

Defining toxicological tipping points in neuronal network development[☆]

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ABSTRACT

Measuring electrical activity of neural networks by microelectrode array (MEA) has recently shown promise for screening level assessments of chemical toxicity on network development and function. Important aspects of interneuronal communication can be quantified from a single MEA recording, including individual firing rates, coordinated bursting, and measures of network synchrony, providing rich datasets to evaluate chemical effects. Further, multiple recordings can be made from the same network, including during the formation of these networks *in vitro*. The ability to perform multiple recording sessions over the *in vitro* development of network activity may provide further insight into developmental effects of neurotoxicants. In the current study, a recently described MEA-based screen of 86 compounds in primary rat cortical cultures over 12 days *in vitro* was revisited to establish a framework that integrates all available primary measures of electrical activity from MEA recordings into a composite metric for deviation from normal activity (total scalar perturbation). Examining scalar perturbations over time and increasing concentration of compound allowed for definition of critical concentrations or “tipping points” at which the neural networks switched from recovery to non-recovery trajectories for 42 compounds. These tipping point concentrations occurred at predominantly lower concentrations than those causing overt cell viability loss or disrupting individual network parameters, suggesting tipping points may be a more sensitive measure of network functional loss. Comparing tipping points for six compounds with plasma concentrations known to cause developmental neurotoxicity *in vivo* demonstrated strong concordance and suggests there is potential for using tipping points for chemical prioritization.

1. Introduction

The lack of information on the potential for tens of thousands of chemicals in the environment to cause developmental neurotoxicity (DNT) is well documented (Grandjean and Landrigan, 2006, 2014). Because guideline DNT studies (US EPA, 1998; OECD, 2007) are expensive, time-consuming and animal intensive (Smirnova et al., 2014), it is now generally recognized that it will be necessary to use a battery of *in vitro* screening assays (Bal-Price et al., 2010, 2015; Fritzsche et al., 2017) to screen thousands of compounds for their potential to cause DNT and thus prioritize them for additional testing. To this end, a wide variety of assays have been developed to determine the effects of compounds on processes critical to the development of the nervous system, including proliferation (Breier et al., 2008), neurite outgrowth

(Radio and Mundy, 2008; Harrill et al., 2010; Stiegler et al., 2011), synaptogenesis (Harrill et al., 2011), transcriptomic responses (Krug et al., 2013), differentiation (Schmuck et al., 2017), and migration (Zimmer et al., 2014).

Formation of interconnected neural networks is critical to development of the nervous system. *In vivo* formation of networks occurs as the structures of the nervous system develop, and connections between different neurons form as axons and dendrites extend using chemical and electrical cues as guidance. Similar *in vitro* network activity arises spontaneously in primary monolayer cultures from many different brain regions, including the cortex, hippocampus, and brainstem as dissociated neurons (re)establish synaptic connections. Both in the brain and in primary cultures, this spontaneous electrical activity becomes patterned and coordinated over time, with synchronous bursting

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observed. *In vivo* synchronous bursting is associated with important nervous system processes including attention, learning, and memory (Buschman and Kastner, 2015; Korte and Schmitz, 2016; Salinas and Sejnowski, 2001). *In vitro*, such activity can be observed in networks of developing neurons using microelectrode array (MEA) recording techniques (Chiappalone et al., 2003; Van Pelt et al., 2004; Wagenaar et al., 2006; Cotterill et al., 2016; Brown et al., 2016). MEAs measure extracellular electrical activity associated with the generation and propagation of action potentials (spikes). Over time, patterned activity in the form of groups of spikes (bursts) from the same neuron, as well as coordinated bursting of multiple neurons across a network are readily observed. Measuring chemical effects on the development of spontaneous network activity has been proposed as a screening assay for DNT (Hogberg et al., 2011; Robinette et al., 2011; Dingemans et al., 2016), and our laboratory recently demonstrated that by using multi-well MEAs (mwMEAs) a medium-throughput assay is feasible (Brown et al., 2016; Frank et al., 2017).

Our laboratory recently reported results using this medium-throughput mwMEA-based DNT assay for 86 compounds, demonstrating that 49 of 60 (82%) compounds with evidence of developmental neurotoxicity in the published literature altered development of at least one network parameter (Frank et al., 2017). These data also were used to develop estimates of potency (effective concentration that caused a 50% change; EC₅₀ value) for those compounds that altered network development (64/86). Although such information is useful for prioritization for additional testing, relating *in vitro* perturbations to *in vivo* adversity is a key challenge for 21st century toxicology (Keller et al., 2012). To address this issue, Shah et al. (2016) developed a novel systems toxicology approach to identify toxicological “tipping points” between adaptation and adversity.

Toxicological “tipping points” are defined by Shah et al. as dose-dependent transitions in cells based on their inability to recover to normal (or basal) functions. To analyze tipping points they examined data on the effects of almost a thousand ToxCast chemicals in HepG2 cells using multiple high-content imaging (HCI) endpoints over time (0, 1, 24 and 72 h). They translated the temporal response of HepG2 cells to each chemical treatment as cell-state trajectories. Interestingly, all chemical effects on cells could be classified into three main groups: no effect, recovering and nonrecovering trajectories. Recovering trajectories were characterized by a return of cells to their normal (or basal) states. On the other hand, nonrecovering trajectories were assumed to be indicative of a loss of cellular homeostatic capacity. They defined the transition from recovering to nonrecovering trajectories as the tipping point, and the corresponding chemical concentration as the critical concentration. This approach for identifying tipping points developed by Shah et al. is quite general and can be readily applied to other *in vitro* models that involve temporal measurements.

In the present study, we used the data from Frank et al., 2017, to determine tipping points for chemical effects on *in vitro* neural network formation, and to estimate the corresponding critical chemical concentrations. The results indicate that by consideration of a unified framework for network formation, critical concentrations can be determined for many compounds that altered neural network development, and that in many cases, these concentrations are lower than the EC₅₀ values for individual parameters describing neural network function or the composite values for effects on all parameters. Further, using high-throughput toxicokinetic estimates, we demonstrate that the tipping point concentrations for a small subset of chemicals are concordant with plasma concentrations *in vivo* that are associated with developmental neurotoxicity. The results indicate that such an analysis can provide a robust and sensitive determination of the point at which compounds begin to alter neural network development *in vitro*.

2. Methods

2.1. Compounds and methods

The present study represents additional analysis of the data

presented in Frank et al., 2017. Briefly, in that study, concentration-dependent (typically 0.03 to 30 μ M) effects of 86 compounds on neural network development were measured in primary cortical cultures prepared from day old rat pups using mwMEA recordings across 4 time points (Days *in vitro* (DIV) 5, 7, 9 and 12). Sixty of the tested compounds had at least limited literature evidence of DNT effects in humans or animal models (Mundy et al., 2015), 4 additional compounds had been previously tested in this assay (Brown et al., 2016) and found to impact network function (bisindolylmaleimide I, loperamide, mevastatin, and sodium orthovanadate), and 21 compounds were classified as unknowns with insufficient evidence for DNT effects in the published literature. Acetaminophen was used as a negative control. Cultures were plated at 150,000 cells/well in 48 well microelectrode array plates from Axion Biosystems (M768-KAP-48) using standardized culture and plating protocols (Cotterill et al., 2016; Brown et al., 2016), and chemical treatment commenced 2 h after plating of the cells and continued throughout the experiment. Chemicals were renewed when media was refreshed on DIV 5, and 9. As described in complete detail in Brown et al., 2016, electrical activity in the cultures was recorded for 15 min on DIV 5, 7, 9 and 12 using an Axion Maestro amplifier, Middleman A/D conversion, and AXiS 1.9 or later software. Axion raw (*.raw) and spike (*.spk) files were saved to a redundant physical hard drive for later analysis. Culture viability was assessed following the final DIV12 recording using total LDH release and CellTiter-Blue assays as described in Frank et al., 2017.

