Certain ortho-hydroxylated brominated ethers are promiscuous kinase inhibitors that impair neuronal signaling and neurodevelopmental processes

Robert G. Poston*, Lillian Murphy^, Ayna Rejepova*, Mina Ghaninejad-Esfahani*, Joshua Segales*, Kimberly Mulligan^, Ramendra N. Saha*†

* Molecular and Cell Biology Department, School of Natural Sciences, University of California, Merced, 5200 North Lake Road, Merced, California, USA, 95343
^ Department of Biological Sciences, Center for Interdisciplinary Molecular Biology: Education, Research and Advancement (CIMERA), California State University, Sacramento, USA, 95819

† To whom correspondence should be addressed: Ramen Saha, Ph.D: University of California, Merced, Room 346 S&E Building 1. 5200 North Lake Road, Merced, CA, USA, 95343; rsaha3@ucmerced.edu; Tel. (209) 228-2425

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Abstract

The developing nervous system is remarkably sensitive to environmental signals, including disruptive toxins, such as polybrominated diphenyl ethers (PBDEs). PBDEs are an environmentally pervasive class of brominated flame retardants whose neurodevelopmental toxicity mechanisms remain largely unclear. Using dissociated cortical neurons from embryonic Rattus norvegicus, we found here that chronic exposure to 6-OH-BDE-47, one of the most prevalent hydroxylated PBDE metabolites, suppresses both spontaneous and evoked neuronal electrical activity. On the basis of our previous work on MAPK/ERK kinase (MEK)–extracellular signal-related kinase (ERK) biology and our observation that 6-OH-BDE-47 is structurally similar to kinase inhibitors, we hypothesized that certain hydroxylated PBDEs mediate neurotoxicity, at least in part, by impairing the MEK–ERK axis of mitogen-activated protein kinase (MAPK) signal transduction. We tested this hypothesis on three experimental platforms: 1) in silico, where modeling ligand–protein docking suggested that 6-OH-BDE-47 is a promiscuous ATP-competitive kinase inhibitor; 2) in vitro in dissociated neurons, where 6-OH-BDE-47 and another specific hydroxylated BDE metabolite similarly impaired phosphorylation of MEK/ERK1/2 and activity-induced transcription of a neuronal immediate early gene; and 3) in vivo in Drosophila melanogaster, where developmental exposures to 6-OH-BDE-47 and a MAPK inhibitor resulted in offspring displaying similarly increased frequency of mushroom–body β–lobe midline crossing, a metric of axonal guidance. Taken together, our results support that certain ortho-hydroxylated PBDE metabolites are promiscuous kinase inhibitors and can cause disruptions of critical neurodevelopmental processes, including neuronal electrical activity, pre-synaptic functions, MEK–ERK signaling, and axonal guidance.

Introduction

Humans around the world face exposure to a vast number of anthropogenic environmental...
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contaminants that are largely untested for safety, many in the form of industrial chemicals. Use of one such class of environmentally widespread flame retardants—polybrominated diphenyl ethers (PBDEs)—began in the 1970s in response to regulatory efforts concerning fire-related safety of consumer products. They are now known to be stable in the environment and highly lipophilic, enabling collection in household dust and persistent bioaccumulation. The primary route of human exposure is ingestion and inhalation of such dust-borne PBDEs (1, 2), with dietary intake from seafood and dairy products accounting for much of the remaining exposures (3). Exposure levels are known to be high in the United States in particular, with infants and toddlers carrying the highest body burdens due to elevated intake rates from household exposures (4).

PBDEs have also been identified as one of a number of environmental pollutants that are considered developmental neurotoxins (5). Concerningly, a substantial amount of research has established the relationship of PBDE exposure levels with behavioral deficits in humans, as well as in animal models. Recently, several large-scale and systematic reviews have concluded that PBDE exposures impact externalizing behaviors and IQ in children and that BDE-47, -99, and -209 affect learning in animal studies (6–9). There is also concern for the relationship between environmental toxins like PBDEs and autism spectrum disorders (ASD), though the evidence is less conclusive, especially in human studies (6, 10–20). Given the prevalence of these compounds and their association with behavioral deficits, it is imperative to understand the molecular and cellular mechanisms underlying PBDE neurotoxicity.

Exposure to PBDEs has been shown to have numerous effects on neural cells, which are thoroughly reviewed elsewhere (21). Several of the major known mechanisms of PBDE toxicity include: 1) interference with endocrine signaling, likely caused by structural similarities to thyroid hormones and other hormone receptors’ ligands (22–25); 2) disruptions of calcium homeostasis/signaling as shown in one of the earliest mechanistic studies of PBDEs (26) and more recently demonstrated in several cell types including human neuronal precursors (27, 28); 3) toxicity produced by mitochondrial disruptions including uncoupling of oxidative phosphorylation (29) and subsequent elevated production of reactive oxygen species (ROS) which can lead to DNA damage and apoptosis; 4) dysregulation of epigenetic mechanisms including DNA methylation, chromatin dynamics, and non-coding RNAs—an emerging point of investigation that we have recently reviewed (30). It is possible that one or more of these molecular mechanisms contribute to exposure-related behavioral abnormalities; however, the extent of their involvement, if any, remains unclear.

The complex nature of the effects of these toxins is likely in part due to their diversity; there are 209 PBDE congeners, many found in commercial flame-retardant mixtures, and these compounds can be further endogenously metabolized in our cells. Such metabolism, mediated in humans by cytochrome P450 enzymes (31–33) and inherent in some other natural contexts (34–37), leads to the production of hydroxylated and methoxylated forms of PBDEs. In addition to parent congeners, toxicity may also arise from these metabolic products of PBDEs, some of which (e.g., 6-OH-BDE-47) have been found to be more toxic than the parent compound (25, 27). Given the numerous PBDEs humans are exposed to, including these metabolized forms, understanding how the effects of exposures to various PBDEs impact neurodevelopmental processes has been a difficult process, and it is likely that unreported mechanisms are involved.

Previously, we have utilized the ability of neurons to induce gene transcription in response to elevated network activity as a tool for assessing developmental PBDE toxicity (38). Specifically, we assayed levels of the neuron-specific activity-induced immediate early gene (IEG) Arc after exposing embryonic cortical neurons across various stages of differentiation and maturation. Our rationale for employing this endpoint was that Arc induction relies on multiple levels of regulation, from electrical activity and synaptic function, to intracellular signaling (particularly Ca2+-dependent mechanisms), to nuclear regulation of gene expression—all of which are critical processes intertwined in regulating
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These processes are largely hardwired by genetic programs but are also modulated by environmental cues. Exposure to toxins like PBDEs represent an environmental challenge that may dysregulate these processes at multiple levels, the cumulative effects of which could be detected as altered transcriptional responses. Results from our previous study showed that chronic exposures to a hydroxylated metabolite of BDE-47, 6-OH-BDE-47 (6-OH), dysregulate basal and activity-dependent neuronal transcription. Among those findings were data which strongly indicated that the attenuated transcriptional response following 6-OH exposure is due to synaptic dysfunction, and that 6-OH is potentially capable of inhibiting intracellular signaling. Either could explain the attenuated transcription of Arc, as its activity-induced transcription relies on synaptic activity as well as signaling via the Mitogen Activated Protein Kinase (MAPK) cascade, one of the major signaling pathways that is important for neuronal plasticity (40) and is necessary for Arc induction (41).

In this study, we propose and test a novel mechanism of PBDE metabolite toxicity: via impairment of MEK-ERK signaling. The notion that a PBDE metabolite could have the capacity to impair MAPK signaling was of particular interest to us for several reasons. The MAPK pathway is a deeply evolutionary conserved intracellular signaling mechanism that supports diverse processes in both prokaryotes and eukaryotes (42–44). In prokaryotes, PBDE-mediated inhibition of MAPK-homologous kinases could explain the natural biogenesis of these compounds as a defense mechanism, known to have antimicrobial activity and to be produced by bacteria in several marine contexts, including sponge-cyanobacterial symbiotic relationships (34–37, 45–48). In eukaryotes, the most well-known role of MAPK is regulation of cell growth, division, and differentiation in many cell types, including stem cells in the developing nervous system (49–51). It is also known to regulate axonal guidance and growth which has been demonstrated in multiple species (52–55). Additionally, there are numerous NDDs which are genetic syndromes involving the dysregulation of Ras and the MAPK pathway, thus known as 'RASopathies,' (56, 57). In mature neurons–post-mitotic non-dividing cells--the pathway regulates synaptogenesis (58, 59), synaptic plasticity (60), activity-induced gene expression (41, 61), and generally communicates information to a large number of cellular substrates. These roles in neural cells make MAPK signaling a concerning target for environmental disruption. However, the MAPK pathway remains largely unexplored as a target for pollutants.

