The Joint Effect of Air Pollution Exposure and Copy Number Variation on Risk for Autism


Autism spectrum disorder is a complex trait with a high degree of heritability as well as documented susceptibility from environmental factors. In this study the contributions of copy number variation, exposure to air pollutants, and the interaction between the two on autism risk, were evaluated in the population-based case-control Childhood Autism Risks from Genetics and Environment (CHARGE) Study. For the current investigation, we included only those CHARGE children (a) who met criteria for autism or typical development and (b) for whom our team had conducted both genetic evaluation of copy number burden and determination of environmental air pollution exposures based on mapping addresses from the pregnancy and early childhood. This sample consisted of 158 cases of children with autism and 147 controls with typical development. Multiple logistic regression models were fit with and without environmental variable-copy number burden interactions. We found no correlation between average air pollution exposure from conception to age 2 years and the child's CNV burden. We found a significant interaction in which a 1SD increase in duplication burden combined with a 1SD increase in oxygen exposure was associated with an elevated autism risk (OR 3.4, P < 0.005) much greater than the increased risks associated with either genomic duplication (OR 1.85, 95% CI 1.25–2.73) or ozone (OR 1.20, 95% CI 0.93–1.54) alone. Similar results were obtained when CNV and ozone were dichotomized to compare those in the top quartile relative to those having a smaller CNV burden and lower exposure to ozone, and when exposures were assessed separately for pregnancy, the first year of life, and the second year of life. No interactions were observed for other air pollutants, even those that demonstrated main effects; ozone tends to be negatively correlated with the other pollutants examined. While earlier work has demonstrated interactions between the presence of a pathogenic CNV and an environmental exposure [Webb et al., 2016], these findings appear to be the first indication that global copy number variation may increase susceptibility to certain environmental factors, and underscore the need to consider both genomics and environmental exposures as well as the mechanisms by which each may amplify the risks for autism associated with the other. Autism Res 2017, 0: 000-000. © 2017 International Society for Autism Research, Wiley Periodicals, Inc.

Introduction

Autism is a behavioral disorder that has been the subject of extensive genetic studies [Jeste & Geschwind, 2014]. The incidence of ASD in the U.S., currently estimated at 1 in 56 children [Developmental Disabilities Monitoring Network Surveillance Year Principal, 2014 #41], makes this set of disorders a public health, educational, and economic concern. While autism has been determined to be highly heritable, recent estimates have elevated the proportion of variance attributable to environment. Early family studies of twins estimated the proportion of genetic contribution to autism as high as 90% [see for example, Bailey et al., 1995; Fols-stein & Rutter, 1977; Lichtenstein, Carlstrom, Rastam, Gillberg, & Anckarsater, 2010]. A more recent assessment of twins, while supporting a substantial genetic component, also indicated significant contributions.
from shared environment [Hallmayer et al., 2011]. A large longitudinal study of all births in Sweden between 1982 and 2006 that included more than 14,000 individuals diagnosed with ASD, provided evidence that genetic and environmental contributors to autism are essentially equal, each accounting for about 50% of the variability [Sandin et al., 2014]. A parallel study of this same cohort that examined SNP variants in some 3,000 subjects, arrived at similar estimates of the heritability proportion (52%), and indicated that common variants comprise the majority of genetic risk for autism [Gaugler et al., 2014]. The best estimates at present, therefore, indicate that the etiological architecture of autism includes essentially equal contributions from environment and DNA sequence variation, and both common and rare genomic variants play a role in disease risk and severity. While these studies have established the importance of both genetic and environmental factors in autism, direct measures of their relative contributions and moreover, of the interactions between them are lacking.