2.2. Initial data processing and normalization

Axion spikelist files with time-stamped spikes were combined with comma-separated plate layout data for conversion to hierarchical data format (.h5) files. Hierarchical data format files were used as input to the ‘meadq’ (<https://github.com/dianaransomhall/meadq>) and ‘sjemea’ (<https://github.com/sje30/sjemea>) R packages, which together generated a set of 16 network measures per well that were used for downstream analyses with date and plate ID tracking information attached. An additional network parameter, normalized mutual information (Ball et al., 2017), a measure of shared information in the network, was computed for each recording separately. Nine of the 17 network parameters (mean firing rate (MFR), number of active electrodes (#AE), burst rate (BR), number of actively bursting electrodes (#ABE), percent of spikes in bursts (%SiB), number of network spikes (#NS), percent of spikes in network spikes (%SiNS), Pearson's correlation (r) between activity on electrodes, and normalized mutual information (MI)) were selected for incorporation into the critical concentration analysis.

To reduce the impact of any batch effects, raw network parameter values were normalized by dividing by the median of untreated control well values located on the same plate and on the same DIV (5, 7, 9, or 12). If the median of same-plate untreated control wells was zero, all values for that parameter on that plate were set to 1 (no change from controls). The fold-change from control median values were then log-transformed to reduce the impact of outlier values. The mean and standard deviation of log-transformed fold change values of all untreated control wells on the same DIV were then used to z-score scale all fold-change values. This resulted in values centered at zero that indicate the number standard deviations away from the untreated control mean (positive or negative).

2.3. Tipping point analysis

We used the methods described previously (Shah et al., 2016) to identify tipping points by calculating the following four quantities from the data: system perturbation (X), total scalar perturbation ($|X|$), velocity (V) and derivative of the velocity with respect to concentration (∂V_c). A conceptual overview of these quantities is provided in Fig. 1. The scripts used for the analysis of tipping points are available at the

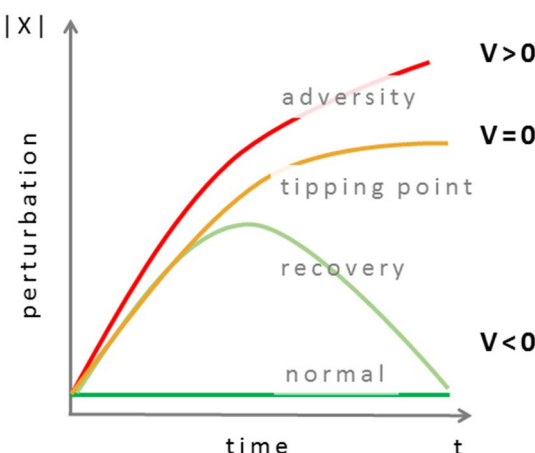


Fig. 1. Understanding tipping points. The graph shows the relationship between scalar system perturbation ($|X|$), time (t) and velocity (V) for different theoretical trajectories. With increasing chemical stimulation, the system deviates from a normal trajectory to recovery ($V < 0$), and then to adversity ($V > 0$). A tipping point is the critical transition between recovery and adversity ($V = 0$).

following doi: <http://dx.doi.org/10.23719/1407690>. First, the z-scores of the nine normalized network parameters (MFR, #AE, BR, #ABE, %SiB, #NS, %SiNS, r and MI) from each well were used to represent the system perturbation at DIV 5, 7, 9, and 12 (denoted as X_5 , X_7 , X_9 , and X_{12} , respectively) for each treatment replicate. Second, total scalar perturbation values ($|X_5|$, $|X_7|$, $|X_9|$, and $|X_{12}|$) were calculated from the system perturbation as the Euclidean norm. This resulted in strictly positive values representative of the total number of standard deviations away from “normal” network recordings at each time point. The difference between scalar perturbation values at DIV 5, 7, or 9 and the terminal DIV 12 value were divided by the number of days elapsed to calculate a numerical derivative of change in scalar perturbation over change in time (denoted as V_5 , V_7 , V_9 , and V_{12}). These values were termed “velocities” of the system and plotted as individual well velocities over increasing concentration of compound. The velocity over concentration relationship for all replicates was approximated with a localized regression function (loess function in R v3.2.5; R Core Team, 2016). The derivative of the velocity with respect to concentration (denoted as ∂V_c) for each compound was then taken to search for a critical concentration at which the slope of velocity changes from negative to positive.

2.4. Critical concentration determination

A search for a critical concentration was triggered when a compound had at least one concentration that produced a terminal DIV 12 scalar perturbation beyond 3 times the median absolute deviation (MAD) of untreated control wells (indicating that the compound perturbed the system beyond random variation). Both negative and positive velocity values along the concentration curve (indicating both recovery and non-recovery trajectories) also were required to identify a critical concentration. If the highest concentration of compound produced negative velocity derivative values, these values were excluded from consideration as they were found to represent a tipping out of scalar perturbation values at total network failure (associated with gross cell loss). The concentration at the last $V = 0$ crossing point of the velocity derivative (going from negative to positive) was then taken as the critical concentration. If multiple $V = 0$ crossings occurred, the greater concentration was selected as the critical concentration to be more conservative in estimating compound effects.

Uncertainty in the critical concentration value was estimated by randomly leaving out one replicate for each concentration of a compound and estimating a new critical concentration for the incomplete

dataset using the above-described method. This process was repeated 1000 times and the 95% confidence interval was taken as the range of 950 out of the 1000 sampled values.

2.5. Network parameter inclusion

In Frank et al., 2017, a total of 17 different parameters were determined from the recordings of network activity. However, 8 of these parameters were derived from nine “empirical” parameters that were collected directly from the recording (e.g. burst duration is undefined in the absence of bursting). To determine whether parameter inclusion would substantially change interpretation of network trajectories, a collection of 9 out of the 17 total network parameters were randomly selected 100 times and scalar perturbations over time were calculated for each compound as described above. Confidence intervals (95%) were then estimated at each time point from the central 95 out of 100 median scalar perturbation trajectories. These ranges were qualitatively assessed to determine whether overall system recovery and non-recovery trends remained present with different network parameter inclusion.

2.6. Comparison to cytotoxicity and individual network parameter effects

The critical concentrations derived herein were compared to the EC₅₀ values for chemical effects on individual network parameters and viability that were published previously in Frank et al., 2017. The EC₅₀ values were determined by calculating the area under the curve (AUC) for each well over time, as described in Brown et al., 2016 and Frank et al., 2017. Briefly, the temporal component of parameters describing network function (e.g. MFR, #AE, etc.) from each well's recordings was reduced to a single area under the curve (AUC) value by calculating the trapezoidal area under the curve for each treatment concentration. Raw AUC values were normalized by same plate untreated control means. For chemicals that produced AUC values exceeding 3 × the MAD of untreated control wells for any concentration, 4-parameter log-logistic curves were fit to model concentration-response relationships.

Raw cell viability assay values from alamar blue and lactate dehydrogenase (LDH) assays were normalized by subtracting blank well controls and dividing by the median of untreated well absorbance. Concentration-response relationships and the concentrations that produced a median 50% or greater reduction in cell viability value were determined. Since these assays were conducted at only a single time point, AUC values were not determined. The lower of the two concentrations identified by cytotoxicity assays (LDH release or reduction of resazurin) was used for comparison with tipping point concentrations.

2.7. Comparison to *in vivo* results using toxicokinetics

The R package “httk” (Pearce et al., 2017) includes both chemical-specific *in vitro* data and models for predicting *in vivo* toxicokinetics (i.e., absorption, distribution, metabolism, and excretion). Six study chemicals with literature *in vivo* data on doses causing neurological effects in laboratory animals (see Table 1) following developmental exposure also were available for simulation with “httk”. These six chemicals had chemical specific *in vitro* toxicokinetic measurements and physico-chemical properties (i.e., molecular weight, hydrophobicity, and ionization equilibria). Where rat-specific *in vitro* toxicokinetic data were available, these were used; otherwise rat physiology and human *in vitro* data were used. Human *in vitro* data were used for all mouse predictions. Version 1.8 was used with non-restrictive metabolism assumed (i.e., that chemical bound to protein rapidly dissociates to be metabolized in the liver). The function “solve_pbtk” was used to model a generic physiologically-based toxicokinetic (PBTK) model featuring liver, kidney, gut, venous, arterial, and rest of body compartments. Time-integrated plasma concentration

Table 1
Studies used for comparison to in vivo data.