With these factors in mind, we sought to expand and clarify our previous findings with the studies reported here. Here, we show that chronic 6-OH exposure suppresses neuronal electrical activity, that 6-OH is a promiscuous kinase inhibitor, and that acute exposures can inhibit MEK-ERK signaling. We also report that in vivo exposures to 6-OH and a MEK inhibitor, PD0325901, similarly alter axonal guidance in the mushroom bodies of D. melanogaster. These new findings corroborate our earlier work, provide evidence for the novel concept that certain hydroxylated PBDEs are promiscuous kinase inhibitors potentially capable of disrupting cellular processes via a range of intracellular signaling mechanisms, and further demonstrate the ability of PBDE metabolites to disrupt neurodevelopmentally relevant neuronal processes.

Results

Chronic 6-OH-BDE-47 exposure suppresses electrical activity in cortical neurons

We have previously shown that chronic exposure to nanomolar concentrations of 6-OH-BDE-47 (6-OH) impairs the ability of primary cortical neurons to induce transcription of the IEG Arc (38). Findings from that work indicated that the effects of the exposure were manifested at the level of synaptic functionality. To test this hypothesis directly, we utilized the same cell culture system, but grew cells on microelectrode arrays (MEAs). Cells were exposed to 500nM 6-OH from the day of plating and spontaneous electrical activity of the networks was recorded daily (Fig. 1A). There were no obvious morphological differences in cell growth. We also previously demonstrated that this exposure level does not significantly impact cell viability.
(38). Spontaneous activity was detected usually starting at 6 days in vitro (DIV), which is consistent with previous reports for cultured cortical neurons (62), and continued to the end of the experiment at DIV14. Networks exposed to 6-OH displayed significant suppression of spontaneous spiking across the recording period which was quantified and is summarized in Fig. 1B. To further clarify the synaptic nature of the exposure effects, we isolated synaptosomes from 6-OH treated cultures (DIV0-10) and evaluated levels of both pre- and post-synaptic protein markers by western blotting. We found a significant decrease in the amount of detectable Synapsin1, but not several other pre-synaptic proteins or the post-synaptic proteins PSD-95 and Gephyrin (Fig. 1C). This suggests that the observed activity deficits induced by chronic 6-OH exposure may be specifically mediated by pre-synaptic dysfunction involving Synapsin, although effects unrelated to the markers assayed here cannot be ruled out. Together, these data support the hypothesis generated from our earlier work that chronic exposure to the BDE-47 hydroxylated metabolite, 6-OH, impairs synaptic function.

Next, we observed the effects of 6-OH on induced neuronal activity. This was partly to explain one of our previous observations where activity-induced expression of the IEG Arc was significantly reduced in networks exposed chronically to 6-OH (38). To test if chronic 6-OH exposure impaired firing patterns during induced activity—which could explain impaired Arc transcription—we evoked activity in unexposed and 6-OH exposed networks using two different bicuculline (Bic) stimulation paradigms (Fig. 2A). The first was continuous treatment with Bic, similar to the approach we previously used to induce Arc transcription (38). The other was a Bic stimulus followed by washout, a treatment paradigm known to produce long-lasting recurrent synchronous bursting that has been previously published as a model of in vitro plasticity (63). Both types of Bic treatments produced characteristic burst firing patterns (Fig. 2B) that were diminished in 6-OH treated cells (Fig. 2C-F). The observed reduction in activity after 15 minutes of continuous Bic treatment (Fig. 2C, D) directly explains our earlier finding that chronic 6-OH exposure impairs activity-induced Arc transcription due to reduced firing of exposed neurons. The strong effect seen under the Bic washout paradigm (Fig. 2E, F) further demonstrates the compromised synaptic functionality of 6-OH treated cells and suggests that 6-OH may specifically impair the molecular mechanisms regulating induction and maintenance of Bic-induced bursting.

6-OH-BDE-47 is a promiscuous kinase inhibitor

The report which previously published the Bic washout paradigm as a model of in vitro plasticity also demonstrated that the effect was regulated by MAPK signaling, and could be blocked with a pharmacological MEK inhibitor (63). We additionally found that chronically treating neurons with low concentrations of a similar MEK inhibitor, PD0325901 (PD), impairs spontaneous spiking as well (Fig. S1). Combined with our previous observation that 6-OH is capable of attenuating MAPK driven gene transcription (38) and the observation that 6-OH shares key structural features with the same class of MEK inhibitors shown to regulate in vitro Bic-induced recurrent bursting (Fig. S2-A), we hypothesized that 6-OH may be capable of directly inhibiting MEK1. To test this hypothesis, and gain insight into the plausibility of 6-OH acting as a small molecule kinase inhibitor, we conducted ligand-protein docking simulations using published crystal structures of MEK1 and 6-OH as well as a commercial inhibitor, PD0325901 (PD) (a type-III non-ATP competitive inhibitor, which are well studied (64, 65)). Utilizing AutoDock Vina (66), which evaluated energetically optimized binding poses for each ligand, we found that both 6-OH and PD were placed around the well-characterized allosteric binding pocket for this type of inhibitor, without providing prior information on the binding surface. Almost all of the generated top poses for both 6-OH and PD clustered around this surface when the simulation was initialized near the catalytic kinase domain (Fig. S2-B). The poses for 6-OH and PD that are closest to published binding modes of MEK1 inhibitors are superimposed in Fig. S2-C. These results were encouraging and suggested that 6-OH is capable of realistically interacting with the type of
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To our surprise, when we attempted to demonstrate direct MEK1-6-OH binding in a cell free kinase-inhibition assay, there was no decrease in MEK1 activity across a range of 6-OH concentrations (Fig. 3B). However, when we screened a range of kinases, representing much of the human kinome, we found that 6-OH directly reduces the activity of a number of other kinases (Fig. 3A). In an attempt to identify shared features that enable the observed promiscuous inhibitory potential of 6-OH, further ligand-docking simulations were conducted with crystal structures of the top hits from the screen (Fig. 3C). Interestingly, the top 6-OH binding poses clustered largely in the well-known ATP-binding pockets of each of the modeled kinases. This suggests that 6-OH may, in fact, be a promiscuous ATP-competitive kinase inhibitor. Two of the top hits from the screen also suggest routes via which 6-OH may indirectly impair MEK-ERK signaling: Dyrk3, which is known to regulate Sirtuin activation thereby potentiating ERK activation by insulin signaling (67–69), and intriguingly, CaMK1, which has been shown to regulate depolarization-induced MEK-ERK activity and ERK-regulated long-term potentiation (LTP) (70–73). Focusing on CaMK1, we found that acute 6-OH exposures completely attenuate phosphorylation levels of a well-known pre-synaptic CaMK1 target, Synapsin1-Serine 9 (Fig. 3A) (74), providing in vitro confirmation of CaMK1 inhibition. We also reproduced an effect from a past report on CaMK1-MEK-ERK signaling, showing that acute exposure to the CaMKK inhibitor STO-609 attenuates depolarization induced phospho-ERK (pERK) levels (Fig. 3A) (70). Given this evidence, we further explored the dynamics of MEK-ERK signaling impairment by hydroxylated PBDE metabolites.

