**PARK2 patient neuroprogenitors show increased mitochondrial sensitivity to copper**

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Abstract:

Poorly-defined interactions between environmental and genetic risk factors underlie Parkinson's disease (PD) etiology. Here we tested the hypothesis that human stem cell derived forebrain neuroprogenitors from patients with known familial risk for early onset PD will exhibit enhanced sensitivity to PD environmental risk factors compared to healthy control subjects without a family history of PD. Two male siblings (SM and PM) with biallelic loss-of-function mutations in *PARK2* were identified. Human induced pluripotent stem cells (hiPSCs) from SM, PM, and four control subjects with no known family histories of PD or related neurodegenerative diseases were utilized. We tested the hypothesis that hiPSC-derived neuroprogenitors from patients with *PARK2* mutations would show heightened cell death, mitochondrial dysfunction, and reactive oxygen species generation compared to control cells as a result of exposure to heavy metals (PD environmental risk factors). We report that *PARK2* mutant neuroprogenitors showed increased cytotoxicity with copper (Cu) and cadmium (Cd) exposure but not manganese (Mn) or methyl mercury (MeHg) relative to control neuroprogenitors. *PARK2* mutant neuroprogenitors also showed a substantial increase in mitochondrial fragmentation, initial ROS generation, and loss of mitochondrial membrane potential following Cu exposure. Our data substantiate Cu exposure as an environmental risk factor for PD. Furthermore, we report a shift in the lowest observable effect level (LOEL) for greater sensitivity to Cu-dependent mitochondrial dysfunction in patients SM and PM relative to controls, correlating with their increased genetic risk for PD.
Introduction

Parkinson's disease (PD) is characterized by progressive motor function decline due to loss of dopaminergic neurons in the substantia nigra (SN). A minority of PD patients (familial PD) have single gene mutations associated with disease etiology (~ 10% of cases). The remaining idiopathic PD cases are influenced by interactions of genetic and environmental risk factors, including exposure to heavy metals and pesticides (Betarbet et al., 2000, Gorell et al., 1999, Gorell et al., 1998 and Jomova et al., 2010). The role of environmental factors is underscored by a high discordance between monozygotic twins (Tanner et al., 1999).

Mutations of PARK2 are one of the most common causes of early onset PD (EOPD) (Kitada et al., 1998). Despite this association, patients with biallelic loss-of-function PARK2 mutations can have incomplete penetrance with high intra-familial and inter-familial variability in ages of onset (Abbas et al., 1999, Deng et al., 2006 and Khan et al., 2003). For example, one study found a 56-year old subject with compound heterozygous mutations in PARK2 exhibiting no evidence of PD despite having four siblings with the same PARK2 mutation that were diagnosed with EOPD at ages from 30 to 38 (Deng et al., 2006). Therefore, we hypothesized that biallelic loss-of-function PARK2 mutations would increase sensitivity to PD-associated environmental risk factors.

Recently, differentiated neural lineages from hiPSCs of patients with PARK2 mutations have been reported to exhibit mitochondrial dysfunction and increased oxidative stress (Imaizumi et al., 2012). Both these phenotypes have separately been associated with exposures to heavy metals (Jomova et al., 2010). Here we performed a proof-of-principle assessment of PD-relevant toxicant vulnerability in two subjects (SM and PM) with compound heterozygous loss-of-function mutations in PARK2 versus control subjects. We utilized hiPSCs that are differentiated into neuroprogenitors that retain the unique genetic information of each human subject. Toxicant sensitivities at this early neurodevelopmental time point are pertinent given the strong evidence for in utero and early life environmental insults contributing to subsequent risk for PD (de la Fuente-Fernández and Calne, 2002 and Landrigan et al., 2005).

