The influence of selected genetic and environmental factors on white matter pathway structure measured with diffusion tensor imaging

Raquel Fenoll Sanguino
Airborne copper exposure in school environments associated with poorer motor performance and altered basal ganglia

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Abstract

Introduction: Children are more vulnerable to the effects of environmental elements. A variety of air pollutants are among the identified factors causing neural damage at toxic concentrations. It is not obvious, however, to what extent the tolerated high levels of air pollutants are able to alter brain development. We have specifically investigated the neurotoxic effects of airborne copper exposure in school environments. Methods: Speed and consistency of motor response were assessed in 2836 children aged from 8 to 12 years. Anatomical MRI, diffusion tensor imaging, and functional MRI were used to directly test the brain repercussions in a subgroup of 263 children. Results: Higher copper exposure was associated with poorer motor performance and altered structure of the basal ganglia. Specifically, the architecture of the caudate nucleus region was less complete in terms of both tissue composition and neural track water diffusion. Functional MRI consistently showed a reciprocal connectivity reduction between the caudate nucleus and the frontal cortex. Conclusions: The results establish an association between environmental copper exposure in children and alterations of basal ganglia structure and function.
Copper is an air pollutant that may potentially interfere with brain development. Although this element is necessary for cellular metabolism, abnormal copper levels lead to relevant brain impairment (Scheiber et al. 2014). The deleterious effect of an excess of copper is well-known from Wilson’s disease, an inherited metabolic disorder affecting the basal ganglia with symptoms usually evident from preadolescence onward and mainly in the form of deficient motor control (Bandmann et al. 2015).

We have tested the potential harmful effects of copper as an air pollutant in children aged from 8 to 12 years using specific motor testing and brain structural and functional imaging. Exposure to airborne copper at school was estimated for 2836 children in the city of Barcelona. Both the speed and consistency of motor response were measured to test the integrity of motor function in the whole sample. High-resolution 3D MRI, DTI (diffusion tensor imaging), and functional MRI were used to directly test the potential repercussion on brain anatomy, architecture, and function in a subgroup of 263 children.

**Methods**

**Participants**

This study was developed in the context of the BREATHE project (The European Commission: FP7-ERC-2010-AdG, ID 268479) aimed at assessing the impact of long-term exposure to traffic-related air pollutants on school children. A total of 2897 children participated in the whole survey from 39 representative schools of the city of Barcelona (Sunyer et al. 2015). A group of 2836 children completed the behavioral testing (mean age at the study end, 9.4 years; SD 0.9; and range 7.9–12.1 years) and served to test the effect of copper on motor function. From this sample, 1564 families were invited to participate in the MRI study via post, e-mail, or telephone, and 810 of them gave an initial positive response. The recruitment of this group was consecutive with the aim of including children from all participating schools. Parents of 491 children were directly contacted. Consent to participate was finally not obtained in 165 cases, 27 children were lost before the assessment and 21 children were not eligible because of dental braces. The finally selected MRI group included 278 participants, 263 of whom completed the imaging protocol (mean age of 9.7 years, SD 0.9 and range, 8.0–12.1 years) and served to test direct effects of copper on brain. Study design and participant selection is fully described in Sunyer et al. (2015). Table 1 reports the characteristics of the study samples.

All parents or tutors signed the informed consent form approved by the Research Ethical Committee (No. 2010/41221/I) of the IMIM-Parc de Salut MAR, Barcelona, Spain, and the FP7-ERC-2010-AdG Ethics Review Committee (268479-22022011).

**Copper exposure**

Each school was measured twice during 1-week periods separated by 6 months, in the warm (year 2012) and cold

<table>
<thead>
<tr>
<th>Table 1. Characteristics of study samples.</th>
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<tbody>
<tr>
<td>Gender</td>
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<tr>
<td>Age, years, mean ± SD (range)</td>
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<tr>
<td>Overall school achievement, 5-point scale</td>
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<tr>
<td>Difficulties score (SDQ), range 0–40</td>
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<tr>
<td>Obesity: Normal</td>
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<tr>
<td>Overweight, BMI ≥ 25–29.9 kg/m²</td>
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<tr>
<td>Obesity, BMI ≥ 30 kg/m²</td>
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<tr>
<td>Father education (5-point scale, 5 = University)</td>
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<tr>
<td>Vulnerability index¹–Home</td>
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<tr>
<td>Vulnerability index¹–School</td>
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<tr>
<td>Public/Nonpublic school</td>
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<tr>
<td>Task performance</td>
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<tr>
<td>Reaction time (mean of medians, msec)</td>
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<tr>
<td>Reaction time standard deviation (msec)</td>
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<td>Commission errors (number)</td>
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</table>

BMI, body mass index; SDQ, Strengths and Difficulties Questionnaire.

¹Neighborhood socioeconomic status vulnerability index based on the level of education, unemployment, and occupation at the census tract (Atlas de vulnerabilidad urbana de España, 2012).
(year 2012/2013) periods. Indoor air in a single classroom and outdoor air in the courtyard were measured simultaneously. Several pollutants were measured during class time (Amato et al. 2014; Rivas et al. 2014; Sunyer et al. 2015). Copper specifically was measured during 8 h (09:00–17:00 h) in particulate matter with an aerodynamic diameter <2.5 μm (PM2.5) collected on filters with high-volume samplers (MCV SA, Spain). After acid digestion of the filter, copper concentrations were determined via ICP-MS (inductively coupled plasma mass spectrometry). Yearly school air pollution levels were obtained by averaging the two 1-week measurements. This study was focused on outdoor (courtyard) copper measurements, as they are more directly related to urban pollution, which is the primary interest of this project. All children had been in the school for more than 18 months (and 98% more than 2 years) at imaging assessment, which was carried out after the pollution measurement campaigns.

Behavioral measurements

Behavioral assessment was focused on testing speed and consistency of motor responses. We used the computerized “Attentional Network Test,” child version (Child ANT) (Rueda et al. 2004), which was developed to specifically assess children’s motor speed during attentional challenge. Among the ANT measurements, we used the overall “reaction time” to measure speed and “reaction time standard deviation” to measure trial-to-trial consistency of motor responses. A higher reaction time standard deviation indicates higher reaction time variability, which increases with reduced executive and attentional resources depending on the integrity of frontal-basal ganglia circuits (Bellgrove et al. 2004; MacDonald et al. 2006; Langner and Eickhoff 2013). Normal children become efficient in sustained motor responses during the preadolescent age period contemplated in our study (MacDonald et al. 2006).

The procedure used to administer the task is fully described in a previous report (Forns et al. 2014). The Child ANT takes approximately 20 min to complete. Groups of 10–20 children were assessed together wearing ear protectors and were supervised by one trained examiner per 3–4 children. The task required to respond to a target (yellow-colored fishes) presented on a computer screen by pressing a key. The whole task contains cues to test different attentional aspects, but outcome measurements for the current study were limited to median reaction time and standard deviation expressed in milliseconds from the correct responses obtained across the 128 trials of the task (four separate blocks of 32 trials each). Commission and omission errors were also registered. Cases with more than 30% commission or omission errors were excluded from further analyses (total nine cases in the whole sample and two cases in the MRI sample).

Contextual behavioral assessment

Sociodemographic factors were measured using a neighborhood socioeconomic status vulnerability index (based on the level of education, unemployment, and occupation at the census tract) (Atlas de Vulnerabilidad Urbana de España, 2012; http://www.fomento.gob.es/.../AtlasVulnerabilidadUrbana/) according to both school and home address. Parental education was registered for both parents using a 5-point scale (1 illiterate/2 less than/3 primary/4 secondary/5 university). Standard measurements of height and weight were performed to define overweight and obesity (de Onis et al. 2009). Parents completed the SDQ (Strengths and Difficulties Questionnaire) on child behavioral problems (Goodman 2001). A “difficulties” score ranging from 0 to 40 was generated. Overall school achievement was rated by teachers using a 5-point scale (from the worse = 1 to the best = 5).

MRI acquisition

A 1.5-Tesla Signa Excite system (General Electric, Milwaukee, WI) equipped with an eight-channel phased-array head coil and single-shot EPI (echoplanar imaging) software was used. The imaging protocol included high-resolution T1-weighted 3D anatomical images, DTI, and a functional MRI sequence acquired in the resting state.

High-resolution 3D anatomical images were obtained using an axial T1-weighted three-dimensional fast spoiled gradient inversion recovery prepared sequence. A total of 134 contiguous slices were acquired with repetition time 11.9 msec, echo time 4.2 msec, flip angle 15°, field of view 30 cm, 256 × 256 pixel matrix, and slice thickness 1.2 mm.

Diffusion tensor imaging was obtained using spin-echo single-shot echo-planar sequences of 25 directions with a B-factor of 1000 sec/mm². Twenty-six slices were acquired with repetition time 8300 msec, echo time 94 msec, thickness 5 mm, no gap, pulse angle 90°, field of view 26 cm, and 128 × 128 acquisition matrix reconstructed into a 256 × 256 matrix.

The functional MRI sequence consisted of gradient recalled acquisition in the steady state with repetition time 2000 msec, echo time 50 msec, pulse angle 90°, field of view 24 cm, 64 × 64-pixel matrix, and slice thickness 4 mm (interslice gap, 1.5 mm). Twenty-two interleaved slices were prescribed parallel to the anterior–posterior commissure line covering the whole brain. A 6-min
continuous resting-state scan was acquired for each participant. Children were instructed to relax, stay awake, and lie still without moving, while keeping their eyes closed throughout. This scan generated 180 whole-brain EPI volumes. The first four (additional) images in each run were discarded to allow magnetization to reach equilibrium.

**Image preprocessing**

**Anatomical 3D**

All the anatomical images were visually inspected before analysis by a trained operator to detect any motion effect. A total of 10 children were discarded as a result of poor quality images and thus the final sample for the 3D anatomical analysis included 253 children. Anatomical 3D data were processed in two separate analyses assessing different anatomical characteristics.

Gray and white matter tissue concentration and volume at a voxel level were measured using Statistical Parametric Mapping (SPM8) (http://www.fil.ion.ucl.ac.uk/spm, Wellcome Department of Cognitive Neurology, London, UK, 2008). SPM VBM (voxel-based morphometry) algorithms with DARTEL registration were used with the following processing steps: segmentation of anatomical images into gray and white matter tissue probability maps in their native space; estimation of the deformations that best align the images together by iteratively registering the segmented images with their average; finally, generating spatially normalized and smoothed segmentations (5 x 5 x 5 FWHM) using the deformations estimated in the previous step. The analyses were performed with scaling by Jacobian determinants (estimates of volume change during the normalization) to consider tissue volume and without Jacobian scaling to assess the relative concentration of gray matter and white matter. Normalized images were finally transformed to the standard SPM template, resliced to 1.5 mm resolution in MNI (Montreal Neurological Institute) space.

Cortical thickness measurements across the whole cortex were obtained using FREESURFER tools (http://surfer.nmr.mgh.harvard.edu/). Processing steps included removal of nonbrain tissue, segmentation of the subcortical white matter and deep gray matter volumetric structures, tessellation of the gray and white matter boundary, registration to a spherical atlas which is based on the individual cortical folding patterns to match cortical geometry across subjects, and creation of a variety of surface-based data. Cortical thickness is calculated as the closest distance from the gray/white boundary to the gray/CSF boundary at each vertex on the tessellated surface (Fischl et al. 1999).

**Diffusion tensor imaging**

Diffusion tensor imaging was processed using FMRIB (functional MRI of the brain) Software Library 5.0 (FSL), developed by the Analysis Group at the Oxford Centre for FMRIB (Smith et al. 2004). Diffusion-weighted images were corrected for motion and eddy current distortions (“Eddy Current Correction” option in the FMRIB Diffusion Toolbox [FDT] version 2.0 in FSL), and a whole-brain mask was applied using the FSL Brain Extracting Tool. A further rigorous image quality control was carried out to identify potential residual effects of head motion, which involved the visual inspection of each DTI slice for all 25 DTI volumes in all participants. Volumes with slices with signal loss (greater than ~10%) or residual artifacts were identified by an expert researcher. DTI full examinations showing one or more degraded images in more than five volumes were discarded. A total of 76 children were removed from the DTI analysis on the basis of this criterion (in addition to 10 cases showing gross image degradation). The final DTI sample involved 177 children showing a mean ± SD of 23.5 (94%) ± 1.9 optimal quality volumes. Subsequently, we estimated FA (fractional anisotropy) maps using FDT in FSL after local fitting of the diffusion tensor model at each voxel (“dtifit”). Next, diffusion data were processed using tract-based spatial statistics (Smith et al. 2006). Each FA dataset was resliced to a 1 mm x 1 mm x 1 mm anatomical resolution and normalized to standard MNI space via the FMRIB58_FA template using the FMRIB’s nonlinear registration tool.

To complement the analysis and assist interpretation of the FA results, we carried out a directional analysis of water diffusion along each (x, y, z) axis separately. We used the diffusion tensor estimated by FSL “dtifit” algorithm to determine diffusion strengths. Geometrically, the procedure involved determining the radius of the arbitrarily oriented ellipsoid along the spatially fixed x, y, and z axes using basic 3D quadratic geometry. As the eigenvectors and eigenvalues were previously normalized to MNI space in all subjects, we were able to carry out group-level voxel-wise analyses.

**Functional MRI**

Preprocessing was carried out using SPM8 and involved motion correction, spatial normalization, and smoothing using a Gaussian filter (full-width half-maximum, 8 mm). Data were normalized to the standard SPM-EPI template and resliced to 2-mm isotropic resolution in MNI space.

The following procedures were adopted to control for potential head motion effects: (1) Conventional SPM time-series alignment to the first image volume in each
subject. (2) Exclusion of 24 children with large head motion (boxplot-defined outliers) (Pujol et al. 2014b). The finally analyzed sample therefore included 239 children. (3) Both motion-related regressors and estimates of global brain signal fluctuations were included as confounding variables in first-level (single-subject) analyses. (4) Within-subject, censoring-based MRI signal artifact removal (Power et al. 2014) (scrubbing) was used to discard motion-affected volumes. For each subject, interframe motion measurements (Pujol et al. 2014b) served as an index of data quality to flag volumes of suspect quality across the run. At points with interframe motion >0.2 mm, that corresponding volume, the immediately preceding and the succeeding two volumes were discarded. Using this procedure, a mean ± SD of 11.2 (6.2%) ± 13.8 volumes of the 180 fMRI sequence volumes were removed in the analyzed sample. (5) Potential motion effects were further removed using a summary measurement for each participant (mean interframe motion across the fMRI run) as a regressor in the second-level (group) analyses in SPM (Pujol et al. 2014b).

Functional connectivity maps were generated using procedures detailed in previous reports (Harrison et al. 2013; Pujol et al. 2014a). Maps representative of frontal lobe functional connectivity were obtained by locating seed regions at the medial-dorsal, lateral (right and left), and medial-anterior aspects of the frontal cortex using coordinates taken from previous literature (Fox et al. 2005) converted to MNI: medial-dorsal (x = −2, y = −2, z = 55), right lateral (x = 45, y = 3, z = 15), left lateral (x = −45, y = 5, z = 9), and medial-anterior (x = 1, y = 54, z = 26). Additional maps were generated from the results obtained in the structural (anatomical and DTI) analyses. The seed regions were centered at the left caudate nucleus at three anterior–posterior levels covering the part of the head of the caudate nucleus showing significant copper effects on its structure (MNI coordinates [x = −12, y = 20, z = 4], [x = −12, y = 14, z = 9], and [x = −12, y = 8, z = 12]).

For each of the striatal locations, the seed region was defined as a 3.5-mm radial sphere (sampling ~ 25 voxels in 2 mm isotropic space). This was performed using MarsBar ROI (region of interest) toolbox in MNI stereotaxic space (Brett et al. 2003). Signals of interest were then extracted for each seed region, respectively, by calculating the mean ROI value at each time point across the time series. To generate the seed maps, the signal time course of a selected seed region was used as a regressor to be correlated with the signal time course of every voxel in the brain in order to generate first-level (single-subject) voxel-wise statistical parametric maps (contrast images). The maps were estimated for each seed separately. A high-pass filter set at 128 sec was used to remove low-frequency drifts below ~0.008 Hz. In addition, we derived estimates of white matter, CSF, and global brain signal fluctuations to include in the regression analyses as nuisance variables.

Statistical analysis

A multiple linear regression was used to estimate the source of copper and the relative contributions were given as standardized β values. A linear regression was used to estimate the association of copper measurements with motor performance. β values are reported as time increments (msec) for each copper measurement unit (ng/m3).

