Localization of FosB in PVN

Based on the FosB quantitation described above, we focused on the medial hypothalamus, specifically the PVN as a major autonomic center, to identify the phenotype(s) of neurons that express FosB following hypoglycemia, as vulnerable neuronal subtypes that might be implicated in the impaired CRR. To do this, we assayed coexpression of FosB with the candidate neuropeptides OT, VP, CRF, and TRH by means of double-label immunofluorescence in saline-treated and RH-treated rats.

Figure 2 shows representative images of sections from RH-treated animals. As in Fig. 1C, RH-induced FosB expression is observed (Fig. 2A). CRF expression in the hypothalamus was found to be limited to the parvocellular PVN region, and immunostaining of both cell bodies and fibers was observed (Fig. 2B). A proportion of FosB-positive neurons were found to coexpress CRF, as demonstrated by antibody colocalization (Fig. 2, C and D, at higher magnification). This subpopulation of neurons exhibited a central nuclear region expressing FosB surrounded by a cytoplasmic limbus of CRF immunoreaction product. Quantitation by means of cell counts revealed that 30% of FosB-positive cells express CRF. It is possible that this may be an underestimate of the FosB expression in CRF neurons, since rats were not colchicine pretreated to limit CRF transport out of the cell body (hence, lower cell body immunostaining).

Figure 3 illustrates FosB (Fig. 3A) and TRH (Fig. 3B) immunoeexpression. The TRH labeling in the somata was seen to be predominantly cytoplasmic, whereas the fiber staining had a highly punctate appearance. In contrast to CRF, and within the limits of detection of TRH immunoreactivity, TRH-positive cell bodies did not appear to coexpress FosB, and clearly both immunoreactive products appeared to label separate neuronal populations (Fig. 3D). The lower-magnification image (Fig. 3C) confirms the presence of TRH-positive cells in the parvocellular region of the PVN, as expected.

Analysis of VP expression revealed confinement of VP-expressing cell bodies to the magnocellular PVN region of the hypothalamus. Densely packed somata along with fiber staining were observed (Fig. 4B). Examination of OT expression in the hypothalamus again revealed a confinement to the magnocellular PVN region with clearly defined subpopulations of neurons displaying immunoreaction product in both cell bodies and fiber terminals (Fig. 4E). These OT-positive fibers displayed puncta-like varicosities along their lengths. However, double-label immunofluorescence revealed no colocalization of either VP or OT immunopositivity with FosB (Fig. 4, C and F, respectively), indicating that both of these neuropeptides, while robustly expressed in the hypothalamic PVN, are not the neuronal subpopulations in which FosB is being expressed.

FosB expression within the parvocellular PVN compartments was observed at the mid-PVN levels in the medial and ventral parvocellular subdivisions and at the caudal PVN levels in the medial and lateral parvocellular subdivisions [Fig. 1, C and D; designations of the PVN subdivisions are based on the work of Lechan and Fekete (38)]. This finding is significant in that neurons in these parvocellular regions have been reported to send projections to preganglionic targets in the brain stem and spinal cord (12, 48, 54). The involvement of putative preautonomic projecting neurons is consistent with the impaired glucagon and epinephrine responses we measure in response to RH.

Finally, we evaluated expression of glutamic acid decarboxylase (GAD), the synthetic enzyme for the inhibitory neurotransmitter γ-aminobutyric acid (GABA). GABAergic innervation of the PVN is extensive and may be both local and extrinsic from regions such as the DMH (30, 31). Kovacs et al. (37) recently reported that, in the parvocellular PVN, 79% of GABAergic boutons terminate on CRF neurons and that these comprise 36% of the total synapses on CRF neurons.
increase in either GABA synthesis or signaling could be one mechanism contributing to the decreased PVN activation with RH. Immunohistochemistry revealed a high intensity of GAD immunoreactivity in the RH group (Fig. 5A) with very bright and widespread GAD-positive puncta, presumably reflecting synaptic terminals. A dense plexus of GAD-positive fiber staining was observed in the area of the PVN and the entire periventricular area, whereas GAD-positive somata were only observed in the area immediately adjacent to the PVN or the “peri-PVN” (see Fig. 5A). Double-label immunofluorescence was performed to examine the distribution and expression of GAD (Fig. 5C) relative to FosB (Fig. 5B). The GAD-positive staining did not appear to colocalize to the same cell bodies as FosB, but the GAD-positive fibers did appear to be impinging on the FosB-positive cells (Fig. 5D) and perhaps making synaptic contacts.