Methods

Human subjects, clinical findings, generation of hiPSC lines

Primary dermal fibroblasts were obtained by skin biopsy from healthy adult subjects (CA, CE, CF) with no known family history or genetic risk factors for PD and two PD patients (PM and SM) after appropriate patient consent/assent under the guidelines of an approved IRB protocol (Vanderbilt #080369). The MRC-5 fibroblasts (obtained from Coriell Institute for Medical
Research and designated CB) were originally derived from a 14-week fetus aborted for maternal psychiatric reasons from a 27 year-old physically healthy woman with no known family history or genetic risk factors for PD. The studied patient PM was diagnosed with EOPD in his 30s and had dystonia from age 12 and resting tremor at age 17. He began having gait difficulties at age 23. Medical genetic analysis was performed, and biallelic mutations in the \textit{PARK2} locus were identified. The studied patient SM, brother of PM, was diagnosed at the Vanderbilt University Medical Center Movement Disorders Clinic with exercise-induced dystonia. Given his family history, SM also underwent genetic testing revealing the same compound heterozygous mutations in \textit{PARK2} as his brother PM. SM had a normal neurological exam by a movement disorders specialist at age 40 – the time of the skin biopsy – with no evidence of baseline dystonia or parkinsonism at that time. However, a clinical DaTScan via SPECT imaging later revealed bilateral dopaminergic denervation in the putamen with relative preservation of caudate and led to his current diagnosis of preclinical PD. Patients SM and PM reported no known exposures to heavy metals or other PD-relevant environmental risk factors such as pesticides. Furthermore, there was no evidence of any additional occupational-related exposures. Fibroblasts and hiPSCs were cultured as described previously (Neely et al., 2012). hiPSC lines CA4, CA6, CB5, SM3, SM4, and SM5 were reprogrammed using viral vectors whereas hiPSC lines CA11, CE6, CF1, SM14, PM12 and PM17 were reprogrammed using an episomal-based method (Okiti et al., 2011 and Takahashi et al., 2007). Line nomenclature follows an alphanumeric sequence, wherein the first two letters designate the human subject they are derived from. The sequential numbers identify individual clones picked from the original reprogramming plate (e.g. SM14 comes from the 14th iPSC colony isolated from this subject). The two-letter subject code does not contain patient identifying characteristics. Validation of most hiPSC lines used in this study has been previously published (Neely et al., 2012 and Srinivasakumar et al., 2013). Additional lines underwent and passed the same validation. For example, see the \textit{Supplemental materials} (Figure S1). Like our previously published lines, new hiPSC lines used in this study had normal euploid karyotypes and validated expression levels of pluripotency markers as assessed by immunocytochemistry, qPCR, and/or by the PluriTest, a bioinformatics assay developed by Muller and colleagues to quantify pluripotency (Muller et al., 2008). Multiple clonal lines for each subject were used in our experiments to decrease the influence of line-specific variability. See \textit{Supplemental materials} for more details.

Differentiation of early forebrain neuroprogenitors

Dual-SMAD inhibition-based forebrain neuroprogenitor differentiation of hiPSC lines was performed using our previously published protocols and based on the work of Studer and colleagues (Chambers et al., 2009 and Neely et al., 2012). These forebrain neuroprogenitors are positive for expression of \textit{PAX6}, \textit{FOXG1}, \textit{OTX2}, and \textit{SOX1} (Neely et al., 2012). See \textit{Supplemental materials} for more details.

Cell viability/cytotoxicity assay
Neuroprogenitors were exposed to the metal toxicants on day 6 of neural induction for a period of 24 or 48 h as indicated. For fibroblast experiments, cells were plated at a density of $5 \times 10^4$/mL and exposed to toxicants the day after plating. Toxicants included heavy metal cations of Cd (Cd$^{2+}$ as CdCl$_2$), Cu (Cu$^{2+}$ as CuSO$_4$), Mn (Mn$^{2+}$ as MnCl$_2$), and MeHg (CH$_3$Hg$^+$ as CH$_3$HgCl$_2$). Cell viability was assessed by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay as previously described with 2 h incubation in neuralization medium containing MTT (Williams et al., 2010).

Alternatively, CellTiter-Blue assay (Promega) was used to measure cell viability. At 2 h prior to the completion of the 24-hour exposure period, 20 μL of CellTiter-Blue reagent (Promega) was added to each well. Prior to this addition, cell lysis buffer was added to several wells to provide an accurate fluorescence background for 0% viable cells. The plates were then incubated for 2 h at 37 °C. Fluorescence was measured using excitation of 570 nm and emission of 600 nm on a POLARstar Omega microplate reader (BMG Labtech). Data were fitted to a normalized inhibitor curve with variable slope using Prism5 (GraphPad).

We also measured the percentage of non-apoptotic cells via forward and side scatter measurements by flow cytometry. Single cell events were also identified and gated by forward and side scatter. For each experiment, the percentage of non-apoptotic cells was calculated by excluding apoptotic cells based on their reduced forward scatter due to cell shrinkage (Healy et al., 1998). Gating was set based on the non-apoptotic cells of the vehicle-treated sample for each genotype and applied to the Cu-exposed cells of that genotype for each experiment (Figure S4). The percentage of non-apoptotic cells is expressed as the percentage of total gated events (Figure S4) in each sample that satisfy the single cell gating parameters.