Specific ortho-hydroxylated PBDE metabolite exposures impair MEK-ERK signaling in vitro

In mature neurons, MEK-ERK signaling conveys information to many subcellular locations and substrates, including targets in synapses and the nucleus. To further test the prediction that 6-OH is capable of disrupting MEK-ERK signaling, we acutely exposed primary cortical neurons to BDE-47 and its hydroxylated metabolites and then stimulated them with a treatment of Bic and 4AP. The recurrent bursting driven by this treatment elevates intracellular calcium and induces the MEK-ERK signaling pathway (driving gene expression and plasticity-related events), which can be detected by measuring the phosphorylation status of both MEK and ERK. In whole cell extracts, we found that activity-induced elevation of both pMEK and pERK was significantly reduced only in cells treated with 6-OH (Fig. 4B), but not the parent compound or its other hydroxylated metabolites. The observed attenuation of pMEK induction is further evidence that 6-OH exerts effects somewhere above MEK in the signaling cascade, which includes CaMK1 as a possibility. As CaMK1 inhibition by 6-OH should only impact depolarization-induced MEK-ERK activation, we assessed the ability of 6-OH exposed cells to induce pERK when the activation was driven synapse-independently. To do this, we also treated cells with TTX and PMA. This treatment, which is previously described (38) and depicted in Fig. 5A, silences propagation of neuronal activity while activating MEK-ERK signaling through Protein Kinase-C (PKC) (which we knew was not inhibited by 6-OH from the kinase screen data). Again, we found that acute exposure only to 6-OH significantly attenuated pERK induction (Fig. 4C), indicating that 6-OH likely also impairs MEK-ERK signaling via synapse-independent mechanisms at high concentrations, though to a lesser extent than synapse-driven signaling. Inhibition of a synaptic signaling mechanism, such as CaMK1, could explain the stronger effect seen on synapse-dependent MEK-ERK induction.

While strong inhibition of MEK-ERK induction was observed with higher 6-OH concentrations that are consistent with the IC50 determination for CaMK1 (Fig. 3B), the lowest exposure level used (500nM) was ineffective. The 500nM dose was of interest to us as we have previously estimated it to be the approximate brain concentration reported from in vivo exposure studies of PBDEs in rodents (38) and is on the same order of magnitude as the EPA reported average for human exposure (4). As we initially did not see a significant decrease in
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pMEK/ERK levels with this environmentally-relevant concentration, we exposed cells to 500nM 6-OH with increasing pre-treatment lengths to test the impact of providing more time for membrane penetration and intracellular accumulation. This consistently yielded qualitatively mild but statistically insignificant inhibition of pERK in whole cell extracts (Fig. S3-A). Considering the subcellular location-specific functions of the MEK-ERK pathway, we next prepared nuclear extracts from neurons treated with 500 nM 6-OH to observe only the amount of pERK entering the nucleus (Fig. S3-B). Again, we observed mild reduction in pERK levels after acute exposure that was insignificant after quantification. At this point, it appeared that acute 500nM 6-OH exposures are not sufficient to detectably inhibit high pERK levels induced by the strong Bic+4AP stimulus. To test the effect of stimulus strength, we activated neurons with 5μM Bic without 4AP. With this weaker stimulus, we found that acute exposure to 500nM 6-OH significantly decreased nuclear levels of pERK (Fig. S3-C).

Next, we measured pre-mRNA levels of the MEK-ERK-dependent and neuron-specific IEG Arc after acute PBDE exposure as a second approach to corroborate the above-mentioned pERK induction data. We acutely exposed neurons to BDE-47 and its hydroxylated metabolites as in Fig. 4B and found significant attenuation of Arc induction only after 6-OH exposure (Fig. 5B). We also found that acute exposure to 6-OH significantly attenuated Arc induction driven synapse-independently (Fig. 5C). Again, the inhibitory effect of 6-OH was stronger on synapse-dependent induction, with significant attenuation of Arc induction at all tested doses. With the synapse-independent treatment, the effect was again insignificant with a 500nM exposure, as with the whole-cell pERK data (Fig. 4), so we again examined whether the strength of the stimulus impacts the extent of inhibition by titrating the concentration of PMA the cells were treated with. As expected, we found that with a lighter stimulus, the extent of 6-OH-induced inhibition was greater, with Arc pre-mRNA levels significantly reduced when 500nM 6-OH exposed cells were stimulated with 0.01μM PMA (Fig. 5D). Taken together these data indicate that acute 6-OH exposure impairs MEK-ERK signaling in neurons in dose- and stimulus strength-dependent manners and suggest that signaling in various subcellular locations may be differentially impacted.

To test the generality of our hypothesis regarding how hydroxylated PBDEs like 6-OH can impair MEK-ERK signaling, we acutely exposed cells to several other ortho-hydroxylated PBDE metabolites, whose possession or lack of the substituents hypothesized to mediate kinase inhibition are depicted in Fig. 6A. These include BDE-99, the second most environmentally prevalent PBDE humans are exposed to (4). It was surprising to find that only one of these ortho-hydroxylates metabolites, 6-OH-BDE-99, reduced pERK induction following Bic treatment (Fig. 6B). This metabolite produced nearly an identical effect on Arc induction as 6-OH-BDE-47, at various concentrations (Fig. 5B & 6C). This result suggests that for efficient MEK-ERK impairment, ortho-hydroxylated PBDE metabolites require a para-substituted halogen on the same ring as the hydroxyl group (Fig. 6A), potentially explaining why only certain ortho-hydroxylated metabolites impair elevation of pERK levels and subsequent Arc transcription.

Axonal guidance is dysregulated in vivo by both 6-OH-BDE-47 and PD0325901

Many developmental and functional processes in neurons are regulated by kinase activity, especially MEK-ERK signaling (54, 61, 75). Specifically, it is known to be involved in axonal guidance, along with CaMK1 (73, 76, 77), and that hydroxylated BDEs impact axonal growth in vitro (the evidence for which is covered in the discussion). We therefore hypothesized that developmental 6-OH exposures would lead to axonal guidance defects in vivo. To test this, we exposed adult flies (D. Melanogaster) and their offspring to BDE-47, two of its hydroxylated metabolites, and PD. We then assessed axonal guidance in the brains of offspring by observing effects on mushroom body ß-lobe axonal midline crossing (see Fig. S4-A for detailed experimental-timeline). The amino acid sequence of CaMK1 is highly conserved between H. Sapiens, R. Norvegicus, and D. melanogaster—including key residues known to be involved in binding ATP-competitive kinase inhibitors (Fig. 7A) (78). In wild type Drosophila,
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β-lobe axons are largely excluded from the midline of the mushroom body, though have been observed to have midline crossing defects at frequencies of approximately 20 percent (79). However, under conditions that disrupt normal axonal growth, increased frequency of midline crossing is observed for these axons (mild, moderate and severe). The qualitative scheme for assessing the extent of midline crossing is depicted in Fig. 7B and the frequency of the different effect severities are summarized in Fig. 7C. We found that 6-OH and PD exposure produced qualitatively similar increases in midline crossing, consistent with the involvements of both CaMK1 and MEK-ERK signaling in axonal growth. More mild effects were observed for BDE-47 and 5-OH-BDE-47 (5-OH) exposures. This became more evident when the distribution of crossing frequencies was rank scored for each exposure and assessed quantitatively (Fig. 7D). Analysis of rank score revealed that 6-OH and PD shift the median effect from “none” (as in the DMSO control) to “mild” while the parent compound and 5-OH did not produce a significant change. We also found that larval 6-OH exposure led to globally reduced levels of ERK activation –measured as the fraction of pERK detectable in total ERK immuno-precipitates from larval tissue samples (Fig. S4-B). These data indicate that 6-OH exposure can dysregulate axonal guidance in vivo and that the effect may be mediated via impairment of developmental CaMK1 and MEK-ERK signaling.

Discussion

PBDEs are well known environmentally pervasive toxins that are concerning for human health, especially considering the growing body of evidence linking their exposure levels to behavioral abnormalities related to NDDs. Much work has been done characterizing the effects and mechanisms of toxicity of PBDEs, but there is still not a clear general understanding of how these compounds compromise the developing nervous system and how the effects are related to adverse phenotypic outcomes. In this study, we propose a novel mechanistic hypothesis regarding how certain hydroxylated metabolites of common PBDEs (including BDE-47 and BDE-99) exert effects in neurons. We corroborate earlier findings indicating that exposures to an ortho-hydroxylated BDE-47 metabolite, 6-OH, impacts synaptic functionality and extended those findings by showing that 6-OH exposure affects neurodevelopmental processes in vivo. We also, for the first time, demonstrate that an ortho-hydroxylated PBDE metabolite is a promiscuous kinase inhibitor and that specific metabolites can impair neuronal MEK-ERK signaling.

