Variations in the Nature of Behavioral Experience Can Differentially Alter the Consequences of Developmental Exposures to Lead, Prenatal Stress, and the Combination

Deborah A. Cory-Slechta,1 Kian Merchant-Borna, Joshua L. Allen, Sue Liu, Douglas Weston, and Katherine Conrad

Department of Environmental Medicine, University of Rochester School of Medicine and Dentistry, Rochester, New York 14642

1To whom correspondence should be addressed at Department of Environmental Medicine, University of Rochester School of Medicine, Box EHSC, Rochester, NY 14642. Fax: (585) 256-2591. E-mail: deborah_cory-slechta@urmc.rochester.edu.

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Behavioral experience (BE) can critically influence later behavior and brain function, but the central nervous system (CNS) consequences of most developmental neurotoxicants are examined in the absence of any such context. We previously demonstrated marked differences in neurotransmitter changes produced by developmental lead (Pb) exposure ± prenatal stress (PS) depending upon whether or not rats had been given BE (Cory-Slechta, D. A., Virgolini, M. B., Rossi-George, A., Weston, D., and Thiruchelvam, M. (2009). The current study examined the hypothesis that the nature of the BE itself would be a critical determinant of outcome in mice that had been continually exposed to 0 or 100 ppm Pb acetate in drinking water alone or in combination with prenatal restraint stress. Half of the offspring in each of the four resulting groups/gender were exposed to positively reinforced (food-rewarded Fixed Interval schedule-controlled behavior) or negatively reinforced (inescapable forced swim) BE. Brain monoamines and amino acids differed significantly in relation to BE, even in control animals, as did the trajectory of effects of Pb ± PS, particularly in frontal cortex, hippocampus (both genders), and midbrain (males). In males, Pb ± PS-related changes in neurotransmitters correlated with behavioral performance. These findings suggest that CNS consequences of developmental toxiciants studied in the absence of a broader spectrum of BEs may not necessarily be predictive of human outcomes. Evaluating the role of specific BEs as a modulator of neurodevelopmental insults offers the opportunity to determine what specific BEs may ameliorate the associated impacts and can assist in establishing underlying neurobiological mechanisms.

Key Words: lead; prenatal stress; fixed interval; forced swim; neurotransmitters; behavioral experience.

Animal models typically evaluate the central nervous system (CNS) effects of developmental insults such as lead (Pb) exposure or prenatal stress (PS) in the absence of any context, even though the human environment is highly dynamic. All organisms inevitably encounter a variety of behavioral experiences (BEs), both positive and negative, over the life span. It is well established, both in human studies and animal models, that early BE can profoundly influence later brain function and behavior. Early educational interventions like Head Start, for example, can enhance later academic success. Such enhancements have included educational and social outcomes (Bierman et al., 2008) that are present even 20 years later (Palfrey et al., 2005; Reynolds et al., 2007). In contrast, early intense negative experiences, such as low perceived parental support, maternal distress, or parental verbal, physical, or emotional abuse, can produce long-standing adverse behavioral consequences. These have included impaired cognition, lower academic function, and behavior problems (Cheatham et al., 2010; Mills et al., 2011) that may, under some conditions, even be cumulative (Jaffee and Maikovich-Fong, 2011; Shonkoff et al., 2012). In contrast, negative experience that is predictable and controllable can be associated with later resiliency (Koolhaas et al., 2011).

In animal studies, positive BE such as “enrichment” (e.g., enriched housing and cognitive training) has been shown to mitigate CNS damage in models that include schizophrenia, Huntington’s disease, Parkinson’s disease, drug abuse liability, depression, and Alzheimer’s disease (Laviola et al., 2008; Nippak et al., 2007). In contrast, severe negative BE can produce sustained detrimental impacts on cognition and other behaviors. For example, male rats exposed to inescapable shock later showed both slower acquisition and reversal on a food-rewarded discrimination reversal task (Rosellini et al., 1982). A single period of immobilization (restraint) stress increased basolateral amygdala spine density in rats and was paralleled by the development of anxiety-like behavior in an elevated plus maze over a 10-day period post-restraint (Mitra et al., 2005). Ten days after rats were subjected to the single prolonged stress paradigm (restraint followed by forced swim [FS] followed by ether), significantly decreased glutamate and glutamine levels were observed in medial prefrontal cortex (Knox et al., 2010), a region critical to learning and executive functions.
Far more underappreciated, however, is the fact that BE can also significantly modify the consequences of developmental insults such as Pb and PS. Both human and animal studies already suggest differential modifications of Pb toxicity in response to different BE. IQ reductions in children with increasing blood Pb levels, for example, were significantly attenuated by higher socioeconomic status (SES; Bellinger et al., 1989). Such findings could suggest that positive BE can lead to a more “resilient” phenotype with improved outcome. This interpretation is supported by animal studies demonstrating that “environmental enrichment” conditions have the capacity to mitigate the effects of developmental Pb exposures (Guilarte et al., 2003; Schneider et al., 2001). An alternative or additional explanation of those findings, however, is that negative experiences associated with low SES enhance the neurotoxicity of Pb. Negative experience, such as embodied by permissive parenting (low parental involvement and stimulation) was reported to exacerbate the inverse association between blood Pb in children and scores on the McCarthy Scales of Children’s Abilities (Hubbs-Tait et al., 2009).