Substance_CASRN	Chemical	DSSTox_Substance_Id	Species	Exposure	Dose_Route	Endpoint	References	Dose
2921-88-2	Chlorpyrifos*	DTXSID4020458	Rat	Postnatal	1–5 mg/kg, s.c.	Neurochemistry, behavior	Dam et al., 2000	3
2921-88-2	Chlorpyrifos*	DTXSID4020458	Mouse	Pre/postnatal	3–6 mg/kg, oral	Behavior	Ricceri et al., 2006	4.5
439-14-5	Diazepam	DTXSID4020406	Rat	Prenatal	2.5 mg/kg, s.c.	Behavior	Nicosia et al., 2003	2.5
439-14-5	Diazepam	DTXSID4020406	Rat	Pre/postnatal	10 mg/kg, s.c.	Behavior	Frieder et al., 1984	10
50-53-3	Chlormazazine	DTXSID0022808	Rat	Prenatal	3 mg/kg, s.c.	Behavior	Clark et al., 1970	3
50-53-3	Chlormazazine	DTXSID0022808	Rat	Pre/postnatal	15 mg/kg, i.m.	Morphology	Hannah et al., 1982	15
56-53-1	Diethylstilbestrol	DTXSID3020465	Mouse	Prenatal	0.1 μg, s.c.	Behavior	Mihalick, 2003	0.1
56-53-1	Diethylstilbestrol	DTXSID3020465	Mouse	Prenatal	0.1 μg/kg, oral	Behavior, neurochemistry	Kaitsuwa et al., 2007	0.1
58-25-3	Chlordiazepoxide*	DTXSID4046022	Rat	Pre/postnatal	1 mg/kg, i.p.	Behavior	Adams, 1982	1
58-25-3	Chlordiazepoxide*	DTXSID4046022	Mouse	Prenatal	10–20 mg/kg, s.c.	Behavior	Kurishigai et al., 1992	15
60-57-1	Dieldrin	DTXSID9020453	Rat	Pre/postnatal	0.35 mg/kg, oral	Behavior	Olson et al., 1980	0.35
60-57-1	Dieldrin	DTXSID9020453	Rat	Prenatal	10 mg/kg, i.p.	Neurochemistry	Brannen et al., 1998	10
Substance_CASRN	Dose_Units	Route	Days	Cmax	AUC	Critical.concentration	Lower.95.CI	Higher.95.CI
2921-88-2	ng/kg	s.c.	4	405.782	0.291206	1.23	0.072	1.828
2921-88-2	ng/kg	oral	4	0.955984	0.119065	1.23	0.072	1.828
439-14-5	ng/kg	s.c.	8	292.2317	0.138648	9.766	7.405	16.267
439-14-5	ng/kg	s.c.	7	1168.927	0.554593	9.766	7.405	16.267
50-53-3	ng/kg	s.c.	1	220.0537	0.102801	0.476	0.267	0.762
50-53-3	ng/kg	i.m.	4	0.514056	0.476	0.267	0.762	0.762
56-53-1	μg	s.c.	10	0.020638	7.79E-06	2.675	0.06	4.015
56-53-1	μg	oral	7	1.63E-07	1.38E-07	2.675	0.06	4.015
58-25-3	ng/kg	i.p.	21	168.5788	0.127206	2.454	1.828	2.83
58-25-3	ng/kg	s.c.	10	1533.209	1.101632	2.454	1.828	2.83
60-57-1	μg/kg	oral	17	1.49E-05	2.56E-06	1.035	0.347	2.177
60-57-1	ng/kg	i.p.	6	1148.892	0.561138	1.035	0.347	2.177

(area under the curve or AUC) was calculated for the species, doses, number of days, and administration routes as indicated by Table 1. Due to software limitations, intraperitoneal, subcutaneous, and intramuscular injection routes were all modeled as intravenous doses.

3. Results

3.1. Calculation of network scalar perturbations

Complex primary cultures isolated from newborn rat cortex exhibit a reproducible developmental trajectory as neurite outgrowth leads to synaptogenesis and increasing network activity over approximately two weeks *in vitro* (Cotterill et al., 2016). mwMEA recordings over this time period provide multiparametric data that includes measures of spike, burst, and network synchrony in each culture. Our previous work demonstrated that many compounds that are developmentally neurotoxic in mammals perturb these parameters (Frank et al., 2017). Because of the temporal nature of our previous data, it was possible to determine a critical concentration at which a given compound began to perturb the developing system beyond recovery. Such concentrations may represent the transition point in the series of events that results in developmental neurotoxicity. To determine these tipping points, it was first necessary to transform network parameter data into values that define a distance from normal activity range. Importantly, normal network values change across developmental time as activity and synchrony increase between DIV5 and DIV12, so a normal range needs to be redefined at each time point. To reduce variation attributed to cortical culture batch and plate-to-plate variability, the 17 individual network parameters quantified from recordings were first divided by the median value of untreated control wells run on the same plate and same DIV to obtain fold-change values. Log-transformed fold-change values were then standardized by subtracting the mean fold-change and standard deviation of the same DIV untreated control wells. Standardizing to the distribution of all untreated control wells on each DIV yielded z-score values that reflected the number of standard deviations away from normal at each time point; ensuring that all network parameters were measured on a common scale (Fig. 2).

Examination of z-score values over time revealed a number of different trajectories a network might take depending on the concentration of compound present. For a compound that did not appreciably impact network activity, such as acetaminophen, all concentrations resulted in network parameter trajectories that vary within the normal range of controls (Fig. 2B). In contrast, for compounds that significantly perturbed normal network development, such as haloperidol, a concentration-response relationship was often observed whereby the z-score perturbation exits the normal range of controls in a positive or negative direction at certain timepoints and this departure becomes more pronounced with increasing concentration (Fig. 2C). Interestingly, it was noted that intermediate concentrations of many compounds, including haloperidol, produce z-score perturbations that exit the normal range at earlier recording time points, but return to fall within normal range by DIV12. This occurred in the presence of compound exposure, which was from 2 h after plating through DIV12.

By integrating data from multiple measures of network activity, it was reasoned a more comprehensive and reliable metric of network function could be defined. To this end, the total perturbation of the cortical network was defined by the sum of individual network parameter perturbations observed. In particular, the Euclidean norm of 9 network parameters was selected at each time point to equally weight values that depart from the normal range in both the positive and negative direction. This resulted in total system scalar perturbation over time trajectories that were visualized for each replicate and summarized by localized regression curves (Fig. 3). For example, the total scalar perturbation of acetaminophen remained below the shaded uncertainty range of $3 \times$ the median absolute deviation (MAD) of untreated control wells across all timepoints (Fig. 3A). In contrast, the scalar perturbation