From our previously published work, we had made two predictions related to the effects of chronic 6-OH exposure that are relevant to the data presented here: 1) effects in developing neurons likely manifest at the level of synaptic functionality and 2) 6-OH may be capable of inhibiting intracellular signaling at or downstream of membrane depolarization and PKC (38). Primary cortical neuronal cultures grown on MEAs (Fig. 1&2) allowed us to directly test the first hypothesis. While some recent studies have begun to employ MEAs to study PBDE developmental neurotoxicity as part of larger screens (80–82), the data presented here are, to our knowledge, the first example of such evidence for hydroxylated PBDE metabolite-induced effects. They demonstrate that chronic nanomolar 6-OH exposures suppress both spontaneous and evoked electrical activity and suggest that the exposures may specifically impact pre-synapses, including synapsin-related signaling and function. Inhibition of CaMK1 may interfere with neurotransmitter vesicle release by preventing phosphorylation of a well-known presynaptic target, Synapsin1-Serine 9 (74). This phosphorylation event is known to promote release of sequestered vesicles for exocytosis at the pre-synaptic membrane. Subsequent ERK activation is also known to be involved in recycling of vesicles via further Synapsin1-phosphorylation-dependent mechanisms (83, 84). 6-OH mediated CaMK1 inhibition and subsequent impairment of ERK induction could thus dysregulate pre-synaptic vesicle cycling, potentially explaining the observed effects of 6-OH on spontaneous electrical activity and Synapsin protein levels (Fig. 1B,C). These pre-synaptic defects and the pronounced effect of 6-OH on evoked activity under the Bic washout treatment paradigm, which has been previously
published as a model of in vitro plasticity (63), may partially explain the reported relationship between learning deficits and BDE-47/99 exposure in animal studies (9), as it is known that their hydroxylated metabolites can be produced endogenously, primarily via CYP2B6, which is dynamically expressed in the brain (31–33, 85–87).

Investigation of the second prediction generated from our previous work was spurred on by several lines of evidence: our previous observation of disrupted MAPK-driven gene induction by 6-OH (38), the known role of MAPK signaling in the effects seen on Bic-induced neuronal activity (Fig. 2) (63), and the serendipitous observation that 6-OH shares key chemical features with a class of inhibitors designed for MEK1, a central kinase in the MAPK signaling cascade (61, 88). The structural similarities and results from ligand-protein docking simulations encouraged us to pursue 6-OH as a potential MEK1 inhibitor (Fig. S2-A). Surprisingly, we came to find that 6-OH does not directly inhibit MEK1/2, but rather appears to be a promiscuous ATP-competitive kinase inhibitor (Fig. 3). We believe this novel finding will be of use in generating future hypotheses and furthering understanding of the complicated and diverse effects of PBDE exposures documented to date. Here we further pursued effects related to MEK-ERK signaling, particularly related to CaMK1, using multiple approaches to activate the pathway, mediated both synaptically and synapse-independently (Fig. 5A). We subsequently found that acute 6-OH exposures impair induction of MEK-ERK signaling and gene transcription (Fig. 4&5). Strong inhibition was found with 6-OH concentrations consistent with the data from the kinase screen (Fig. 3), while the lower, more environmentally relevant doses had milder effects, though still pronounced in the case of Arc transcription (Fig. 5). Additionally, we found that time-of-exposure does not appear to increase the extent of inhibition by these lower concentrations of 6-OH (Fig. S3-A) and that the strength of the stimulus driving MEK-ERK activation affects the extent of 6-OH-mediated impairment, with lighter stimuli leading to stronger impairment for both the synaptic and PMA-driven MEK activation (Fig. S3-C&5D). Overall, it is likely that acute exposure to environmentally relevant levels of MEK-inhibiting ortho-hydroxylated metabolites are not sufficient to widely suppress MEK-ERK signaling throughout the cell (especially given the extensive feedback opposing perturbations), but they may chronically and regionally accumulate to high enough concentrations to inhibit various kinases regulating MEK-ERK signaling in specific subcellular locations, an interesting possibility that is further discussed below.

To generalize our findings, we screened several other ortho-hydroxylated metabolites of BDEs-68/99/123 and found that the metabolite of BDE-99 also impairs MEK-ERK signaling following acute exposure (Fig. 6). While initially surprising, these data indicate that in order for an ortho-hydroxylated PBDE metabolite to be effective at impairing MEK-ERK signaling, it also specifically requires an ortho-substituted halogen that presumably enables binding to susceptible kinases. Taken together, these several lines of molecular evidence demonstrate that specific ortho-hydroxylated PBDE metabolites can inhibit MEK-ERK signaling in vitro. This potential for acute MEK-ERK impairment, combined with chronic-exposure-induced deficits in synaptic activity and composition, corroborates and further explains our previous findings concerning the ability of cortical neurons to induce Arc expression following chronic 6-OH exposure (38).

To validate our in silico and in vitro findings and extend them in vivo, we compared the exposure effects of 6-OH and PD using a neurodevelopmentally-relevant readout in fruit flies (D. melanogaster). Drosophila was chosen due to its amenability as an animal model for screening axonal guidance. Importantly, the Drosophila ortholog of CaMK1 is highly conserved— including the key amino acid residues that mediate binding of ATP-competitive inhibitors (Fig. 7A). Further, it is known that CaMK1 and MEK-ERK signaling play roles in developmental axon growth (89). One study reported that a let-60c mutation (a C. elegans ortholog of KRAS, a membrane GTPase upstream of MEK) led to axonal guidance defects manifested as aberrant ventral midline crossing (52). Another confirmed that MEK-ERK signaling regulates netrin-1 dependent axonal branching in cultured hamster sensorimotor
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cortical neurons (53). In our studies, exposure was accomplished via oral administration to the P1 generation—to ensure embryonic exposure—and also to F1 larva to cover the neurodevelopmental period of the mushroom body formation. The brains of offspring were examined post-eclosion to assess the extent of mushroom body β-lobe axonal midline crossing (see supplemental figure 4-A for detailed timeline). Here, we found that 6-OH and PD exposures produced qualitatively similar increases in midline crossing, with BDE-47 and 5-OH also increasing crossing frequency, but to a lesser extent (Fig. 7C). After applying a numerical scoring scheme and quantifying the effects, we found that 6-OH and PD shift the median effect to ‘mild’ while the parent compound and 5-OH did not produce a significant change. These data represent an in vivo confirmation of the effects of 6-OH exposure on axonal guidance (previously reported in primary hippocampal cultures (90)) and indicate that the effect may be mediated by disruptions to CaMK1 and MEK-ERK signaling. These findings align well with previous reports in C. elegans, where mutants of let-c, an ortholog of an upstream MAPK signaling component, KRAS, exhibit aberrant axonal crossing of the ventral midline (52).

While the evidence we present here demonstrates the promiscuous inhibitory potential of ortho-hydroxylated metabolites towards ATP-dependent kinases and MEK-ERK signaling, we note that such effects are highly nuanced and will require further investigation. MEK-ERK signaling is activated by a variety of stimuli in neurons and is a very dynamic process. Here alone we have identified CaMK1 and DYRK3 disruption as potential avenues of MEK-ERK impairment. Known modulation of Ca\(^{2+}\) homeostasis by 6-OH further complicates interpretation of acute exposures and potentially explains the inhibition of pERK levels by 6-OH following the synapse-independent stimulus paradigm (Fig. 4C). Further examples of nuanced effects include that cultures chronically exposed to 500nM 6-OH were able to synapticly induce cytosolic pERK to the same extent as untreated controls, while significantly less pERK was available in the nucleus (Fig. S3-B&C). This was unexpected, especially given the compromised electrical activity observed for chronically treated cells, but could be explained by the extensive homeostatic feedback mechanisms known to regulate MEK-ERK signaling (91). Such feedback mechanisms may also explain the discrepancy between our findings and those of another recent report which showed that exposures to 6-OH selectively impaired ERK5 phosphorylation, but not ERK1/2, in adult neural stem cells (aNSCs) (92). In this study the authors incubate aNSCs overnight with 5μM 6-OH, followed by a stimulus with EGF/bFGF to induced MAPK, and report no decrease in the amount of pERK1/2. In addition to the data from the present study showing that after prolonged treatment with 6-OH, cortical neurons induce the same level of pERK as untreated controls, it has been shown in mouse embryonic stem cells (ESCs) that acute PD treatment reduces pERK levels, but between 12 and 24 hours later, pERK signal strongly returns (93). Therefore, it is possible that either feedback within the MAPK pathways or differences in signaling dynamics in aNSCs, particularly those that activate MEK-ERK signaling, account for this discrepancy between our observation and that by Li and colleagues.