Our prior studies have shown marked differences in neurotransmitter levels and their alteration by Pb ± PS in animals that were behaviorally tested relative to nonbehaviorally tested littermates (Cory-Slechta et al., 2009), indicating the importance of BE as a determinant of the effects of Pb ± PS. Further, these studies demonstrated that the behavioral toxicity, neurotransmitter changes, and hypothalamic-pituitary-adrenal axis dysfunction associated with Pb±PS can be further enhanced, in an enduring capacity, by subsequent negative uncontrollable, unpredictable stressor exposure (OS). Collectively, such findings suggest that the cumulative “dose” of adversity may be important (e.g., developmental Pb and PS followed by OS) (Virgolini et al., 2008).

This study extends the prior studies in hypothesizing that the nature of the BE itself will be a critical determinant of outcome (Koolhaas et al., 2011; Maier and Watkins, 2010). Specifically, it postulates that different BEs, here “positively reinforced” (food-rewarded operant responding) versus “negatively reinforced” (exposure to inescapable FS) BEs, would differentially alter brain neurotransmitter systems in mice and also possibly modify the trajectory of effects of Pb, PS, and combined Pb + PS. Ultimately, such an understanding could provide a better understanding of mechanisms, particularly gender-dependent, of the trajectory of these developmental insults, as well as a basis for specific forms of behavioral enrichment that may mitigate adverse effects.

**MATERIALS AND METHODS**

**Experimental design.** Figure 1 depicts the overall experimental design. Offspring were generated with developmental exposures to either lead (Pb: 0 or 100 ppm), prenatal restraint stress (PS), or both, leading to 4 groups of 16 dams each: control (0-NS), prenatal stress (0-PS), lead (100-NS), and lead + prenatal stress (100-PS). Trunk blood was collected from a subset of dams immediately after weaning for measurement of blood Pb, corticosterone levels, and neurotransmitter levels. Blood collection during breeding and lactation was avoided to preclude any additional stressors that would confound interpretation of outcomes. Subsets of male and female offspring were sacrificed at 2.5 months of age for determinations of blood Pb and corticosterone.

At ∼7-10 months of age, male and female offspring from these groups were randomly assigned to either a positively reinforced BE (food-rewarded Fixed Interval [FI] schedule-controlled operant behavior) or to a negatively reinforced BE comprised of four sessions of inescapable FS testing, assuring single males and females/dam/group. The four FS tests were spaced to encompass the period of FI BE. Blood was collected after the second FS test from FS-tested offspring for determination of corticosterone. Four days after the completion of BE, trunk blood was collected for blood lead (PbB) and corticosterone determinations and brain extracted for measurement of brain neurotransmitters.

**Breeding and generation of offspring.** After a 1-week habituation period, 28-day-old female C57Bl6 mice (Jackson Laboratories) began exposure to Pb in drinking water. Two months later, to ensure adequate bone Pb levels consistent with human environmental Pb exposure, females were bred with males, and if pregnancy was indicated, were individually housed through weaning. One third of dams from each treatment group were weighed daily over the course of pregnancy. At birth, numbers of male and female offspring were counted; the offspring were weighed and any losses were monitored over the course of the study. Offspring were weaned at 24 days of age.

**Lead exposure.** Lead acetate (0 or 100 ppm) was dissolved in distilled deionized water and provided as the drinking solution to dams. Offspring were continued on the same Pb exposure concentration as the dam. The 100-ppm exposure level was chosen based on an initial pilot study comparing PbB at several different concentrations. Choice of a single Pb exposure concentration for use in this study was to focus on the most human relevant PbB levels, given that the extensive numbers of subjects, groups, and treatments made additional comparisons with higher PbB levels unfeasible.