Semi-quantitative mitochondrial fragmentation analysis

Semi-quantitative mitochondrial fragmentation analysis was done similar to our previously published method, based on previously published methods (Lutz et al., 2009). In brief, on day 6 of neuralization, neuroprogenitors were passaged as described in the Supplemental materials and plated onto 8-well glass slides (Lab-Tek) at a cell density of $1 \times 10^5$ cells/mL. Two mitochondrial staining methods were used in this work: Mitotracker red dye and CellLight Mitochondria-GFP. In the former method, neuroprogenitors were exposed on day 7 of neuralization to either vehicle or Cu for 24 h. Following metal exposure, neuroprogenitors were incubated with 100 nM Mitotracker red dye (Invitrogen) in neuralization medium for 45 min at 37 °C. The dye-containing medium was subsequently taken off and cells were incubated in dye-free neuralization medium for 10 min, washed once with PBS, and fixed with 4% paraformaldehyde for 10–15 min at room temperature. Alternatively, the mitochondrial-specific viral transduction system (CellLight Mitochondria-GFP, BacMam 2.0; Invitrogen) was used to label mitochondria following the manufacturer's recommended protocol. Briefly, the reagent was added to neuroprogenitors on day 6 of neuralization at time of replating at a multiplicity of infection (MOI) of 15. Neuroprogenitors were exposed 24 h later to either vehicle
or Cu for another 24 h then fixed as specified above. Following fixation, immunofluorescence staining was performed with PAX6 antibody (Covance, 1:200 dilution). Then cells were incubated for 10 min with 2 μM TO-PRO-3 for nuclear counterstaining. Mitochondrial fragmentation was analyzed with a confocal microscope (LSM 151 Meta confocal laser scanning microscopy system; Carl Zeiss) using 63 × 1.4 NA objective. Mitochondrial fragmentation in each of forty randomly chosen PAX6-expressing cells (PAX6 +) for each exposure paradigm was assessed at the confocal microscope by a trained observer blinded to cell lineage and exposure based on published methods (Lutz et al., 2009). The number and size of mitochondrial fragments, as well as the interconnectivity of the mitochondrial network, were considered in sum by the human observer to classify mitochondria of each neuroprogenitor into the semi-quantitative scale. Using this method, cells were assigned to one of three categories of mitochondrial fragmentation: mild, moderate, or severe (an example of each category is shown, Fig. 4). Apoptotic and mitotic cells were identified by morphology (shrunken cells with condensed nuclei and cells with nuclear fission, respectively) and excluded from analysis.

Repeated measures general linear model two-way ANOVA (fixed factors of Cu exposure and disease group) was performed using SPSS (IBM), with the fragmentation severity categories as the repeated measure for each independent sample. Post-hoc analysis was performed by two-tail t-test or Bonferroni binary comparison (when the number of comparisons exceeded two) to identify significant differences between groups in each fragmentation severity class.

Quantitative mitochondrial morphology analysis

Mitochondrial morphology was also assessed in a quantitative manner using Mito-Morphology macro as previously described (Dagda et al., 2009). The GFP channel of PAX6 + cells of 10 confocal images from two independent experiments (acquired as above) were converted to RGB images in ImageJ (National Institutes of Mental Health). Images were then thresholded to a binary image mask using the “Process RGB Image” function of the macro then analyzed using the “Measure” function of the macro. Circularity, particle count, and total area of the mitochondria were quantified to assess mitochondrial morphological changes as previously described (Dagda et al., 2009 and Xie and Chung, 2012).

Mitochondrial membrane potential analysis

Neuroprogenitors were exposed on day 7 of neuralization for 24 h and then dissociated using Accutase, centrifuged, and resuspended in neuralization medium containing 5 nM of DiIC1(5) dye for 15 min. Cells were washed once in PBS and then analyzed using flow cytometry (638/658 nm excitation/emission). The samples were analyzed on a Custom Becton Dickson five-laser Fortessa analytical cytometer using BD FACSDiva acquisition (BD Biosciences) and FlowJo analysis software (Tree Star, Inc.). A total of 10,000 events were acquired, and analysis was restricted to non-apoptotic single cell events as determined by light scatter properties (side and forward scatters). Apoptotic cells were identified by their relative higher side scatter and lower forward scatter due to cell shrinkage and were excluded from the analysis. Experiments
were done in pairs (CA line versus an SM line, as indicated), and the average fluorescence intensity of all viable cells in the Cu-treated cells was normalized to the average intensity for the viable vehicle-treated cells for each line.

Reactive oxygen species (ROS) analysis

ROS analysis was done as previously described (Aboud et al., 2012). In brief, neuroprogenitors were passaged on day 6 of neuralization as described in the Supplemental methods. On day 7, the cells were loaded with 2 μM CM-2′,7′-dichlorodihydrofluorescein diacetate (CM-H2DCFDA) dye for 30 min in exposure buffer [25 mM HEPES buffer (pH 7.2), 140 mM NaCl, 5.4 mM KCl, and 5 mM d-glucose] in the dark at room temperature. The cells were washed with exposure buffer to remove excess dye and then exposed to either vehicle (exposure buffer), Cu, or H2O2 in exposure buffer. Fluorescence intensity was measured at 37 °C every 5 min using a Beckman Coulter DTX 880 microplate reader with excitation at 485 nm (filter bandwidth ± 20 nm) and emission at 535 nm (filter bandwidth ± 25 nm). Hydrogen peroxide exposure was used as a positive control in each experiment, and a significant concentration-dependent increase by hydrogen peroxide in fluorescence was seen across the experimental set (repeated measures ANOVA, p < 0.0001, data not shown).

Additional reagent and method information is available in the Supplemental materials.

