A permethrin metabolite is associated with adaptive immune responses in Gulf War Illness


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ABSTRACT

Gulf War Illness (GWI), affecting approximately 30% of veterans from the 1991 Gulf War (GW), is a multi-symptom illness with features similar to those of patients with autoimmune diseases. The objective of the current work is to determine if exposure to GW-related pesticides, such as permethrin (PER), activates peripheral and central nervous system (CNS) adaptive immune responses. In the current study, we focused on a PER metabolite, 3-phenoxybenzoic acid (3-PBA), as this is a common metabolite previously shown to form adducts with endogenous proteins. We observed the presence of 3-PBA and 3-PBA modified lysine of protein peptides in the brain, blood and liver of pyridostigmine bromide (PB) and PER (PB+PER) exposed mice at acute and chronic post-exposure timepoints. We tested whether 3-PBA-haptenated albumin (3-PBA-albumin) can activate immune cells since it is known that chemically haptenated proteins can stimulate immune responses. We detected autoantibodies against 3-PBA-albumin in plasma from PB+PER exposed mice and veterans with GWI at chronic post-exposure timepoints. We also observed that in vitro treatment of blood with 3-PBA-albumin resulted in the activation of B- and T-helper lymphocytes and that these immune cells were also increased in blood of PB+PER exposed mice and veterans with GWI. These immune changes corresponded with elevated levels of infiltrating monocytes in the brain and blood of PB+PER exposed mice which coincided with alterations in the markers of blood-brain barrier disruption, brain macrophages and neuroinflammation. These studies suggest that pesticide exposure associated with GWI may have resulted in the activation of the peripheral and CNS adaptive immune responses, possibly contributing to an autoimmune-type phenotype in veterans with GWI.

1. Introduction

Gulf War Illness (GWI) affects approximately 30% of the US military personnel who were deployed to the 1991 Gulf War (GW) (White et al., 2016). Nearly 30 years later, GW veterans continue to experience debilitating symptoms such as pain, fatigue and memory problems. Clinical studies suggest that immune dysfunction and chronic inflammation are among the key causes of the symptoms experienced by ill GW veterans (White et al., 2016). Studies using the rodent models of GWI implicate both the central nervous system (CNS) and peripheral immune system in contributing to the pathogenesis of this illness (Koo et al., 2018). However, a better understanding of these immune
Elevated proinflammatory cytokines can increase matrix served in the brains of GWI rodent models (Abdullah et al., 2011, 2012, phosphorylation and proinflammatory cytokines are commonly ob-
erved in the brains of GWI rodent models (Abdullah et al., 2011, 2012, 2016; Parihar et al., 2013; Zakirova et al., 2015; Joshi et al., 2018). Elevated proinflammatory cytokines can increase matrix metalloproteinase 9 (MMP9) activity at the BBB and enhance the transmigration process, potentially promoting the infiltration of peripheral immune cells into the CNS (Song et al., 2015). Among these, Ly6C+ monocytes are capable of converting to macrophages upon CNS infiltration (Wohleb et al., 2013; Miró-Mur et al., 2016). Increases in C-C chemokine receptor type 2 (CCR2) and its ligand, CCL2, were ob-
served in the brains of a GWI mouse model of combined PB and PER (PB + PER) exposure (Joshi et al., 2018). In neuroinflammatory conditions, CCL2 released by astroglia recruits CCR2-expressing peripheral monocytes into the parenchyma, leading to the activation of NFκB-mediated inflammatory pathways (Gerard and Rollins, 2001; Jimenez et al., 2010; Bose and Cho, 2013; Harrison-Brown, Liu and Banati, 2016; Parillaud et al., 2017). These studies suggest that peripheral adaptive immune responses could be involved in promoting neuroinflammation in GWI.

We hypothesize that exposure to PER results in 3-PBA-haptenation of proteins and that haptenated proteins presented by antigen-presenting cells (APC) could activate T- and B-cells, resulting in autoimmune production. The presence of haptenated proteins in the CNS may promote infiltration of peripheral immune cells, which would further contribute to neuroinflammation. In this study, we determined the presence of 3-PBA modified lysine (3-PBA-Lys) in proteins and 3-PBA recovered from protein hydrolysis in the brains and livers of a PB + PER GWI mouse model. We examined the presence of autoantibodies against 3-PBA-haptenated albumin in plasma from PB + PER exposed mice and in veterans with GWI. We investigated the ability of 3-PBA-haptenated albumin to activate peripheral T- and B-cells and determined that these immune cells are elevated in the blood of PB + PER exposed mice and in veterans with GWI. We determined that proinflammatory monocytes capable of tissue infiltration are present in the blood of PB + PER exposed mice. We also examined infiltrating monocytes and macrophages derived from these monocytes in the brains of PB + PER exposed mice. Collectively, these studies will help determine the role of PER metabolites in activating adaptive immune responses in GWI.

2. Material and methods

2.1. Human subjects

Plasma and/or peripheral blood mononuclear cells (PBMC) were utilized from 3 different cohorts: (1) GWI and GW control veterans recruited at the Roskamp Institute Clinics; (2) GWI cases and GW controls from the Boston Gulf War Illness Consortium (GWIC); and (3) subjects from Thailand who were farm workers or consumed pyre-
throid-infested fruits and vegetables and either had positive or negative urine 3-PBA measurements. Plasma from age- and gender-matched GWI and control GW veterans were used for this study. The Kansas GWI criteria (Steele, 2000) were used to diagnose GWI. The Kansas GWI criteria require that GW veterans show symptoms in at least 3 of 6 symptom domains (fatigue/sleep problems, somatic pain, neurological/ cognitive/mood symptoms, gastrointestinal symptoms, respiratory symptoms, and skin abnormalities). Study participants were excluded if they reported being diagnosed with another medical condition that could explain their chronic health symptoms. Sample collection at the Roskamp Institute Clinics was approved by the Western Institutional Review Board (WIRB), and all protocols were conducted in accordance with relevant guidelines and regulations. Recruitment of study subjects and blood collection to obtain plasma and PBMC has been described previously (Emmerich et al., 2017; Qiang et al., 2017; Joshi et al., 2018) and demographic details are provided in Table 1A. Subjects were also excluded from the study if they were currently on any steroid medication to minimize interferences from the effects of these medications at the time of the blood draw. Controls were veterans deployed to the 1991 GW who did not meet the Kansas GWI criteria or any of the exclusionary criteria listed above. Plasma and PBMC collection at the Boston GWIC were approved by the Boston University IRB. As described
above, the Kansas GWI criteria (Steele, 2000) were used to diagnose GWI, exclusion criteria are the same as those described for the cohort above. Demographic details for GWIC samples are provided in Table 1B. Self-report of pesticide exposure was based on questions which inquired whether they had worn a uniform saturated with pesticide and/or used pesticide cream/spray. Using these data, a summary variable was generated which assigned yes/no to pesticide exposure. Since the exposure status of GW veterans is based on self-report, we also examined plasma from samples obtained from farm and non-farm workers with documented exposure to pyrethroids and measurements of 3-PBA in urine. These were occupational health worker who were exposed to different pyrethroid pesticides during their occupation, and were used as positive and negative controls for a 3-PBA-albumin autoantibody detection assay. All the participants gave their written consents and the study was approved by their respective IRBs. A detailed demographic table is provided in Table 2. Many of these age- and gender-matched people had been exposed to pyrethroids during their occupation, and half of them had 3-PBA detected in their urine (n = 5 positive and n = 5 negative).