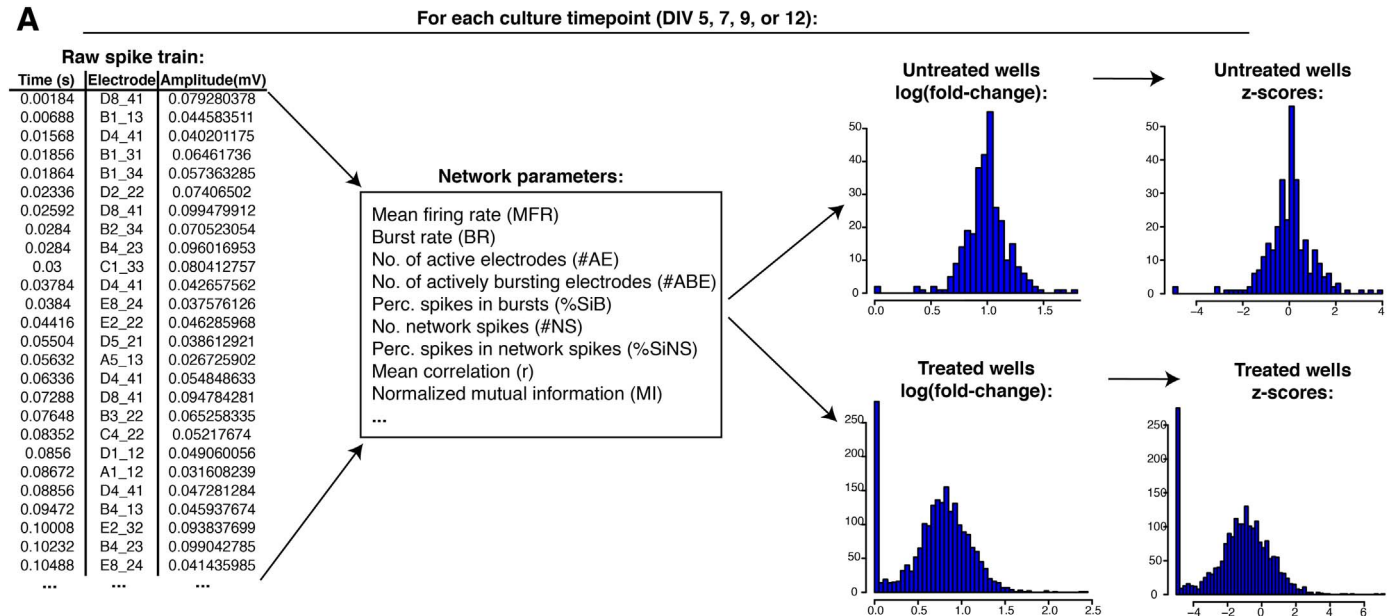
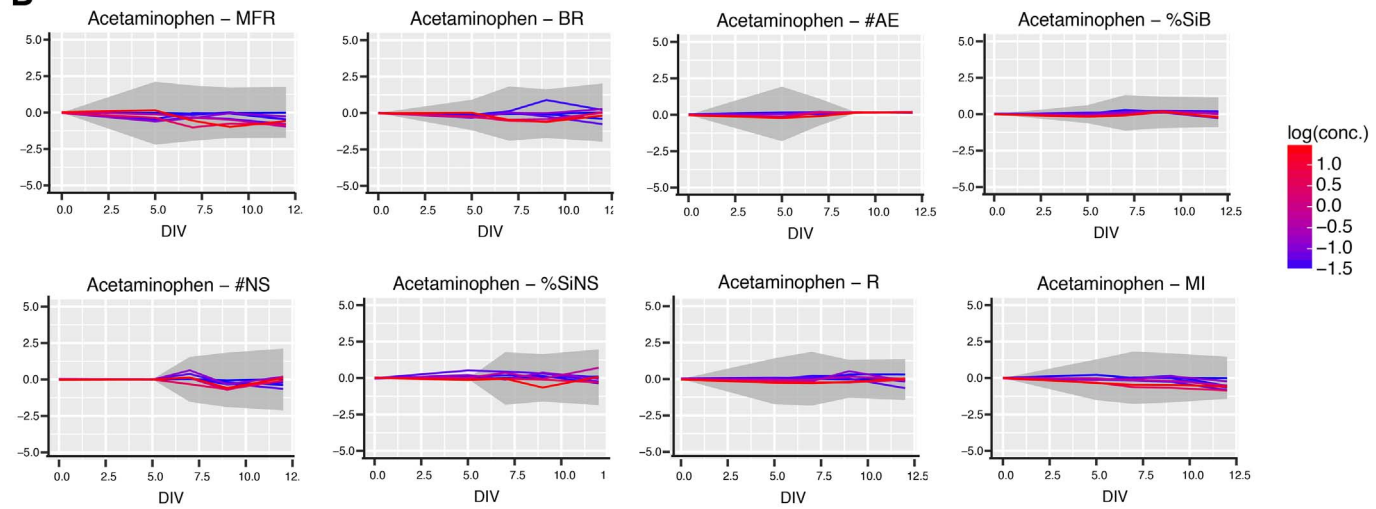
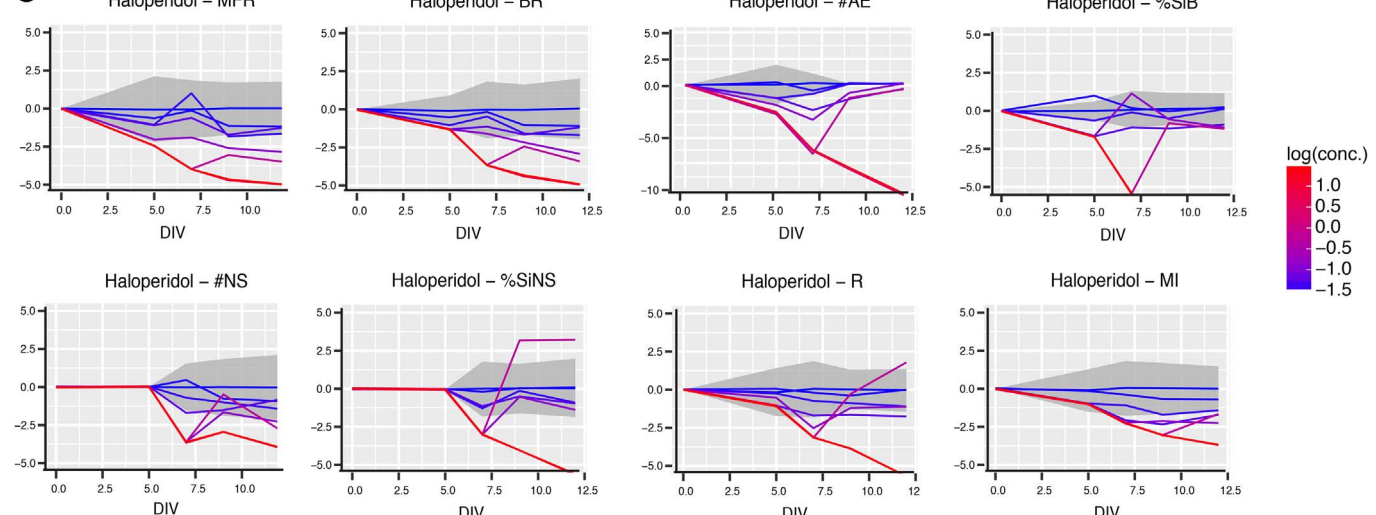
of haloperidol exposure increased with increasing haloperidol concentration and extended well past the $3 \times$ MAD limit (Fig. 3B). Interestingly, the two highest concentrations of haloperidol tested (10 and 30 μ M) caused apparent network failure (maximal scalar perturbation), while lower concentrations (0.3–3 μ M) exhibited some recovery toward the normal range by DIV 12, reflecting the similar trends observed in the individual parameter z-score transformations. Graphs showing Z-scores and total scalar perturbations are available for all compounds at the following doi: <http://dx.doi.org/10.23719/1407690>.

The decision to model system scalar perturbation as a function of 9 of the 17 network parameters measured was made for two reasons. First, the 9 selected parameters (MFR, #AE, BR, #ABE, %SiB, #NS, %SiNS, r, and MI) are directly calculated from the recording spike trains, as opposed to the other 8 network parameters that are dependent upon non-zero values for one of the selected parameters. For example, the mean network spike duration (NSD) is dependent upon the observation of at least one network spike during the recording time, else it remains undefined. At earlier time points in development, the cortical cultures have little or no network spiking and bursting activity and this results in many undefined values for the dependent parameters that could introduce extra uncertainty into the analysis. Second, it was found that random selection of 9 out of the 17 network parameters for calculation of total system perturbation did not qualitatively alter the overall concentration-dependent trajectories of network scalar perturbation (data not shown). That is, concentrations that showed a pattern of escalating scalar perturbation over time or exhibited a recovery trend with the 9 selected parameters, produced a similar pattern with random selection of network parameters. Together, these results suggested that inclusion of the additional 8 network parameters would introduce additional uncertainty into the analysis without contributing to sufficiently different trajectories that would meaningfully change interpretation of recovery and non-recovery trends.

