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Functional MRI approach to developmental methylmercury and polychlorinated biphenyl neurotoxicity

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Abstract

Prenatal and early childhood exposure to methylmercury (MeHg) or polychlorinated biphenyls (PCBs) are associated with deficits in cognitive, sensory, motor and other functions measured by neurobehavioral tests. The main objective of this pilot study was to determine whether functional magnetic resonance imaging (fMRI) is effective for visualization of brain function alterations related to neurobehavior in subjects with high prenatal exposure to the two neurotoxicants, MeHg and PCBs. Twelve adolescents (all boys) from a Faroese birth cohort assembled in 1986-1987 were recruited based on their prenatal exposures to MeHg and PCB. All underwent fMRI scanning during behavioral tasks at age 15 years. Subjects with high mixed exposure to MeHg and PCBs were compared to those with low mixed exposure on fMRI photic stimulation and a motor task. Boys with low mixed exposures showed patterns of fMRI activation during visual and motor tasks that are typical of normal control subjects. However, those with high exposures showed activation in more areas of the brain and different and wider patterns of activation than the low mixed exposure group. The brain activation patterns observed in association with increased exposures to MeHg and PCBs are meaningful in regard to the known neurotoxicity of these substances. This methodology therefore has potential utility in visualizing structural neural system determinants of exposure-induced neurobehavioral dysfunction.

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Keywords

Developmental neurotoxicity; Environmental exposure; Functional magnetic resonance imaging; Methylmercury; Neurotoxicity syndromes; Polychlorinated biphenyls; Prenatal exposure delayed effects

1. Introduction

For the past three decades, the detection of subtle, low-dose effects of exposures to neurotoxicants such as lead (Canfield et al., 2003), methylmercury (MeHg) (Cordier et al., 2002; Debes et al., 2006), and polychlorinated biphenyls (PCBs) (Jacobson et al., 1990) on brain functioning in children has received a great deal of attention (Bellinger et al., 2006; Grandjean and Landrigan, 2006). Neurobehavioral tasks are frequently used in these investigations because they are non-invasive and can be administered in a standardized manner (White, 2004).

Based on a variety of domain-specific neuropsychological tests and omnibus intelligence (IQ) tests, converging evidence shows that developmental exposures to MeHg adversely affects several brain functions, although some uncertainties exist, including impacts of mixed exposures among the populations examined, the extent of counterbalancing effects of seafood nutrients, timing of exposures and effects, and critical exposure indicators (Grandjean et al., 2006). Similarly, PCBs are known through animal and human studies to adversely affect brain function, though there is inconsistency across study results with regard to exposure levels and ages at which effects are detected and the specific test measures indicating neurotoxicity (Rice, 2006). As MeHg and PCBs are both contaminants of marine food, co-exposures occur in some populations, thus possibly leading to combined effects (Grandjean et al., 2001).

Although the precise neural mechanisms by which human brain functions are altered, especially in low-dose exposure scenarios, are poorly understood, these neurotoxicants most likely induce a cascade of complex changes within the developing central nervous system, resulting in the behavioral deficits that can be clinically assessed later on. An ideal investigative tool would allow researchers to visualize the functioning brains of exposed individuals while they are engaged in cognitive tasks that are affected by these toxicants. Functional magnetic resonance imaging provides such a tool (Papanicolaou, 1998) and has been used to assess the effects of lead exposure in children (Yuan et al., 2006).

In this pilot study, fMRI techniques were used to visualize activation of specific anatomical brain areas during behavioral tasks among adolescent children with well-defined prenatal exposures to MeHg and PCBs. The tasks used are associated with functioning of highly specific areas of the brain. This paper compares the findings among adolescents with high and low mixed exposures to MeHg and PCBs on behavioral outcomes assessing the domains of visual processing and manual motor function.