Imaging data were analyzed using SPM. Individual anatomical (Jacobian-modulated and nonmodulated white and gray matter and cortical thickness), DTI, and functional connectivity maps were included in second-level (group) analyses to map voxel-wise the correlation across-subjects between individual brain measurements and individual copper exposure (the measurements obtained in the school of each participant). Results were considered significant with clusters of 1.032 mL (e.g., 129 voxels with a resolution of 2 × 2 × 2 mm) at a height threshold of P < 0.005, which satisfied the FWE (family-wise error) rate correction of P_{FWE} < 0.05 according to recent Monte Carlo simulations (Pujol et al. 2014c). Maps in figures are displayed at t > 2.3.

Results

Copper as an air pollutant

Copper measured in fine particles (PM_{2.5}) from school playgrounds showed a mean of 8.7 ng/m^3 (SD, 3.0; range 3.7–13.8). According to the correlation with specific tracers, the main source of copper was road traffic, but a significant contribution was the result of industrial activity. An additional third source was identified in relation to the close proximity (mean ± SD, 90 ± 58 m) of seven schools to busy overhead-wire railway lines. For instance, a multiple regression accounted for 70% variance of copper measurements including elemental carbon as a single road traffic tracer (Amato et al. 2014) (standardized β = 0.66, P < 0.00001), zinc as a single industry tracer (Amato et al. 2014) (standardized β = 0.31, P = 0.002), and train proximity (standardized β = 0.20, P = 0.037).

Relationship of copper exposure with children’s performance

Higher copper exposure was associated with poorer motor performance in children. Although in terms of the whole sample (n = 2827 after nine exclusions), the
association was significant for motor response speed (reaction time, $\beta = 2.2$ and $P = 0.006$), the effect was more robust on motor response consistency (reaction time standard deviation, $\beta = 2.9$ and $P < 0.00001$). Such a negative relationship with reaction time variability was significant in the group of children receiving MRI ($n = 261$ after two exclusions; $\beta = 4.2$ and $P = 0.026$).

**Neuroimaging results**

Three-dimensional anatomical (T1-weighted) MRI and DTI were used to assess the potential association of copper exposure with alteration in the fine brain structure. Copper was associated with higher gray matter concentration (i.e., a higher proportion of gray matter in the tissue) in the striatum, specifically in the caudate nucleus (Fig. 1 and Table 2), with no effect on tissue volume. This finding potentially expresses a relative reduction of striatum white matter (i.e., of white matter “striae” that actually give the corpus striatum its anatomical name). No other significant alterations were identified with 3D anatomical MRI with the exception of an area of increased cortical thickness in the supplementary motor area of the left hemisphere (data not shown).

Results from the DTI analysis were highly consistent with the anatomical results (Fig. 2, Table 2). Copper was associated with an increase of neural tissue FA. This DTI measurement may express the extent to which an anatomical structure is composed of white matter tracts showing one dominant direction. In brain regions containing tracts with a single direction, FA increases as a result of brain maturation. Nevertheless, in complex structures with tracts crossing in different directions, higher FA may denote less mature or less structured tissue (Douaud et al. 2011; Jones et al. 2013). In our analysis, higher copper levels were associated with higher FA in white matter close to the caudate nucleus and in the caudate nucleus itself. This region is anatomically characterized as showing superior–inferior, posterior–anterior, and lateral–medial crossing white matter tracts (see Fig. 3 and Kotz et al. 2013). Within this scenario, a less mature structure will show higher FA (the effect of copper in our study).

A voxel-wise correlation analysis with behavior measurements helped to establish the nature of the identified changes associated with copper exposure. Reaction time and, mostly, reaction time variability showed significant positive correlation with FA of white matter adjacent to the caudate nucleus (Fig. 2). Thus, in the direction of copper exposure findings, slower children and the children with less consistent motor response exhibited higher FA in this region.

To investigate copper effects on the neural track architecture in the caudate nucleus region, we analyzed tissue diffusion along the three orthogonal ($x$, $y$, $z$) directions separately. Higher copper exposure was associated with a complex combination of diffusion changes in the caudate nucleus region, with distinct sectors showing reduced diffusion in one or more directions (Fig. 4). In other words, copper was related to changes in the complex architecture of neural tissue diffusion, further supporting the notion that white matter pathways in the caudate nucleus region may be affected by copper exposure.

The functional significance of the identified basal ganglia changes was further investigated by assessing functional connectivity in the basal ganglia network. Functional connectivity MRI maps representative of frontal-basal ganglia circuits were generated using coordinates taken from previous works and from the current anatomical and DTI results centered at the caudate nucleus (see Methods). The most relevant finding was the association of higher copper exposure to a reciprocal reduction of functional connectivity between the caudate nucleus and the frontal lobe operculum bilaterally (Fig. 5). This association was consistent with the anatomical and DTI results and notably specific in terms of functional anatomy. Remarkably, reduced caudate-to-frontal operculum connectivity was the only significant finding in three functional connectivity maps (see Table 2).

The effect of potential confounders was tested for each significant finding including age, sex, academic achieve-
ment, academic difficulties score, obesity, parental education, home and school vulnerability index, and public/nonpublic school category as covariates. No single confounder or combination showed a significant effect. That is, decreases in $\beta$ estimates after the inclusion of confounders in a regression model were very small with no variables affecting the primary results with $\beta$ reductions >10%. Finally, potential alterations associated with Mn (manganese) were also investigated, as this element may be associated with basal ganglia alterations. We did not, however, find any significant finding associated with manganese. Moreover, the effect of copper had a tendency to be more robust when adjusted by Mn (e.g., the primary correlation between copper and motor response consistency showing $\beta = 2.9$ in the whole sample, showed $\beta = 3.3$ after entering Mn measurements as a covariate). Similarly, the effect of copper remained (no $\beta$ decrease > 10%) after additionally adjusting by other representative elements (Table 3, i.e., C, Pb, Fe, and Sb).

### Table 2. MRI correlation results.

<table>
<thead>
<tr>
<th></th>
<th>Cluster size, mL</th>
<th>$x$, $y$, $z$</th>
<th>$t$</th>
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<tbody>
<tr>
<td><strong>Copper versus gray matter concentration</strong></td>
<td></td>
<td></td>
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<tr>
<td>R Caudate nucleus-positive correlation</td>
<td>4.09</td>
<td>8, 19, 3</td>
<td>3.12</td>
</tr>
<tr>
<td>L Caudate nucleus-positive correlation</td>
<td>4.09 (same cluster)</td>
<td>$-3$, 16, 6</td>
<td>3.12</td>
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<tr>
<td><strong>Copper versus fractional anisotropy</strong></td>
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<td></td>
<td></td>
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<tr>
<td>R Caudate nucleus-positive correlation</td>
<td>1.40</td>
<td>11, 9, 9</td>
<td>3.8</td>
</tr>
<tr>
<td>L Caudate nucleus-positive correlation</td>
<td>9.23</td>
<td>$-8$, 9, 10</td>
<td>3.9</td>
</tr>
<tr>
<td>L Supracaudate white matter-positive correlation</td>
<td>9.23 (same cluster)</td>
<td>$-17$, 20, 25</td>
<td>3.6</td>
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<tr>
<td>L Suprathalamic white matter-positive correlation</td>
<td>9.23 (same cluster)</td>
<td>$-20$, $-17$, 28</td>
<td>3.4</td>
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<td>R Corpus callosum-positive correlation</td>
<td>1.90</td>
<td>17, $-40$, 13</td>
<td>3.8</td>
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<td><strong>Reaction time versus fractional anisotropy</strong></td>
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<tr>
<td>L Supracaudate white matter-positive correlation</td>
<td>1.46</td>
<td>$-15$, 11, 21</td>
<td>3.6</td>
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<td><strong>Reaction time SD versus fractional anisotropy</strong></td>
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<td>L Supracaudate white matter-positive correlation</td>
<td>4.29</td>
<td>$-17$, 10, 19</td>
<td>3.9</td>
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<tr>
<td>R Supracaudate white matter-positive correlation</td>
<td>3.60</td>
<td>19, 15, 20</td>
<td>4.1</td>
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<td><strong>Copper versus functional connectivity</strong></td>
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<tr>
<td>Left caudate nucleus seed map</td>
<td>R Frontal operculum-negative correlation</td>
<td>5.85</td>
<td>48, 2, 16</td>
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<td></td>
<td>L Frontal operculum-negative correlation</td>
<td>3.30</td>
<td>$-38$, 14, 12</td>
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<tr>
<td>Right frontal operculum seed map</td>
<td>L Caudate nucleus-negative correlation</td>
<td>2.35</td>
<td>$-14$, 22, 8</td>
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<td></td>
<td>R Caudate nucleus-negative correlation</td>
<td>1.39</td>
<td>16, 22, 10</td>
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<td>Left frontal operculum seed map</td>
<td>L Caudate nucleus-negative correlation</td>
<td>1.30</td>
<td>$-16$, 24, 6</td>
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<td>Frontal medial seed map</td>
<td>R Frontal operculum-positive correlation</td>
<td>3.29</td>
<td>$-44$, 32, $-2$</td>
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<td>L Auditory cortex-positive correlation</td>
<td>1.72</td>
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<td>L Medial frontal cortex-positive correlation</td>
<td>1.42</td>
<td>$-12$, 22, 44</td>
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<td></td>
<td>R Visual cortex-negative correlation</td>
<td>12.81</td>
<td>14, $-54$, 2</td>
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<td>Supplementary motor area seed map</td>
<td>L Supramarginal gyrus-positive correlation</td>
<td>1.08</td>
<td>$-56$, $-34$, 24</td>
</tr>
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</table>

SD, standard deviation.

$x$, $y$, $z$ coordinates given in MNI (Montreal Neurological Institute) space. Statistics at corrected threshold $P_{FWE} < 0.05$ estimated using Monte Carlo simulations.

### Discussion

Airborne copper was significantly associated with poorer motor performance and detectable brain damage in developing children. The alterations are highly consistent with the known consequences of copper excess on the brain with basal ganglia as the main target. The associations were demonstrated with copper levels common in urban environments, thus suggesting a risk to large populations with potentially significant implications for public health. Children may be particularly vulnerable to copper as an agent capable of interfering with brain development during critical developmental stages. Consistent with our results, a recent study has reported a significant association between high copper levels in blood and poorer cognitive performance in normal school children (Zhou et al. 2015).

The $\beta$ weights reported in Table 4 may help in determining the biological relevance of behavior and imaging.
changes associated with air copper in our study. While the change related to reaction time was small, the effect on motor response consistency (measured as reaction time standard deviation) was more important. For instance, the increase in one unit (ng/m$^3$) of air copper predicts an increase of 2.9 msec in reaction time standard deviation. Therefore, if the range of copper measurements is 10 units, from the least (3.7 ng/m$^3$) to the most polluted school (13.8 ng/m$^3$), the potential variation predicted is 29 msec, which is approximately one third of the standard deviation of this motor performance measurement (SD, 91 msec). The magnitude of the effect on the caudate nucleus structure and function was of the same order. Overall, our conclusion is that the effect of copper is subtle, but biologically meaningful.

In the current study, we have identified copper as one road traffic-related pollutant. Traffic-related copper is thought to be mostly released from brake pads (Hulskotte et al. 2007) and it was the main source in our study. A significant proportion of copper, however, comes from industrial activity and a third source seems to be the result of railway traffic. This is consistent with reports indicating that the air in busy train stations contains large amounts of copper generated by overhead train wire supplying electrical power (Kim et al. 2010; Loxham et al. 2013).

Although a normal oral diet contains a considerable amount (~1 mg) of copper (Morris et al. 2006), our data suggest that relatively lower levels are neurotoxic in chronic airborne exposures. Ingested copper is mostly incorporated into ceruloplasmin (safe copper) and any excess is removed by excretion into the bile (Madsen and Gitlin 2007). There is evidence indicating that the “toxic copper” is actually the circulating free (i.e., nonceruloplasmin bound) copper (Squitti 2014c; Pal and Prasad 2015), which is the only fraction capable of penetrating the brain parenchyma (Zheng and Monnot 2012). Inhaled copper
may notably circumvent the liver regulation and safe binding to ceruloplasmin. So, nonceruloplasmin-bound copper absorbed in the respiratory tract can more easily enter the brain and achieve higher tissue concentrations.

Environmental copper has been proposed as a risk factor for neurodegenerative disease (Morris et al. 2006; Caudle 2015; Pal and Prasad 2015). The highest exposures occur through the inhalation of fumes generated from welding, as well as metal mining and smelting activities. Long-term (20 years) occupational copper exposure has been shown associated with 2.5-fold increase in risk for Parkinson’s disease (Gorell et al. 2004; Caudle 2015) and with a younger age at onset (46 years) (Racette et al. 2001). However, manifest neurological disease seems not to be a necessary outcome in occupational copper exposure, which suggests different susceptibilities among individuals and the potential interaction between a genetic predisposition and copper availability. This is obvious in the paradigmatic Wilson disease where a similar genetic alteration may correspond to a range of clinical severity and, on the other hand, the neurological disorder may be not expressed if copper intake is properly controlled (Bandmann et al. 2015). Also, there is a possibility that the toxic capacity of environmental copper excess saturates at relatively lower copper levels (i.e., high environmental copper concentrations and very high concentrations could generate similar brain damage if the assumption is correct).

The neurotoxic action of copper via oxidative stress and mitochondrial injury (Eskici and Axelse 2012; Bandmann Figure 4. Water diffusion direction changes related to copper. Correlation of copper measurements with water diffusion along the three (x, y, z) axes separately and superimposed in a single RGB color display (thresholded at $P = 0.001$ in MRicron$^3$). Color regions express sites where higher copper levels were associated with reduced diffusion for one or more directions. The most evident changes were along the right–left (red), anterior–posterior (green) directions, or both (yellow) (note that regions with mixed effects show RGB composite colors). The right hemisphere corresponds to the right side of axial and coronal views. $X$, $Y$, $Z$, denote MNI coordinates.
et al. 2015) may also depend on tissue energy consumption rates. In children, the highly energetic basal ganglia appear to be the main target. This occurs in our study and in Wilson’s disease, in which the lenticular nucleus (putamen and globus pallidus) was originally described with the most dramatic pathological changes (Compston 2009), although the caudate nucleus may show the most severe reduction in glucose consumption (Hermann 2014). By contrast, a more persistent subtle excess of copper throughout life could be responsible for protracted but widespread brain damage. In this context, an increasing amount of evidence indicates that copper may play a causal role in late-onset neurodegenerative disorders (Bush 2003; Bandmann et al. 2015).

For example, nonceruloplasmin-bound copper levels are higher than normal reference values in up to 60% of Alzheimer’s disease patients (Squitti 2014c; Squitti et al. 2014b). Also, high-nonceruloplasmin-bound copper concentrations were associated with an increased rate of mild cognitive impairment conversion to full Alzheimer’s disease (Squitti et al. 2014a). A prospective study revealed an association between a diet simultaneously high in copper and saturated fats and cognitive decline (Morris et al. 2006). In a variety of experimental animal studies, oral copper intake resulted in significant amyloid β deposition and performance decline (Alzheimer’s disease-like pathology), even at low concentrations (Pal and Prasad 2015).

Table 3. Potentially relevant airborne elements and their correlation with Cu.

<table>
<thead>
<tr>
<th>Measures from 39 schools</th>
<th>Mean ± SD</th>
<th>Range</th>
<th>Corr, with Cu r values</th>
<th>Shared variance (adjusted $r^2 \times 100$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon (C), µg/m$^3$</td>
<td>1.5 ± 0.7</td>
<td>0.6–3.9</td>
<td>0.41</td>
<td>15%</td>
</tr>
<tr>
<td>Manganese (Mn), ng/m$^3$</td>
<td>15.3 ± 13.8</td>
<td>3.7–64.8</td>
<td>0.22</td>
<td>2%</td>
</tr>
<tr>
<td>Lead (Pb), ng/m$^3$</td>
<td>8.1 ± 2.8</td>
<td>4.3–16.8</td>
<td>0.46</td>
<td>19%</td>
</tr>
<tr>
<td>Iron (Fe), µg/m$^3$</td>
<td>0.6 ± 0.6</td>
<td>0.1–3.0</td>
<td>0.15</td>
<td>0.4%</td>
</tr>
<tr>
<td>Antimony (Sb), ng/m$^3$</td>
<td>1.1 ± 0.4</td>
<td>0.4–2.4</td>
<td>0.75</td>
<td>55%</td>
</tr>
</tbody>
</table>

Cu, copper; SD, standard deviation.
In the broader context of traffic-related pollution, the effect of air pollutants may be more dramatic when the exposure involves earlier developmental periods. Indeed, Peterson et al. (2015) have provided evidence of brain structural alterations in later childhood associated with prenatal pollutant exposure affecting large areas of the left-hemisphere white matter, and a less severe effect associated with postnatal exposures at age 5 years. On the other hand, recent studies have revealed that long-term ambient air pollution exposure may ultimately affect brain tissue volume in older people (Chen et al. 2015; Wilker et al. 2015).