### Table 1A
Demographics of the Gulf War veteran autoantibody cohort.

<table>
<thead>
<tr>
<th></th>
<th>Controls (GW veteran)</th>
<th>GWI cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>N total</td>
<td>10</td>
<td>14</td>
</tr>
<tr>
<td>Age (Mean ± SEM)</td>
<td>54.3 ± 7.6</td>
<td>52.4 ± 7.5</td>
</tr>
<tr>
<td>Male (%)</td>
<td>79.7%</td>
<td>86%</td>
</tr>
<tr>
<td>Self-report of pesticide exposure during war</td>
<td>95% (12)</td>
<td>95% (12)</td>
</tr>
<tr>
<td>Self-report of PB exposure during war</td>
<td>60% (6)</td>
<td>100% (14)</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>8 (70%)</td>
<td>11 (78.4%)</td>
</tr>
<tr>
<td>African American</td>
<td>2 (14.2%)</td>
<td>2 (14.2%)</td>
</tr>
<tr>
<td>Other/Multiracial</td>
<td>0</td>
<td>1 (7.13%)</td>
</tr>
</tbody>
</table>

There was no significant difference in age among two groups.

### Table 1B
Demographics of the Gulf War veteran Boston consortium cohort.

<table>
<thead>
<tr>
<th></th>
<th>Controls (GW veteran)</th>
<th>GWI cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>N total</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>Age (Mean ± SEM)</td>
<td>51.23 ± 8.56</td>
<td>54.3 ± 7.4</td>
</tr>
<tr>
<td>Male (%)</td>
<td>83.33%</td>
<td>100%</td>
</tr>
<tr>
<td>Self-report of pesticide exposure during war</td>
<td>25% (3)</td>
<td>83.33% (10)</td>
</tr>
<tr>
<td>Self-report of PB exposure</td>
<td>83.33% (10)</td>
<td>100% (12)</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>9 (75%)</td>
<td>10 (83.4%)</td>
</tr>
<tr>
<td>African American</td>
<td>2 (16.67%)</td>
<td>1 (8.3%)</td>
</tr>
<tr>
<td>Other/Multiracial</td>
<td>1</td>
<td>1 (8.3%)</td>
</tr>
</tbody>
</table>

There was no significant difference in age among two groups.

### Table 2
Demographics of the subjects from Thailand.

<table>
<thead>
<tr>
<th></th>
<th>Negative Controls (no urine 3PBA detected)</th>
<th>Positive cases (urine 3-PBA detected)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N total</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Age (Mean ± SEM)</td>
<td>57.3 ± 7.6</td>
<td>61.4 ± 6.33</td>
</tr>
<tr>
<td>Male (%)</td>
<td>80%</td>
<td>80%</td>
</tr>
<tr>
<td>Pesticide exposure</td>
<td>20% (1)</td>
<td>66.6% (3)</td>
</tr>
<tr>
<td>PB exposure</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Occupation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Farm worker</td>
<td>60% (3)</td>
<td>60% (3)</td>
</tr>
<tr>
<td>Non-farm worker</td>
<td>40% (2)</td>
<td>40% (2)</td>
</tr>
</tbody>
</table>

There was no significant difference in age among two groups.

#### 2.2. Animals

All procedures on mice were approved by the Roskamp Institute’s Institutional Animal Care and Use Committee (IACUC) and were conducted in compliance with the Office of Laboratory Animal Welfare and laboratory animal care guidelines as previously described (Abdullah et al., 2011; Zakirova et al., 2015). Mice were maintained with a 12 h light/dark cycle in a temperature-controlled environment (21 ± 2°C). Male C57BL/6 mice (3 months of age, weight 25 g ± 0.7 SD) were co-administered 0.7 mg/kg of pyridostigmine bromide (PB) (Fisher Scientific Hanover Park, IL) and 200 mg/kg of PER (Sigma Aldrich, St. Louis, MO, USA) in a single 50 μL intraperitoneal (i.p.) injection in dimethyl sulfoxide (DMSO) (Sigma Aldrich, St. Louis, MO, USA) or DMSO alone (control) daily for 10 consecutive days. Mice were divided in two groups with the control group receiving vehicle only (DMSO) and the GWI group receiving vehicle along with PB and PER. A timeline of the study procedures is provided as Fig. 1.

#### 2.3. Detection and quantification of 3-PBA using LC/MS reverse-phase analysis:

Levels of 3-PBA in plasma, brain and liver were quantified as described by Ahn et al. (2011). Briefly, 400 mg of liver and 100 mg of brain from both control and PB + PER exposed mice were homogenized in 1 mL of PBS, and a protein crash was then performed via the addition of 4 mL of acetone. Samples were then subjected to acid hydrolysis by
1 mL 6 M HCl at 100 °C for 2 h to hydrolyze covalent bonds between 3-PBA and proteins (Ahn et al., 2011). These samples were subjected to solid-phase extraction (SPE) using BB-S100-UBr Statta™ X 33 µm polymeric reversed phase, 60 mg/3 mL spin columns (Phenomenex, Torrance, CA, USA) for clean-up and spiked with 10 µM phenoxy-13C6-polymeric reversed phase, 60 mg/3 mL spin columns (Phenomenex, Torrance, CA, USA). These samples were subjected to 1 mL 6 M HCL at 100°C for 2 h to hydrolyze covalent bonds between 3-U.

A gradient was run from 90% solvent A in 10% solvent B to 50% B in 6 min with a 4 min hold at 100% solvent B with a flow rate of 100 µL/min and the column temperature was set at 40°C. Mass spectrometry analyses were conducted with high mass accuracy and parallel reaction monitoring (PRM) using a Q-Exactive (QE) hybrid quadrupole-Orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA). Negative ion electrospray spectra were acquired at 17,500 resolution with a scan range from m/z 150 to 600. All spectra were obtained with a 100 ms maximum ion injection time and 5 microscans. The 3-PBA IS (Cambridge Isotope Laboratories, Tewksbury, MA, USA) was monitored with m/z 219.1 as the precursor mass and product ions at m/z 99.053 and 175.084. For 3-PBA, we used m/z 213.1 as the precursor ion and m/z 93.033 and 169.065 as product ions.

