World Trade Center Site Exposure Duration Is Associated with Hippocampal and Cerebral White Matter Neuroinflammation

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Abstract
Responders to the World Trade Center (WTC) attacks on 9/11/2001 inhaled toxic dust and experienced severe trauma for a prolonged period. Studies report that WTC site exposure duration is associated with peripheral inflammation and risk for developing early-onset dementia (EOD). Free Water Fraction (FWF) can serve as a biomarker for neuroinflammation by measuring in vivo movement of free water across neurons. The present case-controlled study aimed to examine associations between WTC site exposure duration as well as EOD status with increased hippocampal and cerebral neuroinflammation. Ninety-nine WTC responders (mean age of 56) were recruited between 2017 and 2019 (N = 48 with EOD and 51 cognitively unimpaired). Participants were matched on age, sex, occupation, race, education, and post-traumatic stress disorder (PTSD) status. Participants underwent neuroimaging using diffusion tensor imaging protocols for FWF extraction. Region of interest (ROI) analysis and correlational tractography explored topographical distributions of FWF associations. Apolipoprotein-e4 allele (APOE4) status was available for most responders (N = 91). Hippocampal FWF was significantly associated with WTC site exposure duration (r = 0.30, p = 0.003), as was cerebral white matter FWF (r = 0.20, p = 0.044). ROI analysis and correlational tractography identified regions within the limbic, frontal, and temporal lobes. Hippocampal FWF and its association with WTC exposure duration were highest when the APOE4 allele was present (r = 0.48, p = 0.039). Our findings demonstrate that prolonged WTC site exposure is associated with increased hippocampal and cerebral white matter neuroinflammation in WTC responders, possibly exacerbated by possession of the APOE4 allele.

Keywords World Trade Center · Exposure · Neuroinflammation · Free water fraction · Diffusion tensor imaging · Cognitive impairment · Early-onset dementia

Introduction
Alzheimer’s disease (AD) and other related dementias (ADRD) are among the most common causes of death in old age [1], with cognitive impairment (CI) as the most common early symptom [2, 3]. ADRD arises from neuropathology [4], including neuroinflammation [5], and is characterized by neurodegeneration in the frontal, parietal, and temporal lobes [6, 7], with the hippocampus as a centrally affected region [8]. Hippocampal degeneration in ADRD is often accompanied with cognitive decline such as hallmark memory disruptions [9, 10]. Other etiologies for ADRD often include risk factors such as older age (> 65 years), family history of dementia and/or APOE4 allele possession, and cardiovascular disease [1]. However, early-onset dementias (EOD) can be sporadic and can manifest with late-life dementia symptoms during midlife (40–65 years on average), potentially leading to more severe outcomes later [11]. Toxic exposures, such as repeated inhalations of nPM < 2.5 μm nano-particulate matter (nPM) pollutants, can also contribute to accelerated cognitive decline [12, 13], and are associated with neuroinflammatory and neurodegenerative processes [14–16]. Furthermore, APOE4 allele possession can compromise blood–brain barrier (BBB)
permeability and vulnerability to nPM infiltration from repeated exposures, leading to worse outcomes [13, 17].

During the attacks on the World Trade Center (WTC) on 9/11/2001, WTC responders, now at midlife, were involved in the search, rescue, and clean-up efforts and endured a multitude of exposures, such as repeated inhalations to toxic aerosolized nPM, such as smoke, dioxins, and polycyclic aromatic hydrocarbons [13, 18, 19]. Two decades since 9/11t1, approximately 14% of WTC responders are experiencing early-onset mild cognitive impairment (MCI) or EOD [20, 21], which is associated with prolonged WTC site exposure duration [18], and carrying the APOE4 allele [20, 22]. More specifically, we identified that WTC responders with EOD had reduced cerebellar [23] and cortical thickness in the frontal, parietal, and temporal lobes [24], while those with MCI had altered white matter integrity [25] and reduced hippocampal volume [26]. Additionally, chronic post-traumatic stress disorder (PTSD) had reduced cortical complexity [27], systemic inflammation [28] with monocytic signaling [29], and positron emission tomography (PET) translocator protein (TSPO) radioligand positive scans that confirmed the presence of activated glia throughout the brain [30]. Taken together, we interpret these findings at large as an inevitable consequence of their novel exposures at the WTC site, insofar that a substantial portion of WTC responders are now experiencing an emerging clinical condition based on an undetermined, but early-onset ADRD type of neurodegenerative process, with a neuroinflammatory basis [20, 23].