Interestingly, chronic 6-OH treatment-induced differences in cytosolic and nuclear pERK inhibition also suggest another nuanced aspect of PBDE toxicity: their sub-cellular localization and localized effects. Due to the highly lipophilic nature of PBDEs, they are likely to be more concentrated within close proximity of lipid membranes. Supporting this notion, it has been demonstrated that several PBDEs localize in higher amounts in mitochondrial and microsomal fractions collected from cerebral granule neurons (CGNs) (94). Accordingly, it will be interesting and necessary to evaluate the potential of ortho-hydroxylated PBDE metabolites to inhibit kinases in specific sub-cellular locations, as well as MEK-ERK signaling events, which are also known to be localized in different cellular compartments, including the cytosol, mitochondria, outgrowing neurites, and synapses (95, 96). Considering the effects on such compartmentalized signaling inhibition may help to generally explain some of the various documented effects of exposure to certain PBDEs, especially well-documented
mitochondrial toxicity (21). Further, mitochondrial toxicity may link PBDE exposure to NDD etiology given the known comorbidity of mitochondrial deficits and autism (12). Pre-synaptic accumulation could also be involved in the strong effect seen on Synapsin (Fig. 1C, 4A), as it and other synaptic proteins are known targets of CaMK1 and synapse localized MAPKs (97, 98). Both CaMK- and MAPK-dependent Synapsin phosphorylation are also known to play a role in pre-synaptic plasticity (59, 74, 84, 99). Local inhibition of CaMK1 and MAPK signaling and subsequent impairment of pre-synaptic vesicle regulation may explain the suppressed electrical activity caused by chronic 6-OH exposure (Fig. 1&2), though more work is needed for confirmation.

Taken together, the findings presented here provide new evidence on the effects of exposure to hydroxylated PBDE metabolites and establish novel insight into the underlying molecular mechanisms that mediate their toxicity. The indication that PBDE metabolites are capable of impairing MEK-ERK signaling sets up several potential avenues for further investigation. One takes into account another aspect of our previously published work (38), that 6-OH exposure dysregulates expression of BAF chromatin remodeling subunits. This evidence, along with a recent preprint report concerning ERK1/2 regulation of neurodevelopmental Polycomb Repressive Complex (PRC) chromatin remodeling that further speculates about additional regulation of BAF complex composition (100), suggests that 6-OH exposure may interact with chromatin remodeling via modulation of MAPK signaling. More closely related to the present study, it will be critical to investigate the role of ortho-hydroxylated PBDEs as promiscuous kinase inhibitors and their links to impairment of MEK-ERK signaling and the many established routes of PBDE toxicity. This will be especially interesting for effects related to mitochondrial function, calcium homeostasis, and regulation of synaptic function. Overall, such future studies may help to clarify the growing evidence linking human PBDE exposures to NDDs and perhaps provide useful insight into these challenging disorders.

Materials and Methods

Primary neuronal culture and cell treatment
Timed pregnant Sprague Dawley rats were obtained from Charles River Laboratories. Dams were consistently delivered two days prior to dissection and housed individually on a 12-hour light-dark cycle with access to food and water ad libitum. On the day of dissection, dams were anesthetized with Euthasol® euthanasia solution (pentobarbital 390mg/mL, phenytoin 50mg/mL) in order to minimize distress. Loss of responsiveness was ensured with the toe pinch method, confirming a lack of reflex prior to decapitation via guillotine. Primary cultures of cortical neurons were then prepared from the embryonic day 18 pups (UC Merced IACUC approval: AUP#16-0004). Cortical hemispheres were isolated and pooled from male and female embryos in magnesium and calcium-containing HBSS (Gibco, #14025092). They were then mechanically dissociated by pipetting with a fire-polished Pasteur pipet following a 7-minute digestion with StemPro® Accutase® (LifeTech, #A1110501). The enzyme was then deactivated by diluting the triturated cell suspension in HBSS lacking calcium and magnesium (Gibco, #14175095). The dissociated cortical neurons were then pelleted by centrifugation, resuspended in plating media, counted using trypan blue and a TC20™ automated cell counter (BIO-RAD, #1450102), and subsequently plated to dishes containing pre-warmed Neurobasal medium (Gibco, #21103049) supplemented with 25 µM glutamate (Sigma-Aldrich, #1446600), 0.5 mM L-glutamine (Sigma-Aldrich, #G8540), and 2% NS21 supplement. Cells were maintained at 37°C in a humidified incubator with 5% CO₂. They were grown in the medium described above without glutamate, replacing half the media every 3-4 days. NS21 was prepared in the laboratory as previously described (101). Neurons were used for various assays between 10–14 days in vitro.
and gene transcription using synaptic activity, neurons were co-treated with 50 or 5μM Bicuculline (Sigma-Aldrich, #14340) with or without 75μM 4-Aminopyridine (Acros Organics, #104571000). To induce gene transcription synapse-independently, membrane activity was blocked with 1μM TTX (Calbiochem, #554412) and the MAPK pathways were activated via PKC with 1μM-1nM phorbol 12-myristate 13-acetate (PMA; Sigma-Aldrich, #P1585). BDE-47 and hydroxylated BDE metabolites used to treat cultures were obtained from AccuStandard (BDE-047N, HBDE-4003N, HBDE-4004N, HBDE-4005N, HBDE-4006N, HBDE-5006-N, HBDE-5011N) and initially dissolved in dimethylsulfoxide (DMSO) at 2mM and 5mM stock concentrations. Final treatment concentrations were produced at the time of treatment by dilution in culture medium. PD0325901 was obtained from AdipoGen (SYN-1059) and dissolved in DMSO at 5mM and similarly diluted at the time of treatment.

Microelectrode array recording and data processing

Neurons from the preparations described above were plated on poly-L-lysine/laminin coated MEAs (60MEA200/30-Ti, Multi Channel Systems (MCS), Reutlingen, Germany) in 500 μL of the NS21-supplemented Neurobasal plating medium described above. When cells were fed, again every 3-4 days, approximately half the media was replaced with NS21-supplemented BrainPhys feeding media (StemCell). This was done to promote optimal conditions for neuronal firing which has been shown to be enhanced in BrainPhys media (102). Recordings were made with an MEA2100-lite system that interfaces with MCS provided Multi Channel Experimenter software. Sampling was conducted at 10-20kHz in 3-minute sessions at room temperature (arrays were covered to prevent contamination). Recording were post-processed in Multi Channel Analyzer with a high-pass 1st order Butterworth filter with 100hz cutoff prior to generation of spike time-stamps. Spikes were detected by an automatic threshold estimator set to 6-8 standard deviations from the baseline signal. To quantify burst properties, the Multi Channel Analyzer burst detection tool was used with the following settings: max. interval to start burst, 25ms; max. interval to end burst, 250ms; min. interval between bursts, 1500ms; min duration of burst, 50ms; min. number of spikes in burst, 5. Raster images from example recordings were generated in NeuroExplorer (RRID: SCR_001818, NexTechnologies, Herdon, VA).

Ligand-Protein docking simulations

For ligand-protein docking simulations, we used AutoDock Vina (RRID: SCR_011958), an open-source molecular docking program made available through the Scripps Research Institute (66). Initially, blind-docking of 6-OH-BDE-47 and PD0325901 was conducted with large areas of a published crystal structure of human MEK1 (PDB ID: 3EQI, (103)). Subsequent simulations searching a smaller grid-space centered around the catalytic binding domain were then conducted. Several of the top hits from the kinase inhibition screen reported here were also used in docking simulations with 6-OH-BDE-47: CAMK1 (4FG7, 4FGB, (104)), DYRK3 (5Y86, (78)), HIPK2 (6P5S, (105)), MAPKAP-K3 (3R1N, (106)), PIM1 (1YXT, (107)). The simulations were run with standard parameters for a rigid receptor and ligands with rotable bonds (108). The binding structures were evaluated visually based on comparison to the well characterized binding modes of allosteric MEK inhibitors, the first example of which was published in 2004 (109), as well as with various ATP-competitive kinase inhibitors crystallized with some of the other kinases.