**Prenatal stress.** Half of the dams in each Pb treatment group were exposed to immobilization restraint stress carried out on approximate gestational days 11–19, a paradigm typical of mouse prenatal restraint stress (Diz-Chaves et al., 2012; Miyagawa et al., 2011). Immobilization was carried out 3× per day for 30 min each time with a 2-h separation between immobilizations. Nontreated dams simply remained in home cages during this period.

**Behavioral experience.** BE was initiated at 7–10 months of age with offspring randomly assigned to either positively reinforced or negatively reinforced BE with no more than a single pup/gender/dam in each treatment group. FI schedule-controlled behavior was carried out in operant chambers containing three response levers and a feeder that delivered 20 mg food pellets. Lever press responding was first autoshaped using procedures previously developed in our laboratory (Cory-Slechta et al., 1985) to a criterion of the delivery of 100 reinforcers on a fixed ratio 1 schedule. A 60-s FI schedule of reinforcement was imposed in the next behavioral test session. On the FI schedule, the first lever press response after the 60-s interval elapsed resulted in food delivery and initiated the next 60-s interval. Sessions were 20 min in duration and carried out 5 days per week (M-F) for a total of 40 sessions. Behavioral measures included overall response rates, run rates, postreinforcement pause, and index of curvature as defined in our previous studies (Rossi-George et al., 2011). Inescapable FS tests were carried out on four occasions using procedures previously described (Porst et al., 1977). For each test, the mouse was placed in a 5×1 glass cylinder of 27°C ± 1°C water filled to a depth of 18 cm for a total of 5 min, and from which escape or use of the tail for balance was not possible. The minimum time between any two FS tests was 2 weeks. Behavior during the tests was video-recorded using a Kodak Zi8 video camera placed 0.5 m above the cylinder and subsequently scored by an investigator blinded to treatment conditions. Dependent measures included time spent immobile, number of immobile bouts, latency to first immobile bout, and time immobile per bout, with immobility defined as cessation of movement. After each test, mice were removed and returned to the home cage.

**Measurement of brain neurotransmitters.** Levels of monoamines as well as glutamate, glutamine, and gamma aminobutyric acid (GABA) were
measured in the right hemisphere of multiple brain regions (frontal cortex, striatum [combined nucleus accumbens and dorsal striatum], midbrain [combined ventral tegmental area and substantia nigra], hypothalamus and olfactory bulb). Levels of dopamine (DA), dihydroxyphenylacetic acid (DOPAC), homovanillic acid (HVA), norepinephrine (NE), serotonin (5-HT), and 5-hydroxyindoleacetic acid (5-HIAA) were analyzed using high-performance liquid chromatography (HPLC) with electrochemical detection as previously detailed (Cory-Slechta et al., 2004, 2009, 2010; Virgolini et al., 2008). Concentrations of neurotransmitters were expressed as ng/mg protein. DA turnover (DA TO) was calculated as the DOPAC/DA ratio.

Levels of glutamate, glutamine, and GABA were assayed in hippocampus and frontal cortex using a modified version of a previously described method (de Freitas Silva et al., 2009). Standards were prepared in 0.1N perchloric acid at concentrations of 6 µg/ml glutamine, 12 µg/ml glutamate, and 1.2 µg/ml GABA. Precolumn derivatization was performed by mixing 100 µl sample or standard solution, 20 µl methanolic OPA (5 mg/ml), 75 µl borate buffer (pH 9.9), and 5 µl MPA. The standard/sample solution was vortexed and injected onto the chromatographic column at a volume of 10 µl after 1 min to allow the derivitization reaction to proceed. The HPLC system consisted of a Waters 2695 Separations Module with a 100-µl sample loop and a Waters 24754 multiwavelength fluorescence detector set at 337 nm for excitation and 454 nm emission wavelengths. A Waters Xbridge C18 3.5 x 4.6 x 150 mm analytical column was used for chromatographic analysis. Mobile phase consisted of 0.05M sodium acetate, tetrahydrofuran, and methanol (50:1.49, vol/vol). Analyses were performed at 25°C ± 2°C. Analytes were isocratically eluted over a 22-min period at a flow rate of 0.8 ml/min. Glutamine, glutamate, and GABA were identified by their retention times (4.5, 7.6, and 18.6 min, respectively) as determined by standard injections. Standards for each analyte were assayed at the beginning of each HPLC run.

