Drosophila melanogaster as a model for lead neurotoxicology and toxicogenomics research

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Drosophila melanogaster is an excellent model animal for studying the neurotoxicology of lead. It has been known since ancient Roman times that long-term exposure to low levels of lead results in behavioral abnormalities, such as what is now known as attention deficit hyperactivity disorder (ADHD). Because lead alters mechanisms that underlie developmental neuronal plasticity, chronic exposure of children, even at blood lead levels below the current CDC community action level (10 μg/dl), can result in reduced cognitive ability, increased likelihood of delinquency, behaviors associated with ADHD, changes in activity level, altered sensory function, delayed onset of sexual maturity in girls, and changes in immune function. In order to better understand how lead affects neuronal plasticity, we will describe recent findings from a Drosophila behavioral genetics laboratory, a Drosophila neurophysiology laboratory, and a Drosophila quantitative genetics laboratory who have joined forces to study the effects of lead on the Drosophila nervous system. Studying the effects of lead on Drosophila nervous system development will give us a better understanding of the mechanisms of Pb neurotoxicity in the developing human nervous system.

Keywords: Drosophila, toxicology, toxicogenomics, behavioral toxicology

PART 1: QTL MAPPING: BEHAVIORS AND TOXINS

Behavior as a QTL Endpoint for Toxicology Studies

There are two main reasons why behavioral assays are so useful in detecting effects of toxins. First is their richness. Behavior can be described as an ongoing, generally complex, spatio-temporal pattern; subtle changes in that pattern can signal the action of very low doses of a toxin well before there are dramatic changes in organ systems: in children overt clinical encephalopathy is associated with blood lead levels of 80–100 μg/dl, while changes in intelligence quotient (IQ) and learning occur at or below a tenth of that dose (UNEP, 2010). Second is that even subtle toxin-dependent changes in behavior can have consequences for well-being which makes them meaningful and relevant.

Since chronic developmental exposure to toxins affects assembly of the neuronal and hormonal systems mediating adult behavior, their effects are generally more pronounced than that following acute adult exposure. Developmental exposure to sub-lethal and sub-teratogenic levels of toxins may alter or degrade physiological and behavioral mechanisms in a quantitative manner. Examples include effects of low doses of lead on behavior in fruit flies (Hirsch et al., 2003), cognitive effects of polychlorobiphenyls in people (Faroon et al., 2000), lead effects on cognitive function in people (Counter et al., 1998), and the effects of aluminum on adult behavior and developmental rate of their offspring in mice (Abu-Taweel et al., 2012). Once toxin-dependent behavioral changes are observed, the next steps may include study of their underlying physiological effects, for example, effects of lead on synaptic function in fruit fly larvae (He et al., 2009).

The bulk of what is known about the effects of various toxins is based on studies focusing on a single toxin; combinations of toxins can have additive, protective, or synergistic effects (Rai et al., 2010; Singh et al., 2010).

DROSOPHILA BEHAVIORAL QTLs

Quantitative trait locus (QTL) mapping is used for unbiased genome-wide screens to identify genetic loci causing variation in a trait, but they are most efficient when large numbers of individual genomes can be assayed. The large sample sizes required for precise QTL analysis favors model organisms that can be bred in the lab easily and inexpensively, and traits that are simple to assay yet rich in their information content and thus provide sensitive measures of effects of toxins are especially desirable.

Drosophila is an ideal model organism for both efficiency and genetic analysis, since they are easy and relatively inexpensive to breed and maintain in large numbers, and have a surprising degree of genetic homology to mammals (Mackay and Anholt, 2006). Because of the small physical size of Drosophila many phenotypic traits can be difficult to assay rapidly or quantitatively. Exceptions include very simple traits such as bristle number, or...
those easy to quantify such as number of offspring, others may be easy to automate given an investment in the required technical support.

Automated phenotypic assays in *Drosophila* tend to favor either gene expression during embryogenesis, where molecular marker extraction and detection can be automated, or locomotor behavior which, by using *Drosophila* Activity Monitors (DAMs), can be recorded relatively easily for large numbers of flies over periods of many days (Figure 1A; TriKinetics Inc., Waltham, MA, USA; Rosato and Kyriacou, 2006; Rosato et al., 2008).