A general limitation when assessing children with MRI is the potential effect of head movements on image quality, particularly on functional MRI and DTI acquisitions. We have considered this issue carefully and adopted several means to rigorously control the effects of motion on functional MRI (see Methods). In the case of DTI, we decided to exclude cases with detectable image degradation, as no correction procedure is wholly efficient. A post hoc analysis on DTI using less rigorous exclusion criteria (n = 242) showed similar but weakened DTI results, indicating that the data obtained in the more selective sample (n = 177) were most probably not the result of motion effects. Also, a higher MRI signal may be obtained using a higher magnetic field (i.e., 3-Tesla magnets). Although we did have the 3-Tesla option, the present study was developed using a 1.5-Tesla magnet following the recommendations of the FP7-ERC Ethics Review Committee to limit magnetic field strength in children.

Conclusions

Our study has revealed that apparently safe school environments may indeed expose developing children to the harmful effects of air pollutants. Tolerated amounts of airborne copper were associated with poorer motor performance in children in the city of Barcelona. The effect on motor performance was directly associated with changes in the structure and function of the basal ganglia, suggesting underdevelopment of the caudate nucleus complex neural connections.

Acknowledgments

This work was supported by the European Research Council under the ERC (grant number 268479) — the BREATHE project. The Agency of University and Research Funding Management of the Catalonia Government participated in the context of Research Group SGR2014-1673. We acknowledge C. Persavento, J. Gonzalez, L. Bouso, M. Lopez, and P. Figueras their contribution to the field work. We also acknowledge all the families and schools participating in the study.

Table 4. Behavior and imaging changes associated with air copper (copper range, 3.7–13.8 ng/m3).

<table>
<thead>
<tr>
<th></th>
<th>Whole sample (n = 2827)</th>
<th>Mean ± SD</th>
<th>Motor speed (reaction time)</th>
<th>Motor response consistency (reaction time SD)</th>
<th>Gray matter concentration L caudate nucleus</th>
<th>Fractional anisotropy DTI L caudate nucleus</th>
<th>Functional connectivity L frontal cortex to L caudate</th>
</tr>
</thead>
<tbody>
<tr>
<td>β Coefficient (adjusted p)</td>
<td>2.9 (4.7) ms/(ng/m3)</td>
<td>0.6 to 3.7 (1.8 to 7.5) ms/(ng/m3)</td>
<td>0.5 to 7.7 (1.7 to 17.6) ms/(ng/m3)</td>
<td>0.1 to 0.9 (0.1 to 0.5) GMc/(ng/m3)</td>
<td>0.1 to 0.4 (0.1 to 0.2) FAi/(ng/m3)</td>
<td>0.1 to 0.2 (0.05 to 0.2) FCi/(ng/m3)</td>
<td></td>
</tr>
<tr>
<td>tP</td>
<td>2.7</td>
<td>0.006</td>
<td>2.2</td>
<td>0.026</td>
<td>0.001</td>
<td>0.001</td>
<td>0.0001</td>
</tr>
<tr>
<td>SD</td>
<td>671 ± 124 msec</td>
<td>225 ± 91 msec</td>
<td>224 ± 91 msec</td>
<td>14.2 ± 3.9 GMc</td>
<td>0.3 ± 1 FAi</td>
<td>0.3 ± 1 FCi</td>
<td></td>
</tr>
<tr>
<td>95% CI (adjusted 95% CI)</td>
<td>4.9</td>
<td>3.9</td>
<td>4.9</td>
<td>0.9 (0.3) GMc</td>
<td>0.06 (0.1) FAi</td>
<td>0.06 (0.1) FCi</td>
<td></td>
</tr>
<tr>
<td>Socioeconomic status</td>
<td>8e-7</td>
<td>3.1</td>
<td>3.1</td>
<td>0.1 (0.1) FAi</td>
<td>0.06 (0.1) FCi</td>
<td>0.06 (0.1) FCi</td>
<td></td>
</tr>
<tr>
<td>1Socioeconomic status</td>
<td>8e-7</td>
<td>3.1</td>
<td>3.1</td>
<td>0.1 (0.1) FAi</td>
<td>0.06 (0.1) FCi</td>
<td>0.06 (0.1) FCi</td>
<td></td>
</tr>
<tr>
<td>2Socioeconomic status</td>
<td>8e-7</td>
<td>3.1</td>
<td>3.1</td>
<td>0.1 (0.1) FAi</td>
<td>0.06 (0.1) FCi</td>
<td>0.06 (0.1) FCi</td>
<td></td>
</tr>
<tr>
<td>3Socioeconomic status</td>
<td>8e-7</td>
<td>3.1</td>
<td>3.1</td>
<td>0.1 (0.1) FAi</td>
<td>0.06 (0.1) FCi</td>
<td>0.06 (0.1) FCi</td>
<td></td>
</tr>
</tbody>
</table>

SD, standard deviation; p, β coefficients from the regressions with copper as the predictor factor; GMc, percentage of gray matter concentration; FAi, anisotropy index expressed in the range of 0–100 units; FCi, strength of functional connectivity expressed in arbitrary units.

n varied in each imaging modality (see Methods section).
Conflict of Interest
We have no competing interests.

References

Traffic pollution exposure is associated with altered brain connectivity in school children

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Abstract

Children are more vulnerable to the effects of environmental elements due to their active developmental processes. Exposure to urban air pollution has been associated with poorer cognitive performance, which is thought to be a result of direct interference with brain maturation. We aimed to assess the extent of such potential effects of urban pollution on child brain maturation using general indicators of vehicle exhaust measured in the school environment and a comprehensive imaging evaluation. A group of 263 children, aged 8 to 12 years, underwent MRI to quantify regional brain volumes, tissue composition, myelination, cortical thickness, neural tract architecture, membrane metabolites, functional connectivity in major neural networks and activation/deactivation dynamics during a sensory task. A combined measurement of elemental carbon and NO2 was used as a putative marker of vehicle exhaust. Air pollution exposure was associated with brain changes of a functional nature, with no evident effect on brain anatomy, structure or membrane metabolites. Specifically, a higher content of pollutants was associated with lower functional integration and segregation in key brain networks relevant to both inner mental processes (the default mode network) and stimulus-driven mental operations. Age and performance (motor response speed) both showed the opposite effect to that of pollution, thus indicating that higher exposure is associated with slower brain maturation. In conclusion, urban air pollution appears to adversely affect brain maturation in a critical age with changes specifically concerning the functional domain.

Keywords:
Brain development
Air pollution
Functional MRI
Functional connectivity

Introduction

Common to living beings, the brain development cycle is characterized by primary growth and subsequent maturation. Maturation changes implicate structure and function with anatomical shaping, progressive myelination of neural tracks and fine-tuning of functional brain networks (Menon, 2013; Pujol et al., 2006; Toga et al., 2006). The highest-order events procure the integration of brain areas into functional systems and the segregation of distinct but interconnected large-scale networks (Uddin et al., 2010; Vogel et al., 2010; Dwyer et al., 2014; Di Martino et al., 2014).

Developing children are at risk due to the potentially hazardous effects of environmental factors (Paus, 2010). Long-term exposure to traffic-related air pollution has been associated with alterations in children’s cognition (Perera et al., 2009; Suglia et al., 2008; Wang et al., 2009). We have recently identified a significant association between general markers of road traffic pollution and slower cognitive growth in a large group of children (Sunyer et al., 2015).

Epidemiological studies, therefore, indicate that high levels of urban air pollution may be dangerous to children, as they presumably interfere with brain maturation processes. This hypothesis is largely supported by a set of studies in both animals and humans showing significant associations of pollutant exposure with...
inflammatory and degenerative brain pathology (Block and Calderón-Garcidueñas, 2009; Calderón-Garcidueñas, 2012). However, such an interference effect on brain development has not been thoroughly investigated. We aimed to assess the extent of potential repercussions of traffic pollution exposure on child brain maturation using a variety of imaging measurements ranging from basic anatomy to high-order functional integration. A group of 263 children, aged 8 to 12 years, recruited from a large study assessing the impact of long-term exposure to urban pollution in Barcelona city school environments (Sunyer et al., 2015) completed the protocol.

Our hypothesis was that the potential brain effects of air pollution will be more evident on the more detectable anatomical and functional maturation processes. Whereas developmental changes in gray matter volume are less evident in this age period, active myelination implicates increases of relative white matter volumes, elevated choline compounds and water diffusion changes within white matter tracts (Bluml et al., 2013; Toga et al., 2006; Yoshida et al., 2013). At the functional domain, preadolescence is critical to the optimal assembling of large-scale functional networks (Menon, 2013). Accordingly, the imaging protocol included a high resolution 3D anatomical acquisition to measure regional volumes, brain tissue composition, myelination levels and cortical thickness. Diffusion tensor imaging (DTI) measurements of fractional anisotropy served to explore white matter tract architecture. In vivo spectroscopy was used to grossly estimate precursors of membrane components in white matter. Finally, functional MRI was used to test the integrity of relevant networks using both resting-state functional connectivity and a task activation/deactivation paradigm.

Selected cognitive assessment was also conducted to determine to what extent potential repercussions were also detectable on children’s performance in the current study sample.

Methods

Participant selection

This study was developed in the context of the BREATHE project (The European Commission: FP7-ERC-2010-AdG, ID 268479). The general project design is fully described in Sunyer et al. (2015). A total of 1564 families, from 39 schools in the city of Barcelona, were invited to participate in the MRI study via post, email or telephone, and 810 of them gave an initial positive response. The study sample was consecutively recruited from this group with the aim of including children to test the integrity of relevant networks using both resting-state functional connectivity and a task activation/deactivation paradigm.

Selected cognitive assessment was also conducted to determine to what extent potential repercussions were also detectable on children’s performance in the current study sample.

Table 1 reports the characteristics of these participants. Additional cases were excluded on the basis of image quality criteria in each specific MRI analysis (see further).

All parents or tutors signed the informed consent form approved by the Reseach Ethical Committee (No. 2010/41221/1) of the IMIM-Parc de Salut Mar, Barcelona, Spain and the FP7-ERC-2010-AdG Ethics Review Committee (268479-22020211).

Pollutant exposure

Each school was measured twice during one-week periods separated by 6 months, in the warm (year 2012) and cold (year 2012/2013) seasons. Indoor air in a single classroom and outdoor air in the playground were measured simultaneously. Pollutants were measured during class-time using methods previously described (Amato et al., 2014; Rivas et al., 2014; Sunyer et al., 2015).

Elemental carbon was measured during 8 h (09:00 to 17:00 h) in particulate matter with an aerodynamic diameter < 2.5 μm (PM2.5) collected on filters with High-Volume samplers (MCE SA, Spain) using a Thermo Optical Transmission method (Sunset Laboratories Inc.). We carefully followed the EUSAAR-2 protocol, TOT Sunset Laboratories measurements, with a detection limit of 0.1 μg/m³ and an uncertainty of ± 5%. The air cleaning effect of High-Volume samplers may underestimate absolute measurements of elemental carbon in poorly ventilated indoors. In our study, however, elemental carbon penetration was almost 1% (indoor/outdoor ratio 94.1% [95% CI 85.7%–102.4%]), which suggests a permanent ventilation of the measured classrooms. Elemental carbon was additionally measured in each classroom using the MicroAeth AE51 (AethLabs). The correlation between elemental carbon measured through High-Volume samplers and with the aethalometer was 0.95, supporting that High-Volume sampler measurements may be adequate estimations of classroom elemental carbon.

Nitrogen dioxide (NO2) was measured with passive dosimeters (Gradko). The dosimeter was exposed during a period of 96 h (4 days) from Monday to Thursday in each school in both the warm and cold campaigns. Weekly data from both seasons were averaged to obtain a single measurement. Prior to the campaigns, we tested Gradko NO2 passive dosimeters in our urban background monitoring station Palau Reial (with relatively low NO2 concentrations) by measuring during 4 days and comparing the results with simultaneous chemiluminescence NO2 online data. The results showed that sampling periods of 4 days were enough for ensuring a good precision. Also, during the whole sampling campaign, NO2 was measured each week (from Monday to Thursday) with both the Gradko passive dosimeter and conventional chemiluminescence analyzers in this reference station. We obtained a correlation of Gradko =

<table>
<thead>
<tr>
<th>Table 1 Characteristics of the study sample (n = 263).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
</tr>
<tr>
<td>Age, years, mean ± SD (range)</td>
</tr>
<tr>
<td>Overall school achievement—5-point scale</td>
</tr>
<tr>
<td>Difficulties Score (SDQ), range 0–40</td>
</tr>
<tr>
<td>Obesity: normal</td>
</tr>
<tr>
<td>Obesity, BMI &lt; 94</td>
</tr>
<tr>
<td>Obesity, BMI ≥ 94</td>
</tr>
<tr>
<td>Mother education (5-point scale. 5 = university)</td>
</tr>
<tr>
<td>Father education (5-point scale. 5 = university)</td>
</tr>
<tr>
<td>Vulnerability indexa—home</td>
</tr>
<tr>
<td>Vulnerability indexa—school</td>
</tr>
<tr>
<td>Public/non-public school</td>
</tr>
<tr>
<td>Task performance, N-back</td>
</tr>
<tr>
<td>Reaction time (ms)</td>
</tr>
<tr>
<td>Reaction time standard deviation (ms)</td>
</tr>
<tr>
<td>Commission errors (number)</td>
</tr>
<tr>
<td>Omission errors (number)</td>
</tr>
<tr>
<td>Alerting (ms)</td>
</tr>
<tr>
<td>Orienting (ms)</td>
</tr>
<tr>
<td>Interference (ms)</td>
</tr>
<tr>
<td>Air pollution measurementsb</td>
</tr>
<tr>
<td>Outdoor elemental carbon (EC) year average (μg/m³)</td>
</tr>
<tr>
<td>Indoor elemental carbon (EC) year average (μg/m³)</td>
</tr>
<tr>
<td>Overall air pollution indicator (EC + NO2 weighted average)</td>
</tr>
</tbody>
</table>

BML, body mass index; SDQ, Strengths and Difficulties Questionnaire.

a Neighborhood socioeconomic status vulnerability index based on level of education, unemployment, and occupation at the census tract (Atlas de vulnerabilidad urbana de España, 2012).
b After excluding 3 children with outlier measurements.
0.85 \times \text{chemiluminescence} + 3.6 \text{ (R2 = 0.7)} \text{ for a mean of 37 \mu g NOx/m3},
and an uncertainty of \pm 17%.

We operationally selected elemental carbon and NO2 to compute a
general traffic pollution indicator given their relation to vehicle exhaust
emissions in the city of Barcelona (Amato et al., 2014). Our interest here
was not to identify specific neurotoxic agents directly responsible for
neuronal damage, but it was merely to use a measurement globally
representing exposure to this sort of air contamination. NO2 is highly
correlated with elemental carbon in Barcelona and shows the two typi-
cal rush hour peaks (Reche et al., 2011). Barcelona has a diesel domi-
nated vehicle fleet (with very high NOx emissions and a high NO2/NOx
rate) with a primary NOx driven daily pattern (Reche et al., 2011). In
Barcelona, 80% of the NOx emissions arise from road traffic (Catalonian
eu/wp-content/uploads/2012/10/Kick-off_Generalitat.pdf), and even if
a fraction of NOx is secondary, this is arising in a large proportion from
NO emitted from road traffic as well (Grice et al., 2009). We did not
use in this study levels of ultrafine particles, which may be more directly
related to the potential neural damage, because in high insolation cities
such as Barcelona, levels of ultrafine particles are highly influenced by
photochemical nucleation usually occurring at midday. This effect pre-
vents a high correlation of ultrafine particle levels with other traffic
tracers (Brines et al., 2015; Reche et al., 2011).

To obtain more representative measurements, elemental carbon and
NO2 were adjusted for temporal variability using whole-year data from
a background monitoring station in Barcelona (Sunyer et al., 2015). A sin-
gle traffic-related pollutant indicator was computed using the weighted
average of elemental carbon and NO2 [(EC/group median) + (NO2/
group median) / 2] of pooled indoor and outdoor measurements from
both cold and warm seasons. All children had been in the school for
more than 18 months (and 98% more than two years) at imaging assess-
ment, which was carried out after the pollution measurement campaigns.