2.1. Participants

The 12 study participants were members of a cohort of children established at the three hospitals in the Faroe Islands by two of the authors (PW, PG). This cohort has been well described elsewhere (Grandjean et al., 1997). Briefly, it consists of 1022 children born in 1986–1987, about 75% of all births. These children were prenatally exposed to MeHg and PCBs due to maternal consumption of seafood, including pilot whale meat and blubber. At parturition, cord blood and maternal hair samples were collected in order to determine Hg concentrations. A piece of umbilical cord tissue was also taken, and half of these samples were later analyzed for PCBs and other major organochlorine compounds. Dose–effect relationships between prenatal exposure to MeHg and cognitive outcomes were seen in several domains at ages 7 and 14, including motor function, reaction time, language, and memory (Debes et al., 2006; Grandjean et al., 1997). The exposure to both substances at the time of clinical assessment was not associated with test performance. Sociodemographic variables affected these associations only marginally.

For the current study, recruitment of participants was designed to select groups of three boys who best represented each of four widely different types of prenatal exposure: high MeHg; high PCB; high mixed exposure; and low mixed exposure. Exposure to MeHg was defined as cord blood mercury concentration (μ g/L). PCB exposure was defined as total cord tissue PCB concentration (ng/g wet weight) based upon the three major congeners: (PCB-138 + $PCB-153 + PCB-180) \times 2$. The high and low mixed exposure groups were identified as the highest and the lowest products of Hg and PCB concentrations, while the highest and lowest ratios of Hg/PCB were used to identify the groups with high MeHg or high PCB exposures, respectively. The cohort members identified in this way ranged in age from 14 years, 9 months to 15 years, 7 months, with the median age being 15 years, 0 months. None of the subjects had any neurological diagnosis that might affect their performance, and all of them were right-handed and had birth weights above 2500 g. Any alcohol use during pregnancy was rare in this cohort and no effects were seen in relation to maternal tobacco smoking. Thus, given the homogeneity of this cohort, no attempt was made to match the subjects on parameters other than MeHg and PCB (to preserve anonymity, details on individual characteristics are not reported here). All 12 boys and their parents agreed to participate in the study. Consent forms for the study methods and for travel to the United States were signed by the parents. The study received ethical review board approval from the Faroese Ethical Review Committee and the McLean Hospital Institutional Review Board. Only boys were included to facilitate ethical review approval and travel permission to the US from the parents.

Results for mercury and PCB exposure for the adolescents are detailed in Table 1. Analysis of the PCB concentration in stored cord blood from the cohort subjects confirmed the high validity of the cord tissue analyses (Grandjean et al., 2001). The products of mercury and PCB in the high mixed exposure group (high Hg and high PCB) averaged approximately 100-fold greater than the products in the low, mixed exposure group (low Hg and low PCB); each of the two neurotoxicants averaged about 10-fold higher in the highly exposed group.

Similarly, ratios of Hg/PCB for the high Hg group were almost 100-fold higher than the ratios of Hg/PCB in the low Hg group. The high-PCB group had exposure levels similar to the high, mixed group, while the high Hg group averaged about half the MeHg exposure of the high-mixed group. Exposure levels, especially for MeHg, were much lower at the time of examination and were therefore not considered. Along with PCB, the pesticide metabolite 2,2-bis(4-chlorophenyl)-1,1-dichloroethene (p,p'-DDE) biomagnifies in pilot whale blubber, and the two persistent pollutants showed similar concentrations in cord tissue from the 12 subjects (average ratio, 0.97), with a correlation coefficient of 0.94.

2.2. Imaging methods

All participants underwent structural and functional MRI studies at the McLean Brain Imaging Center in Belmont, MA. The examiners were blinded to the exposure status of the subjects.

Conventional MRIs were acquired on all participants to ensure there were no structural brain abnormalities. Scans were acquired using a 1.5 Tesla General Electric Signa (modified by Advanced NMR, Milwaukee, WI) whole body imaging device (software level 4.8). A T1weighted mid-sagittal localizer image was followed by double spin-echo (DSE) 3 mm axial slices of whole brain (256×256 matrix, TR = 3000 ms, TE = 30 and 8 ms, 24-cm FOV with two interleaved acquisitions resulting in 108 contiguous DSE slices; voxel dimensions = $0.975 \text{ mm} \times 0.97 \text{ mm} \times 3 \text{ mm}$). A 3D Fourier transform spoiled gradient-recalled acquisition (3 DFT SPGR) was also used to generate images with good contrast between gray matter and white matter (Shenton et al., 1992) (124 1.5 mm thick coronal slices; TR = 35 ms, TE = 5 ms with one repetition; flip angle = 45° ; 256×256 matrix; voxel dimensions = $0.975 \text{ mm} \times 0.975 \text{ mm} \times 1.5 \text{ mm}$).