The richness of the data gathered using DAMs makes it an ideal automated assay for genetic analysis of toxins. DAM data always provides, at a minimum, an estimate of mean locomotor activity level for the duration of the experiment. However, by measuring locomotor activity under the influence of a controlled light–dark cycle the photosensitivity pattern of activity can be analyzed. Using regulated light–dark cycles allows for determining the distribution of activity between the two phases of the photoperiod; the effect of light–dark cycle length other than 24 h; the amplitude of “daily” activity rhythms; the percentage of flies that show significant daily rhythms; and the timing, length, and distribution of sleep periods (Shaw et al., 2000).

Finally, by replacing light–dark cycles with constant darkness the free-running circadian period can be estimated. This provides a direct measure of the output of the biological circadian pacemaker driving the activity rhythm. Toxins that alter the circadian periodicity pattern of locomotor activity are likely, therefore, to perturb the circadian synchronization of many different biological functions regulated by a common central circadian clock. Temporal analysis of behavioral effects of toxins could also suggest the presence of daily rhythms in sensitivity to specific toxins, or in the manifestation of toxic effects. In humans, certain forms of cancer chemotherapy are more effective when administered at specific times of day (Levi et al., 2010; Li et al., 2010); other toxins may also vary in their effects during the rhythm of daytime sunlight and nighttime darkness.

In short, DAMs allow for analysis of toxic effects on both the level of locomotor activity per se and on its temporal pattern of expression (von Mayersbach, 1975; Mayersbach, 1976).

**DROSOPHILA ACTIVITY MONITORS AND ACTIVITY ASSAYS**

Behavioral toxicogenetic analysis is a relatively new application of DAM technology. DAMs were originally devised for genetic analysis of the circadian oscillator controlling locomotor activity. Locomotor activity in fruit flies, similar to running wheel activity in rodents, is an excellent reporter phenotype for underlying circadian clock function (Takahashi et al., 2008). Since precise analysis of circadian rhythms require frequent, continuous long-term data sampling for time-series estimation of periodic oscillations in the 24-h range, DAMs are designed to produce a data-rich profile of the locomotor activity of individual fruit flies. Flies are housed, usually singly, in a 5 mm × 60 mm transparent tube with enough food to last for approximately 2 weeks. A single monitor holds 32 tubes, and a photosensor in the center of each tube tracks locomotor activity by numbers of beam-crossing per unit of time, usually 10-min intervals.

*Drosophila* Activity Monitors permit a variety of experimental designs. It is relatively easy to assay developmental effects of toxins by adding them to the fly culture food during egg, larval, or pupal development, and then monitoring adults. Effects of toxins can also be assayed acutely by adding them to the food in the activity monitor tubes. A single DAM, approximately 5” long, 4” high, and 3” deep holds 32 flies so one refrigerated–sized incubator can house thousands of flies per experiment. The efficiency of DAMs facilitates experimental analysis of sex differences, dose–response curves, sensitive periods during development, interaction among multiple toxins, trans–generational effects, and various types of genetic analysis including behavioral QTL. It is also just as easy to monitor survival time of individual flies to the minute in assays of lethal toxic effects and degrees of resistance to them.

**BEHAVIORAL QTL ANALYSIS OF DEVELOPMENTAL EXPOSURE TO TOXINS**

We have identified a toxin-induced behavioral QTL by assaying locomotor activity levels for a set of recombinant inbred (RI) fly lines raised on medium containing 250 μM lead acetate or control medium made with 250 μM sodium acetate. The variation among the RI fly lines in the difference in mean activity level per line between the two treatments indicated a behavioral QTL in the
3RAB region of the second Drosophila chromosome (Figure 1B; Hirsch et al., 2009). This QTL was independently identified as the site of an expression QTL (eQTL) in response to the same lead treatment in the same set of RI lines (RILs). Using the gene expression assay, we further showed the 30AR eQTL to function as a master modulatory regulator of approximately 70 additional eQTLs throughout the genome in response to developmental lead exposure (Raden et al., 2009). Current research underway in our labs will map these behavioral QTLs in response to developmental lead exposure more precisely.