Corticosterone measurement. Corticosterone levels were measured using an immunosassay kit (Corticosterone EIA kit AC-14F1; IDS Inc.). Following the manufacturer’s protocol, blood was collected and centrifuged for 20 min at 3500 x g. Serum was removed and stored at −20°C. Samples were diluted in 1:10 ratio and all samples were run in duplicate. The corticosterone levels were calculated by comparison with a standard curve ranging from 0 to 192 ng/ml. Optical density values were measured at 450 nm using a microplate reader.

**RESULTS**

**Dam and Pup Measures**

Dam body weights increased significantly across the course of GD1-18 (F(17,238) = 76.13, p < 0.0001) from group mean values ranging from 21.4–23.37 to 29.03–30.56 g. No evidence was found of any effect of Pb, PS, or the combination.

Group mean litter sizes ranged from 5.57 to 6.6 and were not
influenced by Pb, PS, or the combination. Group mean male pup weights ranged from 10.38 to 11.97 g, and group mean female body weights from 10.0 to 11.09, and were not affected by Pb, PS, or the combination. A small (7–9%) but significant increase in the percent of male pups was found (F(1,51) = 4.35, p = 0.042) in response to PS. Values also differed significantly by BE but systematic differences were generally associated with 20–45% lower neurotransmitter levels. In addition, FS experience was associated with 19% reductions in levels of the 5-HT metabolite and 5-HIAA in both hippocampus and striatum.

**Females.** BE-related differences in 0-NS females were also found primarily in frontal cortex and hippocampus (Fig. 3; Tables 3 and 4). Differences in frontal cortex included DOPAC, DA turnover, and NE, where the direction of effects was opposite to that seen in males, that is, FS experience increased neurotransmitter levels relative to FI experience by levels ranging from 17 to 39%. In addition, levels of frontal cortex GABA were significantly increased ~30% by FS relative to FI experience.

**BE-Related Changes in the Effects of Pb, PS, or Pb + PS**

**Monoamines and males.** BE-induced differences in the effects of Pb, PS, or Pb + PS in male offspring were seen in frontal cortex, hippocampus, and striatum, but were particularly prominent in midbrain (Fig. 4; Table 3). In frontal cortex, these included changes in DA turnover and NE. No Pb or PS effects on DA turnover were found after FI experience, but both Pb alone and PS alone increased DA turnover after FS experience. Pb increased NE levels after FI experience, an effect more pronounced in Pb + PS conditions, albeit not significantly, whereas Pb decreased NE after FS experience.

**Effects of BE on Brain Neurotransmitter Levels**

**BE-Related Differences in Control (0-NS) Offspring**

**Males.** Analyses based on comparisons in the 0-NS groups with significant main effects of BE in ANOVAs revealed that the impact of FI versus FS experience in normal males was seen primarily in frontal cortex and hippocampal levels of DA and its metabolites (Fig. 2; Tables 3 and 4). FS experience was generally associated with 20–45% lower neurotransmitter levels. In addition, FS experience was associated with 19% reductions in levels of the 5-HT metabolite and 5-HIAA in both hippocampus and striatum.

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FIG. 2. Group mean ± SE neurotransmitters (ng/mg/protein) levels in male 0-NS offspring of indicated brain regions. FI: fixed interval schedule of reward BE; FS: forced swim BE. *Significant difference in post hoc test of at least $p \leq 0.05$ between FI and FS values following initial ANOVA (Tables 3 and 4), indicated by black bars for FS values. Sample sizes: frontal cortex, 7–17 per group; hippocampus, 8–17 per group; striatum, 7–16 per group.

TABLE 3
Summary of Statistical Analyses of Changes in Monoamine Neurotransmitter Levels