3.2. Determination of critical concentrations

Determination of a critical concentration was based on the premise that a developing cortical network is in some ways capable of adapting to the presence of certain compounds at particular concentrations (*i.e.*, overcoming the initial perturbation). Therefore, after calculation of scalar perturbation values, it was important to first establish that instances of network recovery could be identified in the assay. If the median scalar perturbation of a treatment group exceeded three times the MAD of untreated control wells at DIV 5, 7, or 9, but then returned to a median value closer to zero at the final time point (DIV 12), this was taken as preliminary evidence of system recovery in the presence of a given compound at a defined concentration. Applying these criteria, evidence of system recovery was observed in 54 out of the 86 compounds (62.8%) for at least one concentration (Supp. Table 1). The presence of these scalar perturbation trajectories indicated the assay is capable of modeling treatments where neural networks are disrupted at one stage of *in vitro* development but exhibit a return to normal range at a later time.

To further model the dynamics of system recovery and failure in the presence of tested compounds, the “velocity” of scalar perturbation values was calculated between DIV 5, 7, or 9 and DIV 12 to determine the transition from a system recovery to a non-recovery trajectory. The final recording time point was chosen as a consistent anchor because it provided the strongest evidence in the assay of final system status as network development concluded (*i.e.* DIV12 status best reflected whether the network resembled a normal untreated network at the conclusion of *in vitro* development). System velocities were then plotted over increasing concentration of each compound to visualize the relationship (Fig. 4). The derivative of velocity with respect to concentration was taken to identify the point in the velocity curve where the slope changed from negative to positive, indicating a transition from a system recovery to a non-recovery trajectory. In order to focus

A**B****C**

(caption on next page)

Fig. 2. Conversion of network parameters to normalized z-scores. A) Individual network parameters were calculated from raw spike train data from recordings, normalized to same-plate untreated well medians, then log-transformed and converted to z-scores (± 1 z-score = ± 1 standard deviation from the untreated well mean value) to put all network parameters on the same scale. Example of log-transformed and z-score distributions for untreated and treated wells are shown with histograms on right. B and C) Examples of z-scores over time for acetaminophen (B) and haloperidol (C). In each plot, the shaded gray area represents the variability of untreated control wells (3 times the median absolute deviation) and colored lines show median z-score values over time for varying concentrations of compound for one network parameter. The 8 network parameters shown are mean firing rate (MFR), burst rate (BR), number of active electrodes (#AE), percent of spikes in bursts (%SiB), number of network spikes (#NS), percent of spikes in network spikes (%SiNS), mean correlation (R), and mutual information (MI).

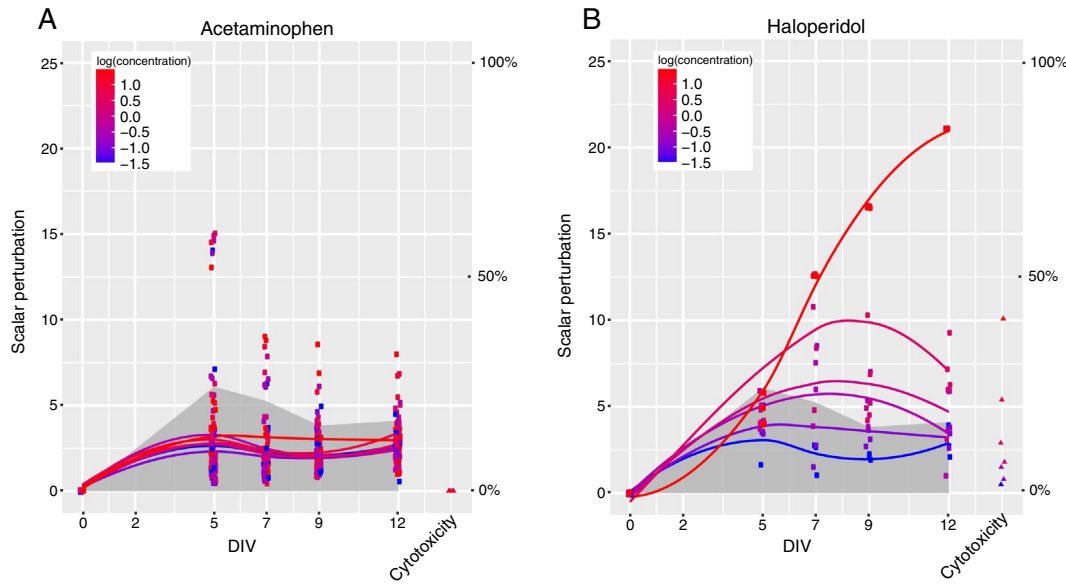


Fig. 3. Total Scalar Perturbations integrate changes across network parameters. Total scalar perturbations (sum of individual network parameter z-scores) for varying concentrations of acetaminophen (A) and haloperidol (B) are shown as individual data points and best fit lines colored by concentration over time *in vitro*. Shaded gray area represents 3 times the median absolute deviation of untreated control wells at each timepoint.

on compounds with a more reproducible impact on network activity and avoid overfitting subtle trajectories, a compound was required to have at least one concentration where the DIV 12 scalar perturbation exceeded three times the MAD of untreated control networks. The minimum concentration derived from the three time windows profiled for each compound was selected to accommodate different trajectory shapes and identify the lowest concentration with evidence of a tipping point and possible adverse effect. In total, a point of zero velocity slope (where trajectory transitioned from recovery to non-recovery) was found for 42 out of the 86 compounds, with critical concentrations ranging from $0.005\text{ }\mu\text{M}$ for bis(*tri-n*-butylin) oxide to $9.77\text{ }\mu\text{M}$ for diazepam (Table 2). Uncertainty in these critical concentrations was estimated by a 95% confidence interval derived from iterative random sampling and recalculation of the critical concentration with available replicates.

The same compound often produced multiple tipping points depending on the time interval considered (e.g. DIV 5–12 vs DIV 9–12). Tipping points with uncertainty ranges spanning two orders of magnitude or greater were discarded as unreliable estimates. These criteria applied to 9 out of the 42 compounds, but in no case did it remove all tipping point estimates for a compound (Supp Table 2). To interrogate the magnitude of tipping point concentration differences between different time windows, the fold-change between the most discordant pair of tipping point estimates was calculated for each compound with more than one estimate. The average fold-difference among the 33 compounds with multiple estimates was found to be 2.21 (Supp Table 2). Considering the concentration series employed in this assay was incremented by half-log units, these results showed relatively high consistency between tipping points derived from different time intervals for the same compound.

3.3. Critical concentration comparison to cytotoxicity and individual network parameters

To evaluate whether the tipping point concentrations determined for the 42 compounds corresponded to treatments that also resulted in high general cytotoxicity, tipping points were compared to compound concentrations that produced a 50% reduction in cell viability levels (EC_{50}) from resazurin reduction and LDH release assays (Frank et al., 2017) (Fig. 5A). Only two out of the 42 compounds (4.8%), emamectin benzoate and mevastatin, exhibited high cytotoxicity at concentrations lower than the tipping point estimate. For all other compounds, there was at least a 4-fold difference between the tipping point and cytotoxicity EC_{50} . The mean difference between 50% cell viability (where defined) and tipping point concentration was 13.8-fold. This suggests that tipping points often occur at concentrations substantially lower than those causing high levels of cell viability loss in the cortical networks.