Copy number variants (CNVs) are one type of genomic change that contributes to autism susceptibility. The pioneering study by Sebat and colleagues found a ten-fold higher frequency of large and rare de novo variants in children with autism compared to control subjects [Sebat et al., 2007]. Subsequent work confirmed these findings [Pinto et al., 2010] and revealed that copy number burden represented in large CNVs correlated with the severity of the phenotypes found in children with neurobehavioral disorders [Girirajan et al., 2011]. Genome-wide analysis of both rare and common CNVs demonstrated that autism is associated with increased levels of copy number load, measured as base pairs of change, with a preponderance of duplications [Girirajan et al., 2013]. In addition, the level of copy number load negatively correlated with measures of communication and social skills [Girirajan et al., 2013]. Further, whole exome sequence analysis has shown that children with autism inherit more CNVs than their unaffected siblings [Krumm et al., 2013]. There is therefore good evidence that total genomic copy number burden is a meaningful measure of genomic change that contributes to autism susceptibility and a good sensor for assessing genome-environment interactions.

Phenotypic variation is an expression of several determinants, including genetic variation, environmental exposure, and the interactions between genetic variants and the environment. The magnitude and frequency of gene–environment interactions are largely unknown, and therefore the degree to which environmental impact coupled with genetic variation can explain the so-called “missing heritability” is unresolved. Current genetic studies of complex disorders often assume no appreciable gene-environment interactions, an assumption that has not been validated by experiment [Gaugler et al., 2014]. The paucity of gene–environment interaction measures has a simple origin, it requires detailed genetic and environmental data for the same group of individuals, an expensive and time consuming endeavor.

A growing body of evidence supports specific environmental contributors to autism susceptibility. In particular, prenatal air pollution exposure has come to the forefront of environmental ASD risk factors as 11 studies from the United States, using different study designs and methods, all suggest increased risk with increasing exposure [Becerra, Wilhelm, Olsen, Cockburn, & Ritz, 2013; Kalkbrenner et al., 2010, 2015; Raz et al., 2015; Roberts et al., 2013; Talbott et al., 2015a, 2015b; Volk, Hertz-Picciootto, Delwiche, Lurmann, & Mcconnell, 2011; Volk, Lurmann, Penfold, Hertz-Picciootto, & Mcconnell, 2013; Von Ehrenstein, Aralis, Cockburn, & Ritz, 2014; Windham, Zhang, Gunier, Croen, & Grether, 2006]. Criteria air pollutants, including nitrogen dioxide (NO2), particulate matter less than 10 and less than 2.5 microns in diameter (PM10, PM2.5), and ozone, are routinely monitored by the Environmental Protection Agency (EPA) and exposure to both NO2 and PM have been associated with ASD in populations from both Northern and Southern California [Becerra et al., 2013; Kalkbrenner et al., 2015; Volk et al., 2013], North Carolina [Kalkbrenner et al., 2010; Kalkbrenner et al., 2015] West Virginia [Kalkbrenner et al., 2010], Pennsylvania [Talbott et al., 2015b], and in the nation-wide Nurses Health Study [Raz et al., 2015]. When studies have attempted to identify critical time periods of exposure, pregnancy, and potentially the latter half of pregnancy appear the most important [Weisskopf, Kioumourtzoglou, & Roberts, 2015]. Few studies have attempted to include multiple criteria pollutants in the same model, though in the analyses that do, effects of NO2, PM2.5, and ozone persist [Becerra et al., 2013; Volk et al., 2013]. We are aware of only one other paper to date, that has examined genetic susceptibility together with air pollution exposure on risk of ASD [Volk et al., 2014].

This study sought to examine the joint effect of genetic susceptibility for ASD, as reflected in copy number variation, and air pollution exposure on risk of ASD in the Childhood Autism Risks from Genetics and Environment (CHARGE) study.