2.4. Quantification of 3-PBA peptide using LC/MS reverse-phase analysis

Levels of 3-PBA bound lysine in plasma, brain and liver were quantified as described by Noort et al. (2008). Briefly, 400 mg of liver and 100 mg of brain from both control and PB + PER exposed mice were homogenized in 1 mL of PBS, and protein was precipitated by adding 4 mL of acetone at −20 °C for 3 hrs. Precipitated protein was digested with 100 µL of pronase solution (10 mg/mL, 50 mM NH4HCO3) for 1 h at 37°C followed by analysis with LC-tandem MS as explained above. In-house synthesis of 3-PBA-albumin (described below) was as standard. The 3-PBA-lysine was monitored with m/z 343.12 as the precursor mass and product ions at m/z 197.06.

2.5. Detection of 3-phenoxybenzoic acid in urine using GC-ECD

The first urine samples of the morning were collected and frozen at −20 °C until they were analyzed for 3-PBA which was detected as described by Pakvila et al. (2014). Briefly, 50 µL of labeled 3-PBA (1 µg/mL), as internal standard, was added to 2 mL of urine sample, then mixed for 30 s. The urine samples were subjected to acid hydrolysis by adding 500 µL of concentrated hydrochloric acid (37%) at 100 °C for 2 h. Each sample was extracted twice with 2 mL of ethyl acetate (EA), followed by centrifugation at 2500 rpm for 5 min. The organic phase was separated, and sample cleanup was then performed on pre-conditioned C18– cartridges. The resulting solution was dried and re-dissolved in acetonitrile. The 3-PBA was analyzed by using a Hewlett-Packard 7890B-electron capture detector (GC-ECD) and Autosampler G4513A (Agilent Technology, CA, USA) equipped with HP-5 (5% phenylmethyl polysiloxane with 30 m x 0.25 mm, 0.25 µm film thickness) after derivatization by 1,1,1,3,3,3-Hexafluoroisopropanol (HFIP) and N,N’-Disopropylcarbodiimide (DIC).

2.6. Preparation of 3-PBA-albumin conjugate

It has been shown that the 3-PBA metabolite of PER forms covalent adducts with albumin, and, due to its high abundance, albumin is often the major target of drug haptenation (Grigoryan et al., 2009; Tailor et al., 2016). We chose albumin to test our hypothesis that 3-PBA can also haptenate protein (3-PBA-albumin) and stimulate immune responses. We prepared a 3-PBA-albumin conjugate as described by Ahn et al. (2007). Briefly, 65 mg of 3-PBA (Sigma Aldrich, St. Louis, MO, USA) and 92 mg 1,3-dicyclohexylcarbodiimide (DCC) (Sigma Aldrich, St. Louis, MO, USA) was dissolved in 4 mL of N,N-Dimethylformamide (DMF) (Sigma Aldrich, St. Louis, MO, USA). The active ester in 3-PBA was formed by adding 52 mg of N-hydroxysuccinimide (NHS) (Thermo Fisher Scientific, Waltham, MA USA) and allowed to react for 5 hrs at room temperature, after which 10 mL of 50 m M bovine or human serum albumin in pH 8 borate buffer was added to the solution and allowed to react overnight. The resulting precipitate containing 3-PBA-albumin was collected and dialyzed (12,000 Da) against PBS for 3 days to remove low molecular weight contaminates. For mouse studies (including blood culture studies), bovine albumin was used as previous studies have shown that mouse and bovine serum albumin (BSA) share 72% amino acid homology, and BSA is immunologically inactive in mice (Di Domizio, Dorta-Estremera and Cao, 2013). For human studies, human albumin was used.

2.7. 3-PBA-albumin autoantibody detection in the plasma

Both free albumin and 3-PBA-albumin were subjected to sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis in 4–20% Criterion™ TGX Stain-Free™ Precast Gels (Bio-Rad, Hercules, CA, USA). These proteins were then transferred to a polyvinylidene difluoride (PVDF) membrane for 4 h at 60 mA. After blocking in 5% milk for 1 h, PB + PER mouse plasma and control mouse plasma were used as a source of antibody. The membrane was incubated in PB + PER mice or control mice plasma (1:300) overnight at 4°C. Membranes were then incubated in HRP-linked mouse secondary anti-IgG antibody, and the blot was developed using SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Fisher Scientific Waltham, MA USA) and visualized using a chemiluminescence imager (Bio-Rad, Hercules, CA, USA) as described previously (Joshi et al., 2018). For human samples, the same procedure was used with the exception that human secondary anti-IgG was used.

2.8. Ex vivo whole blood flow cytometry studies

To test our hypothesis that 3-PBA albumin can stimulate immune response and activate T- and B-lymphocytes, blood from 7-months old control mice (C57BL/6) was collected following a cardiac puncture. Blood was then combined with 2% EDTA and cultured in RPMI media (Gibco, Carlsbad, CA, USA) at 37 °C in a humidified atmosphere containing 5% CO2 supplemented with 1% of penicillin-streptomycin solution and 10% FBS. Cultured blood was then treated with 5 µM 3-PBA-albumin, 5 µM 3-PBA and 5 µM albumin for 12 h followed by red blood cell (RBC) lysis and stained with antibodies against cell surface markers: CD4 (Thermo Fisher Scientific, Waltham, MA, USA), CD8 (Thermo Fisher Scientific, Waltham, MA, USA), CD3 (Thermo Fisher Scientific, Waltham, MA, USA), CD19 (Thermo Fisher Scientific, Waltham, MA, USA), CD27 (Thermo Fisher Scientific, Waltham, MA, USA) and CD45R (B220) (Thermo Fisher Scientific, Waltham, MA, USA). Flow cytometry analyses were done using Attune® NxT Acoustic Focusing Flow Cytometry (Thermo Fisher Scientific, Waltham, MA, USA). Flow cytometry data were presented as total percentage change, and graphs were generated using GraphPad Prism7. CD4 + CD3 + staining was used to identify T-helper cells while CD3 + CD8 + staining was used to identify cytotoxic T-cells (Alb Pers and Johnson, 2002). CD19 + and CD27 + were used to distinguish memory and naïve B-cell population (Santer et al., 2013).