Cognitive performance measurements

General cognitive assessment included working memory, motor re-
sponse speed and attention. Working memory was assessed using a
computerized version of the N-Back task (Anderson, 2002). Children’s
correct detections on 2-back and 3-back loads for number items were
assessed using the computerized version of the N-Back task (Anderson, 2002). Children’s
response speed and attention. Working memory was assessed using a
computerized version of the "Attentional Network Test", child version (Child ANT) (Rueda et al., 2004). We used the overall
"reaction time" to measure speed of motor responses and reaction time
standard deviation to measure trial-to-trial variability. A higher reaction
time standard deviation indicates lower executive and general attention-
alf resources (Langner and Eickhoff, 2013). The ANT task also uses a
set of cued and congruent/incongruent conditions to measure specific
attention features such as ‘alerting’, ‘orienting’ and ‘interference’. De-
tails for administering both ANT and N-Back tasks are fully described in
a previous report (Forns et al., 2014).

Additional contextual assessments

Socio-demographic factors were measured using a neighborhood
socioeconomic status vulnerability index (based on level of education,
Atlas Vulnerabilidad Urbana/) according to both school and home ad-
dress. Distance from home to school was estimated based at the
gecoded postal address of each participant and school. Parental educa-
tion was registered for both parents using a 5-point scale (1 illiterate/2
less than/3 primary/4 secondary/5 university). Standard measurements
of height and weight were performed to define overweight and obesity
(de Onis et al., 2009). Parents completed the Strengths and Difficulties
Questionnaire (SDQ) on child behavioral problems (Goodman, 2001).
A “difficulties” score ranging from 0 to 40 was generated. Overall school
achievement was rated by teachers using a 5-point scale (from the
worse = 1 to the best = 5).

MRI and MR spectroscopy acquisition

A 1.5 Tesla Signa Excite system (General Electric, Milwaukee, WI,
USA) equipped with an eight-channel phased–array head coil and
single-shot echoplanar imaging (EPI) software was used. The imaging
protocol involved an anatomical T1-weighted 3D sequence, diffusion
tensor imaging (DTI), MR proton spectroscopy and functional MRI.

High-resolution 3D anatomical images

High-resolution 3D anatomical images were obtained using an
axial T1-weighted three-dimensional fast spoiled gradient inversion
recovery-prepared sequence. A total of 134 contiguous slices were ac-
quired with inversion time 400 ms; repetition time 11.9 ms; echo
time 4.2 ms; flip angle 15°; field of view 30 cm; 256 × 256 pixel matrix;
slice thickness 1.2 mm.

Diffusion tensor imaging (DTI)

Diffusion tensor imaging (DTI) was obtained using spin-echo
single-shot echo-planar sequences of 25 directions with a B-factor of
1000 s/mm². Twenty-six slices were acquired with repetition time
8300 ms; echo time 94 ms; thickness 5 mm, no gap; pulse angle 90°;
field of view 26 cm; 128 × 128 acquisition matrix reconstructed into a
256 × 256 matrix.

Magnetic resonance spectroscopy

Proton (1 H) spectroscopy was performed using the fully automated
Proton Brain Exam-Single Voxel (PROBE-SV) software package (GE
Medical Systems, Milwaukee, WI) and a Stimulated Echo Acquisition
Mode (STEAM) pulse sequence with TR/TE = 2000/30 ms and 128 sig-
nal averages. Total acquisition time was 5 m 4 s. The voxel showed a di-
mension of 23 × 14 × 14 mm and was always placed in the left frontal
white matter with the aid of high-resolution 3-D images. Orthogonal
projections in the three planes assisted the placement of the voxel.
The major axis (23 mm) was aligned along the anterior–posterior direc-
tion in the frontal white matter. Care was taken to minimally include
gray matter and to place the voxel just above the caudate nucleus be-
tween the cingulate cortex and the frontal cortex at the level of the
precentral sulcus.

The functional MRI sequences

The functional MRI sequences consisted of gradient recalled acqui-
sition in the steady state with repetition time 2000 ms; echo time
50 ms; pulse angle 90°; field of view 24 cm; 64 × 64-pixel matrix;
slice thickness 4 mm (inter-slice gap, 1.5 mm). Twenty-two interleaved
slices were prescribed parallel to the anterior–posterior commissure
line covering the brain. Two fMRI sequences were acquired for each par-
ticipant including a 6-min continuous resting-state scan generating 180
whole-brain EPI volumes, and a 4-min sensory task generating 120
whole-brain EPI volumes. The first four (additional) images in each
fMRI run were discarded to allow magnetization to reach equilibrium.

During the resting-state functional fMRI, children were instructed to
relax, stay awake and lie still without moving, while keeping their
eyes closed throughout. The task-activation paradigm involved an
ABABABAB block design alternating four 30-s periods of rest (visual
fixation to a cross) with four 30-s periods of visual–auditory stimulation
delivered using MRI compatible goggles and headphones (VisuaStim
Digital, Resonance Technology, USA). Subjects were passively
confronted with a set of facial images expressing happiness and with
music showing a rapid tempo (Beethoven Symphony No. 6 “Pastorale”).
Image/spectra processing

3D anatomical images

All the anatomical images were visually inspected before analysis by a trained operator to detect any motion effect. A total of 10 children were discarded as a result of poor quality images and thus the final sample for the 3D anatomical analysis included 253 children. Anatomical 3D data were processed in two separate analyses assessing different anatomical characteristics:

Gray and white matter tissue concentration and volume at a voxel level was measured using Statistical Parametric Mapping (SPM8) (http://www.fil.ion.ucl.ac.uk/spm, Wellcome Department of Cognitive Neurology, London, UK, 2008). SPM voxel-based morphometry (VBM) algorithms with DARTEL registration were used with the following processing steps: segmentation of anatomical images into gray and white matter tissue probability maps in their native space; estimation of the deformations that best align the images together by iteratively registering the segmented images with their average; finally, generating spatially normalized and smoothed segmentations (5 × 5 × 5 FWHM) using the deformations estimated in the previous step. The analyses were performed with scaling by Jacobian determinants (estimates of volume change during the normalization) to consider tissue volume and without Jacobian scaling to assess the relative concentration of gray matter and white matter. Normalized images were finally transformed to the standard SPM template, re-sliced to 1.5 mm resolution in Montreal Neurological Institute (MNI) space.

Cortical thickness measurements across the whole cortex were obtained using FreeSurfer tools (http://surfer.nmr.mgh.harvard.edu/). Processing steps included removal of non-brain tissue, segmentation of the subcortical white matter and deep gray matter volumetric structures, tessellation of the gray and white matter boundary, registration to a spherical atlas which is based on individual cortical folding patterns to match cortical geometry across subjects and creation of a variety of surface based data. Cortical thickness is calculated as the closest distance from the gray/white boundary to the gray/CSF boundary at each vertex on the tessellated surface (Fischl et al., 1999).

Diffusion tensor imaging (DTI)

DTI was processed using Functional MRI of the Brain (FMRIB) Software Library 5.0 (FSL), developed by the Analysis Group at the Oxford Centre for FMRIB (Smith et al., 2004). Diffusion-weighted images were corrected for motion and eddy current distortions (“Eddy Current Correction” option in the FMRIB Diffusion Toolbox [FDT] version 2.0 in FSL), and a whole-brain mask was applied using the FSL Brain Extracting Tool. A further rigorous image quality control was carried out to identify potential residual effects of head motion, which involved the visual inspection of each DTI slice for all 25 DTI volumes in all participants. Volumes with slices with signal loss (greater than ~10%) or residual artifacts were identified by an expert researcher. DTI full examinations showing one or more degraded images in more than 5 volumes were discarded. A total of 76 children were removed from the DTI analysis on the basis of this criterion (in addition to 10 cases showing gross image degradation). The final DTI sample involved 177 children showing a mean ± SD of 23.5 (94%) ± 1.9 optimal-quality volumes. Subsequently, we estimated fractional anisotropy (FA) maps using FDT in FSL after local fitting of the diffusion tensor model at each voxel (“dtifit”). Next, diffusion data were processed using Tract-Based Spatial Statistics (Smith et al., 2006). Each FA data set was re-sliced to a 1 mm × 1 mm × 1 mm anatomical resolution and normalized to standard MNI space via the FMRIB58_FA template using the FMRIB’s Non-linear Registration Tool.

Magnetic resonance spectroscopy

Metabolite relative measurements were performed on the Advantage Windows, v. 4.2, workstation using the PROBE-SV software package, which includes automatic processing of the raw data that permits immediate display and evaluation of spectra. Measurements of choline-containing compounds (choline-to-total creatine ratio) were
used to test the association of air pollution with membrane precursors in white matter (Blüml et al., 2013). N-acetylaspartate (NAA) spectra width and the ratio creatine-to-noise (root-mean-square ‘RMS’ noise) were used to assess spectra quality. The quality criteria to be retained for statistical analyses were NAA linewidth (full width at half maximum of peak) of 0.09 ppm or less and creatine RMS noise of 8 or greater. A total of 34 participants were excluded as a result of poor spectrum quality. The final sample for the spectroscopy analysis included 229 children.

**Functional MRI**

Resting-state and task functional MRI preprocessing was carried out using SPM8 and involved motion correction, spatial normalization and smoothing using a Gaussian filter (full-width half-maximum, 8 mm). Data were normalized to the standard SPM-EPI template and re-sliced to 2 mm isotropic resolution in MNI space.

The following procedures were adopted to control for potential head motion effects: (i) Conventional SPM time-series alignment to the Data were normalized to the standard SPM-EPI template and re-sliced filter (full-width half-maximum, 8 mm). (ii) Exclusion of 24 children in the resting-state analysis and 39 children in the task-activation analysis with large head motion. That is, outliers (and extremes) with regard to mean inter-frame motion were excluded using conventional boxplot criteria (cases beyond the quartile Q3 by one-and-a-half Q3-Q1 inter-quartile range [SPSS 15.0; SPSS Inc., Chicago IL]). The final analyzed sample therefore included 239 children with valid resting-state and 39 children in the task-activation analysis.

The maps obtained using the medial frontal cortex and the posterior cingulate cortex regions typically include all the elements of the Default Mode Network (DMN), which is highly active in inner mental processes and is negatively correlated (anticorrelated) with networks participating in attention-demanding tasks (Kelly et al., 2008). On the other hand, maps from the frontal cortex and supplementary motor area regions both capture networks commonly participating in attention-demanding tasks. As classically described, the dynamic relationship

### Table 2

**Functional MRI results.**

<table>
<thead>
<tr>
<th>Correlation with air pollution-</th>
<th>Non-adjusted</th>
<th>Adj. by age and sex</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ml</td>
<td>x  y  z</td>
</tr>
<tr>
<td>Medial frontal seed map</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L Lateral frontal cortex—positive correlation</td>
<td>5.1</td>
<td>-48 46 16</td>
</tr>
<tr>
<td>R Parietal cortex—negative correlation</td>
<td>2.6</td>
<td>-50 -58 44</td>
</tr>
<tr>
<td>Dorsal frontal seed map</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L Parietal cortex—positive correlation</td>
<td>3.2</td>
<td>-32 -64 32</td>
</tr>
<tr>
<td>R Lateral frontal cortex—negative correlation</td>
<td>3.4</td>
<td>50 24 2</td>
</tr>
<tr>
<td>R Insula—negative correlation</td>
<td>1.4</td>
<td>38 2 -2</td>
</tr>
<tr>
<td>Posterior cingulate cortex seed map</td>
<td>1.5</td>
<td>56 36 2</td>
</tr>
<tr>
<td>Supplementary motor area seed map</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L Prefrontal cortex—positive correlation</td>
<td>6.1</td>
<td>-28 38 28</td>
</tr>
<tr>
<td>R Prefrontal cortex—positive correlation</td>
<td>2.2</td>
<td>22 48 34</td>
</tr>
<tr>
<td>L Parietal cortex—positive correlation</td>
<td>3.5</td>
<td>-58 -46 36</td>
</tr>
<tr>
<td>R Parietal cortex—positive correlation</td>
<td>5.2</td>
<td>58 -46 40</td>
</tr>
<tr>
<td>Anterior cingulate cortex—negative correlation</td>
<td>6.2</td>
<td>-14 34 -12</td>
</tr>
<tr>
<td>Sensory task</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R Somatosensory cortex—positive correlation</td>
<td>6.3</td>
<td>34 -44 64</td>
</tr>
<tr>
<td>L Premotor cortex—positive correlation</td>
<td>1.4</td>
<td>-10 0 60</td>
</tr>
</tbody>
</table>

**Correlation with age**

<table>
<thead>
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<th>ml</th>
<th>x  y  z</th>
<th>t</th>
</tr>
</thead>
<tbody>
<tr>
<td>L Parietal cortex—positive correlation</td>
<td>10.7</td>
<td>-46 -70 52</td>
<td>4.9</td>
</tr>
<tr>
<td>Medial frontal cortex—positive correlation</td>
<td>1.5</td>
<td>-6 52 48</td>
<td>3.3</td>
</tr>
<tr>
<td>Cerebellum—positive correlation</td>
<td>5.8</td>
<td>-22 -48 32</td>
<td>4.6</td>
</tr>
<tr>
<td>L Lateral frontal/insula—negative correlation</td>
<td>9.7</td>
<td>-36 22 0</td>
<td>4.8</td>
</tr>
<tr>
<td>R Lateral frontal/insula—negative correlation</td>
<td>10.4</td>
<td>40 16 2</td>
<td>3.5</td>
</tr>
</tbody>
</table>

**Correlation with performance (motor speed)**

<table>
<thead>
<tr>
<th>Medial frontal seed map</th>
<th>ml</th>
<th>x  y  z</th>
<th>t</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medial frontal cortex—positive correlation</td>
<td>7.6</td>
<td>-10 46 26</td>
<td>3.6</td>
</tr>
<tr>
<td>L Frontal lateral cortex—negative correlation</td>
<td>1.6</td>
<td>-32 44 -4</td>
<td>3.9</td>
</tr>
</tbody>
</table>

x  y  z, coordinates given in Montreal Neurological Institute (MNI) space. Statistics at corrected threshold $P_{FWE} < 0.05$ estimated using Monte Carlo simulations. Cluster size in ml.
between the DMN and the anticorrelated task-related networks may reflect the largest-scale functional organization in the brain (Fox et al., 2005; Kelly et al., 2008). Interestingly, this network interaction matures significantly in the age period targeted in the present study (Sherman et al., 2014).

Functional connectivity maps were generated using procedures detailed in previous reports (Harrison et al., 2013; Pujol et al., 2014a). For each location, the seed region was defined as a 3.5 mm radial sphere (sampling ~25 voxels in 2 mm isotropic space). This was performed using MarsBaR region of interest (ROI) toolbox in MNI stereotaxic space (Brett et al., 2003). Signals of interest were then extracted for each seed region respectively by calculating the mean ROI value at each time point across the time-series. To generate the seed maps, the signal time course of a selected seed region was used as a regressor to be correlated with the signal time course of every voxel in the brain in order to generate first-level (single-subject) voxel-wise statistical

![Fig. 2. Correlations of age with functional connectivity in the medial frontal seed map. Age was associated with higher functional connectivity between the medial frontal (seed) region and the left angular gyrus, medial frontal cortex and cerebellum (left panels), and with lower functional connectivity between the seed and the lateral frontal cortex/insula region bilaterally (right panels). The right hemisphere corresponds to the right side of axial and coronal views.](image1)

![Fig. 3. 3D rendering display of urban pollution and age effects in the medial frontal seed map. Age and air pollution were associated with opposite effects on functional connectivity in notably similar areas, thus indicating that higher exposure may interfere with the normal development of functional connections.](image2)
parametric maps (contrast images). The maps were estimated for each seed separately. A high-pass filter set at 128 s was used to remove low frequency drifts below ~0.008 Hz. In addition, we derived estimates of white matter, CSF and global brain signal fluctuations (using standard masks in MNI space from SPM) to include in the regression analyses as nuisance variables.

**Task-activation fMRI.** Functional interaction between brain systems may also be inferred using fMRI task paradigms by assessing the segregation between activated and deactivated areas during stimulation (Harrison et al., 2008; Pujol et al., 2012). Deactivated areas during our sensory task include part of the DMN, which are generally anticorrelated with networks participating in attention-demanding tasks (Chai et al., 2014; Kelly et al., 2008), although the deactivation pattern is not limited to the DMN. In the task analysis, single-subject (first-level) SPM contrast images were estimated for activations (stimulation condition > rest) and deactivations (rest > stimulation condition). For these analyses, the fMRI signal response at each voxel was modeled using the SPM canonical hemodynamic response function.