Free water fraction (FWF) is an MRI diffusion tensor imaging (DTI) technique that has been previously shown to estimate underlying neuroinflammation in vivo and its association with CI [31–33]. FWF can be derived from diffusion MRI data assuming a two-compartment model—extracellular water and tissue fraction [34–37]. A known byproduct of neuroinflammation is when the brain clears interstitial extraneuronal spaces resulting in localized changes to diffusing free water, which is what FWF measures [38–42]. Thus, there is increasing evidence demonstrating that FWF can be used as a marker for neuroinflammation [35, 41], and that it is often associated with CI [31–33]. Moreover, measuring FWF in the hippocampus can potentially serve as a sensitive biomarker of neuroinflammation for detecting ADRD, even in the absence of volumetric changes [43]. FWF is a superior measure for neuroinflammation, as other diffusion metrics such as fractional anisotropy or mean diffusivity are known to be sensitive to both neuroinflammation and other confounding pathology [38].

Considering that our prior work suggests associations between EOD, neuroinflammation, and WTC site exposure duration in this population, the primary aim of this study was to examine whether FWF in the hippocampus, cerebral white, or cerebral gray matter was associated with WTC site exposure duration. Our secondary aim was to examine whether responders with EOD displayed higher FWF in these regions when compared to cognitively unimpaired (CU) controls. Our third aim was to explore a region of interest (ROI) based topographic distribution of FWF levels to highlight which neuroanatomical regions display higher neuroinflammation.

**Methods**

**Population**

The WTC Health Program (WTCHP) was established in 2002 and is in charge of monitoring over 50,000 responders [44] via a protocol described in depth elsewhere [45]. As previously described [46], all WTC responders are eligible to attend annual health monitoring visits and receive treatment for WTC-certified conditions [47]. Eligible WTC responders continue to participate in an ongoing epidemiologic study of cognitive aging [48]. In the present report, WTC responders were contacted if consent had been previously collected at enrollment to permit for possible recruitment in future studies and if their patient characteristics matched necessary inclusion/exclusion criteria.

We employed a two-by-two study design with four groups including EOD (present/absent) and PTSD (present/absent). Inclusion criteria at the time of screening were ages 44 to 65, fluency in English, current cognitive assessment indicative of cognitive unimpaired (CU), EOD, or PTSD. While not an inclusion/exclusion criterion, all but one responder with CI had functional limitations consistent with mild dementia. Responders whose case status was not confirmed at during re-assessment were excluded from further participation in this study. To control for certain characteristic differences across groups, responder participants were matched on age, sex, race/ethnicity, occupation, and education at the time of screening.

Exclusion criteria included: history of psychosis, history of diagnosed neurological conditions, including major stroke, multiple sclerosis, and Parkinson’s disease; head injury during their WTC response efforts, history of military head trauma including combustive blasts; current liver disease; and current use of cognitively active medications. Subjects also had to satisfy eligibility criteria for MRI scanning including body mass index ≤ 40, no known claustrophobia, and no known metal implants or shrapnel that were not deemed MRI-safe.

In total, 598 WTC responders on file were initially contacted upon matching initial inclusion criteria with 176 of which were scheduled for screening visits after expressing participation interest. Approximately 88% of scheduled responders (n = 156) completed screening visits, wherein 27.6% (n = 43) were ineligible to continue and excluded.
with 7.7% (n = 13) declining further participation. Finally, 100 WTC responders participated in neuroimaging; with only 99 responders completing the scan protocol. The time between screening and the neuroimaging scan was 26 days on average (standard deviation [SD] = 17.2; interquartile range = 14–35 days).

Early-Onset Dementia

Dementia status was diagnosed algorithmically following guidelines by the National Institute on Aging and Alzheimer’s Association [49]. EOD was operationalized using the Montreal Cognitive Assessment (MoCA \( \leq 20 \)) [50], a widely used and validated measure of cognitive status developed to identify age-related CI objectively and reliably. CU status was operationalized when no evidence of CI (MoCA \( \geq 26 \)) was present.

Matching Criteria

PTSD status was operationalized using the Structured Clinical Interview for the DSM-IV [SCID-IV] [51], a semi-structured interview, administered by trained clinical interviewers. Severity for PTSD symptoms such as re-experiencing, avoidance, hyperarousal, and numbing was measured by examining subscale scores in a WTC specific version of the PTSD checklist (PCL17-S). Other matching criteria included age, sex, race/ethnicity (White, Black, Hispanic, Other), occupation (law enforcement versus other), and educational attainment (high school or less, some college, or university degree).