The fMRI study protocol consisted of a series of four behavioral tasks using the same imaging device as above and a quadrature head coil receiver system using a modified echoplanar imaging technique (approximately 15 continuous slices; thickness = 6 mm). A series of 50 sequential images were obtained during the tasks. Images were collected every 3 s using a gradient echo pulse sequence (TE = 40 ms, flip angle = 75°; image matrix = 64 × 128 with a 3 mm × 3 mm in plane resolution). The fMRI protocols were sequential task activation paradigms of 30 s resting baseline followed by alternating active and resting states in 30-s intervals, resulting in total paradigm lengths of 150 s.

The tasks included motor and visual organization tasks, naming, and a passive visual stimulation paradigm. These tasks were selected because extensive prior research has shown a significant relationship between higher exposure to MeHg and/or PCBs and lower performance on these tasks or on similar tests that assess the same neuropsychological domains (Grandjean et al., 2006; Rice, 2006) Although the tasks were adapted for use in the scanning environment, where movement is constricted, the naming and visual organization tasks produced highly complex activation patterns that proved difficult to analyze in the small group of subjects. A major limitation of fMRI is that observed activation may be a result of activation in multiple neural circuits though often interpreted as activation of a single cognitive task. Both naming and visual organization are complex cognitive tasks which likely recruit multiple neural circuits resulting in fMRI activation patterns that require

a larger sample size in order to determine more definitive activation patterns. In this paper, we therefore report the results for two tasks which are less complex, photic stimulation and finger tapping, as processing of each of these tasks is known to be mediated by brain structures that show pathological changes at autopsy in cases of clinical MeHg poisoning (Choi, 1989; Davis et al., 1994; Takeuchi et al., 1962). These two tasks, which are highly robust for use in fMRI (Witt et al., 2008; Gratton et al., 1997), are described below.

Photic stimulation—The participant was asked to open his eyes and lie quietly while data were obtained over a two and a half minute period consisting of two 30-s cycles of darkness alternating with two 30-s cycles of binocular, 8 Hz patterned flash photic stimulation produced by goggles with light emitting diodes. This sequence was included because it requires only passive viewing and does not require movement, motivation or attention.

Motor task – **left and right finger tapping**—The participant was asked to touch each finger to his thumb in a sequential, repetitive procedure during the activation periods. During the relaxation phase, the participant was asked to relax and not move his fingers. The task was performed with the right hand (right finger tapping, RFT) then repeated as a separate task with the left hand (left finger tapping, LFT). This task is robust, has demonstrated clear activation sites, and produces easily detectable signal changes for all participants.

2.3. Data analysis

Statistical parametric maps (SPMs) were generated for each participant using SPM99 (Friston et al., 1995a, b) (from the Wellcome Department of Cognitive Neurology, London, UK) implemented in Matlab (The Mathworks Inc., Sherborn, MA, USA). The functional data sets were motion corrected within SPM99 using the first image as the reference then normalized to a standard template from Montreal Neurological Institute (MNI). The normalized data sets were first resampled to $2 \text{ mm} \times 2 \text{ mm} \times 2 \text{ mm}$ within MNI space using sinc interpolation and then smoothed using a $4 \text{ mm} \times 4 \text{ mm} \times 4 \text{ mm}$ Full Width at Half Maximum Gaussian smoothing kernel (Smith, 2002). A 150-s box-car waveform, convolved with hemodynamic response function, was used as the reference paradigm. Using general linear model and the hemodynamically corrected reference paradigm, the *T*-score values were calculated for each voxel and the SPMs were generated.