2.9. Detection of mouse blood monocytes and lymphocytes (B- and T-cells) using flow cytometry

Whole blood was drawn from mice, and 100 µL was subjected to red blood cell (RBC) lysis using 1 mL of RBC lysis buffer (Thermo Fisher
Germany) according to the manufacturer's protocol. Briefly, cells were then isolated by magnetically activated cell sorting (MACS), using anti-CD45 (Biolegend, San Diego, USA) and CD11b (BD Bioscience, NJ, USA) antibodies for flow cytometry analysis. Details on the antibodies used are provided in Supplementary Table S5. Monocytes were identified by gating cells that were positive for both CD11b and CD115 (Randolph et al., 2008; Greter et al., 2015). Patrolling and infiltrating monocyte populations were identified by CD11b+ CD115+ with Ly6C- and CD11b+ CD115+ with Ly6C+, respectively (Chao et al., 2013). Ly6C+ monocytes are defined as proinflammatory and tissue-infiltrating while Ly6C+ are phagocytic monocytes (Randolph et al., 2008). Flow cytometry analysis was conducted using combinations of these monoclonal antibody using Attune® NxT Acoustic Focusing Flow Cytometer (Thermo Fisher Scientific, Waltham, MA USA).

In another experiment, 150 μL of whole blood from the same mice were lysed via addition of 1500 μL of RBC lysis buffer (Thermo Fisher Scientific, Waltham, MA, USA). The monocytes and lymphocytes were pelleted by centrifugation at 700 × g for 10 min and resuspended in 500 μL of FACS buffer (PBS, 0.5–1% BSA or 5–10% FBS, 0.1%), to which 10 μL of CD4/CD8 antibody cocktail (Thermo Fisher Scientific, Waltham, MA USA) was added. Cells were incubated with the cocktail for 20 min at room temperature. Lymphocyte populations were gated, and CD4+ and CD8+ cells were analyzed using Attune® NxT Acoustic Focusing Flow Cytometer (Thermo Fisher Scientific, Waltham, MA USA). CD3+ CD4+ were used to identify T-helper cells while CD3+ CD8+ cells are used to identify cytotoxic T-cells. These cells were analyzed as described above. For B-lymphocytes, a combination of CD27, CD19 and CD45R (B220) stains were used as described previously; detail on antibodies used is provided above. As described previously, these were used to distinguish memory and naive B-cell population (Santer et al., 2013).

2.10. Labeling of human PBMC for T- and B-lymphocytes

The peripheral mononuclear cells provided by GWIC (Boston) were treated with monoclonal antibodies conjugated to different fluorophores. To find the T-cell population, cells were incubated with Anti-human CD3/CD4/CD8 Cocktail (Biolegend, San Diego, CA USA). To find the memory B-cell population, a combination of CD19 (Thermo Fisher Scientific, Waltham, MA, USA) and CD27 (Thermo Fisher Scientific, Waltham, MA, USA) antibodies were used and analyzed using Attune® NxT Flow Cytometer as described above.

2.11. Flow cytometry studies of the brain myeloid and microglia cells

A single cell suspension was generated from adult mouse brains, using the adult brain dissociation kit from Miltenyi Biotec (Cologne, Germany) and closely following the manufacturer instructions. Briefly, mouse brains were collected in serum-free media and chopped finely with a sterile scalpel blade. The brains were subjected to a combination of enzymatic Papain (1 μg/mL) mechanical digestion (pipetting up and down 20 times) to yield a single cell suspension, the brain homogenate was then passed through a 70 μm strainer, and the cell suspension was centrifuged for 10 min at 1000 × g and 4°C. Cell pellets were then suspended in debris removal solution and carefully overlaid with 4 mL D-PBS as per manufacturer instructions; the samples were then centrifuged at 3000 × g for 10 min to generate a pellet containing a mixed glial cell population. The myeloid populations were then isolated by magnetically activated cell sorting (MACS), using anti-CD11b conjugated magnetic microbeads (Miltenyi Biotec, Cologne, Germany) according to the manufacturer's protocol. Briefly, cells were incubated with 10 μL CD11b coated microbeads for 15 min at 4°C. Cells were then washed and placed on a new LS magnetic column (Miltenyi Biotec, Cologne, Germany) in a magnetic holder. The CD11b negative fraction was washed through the column, and the CD11b positive cell fraction was eluted from the column by removing the column from the magnetic field, adding buffer and emptying the column with a plunger. Cells were incubated with Rat anti-mouse Fe-block (1:100 BD biobioce) for 5 min at room temperature. These cells were then stained with monocolonal antibodies for desired surface markers. The antibodies used were CD11b (BioLegend San Diego, USA), CX3CR1 (BioLegend, San Diego, USA), Ly6c (BioLegend San Diego, USA) and CD206 (BioLegend San Diego, USA). Details on the antibodies used are provided in the Supplementary Table S6. These cells were analyzed as described above using an Attune® NxT Acoustic Focusing Flow Cytometer (Thermo Fisher Scientific, Waltham, MA USA). A combination of CD11b+, CX3CR1+, CD206+, and CD6+ staining was used to analyze brain macrophages, and cells stained with CD11b+, CX3CR1+, and CD206− were classified as microglia (Han et al., 2015; Wong et al., 2017; Zhou et al., 2017; Zarruk et al., 2018).

2.12. Immunostaining for microglia and astrocytes

The left-brain hemispheres were fixed in 4% paraformaldehyde (PFA), dehydrated and embedded in paraffin. Sagittal sections (8 μm) were prepared using Leica RM2235 microtome (Leica, Buffalo grove, IL). Approximately 8 to 10 slices of each mouse brain were incubated in blocking buffer (10% donkey serum and 0.3% Triton X-100 in PBS) for 60 min at room temperature. Samples were then incubated overnight with the primary antibody Rabbit anti-IBA-1 (Wako, Osaka, Japan) and Chicken anti-GFAP (Aves Lab, Tigard, Oregon) at 4°C. The following day, slices were incubated with the appropriate fluorescent probe-conjugated secondary antibodies for 1 hr at room temperature. Nuclei were counterstained with DAPI. The entire DG was scanned using an Axiovert 200 M Zeiss microscope (Zeiss) equipped with AxiosVision 4.8.3.0 software. ImageJ software was used to analyze the stained sagittal sections. The stained areas were calculated and expressed as a percentage of the field. Further detail on primary and secondary antibodies used for brain pathology is provided in Supplementary Table S7.