Western blotting and imaging, and antibodies

Neurons were lysed in ice-cold 1X RIPA buffer (25 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Nonidet P-40) supplemented with 1:100 protease/phosphatase inhibitor cocktail (Thermo, #78442). Nuclear lysates were prepared by first washing cells with ice-cold 1X PBS, applying Plasma Membrane Lysis Buffer (PMLB) (25 mM Tris-HCl, pH 7.6, 10mM KCl, 1.5mM MgCl2, 0.5mM DTT, 0.1% NP-40) for 1 min on ice, washing with PMLB, and then collecting in 1X RIPA buffer (pH 7.4). Lysates were sheared by sonication (low setting; three cycles on Bioruptor®), cell debris pelleted at 15,000 rpm for 5 minutes at 4°C, and clarified supernatant transferred to pre-chilled 1.5 mL microcentrifuge tube. Synaptosomal isolations
were prepared using Syn-PER according to manufacturer instructions (Thermo, #87793). The various cell extracts were denatured at 95°C, for 5 minutes, using either 2X- or 4X-Laemmli sample buffer (BIO-RAD, #1610737 & #1610747). Denatured protein samples were resolved on 4-20%- (BIO-RAD, #4568095) or 4-15%- (BIO-RAD #456-1083) Mini PROTEAN® gels in Tris/Glycine/SDS (BIO-RAD #1610772). Resolved proteins were transferred onto LF PVDF membrane, using the BIO-RAD TBT RTA kit and protocol using either 10%MeOH or 20%EtOH-containing transfer buffer (#1704272). PVDF membranes were incubated at 4°C overnight with appropriate primary antibodies in 1X TBS-T with 1.5% BSA. Primary antibodies included the following antibodies: β–Actin (Thermo #AM4302, RRID: AB_2536382), H4 (CST #2935, RRID: AB_1147658), pERK (CST #4370, RRID: AB_2315112), ERK (CST #4696, RRID: AB_390780), pMEK (CST #9154, RRID: AB_2138017), MEK (CST #2352, RRID: AB_10693778), Synapsin1 (SySy #106011, RRID: AB_2619772), PSD-95 (NeuroMab #75-028, RRID: AB_2292909), Gephyrin (SySy #1471111, RRID: AB_887719), Synaptotagmin6 (NeuroMab #75-271, RRID: AB_11001830), Synaptophysin1 (SySy #101011, RRID: AB_887824). Next day, membranes were washed three times in 1X-TBST for 5 min each, probed with either goat-anti-Mouse-647 (RRID: AB_2535808) or goat-anti-Rabbit-546 Alexa Fluor (RRID: AB_2534093) secondary antibodies (Life Technologies) for 45 minutes at room temperature, washed three times with 1X TBS-T for 5 min each, and imaged using BIO-RAD Multiplex ChemiDocTM Imaging System.

RNA extraction and gene transcription quantitation
Total RNA was isolated from cultured neurons using the illustra RNAspin Mini kit (GE Lifesciences, #25050072). Specific pre-mRNAs from these total RNA samples was initially amplified by cDNA synthesis (14 cycles) using Arc and Gapdh primers overlapping an intron-exon junction and a OneStep RT-PCR kit (Qiagen, #210212). Quantitative real-time PCR (qRT-PCR) was then performed from this cDNA to quantify levels of specific transcripts using PerfeCTa SYBR Green FastMix (QuantaBio, #95072-012) and the BIO-RAD CFX Connect realtime PCR Detection System. Fold-change from control was estimated using the delta-delta Ct method.

Drosophila rearing & chemical exposure
Wild-type Canton-S Drosophila were maintained at 25°C in a humidified incubator on a standard cornmeal diet (Bloomington Drosophila Stock Center standard cornmeal medium). No randomization was performed to allocate individuals in the various treatment groups. For oral administration of PD0325901, BDE-47, 5-OH, and 6-OH, compounds were dissolved in dimethylsulfoxide (DMSO) and then added directly to warm food, prior to solidification. The concentration of DMSO was maintained at 0.02% for all exposures. Drosophila were exposed to PD (N=16) at a concentration of 100nM. For all other chemicals (BDE-47 (N=12), 5-OH (N=14), and 6-OH (N=14), Drosophila were exposed to a concentration of 1μM. Control flies were exposed to DMSO alone (N=14).

Fluorescence labeling and microscopy of Drosophila whole-mount brain preparations.
Whole-mount immunostaining of adult male brains (0-2 days old) was performed as previously described (79), except that brains were fixed for 25 minutes in 4% paraformaldehyde. Anti-Fasciclin II (RRID: AB_528235) was used at a 1:20 dilution. Alexa488-conjugated goat anti-mouse secondary (RRID: AB_2338840) was used at a 1:1000 dilution. Images were captured using an Olympus laser scanning confocal microscope. Separate individuals, who were blinded to the exposure group and file naming system, assessed the images for phenotypic scoring.

Immunoprecipitations (IPs) from Drosophila larval tissue
IPs were conducted by pooling the heads from ten chemically-exposed larvae. Tissue was lysed in ice-cold 1X RIPA buffer (25 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Na-deoxycholate, 0.1% SDS, 0.1% NP-40) supplemented with 1:100 protease/phosphatase inhibitor cocktail (Thermo, #78442). Lysate was added to a total volume of 1mL in IP buffer (0.5% Triton X-100, 2 mM
EDTA, 0.02 M Tris pH 7.75, 150 mM NaCl, and 10% glycerol) with 4 ug total ERK antibody (CST, #4696, RRID: AB_390780) and rotated for 2 hours at room temperature. Protein A/G magnetic beads (Thermo, #88803) were then added to the IPs, followed by rotation at room temperature for 1 hour. IPs were then washed three times on a magnetic rack with 1 mL wash buffer (0.5% Triton X-100, 2 mM EDTA, 0.02 M Tris pH 7.75, 150 mM NaCl, 10% SDS), resuspending the magnetic beads between each wash. After the final wash, the isolated proteins were denatured at 95°C for 5 min prior to gel electrophoresis and western blotting by the procedure described in the ‘Western blotting and imaging’ methods section.

**Study design and statistical analysis**

The study was not pre-registered. Data presented were generated from both hypothesis-confirming and exploratory experiments. For rat primary culture experiments, no sample size calculation was performed. For Drosophila experiments, sample size was estimated based on previously published studies that have examined the mushroom body phenotype (79). There were no pre-determined exclusion criteria for animal work. Statistical analyses were conducted using GraphPad Prism 7 (RRID: SCR_002798, GraphPad software, San Diego, CA). Error bars represent standard error of the mean throughout the article, except in Fig.7D where they represent 95% confidence intervals. Data were assessed for normality with the Shapiro-Wilk normality test. No tests for outliers were conducted, therefore all data-points were included. Effects were determined by t-test or one-/two-way ANOVA with appropriate post-hoc tests for generation of specific P-values, details are indicated throughout in figure legends. The only exception is Fig.7D where the non-parametric Mann-Whitney U test was used to compare the medians of effect score distributions for the various treatment groups. Specific P-values are indicated on figures, * indicates $P$-value <0.05, ** indicates $P$-value <0.01 throughout. Biological replicates are indicated throughout as N in corresponding figure legends. Biological replicates constitute cell culture preparations from the pooled cortices of embryos from independent litters or brains of individual flies.
Data availability: All data are included in the manuscript. Raw datasets are available upon request.

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Conflict of Interest: The authors declare that they have no conflicts of interest with the contents of this article.

Author contributions: RGP and RNS conceptualized the study; RGP, LM, AR, MG-E, and JS conducted experiments; RGP analyzed data and performed in silico modeling; the manuscript was written by RGP (lead) and RNS with feedback from all other authors; KM supervised Drosophila experiments, while RNS supervised the entire project.