Measurement of Medial Hypothalamic SYN After RH

Having identified the CRF neuron as the predominant neuron within the PVN to express FosB, we then turned to examine expression of the putative presynaptic protein SYN as a marker of synaptic changes that might underlie the impaired CRR to RH. SYN expression can be altered (increased or decreased) in acute circumstances of stimulus-induced neuroplasticity (52, 61) (see Discussion). Expression of SYN in medial hypothalamic regions was analyzed by Western Blot (Fig. 6A) to investigate whether SYN changed in association with RH. Although no statistically significant changes were observed in the ARC or DMH/VMH, SYN expression was decreased significantly (P < 0.017) in the PVN with RH, as demonstrated by a decrease in the 37-kDa size band representing SYN in the PVN of RH rats compared with the PVN of saline controls (Fig. 6B).

DISCUSSION

In the present study, we have begun to address the hypothesis that RH leads to a change in transcription of regulatory proteins that may decrease the responsiveness of key CNS regions that mediate the neuroendocrine response. We evaluated one aspect of regulatory protein transcription: changes in the expression of the IEG FosB following RH compared with SH or saline-infused controls. Here we present several novel findings. Relative to saline-infused controls, FosB expression is increased in the PVN, VMH, and DMH hypothalamic subnuclei of rats that experienced multiple bouts of hypoglycemia. We also observed significant FosB expression in the THPV, a limbic structure that modulates the CRR to hypoglycemia, as we and others have recently documented (1, 4). Whereas studies in models of seizure and Parkinson’s disease (2, 13, 44) implicate a role for FosB expression in the amygdala and striatum, our finding represents the first report, to our knowledge, of expression of FosB isoforms in the PVN.
and in response to a metabolic stressor. Collectively, these studies suggest that FosB isoforms may play a role in the development of several neuropathologies, a commonality being substantial insult to the neuron with potential depletion of energy reserves.

Our observation of increased FosB expression in the PVN, a major efferent outflow site that receives multiple inputs from the entire neuraxis, then provided the rationale for examining FosB coexpression with specific neuropeptides, specifically, VP and OT, CRF, and TRH. Although robust expression of all of these neuropeptides in the PVN was observed as expected, our findings demonstrate that a significant proportion of FosB expression occurs within the hypothalamic PVN only in a subpopulation of neurons that express CRF but not VP, OT, or TRH. Furthermore, within the various subcompartments of the parvocellular PVN, we observed FosB expression to be included in ventral, medial, and lateral parvocellular compartments, regions that are reported to send preautonomic projections to the brain stem and spinal cord (12, 48, 54), thus potentially playing a major role in regulating the sympathoadrenal component of the CRR to hypoglycemia.

CRF is a neurotransmitter and hypothalamic neurohormone and functions as a major physiological regulator of the basal and stress-induced release of ACTH, β-endorphin, and other proopiomelanocortin-derived peptides from the pituitary (22, 59). Indeed, CRF appears to play a major role in both the activation and coordination of the stress response at the endocrine, behavioral, autonomic, and possibly even the immune levels (21). There is evidence that the repeated experience of certain stressors results in a process of adaptation such that subsequent exposure to the same stressor produces responses that become smaller and may even disappear. Consistent with this, Ma et al. (43) demonstrated desensitization and a loss in CRF mRNA response following repeated immobilization stress, concomitant with increased arginine VP expression in the parvocellular PVN. Somewhat surprisingly, we did not observe FosB coexpression with VP in our RH model. However, while not directly analogous to their recurrent stress model, the impaired neuroendocrine response to RH does include decreased ACTH release, implicating a change in the CRF-ACTH arm of the CNS response.

The ability of CRF neurons to have experience-specific (RH vs. SH) changes in the expression of a major transcription factor, FosB, is consistent with the reported modification of CRF output in a stressor-dependent manner (8) and, secondarily, with downstream actions of CRF in the CNS. In addition to mediating an acute response to stress, CRF has been shown to mediate long-term impacts of stress in several brain regions. For example, in the hippocampus, CRF signaling via the CRF1 receptor subtype has been reported to modulate dendritic length and branching (16, 55, 56). Thus it is conceivable that CRF-related plasticity may be one cellular mechanism that contributes to the impaired CRR observed in our model of RH.