2.13. Enzyme-linked immunosorbent assay

Using a Dounce homogenizer, brains were homogenized in ice cold Mammalian Protein Extraction (mPER) buffer (Thermo Fisher Scientific, Waltham, MA USA), which contained protease (Roche, Indianapolis, IN, USA) and phosphatase inhibitor (Pierce, Grand Island, NY, USA) cocktails. These whole brain homogenates were centrifuged at 10,000 × g for 10 min, and supernatant was used for ELISA and Western blot. The total protein content of each sample was determined by the bicinchoninic acid (BCA) assay (Pierce, Grand Island, NY, USA). Enzyme-Linked Immunosorbent Assay (ELISA) kits for mouse CCR2 (Lifespan Biosciences, Seattle, WA,USA), CCL2 (Lifespan Biosciences, Seattle, WA, USA), Matrix Metalloproteinase-9 (MMP9) (Bosterbio, Pleasanton, CA, USA) and fractalkine (CX3CL1) protein (MyBiosource, San Diego, CA,USA) were used to study levels of these proteins in the brain using the manufacturer’s protocol. Soluble CX3CL1 (Thermo Fisher Scientific, Waltham, MA USA) in human plasma were quantified using a commercial ELISA kit. The level of IgG protein in blood was quantified by IgG ELISA (Cayan Chemical, Ann Arbor, MI, USA) according to the manufacturer’s instruction. MMP9 is a proteolytic protein capable of degrading tight junction protein leading to BBB impairment (Underly et al., 2016; Schactae et al., 2017). Results were normalized to total protein quantification obtained from BCA. Intra-Assay: CV < 10.6% Inter-Assay: CV < 12.6% for all the ELISAs. There was no reported cross-reactivity with other proteins for the primary antibody used in these kits.
2.14. Western blot analyses

CX3CR1 (Abcam, Cambridge, MA, USA), p-65 (p-NFκB) phosphorylated at Ser536 (Cell Signaling, Danvers, MA, USA), total p65 (NFκB) (Cell Signaling, Danvers, MA, USA), tubulin (Abcam, Cambridge, MA, USA), major histocompatibility complex-II (MHC-II) (Abcam Cambridge, MA, USA), actin (Cell signaling, Danvers, MA, USA) and occludin (Santa Cruz biotechnology, Santa Cruz, CA, USA) were analyzed using Western Blotting. MHC-II are markers of antigen-presenting cells (Lehnardt et al., 2003; Vaure and Liu, 2014). In the brain, microglia and macrophages are the main cells expressing MHC-II (Lehnardt et al., 2003). Occludin is a tight junction protein used to analyzed BBB impairment (Liebner et al., 2000; Yang et al., 2017). Western Blotting was performed as described previously (Joshi et al., 2018). A list of primary antibodies used is provided in the Supplementary Table S8 along with the dilutions and sources. Horseradish peroxidase-conjugated appropriate secondary antibodies and loading control anti-β-actin or tubulin antibody was obtained from Cell Signaling. The blots were analyzed as described previously (Joshi et al., 2018). Briefly after the BCA (as described above), 20 μg of protein from whole brain homogenate were heated with Laemmli buffer containing beta-mercaptoethanol (BioRad, Hercules, CA, USA) and separated by gel electrophoresis using 4–15% Criterion™ TGX Stain-Free™ Protein Gel, (Biorad, Hercules, CA, USA). The membranes were blocked for 1 h in 5% blocking milk (BioRad, Hercules, CA, USA). Then, each membrane was individually immunoprobed with a primary antibody in blocking buffer overnight. After each primary antibody incubation, each membrane was incubated with the recommended dilution (1:5000) of corresponding horseradish peroxidase-conjugated secondary antibody (cell signaling) in blocking buffer at room temperature for one hour. Protein bands were visualized using enhanced chemiluminescence detection reagents (Thermo Scientific, MA USA). Band intensities were analyzed using the ChemiDoc imaging system (Bio-Rad). Results were calculated using Image Lab software and normalized to the expression of actin protein in the samples.

2.15. Multiplex cytokine assay

Selected cytokine levels in the plasma were analyzed using Meso Scale Discovery (MSD) 96-Well MULTI-SPOT® Ultra-Sensitive V-PLEX Proinflammatory cytokine Panel I mouse Kit (MSD, Rockville, MD, USA).
USA), using electrochemiluminescence detection on an MSD Sector Image™ 6000 with Discovery Workbench software (version 3.0.18) (MSD®, Gaithersburg, MD, USA). Cytokines were measured using the TH1/TH2 8-plex kit which included interleukin-1β (IL-1β), IL-6, interferon-γ (IFN-γ) and tumor necrosis factor-α (TNFα). All assays were performed according to the manufacturer’s instructions, in duplicates as described previously (Joshi et al., 2018). Human plasma was analyzed using V-PLEX Proinflammatory cytokine Panel 1 Human Kit (MSD, Rockville, MD, USA) according to manufacturer protocol.

2.16. Statistical analyses

Normally distributed data were analyzed using parametric tests; otherwise non-parametric tests were used. As applicable, the Student’s t-test was used to examine the group differences between PB + PER mice and control mice for cytokines, chemokines, percentage of specific cells identified using flow cytometry and protein levels of p-NFxβ, occludin, MMP9, IgG and MHC-II. Murine cytokine data were analyzed by using the Student’s t-test. Human cytokine data were not normally distributed, so the Kolmogorov-Smirnov test (K-S test) was performed on them to check if they were statistically significant. For autoantibody analysis in the GW veteran cohort, the Chi-square test was performed to determine statistical significance. For multiple groups in blood culture studies, we used one-way analysis of variance (ANOVA) followed by the Tukey post-hoc test. A p value < 0.05 was considered statistically significant. These analyses were performed using GraphPad Prism7 (San Diego, CA, USA), and data were expressed as means ± standard errors. All graphs were generated using GraphPad Prism7 (San Diego, CA, USA).

3. Results

3.1. Detection of 3-PBA released following protein hydrolysis and 3-PBA-modified lysine residues from peptides in the brain, liver and plasma of PB + PER mice

For an acute post-exposure timepoint, we examined 3-PBA modified proteins in liver, brain and plasma of PB + PER exposed mice euthanized at 1-day post-exposure. We also examined a serum sample of a subject positive for 3-PBA in urine as a positive control. Fig. 2 shows PER metabolite structures and detected ions that correspond with the presence of 3-PBA lysine residues after pronase digestion that was performed to release these adducts from peptides (Fig. 2B). We also quantified 3-PBA released post-protein-hydrolysis which showed the presence of 3-PBA in the livers and brains of PB + PER mice at 1.5 months post-exposure (Fig. 2C and 2D). However, free 3-PBA was detected only in the livers but not in the brains of exposed mice. As expected, no 3-PBA was detected in control mice (Fig. 2E and 2F). Neither free nor protein-conjugated 3-PBA was detected in the brains or livers of exposed mice at 3- or 7-months post-exposure.