Results

Sequencing of PARK2 cDNA in the patient derived cells

Prior clinical genetic testing of patients SM and PM had revealed the inheritance of maternal and paternal PARK2 deletion alleles with one allele missing a segment of exon 3 and the other allele missing exons 5 and 6. Amplification of cDNA made from fibroblast mRNA with primers flanking the deleted exons from both alleles revealed mutant transcripts consistent with the genomic deletions identified by clinical genetic testing (Figure S2). Gene sequencing of the mutant alleles revealed the presence of both PARK2 deletion mutations in cDNA from the patient's (SM) primary dermal fibroblasts (40 bp deletion in exon 3; and 200 bp deletion of exons 5 and 6; Figure S2). This sequence analysis confirmed the loss of the exons in the genomic DNA of SM, originally identified by clinical genetic testing. Additionally, the analysis showed that mRNA splicing occurred between the exons adjacent to the deletions in each allele. Both allelic deletions result in a frameshift predicted to cause premature termination of translation. These frameshift mutations are N-terminal to the conserved RING domain, and, thus, would be anticipated to completely abolish ubiquitination function of any potentially translated protein (Deshaies and Joazeiro, 2009). Therefore, SM and PM cells are predicted to be functional null alleles for PARK2.

Differential heavy metal sensitivity of SM and CA neuroprogenitors
We sought to test the hypothesis that PARK2 mutant neuroprogenitors (e.g. patient SM) would show elevated sensitivity to PD-relevant environmental risk factors. We tested the viability of control (e.g. subject CA) versus patient SM derived neuroprogenitors by MTT assay after 48 hour exposure to Mn, Cu, Cd, and MeHg, all of which have been implicated in PD-relevant neurodegenerative processes and/or as PD environmental risk factors (Buzanska et al., 2009, Gorell et al., 1999, Götz et al., 2002, Jomova et al., 2010, Landrigan et al., 2005, Rivera-Mancía et al., 2010, Squitti et al., 2009, Weiss et al., 2002, Willis et al., 2010 and Xu et al., 2011). To control for cell line dependent variability, we used multiple clonal hiPSC lines (lines CA4 and CA6 for subject CA; lines SM3, SM4, and SM5 for subject SM). Repeated measures ANOVA across the concentration response curves demonstrated a heightened sensitivity to Cu and Cd but not to Mn or MeHg in SM neuroprogenitors (lines SM3, SM4 and SM5) when compared to CA neuroprogenitors (lines CA4 and CA6) (Fig. 1A). To determine whether the differential Cu and Cd sensitivity was also observed in non-neural cell types between the same human subjects we generated 48 hour Cu and Cd cytotoxicity concentration–response curves by MTT assay in the primary epidermal fibroblasts used to generate the hiPSC lines. No significant sensitivity differences to Cu or Cd (Fig. 1B) viability by MTT assay were observed, suggesting at least some cell type dependence of the differential metal sensitivity between these subjects.
Fig. 1. SM neuroprogenitors but not fibroblasts show increased sensitivity to Cd and Cu toxicity. (A) Neuroprogenitor survival curves measured by MTT assay after 48 hr exposure to Mn, MeHg, Cu, and Cd (CA: CA4, CA6 at N = 6 or 8; SM: SM3, SM4, SM5 at N = 7 or 8). Error bars represent ± SEM. (B) The primary dermal fibroblast lines were exposed to Cd or Cu, and cell viability was measured by MTT assay (Cu: CA, n = 7; SM, n = 7; Cd: CA, n = 3; SM, n = 3). Statistical analysis for genotype effects was performed by two-way repeated measures ANOVA. Pairwise post-hoc analysis with Bonferroni correction is indicated between CA and SM hiPSC lines as *p < 0.05, ***p < 0.001. Error bars represent ± SEM.
We decided to further examine the genotype-dependent/subject-dependent differences in Cu cytotoxicity, rather than Cd, given the broader concentration response curve and the redox potential of Cu that allows it to participate in Fenton chemistry. We saw similar genotype-dependent differences in viability following 24-hour Cu exposure between CA control cells and the PARK2 mutant neuroprogenitors from both SM and PM (Figure S3) by the CellTiter-Blue assay, which quantifies viable cells by measuring total cytoplasmic, microsomal, and mitochondrial reductase activity (Gonzalez and Tarloff, 2001). Since both MTT and CellTiter-Blue viability assays depend, at least in part, on mitochondrial reductase activity, we sought to confirm if the effects observed by these assays included decreased cell viability, rather than just changes in total mitochondrial reductase activity. Preliminary studies ruled out the use of the lactate dehydrogenase (LDH) assay for this purpose because the presence of Cu in the media (as low as 25 μM) significantly impaired detection of LDH as previously shown in the literature (Pamp et al., 2005). Therefore, we utilized measurement of non-apoptotic cells by flow cytometry as a non-mitochondrial dependent assessment of cytotoxicity following 24-hour vehicle or 50 μM Cu exposure in control CA and PARK2 mutant SM neuroprogenitors (Fig. 2 and Figure S4). This was performed by measuring the percentage of non-apoptotic cells (apoptotic cells have reduced forward scatter) from all single cell events. Two-way ANOVA revealed significant differences in viability due to Cu exposure ($F_{1,16} = 41.76$, $p < 0.001$), genotype ($F_{1,16} = 14.33$, $p = 0.002$), and a Cu × genotype interaction effect ($F_{1,16} = 15.39$, $p = 0.001$). Post-hoc analysis demonstrated a significant decrease in percentage of non-apoptotic Cu-exposed SM neuroprogenitors versus both vehicle ($p < 0.001$) and Cu-exposed CA neuroprogenitors ($p < 0.001$); as well as a significant decrease in viability of Cu-exposed CA neuroprogenitors compared to the vehicle-exposed CA group (Fig. 2). No difference in baseline percentage of non-apoptotic cells was observed between CA and SM. Overall we found a consistent increase in sensitivity of SM neuroprogenitors to Cu cytotoxicity that includes increased apoptosis and loss of cellular reductase activity.