<table>
<thead>
<tr>
<th>Region</th>
<th>DA</th>
<th>DOPAC</th>
<th>HVA</th>
<th>DA TO</th>
<th>NE</th>
<th>5-HT</th>
<th>5-HIAA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frontal cortex</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Male</td>
<td>Pb $\times$ S $\times$ BE, 0.045</td>
<td>BE, 0.001</td>
<td>BE, 0.001</td>
<td>BE, 0.003</td>
<td>Pb $\times$ S $\times$ BE, 0.048</td>
<td>Pb $\times$ BE, 0.012</td>
<td>BE, 0.039</td>
</tr>
<tr>
<td>Female</td>
<td>BE, 0.013</td>
<td>Pb $\times$ S $\times$ BE, 0.016</td>
<td>BE, 0.043</td>
<td>BE, 0.000</td>
<td>Pb $\times$ S $\times$ BE, 0.038</td>
<td></td>
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<tr>
<td>Hippocampus</td>
<td></td>
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</tr>
<tr>
<td>Male</td>
<td>BE, 0.014</td>
<td>Pb $\times$ BE, 0.053</td>
<td>BE, 0.000</td>
<td>BE, 0.000</td>
<td>BE, 0.001</td>
<td>Pb $\times$ S $\times$ BE, 0.01</td>
<td>BE, 0.012</td>
</tr>
<tr>
<td>Female</td>
<td>BE, 0.000</td>
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<td>BE, 0.000</td>
<td>BE, 0.000</td>
<td>Be, 0.000</td>
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<tr>
<td>Striatum</td>
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<tr>
<td>Male</td>
<td>BE, 0.005</td>
<td>Pb $\times$ BE, 0.032</td>
<td>Pb $\times$ BE, 0.031</td>
<td>BE, 0.000</td>
<td>BE, 0.002</td>
<td>Pb $\times$ S $\times$ BE, 0.003</td>
<td>Pb $\times$ BE, 0.032</td>
</tr>
<tr>
<td>Female</td>
<td>S $\times$ BE, 0.05</td>
<td>BE, 0.042</td>
<td>Pb $\times$ BE, 0.03</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Midbrain</td>
<td></td>
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</tr>
<tr>
<td>Male</td>
<td>BE, 0.04</td>
<td>Pb $\times$ BE, 0.032</td>
<td>Pb $\times$ S $\times$ BE, 0.023</td>
<td>BE, 0.038</td>
<td>BE, 0.000</td>
<td>Pb $\times$ S $\times$ BE, 0.023</td>
<td>Pb $\times$ BE, 0.052</td>
</tr>
<tr>
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<td>BE, 0.017</td>
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</table>

Note. Pb = lead, S = prenatal stress.
*Includes outcome (main effects and interactions) of ANOVAs and corresponding $p$ value.
In striatum, BE-related differences in effects of Pb ± PS in male offspring were seen with NE, comprised of selective increases by Pb + PS after FI experience, whereas only PS alone increased levels after FS experience. 5-HIAA levels were not associated with Pb ± PS effects following FI experience, whereas Pb, PS, and Pb + PS all increased levels after FS experience.

BE-related differences in Pb ± PS effects in midbrain included DA, DOPAC, HVA, NE, 5-HT, and 5-HIAA. In most cases, this difference reflected increases in response to Pb ± PS in groups with FS experience, with the exception of a Pb + PS-induced reduction in 5-HIAA with FI experience. Pb alone increased HVA levels after FS experience, with similar but non-significant post hoc trends for DA. In the case of both DOPAC and NE, levels were increased by Pb alone and PS alone with FS experience, but not by the combination. Only PS alone increased 5-HT levels after FI experience.

**Monoamines and males.** For female offspring, BE-related changes in effects of Pb ± PS were less extensive than in males, being seen primarily in frontal cortex (Fig. 5; Table 3). In frontal cortex DOPAC, selective reductions were found only in the Pb alone group after FS experience, whereas no Pb ± PS effects were observed after FI experience. Interactions in the case of 5-HT were due to the selective increases in levels produced by Pb alone and by PS alone after FI experience, whereas no significant Pb ± PS effects were observed after FS experience.

Effects of Pb ± PS on hippocampal or midbrain monoamines were not differentiated by BE in female offspring. BE-related differences in striatum in effects of Pb ± PS were seen in 5-HIAA where no effects of Pb ± PS were found after FI experience, but Pb-related increases were observed after FS experience.

**Amino acids and males.** PS increased levels of frontal cortex GABA after FI experience, whereas Pb ± PS-induced
changes after FS experience were restricted to increased levels of frontal cortex GABA in response to PS only (Fig. 6, Table 4).

Amino acids and females. Frontal cortex GABA levels were increased by Pb alone and PS alone after FI experience, whereas Pb per se increased frontal cortex GABA after FS experience (Fig. 6, Table 4). In hippocampus, significant interactions for glutamine and glutamate were found. Hippocampal glutamine levels were unaffected by Pb ± PS after FI experience, whereas both increased levels after FS experience. Notably, in the case of hippocampal glutamate, Pb increased levels after FS experience, but decreased levels after FI experience.