Previous analyses of these same data used an area under the curve (AUC) metric to evaluate concentration-response relationships for each compound (Frank et al., 2017). The AUC approach determined EC_{50} concentrations where an exposure was sufficient to reduce individual network parameters below 50% of control values. AUC-derived EC_{50} concentrations for the 9 network parameters used for scalar perturbation calculations were compared to tipping point concentrations in two ways. The first approach was to determine where the minimum of the 9 AUC-derived EC_{50} values fell in relation to the tipping point concentration. The second was to compare the median of the 9 AUC-derived EC_{50} values to the critical concentration for each compound. Both values are shown in relation to the tipping point concentration in Fig. 5B. For the majority of compounds (27/42, 64.3%), the tipping

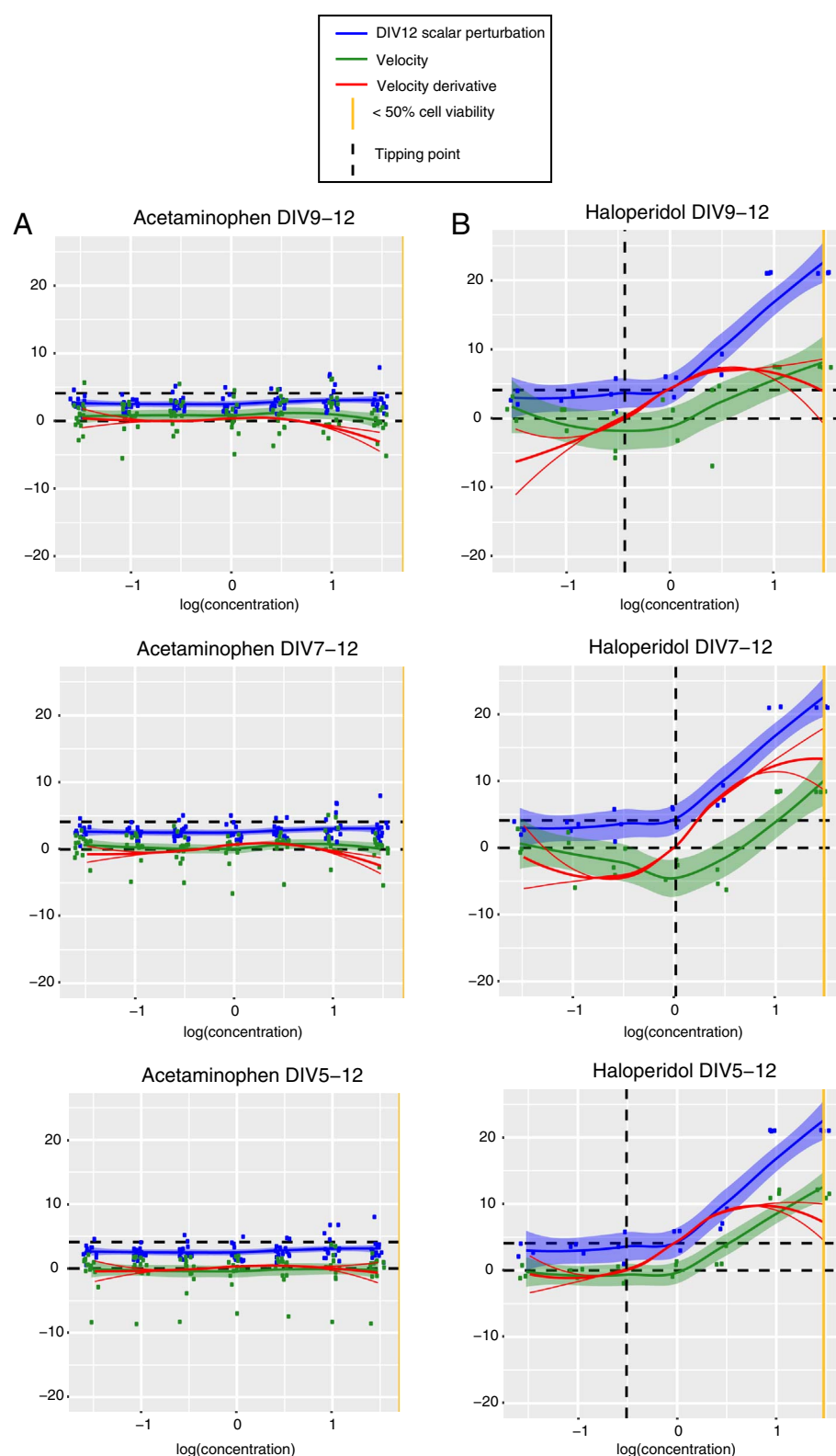


Fig. 4. Examples of Tipping Point determinations for acetaminophen and haloperidol. Plots show DIV 12 scalar perturbation (blue line), as well as velocity of change (green line) and derivative of that velocity curve (dV/dc ; red line) from DIV 9 (top), DIV 7 (middle), or DIV 5 (bottom) for varying concentrations of acetaminophen (A) or haloperidol (B). Horizontal dotted lines demarcate 3 times median absolute deviation of untreated control wells. Vertical dotted line indicates tipping point concentration, defined as the derivative of velocity over concentration transitioning from negative to positive values. For comparison, the yellow vertical line indicates the lower EC_{50} value for cell viability loss. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

point concentration was equal to or lower than the minimum AUC-derived EC_{50} values identified. For all but four compounds (emamectin benzoate, mevastatin, PBDE-47, and diazepam), the critical concentration was below the median AUC-derived EC_{50} values. This result suggests that in most cases, tipping points serve as a more sensitive measure of network developmental effects than looking for individual

network parameter changes. Interestingly, in the cases where the minimum AUC-derived EC_{50} values fell below the critical concentration, 9 out of 15 times it was the number of network spikes parameter that produced the lowest EC_{50} . The number of network spikes may therefore be considered the most sensitive of the 9 parameters examined.

Table 2
Critical concentrations.

Compound	Timepoint comparison	Critical concentration (μM)	Lower 95% CI	Higher 95% CI
Deltamethrin	DIV5 to DIV12	0.05	0.036	0.145
Cytosine Arabinoside	DIV7 to DIV12	0.046	0.036	0.06
DEHP	DIV9 to DIV12	0.11	0.055	0.224
5-Fluorouracil	DIV5 to DIV12	0.142	0.102	0.172
Haloperidol	DIV5 to DIV12	0.309	0.145	0.492
Loperamide	DIV5 to DIV12	0.13	0.111	0.145
Chlorpyrifos	DIV5 to DIV12	1.23	0.072	1.828
Chlorpyrifos oxon	DIV9 to DIV12	0.071	0.039	0.133
Diazepam	DIV5 to DIV12	9.766	7.405	16.267
Thiouracil	DIV9 to DIV12	2.251	0.102	5.696
t-Retinoic acid	DIV5 to DIV12	1.593	0.158	2.83
Trimethyltin	DIV5 to DIV12	0.015	0.012	0.022
Dieldrin	DIV9 to DIV12	1.035	0.347	2.177
Lead acetate	DIV5 to DIV12	2.916	1.828	4.382
Cadmium	DIV5 to DIV12	0.036	0.036	0.043
Methylmercury	DIV5 to DIV12	0.037	0.029	0.041
Bis1	DIV5 to DIV12	0.631	0.015	0.943
Domoic Acid	DIV5 to DIV12	0.205	0.089	0.254
Sodium Orthovanadate	DIV5 to DIV12	4.119	3.371	5.696
Cypermethrin	DIV9 to DIV12	0.283	0.188	0.379
Emamectin benzoate	DIV7 to DIV12	0.078	0.039	0.122
Fipronil	DIV9 to DIV12	0.799	0.158	3.089
Flusilazole	DIV9 to DIV12	0.085	0.043	0.291
Methylchloroisothiazolinone	DIV9 to DIV12	0.436	0.245	0.64
Carbaryl	DIV5 to DIV12	0.238	0.051	1.828
Heptachlor epoxide	DIV5 to DIV12	1.737	1.288	1.995
Lindane	DIV5 to DIV12	0.799	0.267	4.782
Mevastatin	DIV9 to DIV12	0.476	0.145	1.535
Diethylstilbestrol	DIV5 to DIV12	2.675	0.06	4.015
Isoniazid	DIV5 to DIV12	0.13	0.06	0.224
PBDE-47	DIV5 to DIV12	1.155	0.302	2.262
Hexachlorophene	DIV5 to DIV12	0.155	0.188	0.537
Tebuconazole	DIV5 to DIV12	1.341	0.122	4.382
Aminonicotinamide	DIV7 to DIV12	0.733	0.379	1.082
Bis tri n butyl oxide	DIV5 to DIV12	0.005	0.004	0.006
Chlordiazepoxide	DIV5 to DIV12	2.454	1.828	2.83
Permethrin	DIV9 to DIV12	0.871	0.64	1.288
Paraquat	DIV9 to DIV12	0.142	0.102	0.205
Sodium Arsenate	DIV9 to DIV12	0.092	0.055	0.122
Fluoxetine	DIV5 to DIV12	0.565	0.379	0.762
Chlorpromazine	DIV5 to DIV12	0.476	0.267	0.762
Triethyltin bromide	DIV5 to DIV12	0.022	0.017	0.024