Methods
Description of Sample

CHARGE is a population-based case-control study of preschool children being conducted at the University of California Davis MIND (Medical Investigations of Neurodevelopmental Disorders) Institute [Hertz-Picciotto et al., 2006]. Eligibility criteria included: being between the ages of 24 and 60 months at the time of
recruitment, living with a biologic parent who speaks either English or Spanish, having been born in California, and residing within the study catchment area of approximately 1.5 hr drive to the clinic (covering ~20 counties). Our sampling frame for cases consists of children who receive services through the California Department of Developmental Services (DDS), as well as referrals from health and service providers, from other studies at the UC Davis MIND Institute, and through self-referral. Population-based controls are recruited using California Vital Statistics files of births, from which we randomly sample after frequency-matching on sex, age, and broad geographic region. Regions are large, generally encompassing multiple counties, and correspond to the catchment areas of Regional Centers that coordinate services of the California DDS. This sampling strategy ensures that confounding related to regional characteristics is minimized, and simultaneously that the cases and controls are not over-matched with regard to geographically-related exposures. Additional details on study design are provided elsewhere [Hertz-Picciotto et al., 2006]. All autism cases for CHARGE are confirmed on the Autism Diagnostic Observation Schedule (ADOS) and the Autism Diagnostic Interview-Revised (ADI-R). Typically developing controls were children who received a score <15 on the Social Communication Questionnaire and also showed no evidence of other types of developmental delay (composite scores of 70 or greater on Mullen Scales of Early Learning and Vineland Adaptive Behavior Scales). All assessments were conducted at clinics either in the MIND Institute located in Sacramento, CA or the UCLA Neuropsychiatric Institute located in Los Angeles, CA, by trained clinicians with research reliability for the instruments they administered. All components of the data collection were conducted in English or in Spanish by bilingual bicultural staff. For the current investigation, we included only those CHARGE children (a) who met criteria for autism or typical development, (b) who gave blood and agreed to have their biospecimens shared with researchers outside of the original study team, (c) for whom genetic evaluation of copy number burden passed quality control [Girirajan et al., 2013] and (d) for whom assignment of air pollution exposures based on residential addresses was successful [Volk et al., 2013]. These restrictions resulted in 158 cases of children with autism and 147 controls with typical development eligible for this analysis, which is 79% of those that passed QC for CNV calls in our previous publication [Girirajan et al., 2011]. These children were born between 1999 and 2008.

Custom Targeted Hotspot Array

Whole blood was collected for participants’ DNA samples. A custom targeted hotspot Array was used to detect CNVs as described previously [Girirajan et al., 2011]. The hotspot arrays comprised 135,000 probes, with higher density probe coverage (median probe spacing 2.6 kbp) for hotspot genomic regions, flanked by segmental duplications, and a lower probe density across the entire genome (median probe spacing 36 kbp). Hybridization, quality control and segmentation analysis were conducted as previously described [Girirajan et al., 2011, 2013]. Global changes in copy number burden were measured, i.e., bps of duplication or deletion in each individual, or collectively, as total base pairs of altered copy number (i.e., total CNV burden). In the data analysis copy number burden was evaluated as a continuous variable.

Air Pollution Exposure Assignment

Through telephone interviews, we collected demographic characteristics, medical conditions, and environmental exposures, including residential history [Hertz-Picciotto et al., 2006]. Residential histories recorded dates and address locations where the mother lived, beginning before conception through the most recent place of residence, as well as any other place of residence where the child lived. These dates and addresses were used to develop air pollution exposure metrics, as commonly implemented in large epidemiologic studies when direct measurements are not feasible [Hertz-Picciotto et al., 2006].

We used the CALINE4 line-source air quality dispersion model in order to acquire model-based estimates of traffic related air pollution (TRP) exposure derived from freeways, non-freeways, and all roads located within 5 km of each child’s home [Bensen, 1992]. Information on roadway geometry, link-based traffic volumes, period-specific meteorological conditions (wind speed and direction, mixing heights, and atmospheric stability), and vehicle emission rates were all included in the model [Volk et al., 2013]. The CALINE4 model specifically produced estimates of nitrogen oxides (NOx) which were almost perfectly correlated (around 0.99) with estimated concentrations of other traffic-related pollutants, including carbon monoxide and elemental carbon, from this same model. Therefore, our model-based pollutant concentration estimates serve as indicators of the traffic-related air pollutant mixture rather than of any specific pollutant.