**PART 2: SYNAPSES AND Ca2+ REGULATION IN DROSOPHILA**

Early findings suggested that chronic Pb2+ exposure could produce its behavioral effects by altering synaptic development. Mammalian studies reported a variety of alterations in synaptic morphology and physiology in the brains of animals exposed to Pb2+ (Hirsch et al., 1996). Today, the synapse remains a focus for studying the effects of toxins on brain development. Here the Drosophila larval neuromuscular junction (NMJ) offers a distinct advantage since one can compare the same, identified synapse in controls and treated animals (Figure 2).

Note that this neuromuscular system is unique amongst invertebrates because there is a stereotypic pattern of neuromuscular connections where both the presynaptic (motor neuron) and postsynaptic cells (muscle fiber) can be uniquely identified (Keshishian et al., 1996). These neuromuscular synapses have been used to demonstrate the effects of second messengers, cell-adhesion molecules and their modulators, and impulse activity on synaptic development (Budnik, 1996; Budnik et al., 1996; Davis et al., 1996).

We found that indeed one could detect synaptic abnormalities resulting from chronic Pb2+ exposure at the Drosophila larval NMJ. Initial studies found that the motor terminal formed by motoneuron RP3 on Muscle Fiber 6 (MF6) showed greater variability in size for animals raised in media containing Pb2+ compared to controls (Morley et al., 2003). One could imagine that increasing the variability of synaptic size could result in a loss of fine-tuning of synaptic function; this effect, if seen in the brain, would likely be detrimental to circuit dynamics and behavior. It is noteworthy that this synaptic change was subtle and it likely would have been difficult to detect when making comparisons among a large population of synapses in the mammalian brain. Given that the behavioral effects of toxins, such as Pb2+, can be subtle, it seems reasonable that the effects on synapses might also be small.

Pb2+ affects proteins that bind Ca2+ including those that regulate the intracellular Ca2+ concentration ([Ca2+]i), such as Ca2+ channels and Ca2+ pumps. Changes in Ca2+ regulation could be particularly important for the synapse since [Ca2+]i controls multiple steps in synaptic development; e.g., growth cone guidance (Jin et al., 2005), synapse formation (Xu et al., 2009), and synapse elimination and stabilization (Pratt et al., 2003; Lohmann and Bonhoeffer, 2008). In addition, altered Ca2+ regulation could influence transmitter release (Zucker, 1996) and both long-term and short-term forms of synaptic plasticity at the mature synapse (Zucker and Regehr, 2002; MacDonald et al., 2006).

Acute Pb2+ exposure blocks Ca2+ channels. This has been demonstrated for a variety of voltage-dependent Ca2+ channels in both invertebrates (Audesirk and Audesirk, 1989) and mammals (Evans et al., 1991). The acute application of micromolar concentrations of Pb2+ can reduce the activity of the plasma membrane Ca2+ ATPase (PMCA) in humans and rats (Mas-Oliva, 1989; Sandhir and Gill, 1994a,b); however the PMCA can be stimulated by low Pb2+ concentrations (Mas-Oliva, 1989; Campagna et al., 2000). In addition, chronic in vivo Pb2+ exposure has been found to produce a persistent inhibition of PMCA activity in human erythrocytes (Campagna et al., 2000) and rat synaptosomes (Sandhir and Gill, 1994a,b).