3.4. Comparison to *in vivo* results using toxicokinetics

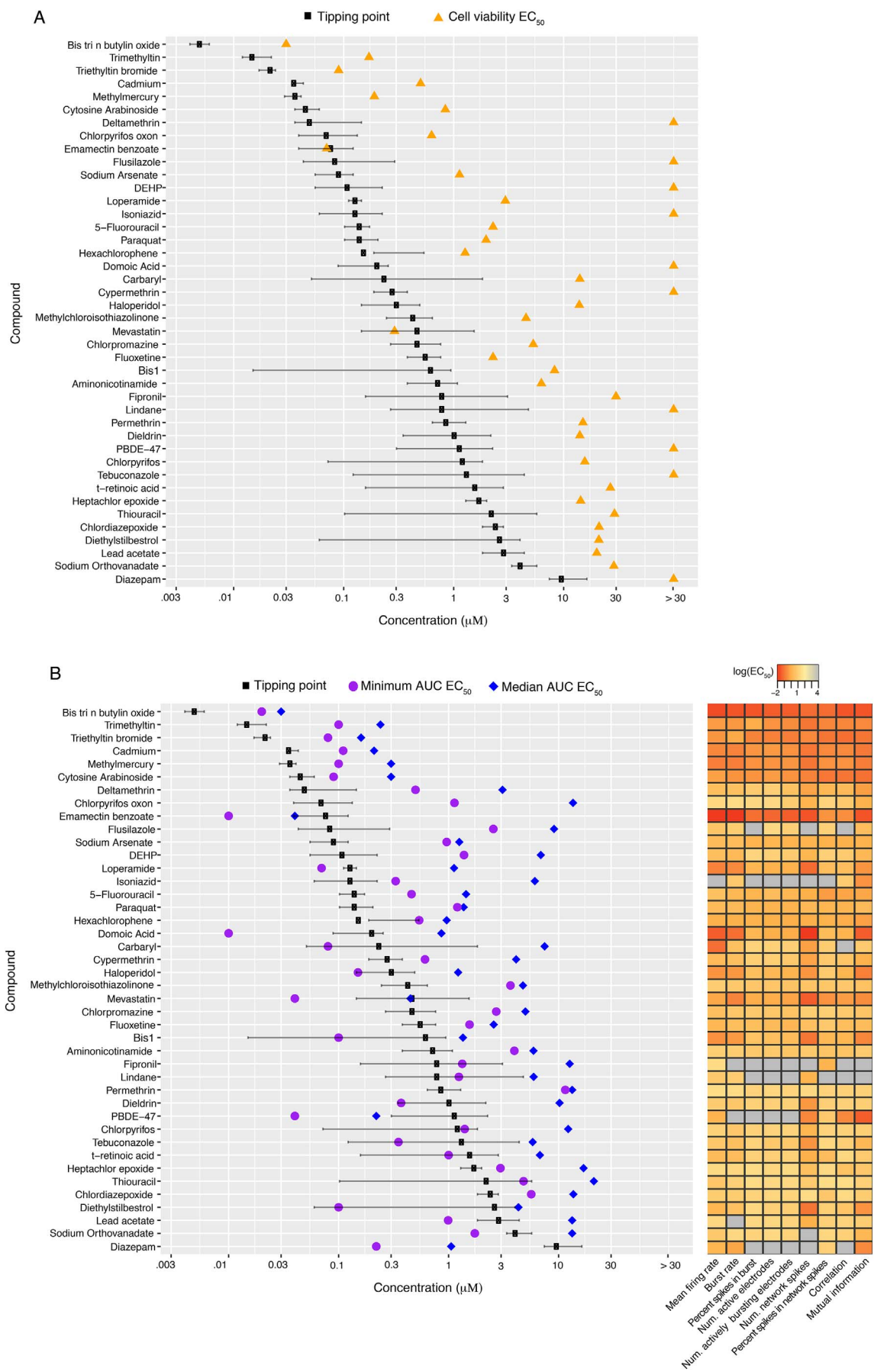
To determine how the tipping point concentrations compare to plasma levels associated with reports of neurotoxicity following exposure during development, *in vitro* concentration levels were converted to estimated plasma levels using the high-throughput toxicokinetic (httk) package in R. For an *in vivo* comparator, studies were selected from Table 1 in Mundy et al., 2015, as the studies in that publication were cited as examples of compounds with evidence for developmental neurotoxicity. There were sufficient *in vitro* toxicokinetic data available for this package for only 6 of the 42 chemicals with tipping point concentrations. Complete information on the compounds, dose levels and model estimates are provided in Table 1. When comparing the *in vivo* AUC concentration estimated with htk to the *in vitro* predicted critical concentration, the data indicate close agreement for five of the six compounds (Fig. 6). The exception was diethylstilbestrol, for which the critical concentration was higher than the *in vivo* estimate. There were also discordant results with dieldrin where the tipping point was less potent than one *in vivo* study; this was due to 3000-fold differences in the *in vivo* doses used.

4. Discussion

The present data demonstrate that, “tipping points” associated with alterations in network formation can be determined for a subset of

compounds recently tested in Frank et al., 2017. For a subset of these compounds where enough data were available, it was possible to use high-throughput toxicokinetic (httk) model to compare *in vitro* tipping points to blood levels associated with neurodevelopmental outcomes in animal models; these comparisons indicated that the *in vitro* alterations in network function occurred at concentrations that are concordant with *in vivo* developmental neurotoxicity following pre- or perinatal exposure to these compounds. Thus, the tipping point concentrations provide a sensitive indicator of biological alterations in developing neural networks that may be relevant to adverse neurodevelopmental outcomes following *in vivo* exposures to chemicals.

The need to develop a screening battery of tests to assess the potential for a compound to cause developmental neurotoxicity is now well recognized (Bal-Price et al., 2010, 2015; Fritzsche et al., 2017). Several *in vitro* assays have now been developed for a number of process critical to nervous system development, including proliferation (Breier et al., 2008), neurite outgrowth and synaptogenesis (Harrill et al., 2010, 2011; Stiegler et al., 2011), migration (Zimmer et al., 2014; Nyffeler et al., 2017), and apoptosis (Druwe et al., 2015). Assessment of function following exposure during *in vitro* development of neural networks using MEAs (Hogberg et al., 2011; Robinette et al., 2011; Brown et al., 2016; Dingemans et al., 2016; Frank et al., 2017) is unique among these assays in that it can provide both functional and temporal measurements from the same network. This facilitates the determination of whether the system recovers or deviates from homeostasis over the



(caption on next page)

Fig. 5. Comparison of tipping points to cytotoxicity and individual network parameters. Tipping points could be determined for 42 compounds, and are compared to A) concentrations that impact cell viability (orange triangles, lower EC₅₀ between total LDH and alamar blue assays) and B) the minimum (purple circles) and median (blue diamonds) EC₅₀ value for individual network parameters based on Area Under the Curve (AUC) calculations. Heatmap on right in panel B shows AUC EC₅₀ values for each network parameter included in tipping point determination for reference. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

measured time span, and thus the resultant data are well-suited for determination of tipping points.

Neural network activity as measured by MEAs can be described by multiple parameters that include measurements of the overall activity (e.g. mean firing rate, number of active electrodes), bursting (e.g. burst duration, burst rate) and coordinated activity (e.g. correlation coefficient, mutual information), most of which show clear “developmental” changes with time *in vitro* (Cotterill et al., 2016). In the present study, tipping points were determined based on chemical effects on 9 of these parameters because they can be derived directly from the spike train. Although other parameters also can be assessed, preliminary analysis using additional parameters indicated that this would have increased the uncertainty in the tipping point analysis. In addition, the parameters selected were among those identified as being more impactful in separating treatments that alter network function from those that do not using Random Forest analyses (Brown et al., 2016; Frank et al., 2017). Thus, the tipping points calculated here represent a composite value for effects on the 9 parameters included. Tipping points could be determined for 42 of the 64 compounds that perturbed activity of neural networks following exposure during network development (Frank et al., 2017). In all but 2 of these cases (mevastatin and emamectin), the calculated tipping point is more potent than the corresponding EC₅₀ value for cytotoxicity, indicating that these changes reflect alterations in network function rather than simply decrements in cellular viability. Further, for 23 of the 42 compounds, the tipping point was lower than the EC₅₀ for the most sensitive of these parameters; while only 4 of the 42 compounds had tipping points that were higher than the median EC₅₀ of individual parameters of network function.