In addition, exposure to PM2.5, PM10, ozone, and NO2, used regional data from the US Environmental Protection Agency Air Quality System (www.epa.gov/ttn/airs/airsaqs) supplemented for Southern California by the University of Southern California’s Children’s Health Study data for 1997 through 2009 [Volk et al., 2013]. The monthly air quality data from up to four monitoring stations located within 50 km of each location were used to create spatially interpolated estimates of exposure to these pollutants.
residence were used for spatial interpolation of ambient concentrations using inverse distance-squared weighting. If one or more stations were located within 5 km of a residence, only data from those stations were used. For PM\textsubscript{2.5}, PM\textsubscript{10}, and NO\textsubscript{2}, measurements were based on 24 hr average concentrations. For ozone, measurements were based on the average range of ozone measurements from 1000 to 1800 hr (reflecting the high 8-hr daytime exposure).

Based on child date of birth, date of conception, and reported start and end dates of each residence, the average air pollution exposure was assigned for the entire pregnancy and for the child’s first and second year of life. We also created an average exposure for each air pollutant for the period from conception to the child’s second birthday.

Ethnicity Estimation

We genotyped 100 SNPs from a custom-designed Illumina based array identified from inherited allele frequencies from four parental populations (Asian, African, European, Amerindian and Indian). To empirically estimate the proportion of ancestry attributable to a particular founding population for each individual we examined these SNPs using the program Structure [Pritchard, Stephens, & Donnelly, 2000] to derive five continuous variables reflecting each parental population. In our analyses, these variables were included as covariates.

Statistical Analysis

In order to assess the main effect contributions of CNV burden and air pollution exposure to autism susceptibility, logistic regression models were used to estimate odds ratios (ORs) and 95% confidence intervals (CIs) for each CNV burden measure and each air pollution exposure adjusted for the other. In initial models, all types of CNV burden (duplication, deletion, or total CNV burden) and air pollutants (TRP, nitrogen dioxide, ozone, PM\textsubscript{2.5}, or PM\textsubscript{10}), measured at each of three different time periods spanning pregnancy and the first 2 years of life were examined as continuous variables, with effect estimates scaled to twice the standard deviation of each exposure distribution. To control for confounding, we included as covariates: maternal education, child’s sex and race/ethnicity characterized by the Ancestry Informative Markers (AIMS).

We also examined the correlations between all pairs of CNV burden measures and air pollution exposures using Spearman’s rank correlations. This analysis aimed to determine the presence of broad evidence supporting a molecular interaction whereby prenatal air pollution might play a role in altering copy number burden.

The main analysis was to evaluate the joint effect of CNV and air pollutants on autism. For this question, we added a product term to assess interaction in the logistic regression model, first using both air pollution exposure and CNV burden as continuous variables. To increase interpretability of these results, as air pollution exposure and CNV burden are measured on vastly different scales, we z-score transformed both variables. Estimates were then scaled to reflect a 1SD increase in exposure. Finally, we reparameterized the above model by dichotomizing both CNV burden measures and air pollution exposure at the 75th percentile (the top quartile (‘high’) vs. the bottom 3 quartiles (‘low’)). This model enabled a comparison of the risk for autism among those in the top quartile of both CNV and air pollutant exposures with the risk among those in the bottom three quartiles of both CNV and air pollutant exposures. Moreover, the analyses of dichotomized CNV and air pollution measures were conducted for each of the three time periods (pregnancy, first year of life, second year of life) separately. Each of these models also adjusted for confounders of maternal education, child sex, child race/ethnicity, and Regional Center.

Results

Interactions between Air Pollution Exposure and CNV Burden Exposure

The correlations between each of the average air pollutant measures across the three time periods and each metric for CNV burden range from −0.088 to +0.081 (Table 1). None were significant, and all of these values are small, indicating that none of our three measures of global CNV has likely been impacted by early life air pollution exposures.