3.2. 3-PBA-albumin autoantibodies in plasma from PB + PER mice and in GWI veterans

Western blot analyses showed that while plasma from control mice did not have immunoreactivity to free albumin or 3-PBA-albumin, plasma from PB + PER exposed mice showed a strong immunoreactivity with 3-PBA-albumin but not with free albumin (Fig. 3A). We then examined if 3-PBA human albumin-reactive autoantibodies were detectable in plasma from GW veterans. Plasma from 10 out of 12 GW veterans diagnosed with GWI showed immunoreactivity to 3-PBA-albumin (χ²=11.73, df = 1, p = 0.001, Fig. 3B). Details of these antibodies detected is provided in Table 1A. None of the plasma samples from GWI veterans showed any immunoreactivity to albumin alone (Fig. 3B). There was a significant association between self-report of pesticide exposure and 3-PBA-albumin autoantibody presence among veterans with GWI (χ²= 9.46, df = 1, p = 0.03). Only one control GW veteran had 3-PBA-albumin autoantibodies and was exposed to pesticides. Additionally, this control GW veteran reported being exposed to PER but not to PB and autoantibodies against 3-PBA-albumin were still detected in plasma. Also, no 3-PBA-albumin autoantibodies were detected in plasma from one GWI case who was exposed to PB but not PER. Two additional controls who didn’t have autoantibodies did report pesticide exposure, where one subject had an exposure to fogged area whereas the other subject did report exposure to pesticide-imbedded uniforms, creams and sprays. In another cohort from Thailand, we detected 3-PBA-albumin autoantibodies in the serum of 5 subjects who had positive 3-PBA urine levels, whereas control serum samples with undetectable urinary 3-PBA levels did not show immunoreactivity to 3-PBA-albumin (Fig. 3C).

Both human and mice secondary antibody alone showed no cross reactivity with albumin or 3-PBA-hapten (Supplementary Fig. S7).

3.3. Peripheral T-cells and B-cells activated following 3-PBA-albumin treatment

Murine blood was cultured overnight to examine if 3-PBA-albumin as compared to albumin alone, 3-PBA alone and control treatments can activate B-cells, T-cells and monocytes. Antigen-responsive CD19 + CD27 + B-cells were significantly different between the four groups (F(3,12)= 12.2, p = 0.001, Fig. 4A), with the 3-PBA-albumin group being significantly elevated compared to all other groups (post-hoc p < 0.05). Consistently, IgG levels were increased by 3-PBA-albumin treatment compared to all other groups (F(3,12)= 221.8, p = 0.001 Supplementary Fig. S1A). Group comparisons for CD3 + CD4 + T-helper cells showed significant differences between treatments (F(3,16)= 61.19, p = 0.001, Fig. 4B) with post-hoc analyses showing increases in CD3 + CD4 + T-helper cells by 3-PBA-albumin treatment as compared to controls, 3-PBA and albumin alone (p < 0.05). There were no differences in CD3 + CD8 + T-lymphocytes when treated with 3-PBA-albumin compared to all other treatments (p > 0.05). Total monocyte population gated by CD11b+ and CD115+ differed significantly between the groups (F(3,12)= 20.48, p = 0.001, Supplementary Fig. S1B), and post-hoc comparisons showed an increase in these cells in the 3-PBA-albumin group compared to others (p < 0.05).

3.4. Activated peripheral immune cells and plasma proinflammatory cytokines are increased in the blood of PB + PER mice and veterans with GWI

An antigen-reactive B-cell population positive for CD45 + CD19 + CD27+ was elevated in the blood of PB + PER mice compared to control mice (t-test(df=6)= 5.7, p = 0.0012, Fig. 5A). However, the CD19 + CD27− population did not differ between the two groups (p > 0.05), suggesting that while total B-cell population doesn’t change, antigen-responsive B-cells proportionally increase. Higher levels of IgG were present in blood from PB + PER exposed compared to control mice (t-test(df=5)= 4.5, p = 0.008, Supplementary Fig. 1C). We analyzed CD4+ and CD8+ markers after gating on the blood lymphocyte population. This study revealed that CD4+ T-helper cell population was significantly higher in the lymphocytes of exposed compared to control mice (t-test(df=6)= 2.7, p = 0.024, Fig. 5B). T-helper cells expressing CD3 + CD4+ were also elevated in the blood of PB + PER exposed mice (t-test(df=5)= 5.3, p = 0.0017, Fig. 5B). Although CD8+ lymphocytes didn’t differ between the two groups (p > 0.05), the ratios of CD4+ to CD8+ cells were elevated in PB + PER exposed compared to control mice (t-test(df=6)= 3.7, p = 0.0014, Fig. 5B). Mouse monoclonies defined by CD11b+ and CD115+ cells were elevated in PB + PER exposed compared to control mice (t-test(df=11)= 4.2, p = 0.0012, Supplementary Fig. S2).
addition, at 7 months post-exposure, several proinflammatory cyto-
kines, including IL-1β, IL-6, IFN-γ and TNF-α and a chemokine CX3CL1 were elevated in the brain and plasma from this PB + PER mouse model (Supplementary Fig. S3).

We also examined peripheral T- and B-cells in PBMCs from GWI and control GW veterans. The memory B-cell population, as defined by CD19+CD27+ cells, was significantly increased in GWI compared to healthy GW veterans ($t$-test ($df=20)=5.01$, $p=0.0014$, Fig. 6A). T-helper cells expressing CD3+CD4+ are also elevated in GWI veterans ($t$-test ($df=28)=8$, $p=0.001$, Fig. 6B). Although CD3+ CD8+ T-cells didn't differ between the two groups ($p > 0.05$), the ratios of CD4+ to CD8+ cells were elevated in GWI compared to control GW veterans ($t$-test ($df=20)=3.7$, $p=0.0014$). Compared to healthy GW veterans, plasma levels of proinflammatory cytokine IL-1β and CX3CL1 were higher in veterans with GWI (Supplementary Fig. S4).

3.5. Disruption of the BBB in PB + PER mice

Since MMP9 is increased during inflammation, we examined MMP9 levels in the brain homogenates from PB + PER mice and control mice. MMP9 was significantly higher in the brains of PB + PER exposed mice ($t$-test ($df=9)=2.7$, $p=0.028$, Fig. 7B). Assessment of BBB integrity examined using occludin showed a significant decrease in PB + PER exposed mice compared to control mice ($t$-test ($df=10)=2.5$, $p=0.029$, Fig. 7A). Increases in brain IgG levels are generally indicative of BBB impairment, and levels of IgG were increased in the brains of PB + PER exposed mice compared to controls ($t$-test ($df=9)=3.71$, $p=0.043$, Fig. 7B).
Fig. 5. Antigen-activated B- and T-cells are increased in blood of PB+PER mice at 7 months post-exposure. Mean ± SE (n = 4/5 per group). (A) Increased staining of antigen-responsive CD19+ CD27+ B-cells were detected in GWI compared to control mice. (B) Panel B shows that CD3+ CD4+ T-cells are increased in blood of PB+PER mice. However, CD8+ T-cells did not differ between the two groups. Ratios of CD4:CD8 was significantly elevated in PB+PER mice. *p < 0.05.