**Fig. 2.** SM neuroprogenitors show a stronger reduction in the non-apoptotic cell population following Cu exposure. Gating for non-apoptotic cells was performed based on the forward and side scatter properties of the vehicle-exposed control group in each independent experiment as shown in Figure S4. Percentage of non-apoptotic cells is plotted. Apoptotic cells have decreased forward scatter properties due to cell shrinkage. Data collected across 5 independent paired
experiments (CA6:SM4 n = 2, CA6:SM5 n = 2, CA11:SM14 n = 1, total n = 5). Statistical analysis was performed by two-way ANOVA and paired two-tailed *-test for post-hoc analysis (** or ^ indicates p < 0.001, absence of a symbol indicates no significant difference p > 0.05). * indicates a significant difference within genotype between exposures, and ^ indicates a significant difference between genotypes with the same exposure. Error bars represent + SEM.

One possible hypothesis to explain the increased sensitivity to Cu cytotoxicity is elevated cellular uptake of Cu in the SM neuroprogenitors. Therefore, we measured the levels of intracellular Cu following a 24-hour exposure to 50 μM Cu by graphite furnace atomic absorption spectroscopy (GFAAS). No differences in intracellular Cu levels in vehicle or Cu treated SM (SM5) and CA (CA6) neuroprogenitors were observed (Fig. 3).

Fig. 3. Total Cu accumulation is not different between SM and CA neuroprogenitors. CA (CA6) and SM (SM5) neuroprogenitors were treated with either vehicle (Veh) or 50 μM Cu (Cu) for 24 h, and total cellular Cu levels were assessed by GFAAS (Vehicle n = 3; Cu n = 5). Two-way ANOVA was used for statistical analysis. Error bars represent + SEM.

Elevated mitochondrial fragmentation due to Cu exposure in SM and PM neuroprogenitors

We hypothesized that the increased sensitivity of SM and PM neuroprogenitors to Cu was due to an enhanced susceptibility of the mitochondria to oxidative damage. To evaluate this, we performed a large study with hiPSC lines from each of three control subjects (CA11, CE6 and CF1) and two lines each from SM and PM (SM3, SM14, PM12, and PM17). We assessed mitochondrial fragmentation by semi-quantitative assessment of severity using a blinded observer in two independent experiments each with 3 control lines (total n = 6 across 2 experimental days) and 4 PARK2 mutant lines (total n = 8 across 2 experimental days) in 24-hour vehicle or 100 μM Cu-exposed PAX6 + neuroprogenitors ( Fig. 4). Cellular mitochondrial fragmentation of individual PAX6 + neuroprogenitors was classified into a 3-level semi-quantitative scale including minimal, moderate, and severe by a trained observer (see examples, Fig. 4A). PAX6 + cells were selected prior to visualizing the mitochondria, and then, stained mitochondria of the selected neuroprogenitor were visualized by scanning through the z-axis to inform the scoring decision. Two-way repeated measures (RM)-ANOVA revealed significant differences in severity of fragmentation due to Cu exposure (F_{2,23} = 86.753,
p < 0.001), genotype (F2,23 = 7.604, p = 0.003), and a Cu × genotype interaction effect (F2,23 = 6.448, p = 0.006) (Fig. 4B). Repeated measures ANOVA of vehicle treated neuroprogenitors found no differences between SM/PM versus control (CA11, CE6, CF1) (F2,11 = 0.314, p = 0.737), while neuroprogenitors exposed to Cu had a significant difference in the severity of fragmentation (F2,11 = 9.630, p = 0.004) between SM/PM versus control. Post-hoc examination of the data demonstrated that both SM (p < 0.05) and PM (p < 0.05) neuroprogenitors showed significantly more mitochondrial fragmentation than control neuroprogenitors following Cu exposure (fewer cells with minimal fragmentation, and more cells with severe fragmentation). No significant differences in fragmentation were observed under vehicle exposure conditions between control and PD lines.
Fig. 4. Enhanced Cu-dependent mitochondrial fragmentation in SM/PM neuroprogenitors. Forty individual PAX6 + neuroprogenitors were scored into three severity classes of mitochondrial fragmentation (minimal, moderate and severe) for each independent sample, and the percentage of cells in each severity category was determined. (A) Representative images of mild, moderate, and severe categories of mitochondrial fragmentation for the semi-quantitative analysis. (B) Quantification of mitochondrial fragmentation in control (CA11, CE6, CF1), SM (SM3, SM14) and PM (PM12 and PM17) neuroprogenitors after a 24 hr exposure to vehicle or 100 μM Cu. Two independent experiments were performed with each line for a total of n = 6 control experiments; n = 8 SM/PM experiments (n = 4 SM; n = 4 PM). Analysis was performed using repeated measures ANOVA (across the three severity categories) and post-hoc t-test (columns are labeled with ‘a’ or ‘b’ to designate significant differences, p < 0.05). Error bars represent ± SEM.