Results for each of the 15 different models representing three metrics of CNV and five air pollutant measures are presented in Table 2. These analyses confirm the strongest main effects from duplications, total CNV, PM10 and PM2.5, as well as two significant interactions, both involving ozone: with duplications and with total CNV. When examining the average air

<table>
<thead>
<tr>
<th>CNV burden</th>
<th>TRP</th>
<th>NO\textsubscript{2}</th>
<th>Ozone</th>
<th>PM2.5</th>
<th>PM10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duplication</td>
<td>0.063</td>
<td>0.081</td>
<td>−0.088</td>
<td>0.009</td>
<td>0.022</td>
</tr>
<tr>
<td>Deletion</td>
<td>−0.007</td>
<td>0.016</td>
<td>0.042</td>
<td>0.079</td>
<td>0.023</td>
</tr>
<tr>
<td>Total CNV</td>
<td>0.070</td>
<td>0.088</td>
<td>−0.055</td>
<td>0.066</td>
<td>0.041</td>
</tr>
</tbody>
</table>

\*All correlation measures were not statistically significant (P value > 0.05).
To visualize the risk associated with duplication burden and ozone exposure in combination we divided individuals into 16 subgroups based on quartiles of duplication burden and ozone exposure. Figure 1 shows the proportion of individuals with autism in each of the quartiles of duplication burden on the y-axis and quartiles of ozone exposure on the x-axis. In our case-control study sample with both CNV and air pollution measures, the overall proportion of cases with autism is 0.52. The subgroup from the top quartiles of both duplication burden and ozone exposure showed the highest proportion of children with autism (0.82) while the subgroup from the lowest quartile of duplication burden and ozone exposure showed the lowest proportion of autism cases (0.08). These representations of the data graphically demonstrate how the compound effect of duplication burden and ozone exposure greatly amplifies the odds of autism.

We then investigated the direction, scale, and magnitude of the CNV burden-air pollution joint effect on autism but now with variables dichotomized to reflect high vs. low exposure for both air pollution and CNV burden. Moreover, we fit models to separately examine air pollutant exposure in each of the time periods. Results from these analyses for all five air pollutants, all three measures of CNV burden, and three time periods (average pregnancy exposure and the first and second year of the child’s life) are summarized in Figure 2 and presented in detail in Table S1. First, we see that when air pollution exposures are low, a high total CNV burden (Fig. 2, estimates in red) is consistently associated with autism (bottom panel for total CNV burden). These estimates reflect increases in autism risk for the top quartile of total CNV burden with a range from OR = 1.5 to 3.5, with most models having OR’s near 2.5. Second, we find that when CNV burden is low, specifically deletions, high exposures to PM2.5 are associated with autism risk (Fig. 2, estimates in blue, for example: comparing high vs. low PM2.5 exposure in postnatal years 1 and 2, respectively, OR = 2.8, 95% CI (1.1–7.0), and OR = 2.8 (1.1, 7.7)). The associations between autism risk and high exposures to PM10 during pregnancy are consistently elevated among those with low CNV (OR’s range from 2.58 to 2.98), whether measured as deletions, duplications, or their combination.

Third, results for interactions were similar to the models based on continuous metrics: joint exposures to both high duplication burden and high ozone, as compared with jointly low exposures to both, were associated with substantially elevated risks for autism (Fig. 2, estimates in purple). For example, children with high duplication burden who were also exposed during the prenatal period to high ozone levels had nearly a three-fold greater risk to develop autism (OR = 2.8 (1.2–6.9)). Even stronger associations were seen for children with...
high ozone exposures during the first and second years of life, with $\text{OR} = 4.2$ (1.5–11.7) and $\text{OR} = 3.4$ (1.4–8.6) (Fig. 2/Table S1). For the other pollutants, little evidence of interaction was found. Thus we observe a consistent pattern for ozone, as compared to sporadic suggestions of slightly elevated risks for joint exposures involving the other pollutants.