Relationship of Neurotransmitter Changes to Behavior
Statistically significant relationships between neurotransmitter changes and behavioral outcomes were seen in males, whereas only trends emerged with females.

FS performance in males. Given that the first exposure to the FS paradigm could be considered most stressful (Koolhaas et al., 2011), simple linear regression analyses were carried out to determine whether outcomes on this test could be related to neurotransmitter changes.

Pb, PS, and Pb + PS reduced total number of immobile bouts (Fig. 7, left; Pb × stress: $F(1,40) = 4.79$, $p = 0.0345$; $p = 0.035$, and $0.004$, respectively vs. 0-NS control). Immobile bouts were significantly correlated with both striatal HIAA (top middle; $p < 0.000$) and frontal cortex GABA (bottom middle; $p = 0.019$). Number of immobile bouts declined with increasing striatal HIAA, and striatal HIAA levels were increased in Pb, PS, and Pb + PS groups (top right; $p = 0.011, 0.001$, and 0.008, respectively) after FS experience. Number of immobile bouts declined with increasing frontal cortex GABA levels, and frontal cortex GABA levels were significantly increased (bottom right) in the 0-PS ($p = 0.002$), with a similar but nonsignificant increase in the 100-NS group.

FI performance in males. A Pb × stress × session interaction ($F(1,1512) = 3.89$, $p < 0.0001$) characterized the changes in male offspring FI overall response rates across sessions (Fig. 8,
FIG. 5. Group mean ± SE monoamine neurotransmitters (ng/mg/protein) levels in female offspring of indicated brain regions. NS: 0-NS, no Pb, no stress; PS: prenatal stress, no Pb exposure; Pb: Pb exposure, no prenatal stress; Pb + PS: Pb exposure and prenatal stress. *Significant difference of at least $p \leq 0.05$ from corresponding 0-NS control group value following initial ANOVA (Table 3). Plots with black FS bars indicate significant interactions of Pb, PS, or both with BE in the overall ANOVA. Sample sizes: frontal cortex, 7–13 per group; hippocampus, 8–13 per group; midbrain, 8–12 per group.

FIG. 6. Group mean ± SE amino acid neurotransmitters (ng/mg/protein) levels in male (left panel) and female (right panel) offspring of indicated brain regions. NS: 0-NS, no Pb, no stress; PS: prenatal stress, no Pb exposure; Pb: Pb exposure, no prenatal stress; Pb + PS: Pb exposure and prenatal stress. *Significant difference of at least $p \leq 0.05$ from corresponding 0-NS control group value following initial ANOVA (Table 3). Plots with black FS bars indicate significant interactions of Pb, PS, or both with BE in the overall ANOVA.
A Bonferroni-Dunnett post hoc test comparing group rates collapsed across sessions confirmed higher rates in the 100-PS group compared with all other groups ($p = 0.000$, $p < 0.000$, and $p < 0.000$ for the 0-NS, 0-PS, and 100-NS comparisons, respectively). In simple linear regression analyses using FI response rates from the final session, significant correlations were found between overall response rates and hippocampal DA turnover (top middle; $p = 0.007$) and NE.

**FIG. 7.** Left: Group mean ± SE number of immobile bouts in male offspring during FS test 1 ($n = 8–16$ per group). Middle: Scatter plot from linear regression of immobile bouts against striatal 5-HIAA (top) and frontal cortex GABA (bottom). Right: Levels (ng/mg protein) of striatal 5-HIAA (top; $n = 7–16$ per group) and frontal cortex GABA (bottom; $n = 8–16$ per group) by treatment group (NS: 0-NS, no Pb, no stress; PS: prenatal stress, no Pb exposure; Pb: Pb exposure, no prenatal stress; Pb + PS: Pb exposure and prenatal stress) and behavioral paradigm (FI: fixed interval schedule of reward BE; FS: forced swim BE). *Significant difference of at least $p \leq 0.05$ from corresponding 0-NS control group value.