Before the statistical analysis, all the processed images (anatomical 3D, DTI, and functional MRI) were visually inspected to detect processing-induced artifacts and verify the accuracy of anatomical segmentation and FreeSurfer surface reconstructions.

**Statistical analysis**

Data was treated as quantitative variables and the analyses involved cross-correlations using SPM. After individual preprocessing of each imaging exam, separate second-level analyses were carried out to map voxel-wise the correlation across-subjects between individual brain images and individual age and pollutant measurements (obtained in the school of each participant). The correlations with pollutant measurements were performed both with and without adjusting by age and sex. The set of individual brain images included whole-brain estimates of regional white matter and gray matter volume and concentration, cortical thickness, DTI fractional anisotropy, resting-state functional connectivity and task-activation/deactivation. The association with choline-to-creatine metabolite ratio was obtained using simple bivariate Pearson correlation. Total creatine was examined first for an association with

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**Fig. 4.** Sensory task results. One-sample group activation (top panel) and deactivation (middle panel) maps showing brain response to passive viewing and listening. Air pollution was associated with lower deactivation in the supplementary motor area and somatosensory cortex included in the deactivation map (bottom panel). The right hemisphere corresponds to the right side of axial and coronal views.
traffic pollution. We found no association. As mentioned, the number of participants finally included in each analysis after exclusions on the basis of image quality criteria was: anatomic T1-weighted images, 253; DTI, 177; MR spectroscopy, 229; resting-state fMRI, 239; and sensory task fMRI 224. Finally, three additional children with outlier pollutant measurements were excluded from the correlation between brain measurements and air pollution.

Results were considered significant with clusters of 1.032 ml (e.g., 129 voxels with a resolution of $2 \times 2 \times 2$ mm) at a height threshold of $p < 0.005$, which satisfied the family-wise error (FWE) rate correction of $P_{\text{FWE}} < 0.05$ according to Monte Carlo simulations (Pujol et al., 2014c). Resting-state fMRI data were additionally adjusted for multiple testing (four functional connectivity maps) using Bonferroni (significant cluster size $\geq 1.4$ ml). Maps in figures are displayed at $t > 2.3$.

Results

Traffic-related pollution at the children's schools was assessed using the weighted average of two reliable vehicle exhaust indicators, namely particulate elemental carbon and NO$_2$ (Methods). Table 1 reports ambient air concentrations of the measured pollutants. According to these data, air pollution levels at school in the city of Barcelona may be considered moderate-to-high when compared with other areas (Cyrys et al., 2012; Eeftens et al., 2012).

This summary pollution index was then whole-brain correlated with several MRI maps characterizing brain maturation. No significant association was identified between air pollution and any anatomical, structural or metabolic brain measurement. By contrast, the functional imaging analysis showed consistent results.

Our functional approach involved the generation of connectivity maps representative of key neural networks using coordinates taken from previous works (see Methods). The map obtained using the medial frontal cortex (seed) region of interest produced the most illustrative results. We found that traffic-related air pollution was significantly associated with weaker functional connectivity between regions belonging to the DMN (i.e., between the medial frontal cortex and the angular gyrus bilaterally), indicating lower intra-network integration (Fair et al., 2009). In addition, pollution also was associated with stronger functional connectivity between the (medial frontal cortex) seed region and the frontal operculum at the lateral boundary of the DMN, indicating lower network segregation (Fair et al., 2009) (Fig. 1 and Table 2).

The analysis of the correlation between children's age and functional connectivity in the DMN was useful in establishing the detrimental nature of the results (Fig. 2). Indeed, the age effect on functional connectivity in this network was the opposite of the pollutant effect. Age was significantly associated with stronger functional connectivity within the elements of the DMN (integration) and significantly associated with weaker functional connectivity with the bordering network (segregation), as shown in a previous longitudinal study (Sherman et al., 2014). Fig. 3 illustrates the opposite effects of age and pollutants using a 3D rendering display.

The analyses based on additional functional connectivity maps mirrored such findings with results in the same direction. The maps included the DMN identified from a seed region located in the posterior cingulate cortex (Table 2) and anticorrelated networks generated from dorsal frontal cortex (Supplementary Fig. 2) and supplementary motor area seeds (Supplementary Fig. 2).

The potential effects of air pollution on brain function were further tested by mapping the correlation between pollutant measurements and fMRI task activations and deactivations. Air pollutants were significantly associated with lower deactivations (rest $\rightarrow$ task map) during passive viewing and listening in the supplementary motor area and somatosensory cortex (Fig. 4). No significant findings were observed in the task $\rightarrow$ rest map.

To test whether measured pollution was associated with cognitive performance, we used children's performance in working memory, motor response speed and attention (Methods). The only significant result involved motor speed. Higher pollution predicted slower reaction time in 248 participants with complete MRI and behavioral testing (standardized $\beta = 0.154$; $p = 0.015$). A further imaging analysis was performed to correlate children's reaction time with functional connectivity in the DMN map (Fig. 5). Interestingly, a faster reaction time was associated with stronger connectivity within the DMN (network integration) and weaker connectivity in the frontal operculum (network segregation).

The effect of potential confounders was tested for each significant finding including age, sex, academic achievement, difficulties scores, obesity, parental education, home and school vulnerability index, distance from home to school and public/non-public school category as co-variates. Each potential confounder was both individually entered into the model and combined with other confounders. No single confounder or combination showed a relevant effect. That is, decreases in $\beta$ estimates after the inclusion of confounders in a regression model were
very small (mean ± SD, 1.2% ± 1.0%) with no variables affecting the primary results with β reductions greater than 7%.

Discussion

Vehicle exhaust-related air pollution exposure was associated with brain changes of a functional nature, with no evident effect on brain anatomy, structure or membrane metabolism. Children from schools with higher traffic-related pollution showed lower functional integration and segregation in key brain networks. Age and performance (i.e., motor response speed) both showed the opposite effect to that of pollution on brain function, thus indicating that higher exposure is associated with slower brain maturation.

The functional findings were highly consistent, as similar effects were observed in different functional networks and the age-sensitive areas notably coincided with the areas showing significant correlation with air pollution. Similarly, the regions identified with the mapping of correlations with motor speed also showed a notable correspondence with the anatomy of findings from both pollutant and age analyses. Nonetheless, despite the evident effect on functional connectivity, the overall brain repercussion may, to some extent, be considered subtle, as changes did not involve any measurement of brain structure. In such a context, one may speculate on the reversibility of the brain damage and the potential effectiveness of actions addressed to reduce pollution. Epidemiological data also support the notion of a subtle repercussion, as large samples are required to demonstrate robust associations (Sunyer et al., 2015).

The effect of air pollution may, however, be more dramatic when the exposure involves early developmental periods. Indeed, Peterson et al. (2015) have provided evidence of brain structural alterations in later childhood associated with prenatal pollutant exposure affecting large areas of the left-hemisphere white matter, and a less severe effect associated with postnatal exposures at age 5 years. Also, recent studies have revealed that long-term ambient air pollution exposure may ultimately affect brain tissue volume in older people (Chen et al., 2015; Wilker et al., 2015).

We have used a general marker of vehicle exhaust traffic-related air pollution based on elemental carbon and NO2. Although this indicator reflects the amount of pollution from the traffic source, carbon and NO2 are not necessarily the agents causing the toxic effect on the brain. Traffic pollution contains a variety of elements with greater potential neurotoxicity, such as manganese, aluminum, lead and copper (Amato et al., 2014). We have recently identified the effects of airborne copper pollution, which is road traffic-related, although a significant proportion also comes from industry, and a third source is the result of railway traffic (submitted). Brain alterations associated with copper were neural system specific and affected the basal ganglia with damage to both structure and function. In contrast, our current study based on a general indicator does not inform on which specific neurotoxicant or a combination thereof may be responsible for the identified effect restricted to functional measurements.

A concern in traffic pollution studies is potential residual confounding by socio-demographic characteristics (e.g., when a relationship exists between proximity to traffic and economically disadvantaged areas). In the city of Barcelona, however, there was a small and inverse relation between air pollution and socioeconomic vulnerability, with higher pollution levels in schools with lower vulnerability (Sunyer et al., 2015). Also, we found no significant associations between school pollution levels and parental education, employment or educational quality. Besides, the associations between air pollution and fMRI remained after adjusting for all the potential confounders (see Results), which is opposed to a potential role of these variables.

A general limitation when assessing children with MRI is the potential effect of head movements on image quality, particularly on functional MRI and DTI acquisitions. We considered this issue carefully and adopted several means to rigorously control the effects of motion (Methods). However, it is relevant to emphasize that a rigorous control of potential head motion may introduce spurious changes (e.g., regressing out global brain signal may introduce negative correlations) or even remove changes related to genuine neural activity (Pujol et al., 2014b). Also, a higher MRI signal may be obtained using a higher magnetic field (i.e., 3-Tesla magnets). Although we did have the 3-Tesla option, the present study was developed using a 1.5-Tesla magnet following the recommendations of the FP7–ERC Ethics Review Committee to limit magnetic field strength in children.

Conclusion

Although children’s brains may be vulnerable at each developmental stage, preadolescence is notably transcend in establishing solid bases for large-scale functional network organization. Urban traffic pollution appears to be capable of affecting the normal development of the proto-adult brain and significantly interfering with functional network maturation.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgments

This work was supported by the European Research Council under the ERC [grant number 268479]—the BREATHE project. The Agency of University and Research Funding Management of the Catalonia Government participated in the context of Research Group SGR2014-1673. We acknowledge Cecilia Persavento, Judit González, Laura Bouso, Mónica López and Pere Figueras for their contribution to the field work. We also acknowledge all the families and schools participating in the study.

Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.neuroimage.2016.01.036.

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Video Gaming in School Children: How Much Is Enough?

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Objective: Despite extensive debate, the proposed benefits and risks of video gaming in young people remain to be empirically clarified, particularly as regards an optimal level of use.

Methods: In 2,442 children aged 7 to 11 years, we investigated relationships between weekly video game use, selected cognitive abilities, and conduct-related problems. A large subgroup of these children (n = 260) was further examined with magnetic resonance imaging approximately 1 year later to assess the impact of video gaming on brain structure and function.

Results: Playing video games for 1 hour per week was associated with faster and more consistent psychomotor responses to visual stimulation. Remarkably, no further change in motor speed was identified in children playing >2 hours per week. By comparison, the weekly time spent gaming was steadily associated with conduct problems, peer conflicts, and reduced prosocial abilities. These negative implications were clearly visible only in children at the extreme of our game-playing distribution, with 9 hours or more of video gaming per week. At a neural level, changes associated with gaming were most evident in basal ganglia white matter and functional connectivity.

Interpretation: Significantly better visuomotor skills can be seen in school children playing video games, even with relatively small amounts of use. Frequent weekly use, by contrast, was associated with conduct problems. Further studies are needed to determine whether moderate video gaming causes improved visuomotor skills and whether excessive video gaming causes conduct problems, or whether children who already have these characteristics simply play more video games.

We performed the current study in 2,442 school-age children to investigate relationships between average weekly video game use and both cognitive performance and conduct-related problems. Considering video gaming as a training exercise based on repetitive use and on the basis of previous research,3 we predicted that video gaming in school children would have a principal beneficial effect on speed of mental processing (eg, reaction time and chronometric measurements of attention) and a marginal influence on more innate cognitive capabilities (eg, working memory).7,8

A subgroup of this population (n = 260) was further examined with magnetic resonance imaging (MRI)
approximately 1 year later to assess the impact of video gaming on brain structure and function. In the brain, video gaming could more notably reinforce neural connections, with a major effect presumably on frontal–basal ganglia circuits, which are central for the acquisition of new skills through practice.9–11

Subjects and Methods

Participants

This study was developed in the context of a large-scale project designed to assess the effects of environmental factors on brain development in children (BREATHE, European Commission FP7-ERC-2010-AdG, ID 268479). Study design and participant selection have been described in full detail elsewhere.12,13 In short, the BREATHE project recruited 2,897 children from 39 schools to form a representative sample of children in Barcelona aged between 7 and 11 years.12 A representative subsample of 278 children was recruited from the larger cohort to participate in neuroimaging. For the present video game study, we included children with complete cognitive and behavioral evaluation, and excluded statistically defined extreme video gamers (n = 23 children playing ≥18 hours per week). The final sample comprised 2,442 children (mean ± standard deviation [SD] age at the baseline = 8.6 ± 0.9 years, range = 7.0–11.1 years, including 1,223 boys and 1,219 girls). The neuroimaging subsample comprised 260 children (mean ± SD age at baseline = 8.4 ± 0.8 years, range = 7.1–10.3 years, including 134 boys and 126 girls) with complete imaging assessment, after the exclusion of statistically defined extreme video gamers (n = 3).

All parents or tutors signed the informed consent form approved by the Research Ethical Committee (No. 2010/41221/I) of the Hospital del Mar Medical Research Institute, Barcelona, Spain and the FP7-ERC-2010-AdG Ethics Review Committee (268479-22022011).

Behavioral Measurements

VIDEO GAME USE. At baseline, parents were asked to estimate how much time (in hours) their child was currently playing video games on an average weekday and weekend. A single score was computed from these estimates to indicate average hours of play per week. To document the video games most frequently used, 100 families from the MRI group completed a follow-up questionnaire indicating which games were most frequently played in the previous year. As shown in Table 1, 96% of children played visuomotor skill-based games (eg, Super Mario Bros), full-body active games (exergames; eg, Wii Sports), or both.

COGNITIVE TESTING. The selected cognitive assessment included motor response speed, attention, and working memory. Speed of motor responses and attention were assessed using the computerized child version of the Attentional Network Test (ANT).14 Overall reaction time was used to index speed of motor responses to visual stimulation and its SD was used to index motor response consistency as a measurement of sustained attention.15 Specific visual attention features additionally measured by

<table>
<thead>
<tr>
<th>Game</th>
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<tr>
<td>Visuomotor skill-based games*</td>
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</tr>
<tr>
<td>Platforms</td>
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<tr>
<td>Mario Bros/Super Mario Bros/</td>
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<tr>
<td>NBA</td>
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<tr>
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<td>48</td>
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</tr>
<tr>
<td>Mario Party</td>
<td>17</td>
</tr>
<tr>
<td>Wii Party</td>
<td>4</td>
</tr>
<tr>
<td>Adventure/strategy games*</td>
<td>49</td>
</tr>
<tr>
<td>Animal Crossing</td>
<td>12</td>
</tr>
<tr>
<td>Pokemon</td>
<td>11</td>
</tr>
<tr>
<td>Minecraft</td>
<td>6</td>
</tr>
<tr>
<td>Doge/Horsez</td>
<td>4</td>
</tr>
<tr>
<td>Others</td>
<td>12</td>
</tr>
<tr>
<td>Visuomotor skill-based games and/or</td>
<td>96</td>
</tr>
<tr>
<td>full-body active games</td>
<td></td>
</tr>
<tr>
<td>Adventure/strategy games alone</td>
<td>4</td>
</tr>
</tbody>
</table>

*Based on reports from 100 families indicating up to 3 games played the most in the year before magnetic resonance imaging. **Alone or combined.
the ANT task are “alerting,” “orienting,” and “interference.” Commission and omission errors were also registered. Cases with >30% commission or omission errors were excluded from further analyses (total 9 cases in the whole sample and 2 cases in the MRI sample). In the final whole sample, commission errors were 3.1% and omission errors were 1.1%.

A computerized version of the N-Back task was used to assess working memory: “Detectability” (normalized hit rate minus normalized false alarm rate) on 2-back and 3-back loads was used to overall index test performance accuracy. See Forns et al for more detail.

**BEHAVIORAL ASSESSMENT.** Parents completed the Strengths and Difficulties Questionnaire (SDQ) of child behavioral problems. The questionnaire includes 25 questions on psychological attributes, some positive and some negative, which are rated from 0 to 2 points each. Responses are divided in 5 separate scales: “emotional symptoms,” “conduct problems,” “inattention/ hyperactivity,” “peer relationship problems,” and “prosocial behavior.” A “difficulties” score ranging from 0 to 40 was generated by summing the scores for the scales 1 to 4. Parents were also asked to report the average daily sleeping time (in hours) of their children. Overall school achievement was rated by teachers using a 5-point scale (from the worse = 1 to the best = 5).