In order to determine the differences in activation between the groups (each compared to the low, mixed group), a second-level paired *t*-test was performed. Since we had only three subjects per group, the SPM group maps were generated using a fixed-effects model with the individual contrast maps (Holmes and Friston, 1998). The resulting maps were thresholded at T = 3.5 (p < 0.0003) for left and right finger tapping paradigms and at T = 5.0 (p < 0.0001) for photic stimulation. The cluster-size threshold was set at 20 voxels. The location, size, and the strength of activation (T-score) were calculated for each activation cluster in SPM99. Anatomical regions were determined for activation maxima using the Talairach Daemon (Lancaster et al., 1997, 2000). The analysis was repeated for each task paradigm.

3. Results

All MRI scans were visually assessed for clinical and structural abnormalities by a neuroradiologist and all were read as structurally normal. Due to the complexity of analyzing and describing all four exposure groups on the behavioral tasks, this presentation focuses on the comparison of the three boys with high exposure to both MeHg and PCB with the three with low, mixed exposure.

Photic stimulation yielded significant activation bilaterally in the occipital lobes for both groups of participants. The group with low exposure showed the normal pattern of greatest activation in the visual association cortex (Brodmann's areas (BAs) 18 and 19) (Fig. 1). In contrast, the group with high, mixed MeHg/PCB exposure showed greatest activation in BA 17, primary visual cortex, and a portion of the mesial aspects of area 18. A paired *t*-test revealed significant bilateral activation within the visual association cortex (BA 19), and to a lesser extent, the right temporal and frontal lobes, for the high exposure group compared to the low exposure group (Table 2). Examination of the fMRI activation for those with only high MeHg (low PCB exposure) versus those with only high PCB exposure (low MeHg exposure) yielded little difference between the two groups though the high PCB group showed greater activation in the right occipital lobe.

Visual inspection of the activation patterns during the finger tapping task performed with the left hand (Fig. 1) in the group with low exposure revealed the typical pattern of activation observed in a simple motor task. There was significant activation in the premotor and motor cortex of the right hemisphere, contralateral to the finger movement. In contrast, the activation pattern visualized for the high, mixed exposure group revealed activation in the same regions, but the activation was bilateral rather than unilateral. The largest differences in activation between the two groups were seen ipsilateral to the side of the finger tapping in the frontal regions, including the motor cortex and the periventricular white matter adjacent to the lateral ventricle that contains fibers of the pyramidal tract. This finding was more dramatic for the left hand than finger tapping with the right hand (Tables 3 and 4). Examination of the fMRI activation for those with only high MeHg versus those with only high PCB group showed comparatively greater activation in the ipsilateral premotor and sensory cortex areas during the RFT task.

4. Discussion

To our knowledge, this is the first attempt to use functional imaging in children with prenatal exposure to MeHg and PCBs. Adolescents with the highest prenatal exposures to MeHg and PCB show significant differences in brain activation patterns compared to those with the lowest exposure. These findings demonstrate the utility of this non-invasive technique and provide confirmatory evidence of the exposure-associated CNS dysfunction that has been documented in the Faroese cohort and other groups of children with exposures to these contaminants (Cordier et al., 2002; Dahl et al., 1996; Grandjean et al., 1997).

In children with no history of toxicant or other brain insults, photic stimulation produces focal activation in the visual association areas (BA 18 and 19) (Kang et al., 2003). As shown in Fig. 1, the pattern of activation demonstrated by the low exposure group for the photic stimulation paradigm was consistent with this expectation. In contrast, the high exposure group showed generally more activation in the occipital lobe and right temporal lobe than the low exposure group, with comparatively less activation in BA 18 and 19 and significantly more activation for passive viewing of photic stimuli, and suggests that there may be toxicant-related damage to visual association areas among children with high exposures, possibly representing brain plasticity. In cases of clinical MeHg poisoning, pathological changes in the brain are especially prominent in the occipital lobes (Choi, 1989; Davis et al., 1994; Takeuchi et al., 1962).