**FIGURE 2** (Ca2+ transients recorded from identified synaptic terminals in control and Pb2+ exposed animals. (Left) The RP3 motor terminals on MF6 were filled with OGB-1 in control larvae and in those exposed to Pb2+. The arrows point to two typical synaptic boutons where Ca2+ transients were measured. (Right) Typical Ca2+ transients produced by single action potentials (AP) and AP trains for control and Pb2+ exposed boutons. For single APs, the Ca2+ transients were similar for control and Pb2+ exposed synaptic boutons; however, the Ca2+ transients produced by AP trains were larger and decayed more slowly in Pb2+ exposed boutons compared to controls. Calibration: single AP – 20% ΔF/ΔF, 0.4 s; AP train – 20% ΔF/ΔF, 2 s. Adapted from Ye et al. (2009).
We studied the effect of chronic Pb\textsuperscript{2+} exposure on presynaptic Ca\textsuperscript{2+} regulation at the RP3-to-MF6 synapse in *Drosophila* larvae (Hö et al., 2009). Ca\textsuperscript{2+} indicators were loaded in these motor terminals and we measured the changes in [Ca\textsuperscript{2+}]\textsubscript{i} produced by single action potentials and action potential trains (Figure 2). We found that chronic exposure to Pb\textsuperscript{2+} resulted in a greater increase in [Ca\textsuperscript{2+}]\textsubscript{i}, during trains of action potentials and a slower decay of post synaptic [Ca\textsuperscript{2+}]\textsubscript{i} at the end of the train. This is likely due to a decrease in the activity of the PMCA and it provides an interesting parallel to the effect of Pb\textsuperscript{2+} exposure on the PMCA in mammals. The direct effect of this increase in [Ca\textsuperscript{2+}]\textsubscript{i} was that the Pb\textsuperscript{2+}-exposed animals showed greater synaptic facilitation. This was consistent with the residual Ca\textsuperscript{2+} model for synaptic facilitation (Zucker and Regehr, 2002) and with mammalian studies showing that knocking down expression of the PMCA resulted in enhanced synaptic facilitation (Emerson et al., 2007). In that study, reduced PMCA expression also produced changes in neuronal structure (Emerson et al., 2007) and it may be that the Pb\textsuperscript{2+}-induced changes in the structure of the larval NMJ resulted from reduced PMCA activity and altered Ca\textsuperscript{2+} regulation.

In summary, identified NMJs in *Drosophila* larvae allow for the detection of subtle changes in synaptic structure and function resulting from developmental exposure to toxins. Our evidence suggests that chronic Pb\textsuperscript{2+} exposure produces parallel synaptic changes in *Drosophila* and mammals; this would not be surprising given that chemical synapses found in vertebrates and invertebrates are very similar.

### PART 3: GENETICAL GENOMICS STUDIES IN *DROSOPHILA*

Developmental neurotoxicology research requires an approach that reduces the candidate toxin-regulated genes to a manageable number. Fortunately, a new multi-dimensional strategy has been developed, called genetical genomics, which identifies master modulatory loci that regulate the expression of hundreds of other genes (Jansen and Nap, 2001; Broman, 2005; de Koning and Haley, 2005; Li and Burmeister, 2005; Rockman and Kruglyak, 2006).

Genetical genomics combines two methodologies, microarray-based whole transcriptome analyses and extracting QTLs by using RILs. Global gene expression levels are determined for each RIL, and then QTL mapping software (e.g., R/QTL; Broman et al., 2003) is used to find above-threshold statistical associations between specific chromosomal loci and transcript levels for all the genes measured on the microarrays. With QTL analysis, we can identify the specific chromosomal loci that regulate genes, which are referred to as eQTLs (Muller et al., 2006; West et al., 2007; Majewski and Pastinen, 2011; Zhang et al., 2012).

Our research is the first that combines genetical genomics with toxicogenomics, an experimental approach we call "genetical toxicogenomics" (Ruden et al., 2009). Li et al. (2008) showed that adding environmental perturbations in genetical genomics studies, as we did in our study (Ruden et al., 2009), allows toxicity responses to be identified. We believe that this genetical toxicogenomics approach will drive the field of toxicology and aid in understanding the effects of toxins such as lead.

Expression QTL analyses were performed for all ∼18,000 genes and other microarray features by treating the expression level of each gene as a quantitative trait. The genetics of gene expression in RILs can be mapped as eQTLs. Flies from each of 75 RILs were fed, from egg to adult, either control food or lead-treated food (made with 250 μM lead acetate). RNA expression analyses of whole adult male flies (5–10 days old) were performed with Affymetrix *Drosophila* whole genome arrays (18,952 probe sets). Loci that are linked to a gene are called locally acting or cis-eQTL, whereas loci that are distant acting are called trans-eQTLs (Broman, 2005).