**FIG. 8.** Left: Group mean ± SE response rates on the FI schedule of food reward across sessions in male offspring ($n = 10–16$ per group). Middle: Scatter plot from linear regression of response rates against hippocampal DA turnover (DOPAC/DA; top) and hippocampal NE (bottom). Right: Levels (ng/mg protein) of hippocampal DA turnover (top; $n = 8–16$ per group) and hippocampal NE (bottom; $n = 8–17$ per group) by treatment group (NS: 0-NS, no Pb, no stress; PS: prenatal stress, no Pb exposure; Pb: Pb exposure, no prenatal stress; Pb + PS: Pb exposure and prenatal stress) and behavioral paradigm (FI: fixed interval schedule of reward BE; FS: forced swim BE). *Significant difference of at least $p \leq 0.05$ from corresponding 0-NS control group value.
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(bottom middle; $p = 0.009$), with increases in both associated with increased overall response rates. Correspondingly, in mice with FI experience, both Pb ($F(1,39) = 22.09, p < 0.000$) and PS ($F(1,39) = 4.31, p = 0.044$) increased hippocampal DA turnover (top right) and Pb increased NE ($F(1,42) = 4.32, p = 0.044$), particularly in the Pb + PS group (bottom right).

DISCUSSION

BE is known to profoundly influence later behavior and brain function (Bierman et al., 2008; Cheatham et al., 2010; Jaffee and Maikovich-Fong, 2011; Mills et al., 2011; Palfrey et al., 2005; Reynolds et al., 2007; Shonkoff et al., 2012). Correspondingly, in a prior study, we demonstrated that brain monoamine levels differed significantly in normal rats depending upon whether or not they had undergone BE (Cory-Slechta et al., 2009). That study additionally demonstrated that Pb ± PS effects on brain monoamines likewise differed, often dramatically, based on BE. Given the evidence that the nature of BE itself is a critical determinant of later outcomes (Koolhaas et al., 2011; Maier and Watkins, 2010), this study examined the hypothesis that positively reinforced food-rewarded responding on an FI schedule versus negatively reinforced inescapable FS experience could also alter the neurochemical effects of Pb ± PS.

Several findings emerged from this study that collectively demonstrate highly different profiles of Pb ± PS-associated neurotransmitter changes after FS versus FI BE. First, levels of several monoaminergic and amino acid neurotransmitters in several brain regions differed significantly in normal non-treated mice, that is, under control (0-NS) conditions. These effects, seen in both genders, were most prominent in regions considered critical to executive functions, that is, frontal cortex and hippocampus (Figs. 2 and 3). Changes in dopaminergic function in frontal cortex were opposite in direction in males versus females, with reduced function after FS experience in males, but with increases in females. Under 0-NS conditions, BE-dependent differences in amino acid neurotransmitters were seen only in females and only in frontal cortex (cf. Figs. 2 and 3). Such differences in both the profile of differences, as well as the directions of changes in neurotransmitter levels are likely determinants of the impact of experience on later behavior and brain function and may also relate to gender differences in behavior.

Findings from this study also are the first to demonstrate that the nature of BE could significantly modify the neurochemical consequences of Pb ± PS (Figs. 4–6), resulting in different overall profiles. These alterations were most pronounced in males, with a profile that included hippocampal dopaminergic changes, norepinephrine across brain regions, and multiple monoamines in midbrain. In females, monoamine effects were not systematic, but were most notable for amino acid neurotransmitters in frontal cortex and hippocampus. In some cases, Pb ± PS-induced changes in neurotransmitter levels were actually in opposite directions with FI versus FS experience, for example, frontal cortex and hippocampal DA and DA turnover in males, and hippocampal glutamate in females. Although PbBs of rodents (Bull et al., 1983; Leasure et al., 2008) and humans (Hertz-Picciotto et al., 2000; Rothenberg et al., 1994) increase over the course of pregnancy and lactation and thus provide spiked exposures at that time, the outcomes here occurred at stable PbBs, the more typically used metric, of ~7–10 μg/dl.

Such differences in the consequences of Pb ± PS-induced changes in neurotransmitter profiles may begin to assist in providing an understanding of mechanisms of gender-related differences in the consequences of Pb ± PS (Cory-Slechta et al., 2004, 2010; Weinstock, 2007). For example, they may relate to the gender differences in the impacts of Pb ± PS on repeated learning that we have observed, where enhanced impairments in accuracy in response to Pb + PS were found in females (Cory-Slechta et al., 2010), whereas enhanced learning occurred in males, a phenomenon we attributed to increased response rates and thus higher reinforcement density in males (Cory-Slechta et al., forthcoming).