Thus, even though they are based on an “averaged” response, these data indicate that the tipping point calculations are sensitive measures of when and at what concentrations network formation begins to be irrevocably altered by chemical exposure.

In Toxicity Testing in the 21st Century: Strategy and Vision, the use of cell-based toxicity testing methods was encouraged (NRC, 2007). Important to this concept was the ability to determine differences in adaptive responses (homeostatic responses) *versus* those responses that led toward a “toxicity pathway”. Tipping points are one method for determining the adaptive *vs* adverse responses, and in the present case, the tipping points represent the concentration at which development of network function no longer proceeds according to its normal trajectory *in vitro*. Although speculative, an additional point to consider in the present study is whether or not the “adaptive” responses just below the tipping points (concentrations at which “recovery” is observed), might also ultimately lead to adverse responses. The importance of electrical activity in development of the nervous system is well established (Kerschensteiner, 2014; Babij and De Marco Garcia, 2016). Further, the timing of electrical activity during nervous system development is also crucial, and delays in electrical activity have been associated with neurodevelopmental disorders such as attention-deficit hyper activity syndrome (McLaughlin et al., 2010). Thus, the possibility exists that even in cases where some recovery of network activity to normal response levels occurs, the excursion out of normal parameters may disrupt the timing of other important event sequences, resulting in latent or delayed effects on nervous system function. Such effects may only become apparent later in life or only when the nervous system is challenged in a way that uncovers the deficit. Of note, the present

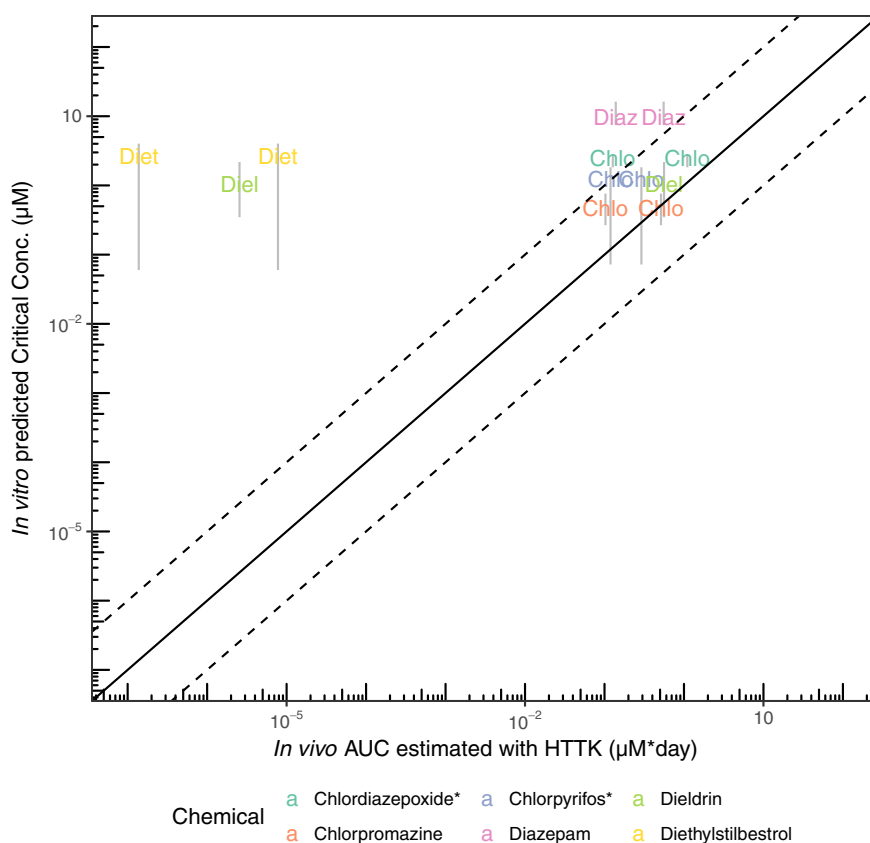


Fig. 6. Comparison between predicted plasma levels for critical concentrations and *in vivo* estimates from the htk model. For those chemicals with 1) *in vitro* predicted critical concentrations, 2) *in vivo* studies indicating neurological effect, and 3) available toxicokinetic data the time-integrated plasma concentration (area under the curve or AUC) was predicted for the LOEL associated with each chemical-specific study. The chemical-specific prediction is indicated by the first four letters of each chemicals name. There were two available studies for each chemical. The identity (“perfect predictor”) line is indicated by a solid black line, while the dashed lines indicate ten-fold above and below perfect prediction. Because all *in vitro* treatments were exposed for the same amount of time, the relationship between nominal *in vitro* concentration and time-integrated concentration is a constant.

experiments stopped on DIV 12, and that network function was not assessed or challenged after this time. As such, it remains to be determined whether or not adaptive responses in the present study represent true recovery, or a change in state that might lead to adverse responses at a later time.

In vitro data will be useful for decision-making beyond screening and prioritization if there is confidence that the concentrations at which effects are observed *in vitro* have relevance to blood levels that cause neurodevelopmental effects *in vivo*. In order to examine how the tipping points determined herein compare to dose levels associated with neurodevelopmental toxicity, compounds from Table 1 in Mundy et al. (2015) were selected as representative *in vivo* exposures that were associated with a DNT outcome. High-throughput toxicokinetics were then used to estimate relevant blood levels for the 6 compounds for which sufficient data existed. When compared to the tipping point concentrations, in 5 of 6 cases, the tipping points for effects on neural network function were equal to or lower than doses associated with DNT outcomes. Interestingly, diethylstilbestrol, the one compound where the tipping point did not match the levels associated with DNT outcomes *in vivo*, is a well-known estrogenic compound. Although acute activity in mature networks does show some responsiveness to estrogenic compounds (Strickland et al., 2017), there is no information on whether those effects are mediated by the estrogen receptor or regarding the role of estrogen in maturation of neural networks *in vitro*. Thus, it could be that the *in vivo* effects of diethylstilbestrol are mediated via mechanisms not present in the *in vitro* assay.

Comparisons to chemical concentrations known or predicted to cause developmental neurotoxicity *in vivo* for all of the compounds tested herein would have been desirable. However, important data were not available for all of the test compounds. Never the less, based on this small sample, calculated tipping points indicate effects at relevant doses *in vivo*. Several caveats should be noted regarding this analysis. First, the *in vivo* dose levels were simply those reported in the literature as cited in Mundy et al., 2015; these were not LOAEL values from guideline studies. Thus, it may be that other events or endpoints are more sensitive to perturbation than those examined here, either *in vivo* or *in vitro*. Second, the httk model predictions in some cases (for example the mouse data) relied on substitution of data from another species (human), as species-specific data were not available, or on other assumptions, for example, regarding chemical binding to protein. Third, it should be noted that a relationship between the *in vitro* alternations of network development and the *in vivo* endpoint reported in the studies used is not implied by these comparisons, simply that dose/concentration levels are concordant. In the future, such associations could be strengthened by linking the *in vitro* effects, through a series of key events, to an adverse neurodevelopmental outcome (e.g. through an adverse outcome pathway or AOP). There is a critical need for the development of adverse outcome pathways relevant to developmental neurotoxicity (Bal-Price and Meek, 2017); only a few (~8) AOPs relevant to DNT have been established in the AOPwiki database (<https://aopwiki.org/>).

In summary, the present data demonstrate that toxicological tipping points can be determined for chemical effects on neural network development, and, based on a small set of compounds, that those tipping point values are relevant to *in vivo* dose levels associated with adverse neurodevelopmental outcomes. As such, this combination of approaches could offer high potential as a powerful tool for assessment of potential developmental neurotoxicity for larger numbers of chemicals.

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.taap.2018.01.017>.

Conflict of interest statement

The authors declare no conflict of interest pertaining to this manuscript. Preparation of this document has been funded by the U.S. Environmental Protection Agency. This document has been subject to

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