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Figure Legends

Figure 1. Chronic 6-OH-BDE-47 exposure suppresses spontaneous neuronal activity and alters pre-synapses. Primary rat cortical neurons were grown on MEAs in order to assess the effects of 6-OH exposure on electrical activity. (A) Examples images of neurons at DIV10 with or without exposure to 500nM 6-OH (left) example traces of recorded spontaneous activity (right) (B) Detectable activity was measured daily with 3 minute recordings and quantified through the first two weeks of growth in vitro, N=3-4. P-values generated with two-way ANOVA with post-hoc FDR method, Time F(8,49) = 2.09, P=0.054, Treatment F(1,49)=36.04, P<0.0001, Interaction F(8,49)=1.503, P=0.1808. (C) Synaptosomes were isolated from DIV10 cultures. Pre- and post-synaptic markers were assessed by western blotting (left) densiometric quantification of blots (right) (note, the band between the Gephyrin and Synaptophysin1 is an additional band from the Gephyrin antibody), N=3-5. P-values generated with one-sample t-tests using a hypothetical mean of 1.

Figure 2. Chronic 6-OH-BDE-47 exposure impairs evoked synaptic activity. Multiple stimulus paradigms were used to assess the effects of chronic 6-OH exposure on evoked electrical activity. (A) Graphical depiction of bicuculline (Bic) stimulus paradigms. (B) Example of characteristic burst activity generate with Bic stimulus. (C&D) Quantification of spikes and bursts detected under continual Bic treatment of neurons chronically exposed to 500nM 6-OH (from time of plating to day of experiment), N=5-8. P-values generated with two-way ANOVA with post-hoc LSD, Spike data: Time F(2,30) = 0.269, P=0.76, Treatment F(1,30)=10.15, P=0.0034, Interaction F(2,30)=0.235, P=0.7920. Burst data: Time F(2,30)=2.356, P=0.11, Treatment F(1,30)=12, P=0.0016, Interaction F(2,30)=0.04056, P=0.9603. (E&F) Same experimental design and presentation as in B., but after brief Bic stimulation and washout instead of continual treatment, N=3-8. Spike data: Time F(2,22) = 4.62, P=0.021, Treatment F(1,22)=24.31, P<0.0001, Interaction F(2,22)=2.8, P=0.0825. Burst data: Time F(2,22)=1.57, P=0.23, Treatment F(1,22)=25.03, P<0.0001, Interaction F(2,22)=1.393, P=0.2695.

Figure 3. 6-OH-BDE-47 is a promiscuous kinase inhibitor. (A) 6-OH-BDE-47 was screened against a panel of 140 kinases representing much of the human kinome ( screened at 10 μM). Bars represent the mean percent activity of each kinase (screened in duplicate), and error bars represent standard deviation. The screening method utilizes a published radioactive filter binding assay using 33P ATP (110, 111), and was conducted by the International Centre for Kinase Profiling (ICKP) at the University of Dundee. Kinases with enlarged names are of particular interest for their relation to MEK–ERK signaling. (B) IC50 determinations were conducted by the ICKP (using the same methods as cited in A.) for indicated compounds with either MEK1 or CaMK1. Error bars represent standard deviation of each data point, which were conducted in duplicate. No inhibition was observed for BDE-47 or 6-OH with MEK1, while the IC50 for PD0325901 with MEK1 was here detected as 103±7nM, and 6.82±0.02μM for 6-OH with CaMK1. It is worth noting that this IC50 value for PD is around an order of magnitude higher than previously published, indicating that the estimations presented here may be affected by the sensitivity of this determination method. (C) Ligand-docking simulation results for the top five hits from the kinase screen. Crystal structures were obtained from the Protein Data Bank and are cited in the methods section. Autodock-Vina was used to find energetically optimal binding poses for 6-OH with each of the kinases, which are displayed in red in each panel. The final panel displays a structural alignment of the five kinases, with the docking results for CaMK1 superimposed. In each case, top binding poses clustered largely to the well-known ATP-binding pockets of the kinases. 6-OH acting as an ATP-competitive kinase inhibitor may help to explain the complex and diverse effects documented for PBDE exposures.

Figure 4. Acute 6-OH-BDE-47 exposure impairs MEK-ERK signaling. (A) Upper: Synaptosomes were prepared from neurons acutely exposed (20 min pre-treatment) to 5μM 6-OH with and without Bic+4AP stimulus (10 min). Levels of the CaMK1 target pSynapsin1(serine 9) were assessed via western blot. N=2. Lower: To reproduce a previously reported effect linking CaMK1 to MEK-ERK activity (70),
neurons were acutely exposed (60 min pre-treatment) to the CaMKK inhibitor STO-609 prior to treatment with Bic+4AP for 10 min (indicated as “+B10’”). Inhibition of pERK induction was assessed by western blot and is quantified below. N=3, P-value generated by unpaired one-tailed t-test. (B) Cells acutely exposed (20 min pre-treatment) to BDE-47 or one of its hydroxylated metabolites were stimulated with Bic+4AP for 10 minutes before collecting whole-cell lysates. Activation of MEK-ERK signaling was assessed by western blot, measuring pERK and pMEK levels. Note, the lower band in the MEK panel is the ERK band visualized on the same blot. Quantification relative to total ERK and MEK shown below representative blots. N=3-6, P-values generated by one-way ANOVA with post-hoc LSD, pERK quantification: Treatment F(7,22)=9.822, P<0.0001. pMEK quantification: Treatment F(7,16)=7.805, P=0.0004 (C) The same experimental design as in B, but neurons were stimulated with TTX+PMA to induce MEK-ERK signaling synapse-independently. N=3, pERK quantification: Treatment F(7,16)=37.95, P<0.0001.

Figure 5. Acute 6-OH-BDE-47 exposure attenuates activity and MEK-ERK dependent Arc transcription. MAPK-dependent gene transcription was evaluated as a measure of 6-OH mediated MEK-ERK inhibition by quantifying levels of Arc pre-mRNA following multiple stimulation approaches. (A) Graphical depiction of the stimulation methods used: Bic+4AP synaptically activates MEK-ERK signaling via depolarization, while TTX+PMA suppresses neuronal activity and activates MEK membrane-independently via PKC. Both ultimately drive induction of Arc gene transcription. (B) Arc pre-mRNA levels were estimated following exposure to BDE-47 or one of its hydroxylated metabolites and treatment with Bic (50μM) and 4AP (75μM), N=3-5 P-values generated by one-way ANOVA with post-hoc LSD, Treatment F(7,24)=25.1, P<0.0001. (C) Similar to B, except that cells were stimulated with TTX (1μM)) and PMA (1μM)), N=3-6. Treatment F(7,21)=12.14, P<0.0001. (D) To assess the effect of stimulus strength on inhibition of Arc induction by 6-OH. Neurons were acutely exposed to 500nM 6-OH before stimulus with a range of PMA concentrations (1μM)-1nM), N=3-6. P-values generated by two-way ANOVA with post-hoc LSD, PMA-Dose F(3,27)=33.16, P<0.0001, 6-OH Exposure F(1,27)=9.969, P=0.0039, Interaction F(3,27)=0.7964, P=0.5066.

Figure 6. Specific ortho-hydroxylated PBDE metabolites impair neuronal MEK-ERK signaling. In an effort to generalize our hypothesis that ortho-hydroxylated PBDE metabolites can inhibit MEK-ERK signaling, both approaches used to evaluate MEK inhibition by BDE-47 and its metabolites (Fig 4 and 5) were re-employed with several additional ortho-hydroxylated metabolites. (A) Chemical structures of PBDEs screened, with key substituents for potential MEK1 binding highlighted, specifically indicating the newly identified ortho-halogen thought to be critical for inhibition of MEK-ERK signaling. (B) Representative western blot measuring pERK levels after exposure to various ortho-hydroxylated PBDEs (top) with quantification of blots summarized (bottom), N=4. P-values generated by one-way ANOVA with post-hoc LSD, Treatment F(3,12)=7.314, P=0.0048. (C) Similar PBDE metabolite exposures as in B, but evaluating Arc pre-mRNA level attenuation by BDE-47 and BDE-99 metabolites, N=4-5. Treatment F(7,31)=8.837, P<0.0001.