We also conducted a sensitivity analysis by adding maternal age to the models with quartiles of CNV, air pollution and the combination of the two. The impact on the OR’s and their 95% CI’s were negligible (<0.5% change), likely because the models were already adjusted for maternal education, child’s race/ethnicity, and Regional Center; maternal age and education are highly correlated in the CHARGE Study.

**Discussion and Conclusions**

**Contributions of Interaction Involving Environmental and Genetic Risk Factors for Autism**

There remains an active debate in the research community on the relative contributions of environmental and genetic factors in autism susceptibility, as well as the degree to which interactions might account for the “missing heritability”, namely, our inability to account for even a majority of autism cases with all known genomic variants. Large studies are converging on estimates of heritability in the ranges of 35–60% [Hallmayer et al., 2011; Rosenberg et al., 2009; Gaugler et al., 2014; Sandin et al., 2014]. As new approaches to environmental exposures, including analyses of biological and environmental samples using non-targeted chemical analyses, are developed and begin to be used in research on ASD etiology, and more comprehensive analyses of interactions are conducted, assessments of the relative contributions from genetics and environment will become feasible. Nevertheless, the question of comparing relative separate contributions from genes and environment may need to be recast when considering a sizable proportion of the risk for autism may be influenced by both types of factors, and moreover, the impacts of environment, broadly defined to include, e.g., nutrition, stress, health of the pregnant mother, the microbiome, and so forth, on gene expression.
mediated by epigenetic modifications are quite likely larger than previously imagined.

Correlations between air pollutant exposure and CNV burden were small. Prenatal exposures could only be expected to influence the de novo mutations, and not those carried by the parents in their somatic cells. However, the proportion of CNVs that are inherited vs. de novo is not clear and we were unable to make estimates, as we did not obtain CNV information from the parents. Hence, we are unable to draw any conclusions regarding the hypothesized pathway in which prenatal air pollution exposures might contribute to altered total CNV burden in the child. If de novo mutations are not a large proportion of the CNV burden, then any impact

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**Figure 2. Estimated effects from genetic copy number variant (CNV) load and from air pollutant exposures, acting separately and in combination, on risk for autism.** Five broad columns represent five different types of air pollutant measurements, and three rows represent three different metrics for assessing CNV burden. For each cell representing a given air pollutant and CNV metric, three time periods for the air pollutant exposures are examined: pregnancy, 1st postnatal year, and 2nd postnatal year. The colored dots and bars are, respectively, the odds ratios and 95% confidence intervals for associations with autism versus typical development: a) comparing high to low environmental air pollutant exposures (blue) among those with low CNV burden; b) comparing high to low CNV burden (red) among those with low air pollutant exposures; and c) comparing the combined high CNV and high air pollutant exposures to the combined low CNV and low air pollutant exposures (purple). For both CNV burden (genetics) and air pollutant exposure (environment), high is defined as the upper quartile and low as the bottom three quartiles. These results from this comprehensive analysis confirm previous findings that the metric for CNV (red) that is most often significant is the total burden summing both deletions and duplications (bottom row of results). The major new findings demonstrate the following clear patterns: First, similar to the models based on continuous metrics for both CNV and air pollution, the interactions (purple) that emerge as most dramatic and with the strongest significance are for ozone, particularly when combined with duplications, but also with total CNV burden. Thus those in the upper quartile of both CNV and ozone exposure are at exceptionally high risk for ASD. Second, the results underscore the impact of taking account of environment on the magnitude of the genetically induced risk; for example, the most significant genetic main effects (red) are in models with PM10 or PM2.5. Among those with low ozone, duplications appear to have virtually no impact on ASD risk, but among those with high ozone exposures, have an exceptionally strong effect. Third, the strongest environmental main effects are: NO2 in the first year of life among those with low duplication burden; ozone during pregnancy among those with low deletions; PM10 during pregnancy in those with any low CNV (deletions, duplications or both combined); and PM2.5 during any time period in those with low deletion burden or low total CNV.