We then applied a quantitative analysis of mitochondrial morphology on a subset of mitochondrial images using Mito-Morphology macro, an ImageJ based program, as previously described (Dagda et al., 2009). 10 images were randomly selected from two independent experiments (5 images from each) for each exposure condition and processed into a binary image mask for analysis by the macro (four representative processed images for each experimental group are provided in Fig. 5A). Automated analysis of the mitochondrial morphology was performed for circularity, segment/particle number, and total area. Two-way ANOVA revealed a significant difference in circularity due to Cu exposure (F_{1,36} = 14.46, p < 0.001), genotype (F_{1,36} = 13.35, p < 0.001), and a Cu × genotype interaction effect (F_{1,36} = 5.946, p = 0.02). Post-hoc analysis demonstrated a significant increase in circularity in Cu-exposed SM neuroprogenitors compared to both vehicle-exposed SM neuroprogenitors (p < 0.001) and Cu-exposed CA neuroprogenitors (p = 0.02) (Fig. 5B). Two-way ANOVA also revealed a significant difference in the number of mitochondrial segments due to Cu exposure (F_{1,36} = 17.21, p < 0.001) and a Cu × genotype interaction effect (F_{1,36} = 6.31, p = 0.02). However, genotype did not have a significant effect. Post-hoc analysis demonstrated a significant increase in the number of mitochondrial segments in Cu-exposed SM neuroprogenitors compared to both vehicle-exposed SM neuroprogenitors (p < 0.001) and Cu-exposed CA neuroprogenitors (p = 0.006) (Fig. 5C). No significant differences were found in circularity or mitochondrial count between vehicle-exposed and Cu-exposed CA neuroprogenitors. No significant change in total mitochondrial area was observed between these groups either (Fig. 5D). Taken together, the automated Mito-Morphology analysis corroborated our findings by semi-quantitative analysis of increased severity of mitochondrial fragmentation by Cu exposure in SM neuroprogenitors relative to control neuroprogenitors.
Fig. 5. Quantitative mitochondrial analysis of Cu-exposed neuroprogenitors. Quantitative analysis of mitochondrial morphology was performed using the ImageJ Mito-Morphology macro. Randomly selected PAX6 + neuroprogenitors were imaged by an observer blinded to genotype or exposure condition from two independent experiments (5 images from each, total n = 10) for both vehicle and Cu exposed CA6 and SM3 neuroprogenitors. (A) Four representative, binary image masks from each treatment category are shown for visual comparison. Quantitative measures were assessed by the macro including (B) mitochondrial
circularity, (C) total number of mitochondrial segments/particles, and (D) total mitochondrial surface area of the binary image masks. Statistical analysis was performed by two-way ANOVA and two-tailed *t*-test for post-hoc analysis (** or ^^^ indicates \( p < 0.001 \); ^^ indicates \( p < 0.01 \)). * symbol indicates significant difference as compared to the vehicle-treated group of the same genotype, and ^ symbol represents significant difference compared to Cu-exposed group of the other genotype. Error bars represent \( \pm \) SEM.

Lower threshold for Cu-dependent mitochondrial fragmentation in SM neuroprogenitors

We hypothesized that the mitochondria of the PD risk group neuroprogenitors are sensitive to lower concentrations of Cu than the control group neuroprogenitors. To evaluate this hypothesis, we assessed mitochondrial fragmentation in PAX6 + neuroprogenitors derived from 3 SM lines (SM3, SM4 and SM5) and 2 control lines (CA6 and CB5) with concentrations of Cu at 10 μM, 25 μM, and 50 μM (Fig. 6 and S5). We performed 8 independent paired experiments, each one with a single SM line and control line. Two-way repeated measures ANOVA of the complete data set revealed significant differences in severity of fragmentation due to Cu-exposure \((F_{6,112} = 17.912, \ p < 0.001)\), genotype \((F_{2,55} = 110.259, \ p < 0.001)\) and a Cu × genotype interaction effect \((F_{6,112} = 6.973, \ p < 0.001)\) similar to what was seen at 100 μM Cu in the experiment above (Fig. 4). Analysis of the 10 μM Cu exposure data revealed significantly increased fragmentation severity in the SM neuroprogenitors but not the controls (Fig. 6). Both 25 and 50 μM Cu caused increased fragmentation in both groups with greater fragmentation severity in SM (Figure S5). Therefore, the lowest observable exposure level (LOEL) to cause alteration in mitochondrial structure is at most 10 μM for SM and ~25 μM for controls. In addition, a subtle but significant difference \((p = 0.017)\) in baseline (vehicle exposure) fragmentation was observed between SM and control neuroprogenitors.