**ADDITIONAL CONTEXTUAL ASSESSMENTS.** A neighborhood socioeconomic status vulnerability index (Urban Vulnerability Atlas of Spain, 2012; http://www.fomento.gob.es/MFOM/LANG_CASTELLANO/DIRECCIONES_GENERALES/ARQ_VIVIENDA/SUELO_Y_POLITICAS/OBSERVATORIO/Atlas_Vulnerabilidad_Urbana/), at both school and home addresses, was used as a composite measure of sociodemographic factors including level of education, unemployment, and occupation at the census tract. Parental education was registered using a 5-point scale (1 = illiterate; 2 = less than primary; 3 = primary; 4 = secondary; 5 = university).

**MRI Acquisition**

MRI was administered a mean ± SD of 1.2 ± 0.4 years after study baseline. A 1.5T Signa Excite system (General Electric, Milwaukee, WI) equipped with an 8-channel phased-array head coil and single-shot echo planar imaging (EPI) software were used. The imaging protocol included high-resolution T1-weighted 3-dimensional (3D) anatomical images, diffusion tensor imaging (DTI), and a 6-minute functional MRI sequence acquired in the resting state with eyes closed. Acquisition parameters for the 3 sequences are fully described in a previous report.

**Image Preprocessing**

**ANATOMICAL 3D.** After visual inspection by a trained operator, images from 9 children were discarded as a result of poor image quality, and thus the final sample for the anatomical analysis included 251 children. Statistical parametric mapping (SPM) voxel-based morphometry algorithms with DARTEL registration were used to measure gray and white matter tissue concentration and volume at a voxel level. The preprocessing steps are detailed elsewhere. Such a preprocessing approach implied the generation of a study-specific template. Normalized images were transformed to the standard SPM template, resliced to 1.5mm resolution in Montreal Neurological Institute (MNI) space.

**DTI.** The Functional MRI of the Brain (FMRIB) Software Library 5.0 (FSL), developed by the Analysis Group at the Oxford Centre for FMRIB, was used to process DTI images. Image quality control was rigorous in this study. A total of 86 children were removed from the DTI analysis on the basis of suboptimal image quality (see Pujol et al). The final DTI sample involved 174 children with a mean ± SD of 23.5 (94%) ± 1.9 optimal-quality volumes. Fractional anisotropy (FA) maps were estimated using FMRIB’s Diffusion Toolbox in FSL after local fitting of the diffusion tensor model at each voxel (“dtifit”). Tract-Based Spatial Statistics was used to process diffusion data. FA data sets were resliced to a 1mm × 1mm × 1 mm anatomical resolution and normalized to standard MNI space via the FMRIB58_FA template and the Non-linear Registration Tool.

**FUNCTIONAL MRI.** Preprocessing was carried out using SPM8 and involved motion correction, spatial normalization, and smoothing using a Gaussian filter (full-width half-maximum = 8mm). Data were normalized to the standard SPM-EPI template and resliced to 2mm isotropic resolution in MNI space.

The comprehensive procedures adopted to control for potential head motion effects are fully described in our previous report. In short, the procedures included: (1) conventional SPM time-series alignment; (2) exclusion of 24 children with outlier head motion (mean interframe motion > 0.12mm; the finally analyzed sample therefore included 236 children); (3) use of both motion-related regressors and estimates of global brain signal fluctuations as confounding variables in first-level analyses; (4) within-subject, censoring-based MRI signal artifact removal (scrubbing); and (5) use of the mean interframe motion across the functional MRI run for each participant as a regressor in the second-level analyses.

Based on the results obtained from the anatomical and DTI assessments, our functional connectivity analysis was focused on the left basal ganglia. To systematically explore basal ganglia, we targeted 4 key subregions: the dorsal and ventral aspects of both caudate nucleus and putamen. Based on a widely applied method for mapping basal ganglia functional connectivity, the regions of interest (or “seeds”) were centered at MNI coordinates (in millimeters): (1) dorsal caudate nucleus (x = −13, y = 15, z = 9); (2) dorsal putamen (x = −28, y = 1, z = 3); (3) ventral caudate nucleus, corresponding approximately to the nucleus accumbens (x = −9, y = 9, z = −8); and (4) ventral putamen (x = −20, y = 12, z = −3). The procedures adopted to generate functional connectivity maps from the selected regions are detailed in our previous report.
Statistical Analysis

BEHAVIORAL DATA. Student t test was used to compare sex and age group means as to hours of play per week. Analysis of variance (ANOVA) adjusted by age and sex (ANCOVA) was used to compare gamers and nongamers as to the behavioral variables. Assumptions under the ANCOVA were met in these analyses including homogeneity of variance, homogeneity of regression slopes of covariates, and noncollinearity. Linear regression adjusted by age and sex was used to assess the relationship between video game playing time and behavioral variables within the user group. Bonferroni correction was applied to account for multiple testing within each analysis. Only probability values < 0.006 (0.05/8) were considered significant. Finally, to easily illustrate the identified associations of gaming with behavior variables, the data were plotted as discrete groups on the basis of playing time in hours (Figs 1 and 2).

IMAGING ANALYSES. ANOVA with age and sex as covariates (ANCOVA) and cross-correlations were estimated using SPM. Separate second-level (group) analyses were carried out for whole-brain estimates of regional white matter and gray matter volume, DTI fractional anisotropy, and resting-state functional connectivity. The behavioral results obtained in the large sample helped to guide the subsequent imaging analysis. Specifically, 3 groups of children were defined on the basis of video game use: 45 nongamers, 113 low-use gamers (playing 1–2 hours per week), and 102 high-use gamers (playing 3 or more hours per week). The cutoff between low and high use corresponds to the median of time of play per week in all gamers. In each case, ANCOVA allowed us to first identify brain areas showing overall differences among these groups. From each analysis, a plot was then generated using brain measurements extracted at representative peak differences to characterize the direction of findings with no bias. That is, the 3-group model allowed us to fairly capture any change combination among the 3 groups (nonplaying, low use, and high use) in a single model sensitive to both the effect of playing versus nonplaying and the effect of low versus high use. Finally, post hoc whole-brain t tests were used to compare nongamers with high-use gamers. In addition, we mapped voxelwise the correlation across subjects between individual brain images and the child’s age at the time of MRI acquisition.

Results

Behavioral Results

The overall study sample comprised 428 nongamers and 2,014 gamers (ie, playing 1 hour or more per week). As a group, gamers played a mean of 4.0 ± 2.9 hours per week (see Fig 1 for participant distribution according to playing time). Boys overall played video games 1.7 hours per week more than girls (95% confidence interval [CI] = 1.4–1.9 hours). There was also a significant effect of age, with older children (above sample median age) playing 0.4 hours per week more than younger children (95% CI = 0.2–0.7 hours). Although the correlation between age and gaming hours was significant in the whole group (r = 0.0004), collinearity between both variables was very low (r = 0.07, tolerance = 0.995).
No significant differences were found between gamers and nongamers as to parents’ education. For example, 59.1% of mothers of gamers had university-level education versus 61.0% in nongamers (chi-square = 0.5, \( p = 0.511 \)). All further statistical analyses were adjusted for sex and age (ie, age and sex were included as covariates in both group comparisons and correlations).

Video gamers showed faster motor response to visual stimulation (ie, shorter reaction time) than nongamers, with a mean group difference of 65 milliseconds (Table 2). Such a difference was statistically robust (\( t = 8.1, \ p = 6e^{-16} \)) and was also significant in the smaller subsample of children who underwent MRI (\( t = 4.5, \ p = 0.0001 \)). Video game use was also associated with higher consistency of motor responses, measured as reaction time SD (see Table 2). However, there were no effects on specific attentional features or working memory. Remarkably, most of the advantage in motor responsiveness was observed in children playing 1 hour per week (mean difference of 52 milliseconds with respect to nongamers), with minimal further change observed after 2 hours per week (illustrated in Fig 2).

By contrast, in the gaming group, the weekly time spent gaming was steadily associated with higher scores in parental ratings of strengths and difficulties (SDQ), particularly in regard to conduct problems (see Fig 2), peer conflicts, and reduced prosocial abilities (see Table 2). In our post hoc analyses, children gaming in the range of 9 to 17 hours per week showed significantly more behavioral problems than nongamers (eg, conduct problems; \( t = 3.3, \ p = 0.001 \)). In addition, the amount of time that was spent gaming was negatively associated with sleeping time. However, as a group, gamers and nongamers did not differ as to strengths and difficulties ratings. Interestingly, gamers showed significantly higher school achievement scores (see Table 2).

A further analysis stratified by sex and age showed that the identified associations were notably consistent across the subgroups. In both boys and girls, most of the attained group difference in motor responsiveness was observed playing 1 hour per week, as in the older children subgroup. It is relevant, however, that further improvement was observed up to 3 to 4 hours of play in the younger children. In each stratified group, differences between gamers and nongamers were highly significant (boys, \( p = 2e^{-10} \); girls, \( p = 5e^{-8} \); younger, \( p = 3e^{-10} \); older, \( p = 9e^{-8} \)). However, such differences between gamers and nongamers were significantly stronger in girls compared to boys (interaction \( t = 2.6, \ p = 0.008 \)). No significant interaction was identified for age. Similarly, gamers showed a significant linear correlation between playing hours and conduct problems in each subgroup analysis (boys, \( p = 0.001 \); girls, \( p = 1e^{-3} \); younger, \( p = 8e^{-6} \); older, \( p = 0.002 \)). No significant interaction was found in the correlation analysis. Also, we found no significant differences between gamers and nongamers in any subgroup (boys, girls, younger, and older children) as to conduct problems. All associations remained highly significant when analyses were further adjusted for other potential confounding variables, including parental education and neighborhood vulnerability index as the most representative indicators of children’s close environment.

**Imaging Results**

Age and sex were included as covariates in all imaging analyses.

**T1-weighted 3D Anatomical Images.** When examining differences among nongamers and low- and high-use gamers in white matter volume measurements (ANCOVA), significant results were observed in a region adjacent to the ventral and lateral aspect of the striatum and extending to the temporal lobe (Fig 3 and Supplementary Table). A plot of white matter volume measurements from the left ventral striatal region revealed a similar increase of white matter volume in both low- and high-user groups compared with nonusers. Figure 3 illustrates this effect in post hoc comparison between nonusers and high users. In addition, the primary ANCOVA identified significant effects in the brainstem (see Supplementary Table), which were not observed in the post hoc analysis. No significant results were obtained for gray matter volumes.

**DTI FA.** ANCOVA consistently showed significant differences among groups in a region adjacent to the lateral aspect of the striatum (see Fig 3, Supplementary Table). The plot of FA measurements from this left striatal region revealed a relevant FA increase, but only in the high-use group (see Fig 3). Direct comparison of nongamers with high-use gamers confirmed the increase of FA in this region and detected additional significant changes in the ventral striatal region bilaterally, as well as the right thalamus and left occipital white matter.

**Basal Ganglia Functional Connectivity.** Video gaming was associated with higher functional connectivity in the putamen and caudate nucleus maps (Fig 4 and Supplementary Table). Specifically in the putamen maps, a significant connectivity increase was identified with the left motor cortex and prefrontal cortex in both low- and high-use groups. In the ventral caudate map, the most relevant finding involved the dorsal anterior cingulate cortex (ACC), which demonstrated a strong effect in the high-use group and an intermediate effect in the low-use group compared with nongamers. The post
hoc analysis confirmed a robust connectivity increase with the ACC and additionally with the left and right anterior insula region (see Fig 4 and Supplementary Table).

**CORRELATION ANALYSIS WITH AGE.** White matter volume increased with age in the visual system and sensory–motor projection tracts at the level of basal ganglia and thalamus, with no areas showing negative correlations with age (Fig 5). FA increased significantly with age mostly in the internal capsule and basal ganglia. Therefore, the identified associations of video gaming with white matter measurements were generally consistent with the effects of age in

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**TABLE 2. Behavioral Results**

<table>
<thead>
<tr>
<th>Measure</th>
<th>Group Comparison</th>
<th>Correlation with Weekly Play Time, Users, n = 2,014</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nonusers, n = 428, Adj. Mean (95% CI)</td>
<td>Users, n = 2,014, Adj. Mean (95% CI)</td>
</tr>
<tr>
<td>Cognitive tests</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reaction time, ms</td>
<td>861 (847 to 875)</td>
<td>796 (790 to 802)</td>
</tr>
<tr>
<td>Reaction time SD, ms</td>
<td>286 (278 to 294)</td>
<td>267 (264 to 271)</td>
</tr>
<tr>
<td>Alerting, ms</td>
<td>43 (35 to 50)</td>
<td>49 (46 to 53)</td>
</tr>
<tr>
<td>Orienting, ms</td>
<td>32 (25 to 40)</td>
<td>35 (31 to 38)</td>
</tr>
<tr>
<td>Interference, ms</td>
<td>66 (60 to 72)</td>
<td>63 (60 to 66)</td>
</tr>
<tr>
<td>2B working memory, d’</td>
<td>2.2 (2.1 to 2.3)</td>
<td>2.3 (2.2 to 2.3)</td>
</tr>
<tr>
<td>3B working memory, d’</td>
<td>1.1 (1.0 to 1.2)b</td>
<td>1.2 (1.2 to 1.3)c</td>
</tr>
<tr>
<td>Behavior</td>
<td></td>
<td></td>
</tr>
<tr>
<td>School achievement</td>
<td>3.41 (3.31 to 3.51)</td>
<td>3.60 (3.55 to 3.64)</td>
</tr>
<tr>
<td>Sleeping time, h</td>
<td>9.63 (9.56 to 9.7)</td>
<td>9.56 (9.53 to 9.59)</td>
</tr>
<tr>
<td>Total SDQ score</td>
<td>8.6 (8.1 to 9.1)</td>
<td>8.3 (8.1 to 8.6)</td>
</tr>
<tr>
<td>Conduct problems</td>
<td>1.7 (1.5 to 1.8)</td>
<td>1.6 (1.6 to 1.7)</td>
</tr>
<tr>
<td>Problems with peer</td>
<td>1.2 (1.0 to 1.3)</td>
<td>1.1 (1.1 to 1.2)</td>
</tr>
<tr>
<td>Emotional symptoms</td>
<td>1.9 (1.7 to 2.1)</td>
<td>1.9 (1.9 to 2.0)</td>
</tr>
<tr>
<td>Inattention/ hyperactivity</td>
<td>3.8 (3.6 to 4.1)</td>
<td>3.7 (3.6 to 3.8)</td>
</tr>
<tr>
<td>Prosocial behavior</td>
<td>8.5 (8.3 to 8.7)</td>
<td>8.5 (8.4 to 8.6)</td>
</tr>
</tbody>
</table>

All the analyses were adjusted by sex and age. 2B and 3B indicate 2-back and 3-back, respectively.

*Significant after Bonferroni correction.

bn = 408.
b* n = 1967.
Adj. = adjusted; CI = confidence interval; d’ = detectability; SD = standard deviation; SDQ = Strengths and Difficulties Questionnaire.
terms of their positive direction and (partial) anatomical overlap. Similarly, age was associated with higher functional connectivity between the putamen and motor cortex in the area associated with video game use, albeit significant only for right hemispheric effects. For the ventral caudate, age was instead negatively associated with its functional connectivity with a medial frontal area adjacent to the ACC region that was positively associated with gaming use (and additionally with the upper brainstem and amygdala).

Discussion

In the cognitive domain, video game use was associated with faster motor response to visual stimulation. Importantly, no further change in motor speed was identified in children playing >2 hours per week, suggesting an early ceiling effect for this measurement. Video gamers, overall, did not demonstrate more problematic behavior that nongamers, although, within the gaming group, we observed a positive association between time weekly spent gaming and conduct problems, peer conflicts, and reduced prosocial abilities, as well as a negative association with sleeping time. At a neural level, structural and functional brain changes associated with gaming use were most evident with respect to basal ganglia circuitry.

Our cognitive assessment selectively measured processing speed, attention, and working memory as primary cognitive domains of interest. In the literature, there is some consensus that gaming may improve speed of information processing and motor response; however, the reported effects on executive function and working memory, as well as general intelligence, have been less consistent. For example, working memory has been shown to improve in some studies, whereas others reported only partial associations or negative results. In our study, larger differences in reaction time than in working memory support the contention that several abilities are more trainable than others with commercial video games.