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**Figure 2**

*Estimated based on CALINE line source dispersion model, described in text.*
on de novo mutations from air pollution might have only a small effect on the total CNV burden, and this study would have had very little power to detect that impact. On the other hand, environmentally-induced de novo CNVs in just a few critical genes (for instance, in synaptogenesis pathways), might have large ramifications for ASD risk. Thus further work, in which de novo and inherited CNV are distinguished is warranted.

Our results do support a second type of interaction, namely, one in which ozone exposure may further exacerbate risk for autism in children with susceptibility arising from genetic instability. This finding needs to be viewed in the context of the full set of analyses (three different measures of CNV and five components of air pollution; air pollutant exposures in several time windows, and both continuous and dichotomized variables for both CNV and air pollution). First, we note that many of the pollutants are correlated, as are duplications and total CNV burden, and therefore the results across models are not independent of each other. For this reason, we have elected to report all of the findings, and to examine the findings that did emerge for consistency, since sporadic significance is more likely to be due to chance. Whether looking at ozone as a continuous measure or comparing the top quartile to the bottom three quartiles combined, the pattern of results is similar. Emphasizing only those results supported in several models, the salient observation from this study of gene-by-environment interaction is an ozone association with autism only among individuals who have a high CNV burden. The ozone association with autism in the presence of greater CNV burden stands in contrast to results from other pollutants, which have tended to show consistent associations in the literature with increased risk of ASD when ignoring genetic susceptibility, i.e., in populations as a whole [Becerra et al., 2013; Kalkbrenner et al., 2010; Kalkbrenner et al., 2015; Roberts et al., 2013; Talbott et al., 2015a,b; Volk et al., 2013; von Ehrenstein et al., 2014]. This distinctive pattern is a coherent one in light of a priori scientific evidence establishing a negative correlation of ozone with the other pollutants examined. Notably, ozone overall, or in the absence of high CNV burden, showed essentially no association with autism risk. An association between ozone exposure and autism risk was not identifiable without taking genetic susceptibility into account. This result highlights how both genes and environment risk can contribute to ASD etiology, and highlights the value of incorporating measures of genomic susceptibility into environmental studies.

Mechanisms of Ozone-Copy Number Burden Interactions

Of the airborne pollutants examined in this study, ozone showed the strongest interactions with CNV burden. What might be the mechanism of this interplay? Ozone is a potent oxidizing agent, known to produce reactive oxygen species (ROS) and cellular damage at exposure levels that can be found in urban environments [Devlin et al., 1991]. In model systems, the oxidative and cellular stress produced by ozone has been shown to compromise neural gene expression, function, and behavior [Rivas-Arancibia et al., 2010]. There is considerable evidence that autism is associated with elevated levels of oxidative stress both in peripheral blood [Gorrindo, Lane, Lee, Mclaughlin, & Levitt, 2013] and in the brain [Rossignol & Frye, 2014]. Reductions in anti-oxidant molecules that serve to scavenge ROS are also associated with autism [Frye et al., 2013], potentially producing a state sensitive to further oxidative stress. Mitochondrial abnormalities, also found associated with autism, provides another mechanism of affecting oxidative stress in these children [Giulivi et al., 2010]. Given the elevated levels of oxidative stress in children with autism it is plausible that ozone exposure potentiates an already compromised metabolic condition, producing a genetic-environment interaction of some magnitude. It is possible that CNVs contributing to autism affect genes altering production of ROS or compromising responses to oxidative stress. For example, a mouse model of Rett Syndrome, shows abnormalities in both oxidative burden and mitochondrial function [Grosser et al., 2012]. In this instance our findings suggest that air pollutant exposure may work as an additional risk factor through one or more pathways affected by CNVs.