![Fig. 6. Decreased Cu-dependent LOEL for mitochondrial fragmentation in SM neuroprogenitors.](image)

Eight independent experiments were performed with pairs of control versus SM neuroprogenitors (CA6:SM3 n = 2, CA6:SM4 n = 2, CA6:SM5 n = 2, CB5:SM5 n = 2, total n = 8) exposed to vehicle or 10 μM Cu. Mitochondrial fragmentation was scored by a blind observer into three severity categories (minimal, moderate, or severe) for 40 PAX6 +
neuroprogenitors for each sample. For concentration comparisons * for p < 0.05, ** p < 0.01, *** p < 0.001 by t-test. Error bars represent + SEM.

Cu toxicity is associated with perturbed mitochondrial membrane potential

As an independent assessment of mitochondrial function, we assessed the mitochondrial membrane potential in CA (lines CA6 and CA11) and SM (lines SM4, SM5, and SM14) neuroprogenitors using flow cytometry. Only live single cells were included in the analysis (Figure S4). We observed that Cu exposure resulted in a decreased mitochondrial membrane potential in the majority of SM neuroprogenitors but only in a minority of CA control neuroprogenitors (see representative experiment, Fig. 7 upper panels). Similar effects were observed in 5 of 5 independent experiments. To quantify this difference in mitochondrial membrane potential, we compared the DiIC1(5) signal intensities and observed significantly decreased membrane potential in Cu-exposed SM neuroprogenitors compared to control neuroprogenitors (Fig. 7 lower panel) (paired t-test p = 0.011).

Fig. 7. Decreased mitochondrial membrane potential by Cu-exposure is more severe in SM neuroprogenitors than control. (Upper two panels) Representative flow cytometry experiment using DiIC1(5) fluorescence as an indicator of mitochondrial membrane potential after a 24 hr exposure to 50 μM Cu in CA6 and SM5. (Lower graph) Mean normalized fluorescence intensity
of Cu-exposed neuroprogenitors. Data represent single live cell events assessed by flow cytometry across 5 independent paired experiments (CA6:SM4 n = 2, CA6:SM5 n = 2, CA11:SM14 n = 1, total n = 5). Statistical analysis was performed by a paired two-tailed t-test (p = 0.011). Error bars represent + SEM.

Increased ROS production in SM neuroprogenitors prior to mitochondrial fragmentation

We hypothesized that the increased mitochondrial fragmentation of SM neuroprogenitors may be caused by increased Cu-dependent generation of reactive oxygen species (ROS). Exposure to Cu at 50, 100, and 200 μM for 30 min resulted in significantly higher levels of ROS in SM (lines SM3 and SM14) compared to CA neuroprogenitors (lines CA6 and CA11) whereas baseline ROS levels were not significantly different between cells from the two subjects (Fig. 8). Because these measurements were taken after only 30 min of Cu exposure, the elevated ROS must occur prior to Cu-induced mitochondrial fragmentation seen at 24 h. This suggests that elevated Cu-induced oxidative stress is a potential mechanism by which SM has increased Cu-dependent mitochondrial fragmentation. The baseline mitochondrial structure differences between SM/PM and control may be the cause for this differential ROS generation.

**Fig. 8.** SM neuroprogenitors have greater ROS production in the presence of Cu. ROS generation in neuroprogenitors was measured using a DCF-dye based assay after a 30 minute exposure to Cu. Paired CA/SM experiments were performed (CA6:SM3 n = 2, CA6:SM14 n = 3, CA11:SM3 n = 2, total n = 7). Statistical analysis for genotype effects was by two-way repeated measures ANOVA. Pairwise post-hoc analysis with Bonferroni correction is indicated between CA and SM hiPSC lines as * p < 0.05, *** p < 0.001. Error bars represent + SEM.