Figure 7. In vivo exposure to 6-OH-BDE-47 and the MEK inhibitor PD0325901 disrupt axonal guidance. Canton-S flies (D. melanogaster) were exposed to BDE-47, 5-OH, 6-OH, or PD via feeding. The mushroom bodies of offspring were then assessed for aberrant midline crossing. (A) Alignment of the amino acid sequences of human CaMK1 and its rat and fly orthologs. The proteins are highly conserved, including the key residues thought to mediate ATP-competitive inhibitor binding (highlighted in yellow). (B) Example images of the effect categories used in the qualitative scoring scheme. (C) Crossing frequency by severity level for each of the treatment groups, N=12-16. (D) Crossing severity was numerically rank scored as follows: No crossing: 4, mild crossing: 3, moderate crossing: 2, severe crossing and/or lobe malformation: 1. Bars represent the median score of each treatment group which was generated from the values of the crossing data from C. after application of the numerical scoring scheme (the score for each fly is additionally plotted as grey circles). Error bars represent 95% confidence
Certain hydroxylated PBDEs impair neuronal MEK-ERK signaling intervals. 

$P$-values generated using exact one-tailed Mann Whitney tests to compare the median of each group with that of the DMSO control, $N=12-16$. 


A. Bicuculline treatment

B. DIV10-14 cortical neurons +/- chronic 6-OH-BDE-47 exposure

C./D.: Bic

C. MEA

D. Time: 0 Final recording (60 min or next day)

E. Time after Bic addition (min)

F. Time after Bic washout (min)
A. 

**pSynapsin1(S9)**

- Un. B10' 6-OH
- Un. B10' 6-OH + B10'

**β-actin**

- Un. B10' 6-OH + B10'
- STO-609 + B10'

**pERK**

- Un. B10' 6-OH
- Un. B10' 6-OH + B10'

**ERK**

- Un. B10' 6-OH
- STO-609 + B10'

**pERK/ERK band intensity**

- Untreated
- B10'
- STO-609 + B10'

B. 

**pERK**

- Un. B10' BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

**ERK**

- Un. B10' BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

**pMEK**

- Un. B10' BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

**MEK**

- Un. B10' BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

**β-actin**

- Un. B10' BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

**pERK/ERK band intensity**

- Untreated
- B10'
- STO-609 + B10'

C. 

**pERK**

- TTX + PMA10'
- 5μM 1μM 500nM

**ERK**

- TTX + PMA10'
- 5μM 1μM 500nM

**pERK/ERK band intensity**

- TTX
- PMA10'
- BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

**pMEK/MEK band intensity**

- TTX
- PMA10'
- BDE-47 3-OH 5-OH 6-OH 6-OH + B10'
- 5μM 1μM 500nM + B10'

Poston et. al. figure 4
**Figure 5**

**A.** GABAergic synapse (inhibitory) with TTX + PMA, Bic + 4AP, 6-OH, PKC, MEK/ERK, CaMK1, and Arc.

**B.** Synaptic-dependent (activity-driven) Arc induction:
- Untreated
- B10 (5 μM BDE-47 + B10)
- B10 (5 μM 3-OH-BDE-47 + B10)
- B10 (5 μM 5-OH-BDE-47 + B10)
- B10 (1 μM 6-OH-BDE-47 + B10)
- 500 nM PD + B10

**C.** Synapse-independent (MAPK-driven) Arc induction:
- TTX
- TTX + PMA (15 μM)
- TTX + PMA (15 μM) + 5 μM BDE-47
- TTX + PMA (15 μM) + 5 μM 3-OH-BDE-47
- TTX + PMA (15 μM) + 5 μM 5-OH-BDE-47
- TTX + PMA (15 μM) + 1 μM 6-OH-BDE-47
- TTX + PMA (15 μM) + 500 nM PD

**D.** Synapse-independent (MAPK-driven) Arc induction (stimulus titration):
- TTX
- TTX + PMA (1 μM)
- TTX + PMA (0.1 μM)
- TTX + PMA (0.01 μM)
- TTX + PMA (0.001 μM)
- 500 nM 6-OH
- 500 nM 6-OH + TTX
- 500 nM 6-OH + TTX + PMA (0.01 μM)
- 500 nM 6-OH + TTX + PMA (0.001 μM)

Poston et. al. figure 5
A. 

BDE-47

3-OH-BDE-47

2'-OH-BDE-68

6'-OH-BDE-99

6-OH-BDE-123

B. 

Un. B10' -99 + B10' -68 + B10' -123 + B10' 50 kDa pERK 37 50 ERK 37

C. 

Arc pre-mRNA relative to Gapdh

Untreated B15' 5μM 6'-OH-BDE-99 + B15' 5μM 6-OH-BDE-68 + B15' 5μM 6'-OH-BDE-123 + B15' 6'-OH-BDE-99 + B15' 500nM PD + B15'

Poston et al. figure 6
A. H. sapiens_CAMK1 --MLGAVEGPRWKQAEDIRDI---------YDFRDVLGTGAFSEVILAEDK-RTQKLVAI
R. norvegicus_CAMK1 --MPGAVEGPRWKQAEDIRDI---------YDFRDVLGTGAFSEVILAEDK-RTQKLVAI
D. melanogaster_CAMK1 MPLFGKKDSGKAKAKDLKELNQVISEEYNYLHGKGATESRLAESKDPGEHFAV

H. sapiens_CAMK1 KCIACKAELGEGSMNEAVLHAKHQPNIANLDDIIYESTGHSLYILQSGELFDVR
R. norvegicus_CAMK1 KCIACKAELGEGSMNEAVLHAKHQPNIANLDDIIYESTGHSLYILQSGELFDVR
D. melanogaster_CAMK1 KIIIDKAKLKGKTEELNEIRVLRLTHPINIVLQLEDEKSKVLYMELTGSGELFDVR

H. sapiens_CAMK1 EKGYTERDASRLIFQVLDAVKYLHDLGIVHRDLKPENNYLSLDIESKMSDSDFGLSKM
R. norvegicus_CAMK1 EKGYTERDASRLIFQVLDAVKYLHDLGIVHRDLKPENNYLSLDIESKMSDSDFGLSKM
D. melanogaster_CAMK1 EKGYTERDASRLIFQVLDAVKYLHDLGIVHRDLKPENNYLSLDIESKMSDSDFGLSKM

H. sapiens_CAMK1 EDPGSVLSTACPQYVAPEVLQPKYSKAVDCWSIGVIAYLLCCYFPYDENDAKLFE
R. norvegicus_CAMK1 EDPGSVLSTACPQYVAPEVLQPKYSKAVDCWSIGVIAYLLCCYFPYDENDAKLFE
D. melanogaster_CAMK1 EDPGSVLSTACPQYVAPEVLQPKYSKAVDCWSIGVIAYLLCCYFPYDENDAKLFE

H. sapiens_CAMK1 QILKAEYEFDSPWDDISDSAKDFIRHLMEKDPEKRFTCEQALQPIAIGATDLKNIHQ
R. norvegicus_CAMK1 QILKAEYEFDSPWDDISDSAKDFIRHLMEKDPEKRFTCEQALQPIAIGATDLKNIHQ
D. melanogaster_CAMK1 QILKAEYEFDSPWDDISDSAKDFIRHLMEKDPEKRFTCEQALQPIAIGATDLKNIHQ

H. sapiens_CAMK1 -PAAGCCCRDCCVEPGTELSPTLPHQL----
R. norvegicus_CAMK1 -PAAGCCCRDCCVEPGTELSPTLPHQL----
D. melanogaster_CAMK1 -PAAGCCCRDCCVEPGTELSPTLPHQL----

Poston et. al. figure 7

B. No Crossing Mild Crossing Moderate Crossing Severe Crossing

C. N =

| Condition         | No Crossing | Mild Crossing | Moderate Crossing | Severe Effects |
|-------------------|-------------|---------------|------------------|---------------|
| 0.02% DMSO        | 14          |               |                  |               |
| 1μM BDE-47        | 12          |               |                  |               |
| 1μM 5-OH-BDE-47   | 14          |               |                  |               |
| 1μM 6-OH-BDE-47   | 14          |               |                  |               |
| 100nM PD0325901   | 16          |               |                  |               |

0.02% DMSO
1μM BDE-47
1μM 5-OH-BDE-47
1μM 6-OH-BDE-47
100nM PD0325901

Effect Score

D. 0.07 0.15 0.01 0.02 *
Certain ortho-hydroxylated brominated ethers are promiscuous kinase inhibitors that impair neuronal signaling and neurodevelopmental processes

Robert G Poston, Lillian Murphy, Ayna Rejepova, Mina Ghaninejad-Esfahani, Joshua Segales, Kimberly Mulligan and Ramendra N Saha

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