Fig. 6. Memory B- and CD4 T-cells are increased in blood of veterans with GWI. Mean ± SE (n = 12 per group). (A) Increased staining CD19+ and CD19+ CD27+ memory B-cells in blood of veterans with GWI compared to healthy GW control veterans. (B) CD3+ CD4+ T_{H} cells are increased in blood of veterans with GWI. However, CD8+ T_{H} cells did not significantly differ between control and GWI veterans. However, there were no group differences for CD3+CD8+ T_{H} cells. *p < 0.05.
3.6. Infiltrating monocyte population in the blood and brain of PB + PER mice

Monocytes staining with CD11b+ CD115+ and Ly6C+ classified as proinflammatory and infiltrating were elevated in PB + PER exposed mice blood compared to control mice \( (t\text{-test}(df=11)=3.7, \ p=0.0032, \ \text{Fig. 8A}) \). Macrophages were detected using CD11b+ CD206+ Ly6C+ positive stains, which were increased in the brains of PB + PER exposed mice compared to control mice \( (t\text{-test}(df=2.8)=4.7, \ p=0.04, \ \text{Fig. 8B}) \). The microglia population characterized by CD11b+, CX3CR1+, CD206− Ly6C− showed no differences between the two groups (Fig. 8C). Infiltrating monocytes detected by CD11b+ CX3CR1− Ly6C+ were significantly increased in the brains of exposed mice compared to controls \( (t\text{-test}(df=4)=4.7, \ p=0.01, \ \text{Fig. 8D}) \). Patrolling monocyte populations (CD11b+ CX3CR1+) were significantly increased in the brain of PB + PER exposed mice \( (t\text{-test}(df=6)=6.2, \ p=0.032, \ \text{Supplementary Fig. 2B}) \). We also observed increased MHC-II protein levels in the brains of PB + PER exposed mice compared to controls \( (t\text{-test}(df=6)=6.2, \ p=0.0007, \ \text{Fig. 8E}) \).

3.7. Chemokines and their receptors that promote CNS and peripheral immune cross-talk are increased in PB + PER mice

Since plasma CX3CL1 levels were increased in PB + PER exposed mice and in veterans with GWI (Supplementary Figs. 3 and 4), we examined its receptor CX3CR1 in the brain homogenates from PB + PER mice and control mice at 7-months post-exposure. Levels of CX3CR1 were elevated in PB + PER mice compared to control mice \( (t\text{-test}(df=5)=3.1, \ p=0.03, \ \text{Fig. 9A}) \). There were no differences in the membrane fraction of the brain homogenates between PB + PER exposed mice and control mice at 7-months post-exposure \( (p=0.86, \ \text{Fig. 9B}) \). Levels of CCR2 \( (t\text{-test}(df=10)=4.71, \ p=0.003, \ \text{Supplementary Fig. 2A}) \) were increased in PB + PER mice compared to controls \( (t\text{-test}(df=10)=4.71, \ p=0.003, \ \text{Supplementary Fig. 2B}) \). We also observed increased MHC-II protein levels in the brains of PB + PER exposed mice compared to controls \( (t\text{-test}(df=6)=6.2, \ p=0.0007, \ \text{Fig. 8E}) \).
Fig. 8. Infiltrating monocytes are increased in blood and brains of PB + PER mice at 7–8 months post-exposure. Mean ± SE (n = 12 per group) (A) Increases in blood monocyte gated by CD11b+ CD115+ and Ly6C+ in PB + PER mice were observed. (B) PB + PER mice brain showed increased macrophage population as detected by CD11b+ CD206+ Ly6C+ but (C) no change in microglia population detected by CD11b+ CX3CR1+ and CD206−. (D) Infiltrating monocytes converting to macrophage (CD11b+ CX3CR1− and Ly6C+) were also increased in the brains of PB + PER mice. (E) Increase in MHC-II in the brain of PB + PER mice was detected by Western Blot *p < 0.05.

Fig. 9. CCR2 and its ligand CCL2 are chronically increased in the brains of PB + PER-exposed mice leading to inflammation. Mean as % control ± SEM (n = 6 per group). (A) Western Blot showing CX3CR1/actin and pNFκB/NFκB is increased in the brains of PB + PER-exposed mice. (B) No change in CX3CL1 level as measured by ELISA. (C) Levels of CCL2 were also increased in PB + PER-exposed mice at 7-months post-exposure. (D) CCR2 levels were increased in the brains of PB + PER mice. *p < 0.05.
p = 0.0008, Fig. 9C) and its ligand CCL2 \((t_{\text{test}(22)} = 3.9, p = 0.003, \text{Fig. 9D})\) were increased in the brains of PB + PER exposed mice compared to control mice. Immunohistochemistry work showed astrogliosis activation in the dentate gyrus and cortex of PB + PER mice compared to control mice (Supplementary Fig. S5). We examined the ratios of phosphorylated p65 (p-NFκB) to total p65 (NFκB) in PB + PER exposed mice to assess inflammation. There was a significant increase in p-NFκB/NFκB ratios \((t_{\text{test}(22)} = 2.2, p = 0.06, \text{Fig. 8A})\). Post-hoc analyses showed that the ratios of p-NFκB/NFκB were increased in PB + PER mice at this 7-months post-exposure timepoint.

4. Discussion

Evidence suggests that overexposure to PB and pesticides, which included pyrethroids and organophosphates, was one of the etiological factors in GWI (Binns et al., 2008; White et al., 2016; Sullivan et al., 2018). Exposure to these compounds has been shown to induce chronic immune dysfunction (Broderick et al., 2013; Craddock et al., 2015; Jokanović, 2018). While clinical symptoms of GWI resemble those of autoimmune diseases, the exact mechanism by which GW chemicals caused immune activation remains unknown. In this translational study, we show that a common pyrethroid metabolite, 3-PBA, can haptenate albumin, which activates CD4+ T-helper and antigen-responsive B-cells. These immune cells were elevated in PB + PER exposed mice and in veterans with GWI. We observed autoantibodies against 3-PBA-albumin in our PB + PER mouse model and in veterans with GWI at chronic post-exposure timepoints. In this PB + PER mouse model, we also show that brain macrophages thought to be derived from peripheral blood monocytes were increased in the brains of PB + PER exposed mice. These studies suggest that activation of autoimmune responses against endogenous proteins modified by 3-PBA corresponds with activation of brain macrophages and neuroinflammation in PB + PER exposed mice. The work described herein could aid in the development of sensitive and specific biomarkers for diagnosing GWI in order to provide appropriate care and treatment to veterans with GWI.