We observed an association between gaming use and changes in basal ganglia circuits in the form of structural (white matter) and functional connectivity increases. These findings are intuitive in many respects, as it is well known that basal ganglia circuits are critical for procedural learning based on the acquisition of new skills through practice. Our results are also generally consistent with other imaging studies reporting significant effects of gaming use on the frontobasal ganglia system structure and function. Specifically, Erickson et al showed that the acquisition of skills on demanding video games may be predicted by variations in the volume of the striatum. Children traditionally acquire procedural skills through action, for instance in relation to sports and outdoor games. Neuroimaging research now suggests that training with desktop virtual environments is also capable of modulating brain systems that support procedural learning. However, the type of video game is likely to be particularly relevant with regard to the modulation of specific neural systems. For example, the use of logic

![Figure 3: Gaming group differences in white matter volume measurements (top left) and diffusion tensor imaging fractional anisotropy (top right). Analysis of variance adjusted by age and sex (ANOVA) indicated significant changes adjacent to the striatum in both analyses. Mean and 95% confidence intervals are plotted for each modality, reflecting regional peak differences among gaming groups. The bottom left and right images illustrate corresponding differences identified between non-gamers versus high-use gamers. The left hemisphere corresponds to the left side of coronal views.]

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and "platform jumping" games has been associated with anatomical changes implicating the hippocampal system in adults.³⁸,³⁹

Reviews of the literature generally conclude that excessive video game use has negative effects on some aspects of behavior in children.¹,³,⁵,⁴⁰ Most of the identified problems relate to the extent to which gaming can become a behavioral addiction, or to the violent content of some of the games themselves. We did not include children with probable gaming addiction, and the amount of time spent on playing in our population (mean 5 4 hours per week, range 0–17 hours) was far less than the gaming level that has been associated with severe psychosocial health disturbances. Despite this, gaming use in our study was positively related to the presence of conduct problems, peer conflicts, and reduced prosocial abilities. This finding is in agreement with the results of a national survey of 10- to 15-year-old children in the United Kingdom, in which gaming use of > 3 hours per day was associated with similar behavioral problems.⁴¹ Thus, frequent video game use does appear to be associated with behavioral problems in

FIGURE 4: Gaming group differences in basal ganglia functional connectivity. Analysis of covariance (ANCOVA) indicated significant changes in functional connectivity between the dorsal putamen (DP) and motor cortex at the level of the frontal operculum. In the ventral caudate (VC) analysis, the most relevant finding involved the anterior cingulate cortex. Mean and 95% confidence intervals are plotted, reflecting regional peak differences among gaming groups. The bottom left and right images illustrate corresponding differences identified between nongamers versus high-use gamers. The left hemisphere corresponds to the left side of axial views.

FIGURE 5: Correlations between age and brain measurements. White matter volume (A) increased significantly with age in projection tracts (and the visual system). Fractional anisotropy (B) increased with age mostly in the internal capsule and basal ganglia. Age was also associated with stronger functional connectivity between the putamen and motor cortex (C) and weaker functional connectivity between the caudate nucleus and the medial frontal cortex (D) and additionally with the upper brainstem (and amygdala).
children even in the absence of a recognized gaming disorder, although the direction of the relationship is less clear. That is, children with peer conflicts and reduced prosocial abilities may show a tendency to isolate and spend more time playing video games. Conversely, highly frequent use likely consumes a proportion of the child’s leisure time and invades other activities that may ultimately affect the normal development of prosocial abilities.

Our study was limited in that we applied a selective cognitive and behavioral assessment that did not comprehensively cover all domains. Also, we estimated the time spent gaming from parental reports but with no formal assessment of the fidelity of these reports in terms of their reliability and accuracy. It is also relevant to mention that the estimation of video game use was based on a single cross-sectional measurement. Finally, the type of video games played varied, and we were unable to perform specific analyses stratified by game content or type.

In conclusion, we have investigated relationships between weekly video game use, selected cognitive abilities, conduct-related problems, and brain correlates in a large group of children. Relatively small amounts of video gaming were associated with better performance in certain visuomotor skills and, by contrast, a frequent use was associated with the presence of conduct problems. Whereas motor response differences between gamers and nongamers were already established with 2 hours per week of video gaming, conduct problems appear to be more common in school children playing 9 hours weekly or more. It is important to emphasize that this is a correlational study and, as such, does not permit direct inferences regarding causal relationships. Further studies will be needed to explore the extent to which better visuomotor skills and conduct problems are a cause or consequence of gaming.

Acknowledgment
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Author Contributions
J.P., J.S., L.B.-H., and J.D. contributed to the study concept and design; J.P., M.A.-P., J.F., S.G.-O., R.F., D.M., and G.M.-V. contributed to data acquisition and analysis; J.P. and B.J.H. contributed to drafting the manuscript and figures.

Potential Conflicts of Interest
Nothing to report.

References


Anomalous White Matter Structure and the Effect of Age in Down Syndrome Patients

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Abstract.
Background: Neural tissue alterations in Down syndrome are fully expressed at relatively late developmental stages. In addition, there is an early presence of neurodegenerative changes in the late life stages.
Objective: The aims of this study were both to characterize white matter abnormalities in the brain of adult Down syndrome patients using diffusion tensor imaging (DTI) and to investigate whether degenerative alterations in white matter structure are detectable before dementia is clinically evident.
Methods: Forty-five adult non-demented Down syndrome patients showing a wide age range (18–52 years) and a matched 45-subject control group were assessed. DTI fractional anisotropy (FA) brain maps were generated and selected cognitive tests were administered.
Results: Compared with healthy controls, non-demented Down syndrome patients showed lower DTI FA in white matter involving the major pathways, but with more severe alterations in the frontal-subcortical circuits. White matter FA decreased with age at a similar rate in both DS and control groups.
Conclusions: Our results contribute to characterizing the expression of white matter structural alterations in adult Down syndrome. However, an accelerated aging effect was not demonstrated, which may suggest that the FA measurements used are not sufficiently sensitive or, alternatively, age-related white matter neurodegeneration is not obvious prior to overt clinical dementia.

Keywords: Accelerated aging, diffusion tensor imaging, magnetic resonance imaging, neurodegeneration

1These authors contributed equally to this work.

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INTRODUCTION

Although research in Down syndrome has substantially progressed in the understanding of basic mechanisms via which gene overexpression interferes with brain development, there is less information as to the general organization of the adult brain and the effect of age [1, 2]. The brain contains billion of neurons that communicate with each other via axons to create complex neural networks. The structural mapping of these networks is essential for understanding brain function [3]. However, although new imaging techniques have emerged in recent years, our knowledge of structural connectivity in Down syndrome is still limited.

Diffusion tensor imaging (DTI) is a noninvasive method that provides information about the microstructural properties of brain tissue by measuring the magnitude and direction of water molecule diffusion [4]. In white matter, the diffusion of water molecules is less restricted along the long axis of a group of aligned tissue fibers than perpendicular to it. This condition of directionally-dependent diffusion is referred to as “anisotropic”. The most commonly used measure for diffusion anisotropy is fractional anisotropy (FA), which serves to characterize white matter tracts by mapping directional diffusion restrictions related mainly to fiber density, axonal diameter and myelination degree [3, 4].

DTI has been used to characterize both age-related changes and disease [5–8]. Results from leading studies have shown that age-related changes are associated with FA decreases [5]. DTI findings have been validated by postmortem histological studies showing that advanced age is linked to alterations of almost all white matter components. Segments of axons degenerate and swell, myelin becomes less compact and glial cells accumulate cellular debris, form glial scars and increase in number [9–11].

In Down syndrome, a number of magnetic resonance imaging (MRI) studies have identified a variety of anatomical [12–17] and functional [18, 19] alterations. The frontal lobes [16, 20] and related circuits [18, 19], as representative of late maturing structures, generally show the most relevant alteration. By contrast, some studies have reported a relative preservation of tissue volume in temporal and parietal regions [16, 20, 21]. Nevertheless, white matter alterations and their functional significance have not been fully characterized using FA measurements. There is one previous study indicating that white matter in adult Down syndrome is abnormal in terms of FA, particularly when dementia is clinically evident [22]. Nevertheless, the number of Down syndrome patients without dementia was small in this study (n = 10) and the study design did not allow a distinction between the effect of aging and pre-existing changes. It is not currently evident whether the neurodegenerative effect on white matter structure is detectable before dementia is expressed. If this were indeed the case, age-related FA changes could serve as early markers of neurodegeneration, which could be of high practical interest given the difficulty in identifying the cognitive deficits related to dementia in a population with significant baseline alteration in cognition [1, 2].

The aims of this study were both to characterize white matter abnormalities in the brain of adult non-demented Down syndrome patients using DTI and to investigate whether degenerative changes in white matter structure are detectable before dementia becomes clinically evident. We predicted that Down syndrome patients would exhibit both widespread FA reduction in white matter and an accelerated aging effect, and that the alteration would be associated with poorer cognitive performance.

METHODS

Participants

Sixty-eight Down syndrome patients were initially recruited in the study. Candidates were recruited from the community via parent organizations and underwent comprehensive medical, psychiatric, neuropsychological, and laboratory evaluation. Individuals with seizure or neurological disease (other than Down syndrome) and non-stable medical conditions were not considered eligible. Participants were selected on the basis of age (18 years old and upwards), Down syndrome confirmed by karyotype, capability to understand MRI instructions, follow commands and keep still, and also optimal attitude and willingness (patients and parents) to participate. Five patients did not complete the image acquisition protocol and a further 18 subjects were ultimately excluded due to head motion during MRI (see below). The final sample, therefore, included 45 relatively high-functioning Down syndrome patients (29 females, 16 males) with genotype-confirmed trisomy 21 and a mean ± SD age of 35.3 ± 10.8 years, range 18–52 (Table 1). The included and excluded patient subgroups did not significantly differ as to age, performance IQ, semantic fluency, and working
Table 1

<table>
<thead>
<tr>
<th>Characteristics of study participants</th>
<th>Down Syndrome</th>
<th>Healthy Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (mean, SD years)(^a)</td>
<td>35.3 (10.8)</td>
<td>34.6 (10.0)</td>
</tr>
<tr>
<td>Gender (men/women)</td>
<td>16/29</td>
<td>19/26</td>
</tr>
<tr>
<td>Medical Background (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>28.9%</td>
<td></td>
</tr>
<tr>
<td>Respiratory</td>
<td>24.4%</td>
<td></td>
</tr>
<tr>
<td>Metabolic / Endocrine</td>
<td>46.7%</td>
<td></td>
</tr>
<tr>
<td>Ophthalmological</td>
<td>66.7%</td>
<td></td>
</tr>
<tr>
<td>Otorhinolaryngological</td>
<td>6.7%</td>
<td></td>
</tr>
<tr>
<td>Disability Levels (%)(^b) DSM - IV - TR</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild</td>
<td>57.8%</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>42.2%</td>
<td></td>
</tr>
<tr>
<td>Severe</td>
<td>0%</td>
<td></td>
</tr>
<tr>
<td>Profound</td>
<td>0%</td>
<td></td>
</tr>
<tr>
<td>Knowledge (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Illiterate</td>
<td>51.1%</td>
<td></td>
</tr>
<tr>
<td>Read/Write</td>
<td>48.9%</td>
<td></td>
</tr>
<tr>
<td>Years of Schooling (mean, SD years)</td>
<td>9.3 (3.9)</td>
<td></td>
</tr>
<tr>
<td>Neuropsychological Assessment:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Performance IQ, K-BIT (mean, SD)(^b)</td>
<td>60.5 (8.0)</td>
<td></td>
</tr>
<tr>
<td>Semantic Fluency (mean, SD)(^c)</td>
<td>9.8 (3.4)</td>
<td></td>
</tr>
<tr>
<td>Digit Span Test (mean, SD)(^d)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Forward Span</td>
<td>2.7 (1.2)</td>
<td></td>
</tr>
<tr>
<td>Backward Span</td>
<td>1.2 (1.1)</td>
<td></td>
</tr>
<tr>
<td>Forward Score</td>
<td>3.3 (1.5)</td>
<td></td>
</tr>
<tr>
<td>Backward Score</td>
<td>1.2 (1.2)</td>
<td></td>
</tr>
<tr>
<td>Total Score</td>
<td>4.5 (2.3)</td>
<td></td>
</tr>
</tbody>
</table>

SD, standard deviation. \(^a\)Between-group differences for age were not significant (\(p=0.76\)).
\(^b\)K-BIT, Kaufman Brief Intelligent Test (2nd edition); matrices test. \(^c\)Semantic fluency, animals in 1 minute. \(^d\)Digit span, verbal short-term memory test.

Cognitive testing

Selected cognitive assessment in all Down syndrome patients included Performance IQ estimated with the Kaufman Brief Intelligent Test, Second Edition (K-BIT) [23] matrices subtest as a general cognitive assessment, the Wechsler Adult Intelligence Scale (WAIS) backward digit span task as one of conventional measurements of working memory [24], and semantic fluency as a sensitive measurement of verbal output [25].

Additionally, each patient underwent comprehensive neurological and psychiatric history and subsequent tailored neuropsychological testing to clinically rule out the presence of dementia (and mild cognitive impairment, MCI) in terms of cognitive deterioration overlapping with developmental cognitive deficits associated with Down syndrome. Clinical diagnosis (or exclusion) of dementia in Down syndrome by experienced clinicians is recommended as being more accurate and reliable than operative diagnostic tools at a relatively early stage of the disease [26]. Nevertheless, apart from expert clinical diagnosis (by SE and SdS), no patient met International

memory. Excluded patients, however, were predominantly males (6 females, 17 males).

A control group of 45 healthy volunteers were selected matched for age with the patient sample (34.6 ± 10.0 years, range 19–51) and showing similar sex distribution (Table 1). Participants were either friends or family of subjects participating in the current and other studies, or were recruited from local advertisements. A complete medical interview was carried out to exclude individuals with relevant medical or neurological disorders, cerebrovascular risk factors, substance abuse, psychiatric disease, or undergoing medical treatment. In all included control subjects, the brain showed a normal appearance on high resolution anatomical MRI scans.

This study was conducted according to the principles expressed in the Declaration of Helsinki. Anxiolytics were not administered in this study. The study protocol was approved by the Clinical Research Ethical Committee of the Parc de Salut Mar of Barcelona. Written informed consent was obtained from parents and control subjects. Verbal or written assent was additionally obtained from Down syndrome patients.
Classification of Diseases (ICD)-10 [27] or Diagnostic and Statistical Manual of Mental Disorder-IV-Text Revision (DSM-IV-TR) [28] criteria for dementia. A total of 3 patients, however, did meet criteria for mild MCI [29], adapted to adults with intellectual disability [30]. The clinical diagnosis of MCI in these 3 cases was established by an experienced neuropsychologist (SEC) on the basis of (i) a report of cognitive impairment by the patient (confirmed by a reliable informant) or by a reliable informant that implies a change from previous capacities, (ii) abnormal performance in the corresponding neuropsychological testing, and (iii) no clinically relevant decline in overall adaptive skills and insufficient ICD-10 and DSM-IV-TR criteria for dementia.

MRI acquisition

A 1.5 Tesla Signa Excite System (General Electric, Milwaukee, WI, USA) equipped with an eight-channel phased-array head coil and single-shot echoplanar imaging (EPI) software was used. Diffusion-weighted scans were obtained using spin-echo single-shot echo-planar sequences of 25 directions with a B-factor of 1000 s/mm². Acquisition parameters were repetition time 8300 ms; echo time 94 ms; thickness 5 mm, no gap; pulse angle 90°; field of view 26 cm; 128 × 128 acquisition matrix reconstructed into a 256 × 256 matrix, and scan duration was 3 m 52 s. Twenty-six slices were prescribed parallel to the anterior-posterior commissure line covering the whole brain. Participants were instructed to relax, stay awake and lie still.

Image preprocessing

DTI was processed using Functional MRI of the Brain (FMRIB) Software Library 5.0 (FSL), developed by the Analysis Group at the Oxford Centre for FMRIB [31]. Diffusion-weighted images were aligned to the B0 image using affine registration and corrected for motion and eddy current distortions (“Eddy Current Correction” option in the FMRIB Diffusion Toolbox [FDT] version 2.0 in FSL). A whole-brain mask, generated with the FSL Brain Extracting Tool, was applied to the DTI images. Subsequently, we estimated FA maps using FDT in FSL by local fitting of the diffusion tensor model at each voxel (“dtifit”). FA maps were then aligned to a common target (FMRIB58_FA template) using Tract-Based Spatial Statistics [32], re-sliced to a 1 mm × 1 mm × 1 mm anatomical resolution and normalized to standard MNI space via the FMRIB58_FA template using the FMRIB’s Non-linear Registration Tool. All the images were visually inspected by a trained researcher before and after the preprocessing steps to avoid the inclusion of poor-quality images. An additional rigorous image quality control was carried out to identify potential effects of head motion on raw images, which involved the visual inspection of each DTI slice for all 25 DTI volumes in all participants. The effect of motion can generally be observed as signal loss in the whole or a part of one slice compared with the other slices. Volumes with slices showing signal loss (greater than ~10% compared with slices normal in signal and measured using the conventional MRICron display tool) or residual artifacts were identified by an expert researcher. DTI full examinations showing more than 5 degraded volumes were discarded. A total of 18 participants were removed from the DTI analysis on the basis of this criterion (in addition to 5 cases showing gross image degradation). The final DTI sample involved 45 patients with a mean ± SD of 23.0 (92%) ± 1.6 optimal-quality volumes.