**Discussion**

Cu has been implicated in the etiology of PD, and its concentration is altered in the blood and substantia nigra of PD patients (Boll et al., 2008, Dexter et al., 1991 and Gorell et al., 1999). Our data show that increased sensitivity to Cu exposure in neuroprogenitors with high genetic risk for PD is associated with elevated ROS generation and a failure to maintain mitochondrial integrity, known toxicological mechanisms of Cu toxicity (Gyulkhandanyan et al., 2003). Our data are
consistent with recent reports examining hiPSCs from PD patients or isogenic lines with mutations in PD-related genes that report mitochondrial deficits and increased sensitivity to other disease-relevant oxidative stressors (e.g. 6-hydroxydopamine, paraquat, mane, and rotenone) in hiPSC derived neural cells (Byers et al., 2011, Cooper et al., 2012, Imaizumi et al., 2012, Nguyen et al., 2011, Ryan et al., 2013, Sánchez-Danés et al., 2012 and Seibler et al., 2011). The heightened vulnerability to Cu we report strongly implicates exposure to this metal (and perhaps Cd as well) as a potential compounding risk factor for patients with a family history of PD. These data, along with recent papers showing alterations in zinc homeostasis coupled with mitochondrial dysfunction in patient derived neuroprogenitors with ATP13A2/PARK9 mutations (Kong et al., 2014, Park et al., 2014 and Tsunemi and Krainc, 2014), strongly implicate alterations in metal biology as a major environmental modifier of genetic risk alleles for PD.

Our data support a hypothesis in which failure to maintain mitochondrial function and integrity during heavy metal exposure may contribute to gene–environment interactions underlying PD associated with mutations in PARK2. These findings are consistent with previous research demonstrating that loss of PARK2 function causes mitochondrial dysfunction and sensitivity to oxidative stress (Grünewald et al., 2010, Gyulkhandanyan et al., 2003, Mortiboys et al., 2008, Pacelli et al., 2011, Pilsl and Winklhofer, 2012, Saini et al., 2010 and Ved et al., 2005). Two studies have investigated the effects of PARK2 mutations in genetic model organisms and did not observe genotype-dependent survival differences due to Cu-supplemented food (Saini et al., 2010 and Ved et al., 2005). However, Saini and colleagues did report exacerbation of a neuronal Cu toxicity assay in Drosophila melanogaster (rough eye phenotype) carrying PARK2 mutations. This difference, relative to our observed increased vulnerability to Cu cytotoxicity in the context of PARK2mutant human neuroprogenitors, may very well be explained by systemic Cu homeostatic processes including gastrointestinal regulation of Cu uptake that would not be accounted for in a cellular model system. Future studies will be needed to resolve the complex role that systemic metal homeostatic control plays in alteration of environmental risk by PARK2 and other PD-related genetic risk factors.

We previously reported a decrease in net cellular Mn uptake following Mn exposure in SM neuroprogenitors versus control, despite equivalent Mn cytotoxicity (Aboud et al., 2012). This lack of difference in Mn cytotoxicity between control and SM neuroprogenitors appears to be due to compensatory neuroprotective changes that decrease Mn uptake in the PARK2 mutant neuroprogenitors (Aboud et al., 2012). As no differences in the accumulation of intracellular Cu were observed, the heightened sensitivity of PARK2mutant neuroprogenitors to Cu likely reflects either decreased capacity to tolerate Cu toxicity (e.g. failure of antioxidant defense mechanisms) or alterations in the biological processes thataugment mechanisms of Cu cytotoxicity (e.g. decreased Cu chaperone capacity leading to increases in free Cu levels). Further study will be needed to investigate whether this gene by toxicant interaction is specific to Cu or if it occurs for a broad range of oxidative stressors, such as iron or mitochondrial inhibitors, which are also
implicated as environmental modifiers or risk factors for parkinsonism (Peng et al., 2010, Saini et al., 2010 and Ved et al., 2005).

Conclusions

We report that neuroprogenitors from an EOPD patient and a preclinical PD patient both carrying biallelic loss-of-function mutations in \textit{PARK2} are highly susceptible to mitochondrial toxicity by a subset of heavy metal toxicants compared to neuroprogenitors from 4 different control human subjects. This is the first report of altered cytotoxic vulnerability of PD patient-derived neuroprogenitors with \textit{PARK2} mutations to widespread environmentally relevant neurotoxicants. Specifically, we found that hiPSC-derived neuroprogenitors from a preclinical PD subject with \textit{PARK2} loss of function alleles (SM) exhibit heightened vulnerability to Cu and Cd cytotoxicity but no difference in sensitivity to Mn or MeHg relative to control neuroprogenitors. The heightened sensitivity to Cu in this patient was also observed in hiPSC-derived neuroprogenitors from his brother (PM), who shares the same \textit{PARK2} susceptibility loci and other familial risk factors and who was diagnosed with EOPD. We further report that the heightened sensitivity of SM neuroprogenitors to Cu resulting in a decreased lowest observable effect level (LOEL) for mitochondrial toxicity by Cu exposure (Fig. 6). Our data support the hypothesis that genetic predisposition to PD can decrease the “safe” exposure threshold to environmental risk factors. Our data also support the potential utility of hiPSC-based neuroprogenitor model systems to define individualized environmental risk. Further research is warranted to determine whether individuals with other genetic PD risk factors may have increased risk if exposed to Cu, Cd, or other heavy metals in their environment.

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Appendix A. Supplementary data

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