Image Acquisition

Three-dimensional T1-weighted magnetization-prepared rapid gradient echo (MPRAGE) images were acquired on a 3 T Siemens Biograph mMR TR/TE/TI = 1900/2.4/900 ms; flip angle = 9°; acquisition matrix = 256 \( \times \) 256; voxel resolution = 1 \( \times \) 1 \( \times \) 1 mm at Icahn School of Medicine, Mount Sinai, NY. For incidental neuropathology screening, T2-weighted anatomical scans used a turbo spin-echo pulse sequence (34 axial slices), TR/TE = 6170/96 ms; flip angle = 150°; acquisition matrix = 320 \( \times \) 320; voxel size = 0.36 \( \times \) 0.36 \( \times \) 3 mm and were read by a board-certified radiologist to determine any incidental neurological findings—no incidental findings were identified. DTI sequences were performed with the following parameters: TE/TR = 87.6/4680 ms, \( b \) value = 1000, 64 diffusion directions, in-plane resolution = 2 \( \times \) 2 mm\(^2\), slice thickness = 2 mm, matrix size = 128 \( \times \) 128, multiband factor = 2.

Image Processing

Following acquisition, DTI images were visually checked for major image or significant motion artifacts, with all 99 images passing inspection. Diffusion data from the DTI sequences were post-processed for each subject and involved motion and eddy correction [52], followed by generation of an FWF map. While the DiPy toolbox [53] has previously been utilized for multi-shell FWF generation [54–56], single-shell FWF images were generated using the recent extension to DiPy proposed by Golub et al. [57]. Our team utilized default initialization for FWF calculation with 200 iterations and a learning rate of 0.0005 as has been done previously [37] with the spatial regularization operator turned off at half iterations. Independent isotropic values (ISO), which measures the density of isotropically diffusing water calculated from q-sampling and is associated with FWF at lower \( b \)-values and represents the density of free water diffusion metrics along the subsection of a fiber pathway [37, 57–60], were calculated to examine subregional results using correlational tractography [61].

Correlational Tractography

Correlational tractography analysis was conducted to measure how ISO in white matter tracts correlated with WTC site exposure duration. Q-space diffeomorphic reconstruction was implemented in DSI-Studio to reconstruct subject data into MNI space and to then calculate a spin distribution function [62]. Nonparametric Spearman’s partial correlations were derived and a t-score threshold of 3 was operationalized as our threshold. Deterministic fiber tracking algorithms were utilized to obtain correlational tractography and whole brain seeding. Tracks were filtered by topology-informed pruning with 4 iterations using a length threshold of 20 voxels.

Region of Interest (ROI) Parcellation

Following linear and affine registration of each scan to a corresponding structural T1 reference image, ROI parcellation was conducted in FreeSurfer 7 using the Desikan Killiany (DK) atlas. A sample of the resulting hippocampal and white matter overlays from the registration pipeline can be seen in Fig. 1. The final output included 68 ROIs with FWF volumes that were estimated using FMRIB Software Library FLIRT tool [63].

WTC Exposure Duration

Upon enrollment into the larger WTC epidemiologic study, all responders undergo a baseline interviewer-administered exposure assessment questionnaire to record exposures to harmful physical and psychological conditions during their WTC recovery efforts [45]. In the present study, we examined the variable of time (hours) spent at the WTC recovery site [47, 64].
APOEε4 Status

To determine APOEε4 status, DNA was extracted from blood collected during visits during study baseline. Single nucleotide polymorphism genotyping was performed using the Agena iPLEX kit with processing completed at Roswell Park Laboratories. Among the 99 responders with imaging data available, the APOEε4 allele was genotyped in 92% of responders (n = 91, 70 APOEε4−, 21 APOEε4 +). APOEε4 + was operationalized when the responder possessed either the homozygous or heterozygous allele.

Statistical Analyses

Descriptive characteristics were provided using mean and standard deviations, or frequencies and percentages were noted. Welch’s t-tests and χ² tests examined differences in matching and diagnostic variables across study groups. A multi-factor analysis of variance (ANOVA) was used to study the association between EOD status, and WTC site exposure duration. Pearson’s correlation was used to study the association between WTC site exposure duration and FWF. Sensitivity analyses examined Spearman’s correlations between FWF and PTSD status, and PTSD symptom severity. Statistical significance was set at α = 0.05, with analyses reporting nominal and false discovery rate (FDR) adjusted p-values were noted [65].