After the full pre-processing, FA maps were transferred to the SPM8 platform and smoothed with an 8 mm Gaussian Kernel to carry out group statistical analyses.

Statistical analysis

Individual FA maps were included in second-level (group) SPM analyses using 2-sample t-test between the 45 Down syndrome patients and 45 healthy controls. Voxel-wise analyses in SPM were also performed to map the correlation between age and whole-brain FA measurements in both groups. Finally, voxel-wise analyses were performed to map the correlation between individual ratings in the selected cognitive tests and FA measurements in the Down syndrome group.

Results were considered significant with clusters of 1.032 ml (1,032 voxels) at a height threshold of p < 0.005, which satisfied the family-wise error (FWE) rate correction of P FWE < 0.05 according to Monte Carlo simulations [33].

RESULTS

Fractional anisotropy differences between Down syndrome and control subjects

Down syndrome patients showed a widespread white matter FA reduction compared with healthy
controls involving parts of the frontal lobes, semioval centers, corpus callosum, external capsule, internal capsule, putamen, thalamus, pyramidal tracts, and brainstem (Table 2). Therefore, the major brain pathways were affected, although the alterations were more severe in the frontal-subcortical circuits (Fig. 1). For example, 21 out of 25 sub-clusters showing the largest between-group differences \( (t > 5) \) involved the frontal-subcortical circuits and only 4 did not \( (\chi^2 = 11.5, p < 0.001) \). In the opposite contrast, a region of significantly larger FA was found in Down syndrome at the temporo-parietal junction (Fig. 1).

**Correlations with cognitive performance in Down syndrome**

No significant correlations were found between FA measurements and both performance IQ (K-BIT matrices) and working memory scores (WAIS digit span). By contrast, FA showed significant positive correlation with semantic fluency (i.e., lower FA values, poorer performance) in a variety of regions involving the frontal lobes, corpus callosum, semioval centers, arcuate fasciculus, caudate nucleus, external capsule, thalamus, and hippocampus (Fig. 2 and Table 2).

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**Table 2**

Diffusion tensor imaging fractional anisotropy results

<table>
<thead>
<tr>
<th></th>
<th>Cluster size, ml</th>
<th>( x ) ( y ) ( z )</th>
<th>( t ) (r)</th>
<th>( t^{<strong>} ) (r)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Down &lt; Controls</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R semioval center</td>
<td>354.0*</td>
<td>18 –7 53</td>
<td>6.3</td>
<td>5.9</td>
</tr>
<tr>
<td>L semioval center</td>
<td>*</td>
<td>–13 2 51</td>
<td>6.1</td>
<td>5.7</td>
</tr>
<tr>
<td>R frontal lobe</td>
<td>*</td>
<td>20 45 15</td>
<td>7.9</td>
<td>7.5</td>
</tr>
<tr>
<td>L frontal lobe</td>
<td>*</td>
<td>–19 47 13</td>
<td>8.2</td>
<td>7.8</td>
</tr>
<tr>
<td>Corpus callosum</td>
<td>*</td>
<td>4 30 –3</td>
<td>4.3</td>
<td>4.0</td>
</tr>
<tr>
<td>R putamen</td>
<td>*</td>
<td>24 13 16</td>
<td>7.4</td>
<td>7.0</td>
</tr>
<tr>
<td>L putamen</td>
<td>*</td>
<td>–21 11 17</td>
<td>8.0</td>
<td>7.7</td>
</tr>
<tr>
<td>R thalamus</td>
<td>*</td>
<td>10 –9 17</td>
<td>7.2</td>
<td>7.0</td>
</tr>
<tr>
<td>L thalamus</td>
<td>*</td>
<td>–11 –11 18</td>
<td>8.4</td>
<td>8.3</td>
</tr>
<tr>
<td>R pyramidal tract</td>
<td>*</td>
<td>19 –18 –7</td>
<td>7.9</td>
<td>7.4</td>
</tr>
<tr>
<td>L pyramidal tract</td>
<td>*</td>
<td>–19 22 –6</td>
<td>9.2</td>
<td>8.7</td>
</tr>
<tr>
<td>Brainstem</td>
<td>*</td>
<td>1 –32 –23</td>
<td>8.3</td>
<td>7.8</td>
</tr>
<tr>
<td><strong>Down &gt; Controls</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporo-parietal junction</td>
<td>6.7</td>
<td>37 –42 12</td>
<td>5.9</td>
<td>5.9</td>
</tr>
<tr>
<td><strong>Correlations with FA</strong></td>
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<td></td>
<td></td>
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<tr>
<td>Semantic fluency in DS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R frontal lobe</td>
<td>148.7*</td>
<td>27 48 4</td>
<td>4.4 (0.6)</td>
<td>4.2 (0.6)</td>
</tr>
<tr>
<td>L frontal lobe</td>
<td>*</td>
<td>–14 24 33</td>
<td>4.9 (0.6)</td>
<td>4.6 (0.5)</td>
</tr>
<tr>
<td>Corpus callosum</td>
<td>*</td>
<td>13 27 3</td>
<td>3.3 (0.5)</td>
<td>3.3 (0.5)</td>
</tr>
<tr>
<td>R semioval center</td>
<td>*</td>
<td>12 2 36</td>
<td>3.5 (0.5)</td>
<td>3.3 (0.5)</td>
</tr>
<tr>
<td>L semioval center</td>
<td>*</td>
<td>–11 –2 43</td>
<td>4.5 (0.6)</td>
<td>4.3 (0.6)</td>
</tr>
<tr>
<td>L arcuate fasciculus</td>
<td>*</td>
<td>–30 13 14</td>
<td>5.4 (0.6)</td>
<td>5.2 (0.6)</td>
</tr>
<tr>
<td>R caudate nucleus</td>
<td>*</td>
<td>13 27 3</td>
<td>3.3 (0.5)</td>
<td>3.3 (0.5)</td>
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<tr>
<td>L caudate nucleus</td>
<td>*</td>
<td>–19 20 18</td>
<td>4.4 (0.6)</td>
<td>4.5 (0.6)</td>
</tr>
<tr>
<td>R external capsule</td>
<td>*</td>
<td>35 1 –2</td>
<td>4.4 (0.6)</td>
<td>4.2 (0.6)</td>
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<tr>
<td>L external capsule</td>
<td>*</td>
<td>–30 13 14</td>
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<td>5.2 (0.6)</td>
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<tr>
<td>R thalamus</td>
<td>*</td>
<td>8 –23 8</td>
<td>3.4 (0.5)</td>
<td>3.8 (0.5)</td>
</tr>
<tr>
<td>L thalamus</td>
<td>*</td>
<td>–3 –21 10</td>
<td>4.0 (0.5)</td>
<td>4.0 (0.5)</td>
</tr>
<tr>
<td><strong>Age in healthy controls</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R frontal lobe</td>
<td>704.1*</td>
<td>24 48 7</td>
<td>4.2 (–0.4)</td>
<td>4.2 (–0.4)</td>
</tr>
<tr>
<td>L frontal lobe</td>
<td>*</td>
<td>–15 46 12</td>
<td>4.4 (–0.4)</td>
<td>4.4 (–0.4)</td>
</tr>
<tr>
<td>Corpus callosum</td>
<td>*</td>
<td>5 30 4</td>
<td>4.0 (–0.4)</td>
<td>4.0 (–0.4)</td>
</tr>
<tr>
<td>Thalamus</td>
<td>*</td>
<td>–1 –20 7</td>
<td>3.8 (–0.4)</td>
<td>3.7 (–0.4)</td>
</tr>
<tr>
<td>R pyramidal tract</td>
<td>*</td>
<td>13 –20 –25</td>
<td>4.0 (–0.4)</td>
<td>4.0 (–0.4)</td>
</tr>
<tr>
<td>L pyramidal tract</td>
<td>*</td>
<td>–15 –26 –13</td>
<td>3.5 (–0.4)</td>
<td>3.5 (–0.4)</td>
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<tr>
<td><strong>Age in DS patients</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R frontal lobe</td>
<td>4.0</td>
<td>19 42 17</td>
<td>3.8 (–0.4)</td>
<td>3.3 (–0.3)</td>
</tr>
<tr>
<td>L arcuate fasciculus</td>
<td>2.4</td>
<td>–31 6 30</td>
<td>3.5 (–0.4)</td>
<td>3.2 (–0.3)</td>
</tr>
<tr>
<td>R external capsule</td>
<td>5.5</td>
<td>35 11 –2</td>
<td>3.0 (–0.3)</td>
<td>3.0 (–0.3)</td>
</tr>
<tr>
<td>Hypothalamus region</td>
<td>1.9</td>
<td>0 6 –2</td>
<td>3.4 (–0.3)</td>
<td>3.4 (–0.4)</td>
</tr>
</tbody>
</table>

DS, Down syndrome; *Same cluster; * without 3 patients with mild cognitive impairment- MCI. x y z coordinates given in Montreal Neurological Institute (MNI) space. Statistics at corrected threshold \( P_{FWE} < 0.05 \) according to Monte Carlo simulations.
Fig. 1. Fractional anisotropy (FA) differences between Down syndrome and control subjects. Down syndrome patients showed a general pattern of lower FA in white matter, although the changes were more severe in the frontal-subcortical circuits (Top). By contrast, regions with significantly higher FA in Down syndrome patients than controls were limited to the temporo-parietal junction (Bottom). The right hemisphere corresponds to the right side of axial and coronal images.

Fig. 2. Significant positive correlation between semantic fluency and fractional anisotropy (FA) measurements in the direction of lower FA values, the poorer performance. The right hemisphere corresponds to the right side of the images.

**Age-related effect on fractional anisotropy**

In both groups, FA decreased as a function of age. Healthy controls showed FA age-related changes in the frontal lobes, corpus callosum, basal ganglia (caudate), thalamus, semioval centers, and pyramidal tracts (Fig. 3 and Table 2). In Down syndrome, significant correlations with age were found in the frontal lobes, left arcuate fasciculus, right external capsule, and hypothalamus (Fig. 3 and Table 2). We found
DISCUSSION

Our results indicate that white matter in Down syndrome patients showed generally lower FA compared with healthy participants. The most affected structures were the frontal lobes, subcortical white matter and some parts of the brainstem. Lower FA in Down syndrome was associated with poorer semantic fluency, which illustrates a degree of correspondence between white matter integrity and performance in patients. White matter FA did indeed decrease with age in both study groups, but we did not find the expected accelerated age effect in the Down syndrome group.

FA is a tissue measurement primarily independent of tissue volume that may generally express the extent to which an anatomical structure is composed of white matter tracts showing one dominant direction. In brain regions containing tracts with a single direction, FA increases as a result of brain maturation [4, 34]. In this context, our results indicate that Down syndrome subjects have a less developed white matter structure than controls in many white matter
tracts. This is a remarkable finding due to the few studies based on DTI in non-demented Down syndrome patients. Our results are consistent with studies of brain anatomy showing generally reduced white matter volumes in Down syndrome compared with healthy controls [13, 16]. However, our data may further contribute to characterizing white matter alterations in Down syndrome in that they would indicate that the changes are not limited to general volume reductions, but may also implicate a less developed structural connectivity pattern.

Although FA abnormalities involved the major brain pathways, the frontal-subcortical circuits showed more severe alterations. As a general trend, the results are consistent with the profile of cognitive deficits in Down syndrome typically progressing with deep impairment in language production and executive functions, which are cognitive domains notably dependent on the frontal lobes [35–37].

We also observed that Down syndrome individuals showed higher FA at the temporo-parietal junction. This observation is partly consistent with studies reporting relatively larger white matter volume in temporal and parietal regions [16, 20, 21], which may reflect a particular abnormality in white matter maturation. Indeed, in complex structures with tracts crossing in different directions (as in the temporo-parietal junction), higher FA may paradoxically denote less mature or less structured tissue [4, 34]. The multimodal temporo-parietal junction is certainly a complex area in terms of white matter connectivity. In Down syndrome, the pattern of connectivity would seem to be incomplete at this level. This event appears in parallel with a more general FA reduction in several other brain structures. In this general effect, the alteration may denote poor development of white matter pathways showing a predominant direction or a simpler structure.

A relevant problem in evaluating cognitive abilities in Down syndrome is the notable lack of consistency and reproducibility of ratings in some neuropsychological tests [38]. We used the K-BIT matrices subtest to estimate general intelligence, the digit span to evaluate working memory and semantic verbal fluency to test verbal output. Only verbal fluency was evaluated to test verbal output. Only verbal fluency was associated with FA alterations in Down syndrome in a rather general manner, thus indicating a certain parallelism between the identified pattern of white matter alterations and cognitive performance. Nevertheless, we failed to find similar associations with performance IQ and working memory. We now, a posteriori, consider that these tests are not perhaps the most suitable to reliably predict brain alterations in Down syndrome. The development of more specific and sensitive tools to evaluate cognition in Down syndrome, as recently proposed [39], could be helpful in future studies.

Our study did not find significant between-group differences in age-related white matter changes. That is to say, although both groups suffered a variation of white matter structure with age, there were no between-group differences. In our Down syndrome group, none of the participants were diagnosed with Alzheimer's disease and only 3 patients had MCI according to clinical criteria. Therefore, we were unable to demonstrate the anticipated premature aging effects on white matter using FA measurements, which contrast with positive findings in studies on familial Alzheimer’s disease (summarized in a recent report [40]). This brings us to the conclusion that DTI FA measurements are perhaps not sufficiently sensitive to capture brain pathology related to the acceleration of aging in subclinical populations. Alternatively, age-related white matter neurodegeneration may be a later event, which is not obvious prior to overt clinical dementia. In the study by Powell et al. [22], FA alterations were indeed more evident when dementia was clinically evident in Down syndrome patients.

It is important to mention, however, that the identified FA decrease with age in the control group is not the expression of the white matter structure involution or degeneration occurring in the senile brain (also expressed in the form of additional FA decrease [41]), as age in the control group showed a mean of 34.6 years and range 19–51 years. Age-related FA changes may better reflect the normal active evolution of white matter tracts in the adult brain. Our findings are consistent with previous studies in normal populations showing FA peaks from early 20 s to late 30 s and subsequent subtle FA decreases starting around mid-adulthood with notably different timings across different tracts [42]. FA studies in normal populations further emphasize early proposals that white matter remodeling is a biological process active over the entire lifespan [43, 44].

One challenge in the assessment of DTI is the control of head-motion effects on the measurement, which may be relevant in low-performance populations. We have considered this issue carefully and adopted several means to rigorously control such effects. We decided to exclude cases with detectable image degradation, as no correction procedure is
wholly efficient once the images have been acquired. The regular use of MRI practice sessions with mock scanners may minimize the problem in future studies. A post-hoc analysis on DTI using less rigorous exclusion criteria \((n=54)\) showed similar but weakened DTI results, indicating that the data obtained in the more selective sample \((n=45)\) was most probably not due to motion effects. Although accurate control of head motion effects may be a strength of the study, it is important to mention that strict participant selection is also a limitation. In this context, our findings cannot generalize to all Down syndrome population, but conclusions should be limited to relatively highly performing individuals. A final limitation relates to using a 1.5-T system, as opposed to a 3-T system with higher MRI signal.

**Conclusions**

Results from our imaging approach indicate that the brain in relatively high-functioning Down syndrome patients shows generally lower FA, suggesting underdevelopment of white matter tracts. The altered white matter structure was associated with poorer performance at neuropsychological assessment. Finally, age-related reduction of FA did not significantly differ between the control group and our non-demented Down syndrome patients. Probably, the adopted DTI approach was not sufficiently sensitive to detect alterations related to the dementia process in subclinical stages of Alzheimer-like pathology. Further studies should focus on the temporal evolution of white matter structure involution and the development of dementia in Down syndrome adults and provide biomarkers for detecting early signs of premature aging.

**ACKNOWLEDGMENTS**

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Authors’ disclosures available online (http://j-alz.com/manuscript-disclosures/161112r1).

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