The first genetical genomics study to identify genes with significant GxE interactions was done in *Caenorhabditis elegans*; the authors identified a group of genes with trans-eQTL that are induced by heat shock, which they called plastic QTL (Li et al., 2006). Smith and Kruglyak (2008) recently performed a detailed analysis of GxE-eQTL in yeast (which they call "geoQTL") grown in either glucose or ethanol as the sole carbon source. Others have identified GxE interactions in which the environment is a different tissue (e.g., brain vs. liver; Hovatta et al., 2007). In our studies, among the 1,389 genes with cis-eQTL, there were 405 genes unique to control flies and 544 genes unique to lead-treated ones (440 genes had the same cis-eQTLs in both samples).

There were 2,396 genes with trans-eQTL that mapped to 12 major hotspots that met statistical significance (p < 0.05, chi-squared test, based on permutation analyses, on 5-cM windows). Unexpectedly, we identified two hotspots, one located on the second chromosome (polyteny region 30AB) and one on the third chromosome (polyteny region 73D), which co-regulate 33 genes, all of which are induced by lead (Figure 3). We have shown by QTL analysis that marker loci 30AB contributes to lead-dependent changes in locomotion, which suggests that the genes in the 30AB hotspot can be used as a functional test to identify both the lead-dependent trans-regulatory factor and the common cis-regulatory motifs (Hirsch et al., 2009). We propose that a trans-regulator located at 73D increases expression of its target genes when it binds lead, regardless of the genotype of the 73D hotspot. In contrast, a second trans-regulator located at 30AB increases expression of the co-regulated target genes only when it has the ORE genotype (Figure 3).

This model can explain the data, but other explanations are possible. For example, the putative trans-regulator could increase the stability of the mRNA of the target genes, or there could be an indirect effect on steady-state mRNA levels. MEME (Multiple Em for Motif Elicitation) analyses of the genes regulated by the 30AB and 73D trans-regulators identified both conserved proximal promoter and 5′-untranslated region (UTR) sequences (data not shown). Fine-mapping of the genes that underlie the QTLs and molecular and biochemical analyses should enable us to determine the mechanism involved.

### FUTURE STUDIES IN *DROSOPHILA* NEUROTOXICOLOGY

In this review, we describe recent results from several laboratories that are collaborating on studying the effects of lead on the developing *Drosophila* nervous system. The Possidente and Hirsch
laboratories study the effects of lead on behavioral quantitative traits. The Lnenicka laboratory studies the effects of lead on the NMJ in the larvae. The Ruden, Lu, and Garfinkel laboratories study the effects of lead on gene expression using quantitative genetic techniques.

Further innovations to the field of genetical genomics make use of next-generation RNA sequencing (RNA-seq) rather than gene expression microarrays. RNA-seq will allow us not only to accurately determine gene expression levels, but also to analyze alternative splicing products of genes, thereby adding a new dimension to these studies. Since over 90% of human genes and a similar percentage of *Drosophila* genes are alternatively spliced, RNA-seq could provide some exciting and novel findings about environmental regulation of alternative mRNA splicing. The potential for RNA-seq to supplant microarrays requires, however, the development of robust statistical tools for analyzing the resulting huge datasets.

Adding metabolomics profiling to these studies will allow us to identify metabolite QTL (mQTL) that are specific for lead. To our knowledge mQTL analyses have never been conducted in any organism, although genome-wide association studies combined with metabolic profiling were conducted in a recent human study (Prakash, 2011; Suhre et al., 2011). The human study identified 37 genetic loci associated with blood metabolite concentrations, of which 25 showed effect sizes of 10–60% which is extraordinarily high for genome-wide association studies (Prakash, 2011; Suhre et al., 2011).

We are now entering the stage of understanding the complex genetic pathways that are affected by developmental lead exposure. The sophisticated genetic analyses that are possible in *Drosophila* will soon allow us to manipulate these pathways to better understand how they are affected by lead. The sequencing of several *Drosophila* strains and studying the effects of lead on neurodevelopment in these strains will allow us to better understand the evolution of lead-sensitive pathways. Together, these studies will provide a better understanding of the health effects of lead in humans.

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