Results

Sample Characteristics

As per our recruitment design, groups stratified by cognitive status did not differ on matching criteria, such as age, sex, race, PTSD status, education, or occupation (see Table 1) thereby negating the need to further control for these variables as potential confounders. Nevertheless, the results with age/sex/education correction are also provided.

FWF and EOD Status

EOD status was not significantly associated with hippocampal mean FWF (p = 0.656, Cohen’s d = −0.09),
nor cerebral gray matter FWF ($p = 0.668$, Cohen’s $d = 0.09$), with cerebral white matter FWF approaching significance ($p = 0.060$, Cohen’s $d = -0.38$). The results stayed the same after age/sex/education corrections; the corresponding $p$-values were 0.314, 0.088, and 0.833.

**FWF and WTC Site Exposure Duration**

Statistically significant correlations were identified between WTC site exposure duration and hippocampal and cerebral white matter FWF, but not cerebral gray matter (see Fig. 2).

**Cerebral White Matter ROI FWF and WTC Site Exposure Duration**

Cerebral white matter was parcellated into unilateral regions from the DK atlas and the mean FWF of several ROIs, including the parahippocampus ($r = 0.28$, $p = 0.005$), were found to be significantly correlated with WTC site exposure duration (hours) (see Table 2 and Fig. 3).

**Correlational Tractography**

Correlational tractography identified a positive association between ISO diffusion and WTC site exposure duration (hours) (T-threshold = 3, FDR = 0.011, with age/sex/education corrected), in ROIs that overlap with ROIs that were significant in Table 2 (see Fig. 3).

**APOEε4 status, FWF, and WTC Site Exposure Duration**

Spearman’s correlations between WTC site exposure duration (hours) and hippocampal and cerebral white and gray matter FWF were performed for the two subgroups with differing APOEε4 status, regardless of cognitive status. Cerebral white and gray matter FWF was not significantly associated with WTC site exposure duration (hours) (data not shown), but hippocampal FWF was, and the correlation coefficients were higher in the APOEε4+ group ($r = 0.48$, $p = 0.039$, $N = 21$) when compared to the APOEε4- group ($r = 0.25$, $p = 0.039$, $N = 70$).

**FWF and PTSD Status**

No significant associations between hippocampal and cerebral white and gray matter FWF were found when examining PTSD status (data not shown).

**Discussion**

As has been previously demonstrated by our team, the WTC population is at high risk of developing cognitive impairment through a neurodegenerative process [20]. We previously suggested that neuroinflammatory mechanisms might help to explain the association between WTC exposures and the risk of MCI in this cohort. The present study primarily...
aimed to test whether WTC site exposure duration, EOD status, and PTSD status, were associated with higher levels of the neuroinflammatory neuroimaging marker FWF, specifically in the hippocampus as well as the cerebral white and gray matter of WTC responders. In summary, we did not identify a significant association of FWF with hippocampal and cerebral white and gray matter FWF with EOD, nor PTSD status. However, we report here, for the first time, a significant correlation between hippocampal and cerebral white matter FWF with WTC site exposure duration that may be exacerbated by possession of the APOEε4 allele. Exploratory ROI-based analysis of cerebral white matter FWF further identified that parahippocampal and entorhinal white matter FWF had the highest correlation coefficients with WTC site exposure duration, followed by the transverse temporal which together with the prior two ROIs remained significant after age/sex/education adjustment, and the pars triangularis regions.

We did not identify a correlation between FWF in the hippocampus and WTC EOD status, which is in line with prior findings of how WTC site exposure was not correlated with MCI after controlling for APOEε4 status [47]. Hippocampal disruptions are a hallmark for patients with AD and certain related dementias [66], and our findings