Although BE has the potential to serve as an intervention strategy to ameliorate Pb ± PS-induced toxicity (Bellinger et al., 1989; Hubbs-Tait et al., 2009), it cannot be ascertained from this study design whether either FI or FS BE served in that capacity. This study served as a “baseline” and rationale from which to address such questions. For example, a next experiment would seek to determine whether the FI or FS behavioral history ameliorates or enhances the impact of Pb ± PS on known behavioral deficits associated with these developmental insults, for example, learning and attention. Indeed, the current findings can provide some specific hypotheses about outcomes of such experiments. For example, would the Pb-induced reductions in NE in frontal cortex after FS experience in males enhance learning (Lapiz and Morilak, 2006)? Or would the suppression of hippocampal glutamate by FI experience in females impair recognition memory in females (Yuen et al., 2012)?

Experimental approaches that systematically compare the spectrum of BEs will ultimately be critical to defining biomarkers of vulnerability versus resiliency (Koolhaas et al., 2011).

A third major finding of this study was the correlations between Pb ± PS-induced changes in neurotransmitter levels and associated behavioral outcomes seen in males (Figs. 7 and 8). PS and Pb both reduced the total number of immobile bouts in FS1 in males, effects that were significantly correlated with increasing levels of striatal 5-HIAA and frontal cortex GABA and corresponded to Pb and PS-induced changes in these neurotransmitters with FS, but not FI, experience. The FS test is used as a model of anxiety induction. Although frontal cortex GABA is a well-known and demonstrated mediator of anxiety (Briones-Aranda et al., 2005; Rodriguez-Landa et al., 2009), the exact nature of the involvement of these neurotransmitters in FS behavior also remains contradictory, given reports that both GABA agonists (Car and Wisniewska, 2006; Frankowska

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et al., 2007) and antagonists (Slattery et al., 2005) can act like antidepressants and reduce immobility time. Increases in FI response rates seen in males correlated with increases in hippocampal DA turnover and NE. Although correlations with frontal cortex or striatal monoamines might have been predicted based on our prior studies in rats (Cory-Slechta et al., 1997, 1998, 2002), these findings suggest additional pathways of involvement in mediation of this behavior.

However, the correlations of behavior with neurotransmitter levels must be viewed with caution. For one thing, $r^2$ values from the linear regressions were modest, indicating much more complex mediation of these behaviors. More importantly, the findings from this study show the dynamic interplay between BE and neurotransmitter function. Specifically, neurotransmitter levels and functions could have changed, for example, over the course of the four FS tests and thus not necessarily be representative of the neurotransmitter levels at the time of the first FS test. Similar considerations apply to the correlations of neurotransmitters with FI response rates. As shown in our prior study (Cory-Slechta et al., 2009), correlations of behavioral outcomes with biochemical or neurochemical outcomes collected postbehavioral testing may not necessarily provide accurate mechanistic insights.

Collectively, this study underscores the need for further assessment of the impact of specific types of BE as modifiers of Pb ± PS-induced CNS changes. Further, it seems highly unlikely that the dynamic effects of BE would be restricted to Pb ± PS, but would also extend to other developmental neurotoxicants. Examples of such phenomena already exist. In animal models, combined cocaine treatment coupled with the rearing environment (natural or cross-fostered) altered maternal behavior of first-generation offspring (Johns et al., 2005). Children exposed to both prenatal alcohol and postnatal traumatic experience showed lower IQ scores and more severe deficits in language, memory, visual processing, and attention, than did traumatized children who did not also have prenatal alcohol exposure (Henry et al., 2007). In another study, prenatal alcohol exposure and subsequent adverse caregiving environments resulted in greater vulnerability to language and social communication deficits (Coggins et al., 2007). For such reasons, it may also be particularly important to consider BE in epidemiological studies where possible, because it may relate to the magnitude of the outcome, as well as assist in explaining individual differences in variability and associated risk.

Given that the current findings demonstrate a critical role for the nature of BE as a determinant of the profile of neurotransmitter alterations of two developmental insults, they have critical implications for both experimental design and interpretation of animal studies. Typical “enrichment” paradigms do provide some BE, although they sometimes include individual housing for the “nonenriched” rodents, which itself can be a stressor. Additionally, rodents are often handled in the course of experiments, which depending upon the specific conditions, may or may not be a stressor. However, these procedures do not encompass the full spectrum, nor do they necessarily provide systematic comparisons, of BEs that would be pertinent. Therefore, the ability to extrapolate outcomes cited in studies of developmental toxicants to human populations and to understand individual differences in vulnerability may be enhanced by a better understanding of BEs during the lifetime. With respect to experimental design, the inclusion of BEs as a component of studies offers the opportunity to determine what specific BEs may ameliorate the impacts of such developmental insults as well as assistance in understanding of the underlying neurobiological mechanisms.

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