Brain activation during the finger tapping task also showed significant differences between the groups. The children with low mixed exposure showed the normal pattern of brain activation, i.e., focal activation in premotor and motor areas of the brain contralateral to the active finger. In contrast, the highly exposed boys showed activation of bilateral motor areas when carrying out finger tapping with one hand. This finding may reflect disordered brain specialization; a finding which has also been seen in children with childhood lead exposure during a test involving language (Yuan et al., 2006). In fatal cases of MeHg poisoning, neuropathological changes include the motor cortex (Takeuchi et al., 1962). Incomplete lateralization has been documented in subjects with prenatal exposure to an endocrine disrupting agent (Smith and Hines, 2000). Both MeHg (Tan et al., 2009) and PCBs (Rasmussen et al., 2003) have been reported to exert endocrine disrupting effects.

Overall, this study showed normal brain activation patterns in the low exposure group, while the highly exposed group revealed different patterns of brain activation and activation of more parts of the brain in response to the tasks. The group with high PCB exposure only tended to show slightly greater activation on tasks of photic stimulation and right finger tapping than did the boys who had high MeHg exposure only. However, any attribution of the effects to MeHg and/or PCBs is not possible due to the small size of the study group and differences in exposure levels. Because p,p'-DDE correlated very closely with PCB and is less clearly documented as a developmental neurotoxicant (Grandjean and Landrigan, 2006), this pollutant was not further considered. Likewise, no attempt was made to link the imaging findings to specific neuropsychological test results of the subjects examined.

Although exposure-related deficits in motor speed and visually mediated tasks have been documented for the full cohort (Debes et al., 2006; Grandjean et al., 1997), which are in accordance with the rather dramatic imaging findings, the boys did not appear to have abnormalities serious enough to affect their visual and motor function in daily living. This suggests that the activation patterns revealed by fMRI may, in part, reflect adaptive brain responses to developmental toxicity on brain structures.

Results must be interpreted with caution due to the small number of participants and a fixedeffects analysis. However, the results are intriguing and statistically significant despite the small group size. They suggest that further fMRI investigations of children exposed to

MeHg and PCBs are warranted and that this technique deserves to be used in neurotoxicity studies in general.

5. Conclusions

This pilot study utilized functional MRI techniques to observe activation of specific brain areas during behavioral tasks in boys with well-characterized prenatal exposure to MeHg and PCB. The method revealed clear differences in brain activation between the boys with high exposure and those with low exposure on tasks requiring visual processing (photic stimulation) and manual motor movement (finger-tapping). The greater and more widespread brain activation in the high exposure group suggests that adolescents with high prenatal exposure required more brain resources to complete these two simple tasks and that differential specialization of brain areas may have occurred following prenatal neurotoxicant exposure. These findings support the utility of fMRI as a probe for CNS injury associated with developmental exposure to neurotoxicants.

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Abbreviations

fMRI	functional magnetic resonance imaging
SPMs	statistical parametric maps
MNI	Montreal Neurological Institute

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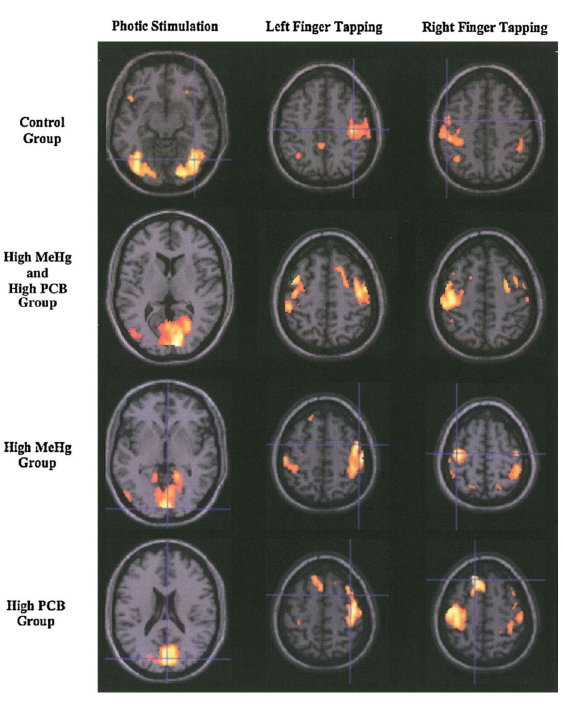


Fig. 1.