| Lobe | Regions of Interest | r   | p    | r#   | P#  |
|------|---------------------|-----|------|------|-----|
| Frontal | Caudal Middle Frontal | 0.19 | 0.302 | 0.09 | 0.430 |
|       | Frontoal Pole        | -0.03 | 0.746 | -0.03 | 0.774 |
|       | Lateral Orbitofrontal | 0.15 | 0.152 | 0.15 | 0.122 |
|       | Medial Orbitofrontal | 0.16 | 0.109 | 0.13 | 0.177 |
|       | Pars Opercularis     | 0.22 | 0.028 | 0.17 | 0.082 |
|       | Pars Orbitalis       | 0.21 | 0.040 | 0.17 | 0.088 |
|       | Pars Triangularis    | 0.27 | 0.008* | 0.23 | 0.021 |
|       | Rostral Middle Frontal | 0.21 | 0.038 | 0.19 | 0.057 |
|       | Superior Frontal     | 0.15 | 0.139 | 0.12 | 0.232 |
|       | Precentral           | 0.23 | 0.026 | 0.15 | 0.135 |
| Limbic | Caudal Anterior Cingulate | 0.15 | 0.059 | 0.19 | 0.053 |
|       | Rostral Anterior Cingulate | 0.15 | 0.153 | 0.11 | 0.288 |
|       | Isthmus Cingulate    | 0.10 | 0.320 | 0.11 | 0.284 |
|       | Insula               | 0.18 | 0.078 | 0.16 | 0.099 |
|       | Parahippocampal      | 0.28 | 0.005* | 0.32 | <0.001 |
|       | Posterior Cingulate  | 0.13 | 0.217 | 0.10 | 0.290 |
| Temporal | Banks Superior Temporal Sulcus | 0.16 | 0.116 | 0.14 | 0.156 |
|       | Entorhinal           | 0.28 | 0.006* | 0.20 | 0.045 |
|       | Inferior Temporal    | 0.21 | 0.043 | 0.18 | 0.063 |
|       | Middle Temporal      | 0.18 | 0.071 | 0.18 | 0.062 |
|       | Superior Temporal    | 0.11 | 0.279 | 0.05 | 0.641 |
|       | Temporal Pole        | 0.27 | 0.007* | 0.24 | 0.014 |
|       | Superior Parietal    | 0.24 | 0.020 | 0.17 | 0.083 |
|       | Paracentral          | 0.18 | 0.181 | 0.09 | 0.343 |
|       | Postcentral          | 0.20 | 0.045 | 0.10 | 0.297 |
|       | Precuneus            | 0.18 | 0.079 | 0.13 | 0.198 |
|       | Superior Parietal    | 0.22 | 0.032 | 0.15 | 0.124 |
|       | Supramarginal        | 0.19 | 0.070 | 0.13 | 0.196 |
|       | Pericalcarine        | 0.06 | 0.533 | 0.02 | 0.826 |
|       | Fusiform             | 0.15 | 0.142 | 0.14 | 0.156 |
|       | Cuneus               | 0.14 | 0.161 | 0.08 | 0.430 |
|       | Lateral Occipital    | 0.08 | 0.435 | 0.04 | 0.725 |
|       | Lingual              | 0.13 | 0.223 | 0.11 | 0.284 |

Correlations between World Trade Center site exposure duration (hours) and cerebral white matter free water fraction in 68 regions of interest (ROIs) from the Desikan-Killiany Atlas. Significant correlations are highlighted in **Bold**; the * denotes ROIs that remained statistically significant following false discovery rate correction (FDR = 0.10); the # denotes age/sex/education corrected
here suggest that WTC responders with EOD are not as significantly affected by hippocampal disruptions as these patients. Further support is suggested from the association between cerebral white matter mean FWF and EOD, which approached significance. Taken together with the association of cerebral FWF with WTC site duration, a larger sample size may reveal that cerebral white matter FWF is also associated with WTC responder EOD status.

Extensive literature has detailed the sensitivity and involvement of hippocampal changes to cognitive decline [9, 26], both through aging and disease progression. In this work, even though hippocampal FWF was not associated with EOD status, a statistically significant positive correlation was found between hippocampal FWF and WTC site exposure duration. Furthermore, this finding is in line with the higher hippocampal glial activation we previously found in WTC responders with MCI [67]. Taken together, this suggests that responders with higher WTC site exposure duration may display more severe hippocampal neuroinflammation, possibly due to prolonged exposures to nPM inhalations at the WTC disaster site [13, 19]. For example, among the nPM present at Ground Zero, thallium—a toxic heavy metal capable of targeting and damaging the nervous system—was existent in concentrations 1,000 times higher than evident in unexposed parts of New York City [68, 69].