Our study expands the limited body of work examining joint environmental and genetic risks for ASD. The joint effects we see between CNV burden and pollution exposure demonstrate an interaction, even when one of the factors has little to no association in the absence of the other. This example demonstrates that exposures with small main effects can also contribute to autism susceptibility via synergism with genetic factors, and that these interactions may be missed when limiting interaction analyses to exposures and genes with significant independent effects. This scenario may be parallel to the numerous examples of genomic regions or loci that showed marginal significance after multiple testing corrections but that have been replicated in larger cohorts or subsequent studies [Hamshere et al., 2013; Athanasius et al., 2010; Denny et al., 2010; Pendergrass et al., 2013; Hall et al., 2014].

It is striking that the only significant GxE interaction found in the quartile analyses was with ozone, which itself showed no main effect in the CHARGE Study, while pollutants with strong main effects showed no interaction. A likely explanation is that for the other pollutants, CNV burden simply did not amplify susceptibility given an already high effect of air pollution on
autism risk. Studies of air pollutants and ASD have not often examined the effect of ozone. To date, ozone has been examined in a report from Taiwan that identified an increased risk for ASD [Jung, Lin, & Hwang, 2013] and was associated with increased ASD risk when modeled along with PM$_{2.5}$ in a large case-control study from Los Angeles County [Becerra et al., 2013]. Investigation of ozone exposure on the brain has been relatively limited, with data largely coming from the field of cognitive aging, and demonstrating effects which implicate poorer performance over time [Chen & Schwartz, 2009; Gatto et al., 2014]. Thus, our findings might represent a chance fluctuation, or it may represent a very novel clue regarding conditions (i.e., vulnerabilities) under which ozone does influence early development and possibly also late cognitive decline.

One limitation to the current study is the small sample size ($n = 305$), which may have limited our statistical power for subtle interactions. Only two pollutants, ozone and PM$_{10}$, demonstrated any joint effect and only an ozone interaction was identified over multiple time points examined and with two measures of CNV burden. It should be acknowledged that ozone measures over time are highly correlated, which could suggest lack of specificity in the importance of timing for such exposures. Similarly, total CNV burden also includes duplication burden making it a less-specific measure of genetic susceptibility. As a first analysis of CNV-by-environment interactions, we selected an objectively measured exposure and controlled for only a few potential confounding factors, namely the sociodemographic factors of maternal education, child’s sex and race/ethnicity, and the Regional Center of their residence. Most factors we screened did not exert any confounding effect and after selection of the final model we conducted a sensitivity analysis and determined that maternal age had virtually no impact on any of our results (main effects or interactions); we recognize that identifying covariates that might be associated with the marginal and/or joint distributions of CNV burden and air pollution exposure, as well as with autism status, is uncharted territory. Hence, we cannot exclude a role for confounding by other known, suspected, or unknown risk factors for autism, which could have either masked an interaction with one or more of the other pollutants, or inflated the interaction with ozone. Another limitation is that this analysis was underpowered for more restricted time windows, such as trimesters of pregnancy. Since brain development requires highly orchestrated sequences of processes, it is possible that only certain critical periods are vulnerable to specific air pollutants. Indeed, in our earlier work, we found air pollution in the third trimester and first year of life to be more strongly related to development of ASD than earlier periods [Volk et al., 2013]. Despite these limitations, our findings can help direct future research confirming and extending these relationships.

Our findings emphasize the value and importance of pursuing gene–environment interactions in both candidate and broad discovery modes, which require collecting both genomic and environmental data for large samples in order to discover new interactions or replicate existing interactions. A broad assessment of environmental factors, together with a comprehensive measurement of diverse genetic variants (SNPs, CNVs, etc.) in data resources with sufficient statistical power will likely be necessary to better quantify the joint effects of susceptibility factors for autism. Overall, the findings reported here show that genetic–environment interactions can be substantial and may contribute significantly to risk for complex disorders.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s website.