During the 1991 GW, a large proportion of veterans were using pyrethroid-based pesticides routinely (Binns et al., 2008; Sullivan et al., 2018). To date, accurate diagnosis of GWI remains an unmet need for ill GW veterans. One reason it is difficult to accurately diagnose GWI is because the diagnosis is based on self-report of symptoms and exposures. Since studies have shown that 3-PBA can form adducts with lysine residues on endogenous proteins (Noort, Benschop and Black, 2002; Noort et al., 2008; Thiphom et al., 2012), we performed this experiment and found to be present in the PB + PER mouse acutely after GW chemical exposure. We also examined 3-PBA recovered after protein hydrolysis and found to be present at 1.5 months after exposure to PB and PER but not at later post-exposure timepoints. This suggests that 3-PBA-haptenated proteins are eventually eliminated. Small molecules which haptenate endogenous proteins can elicit adaptive immune responses, ultimately resulting in B-cell activation and autoantibody production (Naibbit et al., 2002; Palm and Medzhitov, 2009). Since albumin is easily haptenated by drugs due to its abundance in blood and availability of lysine residues that can be covalently modified (Tailor et al., 2016), albumin was chosen for haptenation studies with 3-PBA. Our observations of autoantibodies against 3-PBA-albumin in the periphery suggest that veterans with GWI and pyrethroid-exposed subjects suggest that 3-PBA-albumin is capable of provoking autoimmune responses. These studies for the first time provide evidence of autoantibodies following pyrethroid exposure in veterans with GWI. The specificity of these autoantibodies to pyrethroid exposure is further supported by the observation of 3-PBA-albumin autoantibodies in subjects with detectable urinary 3-PBA levels and observable 3-PBA-lysine adducts in serum. While it remains to be determined if autoantibodies against 3-PBA-albumin have a pathological role in GWI, a higher prevalence of these autoantibodies in GWI veterans compared to control GW veterans suggests that they could be useful biomarkers of GWI.

Our animal and human studies suggest that autoantibodies against 3-PBA-albumin persist for a long time, despite the fact that 3-PBA is not detected in the brain or the liver beyond 1.5 months post-exposure. Studies have shown that proteins haptenated by other chemicals can directly interact with APC, which then travel to the lymphatic system where they present these antigens to T- and B-cells (Karlsson et al., 2018). This may result in the production of autoantibodies against haptenated proteins (Lanzavecchia, 1985; Gefen et al., 2015; Karlsson et al., 2018). While the exact mechanism behind the persistence of autoantibodies against 3-PBA-albumin remains to be investigated, it may involve a similar mechanism which requires memory B-cell activation and subsequent autoantibody production by plasma B-cells (Ochsenbein et al., 2000; Alberts and Johnson, 2002). As mentioned above, we show that autoantibodies against 3-PBA-albumin are generated in our PB + PER mouse model. Furthermore, results of an in vitro blood culture studies suggest that the adaptive immune system is engaged in promoting immune cell activation after 3-PBA-albumin treatment, which is characterized by an elevation of antigen-presenting monocytes, CD3+CD4+ T-helper cells and antigen-responsive CD19+CD27+ B-cells. Interestingly, these cells did not react to 3-PBA or albumin alone, further strengthening the concept that chemical haptenation of albumin is required to elicit adaptive immune responses and that 3-PBA alone cannot promote immune activation. Increases of CD3+CD4+ T-helper and CD19+CD27+ B-cells, together with the presence of autoantibodies against 3-PBA-albumin in PB + PER mice and in GWI veterans, further support an autoimmune-type response in GWI. These studies provide support for an activation of the adaptive immune system in GWI and are consistent with those conducted by Vojdanij and Thrasher (2004), showing an elevation of CD4+ T-helper cells and B-cells in the blood of veterans with GWI. Together, these findings suggest that 3-PBA-albumin can stimulate peripheral immune cell activation.

While most veterans deployed in the GW conflict self-report the use of PB, it is unlikely that PB alone is a major contributing factor in the overactive immune responses noted in veterans with GWI. On the contrary, previous research has suggested that PB is able to suppress inflammation in rodent models (Macht et al., 2019), with exposure to PB alone resulting in decreased production of pro-inflammatory cytokines such as IL-6, IFN-γ and TNF-α at chronic time points (Macht et al., 2018). However, we, along with others, have shown that these cytokines are elevated in plasma from veterans with GWI (Broderick et al., 2011; Khaiboullina et al., 2015). The mechanism of PB is well defined in the literature, acting as a reversible acetylcholinesterase (AChE) inhibitor (Musiek et al., 2011). As such, PB use would likely result in an increase in acetylcholine (ACh) levels in the periphery (Musiek et al., 2011). Recent evidence suggests a role of ACh in stimulating anti-inflammatory responses in the periphery via a direct interaction with nicotinic receptors on peripheral immune cells (Tracey, 2005; Ibrahim et al., 2018). Therefore, one would expect a dampened immune response in PB + PER exposed mice. However, since we observe activated immune responses and the presence of 3-PBA-albumin autoantibodies in plasma of both human and PB + PER exposed mice, it is possible that PB was unable to suppress immune cell activation that occurs in the presence of combined PB + PER exposure (Joshi et al., 2018). Additional research is needed to clarify if PB metabolites may also form haptenated proteins, which may promote synergistic interactions resulting in the observed adverse adaptive immune response.

Under normal physiological conditions, the BBB largely restricts entry of peripheral immune cells into the brain, which helps maintain the immune privilege status of the CNS (Van Dyken and Lacote, 2018). However, peripheral immune cells have been shown to migrate into the brain parenchyma during inflammation resulting from the increases in proinflammatory cytokines and MMP9 activity which weakens the BBB.
5. Conclusions

These explorative studies provide strong evidence that in our PB + PER mouse model of GWI, PER metabolite 3-PBA haptenated proteins and promoted adaptive immune responses. We also observe autoantibodies against 3-PBA-haptenated proteins in PB + PER exposed mice and in veterans with GWI. While the exact mechanisms associated with these aberrant autoimmune responses and their effects on the CNS pathology of GWI remain to be elucidated, the work presented herein suggests a cross-talk between the brain immune cells and peripheral adaptive immune cells, which may ultimately promote the entry of peripheral immune cells into the brain and promote neuroinflammatory responses. We anticipate that these studies will help guide future development of biomarkers and therapies for the CNS pathology of GWI.

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Declaration of Competing Interest

The authors declare no competing interests. The contents do not represent the views of the Department of Veterans Affairs or the United States Government.

Appendix A. Supplementary data

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