Cerebral white matter FWF and WTC site exposure duration were also found to significantly correlated, with several ROIs involved, mostly within the temporal and limbic areas such as the transverse temporal, entorhinal, and parahippocampal formation. Moreover, clusters of correlations with WTC exposure duration were also found in ROIs within the parietal and frontal lobes. The highest correlation was in the pars triangularis, which remained significant after age/sex/education adjustment, followed by the pars orbitalis and opercularis, the precentral, and then the postcentral, inferior, and superior parietal. These ROIs are in line with cortical thinning previously identified in WTC responders with EOD [24], and our work here now suggests that neuroinflammation associated with prolonged WTC site exposure duration may have been a mechanism for neurodegeneration. Taken together, associations of WTC exposure duration with FWF in key cerebral ROIs suggest that prolonged durations at the site may have led to widespread neuroinflammatory activation and downstream outcomes. This finding is also in line with prior literature suggesting toxic effects of nPM on the BBB [70, 71], as well as neuroinflammatory consequences of nPM inhalations [15], which may be placing affected WTC responders with higher WTC exposure duration at increased risk for developing neuropathological phenomena [13].

APOEε4 is known to increase the relative risk for CI and dementia, and we identified that APOEε4 status was associated with hippocampal FWF and WTC site exposure duration, with nearly twice the effect size in APOEε4+ responders (APOEε4+: r = 0.48 versus APOEε4−: r = 0.25). This suggests that the presence of the APOEε4 allele may moderate the correlation between hippocampal FWF and WTC site exposure and is consistent with prior work showing that incidence of MCI was higher among APOEε4+ WTC responders with prolonged WTC exposures [47]. This is in line with studies demonstrating that APOEε4 allele possession can compromise blood–brain barrier (BBB) permeability and vulnerability.
to nPM infiltration from repeated exposures, leading to worse outcomes [13, 17].

WTC responders are known to suffer from chronic PTSD [72], which is also a risk factor for cognitive dysfunction [20]. Approximately half of our study responder participants herein had clinically significant levels of PTSD and related symptoms. However, we found no associations between hippocampal and cerebral white and gray matter FWF and PTSD, suggesting that the effects of PTSD may engage different risk mechanisms for CI in those affected. Indeed, several other studies with WTC responders have similarly shown that PTSD was not associated with degree or type of neurodegeneration after accounting for CI status [23, 24, 26]. Therefore, our present findings suggest that FWF could be an in vivo neuroinflammatory marker for WTC exposure site duration independent of PTSD status, and possibly moderated by APOEε4 status.

Limitations

In this work, we did not have APOEε4 status for all responders and the distribution of APOEε4 ± responders was imbalanced (n = 21 versus n = 70). However, we observed that APOEε4 status may moderate the association between hippocampal neuroinflammation and WTC site exposure. Nevertheless, a future prospective investigation controlling for APOEε4 status is needed to validate this finding. Another limitation of this study are the mathematical limitations of FWF for single-shell diffusion data as prior diffusion studies have suggested to generate FWF images using multi-shell data for the fitting of the two-compartmental bi-tensor model instead [35, 73]. However, other studies have demonstrated reliability across both single-shell and multi-shell algorithms [37, 74], but care is needed to interpret the data [37]. Finally, the present study did not test for network-based structural correlations with WTC site exposure duration, as outlined in this study [75], and although we have performed some diffusion-based studies in prior publications [27, 76], future neuroimaging studies of WTC responders should consider these approaches.

Conclusions

This study identified that hippocampal and cerebral white matter neuroinflammation, as measured by FWF derived from DTI-MRI, is associated with WTC site exposure duration for a sample of 99 WTC responders with and without EOD, regardless of PTSD status. Several key white matter ROIs were involved, suggesting a widespread neuroinflammatory process that may mechanistically underlie our prior findings of gray matter neurodegeneration in those same regions. WTC site exposure-related prolonged inhalations of nPM may be responsible for this neuroinflammation that seems to be independent of PTSD and may be exacerbated by possessing the APOEε4 allele. Further investigation is needed to validate and elaborate on the findings reported herein.

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Data Availability

Medical information for these participants is protected, so only processed de-identified data will be made available upon receipt of reasonable request to the corresponding author.

Declarations

Ethics Approval

The Institutional Review Boards at both Stony Brook University and Icahn School of Medicine at Mount Sinai approved study procedures.

Consent to Participate

All participants in this study provided informed written consent.

Consent for Publication

Not applicable.

Competing Interests

The authors declare no competing interests.

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