Average activation by exposure group during tasks of photic stimulation and finger tapping. The control group refers to the 3 subjects with low MeHg *and* low PCB exposure (left hemisphere is on left side of brain slices).

Table 1

Mercury (μ g/L cord blood) and PCB (ng/g wet weight of cord tissue) concentrations reflecting prenatal exposure levels in four groups of Faroese adolescents examined (three boys in each group).

	Mercury	РСВ
High mercury group	30.70-67.70	0.22-0.36
High PCB group	10.10-13.30	3.93-6.08
High mixed group	81.30-114.00	4.60-6.91
Low mixed group	4.30-13.80	0.40-0.53

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Table 2

Foci of maximally activated brain regions during photic stimulation when the group with high MeHg and high PCB exposure is compared to the group with low MeHg and low PCB exposure.

Region of activation	Brodmann'sarea	Region of activation Brodmann'sarea Number of activated voxels Coordinates	C00	rdinate	S	Max T-score Max Z-score	Max Z-score
			X	X Y Z	Z		
Right cuneus	19	4456	10	4456 10 –90	8	14.53	14.53 Infinity
Right occipital lobe	19	70	40	70 40 -80 28	28	8.33	Infinity
Right uncus	34	99	66 24 4		-32	6.25	6.25 6.05
Right temporal lobe	37	62	60	62 60 -64 -2	-2	6.16	6.16 5.97

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Table 3

Foci of maximally activated brain regions surviving left finger tapping when the group with high MeHg and high PCB exposure is compared to the group with low MeHg and low PCB exposure.

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Region of activation	Brodmann'sarea	Brodmann'sarea Number of activated voxels	Coordinates	linates		Max T-score	Max T-score Max Z-score
			X	Y	Z		
Left frontal lobe	32	468	32	~	32	8.24	7.80
Right frontal lobe		363	-28	-24	32	6.10	5.92
Left cerebellum		348	-24	-62	-30	5.84	5.68
Right premotor cortex	9	226	22	7	62	6.27	6.07
Right cerebellum		199	18	-64	-24	5.75	5.59
Left putamen		167	-20	9	10	6.54	6.32
Left inferior parietal lobe	40	148	-44	-24	22	5.50	5.36
Right premotor cortex	9	143	-40	0	58	7.58	7.24
Left precuneus	7	139	-14	-78	-46	5.66	5.51
Right motor cortex	4	135	10	-30	50	5.10	4.99
Right cingulate gyrus	24	124	10	×	28	4.93	4.83

Table 4

Foci of maximally activated brain regions surviving right finger tapping when the group with high MeHg and high PCB exposure is compared to the group with low MeHg and low PCB exposure.

Region of activation	Brodmann'sarea	Brodmann'sarea Number of activated voxels Coordinates	Coord	linates		Max T-score	Max T-score Max Z-score
			X	Y	Z		
Left precuneus	7	375	-24	-52	48	7.72	7.36
Right cingulate gyrus	31	338	14	-38	40	6.46	6.24
Left cerebellum		291	-30	-42	-28	6.44	6.22
Left thalamus		290	$^{-14}$	-26	0	5.72	5.57
Right cingulate gyrus	32	219	10	9	38	5.53	5.39
Left cerebellum		216	-28	-58	-36	6.42	6.20
Left motor cortex	4	163	-10	-34	60	5.80	5.64
Left lingual gyrus	17	154	0	-86	4	4.68	4.60
Left temporal lobe	36	145	-42	-42	7	5.20	5.08
Left premotor cortex	9	139	-42	8-	58	5.43	5.30
Right motor cortex	4	134	22	0	64	5.58	5.44
Right subcortical white matter at lateral ventricle		125	22	-44	14	6.27	6.06