We speculate that the decreased expression of the synaptic marker SYN specifically in the PVN reflects the initiation of structural changes in association with the impaired CRR response to RH. Functional plasticity in the brain is associated with structural remodeling of synapses (34) and with altered expression or regulation of specific synaptic proteins such as SYN (27). The decrease of SYN expression observed in the PVN supports the idea that initiation of synaptic reorganization there may accompany acute RH (and while we do not address the effect of SH on SYN expression, it is also possible that alterations in SYN expression can occur with SH) and is

Fig. 5. Glutamic acid decarboxylase (GAD) immunoreactivity, a marker of GABAergic transmission, does not colocalize with FosB in the PVN following RH. Immunofluorescence using α-GAD67 antibody revealed intense GAD (green) expression in the hypothalamus. A: GAD-positive staining in the PVN and peri-PVN. Note that the staining is almost exclusively of fibers within the PVN proper (outlined by dotted white line), whereas GAD-positive immunoreactive cell bodies are only observed in the area of the peri-PVN (white arrows). B: FosB-positive cells (red) in the PVN following RH. C: GAD immunoreactivity in the PVN and surrounding area in the same section. The neuropil in the immediate region of the PVN displayed intense GAD-positive fiber staining. Scale bar: 100 μm.

Fig. 6. Synaptophysin (SYN) is decreased in the PVN following RH. A: representative Western blot of SYN expression in the corresponding brain areas of interest (as identified in Fig. 1) from saline and RH-treated animals (n = 4/group). Top: blot probed with α-SYN antibody; bottom: blot probed with α-actin antibody as a control. Separator lines indicate where the image was cropped to include only the brain areas of interest. B: quantification of Western blot SYN protein expression. *A statistically significant (P < 0.017) decrease in SYN expression is observed in the PVN following RH. Protein levels are corrected to actin expression. ARC, arcuate nucleus of the hypothalamus; D/V, combined DMH/VMH.
consistent with reports of decreases in SYN immunoreactivity observed with other models of stress (20, 50). Brief anoxic-hypoglycemic episodes have been reported to affect the structural and functional components of synaptic networks at both the pre- and postsynaptic levels (34). Clinically, hypoglycemia “unawareness” and the blunted epinephrine component of the defective CRR are reversed by 2–3 wk of scrupulous avoidance of hypoglycemia (18). Thus the impaired neuroendocrine CRR is a reversible phenomenon, and it is reasonable to propose that processes of reorganization and plasticity, rather than permanent synaptic or neural destruction per se (45), underlie the impaired CRR to RH.

Local inhibitory GABAergic and local excitatory glutamatergic synaptic inputs to hypothalamic neurosecretory neurons have been reported (10, 11, 28, 29, 30). Both neuroanatomical and electrophysiological studies have confirmed the presence of inhibitory GABAergic inputs to PVN parvocellular neurons (11, 57). Furthermore, in vitro studies have demonstrated that intrinsic GABAergic influences in the PVN impart a tonic inhibitory tone on parvocellular CRF neurons (7, 17, 37). These local networks play an essential role in communicating and integrating extrahypothalamic inputs. We did not observe GAD-positive cell bodies to coexpress FosB. Consistent with other reports in the literature, we found that GAD-positive cell bodies were not located in the PVN proper but appeared to be confined to the immediate perimerid, i.e., the peri-PVN. We observed an intense GAD-positive fiber plexus innervating the PVN and impinging on FosB-expressing cells. Our observation that GAD-positive terminals appear to impinge on FosB-positive cells is suggestive of a potential role for the inhibitory networks in regulating neurons that are engaged in response to hypoglycemia. Within the context of the present study, a potential role for GABAergic circuits projecting to the PVN may include limiting the stress response or modulating it as a function of physiological status. Decreased neuronal responsiveness may serve a protective function in preventing the overactivation of specific circuitry or systems.

Taken together, our results suggest that the medial hypothalamic nuclei, particularly the PVN, may be targets for “reprogramming” in association with RH. We are currently exploring the functional significance of FosB expression on the neuroendocrine CRRs. Finally, our findings lay the groundwork for further studies aimed at the identification of specific proteins that decrease responsivity of PVN neurons, ultimately providing insight into the mechanism(s) underlying hypoglycemia-associated autonomic failure (HAAF).

Perspective

The impaired CRR to RH contributes to the syndrome of HAAF in diabetic individuals; it has been identified as a limiting problem in the use of intensive insulin therapy to minimize or prevent the chronic complications of diabetes. Identifying the CNS mechanisms that underlie the impaired CRR, which develops very acutely, is thus an important clinical goal. The current study extends our previous observations of the CNS efferent coordinator of neuroendocrine and autonomic responsivity, the PVN, as a target of RH. The finding of FosB, a conditionally expressed IEG that regulates transcription of a unique set of neuroregulatory proteins, in the PVN suggests that the PVN is not merely a “passive partner” in the CNS changes of RH. Rather, it implies that, within the PVN, adaptation to RH is an active process and may result in local changes of tone and responsivity to this metabolic stressor.

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